AJCC Cancer Staging Form Supplement

AJCC Cancer Staging Manual, Eighth Edition

Last updated 7 January 2021

AMERICAN JOINT COMMITTEE ON CANCER
Executive Office
633 North Saint Clair Street
Chicago, IL 60611-3211

www.cancerstaging.org ajcc@facs.org





Introduction

The AJCC Cancer Staging Manual, Eighth Edition Staging Form Supplement includes 104 printable staging forms for each distinct staging system published by the American College of Surgeons (ACS).

These printable forms may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; histologic grade; and other important information. These forms may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging forms may be used to document cancer stage at different points in the patient's care and during the course of therapy, including the time before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

See Principles of Cancer Staging¹ (Chapter 1) of the *AJCC Cancer Staging Manual, Eighth Edition*² for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. The staging forms cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

Organization of This Supplement

The staging forms in this supplement are numbered according to their corresponding chapters in the AJCC Cancer Staging Manual, Eighth Edition.² For example, chapter 6, Cervical Lymph Nodes and Unknown Primary Tumors of the Head and Neck is the first chapter in the manual that has data collection items, so it is the first staging form in this supplement.

Some chapters have multiple staging forms as they describe distinct TNM, Prognostic Factors, and AJCC Prognostic Stage Groups for unique topographical sites, histologic types or a combination of the two.

These forms may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

References

- 1. Gress, D.M., Edge, S.B., Gershenwald, J.E., et al. Principles of Cancer Staging. In: Amin, M.B., Edge, S.B., Greene, F.L., et al. (Eds.) AJCC Cancer Staging Manual. 8th Ed. New York: Springer; 2017: 3-30
- 2. Amin, M.B., Edge, S.B., Greene, F.L., et al. (Eds.) AJCC Cancer Staging Manual. 8th Ed. New York: Springer; 2017

Summary of Changes 7 January 2021

| Form Number | Title | Section | Before Correction | After Correction |
|----------------|--------------------|-------------------|---|--|
| | | 4.1 Definition of | Tumor \leq 2 cm, with DOI* > 5 mm and \leq 10 mm or | Tumor ≤ 2 cm, with DOI* > 5 mm or tumor > 2 cm |
| | | Primary Tumor | tumor > 2 cm and ≤ 4 cm, with DOI* ≤ 10 mm | and ≤ 4 cm, with DOI* ≤ 10 mm |
| 7 | Oral Cavity | (T) | , | |
| | Male Penile | | | |
| | Urethra and | | | |
| | Female Urethra: | 5. AJCC | | |
| | Urothelial | Prognostic Stage | Row Omitted | T4 NX M0 Stage IV |
| 63.1 | Carcinomas | Groups | | |
| | Male Penile and | | | |
| | Female Urethra: | | | |
| | Squamous Cell | 5. AJCC | | |
| | Carcinoma and | Prognostic Stage | Row Omitted | T4 NX M0 Stage IV |
| 63.2 | Adenocarcinoma | Groups | | · · |
| | Prostatic Urethra: | 5. AJCC | | |
| | Urothelial | Prognostic Stage | Row Omitted | T4 NX M0 Stage IV |
| 63.3 | Carcinomas | Groups | Now Officied | 14 IVA IVIO Stage IV |
| | Prostatic Urethra: | J. Cups | | |
| | Squamous Cell | 5. AJCC | | |
| | Carcinoma and | Prognostic Stage | Row Omitted | T4 NX M0 Stage IV |
| 63.4 | Adenocarcinoma | Groups | Now officed | Trivino stage iv |
| | Gestational | , | | |
| | Trophoblastic | | Pretreatment hCG (IU/mL) | Pretreatment hCG (mIU/mL) |
| 56 | Neoplasms | 5.1 Risk Score | Treatestiment mes (refiniz) | Trediedament nee (mile) me) |
| | Ovary, Fallopian | | | |
| | Tube and Primary | 4.3 Definition of | | |
| | Peritoneal | Distant | cM1a | Delete |
| 55 | Carcinoma | Metastasis (M) | | |

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6. Cervical Lymph Nodes and Unknown Primary Tumors of the Head and Neck

1 Terms of Use

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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
| | | |
| | | |
| | | |
| | | |

6. Cervical Lymph Nodes and Unknown Primary Tumors of the Head and Neck

4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instruction for clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| I | ✓ | T Category | T Criteria |
|---|---|------------|------------------------------|
| | | T0 | No evidence of primary tumor |

| ✓ | T Suffix | Definition | |
|---|----------|--|--|
| | (m) | m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

These criteria apply to patients who are treated with primary nonsurgical treatment without a cervical lymph node dissection.

| √ | cN Category | cN Criteria |
|----------|--|--|
| | NX | Regional lymph nodes cannot be assessed |
| | NO No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension, ENE(-) |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) with clinically overt ENE(+) (ENE _c) ² |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any node(s) with clinically overt ENE(+) (ENE _c) ² |
| A1-4 | Value Midling and a separate of included and a CNC in defined a impair of this infiltration of a separate data and a table in a first in | |

Notes: Midline nodes are considered ipsilateral nodes. ENE_c is defined as invasion of skin, infiltration of musculature, dense tethering or fixation to adjacent structures, or cranial nerve, brachial plexus, sympathetic trunk, or phrenic nerve invasion with dysfunction.

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |

6. Cervical Lymph Nodes and Unknown Primary Tumors of the Head and Neck

4.2.2 Pathological N (pN)

These criteria apply to patients who are treated surgically with a cervical lymph node dissection.

| ✓ | pN Category | pN Criteria |
|---|-------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); |
| | | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); |
| | | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) |
| | N2a | Metastasis in a single ipsilateral node 3 cm or less in greatest dimension and ENE(+); |
| | | or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); |
| | | or metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | | or multiple ipsilateral, contralateral, or bilateral nodes any size and ENE(+) in any node; |
| | | or a single contralateral node of any size and ENE(+) |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | | or multiple ipsilateral, contralateral, or bilateral nodes any size and ENE(+) in any node; |
| | | or a single contralateral node of any size and ENE(+) |

Note: Midline nodes are considered ipsilateral nodes. ENE detected on histopathologic examination is designated as ENE_{mi} (microscopic $ENE \le 2$ mm) or ENE_{ma} (macroscopic ENE > 2 mm). Both ENE_{mi} and ENE_{ma} qualify as ENE(+) for definition of pN.

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | TO TO | N1 | M0 | III |
| | TO TO | N2 | M0 | IVA |
| | TO | N3 | M0 | IVB |
| | TO TO | Any N | M1 | IVC |

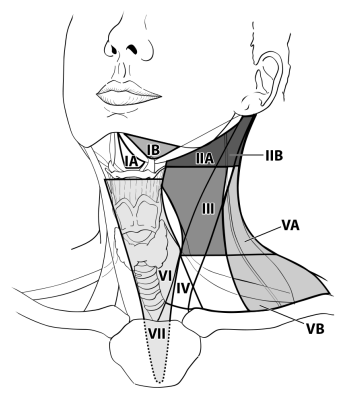
| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
| | |

| 1. Extranodal extension for all anatomic sites with the exception of HPV-related oropharynx cancer, nasopharynx cancer, melanoma sarcoma, and thyroid carcinoma: Yes No 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH Hymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | sarcoma, and thyroid carcinoma: Yes No 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion | | Re | egistry Data Collect | tion Variables |
|---|--|--|------|----------------------------|--|
| 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | | 1. | Extranodal extension for | all anatomic sites with the exception of HPV-related oropharynx cancer, nasopharynx cancer, melanoma |
| 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 2. Size of largest metastatic node: 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | | | sarcoma, and thyroid ca | rcinoma: TYes No |
| 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description O LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 3. Number of metastatic lymph nodes: 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 2. Size of largest metastatic node: | | | |
| 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 4. Laterality of metastatic nodes; note that midline nodes are considered ipsilateral nodes: 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | | | | |
| 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 5. Level of nodal involvement: 6. ENE clinical (select one): Positive (+) Negative (-) 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description Coding 0 LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | | | | |
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| 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description O LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 7. ENE pathological (select one): Positive (+) Negative (-) Lymphovascular Invasion (LVI) Component of LVI Description O LVI not present (absent)/not identified 1 LVI present/identified, NOS 2 Lymphatic and small vessel invasion only (L) 3 Venous (large vessel) invasion only (V) 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion 9 Presence of LVI unknown/indeterminate | 5. Level of nodal involvement: | | | |
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8 Anatomy

FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---------------------|--|--|
| | TX | Primary tumor cannot be assessed | |
| | Tis | Carcinoma in situ | |
| | T1 | Tumor ≤ 2 cm with depth of invasion (DOI)* ≤ 5 mm | |
| | T2 | Tumor ≤ 2 cm with DOI* > 5 mm | |
| | | or tumor > 2 cm and \leq 4 cm with DOI* \leq 10 mm | |
| | T3 | Tumor > 2 cm and ≤ 4 cm with DOI* > 10 mm | |
| | | or tumor > 4 cm with DOI* ≤ 10 mm | |
| | T4 | Moderately advanced or very advanced local disease | |
| | T4a | Moderately advanced local disease | |
| | | Tumor > 4 cm with DOI* > 10 mm | |
| | | or tumor invades adjacent structures only (e.g., through cortical bone of the mandible or maxilla or involves the maxillary sinus or skin of the face) | |
| | | Note: Superficial erosion of bone/tooth socket (alone) by a gingival primary is not sufficient to classify a tumor as T4. | |
| | T4b | Very advanced local disease | |
| | | Tumor invades masticator space, pterygoid plates, or skull base and/or encases the internal carotid artery | |
| | *DOI is depth of in | vasion and <u>not</u> tumor thickness. | |

| ✓ | T Suffix | Definition | |
|---|----------|---|--|
| | (m) | Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | cN Category | cN Criteria |
|-----|-----------------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension ENE(–) |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension, and ENE(-) |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension, and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension, and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension, and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) and clinically overt ENE(+) |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any node(s) and clinically overt ENE(+) |
| Not | e. A designation of " | "I" or "I" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid | |
| | L | Metastasis below the lower border of the cricoid | |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| pN Category | pN Criteria |
|---|--|
| NX | Regional lymph nodes cannot be assessed |
| N0 | No regional lymph node metastasis |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| N2 Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); |
| | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); |
| | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension, ENE(-) |
| N2a | Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+); |
| | or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) |
| N3 | N3: Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); |
| | or metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | or multiple ipsilateral, contralateral, or bilateral nodes, any with ENE(+); |
| | or a single contralateral node of any size and ENE(+) |
| N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| N3b | Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | or multiple ipsilateral, contralateral or bilateral nodes any with ENE(+); |
| | or a single contralateral node of any size and ENE(+) |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid | |
| | L | Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Assignment of the M category for pathological classification may be cM0, cM1, or pM1.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|--------------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | NO | M0 | III |
| | T1,T2,T3 | N1 | M0 | III |
| | T4a | N0,N1 | M0 | IVA |
| | T1,T2,T3,T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

| 6 Registry Data Coll | ection Variables | | | | |
|---|--|---|--|--|--|
| See chapter for more details on | See chapter for more details on these variables. | | | | |
| 1. ENE clinical (select one): | NE clinical (select one): Present/Positive (+) Absent/Negative (-) | | | | |
| 2. ENE pathological (select one) | 2. ENE pathological (select one): Present/Positive (+) Absent/Negative (-) | | | | |
| 3. Extent of microscopic ENE (di tissue): | stance of extension from the native lym | ph node capsule to the farthest point of invasion in the extranodal | | | |
| 4. Perineural invasion: | | ultifocal ultifocal | | | |
| 5. Lymphovascular invasion: Intratumoral: Focal Multifocal Extratumoral: Focal Multifocal | | | | | |
| 6. p16/HPV status: Positiv | • • | | | | |
| 7. Performance status (0-5): | | | | | |
| 8. Tobacco use and pack-year: | 3. Tobacco use and pack-year: □ Never □ ≤ 10 pack-years □ > 10 but ≤ 20 pack-years □ > 20 pack-years | | | | |
| 9. Alcohol use: Number of days drinking per week: Number of drinks per day: | | | | | |
| 10. Depression diagnosis: Previously diagnosed Currently diagnosed | | | | | |
| 11. Depth of invasion (mm): | | | | | |
| Hospital Name/Address | | Patient Name/Information | | | |
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| 7. Oral Cavity | | | | | |
|----------------|--|-----------------------|--|------------|--|
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| 12. | 12. Margin Status: grossly involved | | | | |
| | ☐ microscopic involvement | | | | |
| | | | | | |
| 13. | Distance of | tumor (or moderat | e/severe dysplasia) from closest marg | rin: | |
| | | | | | |
| 14. | WPOI-5 (w | orst patterns of inva | asion): Present No | ot present | |
| | | | | | |
| | | | | | |
| 7 | Histol | ogic Grade (G) | | | |
| - | | | | | |
| ✓ | G | G Definition | | | |
| | GX | Cannot be assesse | | | |
| | G1 | Well differentiate | | | |
| | G2 | Moderately differ | | | |
| | G3 | Poorly differentia | ted | | |
| | | | | | |
| 8 | Lympl | hovascular Inv | asion (LVI) | | |
| | | | | | |
| 1 | Compon | ent of LVI | Description | | |
| | Coding | | | | |
| | 0 | | LVI not present (absent)/not ident | fied | |
| - | 2 | | LVI present/identified, NOS Lymphatic and small vessel invasio | n only (I) | |
| | 3 | | Venous (large vessel) invasion only | | |
| | 4 | | BOTH lymphatic and small vessel A | | |
| | 9 | | Presence of LVI unknown/indetern | | |
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9 Anatomy

FIGURE 7.1. Anatomical sites and subsites of the oral cavity.

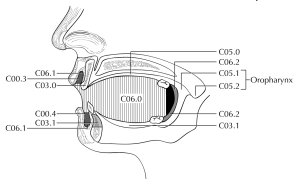


FIGURE 7.2. Anatomical sites and subsites of the oral cavity.

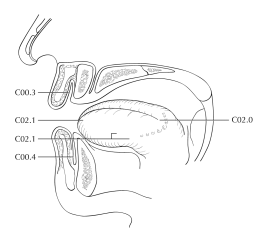


FIGURE 7.3. Anatomical sites and subsites of the oral cavity.

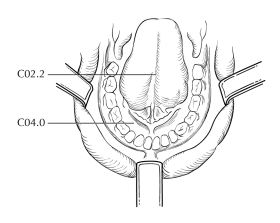
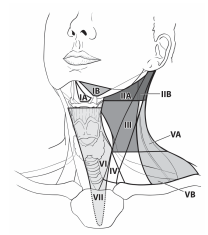


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|--|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ |
| | T1 | Tumor 2 cm or smaller in greatest dimension without extraparenchymal extension* |
| | T2 | Tumor larger than 2 cm but not larger than 4 cm in greatest dimension without extraparenchymal extension* |
| | T3 | Tumor larger than 4 cm and/or tumor having extraparenchymal extension* |
| | T4 | Moderately advanced or very advanced disease |
| | T4a | Moderately advanced disease |
| | | Tumor invades skin, mandible, ear canal, and/or facial nerve |
| | T4b | Very advanced disease |
| | | Tumor invades skull base and/or pterygoid plates and/or encases carotid artery |
| - | * Extraparenchyma | al extension is clinical or macroscopic evidence of invasion of soft tissues. Microscopic evidence alone does not |
| | constitute extraparenchymal extension for classification purposes. | |

| V | T Suffix | Definition |
|----------|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | cN Category | cN Criteria |
|---|--|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(–); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(–) |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) with clinically overt ENE(+) |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any node(s) with clinically overt ENE(+) |
| | Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+). | |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | 1 | Metastasis below the lower horder of the cricoid |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| ✓ | pN Category | pN Criteria | |
|----|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a | Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+) | |
| | | or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); | |
| | | or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | |
| | | or multiple ipsilateral, contralateral, or bilateral nodes any with ENE(+); | |
| | | or a single contralateral node of any size and ENE(+) | |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | |
| | N3b | Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | |
| | | or multiple ipsilateral, contralateral, or bilateral nodes any with ENE(+); | |
| | | or a single contralateral node of any size and ENE(+) | |
| No | Note: A designation of "II" or "I" may be used for any N category to indicate metastasis above the lower horder of the cricoid (II) or below the | | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition | |
|---|--|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |
| | U Metastasis above the lower border of the cricoid | | |
| | L | Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Assignment of the M category for pathological classification may be cM0, cM1, or pM1.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|---------------------|----------|----------|-------------------------|
| | Tis | N0 | MO | 0 |
| | T1 | NO NO | M0 | 1 |
| | T2 | N0 | MO | П |
| | T3 | NO NO | M0 | III |
| | T0, T1, T2, T3 | N1 | MO | III |
| | T4a | N0, N1 | MO | IVA |
| | T0, T1, T2, T3, T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | MO | IVB |
| | Any T | Any N | M1 | IVC |

| 6 Registry Data Coll | ection Variables | | | |
|--|--|--|--|--|
| See chapter for more details on these variables. | | | | |
| 1. ENE clinical (select one): | Present/Positive (+) Absent/Negative (-) | | | |
| 2. ENE pathological (select one) | : Present/Positive (+) Absent/Negative (-) | | | |
| 3. Extent of microscopic ENE (di tissue): | stance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal | | | |
| 4. Perineural invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal | | | |
| 5. Lymphovascular invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal | | | |
| 6. p16/HPV status: Positiv | | | | |
| 7. Performance status (0-5): | | | | |
| 8. Tobacco use and pack-year: | <pre>Never</pre> | | | |
| | of days drinking per week: of drinks per day: | | | |
| 10. Depression diagnosis: | ☐ Previously diagnosed ☐ Currently diagnosed | | | |
| This form continues on the next | page. | | | |
| Hospital Name/Address | Patient Name/Information | | | |
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7 Histologic Grade (G)

There is no uniform grading system for salivary gland.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 8.1. Major salivary glands include the parotid, submandibular, and sublingual glands.

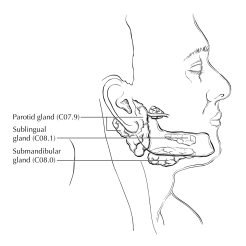
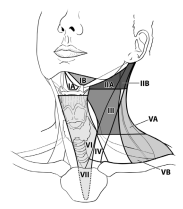


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|---|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No tumor identified, but EBV-positive cervical node(s) involvement |
| | Tis | Tumor in situ |
| | T1 Tumor confined to nasopharynx, or extension to oropharynx and/or nasal cavity without parapharyngeal | |
| | | involvement |
| | T2 | Tumor with extension to parapharyngeal space, and/or adjacent soft tissue involvement (medial pterygoid, lateral |
| | | pterygoid, prevertebral muscles) |
| | T3 | Tumor with infiltration of bony structures at skull base, cervical vertebra, pterygoid structures, and/or paranasal |
| | | sinuses |
| | T4 | Tumor with intracranial extension, involvement of cranial nerves, hypopharynx, orbit, parotid gland, and/or |
| | | extensive soft tissue infiltration beyond the lateral surface of the lateral pterygoid muscle |

| / | T Suffix | Definition | |
|---|----------|---|--|
| | (m) | Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|------------|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Unilateral metastasis in cervical lymph node(s) and/or unilateral or bilateral metastasis in retropharyngeal lymph | |
| | | node(s), 6 cm or smaller in greatest dimension, above the caudal border of cricoid cartilage | |
| | N2 | Bilateral metastasis in cervical lymph node(s), 6 cm or smaller in greatest dimension, above the caudal border of | |
| | | cricoid cartilage | |
| | N3 | Unilateral or bilateral metastasis in cervical lymph node(s), larger than 6 cm in greatest dimension, and/or extension | |
| | | below the caudal border of cricoid cartilage | |

| ✓ | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T1, T0 | N1 | M0 | II |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | II |
| | T1, T0 | N2 | M0 | III |
| | T2 | N2 | M0 | III |
| | T3 | N0 | M0 | III |
| | T3 | N1 | M0 | III |
| | T3 | N2 | M0 | III |
| | T4 | N0 | M0 | IVA |
| | T4 | N1 | M0 | IVA |
| | T4 | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

Beyond the factors required for staging, the authors have not identified any additional registry data collection variables.

7 Histologic Grade (G)

A grading system is not used for NPCs.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of | Description | |
|---|--------------|--|--|
| | LVI Coding | | |
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | ymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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9 Illustrations

FIGURE 9.1. Anatomical sites and subsites of the nasopharynx, oropharynx, hypopharynx, and esophagus.

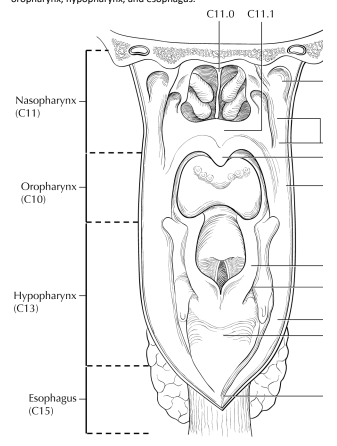
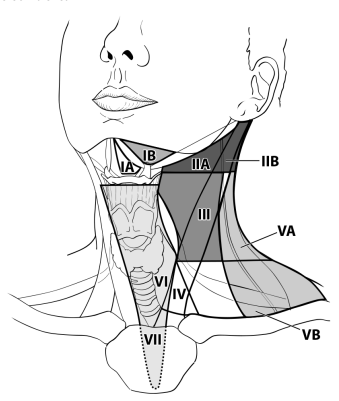


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification Definition | |
|---|--|---|
| workup information, until first treatment, including clinical history and symptoms, physical examination, i endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sar | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information fro diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgi specimens | |
| | ycTNM Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therap before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvat therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | | | |
|-----|--|--|--|--|--|
| | T0 | No primary identified | | | |
| | T1 | Tumor 2 cm or smaller in greatest dimension | | | |
| | T2 | Tumor larger than 2 cm but not larger than 4 cm in greatest dimension | | | |
| | T3 | Tumor larger than 4 cm in greatest dimension or extension to lingual surface of epiglottis | | | |
| | T4 Moderately advanced local disease. | | | | |
| | Tumor invades the larynx, extrinsic muscle of tongue, medial pterygoid, hard palate, or mandible or beyond* | | | | |
| * M | * Mucosal extension to lingual surface of epiglottis from primary tumors of the base of the tongue and vallecula does not constitute invasion of | | | | |
| +60 | lanunu | | | | |

the larynx.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | cN Category | cN Criteria | |
|---|-------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | One or more ipsilateral lymph nodes, none larger than 6 cm | |
| | N2 | Contralateral or bilateral lymph nodes, none larger than 6 cm | |
| | N3 | Lymph node(s) larger than 6 cm | |

| 1 | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.2.2 Pathological N (pN)

| ✓ | pN Category | pN Criteria | |
|---|-------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | pN0 | No regional lymph node metastasis | |
| | pN1 | Metastasis in 4 or fewer lymph nodes | |
| | pN2 | Metastasis in more than 4 lymph nodes | |

| 1 | N Suffix | Definition | |
|---|--|---|--|
| | (sn) | sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

| Patient Name/Information | |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|------------|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | pM1 | Distant metastasis, microscopically confirmed | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of p16/HPV Status

| ✓ | P16/HPV Status | | |
|---|--|--|--|
| | Positive (+) | | |
| | Negative (-) If negative, use staging form for p16- Oropharynx, Chapter 11. | | |
| | Not tested. If not tested, use staging form for p16- Oropharynx, Chapter 11. | | |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6.1 Clinical (cTNM)

| _/ | When p16/HPV | And T is | And N is | And M is | Then the stage |
|----|--------------|----------------------|------------------|----------|----------------|
| | Status is | | | | group is |
| | Positive | T0, T1 or T2 | N0 or N1 | M0 | 1 |
| | Positive | T0, T1 or T2 | N2 | M0 | II |
| | Positive | T3 | N0, N1 or N2 | M0 | II |
| | Positive | T0, T1, T2, T3 or T4 | N3 | M0 | III |
| | Positive | T4 | N0, N1, N2 or N3 | M0 | III |
| | Positive | Any T | Any N | M1 | IV |

6.2 Pathological (pTNM)

| 1 | When p16/HPV Status is | And T is | And N is | And M is | Then the stage group is |
|---|---------------------------|--------------|----------|----------|-------------------------|
| | Positive | T0, T1 or T2 | NO, N1 | M0 | 1 |
| | Positive | T0, T1 or T2 | N2 | M0 | II |
| | Positive | T3 or T4 | N0, N1 | M0 | II |
| | Positive | T3 or T4 | N2 | M0 | III |
| | Positive | Any T | Any N | M1 | IV |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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10. HPV-Mediated (p16+) Oropharyngeal Cancer

| | 201111 V Mediated (P201) Grophary Media Carroer | | | |
|----------|---|---|--|--|
| | | | | |
| 7 | | a Collection Variables | | |
| See | chapter for more de | tails on these variables. | | |
| 4 - | | Landard and all control of the MCC Charles O November 200 | | |
| 1. 11 | umor location: | posterior wall nasopharynx (use AJCC Chapter 9 Nasopharynx) pharyngeal tonsils (use this chapter, AJCC Chapter 10 HPV-Mediated Oropharyngeal Cancer) | | |
| | | pharyngear torisis (use this chapter, Asce chapter 10 fir vivietrated Gropharyngear Cancer) | | |
| 2. N | umber and size of no | odes: | | |
| | | | | |
| 3. P | 3. Perineural invasion: | | | |
| | | ☐ Extratumoral: ☐ Focal ☐ Multifocal | | |
| | | | | |
| 4. Ex | ktranodal extension: | ☐ gross ≥ 2 mm | | |
| | | microscopic | | |
| 5. Ta | obacco use and pack | -year: Never | | |
| | | ☐≤10 pack-years | | |
| | | ☐ > 10 but ≤ 20 pack-years | | |
| | | > 20 pack-years | | |
| | | | | |
| 8 | Histologic Gr | ada (G) | | |
| | Thistologic Gi | auc (6) | | |
| No. | rading system exists | for HPV-mediated oropharyngeal tumors. | | |
| INO 8 | grading system exists | To the v-mediated drophial yrigear tumors. | | |
| 9 | Lymphovasc | ular Invasion (IVI) | | |
| | 7 7 7 7 7 7 7 7 | | | |
| | | | | |
| ✓ | Component of | Description | | |
| ✓ | LVI Coding | Description | | |
| ✓ | LVI Coding | Description LVI not present (absent)/not identified | | |
| ✓ | LVI Coding | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) | | |
| ✓ | 0 1 2 3 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) | | |
| ✓ | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion | | |
| ✓ | 0 1 2 3 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |
| | 0 1 2 3 4 | Description LVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate | | |

| Hospital Name/Address | Patient Name/Information |
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10 Anatomy

FIGURE 10.2. Sagittal view of the face and neck depicting the subdivisions of the pharynx.

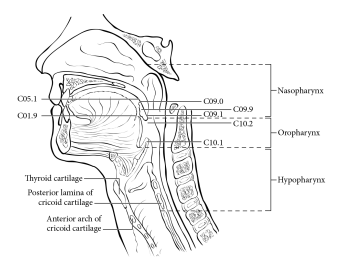
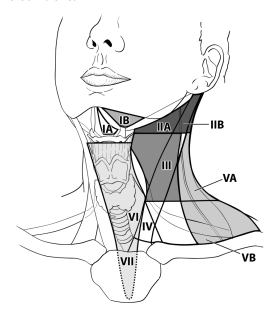


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|---|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor 2 cm or smaller in greatest dimension |
| | T2 | Tumor larger than 2 cm but not larger than 4 cm in greatest dimension |
| | T3 | Tumor larger than 4 cm in greatest dimension or extension to lingual surface of epiglottis |
| | T4 | Moderately advanced or very advanced local disease |
| | T4a | Moderately advanced local disease |
| | | Tumor invades the larynx, extrinsic muscle of tongue, medial pterygoid, hard palate, or mandible* |
| | T4b | Very advanced local disease |
| | | Tumor invades lateral pterygoid muscle, pterygoid plates, lateral nasopharynx, or skull base or encases carotid |
| | | artery |

^{*}Note: Mucosal extension to lingual surface of epiglottis from primary tumors of the base of the tongue and vallecula does not constitute invasion of the larynx.

| 1 | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria | |
|-----|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 | N2 Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2c Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | | |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) and clinically overt ENE(+) | |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | |
| | N3b | Metastasis in any node(s) and clinically overt ENE(+) | |
| Not | te: A designation of | "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information | |
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4.2.2 Pathological N (pN)

| N Category | N Criteria | | |
|--|--|--|---|
| NX | Regional lymph nodes cannot be assessed | | |
| N0 | No regional lymph node metastasis | | |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | | |
| N2 Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension an | | | |
| | | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| | | N2a Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+); | |
| or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | | | |
| N2b Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | | | |
| N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | | |
| N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); | | |
| | or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | |
| | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | | |
| | or a single contralateral node of any size and ENE(+) | | |
| N3a Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | | | |
| N3b Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | | | |
| | | | or a single contralateral node of any size and ENE(+) |
| | NX N0 N1 N2 N2a N2b N2c N3 | | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid | |
| | L | Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of p16/HPV Status

| ✓ | p16/HPV Status | |
|----------|--|--|
| | Negative (-) | |
| | Not tested | |
| | Positive (+) If positive, use staging form for HPV-Mediated (p16+) Oropharyngeal Cancer, Chapter 10. | |

| Hospital Name/Address | Patient Name/Information | |
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6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When p16/HPV Status is | And T is | And N is | And M is | Then the stage group is |
|---|---------------------------|--------------|----------|----------|-------------------------|
| | Negative, not tested | Tis | N0 | M0 | 0 |
| | Negative, not tested | T1 | N0 | M0 | 1 |
| | Negative, not tested | T2 | N0 | M0 | II |
| | Negative, not tested | T3 | N0 | M0 | III |
| | Negative, not tested | T1,T2,T3 | N1 | M0 | III |
| | Negative, not tested | T4a | N0,N1 | M0 | IVA |
| | Negative, not tested | T1,T2,T3,T4a | N2 | M0 | IVA |
| | Negative, not tested | Any T | N3 | M0 | IVB |
| | Negative, not tested | T4b | Any N | M0 | IVB |
| | Negative, not tested | Any T | Any N | M1 | IVC |

| 7 Registry Data Coll | ection Variables | |
|--|--|---|
| See chapter for more details on | these variables. | |
| 1. ENE clinical (select one): | Present/Positive (+) Absent/Negative (-) | |
| 2. ENE pathological (select one) | : Present/Positive (+) Absent/Negative (-) | |
| 3. Extent of microscopic ENE (ditissue): | stance of extension from the native lym | ph node capsule to the farthest point of invasion in the extranodal |
| 4. Perineural invasion: | | ultifocal ultifocal |
| 5. Lymphovascular invasion: | | ultifocal ultifocal |
| · · · = | e (+) (Use AJCC Chapter 10 HPV-Mediat ve (–) (Use this chapter, AJCC Chapter 1: | |
| 7. Performance status (0-5): | | |
| 8. Tobacco use and pack-year: | Never ≤ 10 pack-years > 10 but ≤ 20 pack-years > 20 pack-years | |
| | of days drinking per week: of drinks per day: | |
| 10. Depression diagnosis: | ☐ Previously diagnosed ☐ Currently diagnosed | |
| This form continues on the next | t page. | |
| Hospital Name/Address | | Patient Name/Information |
| | | |

8 Histologic Grade (G)

| ✓ | G | G Definition |
|---|-----------------------------|---------------------------|
| | GX Grade cannot be assessed | |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

9 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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10 Anatomy

FIGURE 11.1. Sagittal view of the face and neck depicting the subdivisions of the pharynx.

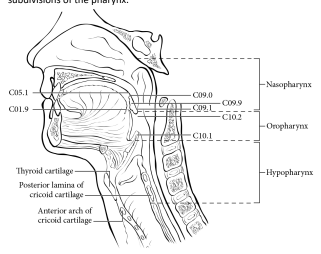
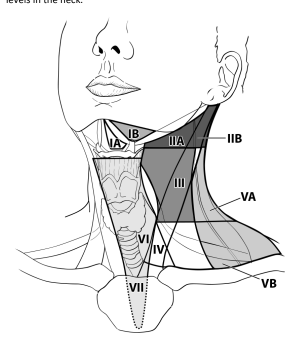


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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1 Terms of Use

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|-----|-----------------------|---|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to one subsite of hypopharynx and/or 2 cm or smaller in greatest dimension |
| | T2 | Tumor invades more than one subsite of hypopharynx or an adjacent site, or measures larger than 2 cm but not |
| | | larger than 4 cm in greatest dimension without fixation of hemilarynx |
| | T3 | Tumor larger than 4 cm in greatest dimension or with fixation of hemilarynx or extension to esophageal mucosa |
| | T4 | Moderately advanced and very advanced local disease |
| | T4a | Moderately advanced local disease |
| | | Tumor invades thyroid/cricoid cartilage, hyoid bone, thyroid gland, esophageal muscle or central compartment |
| | | soft tissue* |
| | T4b | Very advanced local disease |
| | | Tumor invades prevertebral fascia, encases carotid artery, or involves mediastinal structures |
| *No | ote: Central comparti | ment soft tissue includes prelaryngeal strap muscles and subcutaneous fat. |

| Ī | ✓ | T Suffix | Definition |
|---|---|----------|---|
| | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | N Category | N Criteria | |
|--|--|---|--|
| NX Regional lymph nodes cannot be assessed | | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(- | | |
| | N2b Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) N2c Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | | |
| | | | |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) and clinically overt ENE(+) | |
| N3a Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | |
| N3b Metastasis in any node(s) and clinically overt ENE(+) | | Metastasis in any node(s) and clinically overt ENE(+) | |
| Not | e: A designation of | "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition | |
|---|--|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U Metastasis above the lower border of the cricoid | | |
| | L | L Metastasis below the lower border of the cricoid | |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| ✓ | N Category | N Criteria | |
|--|------------|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a | Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+); | |
| or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | | | |
| N2b Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | | | |
| N2c Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and | | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N3 Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | | |
| | | or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | |
| | | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | |
| or a single contralateral node of any size and ENE(+) | | | |
| N3a Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) N3b Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | |
| | | Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | |
| | | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | |
| or a single contralateral node of any size and ENE(+) | | or a single contralateral node of any size and ENE(+) | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) |) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid | |
| | L | Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | ory M Criteria | |
|---|---|-----------------------|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | pM1 Distant metastasis, microscopically confirmed | | |

| Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|--------------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | NO | M0 | III |
| | T1,T2,T3 | N1 | M0 | III |
| | T4a | N0,N1 | M0 | IVA |
| | T1,T2,T3,T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

Registry Data Collection Variables See chapter for more details on these variables. 1. ENE clinical (select one): Present/Positive (+) ☐ Absent/Negative (-) 2. ENE pathological (select one): Present/Positive (+) ☐ Absent/Negative (-) 3. Extent of microscopic ENE (distance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal tissue): ☐ Intratumoral: ■ Multifocal 4. Perineural invasion: ☐ Focal Extratumoral: Focal Multifocal 5. Lymphovascular invasion: Intratumoral: Focal Multifocal Extratumoral: Focal ■ Multifocal 8. Performance status (0-5): 9. Tobacco use and pack-year: □ Never ≤ 10 pack-years _ > 10 but ≤ 20 pack-years > 20 pack-years 10. Alcohol use: Number of days drinking per week: ____ Number of drinks per day: _ 11. Depression diagnosis: Previously diagnosed Currently diagnosed

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 **Anatomy**

FIGURE 11.1. Sagittal view of the face and neck depicting the subdivisions of the pharynx.

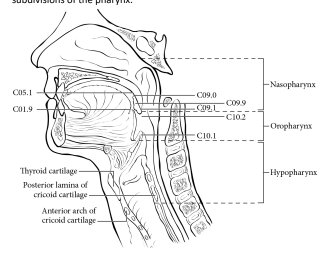
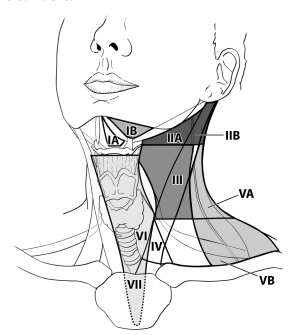


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

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|---------------------|--|
| Physician Signature | Date/Time |

| Hospital Name/Address | Patient Name/Information |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to maxillary sinus mucosa with no erosion or destruction of bone |
| | T2 | Tumor causing bone erosion or destruction including extension into the hard palate and/or middle nasal meatus, except extension to posterior wall of maxillary sinus and pterygoid plates |
| | T3 | Tumor invades any of the following: bone of the posterior wall of maxillary sinus, subcutaneous tissues, floor or medial wall of orbit, pterygoid fossa, ethmoid sinuses |
| | T4 | Moderately advanced or very advanced local disease |
| | T4a | Moderately advanced local disease Tumor invades anterior orbital contents, skin of cheek, pterygoid plates, infratemporal fossa, cribriform plate, sphenoid or frontal sinuses |
| | T4b | Very advanced local disease Tumor invades any of the following: orbital apex, dura, brain, middle cranial fossa, cranial nerves other than maxillary division of trigeminal nerve (V2), nasopharynx, or clivus |

| / | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | N Category | N Criteria |
|-----|--|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) with clinically overt ENE(+) |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any node(s) with clinically overt ENE (ENE _c) |
| Not | Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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4.2.2 Pathological N (pN)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); |
| | | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); |
| | | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) |
| | N2a | Metastasis in single ipsilateral node 3 cm or less in greatest dimension and ENE(+); |
| | | or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); |
| | | or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); |
| | | or a single contralateral node of any size and ENE(+) |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); |
| | | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); |
| | | or a single contralateral node of any size and ENE(+) |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|--------------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | III |
| | T1,T2,T3 | N1 | M0 | III |
| | T4a | N0,N1 | M0 | IVA |
| | T1,T2,T3,T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

| 6 Registry Data Col | lection Variables |
|--|--|
| See chapter for more details or | these variables. |
| 1. ENE clinical (select one): | ☐ Present/Positive (+) ☐ Absent/Negative (-) |
| 2. ENE pathological (select one |): Present/Positive (+) Absent/Negative (-) |
| 3. Extent of microscopic ENE (d tissue): | istance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal |
| 4. Perineural invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 5. Lymphovascular invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 6. Performance status (0-5): | |
| 7. Tobacco use and pack-year: | Never≤ 10 pack-years> 10 but ≤ 20 pack-years> 20 pack-years |
| | of days drinking per week: of drinks per day: |
| 9. Depression diagnosis: | ☐ Previously diagnosed ☐ Currently diagnosed |
| This form continues on the nex | rt page. |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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12.1. Maxillary Sinus

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

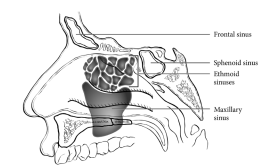
8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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9 Anatomy

FIGURE 12.1. Primary sites of the paranasal sinuses.



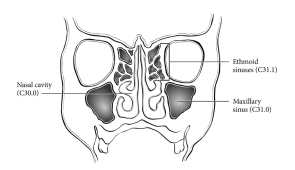
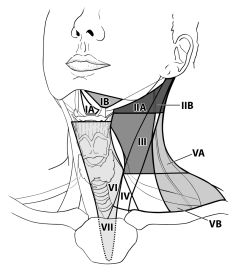


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

1 Terms of Use

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---|--|--|
| | TX | Primary tumor cannot be assessed | |
| | Tis | Carcinoma in situ | |
| | T1 | Tumor restricted to any one subsite, with or without bony invasion | |
| | T2 | Tumor invading two subsites in a single region or extending to involve an adjacent region within the nasoethmoidal complex, with or without bony invasion | |
| | T3 | Tumor extends to invade the medial wall or floor of the orbit, maxillary sinus, palate, or cribriform plate | |
| | T4 Moderately advanced or very advanced local disease | | |
| | T4a | Moderately advanced local disease Tumor invades any of the following: anterior orbital contents, skin of nose or cheek, minimal extension to anterior cranial fossa, pterygoid plates, sphenoid or frontal sinuses | |
| | T4b | Very advanced local disease Tumor invades any of the following: orbital apex, dura, brain, middle cranial fossa, cranial nerves other than (V2), nasopharynx, or clivus | |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | N Category | N Criteria | |
|--|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | | or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–) | | |
| | N2b Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | | |
| N2c Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | | | |
| | N3 Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); | | |
| | | or metastasis in any node(s) with clinically overt ENE(+) | |
| | N3a Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | | |
| | N3b Metastasis in any node(s) with clinically overt ENE (ENE _c) | | |
| Not | Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the | | |
| low | lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(–) or ENE(+). | | |

| 1 | N Suffix | Definition |
|---|----------|--|
| | н эйјјіх | Definition |
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | 1 | Metastasis helow the lower horder of the cricoid |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| N Category | N Criteria | |
|---|--|--|
| NX | Regional lymph nodes cannot be assessed | |
| N0 | No regional lymph node metastasis | |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| N2 Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | | |
| | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N2a Metastasis in single ipsilateral node 3 cm or less in greatest dimension and ENE(+); | | |
| or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(- | | |
| N2b Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | | |
| N2c Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and | | |
| N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); | |
| | or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | |
| | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |
| N3a Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | | |
| N3b Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | |
| | or multiple ipsilateral, contralateral or bilateral nodes, any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |
| | NX N0 N1 N2 N2a N2b N2c N3 | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition | |
|---|--|---|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) | f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid | |
| | L | Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| * | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When Tis | And N is | And M is | Then the stage group is |
|---|--------------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | NO | M0 | II |
| | T3 | NO | M0 | III |
| | T1,T2,T3 | N1 | M0 | III |
| | T4a | N0,N1 | M0 | IVA |
| | T1,T2,T3,T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

| 6 Registry Data Col | lection Variables |
|--|---|
| See chapter for more details or | n these variables. |
| 1. ENE clinical (select one): | ☐ Present/Positive (+) ☐ Absent/Negative (-) |
| 2. ENE pathological (select one |): Present/Positive (+) Absent/Negative (-) |
| 3. Extent of microscopic ENE (d tissue): | listance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal |
| 4. Perineural invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 5. Lymphovascular invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 6. Performance status (0-5): | |
| 7. Tobacco use and pack-year: | Never ≤ 10 pack-years > 10 but ≤ 20 pack-years > 20 pack-years |
| | of days drinking per week: of drinks per day: |
| 9. Depression diagnosis: | Previously diagnosed Currently diagnosed |
| This form continues on the nex | ct page. |

| Hospital Name/Address | Patient Name/Information |
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12.2. Nasal Cavity and Ethmoid Sinus

7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|------------------------------|-----------------------|--|
| | GX Grade cannot be assessed | | |
| | G1 Well differentiated | | |
| | G2 Moderately differentiated | | |
| | G3 | Poorly differentiated | |

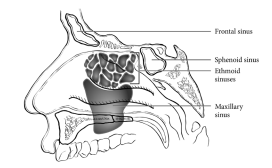
8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| ent Name/Information |
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9 Anatomy

FIGURE 12.1. Primary sites of the paranasal sinuses.



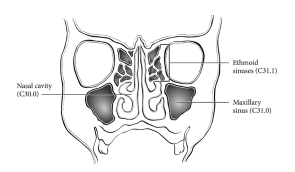
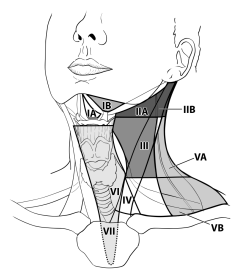


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to one subsite of supraglottis with normal vocal cord mobility |
| | T2 | Tumor invades mucosa of more than one adjacent subsite of supraglottis or glottis or region outside the |
| | | supraglottis (e.g., mucosa of base of tongue, vallecula, medial wall of pyriform sinus) without fixation of the larynx |
| | ТЗ | Tumor limited to larynx with vocal cord fixation and/or invades any of the following: postcricoid area, preepiglottic space, paraglottic space, and/or inner cortex of thyroid cartilage |
| | T4 | Moderately advanced or very advanced |
| | T4a | Moderately advanced local disease Tumor invades through the outer cortex of the thyroid cartilage and/or invades tissues beyond the larynx (e.g., trachea, soft tissues of neck including deep extrinsic muscle of the tongue, strap muscles, thyroid, or esophagus) |
| | T4b | Very advanced local disease Tumor invades prevertebral space, encases carotid artery, or invades mediastinal structures |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| N Category | N Criteria |
|------------|---|
| NX | Regional lymph nodes cannot be assessed |
| N0 | No regional lymph node metastasis |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| N2 | Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(–); or metastasis in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(–) |
| N2a | Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| N3 | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-); or metastasis in any lymph node(s) with clinically overt ENE(+) |
| N3a | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-) |
| N3b | Metastasis in any lymph node(s) with clinically overt ENE(+) |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix Definition | |
|---|--|---|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid |
| | L Metastasis below the lower border of the cricoid | |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| | gory N Criteria | |
|---|--|--|
| NX | Regional lymph nodes cannot be assessed | |
| N0 | No regional lymph node metastasis | |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastasis in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N2a | Metastasis in a single ipsilateral node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | or metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N3 Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-); | | |
| | or metastasis in a single ipsilateral node, larger than 3 cm in greatest dimension and ENE(+); | |
| | or multiple ipsilateral, contralateral, or bilateral lymph nodes any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |
| N3a | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-) | |
| N3b | Metastasis in a single ipsilateral node, larger than 3 cm in greatest dimension and ENE(+); | |
| | or multiple ipsilateral, contralateral, or bilateral lymph nodes any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix Definition | |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U Metastasis above the lower border of the cricoid | |
| | L Metastasis below the lower border of the cricoid | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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This form continues on the next page.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | NO | M0 | III |
| | T1, T2, T3 | N1 | M0 | III |
| | T4a | N0, N1 | M0 | IVA |
| | T1, T2, T3, T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

Registry Data Collection Variables See chapter for more details on these variables. 1. ENE clinical (select one): Present/Positive (+) ☐ Absent/Negative (-) 2. ENE pathological (select one): Present/Positive (+) ☐ Absent/Negative (-) 3. Extent of microscopic ENE (distance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal tissue): ☐ Intratumoral: ■ Multifocal 4. Perineural invasion: ☐ Focal Extratumoral: Focal ■ Multifocal 5. Lymphovascular invasion: Intratumoral: Focal Multifocal Extratumoral: Focal ■ Multifocal 6. Performance status (0-5): 7. Tobacco use and pack-year: □ Never ≤ 10 pack-years _ > 10 but ≤ 20 pack-years > 20 pack-years 8. Alcohol use: Number of days drinking per week: _____ Number of drinks per day: 9. Depression diagnosis: Previously diagnosed Currently diagnosed

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|-----------------------------|---------------------------|
| | GX Grade cannot be assessed | |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 13.1. Anatomical sites and subsites of the three regions of the larynx: supraglottis, glottis, and subglottis. Supraglottis (C32.1) subsites include suprahyoid epiglottis (i), aryepiglottic fold, laryngeal aspect (ii), infrahyoid epiglottis (iv), and ventricular bands or false cords (v).

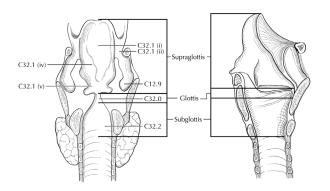


FIGURE 13.2. Anatomical sites and subsites of the supraglottis and glottis. Supraglottis (C32.1) subsites include suprahyoid epiglottis (i), aryepiglottic fold, laryngeal aspect (ii), arytenoids (iii), and ventricular bands or false cords (v). Glottis (C32.0) subsites include vocal cords (i), anterior commissure (ii), and posterior commissure (iii).

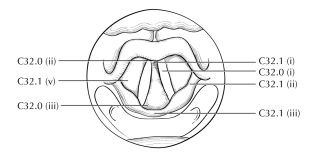
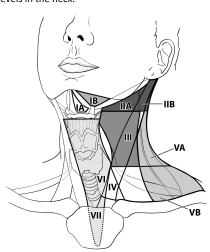


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name | |
|----------|-----------------------|--|
| IA | Submental | |
| IB | Submandibular | |
| IIA, IIB | Upper Jugular | |
| III | Middle Jugular | |
| IV | Lower Jugular | |
| VA, VB | Posterior Triangle | |
| VI | Anterior Compartment | |
| VII | Superior Mediastinal | |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

1 Terms of Use

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2 Instructions

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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to the vocal cord(s) (may involve anterior or posterior commissure) with normal mobility |
| | T1a | Tumor limited to one vocal cord |
| | T1b | Tumor involves both vocal cords |
| | T2 | Tumor extends to supraglottis and/or subglottis, and/or with impaired vocal cord mobility |
| | Т3 | Tumor limited to the larynx with vocal cord fixation and/or invasion of paraglottic space and/or inner cortex of the thyroid cartilage |
| | T4 | Moderately advanced or very advanced |
| | T4a | Moderately advanced local disease Tumor invades through the outer cortex of the thyroid cartilage and/or invades tissues beyond the larynx (e.g., trachea, cricoid cartilage, soft tissues of neck including deep extrinsic muscle of the tongue, strap muscles, thyroid, or esophagus) |
| | T4b | Very advanced local disease Tumor invades prevertebral space, encases carotid artery, or invades mediastinal structures |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria |
|-----|----------------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–); |
| | | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); |
| | | or metastasis in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2a | Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-); |
| | | or metastasis in any lymph node(s) with clinically overt ENE(+) |
| | N3a | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any lymph node(s) with clinically overt ENE(+) |
| Not | te. A designation of | "U" or "I" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

|) | |
|-----------------|--------|
|); | |
| | |
| n and ENE(-); | |
| nension and EN | E(-) |
| | |
| dimension and | ENE(-) |
| (-) | |
| nsion and ENE | (-) |
| | |
| | |
| | |
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| | |
| | |
| | |
| isoid (II) or b | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | NO | M0 | Ш |
| | T1, T2, T3 | N1 | M0 | III |
| | T4a | N0, N1 | M0 | IVA |
| | T1, T2, T3, T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

| 6 | Registry Data Coll | ection Variables |
|----------------|------------------------------|--|
| See c | chapter for more details on | these variables. |
| 1. EN | IE clinical (select one): | Present/Positive (+) Absent/Negative (-) |
| 2. EN | IE pathological (select one) | Present/Positive (+) Absent/Negative (-) |
| 3. Ex | · | stance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal |
| 4. Pe | rineural invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 5. Ly ı | mphovascular invasion: | ☐ Intratumoral: ☐ Focal ☐ Multifocal ☐ Extratumoral: ☐ Focal ☐ Multifocal |
| 6. Pe | rformance status (0-5): | |
| 7. To | bacco use and pack-year: | Never≤ 10 pack-years> 10 but ≤ 20 pack-years> 20 pack-years |
| 8. Ald | | of days drinking per week: of drinks per day: |
| 9. De | epression diagnosis: | ☐ Previously diagnosed ☐ Currently diagnosed |
| Thic | form continues on the nev | · naga |

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|------------------------------|-----------------------|--|
| | GX Grade cannot be assessed | | |
| | G1 | Well differentiated | |
| | G2 Moderately differentiated | | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description | |
|---|-------------------------|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

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| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 13.1. Anatomical sites and subsites of the three regions of the larynx: supraglottis, glottis, and subglottis. Supraglottis (C32.1) subsites include suprahyoid epiglottis (i), aryepiglottic fold, laryngeal aspect (ii), infrahyoid epiglottis (iv), and ventricular bands or false cords (v).

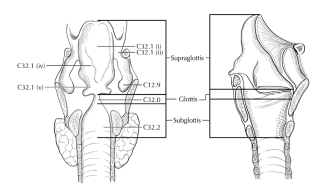


FIGURE 13.2. Anatomical sites and subsites of the supraglottis and glottis. Supraglottis (C32.1) subsites include suprahyoid epiglottis (i), aryepiglottic fold, laryngeal aspect (ii), arytenoids (iii), and ventricular bands or false cords (v). Glottis (C32.0) subsites include vocal cords (i), anterior commissure (ii), and posterior commissure (iii).

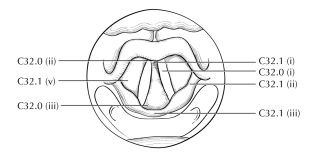
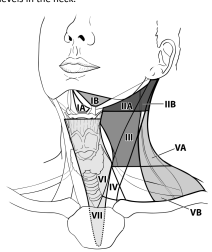


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria : First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to the subglottis |
| | T2 | Tumor extends to vocal cord(s) with normal or impaired mobility |
| | Т3 | Tumor limited to larynx with vocal cord fixation and/or invasion of paraglottic space and/or inner cortex of the thyroid cartilage |
| | T4 | Moderately advanced or very advanced |
| | T4a | Moderately advanced local disease Tumor invades cricoid or thyroid cartilage and/or invades tissues beyond the larynx (e.g., trachea, soft tissues of neck including deep extrinsic muscles of the tongue, strap muscles, thyroid, or esophagus) |
| | T4b | Very advanced local disease Tumor invades prevertebral space, encases carotid artery, or invades mediastinal structures |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria | |
|--|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| | N2 Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | | |
| | | or metastasis in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2a | Metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| | N2c Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) | | |
| | N3 Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-); | | |
| | or metastasis in any lymph node(s) with clinically overt ENE(+) | | |
| | N3a | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-) | |
| | N3b | Metastasis in any lymph node(s) with clinically overt ENE(+) | |
| Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the | | | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information | |
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4.2.2 Pathological N (pN)

| N Category | N Criteria | |
|---|--|--|
| NX | Regional lymph nodes cannot be assessed | |
| N0 | No regional lymph node metastasis | |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| N2 Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | | |
| | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastasis in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N2a | Metastasis in a single ipsilateral node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | or metastasis in a single ipsilateral node, larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(–) | |
| N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N3 Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-); or metastasis in a single ipsilateral node, larger than 3 cm in greatest dimension and ENE(+); | | |
| | | |
| or a single contralateral node of any size and ENE(+) | | |
| N3a | Metastasis in a lymph node, larger than 6 cm in greatest dimension and ENE(-) | |
| N3b Metastasis in a single ipsilateral node, larger than 3 cm in greatest dimension and ENE(+); | | |
| or multiple ipsilateral, contralateral, or bilateral lymph nodes any with ENE(+); | | |
| | or a single contralateral node of any size and ENE(+) | |
| | N0 N1 N2 N2a N2b N2c N3 | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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This form continues on the next page.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | NO | M0 | III |
| | T1, T2, T3 | N1 | M0 | III |
| | T4a | N0, N1 | M0 | IVA |
| | T1, T2, T3, T4a | N2 | M0 | IVA |
| | Any T | N3 | M0 | IVB |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

Registry Data Collection Variables See chapter for more details on these variables. 1. ENE clinical (select one): Present/Positive (+) ☐ Absent/Negative (-) 2. ENE pathological (select one): Present/Positive (+) ☐ Absent/Negative (-) 3. Extent of microscopic ENE (distance of extension from the native lymph node capsule to the farthest point of invasion in the extranodal tissue): ☐ Intratumoral: ■ Multifocal 4. Perineural invasion: ☐ Focal Extratumoral: Focal ■ Multifocal 5. Lymphovascular invasion: Intratumoral: Focal Multifocal Extratumoral: Focal ■ Multifocal 6. Performance status (0-5): 7. Tobacco use and pack-year: □ Never ≤ 10 pack-years __ > 10 but ≤ 20 pack-years > 20 pack-years 8. Alcohol use: Number of days drinking per week: _____ Number of drinks per day: 9. Depression diagnosis: Previously diagnosed Currently diagnosed

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|----|---------------------------|--|
| | GX | Grade cannot be assessed | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 13.1. Anatomical sites and subsites of the three regions of the larynx: supraglottis, glottis, and subglottis. Supraglottis (C32.1) subsites include suprahyoid epiglottis (i), aryepiglottic fold, laryngeal aspect (ii), infrahyoid epiglottis (iv), and ventricular bands or false cords (v).

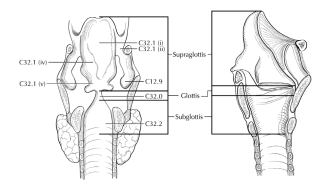
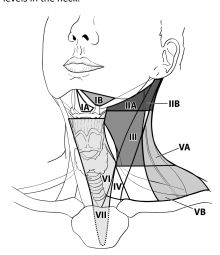


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | T3 | Tumors limited to the mucosa and immediately underlying soft tissue, regardless of thickness or greatest |
| | | dimension; for example, polypoid nasal disease, pigmented or nonpigmented lesions of the oral cavity, pharynx, |
| | | or larynx |
| | T4 | Moderately advanced or very advanced disease |
| | T4a | Moderately advanced disease |
| | | Tumor involving deep soft tissue, cartilage, bone, or overlying skin |
| | T4b | Very advanced disease |
| | | Tumor involving brain, dura, skull base, lower cranial nerves (IX, X, XI, XII), masticator space, carotid artery, |
| | | prevertebral space, or mediastinal structures |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastases |
| | N1 | Regional lymph node metastases present |

| N Suffix Definition | | |
|---------------------|------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no prognostic stage grouping proposed at this time.

| Hospital Name/Address | Patient Name/Information |
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14. Mucosal Melanoma of the Head and Neck

| 6 | Registry Data | a Collection Variables | | |
|------|--|--|--------------------------|--|
| See | See chapter for more information on these variables. | | | |
| 1. S | ize of lymph nodes: | | | |
| 2. E | xtracapsular extensio | on from lymph node for head and neck: | | |
| 3. H | ead and neck lymph | nodes levels: Levels I–III | | |
| 4. H | ead and neck lymph | nodes levels: Levels IV–V | | |
| 5. H | ead and neck lymph | nodes levels: Levels VI–VII | | |
| 6. C | ther lymph node gro | up: | | |
| 7. C | linical location of cer | vical nodes: | | |
| 8. E | NE clinical (select one | e): Present/Positive (+) Absent/Negative (-) | | |
| 9. E | NE pathological (sele | ct one): | | |
| 10. | Tumor thickness: | | | |
| 7 | Histologic Gr | ada (G) | | |
| _ | Histologic di | aue (G) | | |
| The | re is no recommende | d histologic grading system at this time. | | |
| 8 | Lymphovasco | ular Invasion (LVI) | | |
| | | | | |
| ✓ | Component of LVI Coding | Description | | |
| | 0 | LVI not present (absent)/not identified | | |
| | 1 | LVI present/identified, NOS | | |
| | 2 | Lymphatic and small vessel invasion only (L) | | |
| | 3 | Venous (large vessel) invasion only (V) | | |
| | 4 | BOTH lymphatic and small vessel AND venous | (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | | |
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| Hos | pital Name/Address | | Patient Name/Information | |
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9 Anatomy

FIGURE 14.2. T3 is defined as mucosal disease. Involvement of the lateral wall nasal cavity, inferior turbinate is illustrated, as well as septum, hard palate, ethmoid, and nasal vestibule.

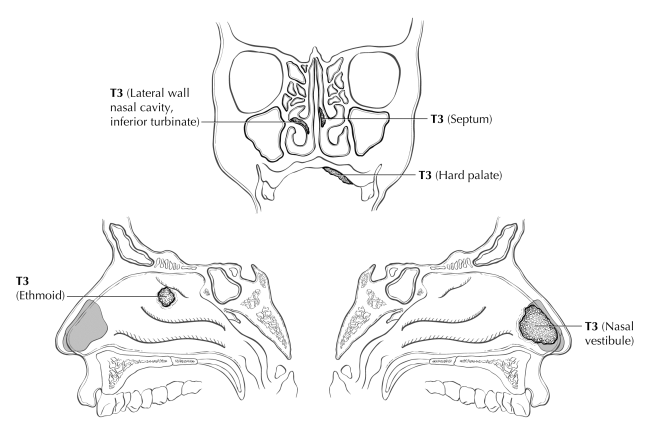
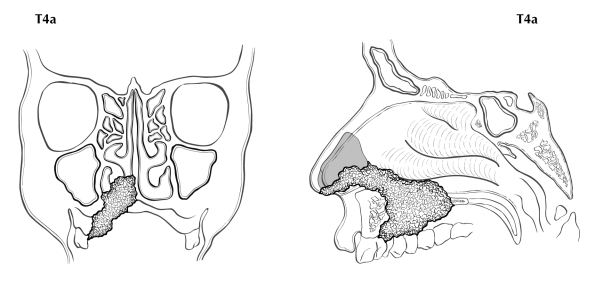


FIGURE 14.3. T4a is defined as moderately advanced disease, with tumor involving deep soft tissue, cartilage, bone, or overlying skin.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 14.4. T4b is defined as very advanced disease, with tumor involving the brain as illustrated, or also involving dura, lower cranial nerves (IX, X, XI, XII), masticator space, carotid artery, prevertebral space, or mediastinal structures.

T4b

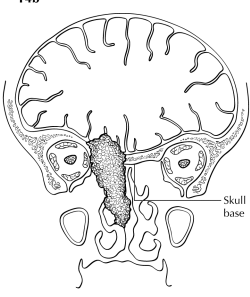
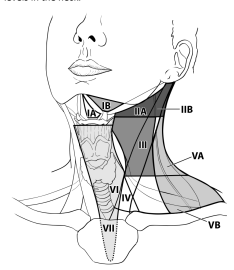


FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ |
| | T1 | Tumor smaller than or equal to 2 cm in greatest dimension |
| | T2 | Tumor larger than 2 cm, but smaller than or equal to 4 cm in greatest dimension |
| | T3 | Tumor larger than 4 cm in maximum dimension or minor bone erosion or perineural invasion or deep invasion* |
| | T4 | Tumor with gross cortical bone/marrow, skull base invasion and/or skull base foramen invasion |
| | T4a | Tumor with gross cortical bone/marrow invasion |
| | T4b | Tumor with skull base invasion and/or skull base foramen involvement |

*Deep invasion is defined as invasion beyond the subcutaneous fat or > 6 mm (as measured from the granular layer of adjacent normal epidermis to the base of the tumor); perineural invasion for T3 classification is defined as tumor cells within the nerve sheath of a nerve lying deeper than the dermis or measuring 0.1 mm or larger in caliber, or presenting with clinical or radiographic involvement of named nerves without skull base invasion or transgression.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | N Category | N Criteria |
|-----|-----------------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) |
| | N2 | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2a | Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) |
| | N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N2c | Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-) |
| | N3 | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); or metastasis in any node(s) and clinically overt ENE [ENE(+)] |
| | N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) |
| | N3b | Metastasis in any node(s) and ENE(+) |
| Not | e: A designation of ' | 'U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the |

lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|----------|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

| Hospital Name/Address | Patient Name/Information |
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4.2.2 Pathological N (pN)

| N Category | N Criteria | |
|--|--|--|
| NX | Regional lymph nodes cannot be assessed | |
| N0 | No regional lymph node metastasis | |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(-) | |
| N2 | Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+); | |
| | or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-); | |
| | or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(-); | |
| | or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension, ENE(-) | |
| N2a | Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+); | |
| | or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(-) | |
| N2b | Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(-) | |
| N2c | Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(-) | |
| N3 Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-); | | |
| or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | |
| | or multiple ipsilateral, contralateral, or bilateral nodes, any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |
| N3a | Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(-) | |
| N3b Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+); | | |
| | or multiple ipsilateral, contralateral, or bilateral nodes, any with ENE(+); | |
| | or a single contralateral node of any size and ENE(+) | |
| | NX N0 N1 N2 N2a N2b N2c N3 | |

Note: A designation of "U" or "L" may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological ENE should be recorded as ENE(-) or ENE(+).

| ✓ | N Suffix | Definition |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |
| | U | Metastasis above the lower border of the cricoid |
| | L | Metastasis below the lower border of the cricoid |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

This form continues on the next page.

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | III |
| | T1 | N1 | M0 | III |
| | T2 | N1 | M0 | III |
| | T3 | N1 | M0 | III |
| | T1 | N2 | M0 | IV |
| | T2 | N2 | M0 | IV |
| | T3 | N2 | M0 | IV |
| | Any T | N3 | M0 | IV |
| | T4 | Any N | M0 | IV |
| | Any T | Any N | M1 | IV |

| 6 Registry Data Coll | ection Variables | | | |
|-----------------------------------|----------------------------|------------------------|-----------------------------------|------------------|
| See chapter for more details on | these variables. | | | |
| 1. ENE clinical (select one): | Present/Positive (+) | ☐ Absent/Negative | (-) | |
| 2. ENE pathological (select one) | Present/Positive (+) | Absent/Negative | (-) | |
| 3. Preoperative clinical tumor di | ameter in millimeters: | | | |
| | | | dermis to the base of the tumor): | |
| and/or tissue levei: | | | | |
| 5. Perineural invasion: | Absent Present, enter size | e in mm: | | |
| 6. Primary site location: | ☐ temple ☐ cheek ☐ ear | ☐ lip, hair-bearing | ☐ lip, vermilion border | |
| 7. High-risk histologic features: | poor differentiation | desmoplasia | sarcomatoid differentiation | undifferentiated |
| 8. Immune status: | munosuppressedimmur | nosuppressed, specify: | | |
| 9. Depression diagnosis: | Previously diagnosed | Currently diagnos | ed | |
| 10. Comorbidities: | and performance sta | atus (0-5): | | |
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| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

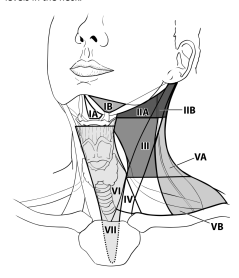
| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description |
|---|-------------------------|--|
| | o Couring | LVI not present (absent)/not identified |
| | U | |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 5.1. Schematic indicating the location of the lymph node levels in the neck.



| Level | Lymph Node Group Name |
|----------|-----------------------|
| IA | Submental |
| IB | Submandibular |
| IIA, IIB | Upper Jugular |
| III | Middle Jugular |
| IV | Lower Jugular |
| VA, VB | Posterior Triangle |
| VI | Anterior Compartment |
| VII | Superior Mediastinal |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
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| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | High-grade dysplasia, defined as malignant cells confined to the epithelium by the basement membrane |
| | T1 | Tumor invades the lamina propria, muscularis mucosae, or submucosa |
| | T1a | Tumor invades the lamina propria or muscularis mucosae |
| | T1b | Tumor invades the submucosa |
| | T2 | Tumor invades the muscularis propria |
| | T3 | Tumor invades adventitia |
| | T4 | Tumor invades adjacent structures |
| | T4a | Tumor invades the pleura, pericardium, azygos vein, diaphragm, or peritoneum |
| | T4b | Tumor invades other adjacent structures, such as the aorta, vertebral body, or airway |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Ī | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in one or two regional lymph nodes |
| | N2 | Metastasis in three to six regional lymph nodes |
| | N3 | Metastasis in seven or more regional lymph nodes |

| ✓ | N Suffix | Definition |
|----------|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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16.1. Esophagus and Esophagogastric Junction: Squamous Cell Carcinoma

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated, undifferentiated |

5.2 Definition of Location (L)

| √ | Location Category | Location Criteria | |
|----------|---|---|--|
| | X | Location Unknown | |
| | Upper | Cervical esophagus to lower border of azygos vein | |
| | Middle | Lower border of azygos vein to lower border of inferior pulmonary vein | |
| | Lower | Lower border of inferior pulmonary vein to stomach, including gastroesophageal junction | |
| Not | Note: Location is defined by the position of the epicenter of the tumor in the esophagus. | | |

| Hospital Name/Address | Patient Name/Information |
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6 AJCC Prognostic Stage Groups

In addition to anatomic tumor depth, nodal status, and metastasis (see Definitions of AJCC TNM), other prognostic factors - grade (G) and location (L) - affect outcome, and therefore staging, of squamous cell carcinoma. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6.1 Clinical (cTNM)

| 1 | When cT is | And cN is | And M is | Then the stage group is |
|---|---------------|--------------|-------------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0-1 | M0 | - |
| | T2 | N0-1 | M0 | II |
| | T3 | N0 | M0 | П |
| | T3 | N1 | M0 | Ш |
| | T1-3 | N2 | M0 | Ш |
| | T4 | N0-2 | M0 | IVA |
| | Any T | N3 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6.2 Pathological (pTNM)

| | When | And | And | And | And | Then |
|---|-------|----------|-----|------|-----------------------|-------|
| | рТ | pΝ | М | G | Location | the |
| ✓ | is | is | is | is | is | stage |
| | | | | | | group |
| | | | | | | is |
| | Tis | N0 | M0 | N/A | Any | 0 |
| | T1a | N0 | M0 | G1 | Any | IA |
| | T1a | N0 | M0 | G2-3 | Any | IB |
| | T1a | N0 | M0 | GX | Any | IA |
| | T1b | N0 | M0 | G1-3 | Any | IB |
| | T1b | N0 | M0 | GX | Any | IB |
| | T2 | N0 | M0 | G1 | Any | IB |
| | T2 | N0 | M0 | G2-3 | Any | IIA |
| | T2 | N0 | M0 | GX | Any | IIA |
| | T3 | N0 | M0 | G1-3 | Lower | IIA |
| | T3 | N0 | M0 | G1 | Upper/middle | IIA |
| | T3 | N0 | M0 | G2-3 | Upper/middle | IIB |
| | T3 | N0 | M0 | GX | Lower/upper middle | IIB |
| | T3 | N0 | M0 | Any | Location X | IIB |
| | T1 | N1 | M0 | Any | Any | IIB |
| | T1 | N2 | M0 | Any | Any | IIIA |
| | T2 | N1 | M0 | Any | Any | IIIA |
| | T2 | N2 | M0 | Any | Any | IIIB |
| | T3 | N1- 2 | M0 | Any | Any | IIIB |
| | T4a | N0- 1 | M0 | Any | Any | IIIB |
| | T4a | N2 | M0 | Any | Any | IVA |
| | T4b | N0- 2 | M0 | Any | Any | IVA |
| | Any T | N3 | M0 | Any | Any | IVA |
| | Any T | Any N | M1 | Any | Any | IVB |

6.3 Postneoadjuvant Therapy (ypTNM)

| | | • | . , ,,, | • |
|---|----------------|---------------|-------------|-------------------------|
| 1 | When ypT is | And ypN is | And M is | Then the stage group is |
| | T0-2 | N0 | M0 | 1 |
| | T3 | N0 | M0 | П |
| | T0-2 | N1 | M0 | IIIA |
| | T3 | N1 | M0 | IIIB |
| | T0-3 | N2 | M0 | IIIB |
| | T4a | N0 | M0 | IIIB |
| | T4a | N1-2 | M0 | IVA |
| | T4a | NX | M0 | IVA |
| | T4b | N0-2 | M0 | IVA |
| | Any T | N3 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

| Hospital Name/Address | Patient Name/Information |
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16.1. Esophagus and Esophagogastric Junction: Squamous Cell Carcinoma

| F | Registry Data Collection Variables | | | | |
|-------|---|--|--|--|--|
| e cha | pter for more de | tails on these variables. | | | |
| 1. | Clinical stagir | ng modalities (endoscopy and biopsy, EUS, EUS-FNA, CT, PET/CT): | | | |
| 2. | Tumor length | 1: | | | |
| 3. | Depth of inva | ision: | | | |
| 4. | Number of no | odes involved, clinical: | | | |
| 5. | Number of no | odes involved, pathological: | | | |
| 6. | Location of n | odal disease, clinical: | | | |
| 7. | Location of n | odal disease, pathological: | | | |
| 8. | Sites of meta | stasis, if applicable: | | | |
| 9. | Presence of s | kip lesions: T(m): | | | |
| 10 |). Perineural inv | vasion: | | | |
| 11 | 1. LVI: lymphatic vascular both | | | | |
| 12 | | | | | |
| 13 | | | | | |
| 14 | I. Chemotherar | by: | | | |
| 15 | 5. Chemoradiat | ion therapy (for ypTNM): | | | |
| 16 | 6. Surgical margin: negative microscopic macroscopic | | | | |
| | | | | | |
| L | Lymphovascular Invasion (LVI) | | | | |
| | | | | | |
| | omponent of /I Coding | Description | | | |
| 0 | | LVI not present (absent)/not identified | | | |
| 1 | | LVI present/identified, NOS | | | |
| 2 | | Lymphatic and small vessel invasion only (L) | | | |
| 3 | | Venous (large vessel) invasion only (V) | | | |
| 4 | | BOTH lymphatic and small vessel AND venous (large vessel) invasion | | | |
| 9 | | Presence of LVI unknown/indeterminate | | | |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 16.1. Anatomy of esophageal cancer primary site, including typical endoscopic measurements of each region measured from the incisors. Exact measurements depend on body size and height. Location of cancer primary site is defined by cancer epicenter. EGJ, esophagogastric junction; LES, lower esophageal sphincter; UES, upper esophageal sphincter.

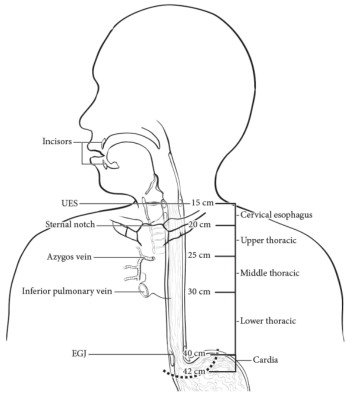
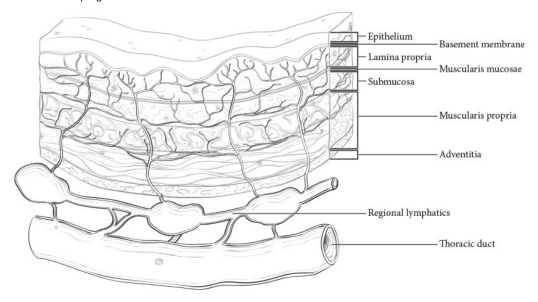
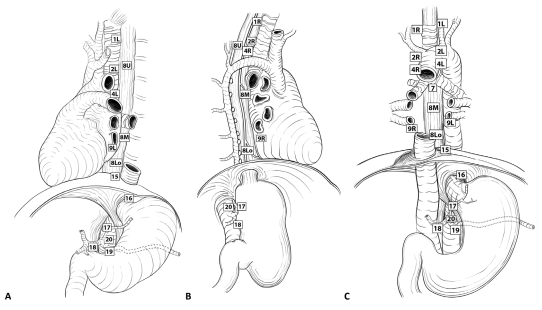


FIGURE 16.2. Esophageal wall.



| Hospital Name/Address | Patient Name/Information | |
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FIGURE 16.3. (A–C) Lymph node maps for esophageal cancer. Regional lymph node stations for staging esophageal cancer from left (A), right (B), and anterior (C). 1R, Right lower cervical paratracheal nodes, between the supraclavicular paratracheal space and apex of the lung. 2R, Right upper paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and the apex of the lung. 2L, Left upper paratracheal nodes, between the top of the aortic arch and the apex of the lung. 4R, Right lower paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and cephalic border of the azygos vein. 4L, Left lower paratracheal nodes, between the top of the aortic arch and the carina. 7, Subcarinal nodes, caudal to the carina of the trachea. 8U, Upper thoracic paraesophageal lymph nodes, from the apex of the lung to the tracheal bifurcation. 8M, Middle thoracic paraesophageal lymph nodes, from the tracheal bifurcation to the caudal margin of the inferior pulmonary vein. 8Lo, Lower thoracic paraesophageal lymph nodes, from the caudal margin of the inferior pulmonary ligament nodes, within the right inferior pulmonary ligament. 9L, Pulmonary ligament nodes, within the left inferior pulmonary ligament. 15, Diaphragmatic nodes, lying on the dome of the diaphragm and adjacent to or behind its crura. 16, Paracardial nodes, immediately adjacent to the gastroesophageal junction. 17, Left gastric nodes, along the course of the left gastric artery. 18, Common hepatic nodes, immediately on the proximal splenic artery. 20, Celiac nodes, at the base of the celiac artery.



| Physician Signature | Date/Time |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, 8th Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---|--|--|
| | TX | Tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | Tis | High-grade dysplasia, defined as malignant cells confined to the epithelium by the basement membrane | |
| | T1 | Tumor invades the lamina propria, muscularis mucosae, or submucosa | |
| | T1a Tumor invades the lamina propria or muscularis mucosae | | |
| | T1b Tumor invades the submucosa | | |
| | T2 Tumor invades the muscularis propria | | |
| | T3 Tumor invades adventitia | | |
| | T4 Tumor invades adjacent structures | | |
| | T4a | Tumor invades the pleura, pericardium, azygos vein, diaphragm, or peritoneum | |
| | T4b Tumor invades other adjacent structures, such as the aorta, vertebral body, or airway | | |

| Ī | ✓ | T Suffix | Definition |
|---|---|----------|---|
| | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|----------|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Metastasis in one or two regional lymph nodes | |
| | N2 Metastasis in three to six regional lymph nodes | | |
| | N3 | Metastasis in seven or more regional lymph nodes | |

| ✓ | N Suffix | Definition | |
|----------|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria | |
|---|------------|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | pM1 | Distant metastasis, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information |
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5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

| ✓ | G | G Definition | |
|---|----|---|--|
| | GX | GX Grade cannot be assessed | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated, undifferentiated | |

6 AJCC Prognostic Stage Groups

The requirements and rules for staging esophageal adenocarcinoma are similar to those for squamous cell carcinoma with regard to determining primary tumor stage, nodal status, and metastasis (see Definitions of AJCC TNM and G for squamous cell carcinoma). Whereas location of tumor is not a prognostic variable in adenocarcinoma of the esophagus, grade significantly affects outcome and therefore staging.

6.1 Clinical (cTNM)

| 1 | When cT is | And cN is | And M is | Then the stage group is |
|---|------------|-----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | IIA |
| | T2 | N0 | M0 | IIB |
| | T2 | N1 | M0 | III |
| | T3 | N0-1 | M0 | III |
| | T4a | N0-1 | M0 | III |
| | T1-T4a | N2 | M0 | IVA |
| | T4b | N0-2 | M0 | IVA |
| | Any T | N3 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6.2 Pathological (pTNM)

| 1 | When pT is | And pN is | And M is | And G is | Then the stage |
|---|------------|-----------|----------|----------|----------------|
| • | | | | | group is |
| | Tis | N0 | M0 | N/A | 0 |
| | T1a | N0 | M0 | G1 | IA |
| | T1a | N0 | M0 | GX | IA |
| | T1a | N0 | M0 | G2 | IB |
| | T1b | N0 | M0 | G1-2 | IB |
| | T1b | N0 | M0 | GX | IB |
| | T1 | N0 | M0 | G3 | IC |
| | T2 | N0 | M0 | G1-2 | IC |
| | T2 | N0 | M0 | G3 | IIA |
| | T2 | N0 | M0 | GX | IIA |
| | T1 | N1 | M0 | Any | IIB |
| | T3 | N0 | M0 | Any | IIB |
| | T1 | N2 | M0 | Any | IIIA |
| | T2 | N1 | M0 | Any | IIIA |
| | T2 | N2 | M0 | Any | IIIB |
| | T3 | N1-2 | M0 | Any | IIIB |
| | T4a | N0-1 | M0 | Any | IIIB |
| | T4a | N2 | M0 | Any | IVA |
| | T4b | N0-2 | M0 | Any | IVA |
| | Any T | N3 | M0 | Any | IVA |
| | Any T | Any N | M1 | Any | IVB |

| Hospital Name/Address | Patient Name/Information |
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6.3 Postneoadjuvant Therapy (ypTNM)

| 1 | When ypT is | And ypN is | And M is | Then the stage group is |
|---|-------------|------------|----------|-------------------------|
| | T0-2 | NO | M0 | 1 |
| | T3 | N0 | M0 | II |
| | T0-2 | N1 | M0 | IIIA |
| | T3 | N1 | M0 | IIIB |
| | T0-3 | N2 | M0 | IIIB |
| | T4a | N0 | M0 | IIIB |
| | T4a | N1-2 | M0 | IVA |
| | T4a | NX | M0 | IVA |
| | T4b | N0-2 | M0 | IVA |
| | Any T | N3 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

7 Registry Data Collection Variables

| | - | | |
|-------------|----------|------------|------------------|
| Saa chantar | tor more | dataile on | these variables. |
| | | | |

| 1. | Clinical staging modalities (endoscopy and biopsy, EUS, EUS-FNA, CT, PET/CT): | | |
|--------------------------|---|---|--|
| 2. | Tumor length: | | |
| 3. | Depth of invasion: | | |
| 4. | Number of no | odes involved, clinical: | |
| 5. | Number of no | odes involved, pathological: | |
| 6. | Location of no | odal disease, clinical: | |
| 7. | Location of no | odal disease, pathological: | |
| 8. | Sites of metas | stasis, if applicable: | |
| 9. | Presence of s | kip lesions: T(m): | |
| 10. | Perineural inv | vasion: | |
| 11. | LVI: | lymphatic vascular both | |
| 12. | . Extranodal extension: yes no | | |
| 13. | 8. HER2 Status: Positive Negative | | |
| 14. | I. Type of surgery: | | |
| 15. | 5. Chemotherapy: | | |
| 16. | Chemoradiati | on therapy (for ypTNM): | |
| 17. | . Surgical margin: negative microscopic macroscopic | | |
| Ly | Lymphovascular Invasion (LVI) | | |
| Component of Description | | Description | |
| LVI | Coding | | |
| 0 | | LVI not present (absent)/not identified | |
| 1 | | LVI present/identified, NOS | |
| 2 | | Lymphatic and small vessel invasion only (L) | |
| 3 | | Venous (large vessel) invasion only (V) | |
| 1 | | POTIL lymphatic and small yessel AND vaneus (large yessel) invasion | |

This form continues on the next page.

Presence of LVI unknown/indeterminate

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 16.1. Anatomy of esophageal cancer primary site, including typical endoscopic measurements of each region measured from the incisors. Exact measurements depend on body size and height. Location of cancer primary site is defined by cancer epicenter. EGJ, esophagogastric junction; LES, lower esophageal sphincter; UES, upper esophageal sphincter.

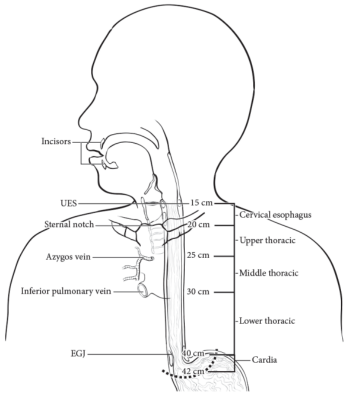
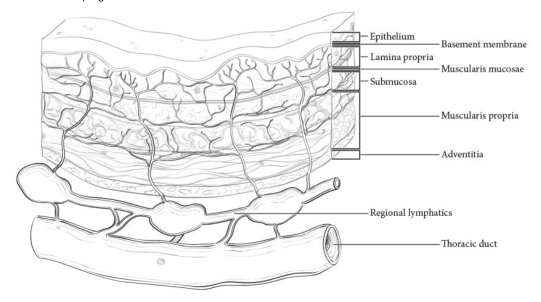
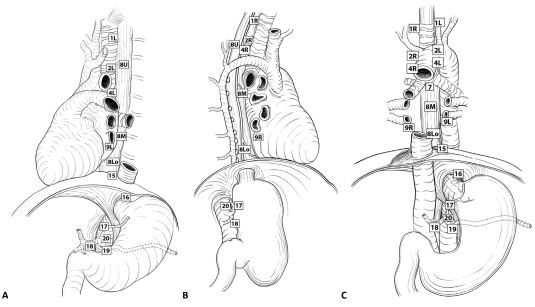


FIGURE 16.2. Esophageal wall.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 16.3. (A–C) Lymph node maps for esophageal cancer. Regional lymph node stations for staging esophageal cancer from left (A), right (B), and anterior (C). 1R, Right lower cervical paratracheal nodes, between the supraclavicular paratracheal space and apex of the lung. 2R, Right upper paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and the apex of the lung. 2L, Left upper paratracheal nodes, between the top of the aortic arch and the apex of the lung. 4R, Right lower paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and cephalic border of the azygos vein. 4L, Left lower paratracheal nodes, between the top of the aortic arch and the carina. 7, Subcarinal nodes, caudal to the carina of the trachea. 8U, Upper thoracic paraesophageal lymph nodes, from the apex of the lung to the tracheal bifurcation. 8M, Middle thoracic paraesophageal lymph nodes, from the tracheal bifurcation to the caudal margin of the inferior pulmonary vein. 8Lo, Lower thoracic paraesophageal lymph nodes, from the caudal margin of the inferior pulmonary ligament nodes, within the right inferior pulmonary ligament. 9L, Pulmonary ligament nodes, within the left inferior pulmonary ligament to or behind its crura. 16, Paracardial nodes, immediately adjacent to the gastroesophageal junction. 17, Left gastric nodes, along the course of the left gastric artery. 18, Common hepatic nodes, immediately on the proximal splenic artery. 20, Celiac nodes, at the base of the celiac artery.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information | |
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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | High-grade dysplasia, defined as malignant cells confined to the epithelium by the basement membrane |
| | T1 | Tumor invades the lamina propria, muscularis mucosae, or submucosa |
| | T1a | Tumor invades the lamina propria or muscularis mucosae |
| | T1b | Tumor invades the submucosa |
| | T2 | Tumor invades the muscularis propria |
| | T3 | Tumor invades adventitia |
| | T4 | Tumor invades adjacent structures |
| | T4a | Tumor invades the pleura, pericardium, azygos vein, diaphragm, or peritoneum |
| | T4b | Tumor invades other adjacent structures, such as the aorta, vertebral body, or airway |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Ī | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in one or two regional lymph nodes |
| | N2 | Metastasis in three to six regional lymph nodes |
| | N3 | Metastasis in seven or more regional lymph nodes |

| ✓ | N Suffix | Definition |
|----------|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated, undifferentiated |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| 6 | AJCC Pi | ognos | tic Stage Groups | | |
|------|--------------------------------------|---|--|--|--|
| The | re is no progi | nostic sta | ge group for other histologies arising in the es | ophagus and esophagogastric junction at this time. | |
| 7 | 7 Registry Data Collection Variables | | | | |
| See | chapter for r | nore deta | ills on these variables. | | |
| | 1. Clinic | Clinical staging modalities (endoscopy and biopsy, EUS, EUS-FNA, CT, PET/CT): | | | |
| | 2. Tumo | r length: | | | |
| | 3. Depth | of invasi | on: | | |
| | 4. Numb | er of nod | les involved, clinical: | | |
| | 5. Numb | er of nod | les involved, pathological: | | |
| | 6. Locat | ion of no | dal disease, clinical: | | |
| | 7. Locat | ion of not | dal disease, pathological: | | |
| | 8. Sites | of metast | asis, if applicable: | | |
| | 9. Prese | nce of ski | p lesions: T(m): | | |
| | 10. Perin | eural inva | sion: | | |
| | 11. LVI: | | ymphatic vascular bo | th | |
| | 12. Extra | nodal exte | ension: yes no | | |
| | 13. HER2 | Status: | Positive Negative | | |
| | 14. Type | of surgery | <i>;</i> : | | |
| | 15. Chem | otherapy | : | | |
| | 16. Chem | oradiatio | n therapy (for ypTNM): | | |
| | 17. Surgio | cal margin | n: negative microscopic | macroscopic | |
| | | | | | |
| 8 | Lymph | ovaccu | lar Invasion (LVI) | | |
| _ | Еупіріі | Jvascu | iai iiivasioii (Evi) | | |
| | Compone | nt of | Description | | |
| ✓ | LVI Codin | g | | | |
| | 0 | | LVI not present (absent)/not identified | | |
| | 2 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) | | |
| | 3 | | Venous (large vessel) invasion only (V) | | |
| | 4 | | BOTH lymphatic and small vessel AND venous | (large vessel) invasion | |
| | 9 | | Presence of LVI unknown/indeterminate | | |
| This | form contin | ues on th | e next page. | | |
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9 Anatomy

FIGURE 16.1. Anatomy of esophageal cancer primary site, including typical endoscopic measurements of each region measured from the incisors. Exact measurements depend on body size and height. Location of cancer primary site is defined by cancer epicenter. EGJ, esophagogastric junction; LES, lower esophageal sphincter; UES, upper esophageal sphincter.

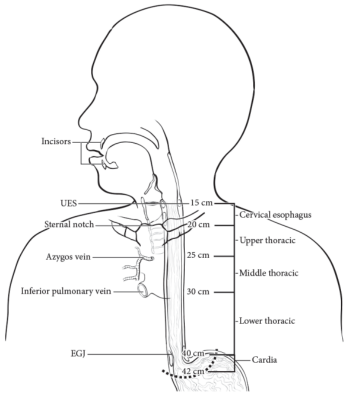
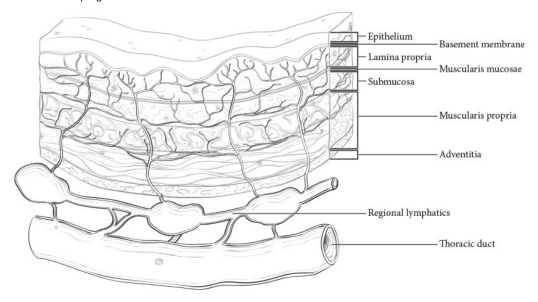
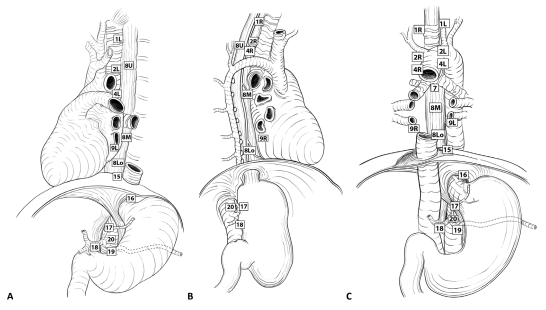


FIGURE 16.2. Esophageal wall.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 16.3. (A–C) Lymph node maps for esophageal cancer. Regional lymph node stations for staging esophageal cancer from left (A), right (B), and anterior (C). 1R, Right lower cervical paratracheal nodes, between the supraclavicular paratracheal space and apex of the lung. 2R, Right upper paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and the apex of the lung. 2L, Left upper paratracheal nodes, between the top of the aortic arch and the apex of the lung. 4R, Right lower paratracheal nodes, between the intersection of the caudal margin of the brachiocephalic artery with the trachea and cephalic border of the azygos vein. 4L, Left lower paratracheal nodes, between the top of the aortic arch and the carina. 7, Subcarinal nodes, caudal to the carina of the trachea. 8U, Upper thoracic paraesophageal lymph nodes, from the apex of the lung to the tracheal bifurcation. 8M, Middle thoracic paraesophageal lymph nodes, from the tracheal bifurcation to the caudal margin of the inferior pulmonary vein. 8Lo, Lower thoracic paraesophageal lymph nodes, from the caudal margin of the inferior pulmonary ligament nodes, within the right inferior pulmonary ligament. 9L, Pulmonary ligament nodes, within the left inferior pulmonary ligament. 15, Diaphragmatic nodes, lying on the dome of the diaphragm and adjacent to or behind its crura. 16, Paracardial nodes, immediately adjacent to the gastroesophageal junction. 17, Left gastric nodes, along the course of the left gastric artery. 18, Common hepatic nodes, immediately on the proximal splenic artery. 20, Celiac nodes, at the base of the celiac artery.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | Tis | Carcinoma in situ: intraepithelial tumor without invasion of the lamina propria, high-grade dysplasia |
| | T1 | Tumor invades the lamina propria, muscularis mucosae, or submucosa |
| | T1a | Tumor invades the lamina propria or muscularis mucosae |
| | T1b | Tumor invades the submucosa |
| | T2 | Tumor invades the muscularis propria* |
| | Т3 | Tumor penetrates the subserosal connective tissue without invasion of the visceral peritoneum or adjacent structures**.** |
| | T4 | Tumor invades the serosa (visceral peritoneum) or adjacent structures **,*** |
| | T4a | Tumor invades the serosa (visceral peritoneum) |
| | T4b | Tumor invades adjacent structures/organs |

^{*} A tumor may penetrate the muscularis propria with extension into the gastrocolic or gastrohepatic ligaments, or into the greater or lesser omentum, without perforation of the visceral peritoneum covering these structures. In this case, the tumor is classified as T3. If there is perforation of the visceral peritoneum covering the gastric ligaments or the omentum, the tumor should be classified as T4.

^{***} Intramural extension to the duodenum or esophagus is not considered invasion of an adjacent structure, but is classified using the depth of the greatest invasion in any of these sites.

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Γ | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph node(s) cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in one or two regional lymph nodes |
| | N2 | Metastasis in three to six regional lymph nodes |
| | N3 | Metastasis in seven or more regional lymph nodes |
| | N3a | Metastasis in seven to 15 regional lymph nodes |
| | N3b | Metastasis in 16 or more regional lymph nodes |

| 1 | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|---|-----------------------|
| | cM0 | No distant metastasis |
| | cM1 Distant metastasis | |
| | pM1 Distant metastasis, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information |
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^{**} The adjacent structures of the stomach include the spleen, transverse colon, liver, diaphragm, pancreas, abdominal wall, adrenal gland, kidney, small intestine, and retroperitoneum.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Clinical (cTNM)

| ✓ | When T is | And N is | And M is | Then the stage group is |
|----------|-----------|---------------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T2 | N0 | M0 | 1 |
| | T1 | N1, N2, or N3 | M0 | IIA |
| | T2 | N1, N2, or N3 | M0 | IIA |
| | T3 | N0 | M0 | IIB |
| | T4a | N0 | M0 | IIB |
| | T3 | N1, N2, or N3 | M0 | III |
| | T4a | N1, N2, or N3 | M0 | III |
| | T4b | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

5.2 Pathological (pTNM)

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | NO | M0 | IA |
| | T1 | N1 | M0 | IB |
| | T2 | NO | M0 | IB |
| | T1 | N2 | M0 | IIA |
| | T2 | N1 | M0 | IIA |
| | T3 | NO NO | M0 | IIA |
| | T1 | N3a | M0 | IIB |
| | T2 | N2 | M0 | IIB |
| | T3 | N1 | M0 | IIB |
| | T4a | NO NO | M0 | IIB |
| | T2 | N3a | M0 | IIIA |
| | T3 | N2 | M0 | IIIA |
| | T4a | N1 | M0 | IIIA |
| | T4a | N2 | M0 | IIIA |
| | T4b | NO NO | M0 | IIIA |
| | T1 | N3b | M0 | IIIB |
| | T2 | N3b | M0 | IIIB |
| | T3 | N3a | M0 | IIIB |
| | T4a | N3a | M0 | IIIB |
| | T4b | N1 | M0 | IIIB |
| | T4b | N2 | M0 | IIIB |
| | T3 | N3b | M0 | IIIC |
| | T4a | N3b | M0 | IIIC |
| | T4b | N3a | M0 | IIIC |
| | T4b | N3b | M0 | IIIC |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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5.3 Postneoadjuvant Therapy (ypTNM)

| ✓ When T is | And N is | And M is | Then the stage group is |
|-------------|----------|----------|-------------------------|
| T1 | NO NO | MO | 1 |
| T2 | NO NO | MO | 1 |
| T1 | N1 | MO | 1 |
| T3 | N0 | M0 | II |
| T2 | N1 | MO | II |
| T1 | N2 | MO | II |
| T4a | N0 | MO | II |
| Т3 | N1 | MO | II |
| T2 | N2 | M0 | II |
| T1 | N3 | MO | II |
| T4a | N1 | M0 | III |
| T3 | N2 | MO | III |
| T2 | N3 | MO | III |
| T4b | N0 | M0 | III |
| T4b | N1 | MO | III |
| T4a | N2 | M0 | III |
| T3 | N3 | MO | III |
| T4b | N2 | M0 | III |
| T4b | N3 | M0 | III |
| T4a | N3 | M0 | III |
| Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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| 6 | | Registry Data Collection Variables | | | |
|-----------------------------|--|---|--|--|--|
| See | See chapter for more details on these variables. | | | | |
| | 1. | Tumor location | on (needed because C16.0 is both cardia and EG | J): | |
| | | _ | | criteria (Use this chapter, AJCC Chapter 17 Stomach) | |
| | | | | | |
| | 2. | EGJ (Use AJCC Chapter 16 Esophagus and Esophagogastric Junction) 2. Serum CEA: | | | |
| | | | | | |
| | 3. | Serum CA 19- | 9: | | |
| | 4. | Clinical stagin | g modalities (endoscopy and biopsy, EUS, EUS- | FNA, CT, PET/CT): | |
| | 5. | Tumor length | : | | |
| | 6. | Depth of inva | sion: | | |
| | 7. | Number of su | spicious malignant lymph nodes on baseline rad | diologic images: | |
| | 8. | Number of su | spicious malignant lymph nodes by EUS assessr | nent: | |
| | 9. | Location of su | ispicious nodes (clinical): | | |
| | 10. | Location of m | alignant nodes (pathological): | | |
| | 11. | Number of tu | mor deposits: | | |
| | 12. | Lymphovascu | lar invasion: | | |
| | 13. | Neural invasion | on: | | |
| | 14. | Extranodal ex | tension: | | |
| | 15. | HER2 status: | positive negative | | |
| | 16. | MSI: | | | |
| | 17. | Surgical marg | in: negative microscopic | macroscopic | |
| | 18. | Sites of metas | stasis, if applicable: | | |
| | 19. | Type of surge | ry: | | |
| 7 | Hi | stologic Gr | ade (G) | | |
| , | | 1 | | | |
| ✓ | G GX | G Defi | nition annot be assessed | | |
| | G1 | | ferentiated | | |
| | G2 | Modera | tely differentiated | | |
| | G3 | Poorly d | lifferentiated, undifferentiated | | |
| 8 | Ly | Lymphovascular Invasion (LVI) | | | |
| | | | | | |
| ✓ | | nponent of Coding | Description | | |
| | 0 | | LVI not present (absent)/not identified | | |
| 1 | | | LVI present/identified, NOS | | |
| 2 | | | Lymphatic and small vessel invasion only (L) | | |
| 3 | | | Venous (large vessel) invasion only (V) | | |
| | 4 | | BOTH lymphatic and small vessel AND venous (large vessel) invasion | | |
| | 9 | 9 Presence of LVI unknown/indeterminate | | | |
| Uar | nital • | Namo / A d d nace | | Patient Name/Information | |
| Hospital Name/Address Patie | | | | i ditent Name, information | |
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9 Anatomy

FIGURE 17.1. Anatomic subsites of the stomach.

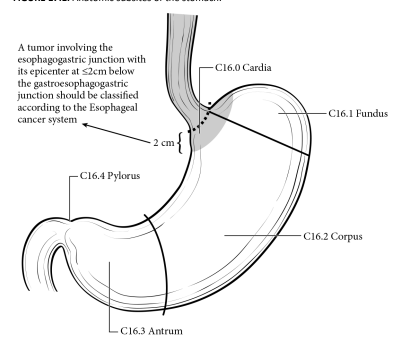
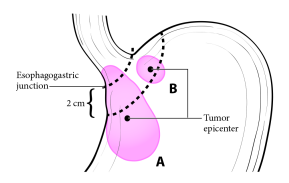
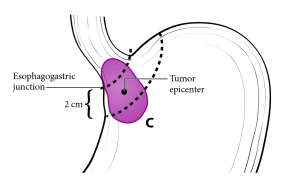


FIGURE 17.2. (A) EGJ tumors with their epicenter located >2 cm into the proximal stomach are staged as stomach cancers. (B) Cardia cancers not involving the EGJ are staged as stomach cancers. (C) Tumors involving the EGJ with thier epicenter <2 cm into the proximal stomach are staged as esophageal cancers.



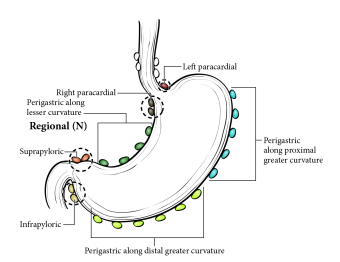
A tumor that has its epicenter located >2 cm from esophagogastric junction (A) or a tumor located within 2 cm of the esophagogastric junction (B) but does not involve the esophagogastric junction is classified as stomach cancer.

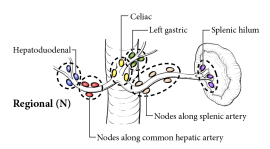


A tumor that has its epicenter located within 2 cm of esophagogastric junction and involves the esophagogatric junction (C) is classified as esophageal cancer.

| Hospital Name/Address | Patient Name/Information |
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FIGURE 17.3. Regional lymph nodes of the stomach.





| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|---|----------------|---|--|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
| | | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy a before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|--|------------|---|
| TX Primary tumor cannot be assessed | | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | High-grade dysplasia/carcinoma in situ |
| | T1 | Tumor invades the lamina propria or submucosa |
| | T1a | Tumor invades the lamina propria |
| T1b Tumor | | Tumor invades the submucosa |
| T2 Tumor invades the muscularis pro | | Tumor invades the muscularis propria |
| T3 Tumor invades through the muscularis prop | | Tumor invades through the muscularis propria into the subserosa, or extends into nonperitonealized |
| | | perimuscular tissue (mesentery or retroperitoneum) without serosal penetration* |
| Tumor perforates the visceral peritoneum or directly invades | | Tumor perforates the visceral peritoneum or directly invades other organs or structures (e.g., other loops of small |
| intestine, mesentery of adjacent loops of bowel, and abdominal wall by way of serosa | | intestine, mesentery of adjacent loops of bowel, and abdominal wall by way of serosa; for duodenum only, |
| invasion of pancreas or bile duct) | | invasion of pancreas or bile duct) |

^{*}Note: For T3 tumors, the nonperitonealized perimuscular tissue is, for the jejunum and ileum, part of the mesentery and, for the duodenum in areas where serosa is lacking, part of the interface with the pancreas.

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ N Category N Criteria | | N Criteria |
|--|----|--|
| NX Regional lymph nodes cannot be assessed | | Regional lymph nodes cannot be assessed |
| NO No regional lymph node metastasis | | No regional lymph node metastasis |
| N1 Metastasis in one or two regional lymph nodes | | Metastasis in one or two regional lymph nodes |
| | N2 | Metastasis in three or more regional lymph nodes |

| | ✓ | N Suffix | Definition | |
|--|---|---|--|--|
| (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|------------|---|--|
| cM0 No distant metastasis | | No distant metastasis | |
| cM1 Distant metastasis | | Distant metastasis | |
| pM1 Distant metastasis, microscopically confirmed | | Distant metastasis, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1-2 | N0 | M0 | 1 |
| | T3 | N0 | M0 | IIA |
| | T4 | N0 | M0 | IIB |
| | Any T | N1 | M0 | IIIA |
| | Any T | N2 | M0 | IIIB |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

- 2. Number of lymph nodes examined:
- 3. Presurgical CEA:
- 4. LVI:
- 5. Microsatellite instability:
- 6. Tumor grade:
- 7. Presence of Crohn's disease:
- 8. Personal or family history of familial GI malignancies (familial adenomatous polyposis, Lynch syndrome, Peutz–Jeghers syndrome):

7 Histologic Grade (G)

| ✓ | G | G Definition |
|------------------------|-----------------------------|---------------------------|
| | GX Grade cannot be assessed | |
| G1 Well differentiated | | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 18.1. Anatomic sites of the small intestine.

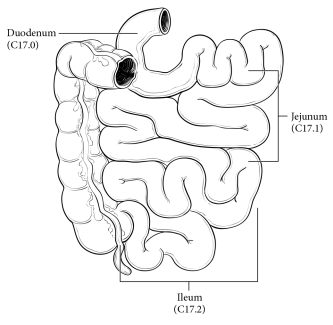
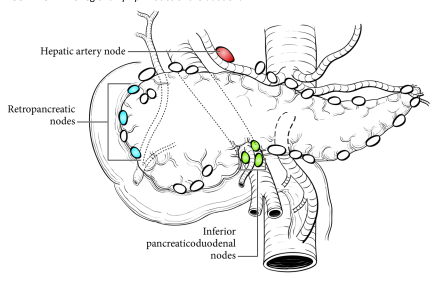


FIGURE 18.2. The regional lymph nodes of the duodenum.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 18.3. The regional lymph nodes of the duodenum.

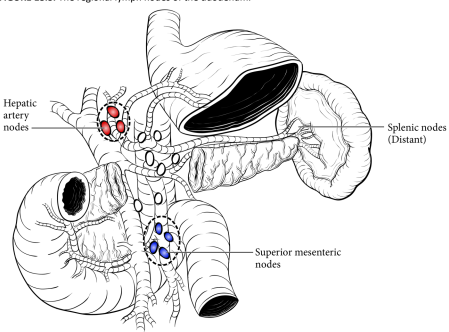
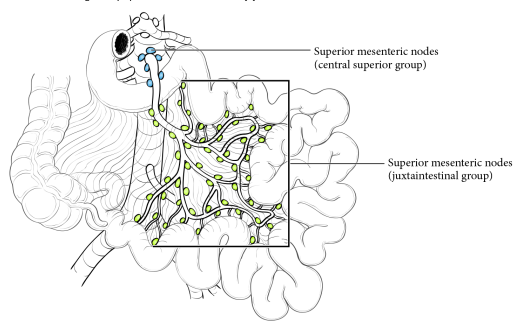


FIGURE 18.4. The regional lymph nodes of the ileum and jejunum.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information |
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| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| Hospital Name/Address | Patient Name/Information |

18.1. Small Intestine: Adenocarcinoma

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | High-grade dysplasia/carcinoma in situ |
| | T1 | Tumor invades the lamina propria or submucosa |
| | T1a | Tumor invades the lamina propria |
| | T1b | Tumor invades the submucosa |
| | T2 | Tumor invades the muscularis propria |
| | T3 | Tumor invades through the muscularis propria into the subserosa, or extends into nonperitonealized perimuscular tissue (mesentery or retroperitoneum) without serosal penetration* |
| | T4 | Tumor perforates the visceral peritoneum or directly invades other organs or structures (e.g., other loops of small intestine, mesentery of adjacent loops of bowel, and abdominal wall by way of serosa; for duodenum only, invasion of pancreas or bile duct) |

^{*}Note: For T3 tumors, the nonperitonealized perimuscular tissue is, for the jejunum and ileum, part of the mesentery and, for the duodenum in areas where serosa is lacking, part of the interface with the pancreas.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in one or two regional lymph nodes |
| | N2 | Metastasis in three or more regional lymph nodes |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| | or printing or the medicagenes (emo) and, or printing see assessment patriological stage grouping. | | |
|---|--|------------|---|
| ſ | ✓ | M Category | M Criteria |
| ſ | | cM0 | No distant metastasis |
| ſ | | cM1 | Distant metastasis |
| ſ | | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

There is no prognostic stage group for non-adenocarcinoma small bowel histologies at this time.

6 Registry Data Collection Variables

- 1. Primary tumor site (duodenum, jejunum, ileum):
- 2. Number of lymph nodes examined:
- 3. Presurgical CEA:
- 4. LVI:
- Microsatellite instability:
- 6. Tumor grade:
- 7. Presence of Crohn's disease:
- 8. Personal or family history of familial GI malignancies (familial adenomatous polyposis, Lynch syndrome, Peutz–Jeghers syndrome):

7 Histologic Grade (G)

| √ | G | G Definition |
|----------|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 18.1. Anatomic sites of the small intestine.

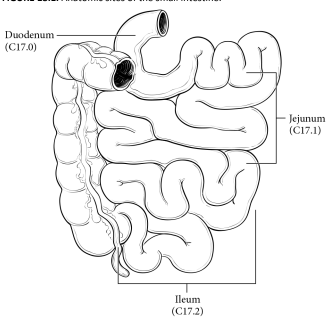
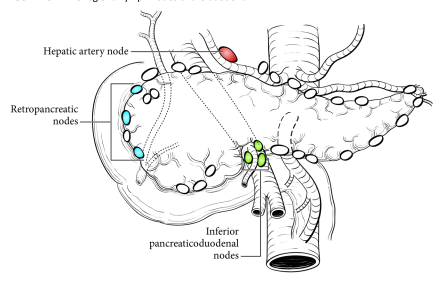


FIGURE 18.2. The regional lymph nodes of the duodenum.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 18.3. The regional lymph nodes of the duodenum.

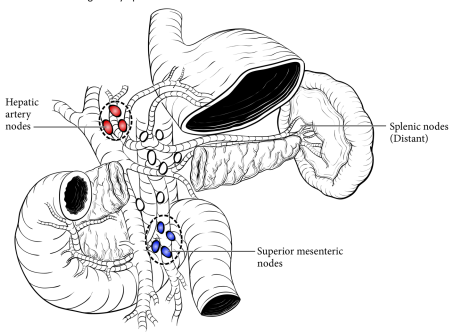
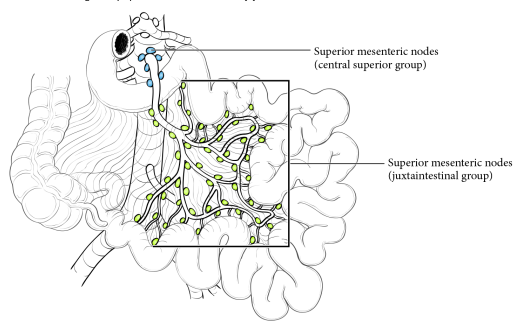


FIGURE 18.4. The regional lymph nodes of the ileum and jejunum.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|--|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma <i>in situ</i> (intramucosal carcinoma; invasion of the lamina propria or extension into but not through the muscularis mucosae) |
| epithelium may invade into the muscularis propria. | | ' ' ' |
| | | T1 and T2 are not applicable to LAMN. Acellular mucin or mucinous epithelium that extends into the subserosa or serosa should be classified as T3 or T4a, respectively. |
| | T1 | Tumor invades the submucosa (through the muscularis mucosa but not into the muscularis propria) |
| | T2 | Tumor invades the muscularis propria |
| | T3 | Tumor invades through the muscularis propria into the subserosa or the mesoappendix |
| | T4 | Tumor invades the visceral peritoneum, including the acellular mucin or mucinous epithelium involving the serosa of the appendix or mesoappendix, and/or directly invades adjacent organs or structures |
| | T4a | Tumor invades through the visceral peritoneum, including the acellular mucin or mucinous epithelium involving the serosa of the appendix or serosa of the mesoappendix |
| | T4b | Tumor directly invades or adheres to adjacent organs or structures |

| ✓ | T Suffix Definition | |
|---|---|--|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|--|---|--|
| | NX Regional lymph nodes cannot be assessed | | |
| | NO No regional lymph node metastasis | | |
| | N1 | One to three regional lymph nodes are positive (tumor in lymph node measuring ≥0.2 mm) or any number of tumor | |
| | | deposits is present, and all identifiable lymph nodes are negative | |
| | N1a | One regional lymph node is positive | |
| | N1b | Two or three regional lymph nodes are positive | |
| | N1c | No regional lymph nodes are positive, but there are tumor deposits in the subserosa or mesentery | |
| | N2 | Four or more regional lymph nodes are positive | |

| ✓ | N Suffix | Definition | |
|---|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|------------|--|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | cM1a | Intraperitoneal acellular mucin, without identifiable tumor cells in the disseminated peritoneal mucinous deposits | |
| | cM1b | Intraperitoneal metastasis only, including peritoneal mucinous deposits containing tumor cells | |
| | cM1c | Metastasis to sites other than peritoneum | |
| | pM1 | Distant metastasis, microscopically confirmed | |
| | pM1a | Microscopically confirmed intraperitoneal acellular mucin, without identifiable tumor cells in the disseminated peritoneal mucinous deposits | |
| | pM1b | Microscopically confirmed intraperitoneal metastasis only, including peritoneal mucinous deposits containing tumor cells | |
| | pM1c | Microscopically confirmed metastasis to sites other than peritoneum | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And G is | Then the stage |
|---|-----------|----------|----------|---------------|----------------|
| • | | | | | group is |
| | Tis | N0 | M0 | Any | 0 |
| | Tis(LAMN) | N0 | M0 | Any | 0 |
| | T1 | N0 | M0 | Any | 1 |
| | T2 | N0 | M0 | Any | I |
| | T3 | N0 | M0 | Any | IIA |
| | T4a | N0 | M0 | Any | IIB |
| | T4b | N0 | M0 | Any | IIC |
| | T1 | N1 | M0 | Any | IIIA |
| | T2 | N1 | M0 | Any | IIIA |
| | T3 | N1 | M0 | Any | IIIB |
| | T4 | N1 | M0 | Any | IIIB |
| | Any T | N2 | M0 | Any | IIIC |
| | Any T | Any N | M1a | Any | IVA |
| | Any T | Any N | M1b | G1 | IVA |
| | Any T | Any N | M1b | G2, G3, or GX | IVB |
| | Any T | Any N | M1c | Any G | IVC |

| Name/Information |
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| 7 | Registry | Data | Collection | Variables |
|---|----------|------|------------|-----------|
|---|----------|------|------------|-----------|

| 1. | Grade: |
|----|--------------------------|
| 2. | CEA levels: |
| 3. | Tumor deposits: |
| 4. | Lymphovascular invasion: |
| 5. | Perineural invasion: |

8 Lymphovascular Invasion (LVI)

| Component of LVI Coding Description | | Description |
|--------------------------------------|---|--|
| | | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 19.1. Anatomic location of the appendix

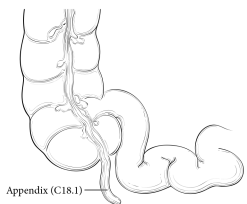
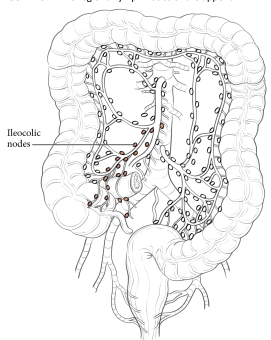


FIGURE 19.2. The regional lymph nodes of the appendix.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ, intramucosal carcinoma (involvement of lamina propria with no extension through muscularis mucosae) |
| | T1 | Tumor invades the submucosa (through the muscularis mucosa but not into the muscularis propria) |
| | T2 | Tumor invades the muscularis propria |
| | T3 | Tumor invades through the muscularis propria into pericolorectal tissues |
| | T4 | Tumor invades* the visceral peritoneum or invades or adheres** to adjacent organ or structure |
| | T4a | Tumor invades* through the visceral peritoneum (including gross perforation of the bowel through tumor and |
| | | continuous invasion of tumor through areas of inflammation to the surface of the visceral peritoneum) |
| | T4b | Tumor directly invades* or adheres** to adjacent organs or structures |

^{*}Direct invasion in T4 includes invasion of other organs or other segments of the colorectum as a result of direct extension through the serosa, as confirmed on microscopic examination (for example, invasion of the sigmoid colon by a carcinoma of the cecum) or, for cancers in a retroperitoneal or subperitoneal location, direct invasion of other organs or structures by virtue of extension beyond the muscularis propria (i.e., respectively, a tumor on the posterior wall of the descending colon invading the left kidney or lateral abdominal wall; or a mid or distal rectal cancer with invasion of prostate, seminal vesicles, cervix, or vagina).

^{**}Tumor that is adherent to other organs or structures, grossly, is classified cT4b. However, if no tumor is present in the adhesion, microscopically, the classification should be pT1-4a depending on the anatomical depth of wall invasion. The V and L classification should be used to identify the presence or absence of vascular or lymphatic invasion whereas the PN prognostic factor should be used for perineural invasion.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | One to three regional lymph nodes are positive (tumor in lymph nodes measuring ≥ 0.2 mm), or any number of tumor deposits are present and all identifiable lymph nodes are negative | |
| | N1a | One regional lymph node is positive | |
| | N1b | Two or three regional lymph nodes are positive | |
| | N1c No regional lymph nodes are positive, but there are tumor deposits in the subserosa | | |
| | mesentery or nonperitonealized pericolic, or perirectal/mesorectal tissues. | | |
| | N2 Four or more regional nodes are positive | | |
| | N2a Four to six regional lymph nodes are positive | | |
| | N2b | Seven or more regional lymph nodes are positive | |

| 1 | N Suffix | fix Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|--|
| | cM0 | No distant metastasis by imaging, etc.; no evidence of tumor in distant sites or organs (This category is not assigned by pathologists.) |
| | cM1 | Metastasis to one or more distant sites or organs or peritoneal metastasis is identified |
| | cM1a | Metastasis to one site or organ is identified without peritoneal metastasis |
| | cM1b | Metastasis to two or more sites or organs is identified without peritoneal metastasis |
| | cM1c | Metastasis to the peritoneal surface is identified alone or with other site or organ metastases |
| | pM1 | Metastasis to one or more distant sites or organs or peritoneal metastasis is identified and microscopically confirmed |
| | pM1a | Metastasis to one site or organ is identified without peritoneal metastasis and microscopically confirmed |
| | pM1b | Metastasis to two or more sites or organs is identified without peritoneal metastasis and microscopically confirmed |
| | pM1c | Metastasis to the peritoneal surface is identified alone or with other site or organ metastases and microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1, T2 | N0 | M0 | 1 |
| | T3 | N0 | MO | IIA |
| | T4a | N0 | M0 | IIB |
| | T4b | N0 | MO | IIC |
| | T1-T2 | N1/N1c | M0 | IIIA |
| | T1 | N2a | M0 | IIIA |
| | T3-T4a | N1/N1c | MO | IIIB |
| | T2-T3 | N2a | M0 | IIIB |
| | T1-T2 | N2b | M0 | IIIB |
| | T4a | N2a | M0 | IIIC |
| | T3-T4a | N2b | MO | IIIC |
| | T4b | N1-N2 | M0 | IIIC |
| | Any T | Any N | M1a | IVA |
| | Any T | Any N | M1b | IVB |
| | Any T | Any N | M1c | IVC |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Tumor deposits:
- 2. CEA levels: preoperative blood level recorded in nanograms per milliliter with fixed decimal point and five numbers (XXXX.X ng/mL):
- 3. Tumor regression score (0-3):
- 4. Circumferential resection margin (in mm):
- 5. Lymphovascular invasion:
- 6. Perineural invasion:
- 7. Microsatellite instability:
- 8. KRAS and NRAS mutation:
- 9. BRAF mutation:

7 Histologic Grade (G)

| ✓ | G | G Definition |
|-----------------------------|----|---------------------------|
| GX Grade cannot be assessed | | Grade cannot be assessed |
| G1 Well differentiated | | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 20.1. Anatomic subsites of the colon.

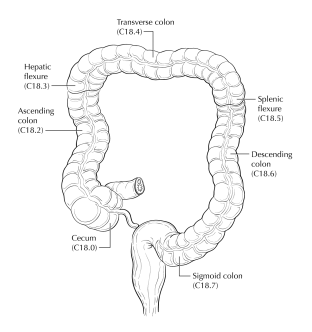


FIGURE 20.2. Anatomic subsites of the rectum.

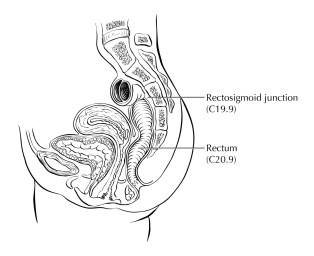
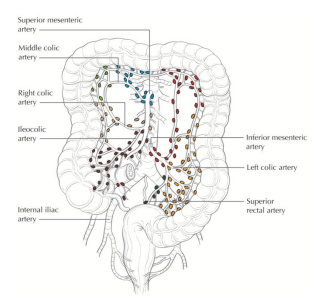


FIGURE 20.4. The regional lymph nodes of the colon and rectum.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor not assessed |
| | TO | No evidence of primary tumor |
| | Tis | High-grade squamous intraepithelial lesion (previously termed carcinoma in situ, Bowen disease, anal intraepithelial neoplasia II–III, high-grade anal intraepithelial neoplasia) |
| | T1 | Tumor ≤2 cm |
| | T2 | Tumor >2 cm but ≤5 cm |
| | T3 | Tumor >5 cm |
| | T4 | Tumor of any size invading adjacent organ(s), such as the vagina, urethra, or bladder |

| ٧ | <u> </u> | T Suffix | Definition |
|---|----------|----------|---|
| | (| (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|----------|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in inguinal, mesorectal, internal iliac, or external iliac nodes |
| | N1a | Metastasis in inguinal, mesorectal, or internal iliac lymph nodes |
| | N1b | Metastasis in external iliac lymph nodes |
| | N1c | Metastasis in external iliac with any N1a nodes |

| | ✓ | N Suffix | Definition |
|---|---|----------|--|
| | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| Г | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | IIIA |
| | T2 | NO | M0 | IIA |
| | T2 | N1 | M0 | IIIA |
| | T3 | NO | M0 | IIB |
| | T3 | N1 | M0 | IIIC |
| | T4 | N0 | M0 | IIIB |
| | T4 | N1 | M0 | IIIC |
| | Any T | Any N | M1 | IV |

| _ | | | . | |
|---|----------|------|------------|-----------|
| 6 | Registry | Data | Collection | Variables |

| 1. | Tumor location: | 🗌 anal 🔲 periai | nal perineal |
|----|-----------------|-----------------|----------------------------|
| | AND | ☐ left ☐ right | anterior posterior lateral |
| 2. | HIV status: | | |
| 3. | Gender: | | |
| 4. | Grade: | | |
| 5. | HPV status: | p16 expression | p18 expression |
| | | | |

7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|--|----------------------------|--|
| | GX | Grade cannot be determined | |
| | G1 Well differentiated (low grade) | | |
| | G2 Moderately differentiated (low grade) | | |
| | G3 Poorly differentiated (high grade) | | |
| | G4 Undifferentiated (high grade) | | |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 21.1A-B. Anal cancer (A–C), perianal cancer (D), and skin cancer (E) as visualized with gentle traction placed on the buttocks.

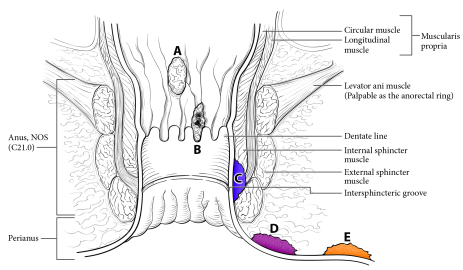
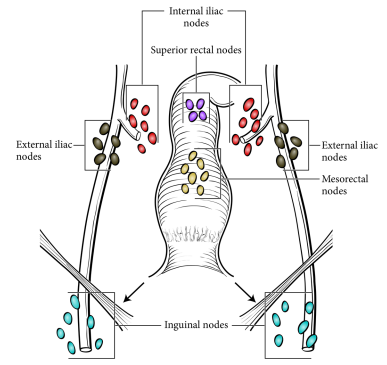


FIGURE 21.3. Regional lymph nodes of the anal canal.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | T1 | Solitary tumor ≤2 cm, or >2 cm without vascular invasion |
| | T1a | Solitary tumor ≤2 cm |
| | T1b | Solitary tumor >2 cm without vascular invasion |
| | T2 | Solitary tumor >2 cm with vascular invasion, or multiple tumors, none >5 cm |
| | T3 | Multiple tumors, at least one of which is >5 cm |
| | T4 | Single tumor or multiple tumors of any size involving a major branch of the portal vein or hepatic vein, |
| | | or tumor(s) with direct invasion of adjacent organs other than the gallbladder or with perforation of visceral |
| | | peritoneum |

| | ✓ | T Suffix Definition | |
|---|---|---|--|
| ſ | | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| 1 | N Category | N Criteria | |
|---|--|------------|--|
| | NX Regional lymph nodes cannot be assessed | | |
| | NO No regional lymph node metastasis | | |
| | N1 Regional lymph node metastasis | | |

| | ✓ | N Suffix | Definition | |
|---|--|----------|------------|--|
| Ī | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| Γ | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | gory M Criteria | |
|---|---|-----------------|--|
| | cM0 No distant metastasis | | |
| | cM1 Distant metastasis | | |
| | pM1 Distant metastasis, microscopically confirmed | | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | IIIA |
| | T4 | N0 | M0 | IIIB |
| | Any T | N1 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

| Hospital Name/Address | Patient Name/Information | |
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6 Registry Data Collection Variables

| 1. | AFP: | |
|----|---|----------------------|
| 2. | Fibrosis score: | Scoring system used: |
| 3. | Hepatitis serology: | |
| 4. | . Creatinine (part of the MELD score): | |
| 5. | Bilirubin (part of the MELD score): | |
| 6. | Prothrombin time (INR; part of the MELD score): | |

7 Histologic Grade (G)

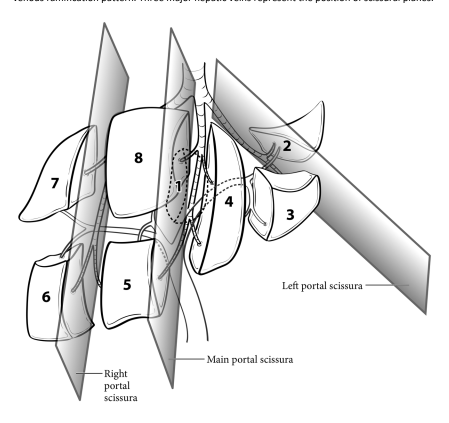
| 1 | G | G Definition |
|-----------------------------|------------------------|---------------------------|
| GX Grade cannot be assessed | | Grade cannot be assessed |
| | G1 Well differentiated | |
| | G2 | Moderately differentiated |
| G3 Poorly differentiated | | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description | |
|---|--|--|--|
| • | LVI Coding | | |
| | 0 LVI not present (absent)/not identified | | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 Venous (large vessel) invasion only (V) | | |
| | 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion | | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 22.1. Couinaud's segmental anatomy of the liver. The liver is divided into two hemilivers and eight segments according to the portal venous ramification pattern. Three major hepatic veins represent the position of scissural planes.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ (intraductal tumor) |
| | T1 | Solitary tumor without vascular invasion, ≤5 cm or >5 cm |
| | T1a | Solitary tumor ≤5 cm without vascular invasion |
| | T1b | Solitary tumor >5 cm without vascular invasion |
| | T2 | Solitary tumor with intrahepatic vascular invasion or multiple tumors, with or without vascular invasion |
| | T3 | Tumor perforating the visceral peritoneum |
| | T4 | Tumor involving local extrahepatic structures by direct invasion |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis present |

| | ✓ | N Suffix | Definition | |
|---|--|----------|------------|--|
| Ī | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| Π | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | IIIA |
| | T4 | N0 | M0 | IIIB |
| | Any T | N1 | M0 | IIIB |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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| 6 Registry Data Collection Var | riables |
|--------------------------------|---------|
|--------------------------------|---------|

See chapter for more details on these variables.

- 1. Presence of nontumoral hepatic parenchymal fibrosis/cirrhosis:
- 2. Primary sclerosing cholangitis:
- 3. Serum CA 19-9 level:
- 4. Tumor growth pattern:

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 23.1. Liver diagram differentiating intrahepatic bile ducts from extrahepatic bile ducts and mass-forming growth pattern (A) from periductal infiltrating growth pattern (B), with associated intrahepatic biliary dilatation.

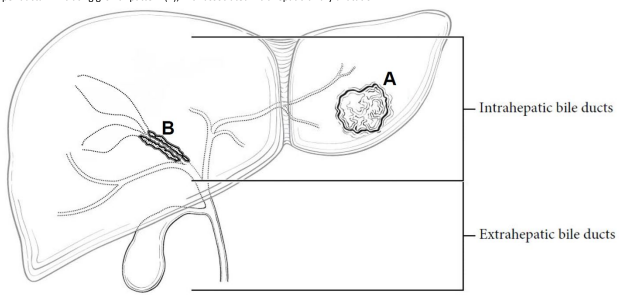
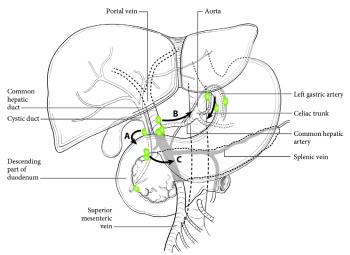


FIGURE 23.2. Differential lymphatic drainage patterns for left and right liver intrahepatic cholangiocarcinomas. Right liver tumors drain to right portal (A) and then portocaval (C) nodal basins, while left liver tumors drain to left gastric and celiac (B) nodal basins.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
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| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ |
| | T1 | Tumor invades the lamina propria or muscular layer |
| | T1a | Tumor invades the lamina propria |
| | T1b | Tumor invades the muscular layer |
| | T2 | Tumor invades the perimuscular connective tissue on the peritoneal side, without involvement of the serosa (visceral peritoneum) |
| | T2- | Or tumor invades the perimuscular connective tissue on the hepatic side, with no extension into the liver |
| | T2a | Tumor invades the perimuscular connective tissue on the peritoneal side, without involvement of the serosa (visceral peritoneum) |
| | T2b | Tumor invades the perimuscular connective tissue on the hepatic side, with no extension into the liver |
| | T3 | Tumor perforates the serosa (visceral peritoneum) and/or directly invades the liver and/or one other adjacent organ or structure, such as the stomach, duodenum, colon, pancreas, omentum, or extrahepatic bile ducts |
| | T4 | Tumor invades the main portal vein or hepatic artery or invades two or more extrahepatic organs or structures |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|----------|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastases to one to three regional lymph nodes |
| | N2 | Metastases to four or more regional lymph nodes |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
| | |

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T2a | NO | M0 | IIA |
| | T2b | N0 | M0 | IIB |
| | T3 | N0 | M0 | IIIA |
| | T1-3 | N1 | M0 | IIIB |
| | T4 | N0-1 | M0 | IVA |
| | Any T | N2 | M0 | IVB |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Specimen type:
- 2. Extent of liver resection:
- 3. Free peritoneal side versus hepatic side for T2 tumors:

7 Histologic Grade (G)

| √ | G | G Definition |
|----------|-----------------------------|---------------------------|
| | GX Grade cannot be assessed | |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

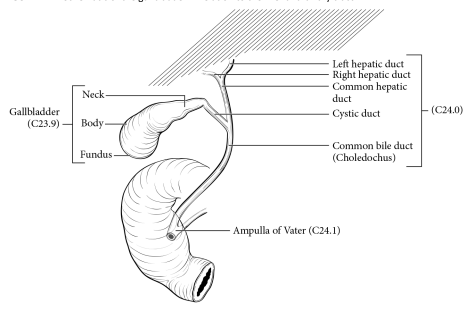
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 24.1. Schematic of the gallbladder in relation to the liver and biliary tract.



| Physician Signature | Date/Time |
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| Patient Name/Information | |
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| | Patient Name/Information |

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | Tis | Carcinoma in situ/high-grade dysplasia |
| | T1 | Tumor confined to the bile duct, with extension up to the muscle layer or fibrous tissue |
| | T2 | Tumor invades beyond the wall of the bile duct to surrounding adipose tissue, |
| | | or tumor invades adjacent hepatic parenchyma |
| | T2a | Tumor invades beyond the wall of the bile duct to surrounding adipose tissue |
| | T2b | Tumor invades adjacent hepatic parenchyma |
| | T3 | Tumor invades unilateral branches of the portal vein or hepatic artery |
| | T4 | Tumor invades the main portal vein or its branches bilaterally, or the common hepatic artery; or unilateral second-order biliary radicals with contralateral portal vein or hepatic artery involvement |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | One to three positive lymph nodes typically involving the hilar, cystic duct, common bile duct, hepatic artery, | |
| | | posterior pancreatoduodenal, and portal vein lymph nodes | |
| | N2 | Four or more positive lymph nodes from the sites described for N1 | |

| ✓ | N Suffix | Definition |
|----------|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | 0 |
| | T1 | NO | M0 | 1 |
| | T2a-b | N0 | M0 | II |
| | T3 | NO | M0 | IIIA |
| | T4 | N0 | M0 | IIIB |
| | Any T | N1 | M0 | IIIC |
| | Any T | N2 | M0 | IVA |
| | Any T | Any N | M1 | IVB |

| Hospital Name/Address | Patient Name/Information |
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| 6 Registry Data Collection Variable | 6 | Registry | Data | Collection | Variable |
|-------------------------------------|---|----------|------|------------|----------|
|-------------------------------------|---|----------|------|------------|----------|

See chapter for more details on these variables.

- 1. Tumor location and extent according to Bismuth–Corlette classification:
- 2. Papillary histology:
- 3. Primary sclerosing cholangitis:

7 Histologic Grade (G)

| 1 | G | G Definition | |
|---|------------------------------|--------------------------|--|
| | GX | Grade cannot be assessed | |
| | G1 | Well differentiated | |
| | G2 Moderately differentiated | | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
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| | |

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | Tis | Carcinoma in situ/high-grade dysplasia |
| | T1 | Tumor invades the bile duct wall with a depth less than 5 mm |
| | T2 | Tumor invades the bile duct wall with a depth of 5–12 mm |
| | T3 | Tumor invades the bile duct wall with a depth greater than 12 mm |
| | T4 | Tumor involves the celiac axis, superior mesenteric artery, and/or common hepatic artery |

| ✓ | T Suffix Definition | |
|---|---|--|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in one to three regional lymph nodes |
| | N2 | Metastasis in four or more regional lymph nodes |

| ✓ | N Suffix | offix Definition | |
|--|----------|--|--|
| (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Category M Criteria | |
|----------|------------|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | pM1 | Distant metastasis, microscopically confirmed | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | IIA |
| | T1 | N2 | M0 | IIIA |
| | T2 | N0 | M0 | IIA |
| | T2 | N1 | M0 | IIB |
| | T2 | N2 | M0 | IIIA |
| | T3 | N0 | M0 | IIB |
| | Т3 | N1 | M0 | IIB |
| | Т3 | N2 | M0 | IIIA |
| | T4 | N0 | M0 | IIIB |
| | T4 | N1 | M0 | IIIB |
| | T4 | N2 | M0 | IIIB |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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This form continues on the next page.

6 **Registry Data Collection Variables**

| See | chap | ter for more details on these variables. |
|-----|------|---|
| | 1. | Tumor location (ICD-O-3 code lacks specificity): |
| | | cystic duct (Use AJCC Chapter 24 Gallbladder) |
| | | perihilar bile ducts (Use AJCC Chapter 25 Perihilar Bile Ducts) |
| | | distal bile duct (use this chapter, AJCC Chapter 26 Distal Bile Duct) |
| | 2. | CEA: |
| | 3. | CA 19-9: |
| 7 | ш: | istologic Grado (G) |

Histologic Grade (G)

| ✓ | G | G Definition | |
|------------------------------|-----------------------------|---------------------------|--|
| | GX Grade cannot be assessed | | |
| | G1 | Well differentiated | |
| G2 Moderately differentiated | | Moderately differentiated | |
| | G3 | Poorly differentiated | |

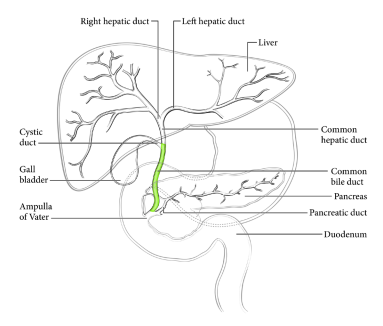
Lymphovascular Invasion (LVI)

| 1 | Component of | Description | |
|---|--|--|--|
| * | LVI Coding | | |
| | 0 LVI not present (absent)/not identified | | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 Venous (large vessel) invasion only (V) | | |
| | 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion | | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 26.1. Diagram highlighting the location of tumors to be staged as distal bile duct tumors. These tumors have an epicenter located between the confluence of the cystic duct and common hepatic duct and the ampulla of Vater (highlighted) (Modified from the College of American Pathologists).



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|--|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ |
| | T1 | Tumor limited to ampulla of Vater or sphincter of Oddi |
| | | or tumor invades beyond the sphincter of Oddi (perisphincteric invasion) and/or into the duodenal submucosa |
| T1a Tumor limited to ampulla of Vater or sphincter of Oddi | | Tumor limited to ampulla of Vater or sphincter of Oddi |
| | T1b | Tumor invades beyond the sphincter of Oddi (perisphincteric invasion) and/or into the duodenal submucosa |
| | T2 Tumor invades into the muscularis propria of the duodenum | |
| T3 Tumor directly invades the pancreas (up to 0.5 cm) | | Tumor directly invades the pancreas (up to 0.5 cm) |
| or tumor extends more than 0.5 cm into the pancreas, or extends into peripancreatic or periduod | | or tumor extends more than 0.5 cm into the pancreas, or extends into peripancreatic or periduodenal tissue or |
| | | duodenal serosa without involvement of the celiac axis or superior mesenteric artery |
| | T3a | Tumor directly invades pancreas (up to 0.5 cm) |
| | T3b | Tumor extends more than 0.5 cm into the pancreas, or extends into peripancreatic tissue or periduodenal tissue |
| | | or duodenal serosa without involvement of the celiac axis or superior mesenteric artery |
| | T4 | Tumor involves the celiac axis, superior mesenteric artery, and/or common hepatic artery, irrespective of size |

| | ✓ | T Suffix | Definition |
|---|---|---|------------|
| Ī | | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | Criteria | | | |
|---|--|---|--|--|--|
| | NX Regional lymph nodes cannot be assessed | | | | |
| | N0 | No regional lymph node metastasis | | | |
| | N1 Metastasis to one to three regional lymph nodes | | | | |
| | N2 | Metastasis to four or more regional lymph nodes | | | |

| √ | N Suffix | Definition | | |
|----------|--|---|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria | |
|---|------------|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | pM1 | Distant metastasis, microscopically confirmed | |

| Patient Name/Information | | |
|--------------------------|--|--|
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1a | NO | M0 | IA |
| | T1a | N1 | M0 | IIIA |
| | T1b | N0 | M0 | IB |
| | T1b | N1 | M0 | IIIA |
| | T2 | NO | M0 | IB |
| | T2 | N1 | M0 | IIIA |
| | T3a | N0 | M0 | IIA |
| | T3a | N1 | M0 | IIIA |
| | T3b | N0 | M0 | IIB |
| | T3b | N1 | M0 | IIIA |
| | T4 | Any N | M0 | IIIB |
| | Any T | N2 | M0 | IIIB |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

| See | chapter | for | more | details | on | these | variables. |
|-----|---------|-----|------|---------|----|-------|------------|
| | | | | | | | |

- 1. Tumor size:
- 2. Lymph node status:
- 3. Margin status:
- 4. Histologic differentiation:
- 5. Histologic subtype:
- 6. Preoperative or pretreatment CEA:
- 7. Preoperative or pretreatment CA 19-9:
- 8. Adjuvant therapy:

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

| Hospital Name/Address | Patient Name/Information |
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8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description | |
|---|---|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 LVI present/identified, NOS | | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 Venous (large vessel) invasion only (V) | | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|---|---|--|--|--|
| | cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagn workup information, until first treatment, including clinical history and symptoms, physical examination, ir endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sam regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and relevant examinations | | | |
| | | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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| | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|--|--|--|
| | TX | Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | Tis | Carcinoma in situ. This includes high-grade pancreatic intraepithelial neoplasia (Panln-3), intraductal papillary mucinous neoplasm with high-grade dysplasia, intraductal tubulopapillary neoplasm with high-grade dysplasia, and mucinous cystic neoplasm with high-grade dysplasia. | |
| T1 Tumor ≤2 cm in greatest dimension | | Tumor ≤2 cm in greatest dimension | |
| T1a Tumor ≤0.5 cm in greatest dimension | | Tumor ≤0.5 cm in greatest dimension | |
| T1b Tumor >0.5 cm and <1 cm in greatest dimension | | Tumor >0.5 cm and <1 cm in greatest dimension | |
| | T1c | Tumor 1–2 cm in greatest dimension | |
| | T2 Tumor >2 cm and ≤4 cm in greatest dimension | | |
| | T3 | Tumor >4 cm in greatest dimension | |
| | T4 | Tumor involves celiac axis, superior mesenteric artery, and/or common hepatic artery, regardless of size | |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| (m) Select if synchronous primary tumors are found in single organ. | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastases |
| | N1 | Metastasis in one to three regional lymph nodes |
| | N2 | Metastasis in four or more regional lymph nodes |

| ✓ | N Suffix | Definition | |
|--|----------|--|--|
| (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria | |
|----------|---|------------|--|
| | cMO No distant metastasis | | |
| | cM1 Distant metastasis | | |
| | pM1 Distant metastasis, microscopically confirmed | | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | IA |
| | T1 | N1 | M0 | IIB |
| | T1 | N2 | M0 | III |
| | T2 | N0 | M0 | IB |
| | T2 | N1 | M0 | IIB |
| | T2 | N2 | M0 | III |
| | T3 | NO | M0 | IIA |
| | T3 | N1 | M0 | IIB |
| | T3 | N2 | M0 | III |
| | T4 | Any N | M0 | III |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Preoperative CA 19-9:
- 2. Preoperative carcinoembryonic antigen (CEA):

7 Histologic Grade (G)

| 1 | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

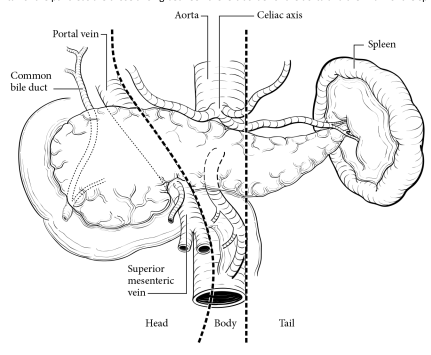
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 28.1. Tumors of the head of the pancreas are those arising to the right of the superior mesenteric-portal vein confluence. Tumors of the body of the pancreas are those arising between the left border of the superior mesenteric vein and the left border of the aorta. Tumors of the tail of the pancreas are those arising between the left border of the aorta and the hilum of the spleen.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1* | Invades the lamina propria or submucosa and less than or equal to 1 cm in size |
| | T2* | Invades the muscularis propria or greater than 1 cm in size |
| | T3* | Invades through the muscularis propria into subserosal tissue without penetration of overlying serosa |
| | T4* | Invades visceral peritoneum (serosa) or other organs or adjacent structures |

^{*}Note: For any T, add (m) for multiple tumors [TX(#) or TX(m), where X = 1–4 and # = number of primary tumors identified**]; for multiple tumors with different Ts, use the highest.

^{**}Example: If there are two primary tumors, one of which penetrates only the subserosa, we define the primary tumor as either T3(2) or T3(m).

| 1 | T Suffix | Definition |
|---|----------|---|
| (m) Select if synchronous primary tumors are found in single organ. | | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| 1 | N Suffix | Definition | |
|---|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Metastasis confined to liver |
| | cM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) |
| | cM1c | Both hepatic and extrahepatic metastases |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Metastasis confined to liver, microscopically confirmed |
| | pM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone), |
| | | microscopically confirmed |
| | pM1c | Both hepatic and extrahepatic metastases, microscopically confirmed |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When Tis | And N is | And M is | Then the stage group is |
|---|----------|------------|----------|-------------------------|
| | TX, T0 | NX, N0, N1 | M1 | IV |
| | T1 | NO | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T1 | NX, N0, N1 | M1 | IV |
| | T2 | NO | M0 | 11 |
| | T2 | N1 | M0 | III |
| | T2 | NX, N0, N1 | M1 | IV |
| | T3 | NO | M0 | II |
| | T3 | N1 | M0 | III |
| | T3 | NX, N0, N1 | M1 | IV |
| | T4 | NO | M0 | III |
| | T4 | N1 | M0 | III |
| | T4 | NX, N0, N1 | M1 | IV |

6 Registry Data Collection Variables

| chapt | er for more details on these variables. |
|-------|--|
| 1. | Size of tumor (value or unknown): |
| 2. | Depth of invasion: |
| 3. | Nodal status and number of nodes involved, if applicable: |
| 4. | Sites of metastasis, if applicable: |
| 5. | Ki-67 index: |
| 6. | Mitotic count: |
| 7. | Histologic grading (from Ki-67 and mitotic count): GX G1 G2 G3 |
| 8. | Preoperative pancreastatin level: |
| 9. | Preoperative gastrin level: |
| 10. | Preoperative CgA level: |
| 11. | Type of gastric NET: |

7 Histologic Grade (G)

| ✓ | G | G Definition |
|-----|------------|---|
| | GX | Grade cannot be assessed |
| | G1 | Mitotic count (per 10 HPF)* < 2 and Ki-67 index (%)** < 3 |
| | G2 | Mitotic count (per 10 HPF) = 2–20 or Ki-67 index (%)** = 3–20 |
| | G3 | Mitotic count (per 10 HPF) > 20 or Ki-67 index (%)** > 20 |
| *10 | *40.1105 2 | |

^{*10} HPF = 2 mm²; at least 50 HPF (at 40× magnification) must be evaluated in areas of highest mitotic density in order to adhere to WHO 2010 criteria.

In cases of disparity between Ki-67 proliferative index and mitotic count, the result that indicates a higher-grade tumor should be selected as the final grade. For example, a mitotic count of 1 per 10 HPF and a Ki-67 of 12% should be designated as a G2 NET.

| Hospital Name/Address | Patient Name/Information |
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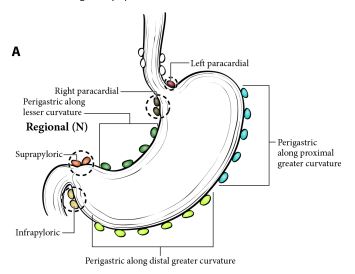
^{**}MIB1 antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling.

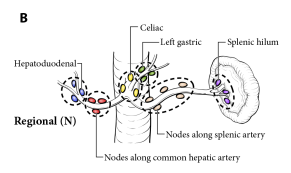
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 29.1. The regional lymph nodes of the stomach for neuroendocrine tumors.





| Physician Signature | Date/Time |
|---------------------|-----------|

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | yp TNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria | |
|---|--|---|--|
| | TX | Primary tumor cannot be assessed | |
| | T1 | T1 Tumor invades the mucosa or submucosa only and is ≤1 cm (duodenal tumors); | |
| | | Tumor ≤1 cm and confined within the sphincter of Oddi (ampullary tumors) | |
| | T2 | Tumor invades the muscularis propria or is >1 cm (duodenal); | |
| | Tumor invades through sphincter into duodenal submucosa or muscularis propria, or is >1 cm (ampullary) | | |
| | T3 | Tumor invades the pancreas or peripancreatic adipose tissue | |
| | T4 | Tumor invades the visceral peritoneum (serosa) or other organs | |
| | | | |

Note: Multiple tumors should be designated as such (and the largest tumor should be used to assign the T category):

- If the number of tumors is known, use T(#); e.g., pT3(4)N0M0.
- If the number of tumors is unavailable or too numerous, use the suffix m—T(m)—e.g., pT3(m)N0M0.

| ~ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | V Criteria | |
|---|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | o regional lymph node involvement | |
| | N1 | Regional lymph node involvement | |

| ✓ | N Suffix | efinition | |
|---|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria | |
|---|---|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastases | |
| | cM1a | Metastasis confined to liver | |
| | cM1b Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) | | |
| | cM1c | Both hepatic and extrahepatic metastases | |
| | pM1 Distant metastases, microscopically confirmed | | |
| | pM1a Metastasis confined to liver, microscopically confirmed | | |
| | pM1b Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) | | |
| | microscopically confirmed | | |
| | pM1c | Both hepatic and extrahepatic metastases, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | II |
| | T4 | N0 | M0 | III |
| | Any T | N1 | M0 | III |
| | Any T | Any N | M1 | IV |

| 5 | Re | legistry Data Collection Variables | | | | |
|------|--|---|---------------------------|--|--|--|
| See | chapt | apter for more details on these variables. | | | | |
| | 1. | Size of tumor (value): | | | | |
| | 2. | Maximum depth of invasion (microscopic tumor extension): | | | | |
| | | Small intestine (including duodenum): | uctures | | | |
| | 3. | Number of tumors (multicentric disease at primary site): | | | | |
| | 4. | Lymph node status (including number of nodes assessed and n | umber of positive nodes): | | | |
| | 5. | Grade (based on Ki-67 and mitotic count: GX (unknown) | ☐ G1 ☐ G2 ☐ G3 | | | |
| | 6. | Mitotic count (value): | | | | |
| | 7. | Ki-67 Labeling Index (value): | | | | |
| | 8. | Perineural invasion: Yes No | | | | |
| | 9. | Lymphovascular invasion: Yes No | | | | |
| | 10. Margin status: Positive (+) Negative (-) | | | | | |
| This | form | continues on the next page. | | | | |
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| 30. | Ne | uroendocrine | : Tumor | s of the | e Duode | enum and Ampulla | a of Vater | |
|-------|---|--|----------------|---------------|----------------|---|-------------------|-------------------------------|
| | 11. | Functional status: | ∏Yes | □No | If ves the | n select type of syndrome: | | |
| | | Tarrettorial Status. | | | | | | |
| | | | | | Function | _ | | |
| | | | | | | Gastrininoma (ZES) | | |
| | | | | | | Somatostatinoma | | |
| | | | | | | NET causing carcinoid | syndrome (5HIA | A, serotonin excess) |
| | | | | | | Other: | | |
| | | | | | Nonfu | nctional | | |
| | | | | | Unkno | wn/unable to assess | | |
| | 12. | Genetic syndrome: | Yes | ☐ No | If yes, typ | e of syndrome: | | |
| | | | | | ☐ MEN1 | | | |
| | | | | | = | | | |
| | | | | | = | ppel-Lindau disease | | |
| | | | | | NF1 | | | |
| | | | | | U Other | syndrome, NOS | | |
| | 13. | Location in duodent | ım: | first p | ortion | second portion t | third portion | fourth portion |
| | | | | ampu | lla of Vater | | | |
| | 14. | Type of surgery: | ☐ EMR | | | | | |
| | | Type or surgery. | _ | | | | | |
| | | | Pancre | eaticoduod | enectomy: | ☐ partial ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ | complete | with partial gastrectomy |
| | | | □ Whin | ole procedu | ıre | without partial gastret | ctomy | |
| | | | | llectomy | | | | |
| | | | | - | المسم مما | startina | | |
| | | | = - | | ion, small ir | itestine | | |
| | | | Unkno | | | | | |
| | | | Other | | | | | |
| | 15. | Preoperative CgA le | | | • | LU NIV. | | |
| | 16. | Preoperative pancre | :astatin ieve | ei (absolute | value with | OLN): | | |
| | 17. | Preoperative neurol | kinin level (a | absolute va | lue with UL | N): | | |
| | 18. | Age of patient: | | | | | | |
| | 19. | Histologic variants: | ☐ Well-d | differentiate | ed NET | Glandular duodenal NE | ET (somatostatin | oma) |
| | | | Gangli | ocytic parag | ganglioma | | | |
| | | | | | | | | |
| 7 | His | stologic Grade | (G) | | | | | |
| | | C D-ft : tt | | | | | | |
| • | G CY | G Definition Grade cannot | | 1 | | | | |
| | GX G1 | Mitotic count | | | i 67 inday / | 2/** ~2 | | |
| | G2 | | | | | ex (%)** = 3–20 | | |
| | G2 G3 | | •• | | | | | |
| *10 | | G3 Mitotic count (per 10 HPF) >20 and Ki-67 index (%)** >20 IPF = 2 mm ² ; at least 50 HPF (at 40× magnification) must be evaluated in areas of highest mitotic density in order to match WHO 2010 | | | | | | |
| crite | | Z IIIII , at least 50 II | 11 (40 40 / 11 | nagimicatio | iii) iiiust be | evaluated in areas of flights | st mitotic densit | y in order to match wino 2010 |
| | | ntibody; % of 500–2,0 | 000 tumor c | ells in areas | s of highest | nuclear labeling. | | |
| | **MIB1 antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling. n cases of disparity between Ki-67 proliferative index and mitotic count, the result that indicates a higher-grade tumor should be selected as | | | | | | | |
| | | | | | | a Ki-67 of 12% should be de | | |
| | | , , , | | | | | = | |
| This | form | continues on the nex | t page. | | | | | |
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| Hospital Name/Address | Patient Name/Information |
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8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description | |
|---|-------------------------|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 | present/identified, NOS | |
| | 2 | mphatic and small vessel invasion only (L) | |
| | 3 | enous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 30.1. Anatomic sites used in the staging of tumors of the duodenum and ampulla of Vater.

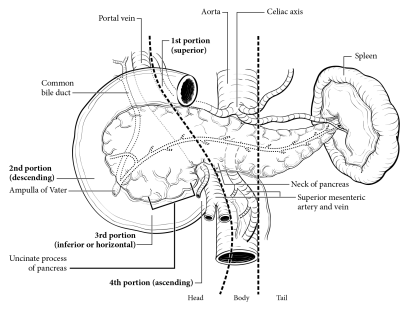
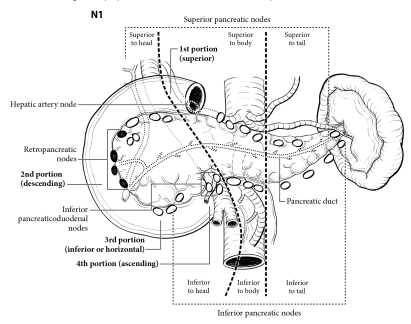


FIGURE 30.2. Regional lymph nodes of the duodenum and ampulla of Vater.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|--|---|---|--|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | |
| | ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoad therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | | |
| rTNM Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression unt treatment is initiated. | | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at auto and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient wit previously diagnosed cancer). | | | |

| Hospital Name/Address | Patient Name/Information | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | ⁻ Criteria | |
|---|---|--|--|
| | TX | Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | T1* | Invades lamina propria or submucosa and less than or equal to 1 cm in size | |
| | T2* Invades muscularis propria or greater than 1 cm in size | | |
| | T3* Invades through the muscularis propria into subserosal tissue without penetration of overlying serosa | | |
| | T4* Invades visceral peritoneum (serosal) or other organs or adjacent structures | | |

^{*}Note: For any T, add (m) for multiple tumors [TX(#) or TX(m), where X = 1–4, and # = number of primary tumors identified**]; for multiple tumors with different T, use the highest.

^{**}Example: If there are two primary tumors, only one of which invades through the muscularis propria into subserosal tissue without penetration of overlying serosa (jejunal or ileal), we define the primary tumor as either T3(2) or T3(m).

| ✓ | T Suffix | Definition |
|----------|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis has occurred | |
| | N1 | Regional lymph node metastasis less than 12 nodes | |
| | N2 Large mesenteric masses (>2 cm) and/or extensive nodal deposits (12 or greater), especially those that encase t | | |
| | | superior mesenteric vessels | |

| , | / N | N Suffix | Definition |
|--|--|--|------------|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category M Criteria | |
|--|-----------------------|---|
| cM0 No distant metastasis | | No distant metastasis |
| cM1 Distant metastasis | | Distant metastasis |
| | cM1a | Metastasis confined to liver |
| | cM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) |
| | cM1c | Both hepatic and extrahepatic metastases |
| | pM1 | Distant metastasis, microscopically confirmed |
| pM1a Metastasis confined to liver, microscopically confirmed | | Metastasis confined to liver, microscopically confirmed |
| pM1b Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum microscopically confirmed | | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone), microscopically confirmed |
| pM1c Both hepatic and extrahepatic metastases, microscopically confirmed | | Both hepatic and extrahepatic metastases, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When Tis | And N is | And Mis | Then the stage group is |
|---|----------|----------------|---------|-------------------------|
| | TX, T0 | NX, N0, N1, N2 | M1 | IV |
| | T1 | NO | M0 | 1 |
| | T1 | N1, N2 | M0 | III |
| | T1 | NX, N0, N1, N2 | M1 | IV |
| | T2 | NO | M0 | II |
| | T2 | N1, N2 | MO | III |
| | T2 | NX, N0, N1, N2 | M1 | IV |
| | T3 | NO | M0 | II |
| | T3 | N1, N2 | M0 | III |
| | T3 | NX, N0, N1, N2 | M1 | IV |
| | T4 | NO | M0 | III |
| | T4 | N1, N2 | M0 | III |
| | T4 | NX, N0, N1, N2 | M1 | IV |

For multiple synchronous tumors, the highest T category should be used and the multiplicity or the number of tumors should be indicated in parenthesis: e.g., T3(2) or T3(m).

| o Registry Data Concetton variable | 6 | Registry | Data | Collection | Variable |
|------------------------------------|---|----------|------|------------|----------|
|------------------------------------|---|----------|------|------------|----------|

| _ | • | 8.51.7 2 4.4 40.104.101.141.145.165 |
|-----|---|--|
| See | chapt | er for more details on these variables. |
| | 1. | Size of tumor (value): |
| | 2. | Tumor focality (unifocal or multifocal): |
| | 3. | Depth of Invasion: |
| | 4. | Nodal status and number of nodes involved, if applicable: |
| | 5. | Sites of metastasis, if applicable: |
| | 6. | NKA level: |
| | 7. | Pancreastatin level: |
| | 8. | Ki-67 index: |
| | 9. | Mitotic count: |
| | 10. | Histologic grading (from Ki-67 and mitotic count): GX G1 G2 G3 |
| 7 | Hi | stologic Grade (G) |

| ✓ | G G Definition | |
|--|--|---|
| | GX | Grade cannot be assessed |
| | G1 Mitotic count (per 10 HPF)* < 2 and Ki-67 index (%)** < 3 | |
| G2 Mitotic count (per 10 HPF) = 2–20 or Ki-67 index (%)** = 3–20 | | |
| G3 Mitotic count (per 10 HPF) > 20 or Ki-67 index (%)** > 20 | | Mitotic count (per 10 HPF) > 20 or Ki-67 index (%)** > 20 |

^{*10} HPF = 2 mm²; at least 50 HPFs (at 40× magnification) must be evaluated in areas of highest mitotic density in order to adhere to WHO 2010 criteria.

In cases of disparity between Ki-67 proliferative index and mitotic count, the result indicating a higher-grade tumor should be selected as the final grade. For example, a mitotic count of 1 per 10 HPF and a Ki-67 of 12% should be designated as a G2 NET.

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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^{**}MIB1 antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling.

31. Neuroendocrine Tumors of the Jejunum and Ileum

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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9 Anatomy

FIGURE 31.1. Anatomic sites of the small intestine. This chapter stages neuroendocrine tumors of the jejunum and ileum. See chapter 30 for more information about staging neuroendocrine tumors of the duodenum.

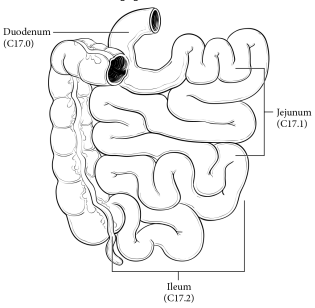
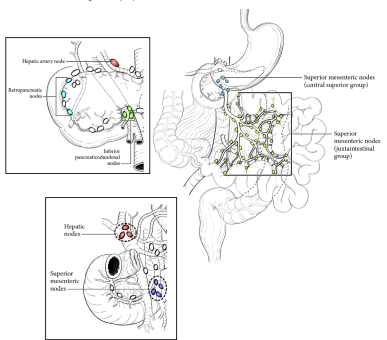


FIGURE 31.2. The regional lymph nodes of the small intestine for neuroendocrine tumors



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
|--------------------------|--------------------------|
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| | |
| | Patient Name/Information |

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor 2 cm or less in greatest dimension |
| | T2 | Tumor more than 2 cm but less than or equal to 4 cm |
| | T3 | Tumor more than 4 cm or with subserosal invasion or involvement of the mesoappendix |
| | T4 | Tumor perforates the peritoneum or directly invades other adjacent organs or structures (excluding direct mural |
| | | extension to adjacent subserosa of adjacent bowel), e.g., abdominal wall and skeletal muscle |

| ✓ | T Suffix | Definition |
|----------|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Metastasis confined to liver |
| | cM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) |
| | cM1c | Both hepatic and extrahepatic metastases |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Metastasis confined to liver, microscopically confirmed |
| | pM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone), microscopically confirmed |
| | pM1c | Both hepatic and extrahepatic metastases, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|------------|----------|-------------------------|
| | TX, TO | NX, N0, N1 | M1 | IV |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T1 | NX, N0, N1 | M1 | IV |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | III |
| | T2 | NX, N0, N1 | M1 | IV |
| | T3 | N0 | M0 | II |
| | T3 | N1 | M0 | III |
| | T3 | NX, N0, N1 | M1 | IV |
| | T4 | N0 | M0 | III |
| | T4 | N1 | M0 | III |
| | T4 | NX, N0, N1 | M1 | IV |

6 **Registry Data Collection Variables**

| hapt | er for more details on these variables. | | |
|------|--|--|--|
| 1. | Size of tumor: | | |
| 2. | Depth of invasion: | | |
| 3. | . Invasion of mesoappendix: | | |
| 4. | Number of nodes involved, mesenteric mass, mesenteric vessel encasement: | | |
| 5. | Perineural invasion: | | |
| 6. | Lymphovascular invasion: | | |
| 7. | Sites of metastasis, if applicable: | | |
| 8. | Type of surgery: | | |
| 9. | Ki-67 proliferative index: | | |
| 10. | Mitotic count: | | |
| 11. | Histologic grading (from Ki-67 and mitotic count): GX G1 G2 G3 | | |

Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Mitotic count (per 10 HPF)* < 2 and Ki-67 index (%)** < 3 |
| | G2 | Mitotic count (per 10 HPF) = 2–20 or Ki-67 index (%)** = 3–20 |
| | G3 | Mitotic count (per 10 HPF) > 20 or Ki-67 index (%)** > 20 |
| *10 HPF = 2 mm²; at least 50 HPFs (at 40× magnification) must be evaluated in areas of highest mitotic density in order to match WHO 2010 | | |

In cases of disparity between Ki-67 (proliferative index) and mitotic count, the result indicating a higher-grade tumor should be selected as the final grade. For example, a mitotic count of 1 per 10 HPF and a Ki-67 of 12% should be designated as a G2 NET.

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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^{**}MIB1 antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling.

32. Neuroendocrine Tumors of the Appendix

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|----------|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| | T |
|-----------------------|--------------------------|
| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 32.1. Anatomic location of the appendix

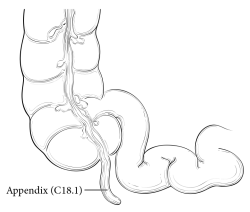
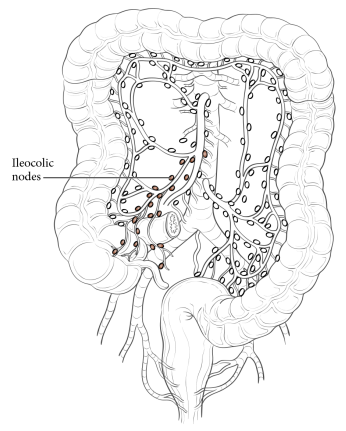


FIGURE 32.2. The regional lymph nodes of the appendix.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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1 Terms of Use

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|---|--|---|--|--|
| workup information, until first treatment, including clinical history and symptoms, physical examina endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resectic relevant examinations | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy | | |
| ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neo therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | |
| | rTNM Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | | |
| | | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---|--|--|
| | TX Primary tumor cannot be assessed | | |
| | TO No evidence of primary tumor | | |
| | T1 | Tumor invades the lamina propria or submucosa and is ≤2 cm | |
| | T1a Tumor <1 cm in greatest dimension | | |
| | T1b Tumor 1–2 cm in greatest dimension | | |
| | T2 Tumor invades the muscularis propria or is >2 cm with invasion of the lamina propria or submucosa | | |
| | T3 Tumor invades through the muscularis propria into subserosal tissue without penetration of overlying seros | | |
| | T4 Tumor invades the visceral peritoneum (serosa) or other organs or adjacent structures | | |

^{*}Note: For any T, add "(m)" for multiple tumors [TX(#) or TX(m), where X = 1–4 and # = number of primary tumors identified**]; for multiple tumors with different T, use the highest.

^{**}Example: If there are two primary tumors, only one of which invades through the muscularis propria into the subserosal tissue without penetration of the overlying serosa, we define the primary tumor as either T3(2) or T3(m).

| ✓ | T Suffix Definition | |
|---|---------------------|--|
| (m) Select if synchronous primary tumors are found in single organ. | | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|---|------------|--|
| | NX Regional lymph nodes cannot be assessed | | |
| | NO No regional lymph node metastasis has occurred | | |
| | N1 Regional lymph node metastasis | | |

| ✓ | ✓ N Suffix Definition | |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria | |
|---|------------|---|--|
| cM0 No distant metastasis | | No distant metastasis | |
| cM1 Distant metastasis | | Distant metastasis | |
| | cM1a | Metastasis confined to liver | |
| | cM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) | |
| | cM1c | Both hepatic and extrahepatic metastases | |
| pM1 Distant metastasis, microscopically confirmed | | Distant metastasis, microscopically confirmed | |
| pM1a Metastasis confined to liver, microscopically confirmed | | Metastasis confined to liver, microscopically confirmed | |
| pM1b Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, periton microscopically confirmed | | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone), microscopically confirmed | |
| | pM1c | Both hepatic and extrahepatic metastases, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information |
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| | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | TX, T0 | Any N | M1 | IV |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | IIIB |
| | T1 | Any N | M1 | IV |
| | T2 | N0 | M0 | IIA |
| | T2 | N1 | M0 | IIIB |
| | T2 | Any N | M1 | IV |
| | T3 | N0 | M0 | IIB |
| | T3 | N1 | M0 | IIIB |
| | T3 | Any N | M1 | IV |
| | T4 | N0 | M0 | IIIA |
| | T4 | N1 | M0 | IIIB |
| | T4 | Any N | M1 | IV |

Note: For multiple synchronous tumors, the highest T category should be used and the multiplicity or the number of tumors should be indicated in parenthesis, e.g., T3(2) or T3(m).

| o Registry Data Concetton variable | 6 | Registry | Data | Collection | Variable |
|------------------------------------|---|----------|------|------------|----------|
|------------------------------------|---|----------|------|------------|----------|

| See | ee chapter for more details on these variables. | | | | | | |
|-----|---|--|--|--|--|--|--|
| | 1. | Tumor site: | | | | | |
| | 2. | Size of tumor (value): | | | | | |
| | 3. | Depth of invasion: | | | | | |
| | 4. | Nodal status and number of nodes involved, if applicable: | | | | | |
| | 5. | Sites of metastasis, if applicable: | | | | | |
| | 6. | Ki-67 index: | | | | | |
| | 7. | Mitotic count: | | | | | |
| | 8. | Histologic grade (from Ki-67 and mitotic count): GX G1 G2 G3 | | | | | |

7 Histologic Grade (G)

| ✓ | G | G Definition |
|-----|---|---|
| | GX | Grade cannot be assessed |
| | G1 Mitotic count (per 10 HPF)* <2 and Ki-67 Index (%)** <3 | |
| | G2 Mitotic count (per 10 HPF) = 2–20 or Ki-67 index (%)** = 3–20 | |
| | G3 | Mitotic count (per 10 HPF) >20 or Ki-67 index (%)** >20 |
| *10 | *10 HPF = 2 mm ² ; at least 50 HPF (at 40× magnification) must be evaluated in areas of highest mitotic density in order to adhere to WHO 2010 | |

^{*10} HPF = 2 mm*; at least 50 HPF (at 40× magnification) must be evaluated in areas of nignest mitotic density in order to adhere to WHO 2010 criteria.

In cases of disparity between Ki-67 proliferative index and mitotic count, the result indicating a higher-grade tumor should be selected as the final grade. For example, a mitotic count of 1 per 10 HPF and a Ki-67 of 12% should be designated as a G2 NET.

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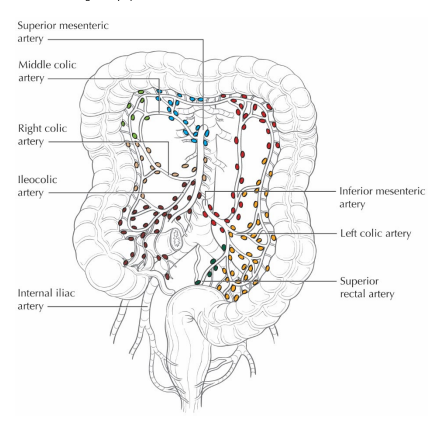
 $[\]ensuremath{^{**}\text{MIB1}}$ antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | LVI County | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 33.1. Regional lymph nodes for NETs of the colon and rectum.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Tumor cannot be assessed |
| | T1 | Tumor limited to the pancreas,* <2 cm |
| | T2 | Tumor limited to the pancreas,* 2–4 cm |
| | T3 | Tumor limited to the pancreas,* >4 cm; or tumor invading the duodenum or common bile duct |
| | T4 | Tumor invading adjacent organs (stomach, spleen, colon, adrenal gland) or the wall of large vessels (celiac axis or |
| | | the superior mesenteric artery) |

*Limited to the pancreas means there is no invasion of adjacent organs (stomach, spleen, colon, adrenal gland) or the wall of large vessels (celiac axis or the superior mesenteric artery). Extension of tumor into peripancreatic adipose tissue is NOT a basis for staging.

Note: Multiple tumors should be designated as such (the largest tumor should be used to assign T category):

- If the number of tumors is known, use T(#); e.g., pT3(4) N0 M0.
- If the number of tumors is unavailable or too numerous, use the m suffix, T(m); e.g., pT3(m) N0 M0.

| Ī | ✓ | T Suffix | Definition |
|---|---|----------|---|
| ſ | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node involvement |
| | N1 | Regional lymph node involvement |

| √ | N Suffix | Definition | |
|----------|--|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastases |
| | cM1a | Metastasis confined to liver |
| | cM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone) |
| | cM1c | Both hepatic and extrahepatic metastases |
| | pM1 | Distant metastases, microscopically confirmed |
| | pM1a | Metastasis confined to liver, microscopically confirmed |
| | pM1b | Metastases in at least one extrahepatic site (e.g., lung, ovary, nonregional lymph node, peritoneum, bone), |
| | | microscopically confirmed |
| | pM1c | Both hepatic and extrahepatic metastases, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | II |
| | T4 | N0 | M0 | III |
| | Any T | N1 | M0 | III |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

| | See chapter for more details on these variables. | | |
|--|--|--|--|
| See chapter for more details on these variables. | | | |
| 1. | Size of tumor (value): | | |
| 2. | Presence of invasion into adjacent organs/structures: Yes No | | |
| | If yes, which ones (pick all that apply): | | |
| | ☐ Stomach ☐ Duodenum ☐ Spleen ☐ Colon ☐ Other: | | |
| | If yes, were multiple adjacent organs involved? | | |
| | ☐ Yes ☐ No | | |
| 3. | Presence of necrosis: | | |
| 4. | Number of tumors (multicentric disease at primary site): | | |
| 5. | Lymph node status (including number of lymph nodes assessed and number of positive nodes): | | |
| 6. | Grade (based on Ki-67 and/or mitotic count): GX G1 G2 G3 | | |
| 7. | Mitotic count (value): | | |
| 8. | Ki-67 Labeling Index (value): | | |
| 9. | Perineural invasion: Yes No | | |
| 10. | Lymphovascular invasion: Yes No | | |
| 11. | Margin status: Positive (+) Negative (-) | | |
| 12. | Functional status: Yes No If yes, type of syndrome: | | |
| 13. | Genetic syndrome: Yes No If yes, type of syndrome: | | |
| 14. | Location in pancreas: head tail body junction body/tail junction body/head unknown | | |
| 15. | Type of surgery: enucleation distal pancreatectomy with splenectomy | | |
| | distal pancreatectomy without splenectomy central pancreatectomy | | |
| | pancreaticoduodenectomy (Whipple procedure) unknown other | | |
| 16. | Preoperative CgA level (absolute value with ULN): | | |
| 17. | Preoperative pancreastatin level (absolute value with ULN): | | |
| 18. | Age of patient: | | |
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| i iospital i | Name/Address Patient Name/Information | | |
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7 Histologic Grade (G)

| √ | G | G Definition |
|----------|--|---|
| | GX | Grade cannot be assessed |
| | G1 | Mitotic count (per 10 HPF)* <2 and Ki-67 index (%)** <3 |
| | G2 | Mitotic count (per 10 HPF) = 2–20 or Ki-67 index (%)** = 3–20 |
| | G3 | Mitotic count (per 10 HPF) >20 or Ki-67 index (%)** >20 |
| *10 | *10 UDF = 2 mm ² ; at least FO UDF (at 40); magnification) must be evaluated in areas of highest mitatic density in order to match WUO 2010 | |

^{*10} HPF = 2 mm²; at least 50 HPF (at 40× magnification) must be evaluated in areas of highest mitotic density in order to match WHO 2010 criteria.

In cases of disparity between Ki-67 proliferative index and mitotic count, the result that indicates a higher-grade tumor should be selected as the final grade. For example, a mitotic count of 1 per 10 HPF and a Ki-67 of 12% should be designated as a G2 NET.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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^{**}MIB1 antibody; % of 500–2,000 tumor cells in areas of highest nuclear labeling.

9 Anatomy

FIGURE 34.1. Anatomy of the pancreas.

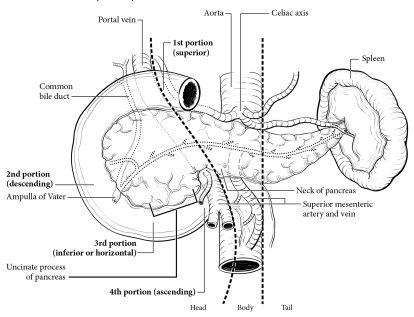
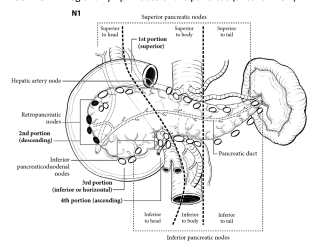


FIGURE 34.2. Regional lymph nodes of the pancreas (anterior view).



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor encapsulated or extending into the mediastinal fat; may involve the mediastinal pleura |
| | T1a | Tumor with no mediastinal pleura involvement |
| | T1b | Tumor with direct invasion of mediastinal pleura |
| | T2 | Tumor with direct invasion of the pericardium (either partial or full thickness) |
| | T3 | Tumor with direct invasion into any of the following: lung, brachiocephalic vein, superior vena cava, phrenic |
| | | nerve, chest wall, or extrapericardial pulmonary artery or veins |
| | T4 | Tumor with invasion into any of the following: aorta (ascending, arch, or descending), arch vessels, |
| | | intrapericardial pulmonary artery, myocardium, trachea, esophagus |

^{*}Involvement must be microscopically confirmed in pathological staging, if possible.

^{**}T categories are defined by "levels" of invasion; they reflect the highest degree of invasion regardless of how many other (lower-level) structures are invaded. T1, level 1 structures: thymus, anterior mediastinal fat, mediastinal pleura; T2, level 2 structures: pericardium; T3, level 3 structures: lung, brachiocephalic vein, superior vena cava, phrenic nerve, chest wall, hilar pulmonary vessels; T4, level 4 structures: aorta (ascending, arch, or descending), arch vessels, intrapericardial pulmonary artery, myocardium, trachea, esophagus.

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| ſ | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria | |
|----------|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | NO No regional lymph node metastasis | | |
| | N1 Metastasis in anterior (perithymic) lymph nodes | | |
| | N2 Metastasis in deep intrathoracic or cervical lymph nodes | | |
| *Inv | *Involvement must be microscopically confirmed in pathological staging, if possible. | | |

| ✓ | N Suffix | V Suffix Definition | |
|---|--|---------------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|---|------------|--|
| | cMO No pleural, pericardial, or distant metastasis | | |
| | cM1 Pleural, pericardial, or distant metastasis | | |
| | cM1a Separate pleural or pericardial nodule(s) | | |
| | cM1b Pulmonary intraparenchymal nodule or distant organ metastasis | | |
| | pM1 Pleural, pericardial, or distant metastasis, microscopically confirmed | | |
| | pM1a Separate pleural or pericardial nodule(s), microscopically confirmed | | |
| | pM1b Pulmonary intraparenchymal nodule or distant organ metastasis, microscopically confirmed | | |

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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1a,b | N0 | M0 | |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | IIIA |
| | T4 | NO | M0 | IIIB |
| | Any T | N1 | M0 | IVA |
| | Any T | N0,1 | M1a | IVA |
| | Any T | N2 | M0,M1a | IVB |
| | Any T | Any N | M1b | IVB |

6 Registry Data Collection Variables

Beyond the factors required for staging, the authors have not noted any registry data collection variables.

7 Histologic Grade (G)

There is no recommended histologic grading system at this time.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | M Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed, or tumor proven by the presence of malignant cells in sputum or bronchial |
| | | washings but not visualized by imaging or bronchoscopy |
| | T0 | No evidence of primary tumor |
| | Tis | Carcinoma in situ |
| | | Squamous cell carcinoma in situ (SCIS) |
| | | Adenocarcinoma in situ (AIS): adenocarcinoma with pure lepidic pattern, ≤3 cm in greatest dimension |
| | T1 | Tumor ≤3 cm in greatest dimension, surrounded by lung or visceral pleura, without bronchoscopic evidence of |
| | | invasion more proximal than the lobar bronchus (i.e., not in the main bronchus) |
| | T1mi | Minimally invasive adenocarcinoma: adenocarcinoma (≤3 cm in greatest dimension) with a predominantly lepidic |
| | | pattern and ≤5 mm invasion in greatest dimension |
| | T1a | Tumor ≤1 cm in greatest dimension. A superficial, spreading tumor of any size whose invasive component is |
| | | limited to the bronchial wall and may extend proximal to the main bronchus also is classified as T1a, but these |
| | | tumors are uncommon. |
| | T1b | Tumor >1 cm but ≤2 cm in greatest dimension |
| | T1c | Tumor >2 cm but ≤3 cm in greatest dimension |
| | T2 | Tumor >3 cm but ≤5 cm or having any of the following features: |
| | | Involves the main bronchus regardless of distance to the carina, but without involvement of the carina |
| | | Invades visceral pleura (PL1 or PL2) |
| | | Associated with atelectasis or obstructive pneumonitis that extends to the hilar region, involving part or all of the lung |
| | | T2 tumors with these features are classified as T2a if ≤4 cm or if the size cannot be determined and T2b if >4 cm |
| | | but ≤5 cm. |
| | T2a | Tumor >3 cm but ≤4 cm in greatest dimension |
| T2b Tumor >4 cm but ≤5 cm in greatest dimension | | Tumor >4 cm but ≤5 cm in greatest dimension |
| | T3 | Tumor >5 cm but ≤7 cm in greatest dimension or directly invading any of the following: parietal pleura (PL3), |
| | | chest wall (including superior sulcus tumors), phrenic nerve, parietal pericardium; or separate tumor nodule(s) in |
| | | the same lobe as the primary |
| | T4 | Tumor >7 cm or tumor of any size invading one or more of the following: diaphragm, mediastinum, heart, great |
| | | vessels, trachea, recurrent laryngeal nerve, esophagus, vertebral body, or carina; separate tumor nodule(s) in an |
| | | ipsilateral lobe different from that of the primary |

| ١ | / | T Suffix | Definition |
|---|---|----------|---|
| | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|---|--|--|
| | NX Regional lymph nodes cannot be assessed | | |
| | NO No regional lymph node metastasis | | |
| | N1 Metastasis in ipsilateral peribronchial and/or ipsilateral hilar lymph nodes and intrapulmonary nodes, includi involvement by direct extension | | |
| | N2 Metastasis in ipsilateral mediastinal and/or subcarinal lymph node(s) | | |
| | N3 | Metastasis in contralateral mediastinal, contralateral hilar, ipsilateral or contralateral scalene, or supraclavicular lymph node(s) | |

| 1 | N Suffix | Definition |
|---|--|---|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|--|---|
| | cM0 | No distant metastasis |
| | cM1 Distant metastasis | |
| | cM1a | Separate tumor nodule(s) in a contralateral lobe; tumor with pleural or pericardial nodules or malignant pleural or |
| | | pericardial effusion. Most pleural (pericardial) effusions with lung cancer are a result of the tumor. In a few patients, |
| | | however, multiple microscopic examinations of pleural (pericardial) fluid are negative for tumor, and the fluid is |
| | | nonbloody and not an exudate. If these elements and clinical judgment dictate that the effusion is not related to the |
| | | tumor, the effusion should be excluded as a staging descriptor. |
| cM1b Single extrathoracic metastasis in a single organ (including involvement of a single nonregional node) | | Single extrathoracic metastasis in a single organ (including involvement of a single nonregional node) |
| | cM1c Multiple extrathoracic metastases in a single organ or in multiple organs pM1 Distant metastasis, microscopically confirmed | |
| | | |
| | pM1a | Separate tumor nodule(s) in a contralateral lobe; tumor with pleural or pericardial nodules or malignant pleural or |
| | | pericardial effusion, microscopically confirmed. Most pleural (pericardial) effusions with lung cancer are a result of |
| | | the tumor. In a few patients, however, multiple microscopic examinations of pleural (pericardial) fluid are negative |
| | | for tumor, and the fluid is nonbloody and not an exudate. If these elements and clinical judgment dictate that the |
| | | effusion is not related to the tumor, the effusion should be excluded as a staging descriptor. |
| | pM1b | Single extrathoracic metastasis in a single organ (including involvement of a single nonregional node), |
| | | microscopically confirmed |
| | pM1c | Multiple extrathoracic metastases in a single organ or in multiple organs, microscopically confirmed |

 TABLE 36.12. Guide to uniform categorization of situations beyond the standard descriptors

| Situation | Category |
|--|------------|
| Direct invasion of an adjacent lobe, across the fissure or directly if the fissure is incomplete, unless other criteria assign | |
| a higher T | T2a |
| Invasion of phrenic nerve | T3 |
| Paralysis of the recurrent laryngeal nerve, superior vena caval obstruction, or compression of the trachea or esophagus related to direct extension of the primary tumor | |
| · | T4 |
| Paralysis of the recurrent laryngeal nerve, superior vena caval obstruction, or compression of the trachea or esophagus related to lymph node involvement | |
| | N2 |
| Involvement of great vessels: aorta, superior vena cava, inferior vena cava, main pulmonary artery (pulmonary trunk), intrapericardial portions of the right and left pulmonary artery, intrapericardial portions of the superior and inferior | |
| right and left pulmonary veins | T4 |
| Pancoast tumors with evidence of invasion of the vertebral body or spinal canal, encasement of the subclavian vessels, or unequivocal involvement of the superior branches of the brachial plexus (C8 or above) | |
| | T4 |
| Pancoast tumors without the criteria for T4 classification | T3 |
| Direct extension to parietal pericardium | T3 |
| Direct extension to visceral pericardium | T4 |
| Tumor extending to rib | T3 |
| Invasion into hilar fat, unless other criteria assign a higher T | T2a |
| Invasion into mediastinal fat | T4 |
| Discontinuous tumor nodules in the ipsilateral parietal or visceral pleura | M1a |
| Discontinuous tumor nodules outside the parietal pleura in the chest wall or in the diaphragm | M1b or M1c |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ When T is | And N is | And M is | Then the stage group is |
|-------------|----------|----------|-------------------------|
| TX | N0 | M0 | Occult carcinoma |
| Tis | N0 | M0 | 0 |
| T1mi | N0 | MO | IA1 |
| T1a | N0 | M0 | IA1 |
| T1a | N1 | M0 | IIB |
| T1a | N2 | MO | IIIA |
| T1a | N3 | M0 | IIIB |
| T1b | N0 | MO | IA2 |
| T1b | N1 | MO | IIB |
| T1b | N2 | M0 | IIIA |
| T1b | N3 | M0 | IIIB |
| T1c | N0 | M0 | IA3 |
| T1c | N1 | M0 | IIB |
| T1c | N2 | MO | IIIA |
| T1c | N3 | M0 | IIIB |
| T2a | N0 | MO | IB |
| T2a | N1 | M0 | IIB |
| T2a | N2 | M0 | IIIA |
| T2a | N3 | M0 | IIIB |
| T2b | N0 | M0 | IIA |
| T2b | N1 | M0 | IIB |
| T2b | N2 | MO | IIIA |
| T2b | N3 | MO | IIIB |
| Т3 | N0 | MO | IIB |
| T3 | N1 | M0 | IIIA |
| T3 | N2 | M0 | IIIB |
| Т3 | N3 | MO | IIIC |
| T4 | N0 | M0 | IIIA |
| T4 | N1 | M0 | IIIA |
| T4 | N2 | MO | IIIB |
| T4 | N3 | MO | IIIC |
| Any T | Any N | M1 | IV |
| Any T | Any N | M1a | IVA |
| Any T | Any N | M1b | IVA |
| Any T | Any N | M1c | IVB |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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6 Registry Data Collection Variables

See chapter for more details on these variables.

For data collection, all T, N, and M descriptors and at least the prognostic factors considered essential and additional in Additional Factors Recommended for Clinical Care should be collected.

| 1. | Patient related | | |
|-----|--------------------------|-------------------------------------|--|
| | a. | Gender: | |
| | b. | Age: | |
| | c. | Weight loss: | |
| | d. | Performance status: | |
| 2. | Environm | nent related | |
| | a. | Resection margins: | |
| | b. | Adequacy of mediastinal dissection: | |
| For | advanced : | non–small cell lung cancer | |
| 1. | Tumor re | lated | |
| | a. | EGFR mutation: | |
| | b. | ALK gene rearrangement: | |
| 2. | Patient related | | |
| | a. | Gender: | |
| | b. | Symptoms: | |
| | c. | Weight loss: | |
| | d. | Performance status: | |
| 3. | Environm | nent related | |
| | a. | Chemoradiotherapy: | |
| | b. | Chemotherapy: | |
| For | r small cell lung cancer | | |
| 1. | Patient related | | |
| | a. | Performance status: | |
| | b. | Age: | |
| | c. | Comorbidity: | |
| 2. | Environment related | | |
| | a. | Chemotherapy: | |
| | b. | Thoracic radiotherapy: | |
| | c. | Prophylactic cranial radiotherapy: | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade of differentiation cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description | |
|---|-------------------------|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Physician Signature | Date/Time |
|----------------------|------------|
| i nysician signature | Date/ Time |

| ent Name/Information |
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1 Terms of Use

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | | |
|---|--|---|--|--|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | | |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information findiagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgespecimens | | | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | | |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor limited to the ipsilateral parietal pleura with or without involvement of |
| | | visceral pleura |
| | | mediastinal pleura |
| | | diaphragmatic pleura |
| | T2 | Tumor involving each of the ipsilateral pleural surfaces (parietal, mediastinal, diaphragmatic, and visceral pleura) |
| | | with at least one of the following features: |
| | | involvement of diaphragmatic muscle |
| | | extension of tumor from visceral pleura into the underlying pulmonary parenchyma |
| | T3 | Describes locally advanced but potentially resectable tumor. |
| | | Tumor involving all the ipsilateral pleural surfaces (parietal, mediastinal, diaphragmatic, and visceral pleura) with |
| at least one of the following features: | | <u> </u> |
| involvement of the endothoracic fascia | | |
| | | extension into the mediastinal fat |
| | | solitary, completely resectable focus of tumor extending into the soft tissues of the chest wall |
| | | nontransmural involvement of the pericardium |
| | T4 | Describes locally advanced technically unresectable tumor. |
| | | Tumor involving all the ipsilateral pleural surfaces (parietal, mediastinal, diaphragmatic, and visceral pleura) with at least one of the following features: |
| | | diffuse extension or multifocal masses of tumor in the chest wall, with or without associated rib |
| · · · · · · · · · · · · · · · · · · · | | destruction |
| | | direct transdiaphragmatic extension of tumor to the peritoneum |
| | | direct extension of tumor to the contralateral pleura |
| | | direct extension of tumor to mediastinal organs |
| | | direct extension of tumor into the spine |
| | | tumor extending through to the internal surface of the pericardium with or without a pericardial effusion; or tumor involving the myocardium |

| ~ | T Suffix | Definition | |
|----------|----------|---|--|
| | (m) | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | | |
|---|------------|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | | |
| | N0 | No regional lymph node metastases | | |
| | N1 | Metastases in the ipsilateral bronchopulmonary, hilar, or mediastinal (including the internal mammary, | | |
| | | peridiaphragmatic, pericardial fat pad, or intercostal) lymph nodes | | |
| | N2 | Metastases in the contralateral mediastinal, ipsilateral, or contralateral supraclavicular lymph nodes | | |

| 1 | N Suffix | Definition | | |
|---|--|------------|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | | |

| Patient Name/Information | |
|--------------------------|--|
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37. Malignant Pleural Mesothelioma

4.3 **Definition of Distant Metastasis (M)**

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|---|------------|--|
| | cMO No distant metastasis | | |
| | cM1 Distant metastasis | | |
| | pM1 Distant metastasis, microscopically confirmed | | |

AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | N0 | M0 | IA |
| | T2 or T3 | N0 | M0 | IB |
| | T1 | N1 | M0 | II |
| | T2 | N1 | M0 | II |
| | T3 | N1 | M0 | IIIA |
| | T1-3 | N2 | M0 | IIIB |
| | T4 | Any N | M0 | IIIB |
| | Any T | Any N | M1 | IV |

| | Registry Data Collection Variables chapter for more details on these variables. | | |
|----|--|--|--|
| 1. | Histologic type: | | |
| 2. | Sex: | | |
| 3. | Age: | | |
| 4. | Performance status: | | |
| 5. | Laboratory parameters including | | |
| | a. WBC: | | |
| | b. Platelet count: | | |
| | c. Hemoglobin: | | |
| 6. | Surgical resection with curative intent: | pleurectomy/decortications extended pleurectomy/decortications | |
| | | extrapleural pneumonectomy | |
| 7. | For patients undergoing multimodality th | erapy, use of chemotherapy and/or radiotherapy: | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

37. Malignant Pleural Mesothelioma

7 Histologic Grade (G)

| ✓ | G G Definition | |
|---|--|---------------------------|
| | GX Grade of differentiation cannot be assessed | |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

9 Anatomy

FIGURE 37.9. Anatomy of the pleura.

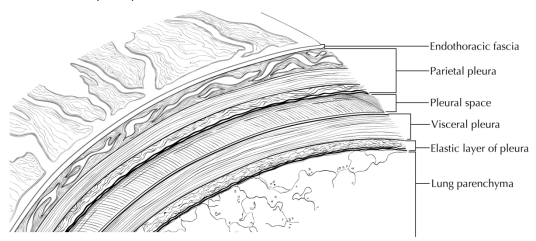
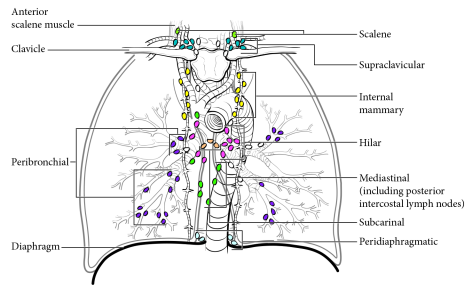


FIGURE 37.10. Regional lymph nodes of the pleura.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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1 Terms of Use

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
| | | |
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| | | |
| | | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤8 cm in greatest dimension |
| | T2 | Tumor >8 cm in greatest dimension |
| | T3 | Discontinuous tumors in the primary bone site |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed. |
| | | Because of the rarity of lymph node involvement in bone sarcomas, the designation NX may not be appropriate, and cases should be considered N0 unless clinical node involvement clearly is evident. |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition | |
|----------|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|------------|--|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | cM1a | Lung | |
| | cM1b | Bone or other distant sites | |
| | pM1 | Distant metastasis, microscopically confirmed | |
| | pM1a | M1a Lung, microscopically confirmed | |
| | pM1b | Bone or other distant sites. Microscopically confirmed | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

| ✓ | G | G Definition | |
|---|--|--------------------------------|--|
| | GX Grade cannot be assessed | | |
| | G1 | Well differentiated, low grade | |
| | G2 Moderately differentiated, high grade | | |
| | G3 Poorly differentiated, high grade | | |

| Name/Information |
|------------------|
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AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And G is | Then the stage |
|---|-----------|----------|----------|----------|----------------|
| • | | | | | group is |
| | T1 | N0 | M0 | G1 or GX | IA |
| | T2 | N0 | M0 | G1 or GX | IB |
| | T3 | N0 | M0 | G1 or GX | IB |
| | T1 | N0 | M0 | G2 or G3 | IIA |
| | T2 | N0 | M0 | G2 or G3 | IIB |
| | T3 | N0 | M0 | G2 or G3 | III |
| | Any T | N0 | M1a | Any G | IVA |
| | Any T | N1 | Any M | Any G | IVB |
| | Any T | Any N | M1b | Any G | IVB |

Registry Data Collection Variables

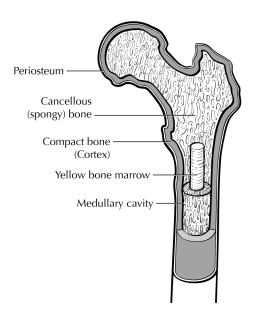
| | | • | | | |
|--|----|---|--|--|--|
| See chapter for more details on these variables. | | | | | |
| | 1. | Grade: | | | |
| | 2. | Three dimensions of tumor size: | | | |
| | 3. | Percentage of necrosis after neoadjuvant systemic therapy, from pathology report: | | | |
| | 4. | Number of resected pulmonary metastases, from pathology report: | | | |
| 8 | Ly | mphovascular Invasion (LVI) | | | |

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 38.12. The anatomic subsites of the bone.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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38.2. Bone: Spine

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3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|--|----------------|---|--|
| workup information, until first treatment, including clinical history and symptoms, physical examinati endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | TNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information fro diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgic specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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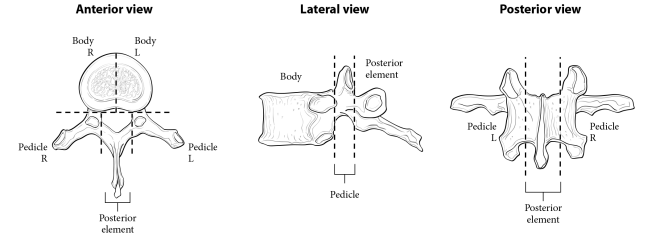
Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria | |
|---|---------------------------------|---|--|
| | TX | TX Primary tumor cannot be assessed | |
| | TO No evidence of primary tumor | | |
| | T1 | Tumor confined to one vertebral segment or two adjacent vertebral segments | |
| | T2 | Tumor confined to three adjacent vertebral segments | |
| | T3 | Tumor confined to four or more adjacent vertebral segments, or any nonadjacent vertebral segments | |
| | T4 | Extension into the spinal canal or great vessels | |
| | T4a | Extension into the spinal canal | |
| | T4b | Evidence of gross vascular invasion or tumor thrombus in the great vessels | |

| / | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

FIGURE 38.1. Spine segments for staging.



4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria |
|----------|------------|--|
| | NX | Regional lymph nodes cannot be assessed. |
| | | Because of the rarity of lymph node involvement in bone sarcomas, the designation NX may not be appropriate, and |
| | | cases should be considered N0 unless clinical node involvement clearly is evident. |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| 1 | N Suffix | Definition |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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38.2. Bone: Spine

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|--|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Lung |
| | cM1b | Bone or other distant sites |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Lung, microscopically confirmed |
| | pM1b | Bone or other distant sites. Microscopically confirmed |

| 5 | AJCC | Prognostic | Stage | Groups |
|---|------|-------------------|-------|--------|
|---|------|-------------------|-------|--------|

9

| There is no AJCC Prognostic Stage Grouping for spine. | Always refer to the specific chapter for rules on clinical and pathological classification | of |
|---|--|----|
| this disease. | | |

| 6 | Re | egistry Data Collection Variables |
|-----|-------|---|
| See | chapt | ter for more details on these variables. |
| | 1. | Grade: GX G1 G2 G3 |
| | 2. | Three dimensions of tumor size: |
| | 3. | Percentage of necrosis after neoadjuvant systemic therapy, from pathology report: |
| | 4. | Number of resected pulmonary metastases, from pathology report: |
| 7 | Ly | mphovascular Invasion (LVI) |

Component of LVI Coding Description UVI Coding UVI not present (absent)/not identified LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion

Presence of LVI unknown/indeterminate

| Physician Signature | Date/Time |
|---------------------|-----------|
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| Hospital Name/Address | Patient Name/Information |
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38.3. Bone: Pelvis

1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|--|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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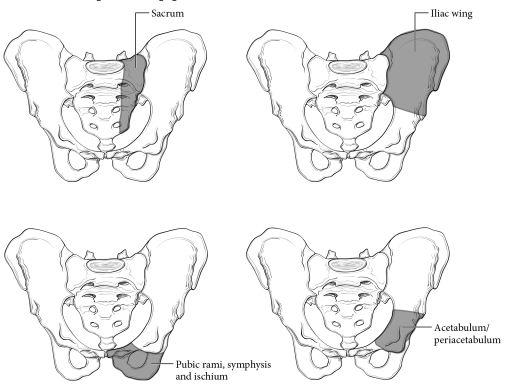
Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | T1 | Tumor confined to one pelvic segment with no extraosseous extension |
| | T1a | Tumor ≤8 cm in greatest dimension |
| | T1b | Tumor >8 cm in greatest dimension |
| | T2 | Tumor confined to one pelvic segment with extraosseous extension or two segments without extraosseous |
| | | extension |
| | T2a | Tumor ≤8 cm in greatest dimension |
| | T2b | Tumor >8 cm in greatest dimension |
| | T3 | Tumor spanning two pelvic segments with extraosseous extension |
| | T3a | Tumor ≤8 cm in greatest dimension |
| | T3b | Tumor >8 cm in greatest dimension |
| | T4 | Tumor spanning three pelvic segments or crossing the sacroiliac joint |
| | T4a | Tumor involves sacroiliac joint and extends medial to the sacral neuroforamen |
| | T4b | Tumor encasement of external iliac vessels or presence of gross tumor thrombus in major pelvic vessels |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

FIGURE 38.2. Pelvic segments for staging.



| Hospital Name/Address | Patient Name/Information | |
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38.3. Bone: Pelvis

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Regional lymph nodes cannot be assessed. |
| | | Because of the rarity of lymph node involvement in bone sarcomas, the designation NX may not be appropriate, and |
| | | cases should be considered N0 unless clinical node involvement clearly is evident. |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| | h | |
|---|------------|--|
| ✓ | M Category | M Criteria |
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Lung |
| | cM1b | Bone or other distant sites |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Lung, microscopically confirmed |
| | pM1b | Bone or other distant sites. Microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no AJCC Prognostic Stage Grouping for pelvis. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 Registry Data Collection Variables

| О | K | egistry Data Collection Variables |
|-----|------|---|
| See | chap | ter for more details on these variables. |
| | 1. | Grade: GX G1 G2 G3 |
| | 2. | Three dimensions of tumor size: |
| | 3. | Percentage of necrosis after neoadjuvant systemic therapy, from pathology report: |
| | 4. | Number of resected pulmonary metastases, from pathology report: |
| | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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7 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | Patient Name/Information | |
|--------------------------|--------------------------|--|
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| | | |
| | | |
| | Patient Name/Information | |

1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T1 | Tumor ≤2 cm |
| | T2 | Tumor >2 to ≤4 cm |
| | T3 | Tumor >4 cm |
| | T4 | Tumor with invasion of adjoining structures |
| | T4a | Tumor with orbital invasion, skull base/dural invasion, invasion of central compartment viscera, involvement of |
| | | facial skeleton, or invasion of pterygoid muscles |
| | T4b | Tumor with brain parenchymal invasion, carotid artery encasement, prevertebral muscle invasion, or central |
| | | nervous system involvement via perineural spread |

| ✓ | T Suffix | Definition |
|----------|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| N Category | N Criteria |
|------------|--|
| N0 | No regional lymph node metastases or unknown lymph node status |
| N1 | Regional lymph node metastasis |

| | ✓ | N Suffix | Definition |
|---|---|----------|--|
| | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| Γ | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

This is a new classification that needs data collection before defining a stage grouping for head and neck sarcomas. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| Hospital Name/Address | Patient Name/Information |
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6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Bone invasion as determined by imaging:
- 2. If pM1, source of pathological metastatic specimen:
- 3. Additional dimensions of tumor size:
- 4. FNCLCC grade:
- 5. Central nervous system extension (head and neck primaries):

7 FNCLCC Histologic Grade (G)

The FNCLCC grade is determined by three parameters: differentiation, mitotic activity, and extent of necrosis. Each parameter is scored as follows: differentiation (1–3), mitotic activity (1–3), and necrosis (0–2). The scores are added to determine the grade.

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Total differentiation, mitotic count and necrosis score of 2 or 3 |
| | G2 | Total differentiation, mitotic count and necrosis score of 4 or 5 |
| | G3 | Total differentiation, mitotic count and necrosis score of 6, 7, or 8 |

7.1 Tumor Differentiation

Tumor differentiation is histology specific (see chapter 39, table 39.1) and is generally scored as follows:

| ✓ | Differentiation Score | Definition |
|---|-----------------------|--|
| | 1 | Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) |
| | 2 | Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) |
| | 3 | Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, |
| | | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue |

7.2 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at 400× magnification = 0.1734 mm²) are assessed using a 40× objective.

| | • . | |
|----------|---------------------|--------------------------|
| ✓ | Mitotic Count Score | Definition |
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

7.3 Tumor Necrosis

Evaluated on gross examination and validated with histologic sections.

| ✓ | Necrosis | Definition |
|---|----------|---------------------|
| | Score | |
| | 0 | No necrosis |
| | 1 | <50% tumor necrosis |
| | 2 | ≥50% tumor necrosis |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor 5 cm or less in greatest dimension |
| | T2 | Tumor more than 5 cm and less than or equal to 10 cm in greatest dimension |
| | T3 | Tumor more than 10 cm and less than or equal to 15 cm in greatest dimension |
| | T4 | Tumor more than 15 cm in greatest dimension |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria |
|----------|------------|--|
| | N0 | No regional lymph node metastases or unknown lymph node status |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | сМ0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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5 Prognostic Factors Required for Stage Grouping

5.1 Definition of FNCLCC Histologic Grade (G)

The FNCLCC grade is determined by three parameters: differentiation, mitotic activity, and extent of necrosis. Each parameter is scored as follows: differentiation (1–3), mitotic activity (1–3), and necrosis (0–2). The scores are added to determine the grade.

| ✓ | G | G Definition |
|---|-----------------------------|---|
| | GX Grade cannot be assessed | |
| | G1 | Total differentiation, mitotic count and necrosis score of 2 or 3 |
| | G2 | Total differentiation, mitotic count and necrosis score of 4 or 5 |
| | G3 | Total differentiation, mitotic count and necrosis score of 6, 7, or 8 |

5.1.1 Tumor Differentiation

Tumor differentiation is histology specific (see chapter 39, table 39.1) and is generally scored as follows:

| ✓ | Differentiation Score | Definition |
|---|-----------------------|--|
| | 1 | Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) |
| | 2 | Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) |
| | 3 | Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, |
| | | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue |

5.1.2 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at 400× magnification = 0.1734 mm²) are assessed using a 40× objective.

| ✓ | Mitotic Count Score | Definition |
|---|---------------------|--------------------------|
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

5.1.3 Tumor Necrosis

Evaluated on gross examination and validated with histologic sections.

| 1 | Necrosis | Definition |
|---|----------|---------------------|
| • | Score | |
| | 0 | No necrosis |
| | 1 | <50% tumor necrosis |
| | 2 | ≥50% tumor necrosis |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And G is | Then the stage |
|---|------------|----------|----------|----------|----------------|
| | | | | | group is |
| | T1 | N0 | M0 | G1, GX | IA |
| | T2, T3, T4 | N0 | M0 | G1, GX | IB |
| | T1 | N0 | M0 | G2, G3 | II |
| | T2 | N0 | M0 | G2, G3 | IIIA |
| | T3, T4 | N0 | M0 | G2, G3 | IIIB |
| | Any T | N1 | M0 | Any G | IV |
| | Any T | Any N | M1 | Any G | IV |

| Hospital Name/Address | Patient Name/Information | |
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41. Soft Tissue Sarcoma of the Trunk and Extremities

| 7 | Re | egistry Data | a Collection Variables | | | |
|------|--|----------------------|--|--------------------------|--|--|
| See | See chapter for more details on these variables. | | | | | |
| | 1. | Bone invasion | n as determined by imaging: | | | |
| | 2. | If pM1, source | e of pathological metastatic specimen: | | | |
| | 3. | Additional dir | mensions of tumor size: | | | |
| | 4. | FNCLCC grade |): | | | |
| 8 | Ly | mphovasci | ular Invasion (LVI) | | | |
| ✓ | | nponent of Coding | Description | | | |
| | 0 | | LVI not present (absent)/not identified | | | |
| | 1 | | LVI present/identified, NOS | | | |
| | 2 | | Lymphatic and small vessel invasion only (L) | | | |
| | 3 | | Venous (large vessel) invasion only (V) | | | |
| | 4 | | BOTH lymphatic and small vessel AND venous | (large vessel) invasion | | |
| | 9 | | Presence of LVI unknown/indeterminate | | | |
| | | | | | | |
| | | | | | | |
| Phy | sician | Signature | | Date/Time | | |
| | | | | | | |
| Hos | pital ! | Name/Address | | Patient Name/Information | | |
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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| | |
| | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T1 | Organ confined |
| | T2 | Tumor extension into tissue beyond organ |
| | T2a | Invades serosa or visceral peritoneum |
| | T2b | Extension beyond serosa (mesentery) |
| | T3 | Invades another organ |
| | T4 | Multifocal involvement |
| | T4a | Multifocal (2 sites) |
| | T4b | Multifocal (3-5 sites) |
| | T4c | Multifocal (> 5 sites) |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | N0 | No regional lymph node metastases or unknown lymph node status |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no recommended prognostic stage grouping at this time. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Bone invasion as determined by imaging:
- 2. If pM1, source of pathological metastatic specimen:
- 3. Additional dimensions of tumor size:
- 4. FNCLCC grade:
- 5. Evidence of multifocality (number of sites):

7 FNCLCC Histologic Grade (G)

The FNCLCC grade is determined by three parameters: differentiation, mitotic activity, and extent of necrosis. Each parameter is scored as follows: differentiation (1–3), mitotic activity (1–3), and necrosis (0–2). The scores are added to determine the grade.

| ✓ | G | G Definition |
|---|-----------------------------|---|
| | GX Grade cannot be assessed | |
| | G1 | Total differentiation, mitotic count and necrosis score of 2 or 3 |
| | G2 | Total differentiation, mitotic count and necrosis score of 4 or 5 |
| | G3 | Total differentiation, mitotic count and necrosis score of 6, 7, or 8 |

7.1 Tumor Differentiation

Tumor differentiation is histology specific (see chapter 39, table 39.1) and is generally scored as follows:

| √ | Differentiation Score | Definition |
|----------|-----------------------|--|
| | 1 | Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) |
| | 2 | Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) |
| | 3 | Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, |
| | | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue |

7.2 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at $400 \times \text{magnification} = 0.1734 \text{ mm}^2$) are assessed using a $40 \times \text{objective}$.

| √ | Mitotic Count Score | Definition |
|----------|---------------------|--------------------------|
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

7.3 Tumor Necrosis

Evaluated on gross examination and validated with histologic sections.

| _ | | | |
|---|---|----------|---------------------|
| | 1 | Necrosis | Definition |
| | • | Score | |
| | | 0 | No necrosis |
| Γ | | 1 | <50% tumor necrosis |
| | | 2 | ≥50% tumor necrosis |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | · · |

42. Soft Tissue Sarcoma of the Abdomen and Thoracic Visceral Organs

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | ! |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor 2 cm or less |
| | T2 | Tumor more than 2 cm but not more than 5 cm |
| | T3 | Tumor more than 5 cm but not more than 10 cm |
| | T4 | Tumor more than 10 cm in greatest dimension |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | N0 | No regional lymph node metastasis or unknown lymph node status |
| | N1 | Regional lymph node metastasis |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Mitotic Rate

| ✓ | Mitotic rate | Definition |
|---|--------------|--|
| | Low | 5 or fewer mitoses per 5 mm ² |
| | High | Over 5 mitoses per 5 mm ² |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
| | |

6 AJCC Prognostic Stage Groups

Hospital Name/Address

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And Mitotic Rate | Then the stage |
|---|-----------|----------|----------|------------------|----------------|
| • | | | | is | group is |
| | T1 or T2 | N0 | M0 | Low | IA |
| | T3 | N0 | M0 | Low | IB |
| | T1 | N0 | M0 | High | II |
| | T2 | N0 | M0 | High | II |
| | T4 | N0 | M0 | Low | II |
| | T3 | N0 | M0 | High | IIIA |
| | T4 | N0 | M0 | High | IIIB |
| | Any T | N1 | M0 | Any | IV |
| | Any T | Any N | M1 | Any | IV |

| 7 | Re | egistry Data | a Collection Variables |
|----------|-------------|----------------------|---|
| See | chap | ter for more de | tails on these variables. |
| | 1. | Tumor size: | |
| | 2. | Tumor site: | ☐ esophagus ☐ stomach ☐ duodenum ☐ jejunum/ileum ☐ rectum ☐ extraintestinal |
| | 3. | Tumor mitoti | c rate: |
| | 4. | Tumor ruptur | e: |
| | 5. | Tumor metas | tasis: |
| | 6. | Tumor KIT im | munohistochemistry: |
| | 7. | Tumor mutat | ional status of KIT, PDGFRA (if known): |
| | | | |
| 8 | | stologic Gr | |
| Gra | ding f | or GIST is depe | ndent on mitotic rate. |
| | | | |
| 9 | Ly | mphovasc | ular Invasion (LVI) |
| | | | |
| ✓ | | nponent of Coding | Description |
| | 0 | | |
| | | | LVI not present (absent)/not identified |
| | 1 | | LVI present/identified, NOS |
| | 2 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) |
| | | | LVI present/identified, NOS |
| | 2 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) |
| | 2 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) |
| | 2 3 4 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 2 3 4 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 2 3 4 | | LVI present/identified, NOS Lymphatic and small vessel invasion only (L) Venous (large vessel) invasion only (V) BOTH lymphatic and small vessel AND venous (large vessel) invasion |

Patient Name/Information

43.2. Gastrointestinal Stromal Tumor: Small Intestinal, Esophageal, Colorectal, Mesenteric, and Peritoneal GIST

1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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43.2. Gastrointestinal Stromal Tumor: Small Intestinal, Esophageal, Colorectal, Mesenteric, and Peritoneal GIST

4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor 2 cm or less |
| | T2 | Tumor more than 2 cm but not more than 5 cm |
| | T3 | Tumor more than 5 cm but not more than 10 cm |
| | T4 | Tumor more than 10 cm in greatest dimension |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category N Criteria | |
|---|-----------------------|--|
| | N0 | No regional lymph node metastasis or unknown lymph node status |
| | N1 | Regional lymph node metastasis |

| | ✓ | N Suffix | uffix Definition | |
|--|---|--|------------------|--|
| (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Mitotic Rate

| 1 | Mitotic rate | Definition |
|---|--------------|--|
| | Low | 5 or fewer mitoses per 5 mm ² |
| | High | Over 5 mitoses per 5 mm ² |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
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43.2. Gastrointestinal Stromal Tumor: Small Intestinal, Esophageal, Colorectal, Mesenteric, and Peritoneal GIST

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And Mitotic Rate | Then the stage |
|---|-----------|----------|----------|------------------|----------------|
| • | | | | is | group is |
| | T1 or T2 | N0 | M0 | Low | 1 |
| | T3 | N0 | M0 | Low | II |
| | T1 | N0 | M0 | High | IIIA |
| | T4 | N0 | M0 | Low | IIIA |
| | T2 | N0 | M0 | High | IIIB |
| | T3 | N0 | M0 | High | IIIB |
| | T4 | N0 | M0 | High | IIIB |
| | Any T | N1 | M0 | Any rate | IV |
| | Any T | Any N | M1 | Any rate | IV |

| | Ally | 1 | Ally IV | IVII | Ally | Tate | 1 V |
|----------|-------------------------------------|-----------------|-----------------------------|-----------------------|-------------------------|---------------|----------|
| | | | | | | | |
| 7 | Re | egistry Data | Collection Variable | es | | | |
| See | chapt | er for more de | tails on these variables. | | | | |
| | | | | | | | |
| | 1. | Tumor size: | | | | | |
| | 2. | Tumor site: | esophagus | stomach | duodenum | jejunum/ileum | n rectum |
| | | | extraintestinal | | | | |
| | 3. | Tumor mitoti | crate: | | | | |
| | 4. | Tumor ruptur | e: | | | | |
| | 5. | Tumor metas | tasis: liver per | itoneum o | her | | |
| | 6. | Tumor KIT im | munohistochemistry: | | | | |
| | 7. | Tumor mutat | ional status of KIT, PDGFRA | A (if known): | | | |
| | | | | | | | |
| 8 | ш: | stologic Gr | ada (G) | | | | |
| | | stologic Gr | | | | | |
| Gra | aing to | or GIST is depe | ndent on mitotic rate. | | | | |
| | | | | | | | |
| 9 | 9 Lymphovascular Invasion (LVI) | | | | | | |
| | | | | | | | |
| ✓ | Component of Description LVI Coding | | | | | | |
| | 0 | | LVI not present (absent), | not identified | | | |
| | 1 | | LVI present/identified, N | OS | | | |
| | 2 | | Lymphatic and small ves | | | | |
| | 3 | | Venous (large vessel) inv | | | | |
| | 4 | | BOTH lymphatic and sma | III vessel AND venous | (large vessel) invasion | | |
| | 9 | | Presence of LVI unknown | /indeterminate | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| Phy | hysician Signature Date/Time | | | | | | |
| | | | | | | | |
| | | | | | | | |
| Hos | pital I | Name/Address | | | Patient Name/Inforr | nation | |
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|--|----------------|---|
| workup information, until first treatment, including clinical history and symptoms, physical examination endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sentinely | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | Patient Name/Information | |
|-----------------------|--------------------------|--------------------------|--|
| | | | |
| | | | |
| | | | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | Category T Criteria | |
|---------------------------------|--|--|--|
| | TX Primary tumor cannot be assessed | | |
| TO No evidence of primary tumor | | No evidence of primary tumor | |
| | T1 Tumor 5 cm or less in greatest dimension | | |
| | T2 | Tumor more than 5 cm and less than or equal to 10 cm in greatest dimension | |
| | T3 Tumor more than 10 cm and less than or equal to 15 cm in greatest dimension | | |
| | T4 Tumor more than 15 cm in greatest dimension | | |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|------------|--|--|
| NO No regional lymph node metastases or unknown lymph node status | | No regional lymph node metastases or unknown lymph node status | |
| | N1 | Regional lymph node metastasis | |

| ✓ | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| • | M Category | M Criteria |
|---|---|------------|
| | cM0 No distant metastasis | |
| | cM1 Distant metastasis | |
| | pM1 Distant metastasis, microscopically confirmed | |

| Hospital Name/Address | Patient Name/Information |
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| | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of FNCLCC Histologic Grade (G)

The FNCLCC grade is determined by three parameters: differentiation, mitotic activity, and extent of necrosis. Each parameter is scored as follows: differentiation (1–3), mitotic activity (1–3), and necrosis (0–2). The scores are added to determine the grade.

| ✓ | G | G Definition | |
|--|--|--------------|--|
| | GX Grade cannot be assessed | | |
| | G1 Total differentiation, mitotic count and necrosis score of 2 or 3 | | |
| | G2 Total differentiation, mitotic count and necrosis score of 4 or 5 | | |
| G3 Total differentiation, mitotic count and necrosis score of 6, 7, or 8 | | | |

5.1.1 Tumor Differentiation

Tumor differentiation is histology specific (see chapter 39, table 39.1) and is generally scored as follows:

| ✓ | Differentiation Score | Definition | |
|---|--|------------|--|
| | 1 Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) | | |
| | 2 Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) | | |
| | 3 Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, | | |
| | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue | | |

5.1.2 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at 400× magnification = 0.1734 mm²) are assessed using a 40× objective.

| ✓ | Mitotic Count Score | Definition |
|---|---------------------|--------------------------|
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

5.1.3 Tumor Necrosis

Evaluated on gross examination and validated with histologic sections.

| 1 | Necrosis | Definition |
|---------------|----------|---------------------|
| • | Score | |
| 0 No necrosis | | No necrosis |
| | 1 | <50% tumor necrosis |
| | 2 | ≥50% tumor necrosis |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And G is | Then the stage |
|---|------------|----------|----------|----------|----------------|
| | | | | | group is |
| | T1 | N0 | M0 | G1, GX | IA |
| | T2, T3, T4 | N0 | M0 | G1, GX | IB |
| | T1 | N0 | M0 | G2, G3 | II |
| | T2 | N0 | M0 | G2, G3 | IIIA |
| | T3, T4 | N0 | M0 | G2, G3 | IIIB |
| | Any T | N1 | M0 | Any G | IIIB |
| | Any T | Any N | M1 | Any G | IV |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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44. Soft Tissue Sarcoma of the Retroperitoneum

| ٠. | | sardonna or the metroperitories | **** |
|---|----------------------------|---|---------------------------|
| 7 | Registry Da | ta Collection Variables | |
| See | chapter for more o | letails on these variables. | |
| | Bone invasi | on as determined by imaging: | |
| If pM1, source of pathological metastatic specimen: | | | |
| | 3. Additional o | limensions of tumor size: | |
| | 4. FNCLCC gra | de: | |
| 8 | Lymphovas | cular Invasion (LVI) | |
| | | T. | |
| ✓ | Component of LVI Coding | Description | |
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | (learness) in anima |
| | 4 | BOTH lymphatic and small vessel AND venous Presence of LVI unknown/indeterminate | (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate | |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|--|--|---|--|
| workup information, until first treatment, including clinical history and symptoms, p endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of informal diagnostic workup from clinical staging combined with operative findings, and pathology review of resecte specimens | | |
| ycTNM Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvan before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neo therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information | |
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4 AJCC Prognostic Stage Groups

There is no prognostic stage grouping for unusual soft tissue sarcoma histologies. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Bone invasion as determined by imaging:
- 2. If pM1, source of pathological metastatic specimen:
- 3. Additional dimensions of tumor size:
- 4. FNCLCC grade:
- Multifocality and number of sites, when noted:

6 FNCLCC Histologic Grade (G)

The FNCLCC grade is determined by three parameters: differentiation, mitotic activity, and extent of necrosis. Each parameter is scored as follows: differentiation (1–3), mitotic activity (1–3), and necrosis (0–2). The scores are added to determine the grade.

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Total differentiation, mitotic count and necrosis score of 2 or 3 |
| | G2 | Total differentiation, mitotic count and necrosis score of 4 or 5 |
| | G3 | Total differentiation, mitotic count and necrosis score of 6, 7, or 8 |

6.1 Tumor Differentiation

Tumor differentiation is histology specific (see chapter 39, table 39.1) and is generally scored as follows:

| 1 | Differentiation Score | Definition |
|---|---|--|
| | 1 | Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) |
| | 2 Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) | |
| | 3 | Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, |
| | | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue |

6.2 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at 400× magnification = 0.1734 mm²) are assessed using a 40× objective.

| ✓ | Mitotic Count Score | Definition |
|---|---------------------|--------------------------|
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

6.3 Tumor Necrosis

Evaluated on gross examination and validated with histologic sections.

| 1 | Necrosis | Definition |
|---|----------|---------------------|
| Y | Score | |
| | 0 | No necrosis |
| | 1 | <50% tumor necrosis |
| | 2 | ≥50% tumor necrosis |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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7 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed (e.g., curetted) |
| | T0 | No evidence of primary tumor |
| | Tis | In situ primary tumor |
| | T1 | Maximum clinical tumor diameter ≤2 cm |
| | T2 | Maximum clinical tumor diameter >2 but ≤5 cm |
| | T3 | Maximum clinical tumor diameter >5 cm |
| | T4 | Primary tumor invades fascia, muscle, cartilage, or bone |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | N | N Criteria | | | | |
|---|----------|--|--|--|--|--|
| • | Category | | | | | |
| | NX | Regional lymph nodes cannot be clinically | | | | |
| | | assessed (e.g., previously removed for | | | | |
| | | another reason, or because of body habitus) | | | | |
| | N0 | No regional lymph node metastasis detected | | | | |
| | | on clinical and/or radiologic examination | | | | |
| | N1 | Metastasis in regional lymph node(s) | | | | |
| | N2 | In-transit metastasis (discontinuous from | | | | |
| | | primary tumor; located between primary | | | | |
| | | tumor and draining regional nodal basin, or | | | | |
| | | distal to the primary tumor) without lymph | | | | |
| | | node metastasis | | | | |
| | N3 | In-transit metastasis (discontinuous from | | | | |
| | | primary tumor; located between primary | | | | |
| | | tumor and draining regional nodal basin, or | | | | |
| | | distal to the primary tumor) with lymph node | | | | |
| | | metastasis | | | | |

| | N | | | | |
|---|--------|---|--|--|--|
| ✓ | Suffix | Definition | | | |
| | (sn) | Select if regional lymph node metastasis identified | | | |
| | | by SLN biopsy only. | | | |
| | (f) | Select if regional lymph node metastasis identified | | | |
| | | by FNA or core needle biopsy only. | | | |

4.2.2 Pathological N (pN)

| N | N Criteria | |
|----------|--|--|
| Category | | |
| pNX | Regional lymph nodes cannot be assessed | |
| | (e.g., previously removed for another reason | |
| | or not removed for pathological evaluation) | |
| pN0 | No regional lymph node metastasis detected | |
| | on pathological evaluation | |
| pN1 | Metastasis in regional lymph node(s) | |
| pN1a(sn) | Clinically occult regional lymph node | |
| | metastasis identified only by sentinel lymph | |
| | node biopsy | |
| pN1a | Clinically occult regional lymph node | |
| | metastasis following lymph node dissection | |
| pN1b | Clinically and/or radiologically detected | |
| | regional lymph node metastasis | |
| pN2 | In-transit metastasis (discontinuous from | |
| | primary tumor; located between primary | |
| | tumor and draining regional nodal basin, or | |
| | distal to the primary tumor) without lymph | |
| | node metastasis | |
| pN3 | In-transit metastasis (discontinuous from | |
| | primary tumor; located between primary | |
| | tumor and draining regional nodal basin, or | |
| | distal to the primary tumor) with lymph node | |
| | metastasis | |
| | pNX pN0 pN1 pN1a(sn) pN1a pN1a pN1b pN2 | |

| | N | |
|---|--------|---|
| ✓ | Suffix | Definition |
| | (sn) | Select if regional lymph node metastasis identified |
| | | by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified |
| | | by FNA or core needle biopsy only. |

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|--|
| | cM0 | No distant metastasis detected on clinical and/or radiologic examination |
| | cM1 | Distant metastasis detected on clinical and/or radiologic examination |
| | cM1a | Metastasis to distant skin, distant subcutaneous tissue, or distant lymph node(s) |
| | cM1b | Metastasis to lung |
| | cM1c | Metastasis to all other visceral sites |
| | pM1 | Distant metastasis microscopically confirmed |
| | pM1a | Metastasis to distant skin, distant subcutaneous tissue, or distant lymph node(s), microscopically confirmed |
| | pM1b | Metastasis to lung, microscopically confirmed |
| | pM1c | Metastasis to all other distant sites, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Clinical (cTNM)

| | When T | And N is | And M | Then the |
|---|--------|----------|-------|----------|
| ✓ | is | | is | stage |
| | | | | group is |
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | _ |
| | T2-3 | N0 | M0 | IIA |
| | T4 | N0 | M0 | IIB |
| | T0-4 | N1-3 | M0 | Ш |
| | T0-4 | Any N | M1 | IV |

5.2 Pathological (pTNM)

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|--------------|-------------------|-------------|-------------------------|
| | Tis | N0 | M0 | 0 |
| | T1 | N0 | M0 | 1 |
| | T2-3 | N0 | M0 | IIA |
| | T4 | N0 | M0 | IIB |
| | T1-4 | N1a(sn) or N1a | M0 | IIIA |
| | T0 | N1b | M0 | IIIA |
| | T1-4 | N1b-3 | M0 | IIIB |
| | T0-4 | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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| | er for more det | tails on these varia | bles. | | | | |
|----------------------|--|---|---|--------------------------------|------------------|--------------------------|-------------------------|
| 1. | Largest tumoi | r diameter (in milli | meters): | | Птеая | sured clinically | measured histologically |
| | | | | logically or n | | area cililically | |
| 2. | | al status (examined | | | either): | | |
| 3. | Unknown prir | nary status: | ∐ yes | o | | | |
| 4. | Tumor thickne | ess (whole millime | ters): | | | | |
| 5. | Excision marg | in status (tumor b | ase transected or | not transecte | ed): | | |
| 6. | Profound imn | nunosuppression: | no immunos | uppressive co | nditions | HIV/AIDS | |
| | | | solid organ t | ransplant reci | ipient | chronic lymphocy | rtic leukemia |
| | | | non-Hodgkin | lymphoma | | multiple conditio | ns |
| | | | condition NC | | | | |
| 7. | LVI: | nrocont | _ | | annant bu | noth alogist | |
| | | present | absent | | comment by | patriologist | |
| 8. | MCPyV-positi | ve staining by IHC: | : yes n | o ∐not | applicable | | |
| 9. | p63-positive s | staining by IHC (if a | applicable): | es no | | | |
| 10. | Tumor-infiltra | ting lymphocytes | in primary tumor | : not | present | present, nonbris | k present, brisk |
| | | | | pres | sent, NOS | | |
| 11. | Growth patte | rn of primary tume | or: ci | rcumscribed/ | nodular 'nodular | infiltrative | |
| 12. | Extranodal ex | tension in regiona | l lymph node(s): | yes | □no | | |
| 13. | | | | | | ggregate in millimeters) | |
| | | | | | | ggregate in millimeters, | · |
| 14. | 14. Isolated tumor cells in regional lymph node(s): | | | | | | |
| 15. | 15. Eyelid tumor involving the upper or lower eyelid, or both: | | | | | | |
| 16. | Eyelid tumor involving the eyelid margin, defined as the juncture of eyelid skin and tarsal plate at the lash line: yes no | | | | | | |
| | If p | oresent, is the eye | lid margin involve | ment full thic | ckness? | full thickness | not full thickness |
| | | | | | | | |
| | | | | | | | |
| His | stologic di | ade (G) | | | | | |
| | | | ng system at this t | ime | | | |
| | | ade (G) | ng system at this t | ime. | | | |
| e is no | o recommende | ed histologic gradir | · , | ime. | | | |
| e is no | o recommende | | · , | ime. | | | |
| e is no | o recommende | ed histologic gradir | · , | ime. | | | |
| Lyi | o recommende | d histologic gradin | · , | ime. | | | |
| Lyı Com | orecommende mphovascu | ular Invasion Description LVI not present (| (LVI) absent)/not ident | | | | |
| Lyi Com LVI 0 1 | orecommende mphovascu | ular Invasion Description LVI not present (LVI present/iden | (LVI) absent)/not identified, NOS | tified | | | |
| Lyi Com LVI 0 1 2 | orecommende mphovascu | Description LVI not present (LVI present/iden Lymphatic and si | (LVI) absent)/not identified, NOS mall vessel invasion | tified on only (L) | | | |
| Lyi Com LVI 0 1 | orecommende mphovascu | Description LVI not present (LVI present/iden Lymphatic and si Venous (large ve | (LVI) absent)/not identified, NOS | tified on only (L) y (V) | large vessell | invasion | |

9 Anatomy

FIGURE 46.1. Regional lymph nodes for skin sites of the head and neck.

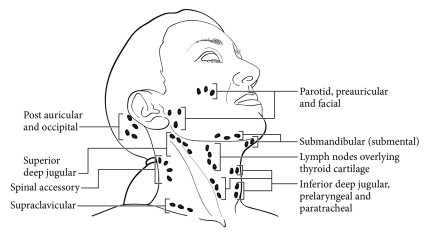


FIGURE 46.7. Merkel cell carcinoma in situ (Tis).

Tis

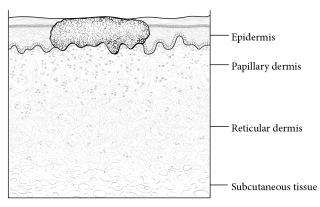
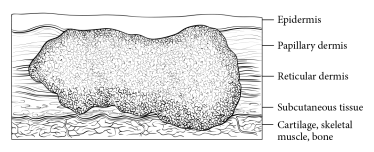


FIGURE 46.11. T4 is defined as a primary tumor invading fascia, muscle, cartilage, or bone.

T4



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|--|--|--|
| | cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imagin endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and othe relevant examinations | | |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgespecimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy arbefore planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | Criteria/Thickness | Criteria/Ulceration Status |
|---|------------|---|----------------------------|
| | TX | Primary tumor thickness cannot be assessed (e.g., diagnosis by curettage) | Not applicable |
| | ТО | No evidence of primary tumor (e.g., unknown primary or completely regressed melanoma) | Not applicable |
| | Tis | Melanoma in situ | Not applicable |
| | T1 | ≤1.0 mm | Unknown or unspecified |
| | T1a | <0.8 mm | Without ulceration |
| | T1b | <0.8 mm | With ulceration |
| | T1b | 0.8–1.0 mm | With or without ulceration |
| | T2 | >1.0–2.0 mm | Unknown or unspecified |
| | T2a | >1.0–2.0 mm | Without ulceration |
| | T2b | >1.0–2.0 mm | With ulceration |
| | T3 | >2.0-4.0 mm | Unknown or unspecified |
| | T3a | >2.0–4.0 mm | Without ulceration |
| | T3b | >2.0–4.0 mm | With ulceration |
| | T4 | >4.0 mm | Unknown or unspecified |
| | T4a | >4.0 mm | Without ulceration |
| | T4b | >4.0 mm | With ulceration |

| ✓ | T Suffix | Definition |
|----------|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

| Hospital Name/Address | Patient Name/Information |
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4.2 Definition of Regional Lymph Node (N)

| | | Extent of regional lymph node and/or lymphatic metastasis | | | |
|---|------------|---|--|--|--|
| ✓ | N Category | Number of tumor-involved regional lymph nodes | Presence of in-transit, satellite, and/or microsatellite metastases | | |
| | NX | Regional nodes not assessed (e.g., SLN biopsy not performed, regional nodes previously removed for another reason) Exception: pathological N category is not required for T1 melanomas, use cN. | No | | |
| | N0 | No regional metastases detected | No | | |
| | N1 | One tumor-involved node or in-transit, satellite, and/or microsatellite metastases with no tumor-involved nodes | One tumor-involved node or in-transit, satellite, and/or microsatellite metastases with no tumor-involved nodes | | |
| | N1a | One clinically occult (i.e., detected by SLN biopsy) | No | | |
| | N1b | One clinically detected | No | | |
| | N1c | No regional lymph node disease | Yes | | |
| | N2 | Two or three tumor-involved nodes or in-transit, satellite, and/or microsatellite metastases with one tumor-involved node | Two or three tumor-involved nodes or in- transit, satellite, and/or microsatellite metastases with one tumor-involved node | | |
| | N2a | Two or three clinically occult (i.e., detected by SLN biopsy) | No | | |
| | N2b | Two or three, at least one of which was clinically detected | No | | |
| | N2c | One clinically occult or clinically detected | Yes | | |
| | N3 | Four or more tumor-involved nodes or in-transit, satellite, and/or microsatellite metastases with two or more tumor-involved nodes, or any number of matted nodes without or with in-transit, satellite, and/or microsatellite metastases | Four or more tumor-involved nodes or intransit, satellite, and/or microsatellite metastases with two or more tumor-involved nodes, or any number of matted nodes without or with in-transit, satellite, and/or microsatellite metastases | | |
| | N3a | Four or more clinically occult (i.e., detected by SLN biopsy) | No | | |
| | N3b | Four or more, at least one of which was clinically detected, or presence of any number of matted nodes | No | | |
| | N3c | Two or more clinically occult or clinically detected and/or presence of any number of matted nodes | Yes | | |

| ~ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| CM0 CM1 CM1a CM1a(0) CM1b CM1b(0) CM1cM1c | Anatomic Site No evidence of distant metastasis Evidence of distant metastasis Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | LDH Level Not applicable Any Not recorded or unspecified Not elevated Elevated Not recorded or unspecified Not recorded or unspecified Not elevated Elevated Elevated |
|--|---|---|
| cM1 cM1a cM1a(0) cM1a(1) cM1b cM1b(0) cM1b(1) | Evidence of distant metastasis Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Any Not recorded or unspecified Not elevated Elevated Not recorded or unspecified Not elevated |
| cM1a cM1a(0) cM1a(1) cM1b cM1b(0) cM1b(1) | Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Not recorded or unspecified Not elevated Elevated Not recorded or unspecified Not elevated |
| cM1a(0) cM1a(1) cM1b cM1b(0) cM1b(1) | node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Not elevated Elevated Not recorded or unspecified Not elevated |
| cM1a(1) cM1b cM1b(0) cM1b(1) | node Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Elevated Not recorded or unspecified Not elevated |
| cM1b cM1b(0) cM1b(1) | node Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Not recorded or unspecified Not elevated |
| cM1b(0) cM1b(1) | Distant metastasis to lung with or without M1a sites of disease Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Not elevated |
| cM1b(1) | Distant metastasis to lung with or without M1a sites of disease Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | |
| ` ′ | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of | Elevated |
| cM1c | | |
| | disease | Not recorded or unspecified |
| cM1c(0) | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of disease | Not elevated |
| cM1c(1) | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of disease | Elevated |
| cM1d | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease | Not recorded or unspecified |
| cM1d(0) | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease | Not elevated |
| cM1d(1) | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease | Elevated |
| pM1 | Evidence of distant metastasis, microscopically proven | Any |
| pM1a | Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node, microscopically proven | Not recorded or unspecified |
| pM1a(0) | Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node, microscopically proven | Not elevated |
| pM1a(1) | Distant metastasis to skin, soft tissue including muscle, and/or nonregional lymph node, microscopically proven | Elevated |
| pM1b | Distant metastasis to lung with or without M1a sites of disease, microscopically proven | Not recorded or unspecified |
| pM1b(0) | Distant metastasis to lung with or without M1a sites of disease, microscopically proven | Not elevated |
| pM1b(1) | Distant metastasis to lung with or without M1a sites of disease, microscopically proven | Elevated |
| pM1c | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of disease, microscopically proven | Not recorded or unspecified |
| pM1c(0) | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of disease, microscopically proven | Not elevated |
| pM1c(1) | Distant metastasis to non-CNS visceral sites with or without M1a or M1b sites of disease, microscopically proven | Elevated |
| pM1d | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease, microscopically proven | Not recorded or unspecified |
| pM1d(0) | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease, microscopically proven | Not elevated |
| pM1d(1) | Distant metastasis to CNS with or without M1a, M1b, or M1c sites of disease, microscopically proven | Elevated |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Clinical (cTNM)

Clinical staging includes microstaging of the primary melanoma and clinical/radiologic/biopsy evaluation for metastases. By convention, clinical staging should be used after biopsy of the primary melanoma, with clinical assessment for regional and distant metastases. Note that pathological assessment of the primary melanoma is used for both clinical and pathological classification. Diagnostic biopsies to evaluate possible regional and/or distant metastasis also are included. Note there is only one stage group for clinical Stage III melanoma.

| 1 | When T is | And N is | And M is | Then the stage |
|---|------------|----------|----------|----------------|
| • | | | | group is |
| | Tis | N0 | M0 | 0 |
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2a | N0 | M0 | IB |
| | T2b | N0 | M0 | IIA |
| | T3a | N0 | M0 | IIA |
| | T3b | N0 | M0 | IIB |
| | T4a | N0 | M0 | IIB |
| | T4b | N0 | M0 | IIC |
| | Any T, Tis | ≥N1 | M0 | III |
| | Any T | Any N | M1 | IV |

5.2 Pathological (pTNM)

Pathological staging includes microstaging of the primary melanoma, including any additional staging information from the wide-excision (surgical) specimen that constitutes primary tumor surgical treatment and pathological information about the regional lymph nodes after SLN biopsy or therapeutic lymph node dissection for clinically evident regional lymph node disease.

| ✓ When T is | And N is | And M is | Then the stage group is |
|-------------------|------------------|----------|-------------------------|
| Tis | N0 | M0 | 0 |
| T1a | N0 | M0 | IA |
| T1b | N0 | M0 | IA |
| T2a | N0 | M0 | IB |
| T2b | N0 | M0 | IIA |
| T3a | N0 | M0 | IIA |
| T3b | N0 | MO | IIB |
| T4a | N0 | MO | IIB |
| T4b | N0 | M0 | IIC |
| ТО | N1b, N1c | MO | IIIB |
| TO | N2b/c, N3b/c | M0 | IIIC |
| T1a/b, T2a | N1a, N2a | MO | IIIA |
| T1a/b, T2a | N1b/c, N2b | M0 | IIIB |
| T2b, T3a | N1a/b/c, N2a/b | M0 | IIIB |
| T1a/b, T2a/b, T3a | N2c, N3a/b/c | MO | IIIC |
| T3b, T4a | Any N≥N1 | MO | IIIC |
| T4b | N1a/b/c, N2a/b/c | MO | IIIC |
| T4b | N3a/b/c | MO | IIID |
| Any T, Tis | Any N | M1 | IV |

Pathological Stage 0 (melanoma in situ) and T1 do not require pathological evaluation of lymph nodes to complete pathological staging; use cN information to assign their pathological stage.

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Breslow tumor thickness (xx.x mm)
- 2. Primary tumor ulceration (yes/no)
- 3. Mitotic rate (whole number per square millimeter [mm²])
- 4. Microsatellites (pathologically detected satellites, not clinically apparent) (yes/no)
- 5. Tumor-infiltrating lymphocytes (absent, nonbrisk, or brisk)
- 6. Clark level of invasion (I–V)
- 7. Regression (yes/no)
- 8. Neurotropism (present or absent)
- 9. Lymphovascular invasion (present or absent)
- 10. In-transit and/or satellite metastasis (in-transit, satellite, both)
- 11. Regional lymph node clinically or radiologically detected (yes/no)
- 12. Microscopic confirmation of tumor metastasis in any regional lymph node that was clinically or radiologically detected (yes/no)
- 13. SLN biopsy performed (yes/no)
- 14. Number of nodes examined from sentinel node procedure (whole number)
- 15. Number of tumor-involved nodes from sentinel node procedure (whole number)
- 16. Sentinel node tumor burden (largest dimension of largest discrete deposit in xx.x mm)
- 17. ENE in any tumor-involved regional lymph node (sentinel or clinically detected) (present or absent)
- 18. Completion or therapeutic lymph node dissection performed (yes/no)
- 19. Number of lymph nodes examined from completion or therapeutic lymph node dissection (whole number)
- 20. Number of lymph nodes involved with tumor from completion or therapeutic lymph node dissection (whole number)
- 21. Matted nodes (yes/no)
- 22. Distant metastasis to skin, soft tissue, or distant nodes (yes/no)
- 23. Distant metastasis to lung (yes/no)
- 24. Distant metastasis to non-CNS viscera (yes/no)
- 25. Distant metastasis to CNS (yes/no)
- 26. Serum LDH level (xx,xxx U/L) and serum LDH level upper limit of normal from laboratory reference range (Note serum LDH recorded for Stage IV only)

7 Histologic Grade (G)

There is no recommended histologic grading system at this time.

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 47.1. T1a melanoma. T1a is defined as invasive melanoma <0.8 mm in thickness without ulceration. Tumor thickness is measured from the top of the granular layer of the epidermis to the deepest invasive cell across the broad base of the tumor.

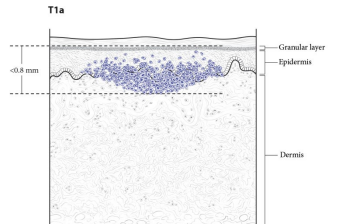
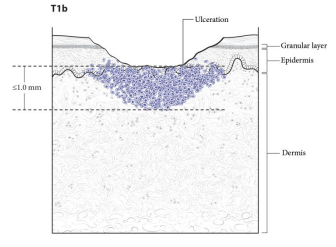


FIGURE 47.2. T1b melanoma. T1b is defined as melanoma 0.8 to 1 mm in thickness regardless of ulceration status OR ulcerated melanoma <0.8 mm in thickness. Tumor thickness is measured from the top of the granular layer of the epidermis (or, if the surface overlying the entire dermal component is ulcerated, from the base of the ulcer) to the deepest invasive cell across the broad base of the tumor.



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FIGURE 47.3. T2a melanoma. T2a is defined as invasive melanoma >1.0 to 2.0 mm in thickness without ulceration. Tumor thickness is measured from the top of the granular layer of the epidermis to the deepest invasive cell across the broad base of the tumor.

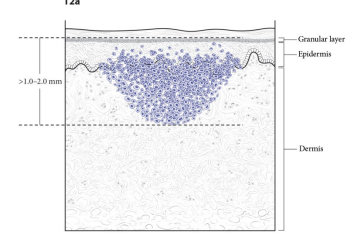


FIGURE 47.4. T2b melanoma. T2b is defined as ulcerated melanoma >1.0 to 2.0 mm in thickness. Tumor thickness is measured from the base of the ulcer to the deepest invasive cell across the broad base of the tumor.

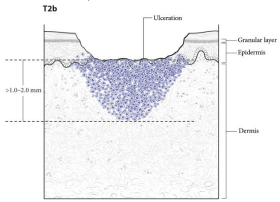
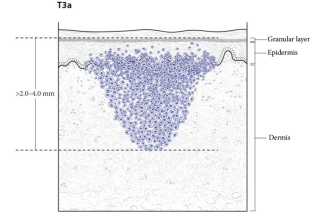


FIGURE 47.5. T3a melanoma. T3a is defined as invasive melanoma >2.0 to 4.0 mm in thickness without ulceration. Tumor thickness is measured from the top of the granular layer of the epidermis to the deepest invasive cell across the broad base of the tumor.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 47.6. T3b melanoma. T3b is defined as ulcerated melanoma >2.0 to 4.0 mm in thickness. Tumor thickness is measured from the base of the ulcer to the deepest invasive cell across the broad base of the tumor.

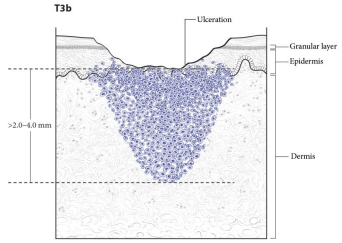
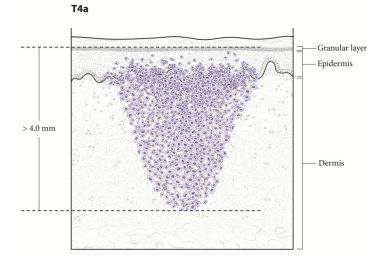
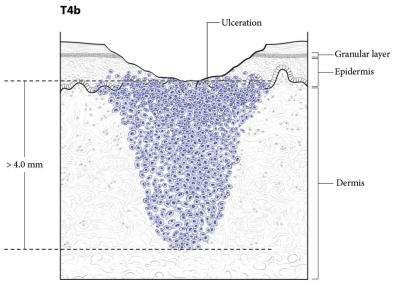


FIGURE 47.7. T4a melanoma. T4a is defined as invasive melanoma >4.0 mm in thickness without ulceration. Tumor thickness is measured from the top of the granular layer of the epidermis to the deepest invasive cell across the broad base of the tumor.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 47.8. T4b melanoma. T4b is defined as ulcerated melanoma >4.0 mm in thickness. Tumor thickness is measured from the base of the ulcer to the deepest invasive cell across the broad base of the tumor.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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1 Terms of Use

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic | |
| | | workup information, until first treatment, including clinical history and symptoms, physical examination, imaging | |
| | | endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of | |
| | | regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other | |
| | | relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from | |
| | | diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical | |
| | | specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy an | |
| | | before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant | |
| | | therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until | |
| | | treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, | |
| | | and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a | |
| | | previously diagnosed cancer). | |

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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|-----|-----------------------|---|
| | TX | Primary tumor cannot be assessed |
| | Т0 | No evidence of primary tumor |
| | Tis(DCIS)* | Ductal carcinoma in situ |
| | Tis(Paget) | Paget disease of the nipple NOT associated with invasive carcinoma and/or carcinoma in situ (DCIS) in the underlying breast parenchyma. Carcinomas in the breast parenchyma associated with Paget disease are categorized based on the size and characteristics of the parenchymal disease, although the presence of Paget disease should still be noted. |
| | T1 | Tumor ≤ 20 mm in greatest dimension |
| | T1mi | Tumor ≤ 1 mm in greatest dimension |
| | T1a | Tumor > 1 mm but \leq 5 mm in greatest dimension (round any measurement >1.0–1.9 mm to 2 mm). |
| | T1b | Tumor > 5 mm but ≤ 10 mm in greatest dimension |
| | T1c | Tumor > 10 mm but ≤ 20 mm in greatest dimension |
| | T2 | Tumor > 20 mm but ≤ 50 mm in greatest dimension |
| | T3 | Tumor > 50 mm in greatest dimension |
| | T4 | Tumor of any size with direct extension to the chest wall and/or to the skin (ulceration or macroscopic nodules); invasion of the dermis alone does not qualify as T4 |
| | T4a | Extension to the chest wall; invasion or adherence to pectoralis muscle in the absence of invasion of chest wall structures does not qualify as T4 |
| | T4b | Ulceration and/or ipsilateral macroscopic satellite nodules and/or edema (including peau d'orange) of the skin that does not meet the criteria for inflammatory carcinoma |
| | T4c | Both T4a and T4b are present |
| | T4d | Inflammatory carcinoma (see "Rules for Classification") |
| * N | ote: Lobular carcinon | na <i>in situ</i> (LCIS) is a benign entity and is removed from TNM staging in the AJCC Cancer Staging Manual, 8 th Edition. |

| ~ | T Suffix | Definition |
|----------|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria | |
|---|------------|--|--|
| | cNX* | Regional lymph nodes cannot be assessed (e.g., previously removed) | |
| | cN0 | No regional lymph node metastases (by imaging or clinical examination) | |
| | cN1 | Metastases to movable ipsilateral Level I, II axillary lymph node(s) | |
| | cN1mi** | Micrometastases (approximately 200 cells, larger than 0.2 mm, but none larger than 2.0 mm) | |
| | cN2 | Metastases in ipsilateral Level I, II axillary lymph nodes that are clinically fixed or matted; | |
| | | or in ipsilateral internal mammary nodes in the absence of axillary lymph node metastases | |
| | cN2a | Metastases in ipsilateral Level I, II axillary lymph nodes fixed to one another (matted) or to other structures | |
| | cN2b | Metastases only in ipsilateral internal mammary nodes in the absence of axillary lymph node metastases | |
| | cN3 | Metastases in ipsilateral infraclavicular (Level III axillary) lymph node(s) with or without Level I, II axillary lymph node | |
| | | involvement; | |
| | | or in ipsilateral internal mammary lymph node(s) with Level I, II axillary lymph node metastases; | |
| | | or metastases in ipsilateral supraclavicular lymph node(s) with or without axillary or internal mammary lymph node | |
| | | involvement | |
| | cN3a | Metastases in ipsilateral infraclavicular lymph node(s) | |
| | cN3b | Metastases in ipsilateral internal mammary lymph node(s) and axillary lymph node(s) | |
| | cN3c | Metastases in ipsilateral supraclavicular lymph node(s) | |

Note: (sn) and (f) suffixes should be added to the N category to denote confirmation of metastasis by sentinel node biopsy or fine needle aspiration/core needle biopsy respectively

^{**} cN1mi is rarely used but may be appropriate in cases where sentinel node biopsy is performed before tumor resection, most likely to occur in cases treated with neoadjuvant therapy.

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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^{*} The cNX category is used sparingly in cases where regional lymph nodes have previously been surgically removed or where there is no documentation of physical examination of the axilla.

4.2.2 Pathological N (pN)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | pNX | Regional lymph nodes cannot be assessed (e.g., not removed for pathological study or previously removed) |
| | pN0 | No regional lymph node metastasis identified or ITCs only |
| | pN0(i+) | ITCs only (malignant cell clusters no larger than 0.2 mm) in regional lymph node(s) |
| | pN0(mol+) | Positive molecular findings by reverse transcriptase polymerase chain reaction (RT-PCR); no ITCs detected |
| | pN1 | Micrometastases; or metastases in 1–3 axillary lymph nodes; and/or clinically negative internal mammary nodes |
| | | with micrometastases or macrometastases by sentinel lymph node biopsy |
| | pN1mi | Micrometastases (approximately 200 cells, larger than 0.2 mm, but none larger than 2.0 mm) |
| | pN1a | Metastases in 1–3 axillary lymph nodes, at least one metastasis larger than 2.0 mm |
| | pN1b | Metastases in ipsilateral internal mammary sentinel nodes, excluding ITCs |
| | pN1c | pN1a and pN1b combined |
| | pN2 | Metastases in 4–9 axillary lymph nodes; or positive ipsilateral internal mammary lymph nodes by imaging in the |
| | | absence of axillary lymph node metastases |
| | pN2a | Metastases in 4–9 axillary lymph nodes (at least one tumor deposit larger than 2.0 mm) |
| | pN2b | Metastases in clinically detected internal mammary lymph nodes with or without microscopic confirmation; with |
| | | pathologically negative axillary nodes |
| | pN3 | Metastases in 10 or more axillary lymph nodes; |
| | | or in infraclavicular (Level III axillary) lymph nodes; |
| | | or positive ipsilateral internal mammary lymph nodes by imaging in the presence of one or more positive Level I, II |
| | | axillary lymph nodes; |
| | | or in more than three axillary lymph nodes and micrometastases or macrometastases by sentinel lymph node biops |
| | | in clinically negative ipsilateral internal mammary lymph nodes; |
| | | or in ipsilateral supraclavicular lymph nodes |
| | pN3a | Metastases in 10 or more axillary lymph nodes (at least one tumor deposit larger than 2.0 mm); |
| | | or metastases to the infraclavicular (Level III axillary lymph) nodes |
| | pN3b | pN1a or pN2a in the presence of cN2b (positive internal mammary nodes by imaging); |
| | | or pN2a in the presence of pN1b |
| _ | pN3c | Metastases in ipsilateral supraclavicular lymph nodes |

biopsy respectively, with NO further resection of nodes.

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|--|---|---|
| | cM0 | No clinical or radiographic evidence of distant metastases* |
| | cM0(i+) | No clinical or radiographic evidence of distant metastases in the presence of tumor cells or deposits no larger than 0.2 mm detected microscopically or by molecular techniques in circulating blood, bone marrow, or other nonregional nodal tissue in a patient without symptoms or signs of metastases |
| cM1 Distant metastases detected by clinical and radiographic means | | Distant metastases detected by clinical and radiographic means |
| | pM1 | Any histologically proven metastases in distant organs; or if in non-regional nodes, metastases greater than 0.2 mm |
| * N | * Note that imaging studies are not required to assign the cM0 category | |

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5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Histologic Grade (G)

5.1.1 Invasive Carcinoma

All invasive breast carcinomas should be assigned a histologic grade. The Nottingham combined histologic grade (Nottingham modification of the SBR grading system) is recommended and is stipulated for use by the College of American Pathologists (see www.cap.org). ¹⁻³ The grade for a tumor is determined by assessing morphologic features (tubule formation, nuclear pleomorphism, and calibrated mitotic count), assigning a value from 1 (favorable) to 3 (unfavorable) for each feature, and totaling the scores for all three categories. A combined score of 3–5 points is designated as grade 1; a combined score of 6–7 points is grade 2; a combined score of 8–9 points is grade 3. The use of subjective grading alone is discouraged.

| ✓ | G | G Definition |
|---|----|--|
| | GX | Grade cannot be assessed |
| | G1 | Low combined histologic grade (favorable), SBR score of 3–5 points |
| | G2 | Intermediate combined histologic grade (moderately favorable); SBR score of 6–7 points |
| | G3 | High combined histologic grade (unfavorable); SBR score of 8–9 points |

5.1.2 Carcinoma in situ

The grade that should be used for in situ carcinomas is nuclear grade (see www.cap.org).

| ✓ | G | G Definition |
|---|----|----------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Low nuclear grade |
| | G2 | Intermediate nuclear grade |
| | G3 | High nuclear grade |

5.2 Definition of HER2 Status

The measurement of Human Epidermal Growth Factor Receptor-2 (HER2) is primarily by either IHC to assess expression of the HER2 protein or by *in situ* hybridization (ISH) - most commonly by fluorescent labeled probes (FISH) or chromogenic labeled probes (CISH) to assess gene copy number.

| ✓ | HER2 Status | | |
|---|---|--|--|
| | Positive | | |
| | Negative | | |
| | Equivocal (use negative category for prognostic stage group assignment) | | |

5.3 Definition of ER Status

Estrogen receptor (ER) expression is measured primarily by IHC. Any staining of 1% of cells or more is considered positive for both ER and PR.

| ✓ | ER Status |
|---|-----------|
| | Positive |
| | Negative |

5.4 Definition of PR status

Progesterone receptor (PR) expression is measured primarily by IHC. Any staining of 1% of cells or more is considered positive for both ER and PR.

| | • • • | | | | |
|---|-------------|--|--|--|--|
| ✓ | ✓ PR Status | | | | |
| | Positive | | | | |
| | Negative | | | | |

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6 Additional Factors Recommended for Clinical Care

6.1 Definition of Oncotype Dx® Recurrence Score

Oncotype Dx® is a genomic test based on the assessment of 21 genes; the result is the outcome of a mathematical formula of the weighted expression of each gene combined into a single score. It is measured and reported by RT-PCR, with recurrence score of < 11 the most pertinent cutoff value. Oncotype Dx® is required only for assigning prognostic stage group to patients with T1–2 N0 M0, ER-positive, HER2-negative cancers. If OncotypeDx® is not performed, not available, or if the OncotypeDx® score is 11 or greater for patients with T1-2 N0 M0 HER2 negative ER positive cancer, then the Prognostic Stage Group is assigned based on the remaining anatomic and biomarker categories. OncotypeDx® is the only multigene panel included to classify Prognostic Stage because prospective Level I data supports this use for patients with a score <11. Future updates may include results from other multigene panels to assign cohorts of patients to prognostic stage groups when there are high level data to support these assignments.

| ✓ | Oncotype Dx® Recurrence Score |
|---|-------------------------------|
| | Less than 11 |
| | 11 or greater |
| | Not performed |
| | Not available |

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7 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

7.1 Clinical Prognostic Stage

Clinical Prognostic Stage applies to ALL patients with breast cancer for clinical classification and staging. It uses clinical tumor (T), node (N) and metastases (M) information based on history, physical examination, any imaging performed (not necessary for clinical staging) and relevant biopsies. Genomic profile information is not included in Clinical Prognostic Stage as pathologic information from surgery is necessary to ascertain the prognosis using these tools.

| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Clinical Prognostic Stage Group is | 1 |
|---------------------------|--------------|-----------------------|------------------|------------------|---|----|
| Tis N0 M0 | Any | Any | Any | Any | 0 | |
| | | | Basili a | Positive | IA | |
| | | Basili a | Positive | Negative | IA | |
| | | Positive | Namativa | Positive | IA | |
| | 61 | | Negative | Negative | IA | |
| G1 | G1 | | Basili a | Positive | IA | |
| | | Nagativa | Positive | Negative | IA | |
| | Negative | Nevel | Positive | IA | | |
| | | | Negative | Negative | IB | |
| | | Positive | Positive | Positive | IA | |
| | | | | Negative | IA | |
| | | | | Nevella | Positive | IA |
| T1* N0 M0 | G2 | | Negative | Negative | IA | |
| T0 N1mi M0 T1* N1mi M0 | G2 | Negative | Positive | Positive | IA | |
| | | | | Negative | IA | |
| | | | Negative | Positive | IA | |
| | | | | Negative | IB | |
| | | | Danition | Positive | IA | |
| | | Positive | Positive | Negative | IA | |
| | | Positive | Namativa | Positive | IA | |
| | 63 | | Negative | Negative | IA | |
| | G3 | | Positive | Positive | IA | |
| | | Negative | Positive | Negative | IB | |
| | | Negative | Negative | Positive | IB | |
| | | Negative | Negative | IB | | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Clinical Prognostic Stage Group is | ✓ |
|---------------------------|--------------|--------------------|---------------------|---------------------|---|----------|
| | | | | Positive | IB | |
| | | | Positive | Negative | IIA | |
| | | Positive | | Positive | IIA | |
| | | | Negative | Negative | IIA | |
| | G1 | | | Positive | IB | |
| | | | Positive | Negative | IIA | |
| | | Negative | | Positive | IIA | |
| | | | Negative | Negative | IIA | |
| | | Positive | Positive | Positive | IB | |
| | | | | Negative | IIA | |
| | | | Negative | Positive | IIA | |
| T0 N1** M0 T1* N1** M0 | G2 | | | Negative | IIA | |
| T2 N0 M0 | G2 | | Positive | Positive | IB | |
| | | Negative | Positive | Negative | IIA | |
| | | | Negative | Positive | IIA | |
| | | | Negative | Negative | IIB | |
| | | | Positive | Positive | IB | |
| | | Positive | Positive | Negative | IIA | |
| | | Positive | Negative | Positive | IIA | |
| G3 | | Negative | Negative | IIA | | |
| | 43 | | Positive | Positive | IIA | |
| | | Negative | Positive | Negative | IIB | |
| | | ivegative | Negative | Positive | IIB | |
| | | | ivegative | Negative | IIB | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Clinical Prognostic Stage Group is | ✓ |
|-------------|--------------|--------------------|---------------------|---------------------|---|----------|
| | | | | Positive | IB | |
| | | Don't c | Positive | Negative | IIA | |
| | | Positive | No. at the | Positive | IIA | |
| | G1 | | Negative | Negative | IIB | |
| | GI | | Daniki | Positive | IIA | |
| | | Nagativa | Positive | Negative | IIB | |
| | | Negative | Negative | Positive | IIB | |
| | | | Negative | Negative | IIB | |
| | | Positive | Positive | Positive | IB | |
| | | | Positive | Negative | IIA | |
| | | | Negative | Positive | IIA | |
| T2 N1*** M0 | G2 | | Negative | Negative | IIB | |
| T3 N0 M0 | G2 | Negative | Positive | Positive | IIA | |
| | | | Positive | Negative | IIB | |
| | | | Negative | Positive | IIB | |
| | | | Negative | Negative | IIIB | |
| | | | Positive | Positive | IB | |
| | | Positive | Positive | Negative | IIB | |
| | | Positive | Negative | Positive | IIB | |
| G3 | | Negative | Negative | IIB | | |
| | d5 | | Positive | Positive | IIB | |
| | | Nogativo | rositive | Negative | IIIA | |
| | | Negative - | Negative | Positive | IIIA | |
| | | | ivegative | Negative | IIIB | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Clinical Prognostic Stage Group is | ✓ |
|-----------------------|--------------|--------------------|---------------------|---------------------|---|----------|
| | | | | Positive | IIA | |
| | | | Positive | Negative | IIIA | |
| | | Positive | | Positive | IIIA | |
| | 64 | | Negative | Negative | IIIA | |
| | G1 | | Desitive | Positive | IIA | |
| | | Nagativa | Positive | Negative | IIIA | |
| | | Negative | Negative | Positive | IIIA | |
| | | | Negative | Negative | IIIB | |
| | | Positive | Positive | Positive | IIA | |
| | | | | Negative | IIIA | |
| T0 N2 M0 | | | Negative | Positive | IIIA | |
| T1* N2 M0 T2 N2 M0 | G2 | | Negative | Negative | IIIA | |
| T3 N1*** M0 | G2 | Negative | Positive | Positive | IIA | |
| T3 N2 M0 | | | | Negative | IIIA | |
| | | | Negative | Positive | IIIA | |
| | | | Negative | Negative | IIIB | |
| | | | Positive | Positive | IIB | |
| | | Positive | Positive | Negative | IIIA | |
| | | Positive | Nogativo | Positive | IIIA | |
| | G3 | | Negative | Negative | IIIA | |
| | ds | | Positive | Positive | IIIA | |
| | | Negative | rositive | Negative | IIIB | |
| | | ivegative | Negative | Positive | IIIB | |
| | | | ivegative | Negative | IIIC | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Clinical Prognostic Stage Group is | ~ |
|-------------------------|--------------|-----------------------|---------------------|---------------------|---|---|
| | | | | Positive | IIIA | |
| | | | Positive | Negative | IIIB | |
| | | Positive | | Positive | IIIB | |
| | | | Negative | Negative | IIIB | |
| | G1 | | | Positive | IIIB | |
| | | | Positive | Negative | IIIB | |
| | | Negative | | Positive | IIIB | |
| | | | Negative | Negative | IIIC | |
| | | | | Positive | IIIA | |
| | | Positive | Positive | Negative | IIIB | |
| | | | Negative | Positive | IIIB | |
| T4 N0 M0 T4 N1*** M0 | 62 | | | Negative | IIIB | |
| T4 N2 M0 | G2 | G2 Negative | Positive | Positive | IIIB | |
| Any T N3 M0 | | | | Negative | IIIB | |
| | | | Nogativo | Positive | IIIB | |
| | | | Negative | Negative | IIIC | |
| | | | Positive | Positive | IIIB | |
| | | Positive | | Negative | IIIB | |
| | | Positive | Negative | Positive | IIIB | |
| | G3 | | Negative | Negative | IIIB | |
| 63 | | Positive | Positive | IIIB | | |
| | | Negative | Positive | Negative | IIIC | |
| | | ivegative | Negative | Positive | IIIC | |
| | | | ivegative | Negative | IIIC | |
| Any T Any N M1 | Any | Any | Any | Any | IV | |

^{*} T1 Includes T1mi

- Because N1mi categorization requires evaluation of the entire node, and cannot be assigned on the basis of an FNA or core biopsy, N1mi can only be used with Clinical Prognostic Staging when clinical staging is based on a resected lymph node in the absence of resection of the primary cancer, such as the situation where sentinel node biopsy is performed prior to receipt of neoadjuvant chemotherapy or endocrine therapy.
- 2. For cases where HER2 is determined to be "equivocal" by ISH (FISH or CISH) testing under the 2013 ASCO/CAP HER2 testing guidelines, the HER2 "negative" category should be used for staging in the Clinical Prognostic Stage Group table. 4,5
- 3. The prognostic value of these Prognostic Stage Groups is based on populations of persons with breast cancer that have been offered and mostly treated with appropriate endocrine and/or systemic chemotherapy (including anti-HER2 therapy).

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^{**} N1 does not include N1mi. T1 N1mi M0 and T0 N1mi M0 cancers are included for prognostic staging with T1 N0 M0 cancers of the same prognostic factor status.

^{***} N1 includes N1mi. T2, T3, and T4 cancers and N1mi are included for prognostic staging with T2 N1, T3 N1 and T4 N1, respectively . Notes:

7.2 Pathological Prognostic Stage

Pathological Prognostic Stage applies to patients with breast cancer treated with surgery as the initial treatment. It includes all information used for clinical staging plus findings at surgery and pathological findings from surgical resection. Pathological Prognostic Stage does not apply to patients treated with systemic or radiation prior to surgical resection (neoadjuvant therapy).

| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | ~ |
|-------------------------|--------------|--------------------|---------------------|------------------|--|----------|
| Tis N0 M0 | Any | Any | Any | Any | 0 | |
| | | | Davili a | Positive | IA | |
| | | Danitiva | Positive | Negative | IA | |
| | | Positive | Nasativa | Positive | IA | |
| | G1 | | Negative | Negative | IA | |
| GI | GI | | Positive | Positive | IA | |
| | | Namativa | Positive | Negative | IA | |
| | Negative | Negative | Positive | IA | | |
| | | | Negative | Negative | IA | |
| | | Positive | Positive | Positive | IA | |
| | | | | Negative | IA | |
| | | | | Positive | IA | |
| T1* N0 M0 T0 N1mi M0 | G2 | | Negative | Negative | IA | |
| T1* N1mi M0 | G2 | | Positive | Positive | IA | |
| | | Nanativa | Positive | Negative | IA | |
| | | Negative | Nanativa | Positive | IA | |
| | | | Negative | Negative | IB | |
| | | | Dooiting | Positive | IA | |
| | | Danitiva | Positive | Negative | IA | |
| | | Positive | Nasativa | Positive | IA | |
| C | 63 | | Negative | Negative | IA | |
| | G3 | | Desitive | Positive | IA | |
| | | Nogativo | Positive | Negative | IA | |
| | | Negative | Nogativa | Positive | IA | |
| | | | Negative | Negative | IB | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | ✓ |
|---------------------------|--------------|--------------------|---------------------|------------------|--|----------|
| | | | Positive | Positive | IA | |
| | | Docitivo | Positive | Negative | IB | |
| | | Positive | Negative | Positive | IB | |
| | G1 | | Negative | Negative | IIA | |
| | GI | | Positive | Positive | IA | |
| | | Negative | Positive | Negative | IB | |
| | | Negative | Negative | Positive | IB | |
| | | | Negative | Negative | IIA | |
| | | | Positive | Positive | IA | |
| | G2 | Positive | FOSITIVE | Negative | IB | |
| | | | Negative | Positive | IB | |
| T0 N1** M0 T1* N1** M0 | | | | Negative | IIA | |
| T2 N0 M0 | | Negative | Positive | Positive | IA | |
| | | | | Negative | IIA | |
| | | | Negative | Positive | IIA | |
| | | | Negative | Negative | IIA | |
| | | | Positive | Positive | IA | |
| | | Positive | Positive | Negative | IIA | |
| | | Positive | Negative | Positive | IIA | |
| | G3 | | ivegative | Negative | IIA | |
| | G5 | | Positive | Positive | IB | |
| | | Negative | Positive | Negative | IIA | |
| | | ivegative | Negative | Positive | IIA | |
| | | | ivegative | Negative | IIA | |

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| Hospital Name/Address | Patient Name/Information |
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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | √ |
|-------------|--------------|--------------------|------------------|------------------|--|----------|
| | | | Positive | Positive | IA | |
| | | Positive | Positive | Negative | IIB | |
| | | Positive | Negative | Positive | IIB | |
| | G1 | | Negative | Negative | IIB | |
| | GI | | Positive | Positive | IA | |
| | | Negative | Positive | Negative | IIB | |
| | | Negative | Negative | Positive | IIB | |
| | | | Negative | Negative | IIB | |
| | | Positive | Positive | Positive | IB | |
| | G2 | | Positive | Negative | IIB | |
| | | | Negative | Positive | IIB | |
| T2 N1*** M0 | | | Negative | Negative | IIB | |
| T3 N0 M0 | | Negative | Positive | Positive | IB | |
| | | | | Negative | IIB | |
| | | | Negative | Positive | IIB | |
| | | | Negative | Negative | IIB | |
| | | | Positive | Positive | IB | |
| | | Positive | Positive | Negative | IIB | |
| | | Positive | Negative | Positive | IIB | |
| | G3 | | Negative | Negative | IIB | |
| | 3 | | Positive | Positive | IIA | |
| | | Negative | rositive | Negative | IIB | |
| | | ivegative | Nogative | Positive | IIB | |
| | | | Negative | Negative | IIIA | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | 1 |
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| | | | Do sitii ve | Positive | IB | |
| | | Positive | Positive | Negative | IIIA | |
| | | Positive | Namakiya | Positive | IIIA | |
| | G1 | | Negative | Negative | IIIA | |
| | GI | | Positive | Positive | IB | |
| | | Nanativa | Positive | Negative | IIIA | |
| | | Negative | Negative | Positive | IIIA | |
| | | | Negative | Negative | IIIA | |
| | | | Positive | Positive | IB | |
| | G2 | Positive | Positive | Negative | IIIA | |
| T0 N2 M0 | | | Negative | Positive | IIIA | |
| T1* N2 M0 T2 N2 M0 | | | | Negative | IIIA | |
| T3 N1*** M0 | | Negative | Positive | Positive | IB | |
| T3 N2 M0 | | | | Negative | IIIA | |
| | | | Negative | Positive | IIIA | |
| | | | | Negative | IIIB | |
| | | | Positive | Positive | IIA | |
| | | Danitiva | Positive | Negative | IIIA | |
| | | Positive | Namakiya | Positive | IIIA | |
| | G3 | | Negative | Negative | IIIA | |
| | GS | | Positive | Positive | IIB | |
| | | Nogotivo | Positive | Negative | IIIA | |
| | | Negative | Negative | Positive | IIIA | |
| | | | | Negative | IIIC | |

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| When TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | 1 |
|-------------------------|--------------|--------------------|------------------|------------------|--|---|
| | | | 0 | Positive | IIIA | |
| | | De alli | Positive | Negative | IIIB | |
| | | Positive | Nagativa | Positive | IIIB | |
| | G1 | | Negative | Negative | IIIB | |
| | GI | | Positive | Positive | IIIA | |
| | | Negative | Positive | Negative | IIIB | |
| | | Negative | Negative | Positive | IIIB | |
| | | | Negative | Negative | IIIB | |
| | | Positive | Positive | Positive | IIIA | |
| | | | Positive | Negative | IIIB | |
| T4 NO N40 | | | Negative | Positive | IIIB | |
| T4 N0 M0 T4 N1*** M0 | G2 | | | Negative | IIIB | |
| T4 N2 M0 | G2 | Negative | Positive | Positive | IIIA | |
| Any T N3 M0 | | | | Negative | IIIB | |
| | | | Negative | Positive | IIIB | |
| | | | Negative | Negative | IIIC | |
| | | | Positive | Positive | IIIB | |
| | | Donitivo | Positive | Negative | IIIB | |
| | | Positive | Negative | Positive | IIIB | |
| | G3 | | Negative | Negative | IIIB | |
| | 63 | | Da sitii | Positive | IIIB | |
| | | Negative | Positive | Negative | IIIC | |
| | | ivegative | Negative | Positive | IIIC | |
| | | | Negative | Negative | IIIC | |
| Any T Any N M1 | Any | Any | Any | Any | IV | |

^{*}T1 includes T1mi.

- 1. For cases where HER2 is determined to be "equivocal" by ISH (FISH or CISH) testing under the 2013 ASCO/CAP HER2 testing guidelines, HER2 "negative" category should be used for staging in the Pathological Prognostic Stage Group Table. 4,5
- The prognostic value of these Prognostic Stage Groups is based on populations of persons with breast cancer that have been offered and mostly treated with appropriate endocrine and/or systemic chemotherapy (including anti-HER2 therapy).

| Hospital Name/Address | Patient Name/Information | | |
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^{**} N1 does not include N1mi. T1 N1mi M0 and T0 N1mi M0 cancers are included for prognostic staging with T1 N0 M0 cancers of the same prognostic factor status.

^{***} N1 includes N1mi. T2, T3, and T4 cancers and N1mi are included for prognostic staging with T2 N1, T3 N1 and T4 N1, respectively. Nates:

7.2.1 Genomic Profile for Pathological Prognostic Staging

When Oncotype Dx Score is less than 11...

| And TNM is | And Grade is | And HER2 Status is | And ER Status is | And PR Status is | Then the Pathological Prognostic Stage Group is | 1 |
|----------------------|--------------|--------------------|------------------|---------------------|--|---|
| T1 N0 M0 T2 N0 M0 | Any | Negative | Positive | Any | IA | |

Notes

- 1. Obtaining genomic profiles is NOT required for assigning Pathological Prognostic Stage. However genomic profiles may be performed for use in determining appropriate treatment. If the OncotypeDx® test is performed in cases with a T1N0M0 or T2N0M0 cancer that is HER2-negative and ER-positive, and the recurrence score is less than 11, the case should be assigned Pathological Prognostic Stage Group IA.
- 2. If OncotypeDx® is not performed, or if it is performed and the OncotypeDx® score is not available, or is 11 or greater for patients with T1-2 N0 M0 HER2—negative, ER-positive cancer, then the Prognostic Stage Group is assigned based on the anatomic and biomarker categories shown above.
- 3. OncotypeDx® is the only multigene panel included to classify Pathologic Prognostic Stage because prospective Level I data supports this use for patients with a score <11. Future updates to the staging system may include results from other multigene panels to assign cohorts of patients to Prognostic Stage Groups based on the then available evidence. Inclusion or exclusion in this staging table of a genomic profile assay is not an endorsement of any specific assay and should not limit appropriate clinical use of any genomic profile assay based on evidence available at the time of treatment.

| Hospital Name/Address | Patient Name/Information | | |
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| 8 | Re | egistry Data | a Collection \ | /ariables | | |
|------|---------|-----------------------------------|---------------------|--|-----------------------------|-----------------------------------|
| See | chapt | ter for more de | tails on these vari | ables. | | |
| | 1. | ER: | positive | negative | percent positive: | Allred score, if available: |
| | 2. | P. PR: ☐ positive ☐ negative pero | | percent positive: | Allred score, if available: | |
| | 3. | HER2—IHC: | O | 1+2+ | 3+ unkno | own not performed |
| | 4. | HER2—FISH: | negat | ive positive | e equivo | ocal unknown not performed |
| | | | HER2:CE | P17 ratio: | HER2 copy number, | if available: |
| | 5. | HER2: | Overall result | negative | positive | unknown if done not performed |
| | 6. | Nottingham h | nistologic grade: | low (1) | intermediate (2) | ☐ high (3) |
| | 7. | Ki-67, if availa | able – percent pos | itive: | | |
| | 8. | Oncotype Dx | erecurrence score | e (numeric score preferr | ed over risk level): | |
| | 9. | Oncotype Dx | DCIS recurrence | score (numeric score pr | referred over risk leve | el): |
| | 10. | Mammaprint | ® (numeric score ¡ | oreferred over risk level |): | |
| | 11. | ProSigna [®] PA | M50 intrinsic subt | ypes and Risk of Recurr | ence score (numeric s | score preferred over risk level): |
| | 12. | Breast Cancer | r Index (numeric s | core preferred over risk | (level): | |
| | 13. | EndoPredict (| numeric score pre | eferred over risk level): | , | |
| | 14. | | c score preferred | , | | |
| | 15. | • | • | or (uPA) and plasminoge | en activator inhibitor | type 1 (PAI-1) ⁶ : |
| | 16. | Response to t | | ☐ CR ☐ PR | □NR | |
| | | Tresponse to t | cuciniciic. | | | |
| _ | | _ | | (a. a. a.) | | |
| 9 | Ly | mphovasc | ular Invasion | (LVI) | | |
| | Con | nponent of | Description | | | |
| ✓ | | Coding | Description | | | |
| | 0 | | · | (absent)/not identified | | |
| | 1 | | LVI present/ider | · | | |
| | 2 | | | small vessel invasion onl | ly (L) | |
| | 3 | | | essel) invasion only (V) | | |
| | 9 | | | and small vessel AND vunknown/indeterminat | | nvasion |
| | | | Tresence of Evi | unknownymaetermmat | | |
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| Phys | sician | Signature | | | | Date/Time |
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10 Anatomy

FIGURE 48.1. Anatomic sites and subsites of the right breast.

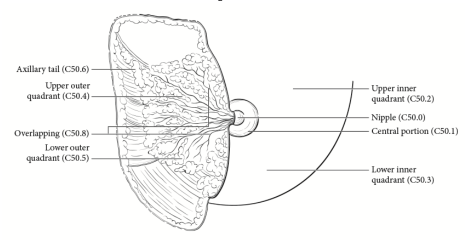
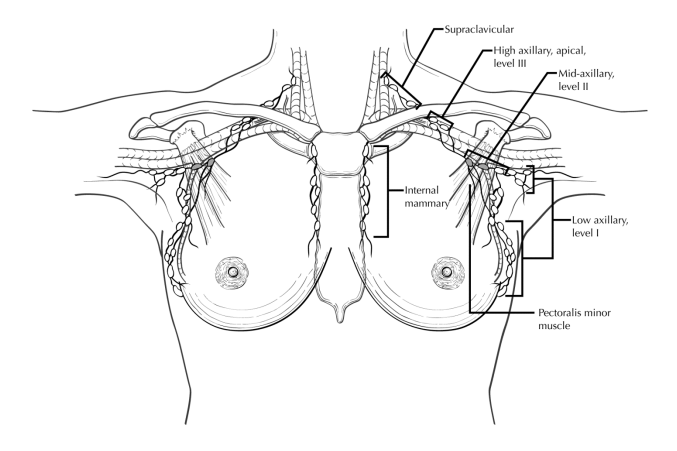


FIGURE 48.2. Schematic diagram of the breast and regional lymph nodes.



| Hospital Name/Address | Patient Name/Information |
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11 Bibliography

- 1. Scarff R, Handley R. Prognosis in carcinoma of the breast. *The Lancet*. 1938;232(6001):582-583.
- 2. Black MM. Survival in breast cancer cases in relation to the structure of the primary tumor and regional lymphnodes. *Surg Gynecol Obstet.* 1955;100:543-551.
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1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|--|--|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information diagnostic workup from clinical staging combined with operative findings, and pathology review of resected suspecimens | |
| | ycTNM Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant the before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neo therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | FIGO Stage | T Criteria |
|---|------------|------------|--|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | I | Tumor confined to the vulva and/or perineum |
| | | | Multifocal lesions should be designated as such. The largest lesion or the lesion with the greatest depth of invasion will be the target lesion identified to address the highest pT stage. |
| | | | Depth of invasion is defined as the measurement of the tumor from the epithelial–stromal junction of the adjacent most superficial dermal papilla to the deepest point of invasion. |
| | T1a | IA | Lesions 2 cm or less, confined to the vulva and/or perineum, and with stromal invasion of 1.0 mm or less |
| | T1b | IB | Lesions more than 2 cm, <i>or</i> any size with stromal invasion of more than 1.0 mm, confined to the vulva and/or perineum |
| | T2 | II | Tumor of any size with extension to adjacent perineal structures (lower/distal third of the urethra, lower/distal third of the vagina, anal involvement) |
| | Т3 | IVA | Tumor of any size with extension to any of the following—upper/proximal two thirds of the urethra, upper/proximal two thirds of the vagina, bladder mucosa, or rectal mucosa—or fixed to pelvic bone |

| * | / | T Suffix | Definition |
|----------|---|----------|---|
| | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria |
|---|------------|------------|---|
| | NX | | Regional lymph nodes cannot be assessed |
| | N0 | | No regional lymph node metastasis |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm |
| | N1 | III | Regional lymph node metastasis with one or two lymph node metastases each less than 5 mm, or one lymph node metastasis ≥5 mm |
| | N1a* | IIIA | One or two lymph node metastases each less than 5 mm |
| | N1b | IIIA | One lymph node metastasis ≥5 mm |
| | N2 | | Regional lymph node metastasis with three or more lymph node metastases each less than 5 mm, or two or more lymph node metastases ≥5 mm, or lymph node(s) with extranodal extension |
| | N2a* | IIIB | Three or more lymph node metastases each less than 5 mm |
| | N2b | IIIB | Two or more lymph node metastases ≥5 mm |
| | N2c | IIIC | Lymph node(s) with extranodal extension |
| | N3 | IVA | Fixed or ulcerated regional lymph node metastasis |

Note: The site, size, and laterality of lymph node metastases should be recorded.

| ٧ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | FIGO Stage | M Criteria |
|----------|------------|------------|--|
| | cM0 | | No distant metastasis (no pathological M0; use clinical M to complete stage group) |
| | cM1 | IVB | Distant metastasis (including pelvic lymph node metastasis) |
| | pM1 | IVB | Distant metastasis (including pelvic lymph node metastasis), microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2 | N0 | M0 | II |
| | T1-T2 | N1-N2c | M0 | III |
| | T1-T2 | N1 | M0 | IIIA |
| | T1-T2 | N2a, N2b | M0 | IIIB |
| | T1-T2 | N2c | M0 | IIIC |
| | T1-T3 | N3 | M0-M1 | IV |
| | T1-T2 | N3 | M0 | IVA |
| | T3 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

| See | chap | ter for more details on these variables. |
|-----|------|--|
| | 1. | FIGO stage: |
| | 2. | Size of regional lymph node metastasis: |
| | 3. | Laterality of regional node metastasis: |
| | 4. | Femoral–inguinal nodal spread identified on imaging: |
| | 5. | Pelvic nodes identified on imaging: Yes No |
| | 6. | p16: immunohistochemistry? |
| | | |

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|-----------------------------|---------------------------|--|
| | GX Grade cannot be assessed | | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

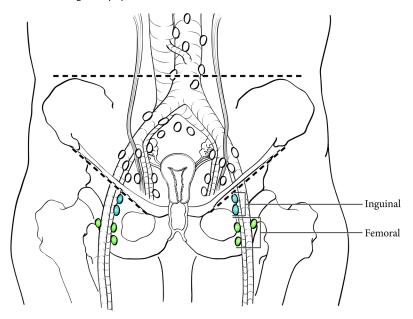
| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 50.1. Vulva and perineum lesions, from top to bottom: the lesion at the top is vulvar, the middle two lesions are perineal, and the lesion at the bottom is considered perianal.



FIGURE 50.2. Regional lymph nodes of the vulva.



| Physician Signature | Date/Time |
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| Patient Name/Information | |
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| | Patient Name/Information |

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | FIGO Stage | T Criteria |
|---|------------|------------|---|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | 1 | Tumor confined to the vagina |
| | T1a | 1 | Tumor confined to the vagina, measuring ≤2.0 cm |
| | T1b | 1 | Tumor confined to the vagina, measuring >2.0 cm |
| | T2 | II | Tumor invading paravaginal tissues but not to pelvic sidewall |
| | T2a | II | Tumor invading paravaginal tissues but not to pelvic wall, measuring ≤2.0 cm |
| | T2b | II | Tumor invading paravaginal tissues but not to pelvic wall, measuring >2.0 cm |
| | T3 | III | Tumor extending to the pelvic sidewall* and/or causing hydronephrosis or nonfunctioning kidney |
| | T4 | IVA | Tumor invading the mucosa of the bladder or rectum and/or extending beyond the true pelvis (bullous edema is not sufficient evidence to classify a tumor as T4) |

^{*}Pelvic sidewall is defined as the muscle, fascia, neurovascular structures, or skeletal portions of the bony pelvis. On rectal examination, there is no cancer-free space between the tumor and pelvic sidewall.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria |
|---|------------|------------|---|
| | NX | | Regional lymph nodes cannot be assessed |
| | N0 | | No regional lymph node metastasis |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm |
| | N1 | Ш | Pelvic or inguinal lymph node metastasis |

| ✓ | N Suffix | Definition | |
|---|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | FIGO Stage | M Criteria |
|----------|------------|------------|---|
| | cM0 | | No distant metastasis |
| | cM1 | IVB | Distant metastasis |
| | pM1 | IVB | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| √ | When T is | And N is | And M is | Then the stage group is |
|----------|-----------|----------|----------|-------------------------|
| | T1a | NO | M0 | IA |
| | T1b | NO | M0 | IB |
| | T2a | N0 | M0 | IIA |
| | T2b | N0 | M0 | IIB |
| | T1-T3 | N1 | M0 | III |
| | T3 | N0 | M0 | III |
| | T4 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

| ee | chapt | er for more details on these variables. | | | | | | |
|----|-------|--|-----|------|------|--|--|--|
| | 1. | FIGO stage: | | | | | | |
| | 2. | Pelvic nodes identified on imaging: | Yes | ☐ No | | | | |
| | 3. | Para-aortic nodes identified on imaging: | Yes | ☐ No | | | | |
| | 4. | l. Distant (mediastinal, scalene) nodes identified on imaging: | | Yes | ☐ No | | | |

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

8 Lymphovascular Invasion (LVI)

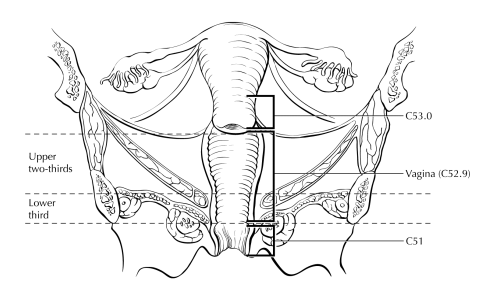
| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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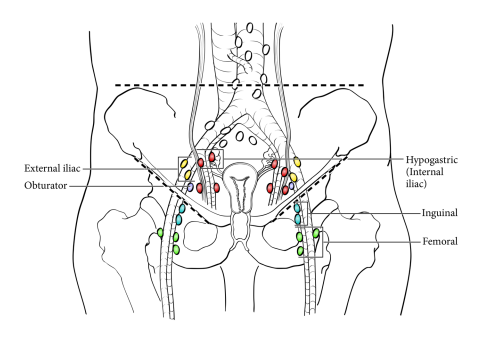
9 Anatomy

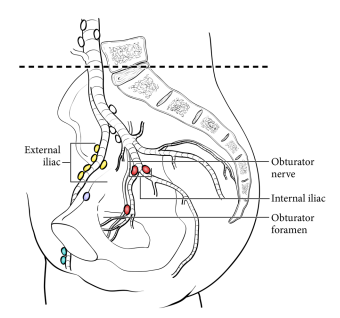
FIGURE 51.1 Anatomic sites and subsites of the vagina.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 51.2. Regional lymph nodes for the vagina.





| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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Cervix Uteri Version 9

- For cases diagnosed 1/1/21 use Protocol for Cancer Staging Documentation: Cervix Uteri Version 9
- Cervix Uteri Version 9 is available on Kindle
- We will not be providing a staging form for Version 9
- For any questions regarding Cervix Uteri Version 9 please contact ajcc@facs.org

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | FIGO Stage | T Criteria |
|---|------------|------------|---|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | 1 | Cervical carcinoma confined to the uterus (extension to corpus should be disregarded) |
| | T1a | IA | Invasive carcinoma diagnosed only by microscopy. Stromal invasion with a maximum depth of 5.0 mm measured from the base of the epithelium and a horizontal spread of 7.0 mm or less. Vascular space involvement, venous or lymphatic, does not affect classification. |
| | T1a1 | IA1 | Measured stromal invasion of 3.0 mm or less in depth and 7.0 mm or less in horizontal spread |
| | T1a2 | IA2 | Measured stromal invasion of more than 3.0 mm and not more than 5.0 mm, with a horizontal spread of 7.0 mm or less |
| | T1b | IB | Clinically visible lesion confined to the cervix or microscopic lesion greater than T1a/IA2. Includes all macroscopically visible lesions, even those with superficial invasion. |
| | T1b1 | IB1 | Clinically visible lesion 4.0 cm or less in greatest dimension |
| | T1b2 | IB2 | Clinically visible lesion more than 4.0 cm in greatest dimension |
| | T2 | II | Cervical carcinoma invading beyond the uterus but not to the pelvic wall or to lower third of the vagina |
| | T2a | IIA | Tumor without parametrial invasion |
| | T2a1 | IIA1 | Clinically visible lesion 4.0 cm or less in greatest dimension |
| | T2a2 | IIA2 | Clinically visible lesion more than 4.0 cm in greatest dimension |
| | T2b | IIB | Tumor with parametrial invasion |
| | Т3 | III | Tumor extending to the pelvic sidewall* and/or involving the lower third of the vagina and/or causing hydronephrosis or nonfunctioning kidney |
| | T3a | IIIA | Tumor involving the lower third of the vagina but not extending to the pelvic wall |
| | T3b | IIIB | Tumor extending to the pelvic wall and/or causing hydronephrosis or nonfunctioning kidney |
| | T4 | IVA | Tumor invading the mucosa of the bladder or rectum and/or extending beyond the true pelvis (bullous edema is not sufficient to classify a tumor as T4) |

^{*}The pelvic sidewall is defined as the muscle, fascia, neurovascular structures, and skeletal portions of the bony pelvis. On rectal examination, there is no cancer-free space between the tumor and pelvic sidewall.

| ✓ | T Suffix | Definition | |
|----------|----------|---|--|
| | (m) | Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| 1 | N Category | FIGO Stage | N Criteria | |
|---|------------|------------|---|--|
| | NX | | legional lymph nodes cannot be assessed | |
| | N0 | | No regional lymph node metastasis | |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm | |
| | N1 | | Regional lymph node metastasis | |

| 1 | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

| Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | FIGO Stage | M Criteria | |
|---|------------|------------|--|--|
| | cM0 | | No distant metastasis | |
| | cM1 | IVB | Distant metastasis (including peritoneal spread or involvement of the supraclavicular, mediastinal, or distant lymph nodes; lung; liver; or bone) | |
| | pM1 | IVB | Distant metastasis (including peritoneal spread or involvement of the supraclavicular, mediastinal, or distant lymph nodes; lung; liver; or bone), microscopically confirmed | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | Any N | M0 | I |
| | T1a | Any N | M0 | IA |
| | T1a1 | Any N | M0 | IA1 |
| | T1a2 | Any N | M0 | IA2 |
| | T1b | Any N | M0 | IB |
| | T1b1 | Any N | M0 | IB1 |
| | T1b2 | Any N | M0 | IB2 |
| | T2 | Any N | M0 | II |
| | T2a | Any N | M0 | IIA |
| | T2a1 | Any N | M0 | IIA1 |
| | T2a2 | Any N | M0 | IIA2 |
| | T2b | Any N | M0 | IIB |
| | T3 | Any N | M0 | III |
| | T3a | Any N | M0 | IIIA |
| | T3b | Any N | M0 | IIIB |
| | T4 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

| | n these variables. |
|--|--------------------|
| | |
| | |

| 1. | FIGO stage: |
|----|--|
| 2. | Pelvic nodal status and method of assessment (microscopic, CT, PET, MR imaging): |
| 3. | Para-aortic nodal status and method of assessment: |
| 4. | Distant (mediastinal, scalene) nodal status and method of assessment: |
| 5. | P16 status: |
| 6. | HIV status: |

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

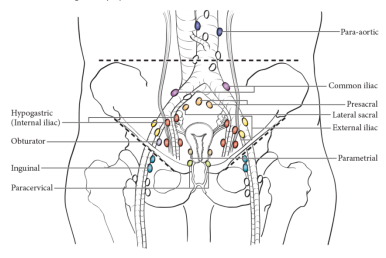
| 1 | G | G Definition | |
|---|----|---------------------------|--|
| | GX | Grade cannot be assessed | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 52.1. Regional lymph nodes for the cervix uteri.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | | |
|---|----------------|---|--|--|--|
| workup information, until first treatment, including clinical history and symptoms, physic endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sent | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information fro diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgispecimens | | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy a before planned surgery. Criteria : First therapy is systemic and/or radiation therapy | | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | | |

| Hospital Name/Address | Patient Name/Information |
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| | ! |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | FIGO Stage | T Criteria | |
|---|------------|------------|---|--|
| | TX | | Primary tumor cannot be assessed | |
| | T0 | | No evidence of primary tumor | |
| | T1 | 1 | Tumor confined to the corpus uteri, including endocervical glandular involvement | |
| | T1a | IA | Tumor limited to the endometrium or invading less than half the myometrium | |
| | T1b | IB | Tumor invading one half or more of the myometrium | |
| | T2 | Ш | Tumor invading the stromal connective tissue of the cervix but not extending beyond the uterus. | |
| | | | Does NOT include endocervical glandular involvement. | |
| | T3 | Ш | Tumor involving serosa, adnexa, vagina, or parametrium | |
| | T3a | IIIA | Tumor involving the serosa and/or adnexa (direct extension or metastasis) | |
| | T3b | IIIB | Vaginal involvement (direct extension or metastasis) or parametrial involvement | |
| | T4 | IVA | Tumor invading the bladder mucosa and/or bowel mucosa (bullous edema is not sufficient to | |
| | | | classify a tumor as T4) | |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria | |
|-----|----------------------|--------------------|--|--|
| | NX | | Regional lymph nodes cannot be assessed | |
| | N0 | | No regional lymph node metastasis | |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm | |
| | N1 | IIIC1 | Regional lymph node metastasis to pelvic lymph nodes | |
| | N1mi | IIIC1 | Regional lymph node metastasis (greater than 0.2 mm but not greater than 2.0 mm in diameter) to pelvic lymph nodes | |
| | N1a | IIIC1 | Regional lymph node metastasis (greater than 2.0 mm in diameter) to pelvic lymph nodes | |
| | N2 | IIIC2 | Regional lymph node metastasis to para-aortic lymph nodes, with or without positive pelvic lymph nodes | |
| | N2mi | IIIC2 | Regional lymph node metastasis (greater than 0.2 mm but not greater than 2.0 mm in diameter) to para-aortic lymph nodes, with or without positive pelvic lymph nodes | |
| | N2a | IIIC2 | Regional lymph node metastasis (greater than 2.0 mm in diameter) to para-aortic lymph nodes, with or without positive pelvic lymph nodes | |
| Suf | fix (sn) is added to | the N category whe | en metastasis is identified only by sentinel lymph node biopsy. | |

| | ✓ | N Suffix | efinition | |
|---|---|----------|--|--|
| I | | (sn) | elect if regional lymph node metastasis identified by SLN biopsy only. | |
| Γ | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | FIGO Stage | M Criteria |
|---|------------|------------|--|
| | cM0 | | No distant metastasis |
| | cM1 | IVB | Distant metastasis (includes metastasis to inguinal lymph nodes, intraperitoneal disease, lung, liver, or bone) |
| | | | (It excludes metastasis to pelvic or para-aortic lymph nodes, vagina, uterine serosa, or adnexa.) |
| | pM1 | IVB | Distant metastasis (includes metastasis to inguinal lymph nodes, intraperitoneal disease, lung, liver, or bone), microscopically confirmed |
| | | | (It excludes metastasis to pelvic or para-aortic lymph nodes, vagina, uterine serosa, or adnexa.) |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|-------------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2 | NO | M0 | II |
| | T3 | N0 | M0 | III |
| | T3a | NO | M0 | IIIA |
| | T3b | NO | M0 | IIIB |
| | T1-T3 | N1/N1mi/N1a | M0 | IIIC1 |
| | T1-T3 | N2/N2mi/N2a | M0 | IIIC2 |
| | T4 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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| 6 | 6 Registry Data Collection Variables | | | | | |
|-----------------|---|-------------|---|---|--|--|
| | See chapter for more details on these variables. | | | | | |
| | 1. | FIGO stage: | | | | |
| | 2. | | | | | |
| | Lymphovascular space invasion: | | | | | |
| | 4. Peritoneal cytology results: Collected? Yes No No | | | | | |
| | | | If yes: Positive | Negative | | |
| | 5. | Estro | ogen and progesterone receptor status: | | | |
| | 6. | Tum | or suppressor and oncogene expression: | s No | | |
| | 7. | Pelvi | ic nodal dissection with number of nodes positive/examir | ned: | | |
| | 8. | Para | -aortic nodal dissection with number of nodes positive/e | xamined: | | |
| | 9. | Perc | entage of nonendometrioid cell type in mixed-histology t | rumors: | | |
| | 10. | Ome | entectomy performed: Yes No | | | |
| | 11. | Mor | cellation: Yes No | | | |
| _ | | -4-1- | :- C | | | |
| <u>7</u> | HI | Stoic | ogic Grade (G) | | | |
| 1 | G | 1 | C Definition | | | |
| • | GX | | G Definition Grade cannot be assessed | | | |
| | G1 | | Well differentiated | | | |
| | G2 | | Moderately differentiated | | | |
| | G3 | | Poorly differentiated | | | |
| 7.1 Case | | | pathology: Degree of Differentiation oma of the corpus uteri should be grouped according to ti | he degree of differentiation of the endometrioid adenocarcinoma: | | |
| ✓ | G | | G Definition | | | |
| | G1 | | 5% or less of a nonsquamous or nonmorular solid growt | | | |
| | G2 G3 | | 6–50% of a nonsquamous or nonmorular solid growth p More than 50% of a nonsquamous or nonmorular solid a | | | |
| | 00 | | Papillary serous, clear cell, and carcinosarcoma are cons | | | |
| | | | | | | |
| Not | | Patho | logical Grading | voyageted for the explitectural grade increases the turner grade by 1 | | |
| | 1. | | (i.e., 1 to 2 and 2 to 3). | y expected for the architectural grade increases the tumor grade by 1 | | |
| | 2. | | Serous, clear cell, and mixed mesodermal tumors are <i>hi</i> | gh risk and considered grade 3. | | |
| | 3. | | Adenocarcinomas with benign squamous elements (squ | amous metaplasia) are graded according to the nuclear grade of the | | |
| | | | glandular component. | | | |
| This | form | conti | nues on the next page. | | | |
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| Hos | Hospital Name/Address Patient Name/Information | | | | | |
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8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description | |
|---|--|--|--|
| • | LVI Coding | | |
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 BOTH lymphatic and small vessel AND venous (large vessel) invasion | | |
| | 9 Presence of LVI unknown/indeterminate | | |

9 Anatomy

FIGURE 53.1. Anatomic sites and subsites of the corpus uteri.

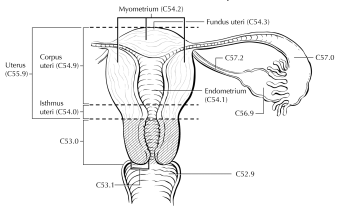
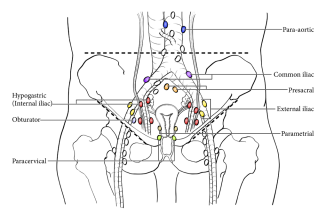


FIGURE 53.2. Regional lymph nodes of the corpus uteri.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | |
| | |
| | Patient Name/Information |

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2 Instructions

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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | FIGO Stage | T Criteria | |
|---|------------|------------|---|--|
| | TX | | Primary tumor cannot be assessed | |
| | TO | | No evidence of primary tumor | |
| | T1 | 1 | Tumor limited to the uterus | |
| | T1a | IA | Tumor 5 cm or less in greatest dimension | |
| | T1b | IB | Tumor more than 5 cm | |
| | T2 | II | Tumor extends beyond the uterus, within the pelvis | |
| | T2a | IIA | Tumor involves adnexa | |
| | T2b | IIB | Tumor involves other pelvic tissues | |
| | T3 | III | Tumor infiltrates abdominal tissues | |
| | T3a | IIIA | Tumor infiltrates abdominal tissues in one site | |
| | T3b | IIIB | Tumor infiltrates abdominal tissues in more than one site | |
| | T4 | IVA | Tumor invades bladder or rectum | |

| | ✓ | T Suffix | uffix Definition | |
|---|---|---|------------------|--|
| ſ | | (m) Select if synchronous primary tumors are found in single organ. | | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria |
|---|------------|------------|---|
| | NX | | Regional lymph nodes cannot be assessed |
| | N0 | | No regional lymph node metastasis |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm |
| | N1 | IIIC | Regional lymph node metastasis |

| ✓ | N Suffix | Definition | |
|----------|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | FIGO Stage | M Criteria |
|---|------------|------------|---|
| | cM0 | | No distant metastasis |
| | cM1 | IVB | Distant metastasis (excluding adnexa, pelvic, and abdominal tissues) |
| | pM1 | IVB | Distant metastasis (excluding adnexa, pelvic, and abdominal tissues), microscopically |
| | | | confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| √ | When T is | And N is | And M is | Then the stage group is |
|----------|-----------|----------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T1a | NO | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T2 | NO | M0 | II |
| | T3a | NO | M0 | IIIA |
| | T3b | NO | M0 | IIIB |
| | T1-3 | N1 | M0 | IIIC |
| | T4 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

Peritoneal washings, if recorded:

| See chapter for more details on these varial | ວles. |
|--|-------|
|--|-------|

| 1. | Lymphovascular space involven | nent: | |
|----|--|-----------|-----------------------------|
| 2. | Pelvic nodal dissection, with number of nodes positive/examined: | | |
| 3. | Para-aortic nodal dissection, wi | th number | of nodes positive/examined: |
| 4. | Omentectomy performed: | Yes | □No |
| 5. | Morcellation performed: | Yes | □No |
| 6 | Cytogenetic analysis (ESS only): | | |

Histologic Grade (G)

| ✓ | G | G Definition | |
|---|-----------------------------|---|--|
| | GX Grade cannot be assessed | | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated or undifferentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 53.1. Anatomic sites and subsites of the corpus uteri.

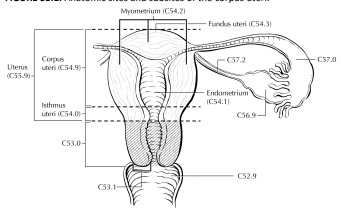
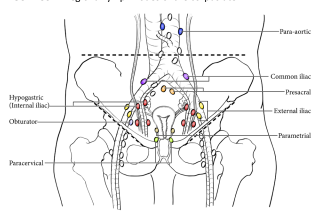


FIGURE 53.2. Regional lymph nodes of the corpus uteri.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
|--------------------------|--------------------------|
| | |
| | |
| | |
| | Patient Name/Information |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information fro diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgi specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | FIGO Stage | T Criteria |
|---|------------|------------|---|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | 1 | Tumor limited to the uterus |
| | T1a | IA | Tumor limited to the endometrium/endocervix |
| | T1b | IB | Tumor invades to less than half of the myometrium |
| | T1c | IC | Tumor invades one half or more of the myometrium |
| | T2 | II | Tumor extends beyond the uterus, within the pelvis |
| | T2a | IIA | Tumor involves adnexa |
| | T2b | IIB | Tumor involves other pelvic tissues |
| | T3 | III | Tumor infiltrates abdominal tissues |
| | T3a | IIIA | Tumor infiltrates abdominal tissues in one site |
| | T3b | IIIB | Tumor infiltrates abdominal tissues in more than one site |
| | T4 | IVA | Tumor invades bladder or rectum |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria |
|---|------------|------------|---|
| | NX | | Regional lymph nodes cannot be assessed |
| | N0 | | No regional lymph node metastasis |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm |
| | N1 | IIIC | Regional lymph node metastasis |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | FIGO Stage | M Criteria |
|---|------------|------------|---|
| | cM0 | | No distant metastasis |
| | cM1 | IVB | Distant metastasis (excluding adnexa, pelvic, and abdominal tissues) |
| | pM1 | IVB | Distant metastasis (excluding adnexa, pelvic, and abdominal tissues), microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T1a | N0 | M0 | IA |
| | T1b | N0 | M0 | IB |
| | T1c | N0 | M0 | IC |
| | T2 | N0 | M0 | II |
| | T3a | N0 | M0 | IIIA |
| | T3b | N0 | M0 | IIIB |
| | T1-3 | N1 | M0 | IIIC |
| | T4 | Any N | M0 | IVA |
| | Any T | Any N | M1 | IVB |

6 Registry Data Collection Variables

| See | chanter | for | more | details | οn | these | variables. |
|-----|---------|-----|------|---------|----|-------|------------|
| | | | | | | | |

| 1. | Lymphovascular space involve | ment: | | | |
|----|---------------------------------|-------------|---------------|------------------|--|
| 2. | Pelvic nodal dissection, with n | umber of no | odes positive | /examined: | |
| 3. | Para-aortic nodal dissection, v | vith number | of nodes po | sitive/examined: | |
| 4. | Omentectomy performed: | Yes | ☐ No | | |
| 5. | Morcellation: Yes | ☐ No | | | |
| 6. | Presence of sarcomatous over | growth: | | | |
| 7. | Peritoneal washings, if record | ed: | | | |

7 Histologic Grade (G)

| 1 | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated or undifferentiated |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 53.1. Anatomic sites and subsites of the corpus uteri.

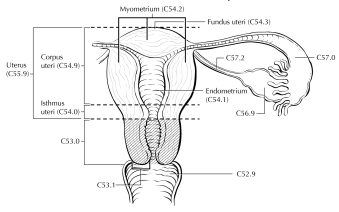
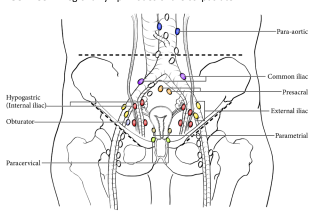


FIGURE 53.2. Regional lymph nodes of the corpus uteri.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | | |
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| | | |
| | Patient Name/Information | |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|---|----------------|---|--|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | FIGO Stage | T Criteria |
|---|------------|------------|---|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | 1 | Tumor limited to ovaries (one or both) or fallopian tube(s) |
| | T1a | IA | Tumor limited to one ovary (capsule intact) or fallopian tube, no tumor on ovarian or fallopian tube surface; no malignant cells in ascites or peritoneal washings |
| | T1b | IB | Tumor limited to both ovaries (capsules intact) or fallopian tubes; no tumor on ovarian or fallopian tube surface; no malignant cells in ascites or peritoneal washings |
| | T1c | IC | Tumor limited to one or both ovaries or fallopian tubes, with any of the following: |
| | T1c1 | IC1 | Surgical spill |
| | T1c2 | IC2 | Capsule ruptured before surgery or tumor on ovarian or fallopian tube surface |
| | T1c3 | IC3 | Malignant cells in ascites or peritoneal washings |
| | T2 | II | Tumor involves one or both ovaries or fallopian tubes with pelvic extension below pelvic brim or primary peritoneal cancer |
| | T2a | IIA | Extension and/or implants on the uterus and/or fallopian tube(s) and/or ovaries |
| | T2b | IIB | Extension to and/or implants on other pelvic tissues |
| | Т3 | III | Tumor involves one or both ovaries or fallopian tubes, or primary peritoneal cancer, with microscopically confirmed peritoneal metastasis outside the pelvis and/or metastasis to the retroperitoneal (pelvic and/or para-aortic) lymph nodes |
| | T3a | IIIA2 | Microscopic extrapelvic (above the pelvic brim) peritoneal involvement with or without positive retroperitoneal lymph nodes |
| | T3b | IIIB | Macroscopic peritoneal metastasis beyond pelvis 2 cm or less in greatest dimension with or without metastasis to the retroperitoneal lymph nodes |
| | T3c | IIIC | Macroscopic peritoneal metastasis beyond the pelvis more than 2 cm in greatest dimension with or without metastasis to the retroperitoneal lymph nodes (includes extension of tumor to capsule of liver and spleen without parenchymal involvement of either organ) |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | FIGO Stage | N Criteria | |
|---|------------|------------|---|--|
| | NX | | Regional lymph nodes cannot be assessed | |
| | N0 | | No regional lymph node metastasis | |
| | N0(i+) | | Isolated tumor cells in regional lymph node(s) no greater than 0.2 mm | |
| | N1 | IIIA1 | Positive retroperitoneal lymph nodes only (histologically confirmed) | |
| | N1a | IIIA1i | Metastasis up to and including 10 mm in greatest dimension | |
| | N1b | IIIA1ii | Metastasis more than 10 mm in greatest dimension | |

| | ✓ | N Suffix | Definition | |
|---|---|--|--|--|
| Ī | | (sn) | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| I | | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | FIGO Stage | M Criteria |
|---|------------|------------|--|
| | cM0 | | No distant metastasis |
| | cM1 | IV | Distant metastasis, including pleural effusion with positive cytology; liver or splenic parenchymal metastasis; metastasis to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside the abdominal cavity); and transmural involvement of intestine |
| | cM1b | IVB | Liver or splenic parenchymal metastases; metastases to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside the abdominal cavity); transmural involvement of intestine |
| | pM1 | IV | Distant metastasis, including pleural effusion with positive cytology; liver or splenic parenchymal metastasis; metastasis to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside the abdominal cavity); and transmural involvement of intestine, microscopically confirmed |
| | pM1a | IVA | Pleural effusion with positive cytology, microscopically confirmed |
| | pM1b | IVB | Liver or splenic parenchymal metastases; metastases to extra-abdominal organs (including inguinal lymph nodes and lymph nodes outside the abdominal cavity); transmural involvement of intestine, microscopically confirmed |

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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|------------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T1a | NO | MO | IA |
| | T1b | NO | M0 | IB |
| | T1c | NO | M0 | IC |
| | T2 | NO | MO | II |
| | T2a | NO | M0 | IIA |
| | T2b | NO | M0 | IIB |
| | T1/T2 | N1 | MO | IIIA1 |
| | T3a | NX, N0, N1 | MO | IIIA2 |
| | T3b | NX, N0, N1 | MO | IIIB |
| | T3c | NX, N0, N1 | M0 | IIIC |
| | Any T | Any N | M1 | IV |
| | Any T | Any N | M1a | IVA |
| | Any T | Any N | M1b | IVB |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. FIGO stage:
- 2. Preoperative CA-125 level:
- 3. Gross residual tumor after primary cytoreductive surgery:
- 4. Residual tumor volume after primary cytoreductive surgery:
- 5. Residual tumor location following primary cytoreductive surgery:

7 Histologic Grade (G)

| √ | G | G Definition | |
|----------|------------------------------|---|--|
| | GX | GX Grade cannot be assessed | |
| | GB Borderline tumor | | |
| | G1 | Well differentiated | |
| | G2 Moderately differentiated | | |
| | G3 | Poorly differentiated or undifferentiated | |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description | |
|---|-------------------------|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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9 Anatomy

FIGURE 55.1. Anatomic sites of the ovary (C56.9), fallopian tube (C57.0) and primary peritoneum (C48).

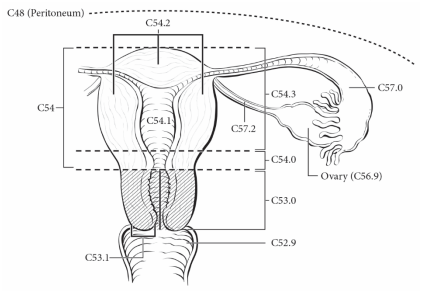
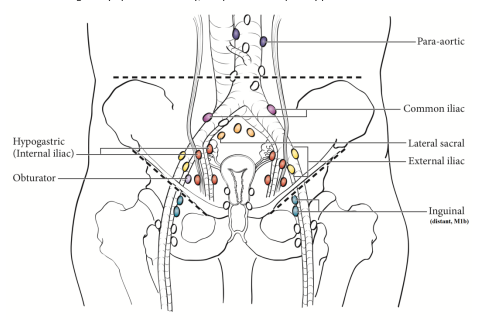


FIGURE 55.2. Regional lymph nodes of ovary, fallopian tube and primary peritoneal carcinomas.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | | | | |
|--|---|--|--|--|--|--|
| | cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnormation, until first treatment, including clinical history and symptoms, physical examination, in endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or same regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and relevant examinations | | | | | |
| pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of info diagnostic workup from clinical staging combined with operative findings, and pathology review of res specimens | | | | | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy a before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | | | |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | FIGO Stage | T Criteria |
|---|------------|------------|--|
| | TX | | Primary tumor cannot be assessed |
| | TO | | No evidence of primary tumor |
| | T1 | 1 | Tumor confined to uterus |
| | T2 | II | Tumor extends to other genital structures (ovary, tube, vagina, broad ligaments) by metastasis or direct extension |

| • | / 1 | T Suffix | Definition |
|---|---|----------|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | | |

4.2 Definition of Regional Lymph Node (N)

Nodal involvement in gestational trophoblastic neoplasia is uncommon (0.5%), but reportedly occurs in 6–16% of PSTTs.

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | FIGO Stage | M Criteria | |
|---|------------|------------|---|--|
| | cM0 | | No distant metastasis | |
| | cM1 | | Distant metastasis | |
| | cM1a | III | Lung metastasis | |
| | cM1b | IV | All other distant metastases | |
| | pM1 | | Distant metastasis, microscopically confirmed | |
| | pM1a | III | Lung metastasis, microscopically confirmed | |
| | pM1b | IV | All other distant metastases, microscopically confirmed | |

5 Prognostic Factors Required for Stage Grouping

5.1 Risk Score

Enter score for each factor and add scores together for total risk score.

| | Risk Score | | | | |
|---|-------------------|-------------------------------------|-------------------------------------|-------------------|--|
| Prognostic Factor | 0 1 2 | | 4 | Factor Score | |
| Age (years) | <40 | ≥40 | | | |
| Antecedent pregnancy | Hydatidiform mole | Abortion | Term pregnancy | | |
| Interval months from index pregnancy | <4 | 4–6 | 7–12 | >12 | |
| Pretreatment hCG (mIU/mL) | <103 | 10 ³ to <10 ⁴ | 10 ⁴ to <10 ⁵ | ≥10⁵ | |
| Largest tumor size, including uterus (cm) | <3 | 3–5 | >5 | | |
| Site of metastases | Lung | Spleen, kidney | Gastrointestinal tract | Brain, liver | |
| Number of metastases identified | | 1–4 | 5–8 | >8 | |
| Previous failed chemotherapy | | | Single drug | Two or more drugs | |
| Total Risk Score | | | | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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6 AJCC Prognostic Stage Groups

In 2000, FIGO combined its anatomic staging system with the modified WHO risk factor scoring system. In 2002, FIGO changed the WHO risk factor score cutoff for low-risk disease to <6, with high-risk disease >7, thus eliminating intermediate-risk disease. The current FIGO classification includes an anatomic stage designated by Roman numeral I, II, III, or IV, followed by the risk factor score expressed in Arabic numerals (e.g., Stage II: 4, Stage IV: 9).

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And Risk Score is | Then Stage is |
|---|-----------|----------|----------|-------------------|---------------|
| | T1 | n/a | M0 | 0 | 1:0 |
| | T1 | n/a | M0 | 1 | I:1 |
| | T1 | n/a | M0 | 2 | I:2 |
| | T1 | n/a | M0 | 3 | 1:3 |
| | T1 | n/a | M0 | 4 | 1:4 |
| | T1 | n/a | M0 | 5 | I:5 |
| | T1 | n/a | M0 | 6 | 1:6 |
| | T1 | n/a | M0 | 7 | 1:7 |
| | T1 | n/a | M0 | 8 | 1:8 |
| | T1 | n/a | M0 | 9 | 1:9 |
| | T1 | n/a | M0 | 10 | I:10 |
| | T1 | n/a | M0 | 11 | I:11 |
| | T1 | n/a | M0 | 12 | I:12 |
| | T1 | n/a | M0 | 13 | I:13 |
| | T1 | n/a | M0 | 14 | I:14 |
| | T1 | n/a | M0 | 15 | I:15 |
| | T1 | n/a | M0 | 16 | I:16 |
| | T1 | n/a | M0 | 17 | I:17 |
| | T1 | n/a | M0 | 18 | I:18 |
| | T1 | n/a | M0 | 19 | I:19 |
| | T1 | n/a | M0 | 20 | 1:20 |
| | T1 | n/a | M0 | 21 | I:21 |
| | T1 | n/a | M0 | 22 | 1:22 |
| | T1 | n/a | M0 | 23 | 1:23 |
| | T1 | n/a | M0 | 24 | 1:24 |
| | T1 | n/a | M0 | 25 | 1:25 |
| | T1 | n/a | M1a | 0 | III:0 |
| | T1 | n/a | M1a | 1 | III:1 |
| | T1 | n/a | M1a | 2 | III:2 |
| | T1 | n/a | M1a | 3 | III:3 |
| | T1 | n/a | M1a | 4 | III:4 |
| | T1 | n/a | M1a | 5 | III:5 |

| Hospital Name/Address | Patient Name/Information |
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| | |

| 1 | When T is | And N is | And M is | And Risk Score is | Then Stage is |
|---|-----------|----------|----------|-------------------|---------------|
| | T1 | n/a | M1a | 6 | III:6 |
| | T1 | n/a | M1a | 7 | III:7 |
| | T1 | n/a | M1a | 8 | III:8 |
| | T1 | n/a | M1a | 9 | III:9 |
| | T1 | n/a | M1a | 10 | III:10 |
| | T1 | n/a | M1a | 11 | III:11 |
| | T1 | n/a | M1a | 12 | III:12 |
| | T1 | n/a | M1a | 13 | III:13 |
| | T1 | n/a | M1a | 14 | III:14 |
| | T1 | n/a | M1a | 15 | III:15 |
| | T1 | n/a | M1a | 16 | III:16 |
| | T1 | n/a | M1a | 17 | III:17 |
| | T1 | n/a | M1a | 18 | III:18 |
| | T1 | n/a | M1a | 19 | III:19 |
| | T1 | n/a | M1a | 20 | III:20 |
| | T1 | n/a | M1a | 21 | III:21 |
| | T1 | n/a | M1a | 22 | III:22 |
| | T1 | n/a | M1a | 23 | III:23 |
| | T1 | n/a | M1a | 24 | III:24 |
| | T1 | n/a | M1a | 25 | III:25 |
| | T1 | n/a | M1b | 0 | IV:0 |
| | T1 | n/a | M1b | 1 | IV:1 |
| | T1 | n/a | M1b | 2 | IV:2 |
| | T1 | n/a | M1b | 3 | IV:3 |
| | T1 | n/a | M1b | 4 | IV:4 |
| | T1 | n/a | M1b | 5 | IV:5 |
| | T1 | n/a | M1b | 6 | IV:6 |
| | T1 | n/a | M1b | 7 | IV:7 |
| | T1 | n/a | M1b | 8 | IV:8 |
| | T1 | n/a | M1b | 9 | IV:9 |
| | T1 | n/a | M1b | 10 | IV:10 |
| | T1 | n/a | M1b | 11 | IV:11 |
| | T1 | n/a | M1b | 12 | IV:12 |
| | T1 | n/a | M1b | 13 | IV:13 |
| | T1 | n/a | M1b | 14 | IV:14 |
| | T1 | n/a | M1b | 15 | IV:15 |
| | T1 | n/a | M1b | 16 | IV:16 |
| | T1 | n/a | M1b | 17 | IV:17 |

| Hospital Name/Address | Patient Name/Information |
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| ✓ | When T is | And N is | And M is | And Risk Score is | Then Stage is |
|----------|-----------|----------|----------|-------------------|---------------|
| | T1 | n/a | M1b | 18 | IV:18 |
| | T1 | n/a | M1b | 19 | IV:19 |
| | T1 | n/a | M1b | 20 | IV:20 |
| | T1 | n/a | M1b | 21 | IV:21 |
| | T1 | n/a | M1b | 22 | IV:22 |
| | T1 | n/a | M1b | 23 | IV:23 |
| | T1 | n/a | M1b | 24 | IV:24 |
| | T1 | n/a | M1b | 25 | IV:25 |
| | T2 | n/a | M0 | 0 | II:0 |
| | T2 | n/a | M0 | 1 | II:1 |
| | T2 | n/a | M0 | 2 | II:2 |
| | T2 | n/a | M0 | 3 | II:3 |
| | T2 | n/a | M0 | 4 | II:4 |
| | T2 | n/a | M0 | 5 | II:5 |
| | T2 | n/a | M0 | 6 | II:6 |
| | T2 | n/a | M0 | 7 | II:7 |
| | T2 | n/a | M0 | 8 | II:8 |
| | T2 | n/a | M0 | 9 | II:9 |
| | T2 | n/a | M0 | 10 | II:10 |
| | T2 | n/a | M0 | 11 | II:11 |
| | T2 | n/a | M0 | 12 | II:12 |
| | T2 | n/a | M0 | 13 | II:13 |
| | T2 | n/a | M0 | 14 | II:14 |
| | T2 | n/a | M0 | 15 | II:15 |
| | T2 | n/a | M0 | 16 | II:16 |
| | T2 | n/a | M0 | 17 | II:17 |
| | T2 | n/a | M0 | 18 | II:18 |
| | T2 | n/a | M0 | 19 | II:19 |
| | T2 | n/a | M0 | 20 | II:20 |
| | T2 | n/a | M0 | 21 | II:21 |
| | T2 | n/a | M0 | 22 | II:22 |
| | T2 | n/a | M0 | 23 | II:23 |
| | T2 | n/a | M0 | 24 | II:24 |
| | T2 | n/a | M0 | 25 | II:25 |
| | T2 | n/a | M1a | 0 | III:0 |
| | T2 | n/a | M1a | 1 | III:1 |
| | T2 | n/a | M1a | 2 | III:2 |
| | T2 | n/a | M1a | 3 | III:3 |

| Hospital Name/Address | Patient Name/Information |
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| √ | When T is | And N is | And M is | And Risk Score is | Then Stage is |
|----------|-----------|----------|----------|-------------------|---------------|
| | T2 | n/a | M1a | 4 | III:4 |
| | T2 | n/a | M1a | 5 | III:5 |
| | T2 | n/a | M1a | 6 | III:6 |
| | T2 | n/a | M1a | 7 | III:7 |
| | T2 | n/a | M1a | 8 | III:8 |
| | T2 | n/a | M1a | 9 | III:9 |
| | T2 | n/a | M1a | 10 | III:10 |
| | T2 | n/a | M1a | 11 | III:11 |
| | T2 | n/a | M1a | 12 | III:12 |
| | T2 | n/a | M1a | 13 | III:13 |
| | T2 | n/a | M1a | 14 | III:14 |
| | T2 | n/a | M1a | 15 | III:15 |
| | T2 | n/a | M1a | 16 | III:16 |
| | T2 | n/a | M1a | 17 | III:17 |
| | T2 | n/a | M1a | 18 | III:18 |
| | T2 | n/a | M1a | 19 | III:19 |
| | T2 | n/a | M1a | 20 | III:20 |
| | T2 | n/a | M1a | 21 | III:21 |
| | T2 | n/a | M1a | 22 | III:22 |
| | T2 | n/a | M1a | 23 | III:23 |
| | T2 | n/a | M1a | 24 | III:24 |
| | T2 | n/a | M1a | 25 | III:25 |
| | T2 | n/a | M1b | 0 | IV:0 |
| | T2 | n/a | M1b | 1 | IV:1 |
| | T2 | n/a | M1b | 2 | IV:2 |
| | T2 | n/a | M1b | 3 | IV:3 |
| | T2 | n/a | M1b | 4 | IV:4 |
| | T2 | n/a | M1b | 5 | IV:5 |
| | T2 | n/a | M1b | 6 | IV:6 |
| | T2 | n/a | M1b | 7 | IV:7 |
| | T2 | n/a | M1b | 8 | IV:8 |
| | T2 | n/a | M1b | 9 | IV:9 |
| | T2 | n/a | M1b | 10 | IV:10 |
| | T2 | n/a | M1b | 11 | IV:11 |
| | T2 | n/a | M1b | 12 | IV:12 |
| | T2 | n/a | M1b | 13 | IV:13 |
| | T2 | n/a | M1b | 14 | IV:14 |
| | T2 | n/a | M1b | 15 | IV:15 |

| Hospital Name/Address | Patient Name/Information |
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56. Gestational Trophoblastic Neoplasms

| √ | When T is | And N is | And M is | And Risk Score is | Then Stage is |
|----------|-----------|----------|----------|-------------------|---------------|
| | T2 | n/a | M1b | 16 | IV:16 |
| | T2 | n/a | M1b | 17 | IV:17 |
| | T2 | n/a | M1b | 18 | IV:18 |
| | T2 | n/a | M1b | 19 | IV:19 |
| | T2 | n/a | M1b | 20 | IV:20 |
| | T2 | n/a | M1b | 21 | IV:21 |
| | T2 | n/a | M1b | 22 | IV:22 |
| | T2 | n/a | M1b | 23 | IV:23 |
| | T2 | n/a | M1b | 24 | IV:24 |
| | T2 | n/a | M1b | 25 | IV:25 |

7 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Risk score:
- 2. FIGO stage:

8 Histologic Grade G)

Histologic grade is not applicable to GTNs.

9 Lymphovascular Invasion LVI)

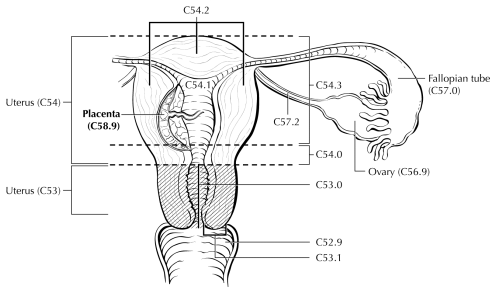
| ✓ | Component of LVI Coding | Description |
|---|-------------------------|---|
| | 0 | LVI not present absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only L) |
| | 3 | Venous large vessel) invasion only V) |
| | 4 | BOTH lymphatic and small vessel AND venous large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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10 Anatomy

FIGURE 56.1. Anatomic site of the placenta for gestational trophoblastic tumors.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
|--------------------------|--------------------------|
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| | |
| | |
| | Patient Name/Information |

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|--|------------|---|
| | TX | Primary tumor cannot be assessed |
| TO No evidence of primary tumor | | No evidence of primary tumor |
| | Tis | Carcinoma in situ (Penile intraepithelial neoplasia [PeIN]) |
| | Та | Noninvasive localized squamous cell carcinoma |
| | T1 | Glans: Tumor invades lamina propria |
| | | Foreskin: Tumor invades dermis, lamina propria, or dartos fascia |
| | | Shaft: Tumor invades connective tissue between epidermis and corpora regardless of location |
| | | All sites with or without lymphovascular invasion or perineural invasion and is or is not high grade |
| T1a Tumor is without lymphovascular invasion or perineural invasion and is not high grade | | Tumor is without lymphovascular invasion or perineural invasion and is not high grade (i.e., grade 3 or |
| | | sarcomatoid) |
| T1b Tumor exhibits lymphovascular invasion and/or perineural invasion or is hip | | Tumor exhibits lymphovascular invasion and/or perineural invasion or is high grade (i.e., grade 3 or sarcomatoid) |
| T2 Tumor invades into corpus spongiosum (either glans or ventral shaft) with or without ur | | Tumor invades into corpus spongiosum (either glans or ventral shaft) with or without urethral invasion |
| | T3 | Tumor invades into corpora cavernosum (including tunica albuginea) with or without urethral invasion |
| T4 Tumor invades into adjacent structures (i.e., scrotum, prostate, pubic bone) | | Tumor invades into adjacent structures (i.e., scrotum, prostate, pubic bone) |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria | | | | |
|---|---|--|--|--|--|--|
| | cNX | NX Regional lymph nodes cannot be assessed | | | | |
| | cNO No palpable or visibly enlarged inguinal lymph nodes | | | | | |
| | cN1 | Palpable mobile unilateral inguinal lymph node | | | | |
| | cN2 Palpable mobile ≥ 2 unilateral inguinal nodes or bilateral inguinal lymph nodes | | | | | |
| | cN3 | Palpable fixed inguinal nodal mass or pelvic lymphadenopathy unilateral or bilateral | | | | |

| 1 | N Suffix | Definition | | |
|---|--|--|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.2.2 Pathological N (pN)

| | The state of the s | | | | | |
|----------|--|-----------------|--|--|--|--|
| √ | N Category | gory N Criteria | | | | |
| | pNX Lymph node metastasis cannot be established | | | | | |
| | pNO No lymph node metastasis | | | | | |
| | pN1 ≤ 2 unilateral inguinal metastases, no ENE | | | | | |
| | pN2 ≥ 3 unilateral inguinal metastases or bilateral metastases, no ENE | | | | | |
| | pN3 ENE of lymph node metastases or pelvic lymph node metastases | | | | | |

| ~ | N Suffix | Definition | | |
|---|--|------------|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | | |

| Patient Name/Information | | |
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4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---------------------------|---|-----------------------|
| cM0 No distant metastasis | | No distant metastasis |
| | cM1 Distant metastasis | |
| | pM1 Distant metastasis, microscopically confirmed | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | MO | Ois |
| | Та | N0 | MO | 0a |
| | T1a | N0 | M0 | 1 |
| | T1b | N0 | M0 | IIA |
| | T2 | N0 | MO | IIA |
| | T3 | N0 | M0 | IIB |
| | T1-3 | N1 | M0 | IIIA |
| | T1-3 | N2 | M0 | IIIB |
| | T4 | Any N | M0 | IV |
| | Any T | N3 | M0 | IV |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

| C | ale a a tara | ۲ | | detection. | a collection | |
|-----|--------------|-----|------|------------|--------------|---------------|
| see | cnapter | tor | more | details | on the | se variables. |

- 1. Histologic subtype:
- 2. Size of largest nodal metastasis:
- 3. Total number of lymph nodes removed:
- 4. High-risk HPV expression:
- 5. p16 immunohistochemical expression:
- 6. Urethral mucosal invasion:

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|----------------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated/high grade |

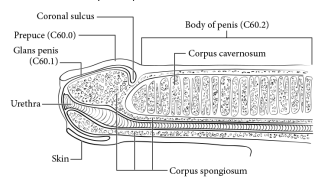
| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 57.1. Anatomy of the penis.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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| | | |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

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3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

4.1.1 Clinical T (cT)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | T1 | Clinically inapparent tumor that is not palpable |
| | T1a | Tumor incidental histologic finding in 5% or less of tissue resected |
| | T1b | Tumor incidental histologic finding in more than 5% of tissue resected |
| | T1c | Tumor identified by needle biopsy found in one or both sides, but not palpable |
| | T2 | Tumor is palpable and confined within prostate |
| | T2a | Tumor involves one-half of one side or less |
| | T2b | Tumor involves more than one-half of one side but not both sides |
| | T2c | Tumor involves both sides |
| | T3 | Extraprostatic tumor that is not fixed or does not invade adjacent structures |
| | T3a | Extraprostatic extension (unilateral or bilateral) |
| | T3b | Tumor invades seminal vesicle(s) |
| | T4 | Tumor is fixed or invades adjacent structures other than seminal vesicles such as external sphincter, rectum, |
| | | bladder, levator muscles, and/or pelvic wall |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Ī | | (m) | Select if synchronous primary tumors are found in single organ. |

4.1.2 Pathological T (pT)

| ✓ | T Category | T Criteria |
|-----|--|---|
| | T2 | Organ confined |
| | T3 | Extraprostatic extension |
| | T3a | Extraprostatic extension (unilateral or bilateral) or microscopic invasion of bladder neck |
| | T3b | Tumor invades seminal vesicle(s) |
| | T4 | Tumor is fixed or invades adjacent structures other than seminal vesicles such as external sphincter, rectum, |
| | | bladder, levator muscles, and/or pelvic wall |
| Not | e: There is no pathologic | al T1 classification. |
| Not | Note: Positive surgical margin should be indicated by an R1 descriptor, indicating residual microscopic disease. | |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No positive regional nodes |
| | N1 | Metastases in regional node(s) |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
| | | |
| | | |
| | | |
| | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|-----|---|---|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | cM1a | Nonregional lymph node(s) | |
| | cM1b | Bone(s) | |
| | cM1c | Other site(s) with or without bone disease | |
| | pM1 | Distant metastasis, microscopically confirmed | |
| | pM1a Nonregional lymph node(s), microscopically confirmed | | |
| | pM1b | Bone(s), microscopically confirmed | |
| | pM1c | Other site(s) with or without bone disease, microscopically confirmed | |
| Not | Note: When more than one site of metastasis is present, the most advanced category is used. M1c is most advanced. | | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Prostate-Specific Antigen (PSA)

PSA values are used to assign this category.

| ✓ | PSA values |
|---|------------|
| | < 10 |
| | ≥ 10 < 20 |
| | < 20 |
| | ≥ 20 |
| | Any value |

5.2 Definition of Histologic Grade Group (G)

| 1 | Grade Group (G) | Gleason Score | Gleason Pattern |
|---|-----------------|---------------|------------------|
| | 1 | ≤ 6 | ≤ 3+3 |
| | 2 | 7 | 3+4 |
| | 3 | 7 | 4+3 |
| | 4 | 8 | 4+4, 3+5, 5+3 |
| | 5 | 9 or 10 | 4+5, 5+4, or 5+5 |

| Hospital Name/Address | Patient Name/Information |
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6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And PSA is | And Grade | Then the stage |
|---|-------------------|----------|----------|------------|-----------|----------------|
| • | | | | | Group is | group is |
| | cT1a-c, cT2a | N0 | M0 | < 10 | 1 | 1 |
| | pT2 | N0 | M0 | < 10 | 1 | 1 |
| | cT1a-c, cT2a, pT2 | N0 | M0 | ≥ 10 < 20 | 1 | IIA |
| | cT2b-c | N0 | M0 | < 20 | 1 | IIA |
| | T1-2 | N0 | M0 | < 20 | 2 | IIB |
| | T1-2 | N0 | M0 | < 20 | 3 | IIC |
| | T1-2 | N0 | M0 | < 20 | 4 | IIC |
| | T1-2 | N0 | M0 | ≥ 20 | 1–4 | IIIA |
| | T3-4 | N0 | M0 | Any | 1–4 | IIIB |
| | Any T | N0 | M0 | Any | 5 | IIIC |
| | Any T | N1 | M0 | Any | Any | IVA |
| | Any T | Any N | M1 | Any | Any | IVB |

Note: When either PSA or Grade Group is not available, grouping should be determined by T category and/or either PSA or Grade Group as available.

| 7 | Registry | Data | Col | lection | Variak | oles |
|---|----------|------|-----|---------|--------|------|
|---|----------|------|-----|---------|--------|------|

| See chap | oter for more details on these variables. | | |
|----------|---|--|--|
| 1. | Pretreatment serum PSA levels lab value (in tenths, highest value XXX.X, last pre-diagnosis value): | | |
| 2. | Grade Group for clinical stage: | | |
| 3. | Gleason score for clinical stage: | | |
| 4. | Gleason patterns for clinical stage: | | |
| 5. | Grade Group for pathological stage: | | |
| 6. | 6. Gleason score for pathological stage: | | |
| 7. | 7. Gleason patterns for pathological stage: | | |
| 8. | Tertiary Gleason pattern on prostatectomy: | | |
| 9. | Number of cores examined: | | |
| 10 | 10. Number of cores positive: | | |
| 11 | Needle core biopsies positive: in one side in both sides beyond prostate | | |
| 12 | Metastatic sites: | | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 58.1. Anatomy of the prostate.

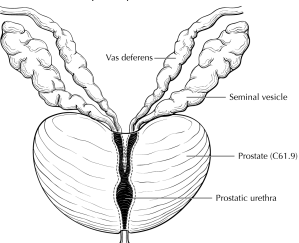
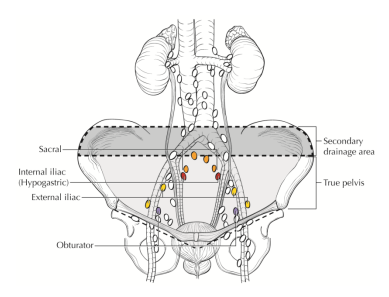


FIGURE 58.2. Lymph nodes of the prostate. The shaded area represents distribution of regional lymph nodes. The non-shaded area indicates nodes outside of regional distribution.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information | |
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria : First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

4.1.1 Clinical T (cT)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | cTX | Primary tumor cannot be assessed |
| | сТ0 | No evidence of primary tumor |
| | cTis | Germ cell neoplasia in situ |
| | cT4 | Tumor invades scrotum with or without vascular/lymphatic invasion |

Note: Except for Tis confirmed by biopsy and T4, the extent of the primary tumor is classified by radical orchiectomy. TX may be used for other categories for clinical staging.

| ✓ | T Suffix Definition | |
|---|---|--|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.1.2 Pathological T (pT)

| ✓ | T Category | T Criteria |
|-----|---------------------------|--|
| | pTX | Primary tumor cannot be assessed |
| | pT0 | No evidence of primary tumor |
| | pTis | Germ cell neoplasia in situ |
| | pT1 | Tumor limited to testis (including rete testis invasion) without lymphovascular invasion |
| | pT1a* | Tumor smaller than 3 cm in size |
| | pT1b* | Tumor 3 cm or larger in size |
| | pT2 | Tumor limited to testis (including rete testis invasion) with lymphovascular invasion OR |
| | | Tumor invading hilar soft tissue or epididymis or penetrating visceral mesothelial layer covering the external |
| | | surface of tunica albuginea with or without lymphovascular invasion |
| | pT3 | Tumor directly invades spermatic cord soft tissue with or without lymphovascular invasion |
| | pT4 | Tumor invades scrotum with or without lymphovascular invasion |
| *Su | bclassification of pT1 ap | plies only to pure seminoma. |

| / | T Suffix | Definition | |
|---|----------|---|--|
| | (m) | Select if synchronous primary tumors are found in single organ. | |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information |
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4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| ✓ | N Category | N Criteria | | |
|---|------------|--|--|--|
| | cNX | Regional lymph nodes cannot be assessed | | |
| | cN0 | No regional lymph node metastasis | | |
| | cN1 | Metastasis with a lymph node mass 2 cm or smaller in greatest dimension | | |
| | | DR . | | |
| | | Multiple lymph nodes, none larger than 2 cm in greatest dimension | | |
| | cN2 | Metastasis with a lymph node mass larger than 2 cm but not larger than 5 cm in greatest dimension | | |
| | | OR | | |
| | | Multiple lymph nodes, any one mass larger than 2 cm but not larger than 5 cm in greatest dimension | | |
| | cN3 | Metastasis with a lymph node mass larger than 5 cm in greatest dimension | | |

| ✓ | N Suffix | Definition | |
|---|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.2.2 Pathological N (pN)

| N Category | N Criteria | |
|--|--|--|
| pNX | Regional lymph nodes cannot be assessed | |
| pN0 | No regional lymph node metastasis | |
| pN1 | Metastasis with a lymph node mass 2 cm or smaller in greatest dimension and less than or equal to five nodes | |
| | positive, none larger than 2 cm in greatest dimension | |
| pN2 Metastasis with a lymph node mass larger than 2 cm but not larger than 5 cm in greatest dimension; or mo | | |
| | five nodes positive, none larger than 5 cm; or evidence of extranodal extension of tumor | |
| pN3 | Metastasis with a lymph node mass larger than 5 cm in greatest dimension | |

| 1 | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|---|---|--|
| | cM0 | No distant metastases | |
| | cM1 | Distant metastases | |
| | cM1a | Non-retroperitoneal nodal or pulmonary metastases | |
| | cM1b | Non-pulmonary visceral metastases | |
| | pM1 Distant metastases, microscopically confirmed | | |
| | pM1a Non-retroperitoneal nodal or pulmonary metastases, microscopically confirmed | | |
| | pM1b Non-pulmonary visceral metastases, microscopically confirmed | | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Serum Markers (S)

| 1 | S | S Criteria | | |
|----|---|---|--|--|
| • | Category | | | |
| | SX | Marker studies not available or not performed | | |
| | SO Marker study levels within normal limits | | | |
| | S1 | LDH < 1.5 × N*and hCG (mIU/mL) < 5,000 and AFP (ng/mL) < 1,000 | | |
| | S2 | LDH 1.5–10 × N*or hCG (mIU/mL) 5,000-50,000 or AFP (ng/mL) 1,000–10,000 | | |
| | S3 | LDH > 10 × N*or hCG (mIU/mL) >50,000 or AFP (ng/mL) > 10,000 | | |
| *N | *N indicates the upper limit of normal for the LDH assay. | | | |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | And S is | Then the stage |
|---|-----------|----------|----------|----------|----------------|
| • | | | | | group is |
| | pTis | N0 | M0 | SO SO | 0 |
| | pT1-T4 | N0 | M0 | SX | 1 |
| | pT1 | N0 | M0 | S0 | IA |
| | pT2 | N0 | M0 | SO SO | IB |
| | pT3 | N0 | M0 | SO SO | IB |
| | pT4 | N0 | M0 | SO SO | IB |
| | Any pT/TX | N0 | M0 | S1-3 | IS |
| | Any pT/TX | N1-3 | M0 | SX | II |
| | Any pT/TX | N1 | M0 | SO SO | IIA |
| | Any pT/TX | N1 | M0 | S1 | IIA |
| | Any pT/TX | N2 | M0 | SO SO | IIB |
| | Any pT/TX | N2 | M0 | S1 | IIB |
| | Any pT/TX | N3 | M0 | SO SO | IIC |
| | Any pT/TX | N3 | M0 | S1 | IIC |
| | Any pT/TX | Any N | M1 | SX | III |
| | Any pT/TX | Any N | M1a | SO SO | IIIA |
| | Any pT/TX | Any N | M1a | S1 | IIIA |
| | Any pT/TX | N1-3 | M0 | S2 | IIIB |
| | Any pT/TX | Any N | M1a | S2 | IIIB |
| | Any pT/TX | N1-3 | M0 | S3 | IIIC |
| | Any pT/TX | Any N | M1a | S3 | IIIC |
| | Any pT/TX | Any N | M1b | Any S | IIIC |

| Hospital Name/Address | Patient Name/Information |
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7 Registry Data Collection Variables

See chapter for more details on these variables.

Clinical stage grouping

- 1. Serum tumor markers (S) for clinical stage grouping:
- 2. Alpha fetoprotein (AFP) for clinical stage grouping (xx,xxx ng/mL):
- 3. Human chorionic gonadotropin (hCG) for clinical stage grouping (xx,xxx mIU/ml):
- 4. Lactate dehydrogenase (LDH) for clinical stage grouping (xx,xxx U/L):

Pathological stage grouping

- 5. Serum tumor markers (S) for pathological stage grouping:
- 6. Alpha fetoprotein (AFP) for pathological stage grouping (xx,xxx ng/mL):
- 7. Human chorionic gonadotropin (hCG) for pathological stage grouping (xx,xxx mIU/mI):
- 8. Lactate dehydrogenase (LDH) for pathological stage grouping (xx,xxx U/L):

8 Histologic Grade (G)

Germ cell tumors are not graded.

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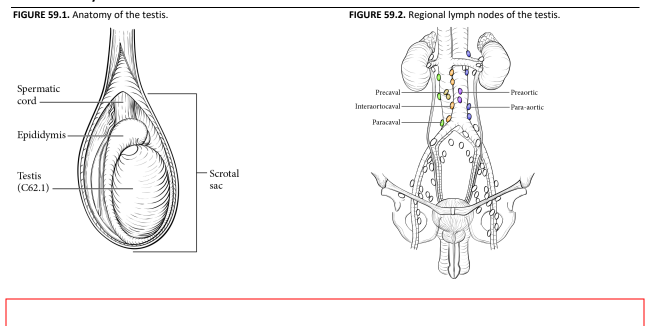
9 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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10 Anatomy

Physician Signature



Date/Time

| Hospital Name/Address | Patient Name/Information | Patient Name/Information | |
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | ntion Definition | | | | |
|--|---|--|--|--|--|--|
| | cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imagi endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or samplin regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | | | | |
| pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of informa diagnostic workup from clinical staging combined with operative findings, and pathology review of resected specimens | | | | | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | | | |
| | ypTNM Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoa therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | | | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | | | |

| Hospital Name/Address | Patient Name/Information | |
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| | ! | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|--|-------------------------------------|--|
| | TX Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤7 cm in greatest dimension, limited to the kidney |
| | T1a | Tumor ≤4 cm in greatest dimension, limited to the kidney |
| | T1b | Tumor >4 cm but ≤7 cm in greatest dimension limited to the kidney |
| T2 Tumor >7 cm in greatest dimension, limited to the kidney | | Tumor >7 cm in greatest dimension, limited to the kidney |
| T2a Tumor >7 cm but ≤10 cm in g | | Tumor >7 cm but ≤10 cm in greatest dimension, limited to the kidney |
| T2b Tumor >10 cm, limited to the kidney | | Tumor >10 cm, limited to the kidney |
| | T3 | Tumor extends into major veins or perinephric tissues, but not into the ipsilateral adrenal gland and not beyond Gerota's fascia |
| | T3a | Tumor extends into the renal vein or its segmental branches, or invades the pelvicalyceal system, or invades perirenal and/or renal sinus fat but not beyond Gerota's fascia |
| T3b Tumor extends into the vena cava below the diaphragm | | Tumor extends into the vena cava below the diaphragm |
| | T3c | Tumor extends into the vena cava above the diaphragm or invades the wall of the vena cava |
| T4 Tumor invades beyond Gerota's fascia (including contiguous extension into the ipsilateral adrenal gland | | Tumor invades beyond Gerota's fascia (including contiguous extension into the ipsilateral adrenal gland) |

| • | ✓ | T Suffix | Definition |
|---|---|----------|---|
| | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|--|--------------------------------------|--|
| | NX Regional lymph nodes cannot be assessed | | |
| | NO No regional lymph node metastasis | | |
| | N1 | Metastasis in regional lymph node(s) | |

| 1 | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | rtegory M Criteria | | |
|---|------------|---|--|--|
| | cM0 | No distant metastasis | | |
| | cM1 | Distant metastasis | | |
| | pM1 | Distant metastasis, microscopically confirmed | | |

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | NO | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | NX, NO | M0 | III |
| | T3 | N1 | M0 | III |
| | T4 | Any N | M0 | IV |
| | Any T | Any N | M1 | IV |

| 6 | Registry | Data | Collection | Variables |
|---|----------|------|------------|-----------|
|---|----------|------|------------|-----------|

| Histologic subtype: WHO/ISUP grade: Tumor size: | |
|--|--|
| | |
| Tumor size: | |
| | |
| Invasion into perinephric fat or sing | us tissues: |
| Venous involvement with specific renal vein involvement, or IVC invo | mention of intra-renal lymphovascular invasion, branches of renal vein in the renal sinus invasion, olvement: |
| Lymphovascular invasion (LVI): | |
| Adrenal gland involvement by dire | ct extension (T4) or as a separate nodule (M1): |
| Sarcomatoid features: | absent present; percentage: |
| Rhabdoid differentiation: | absent present |
| Histologic tumor necrosis: | |
| r L | Venous involvement with specific renal vein involvement, or IVC involvement, or IVC involvement with specific renal vein involvement, or IVC involvement by direction and involvement by direction and involvement with specific renal gland involvement by direction and involvement by direction and involvement with specific renal vein involvement, or IVC involvement with specific renal vein involvement with specif |

7 Histologic Grade (G)

The Fuhrman grading system, published in 1982, has been widely utilized. It is a four-tier system based on nuclear size, nuclear shape, and nucleolar prominence. Despite the widespread usage of Fuhrman grading, serious problems are associated with its implementation, reproducibility, and outcome prediction. As a result, a modified grading system has been proposed to be based on nucleolar prominence for the first three grading categories, while grade 4 is based on the presence of marked nuclear pleomorphism, which may include tumor giant cells or sarcomatoid and/or rhabdoid differentiation. Known as the WHO/ISUP grade, this grading system was validated for clear cell and papillary RCC, but was shown not to be useful for chromophobe RCC and has not been validated in other RCC histologic subtypes.

| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Nucleoli absent or inconspicuous and basophilic at 400x magnification |
| | G2 | Nucleoli conspicuous and eosinophilic at 400x magnification, visible but not prominent at 100x magnification |
| | G3 | Nucleoli conspicuous and eosinophilic at 100x magnification |
| | G4 | Marked nuclear pleomorphism and/or multinucleate giant cells and/or rhabdoid and/or sarcomatoid differentiation |

| Hospital Name/Address | Patient Name/Information |
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8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 60.1. Anatomic sites and subsites of the kidney.

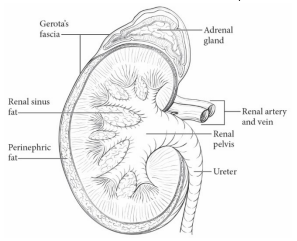


FIGURE 60.2. Regional lymph nodes of the kidney.

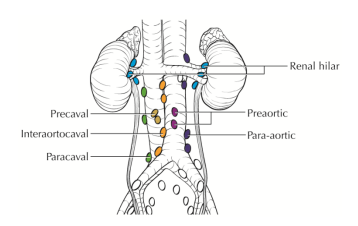
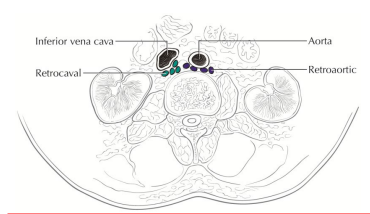


FIGURE 60.3. Regional lymph nodes of the kidney.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information | |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Papillary noninvasive carcinoma |
| | Tis | Carcinoma in situ |
| | T1 | Tumor invades subepithelial connective tissue |
| | T2 | Tumor invades the muscularis |
| | T3 | For renal pelvis only: Tumor invades beyond muscularis into peripelvic fat or into the renal parenchyma |
| | | For ureter only: Tumor invades beyond muscularis into periureteric fat |
| | T4 | Tumor invades adjacent organs, or through the kidney into the perinephric fat |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis ≤2 cm in greatest dimension, in a single lymph node |
| | N2 | Metastasis >2 cm, in a single lymph node; or multiple lymph nodes |

| ✓ | N Suffix | Definition | |
|----------|----------|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Та | N0 | M0 | 0a |
| | Tis | N0 | M0 | Ois |
| | T1 | N0 | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | III |
| | T4 | NX, NO | M0 | IV |
| | Any T | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Presence or absence of extranodal extension:
- 2. Size of the largest tumor deposit in the lymph nodes:
- 3. Total number of lymph nodes dissected:
- 4. Presence of urothelial carcinoma in situ (Tis) with other tumors:
- 5. Presence of papillary noninvasive carcinoma (Ta) with other tumors:
- 6. Lymphovascular invasion:
- 7. World Health Organization/International Society of Urologic Pathology (WHO/ISUP) grade:
- 8. Grade 1–3 for squamous and adenocarcinoma:
- 9. Intratubular spread of Tis urothelial carcinoma (involvement of renal collecting tubules without stromal invasion):

7 Histologic Grade (G)

For urothelial histologies, a low- and high-grade designation is used to match the current WHO/ISUP recommended grading system.

| ✓ | G | G Definition (Urothelial Carcinoma) |
|---|----|-------------------------------------|
| | LG | Low grade |
| | HG | High grade |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 60.1. Anatomic sites and subsites of the kidney.

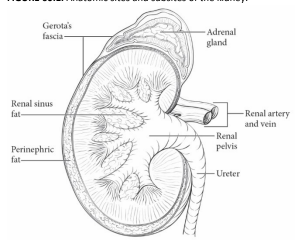


FIGURE 61.1. The regional lymph nodes of the renal pelvis.

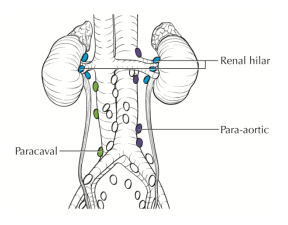
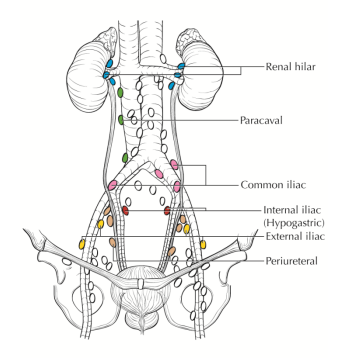


FIGURE 61.2. The regional lymph nodes of the ureter.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Papillary noninvasive carcinoma |
| | Tis | Carcinoma in situ |
| | T1 | Tumor invades subepithelial connective tissue |
| | T2 | Tumor invades the muscularis |
| | T3 | For renal pelvis only: Tumor invades beyond muscularis into peripelvic fat or into the renal parenchyma |
| | | For ureter only: Tumor invades beyond muscularis into periureteric fat |
| | T4 | Tumor invades adjacent organs, or through the kidney into the perinephric fat |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis ≤2 cm in greatest dimension, in a single lymph node |
| | N2 | Metastasis >2 cm, in a single lymph node; or multiple lymph nodes |

| ✓ | N Suffix | Definition | |
|----------|--|---|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|----------|-----------|----------|----------|-------------------------|
| | Ta | N0 | M0 | 0a |
| | Tis | N0 | M0 | Ois |
| | T1 | NO | M0 | 1 |
| | T2 | NO | M0 | II |
| | T3 | N0 | M0 | III |
| | T4 | NX, NO | M0 | IV |
| | Any T | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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61.2 Renal Pelvis and Ureter: Squamous Cell Carcinoma and Adenocarcinoma

| 6 Registry Data Collection Var | riables |
|--------------------------------|---------|
|--------------------------------|---------|

See chapter for more details on these variables.

- 1. Presence or absence of extranodal extension:
- 2. Size of the largest tumor deposit in the lymph nodes:
- 3. Total number of lymph nodes dissected:
- 4. Presence of urothelial carcinoma in situ (Tis) with other tumors:
- 5. Presence of papillary noninvasive carcinoma (Ta) with other tumors:
- 6. Lymphovascular invasion:
- 7. World Health Organization/International Society of Urologic Pathology (WHO/ISUP) grade:
- 8. Grade 1–3 for squamous and adenocarcinoma:
- 9. Intratubular spread of Tis urothelial carcinoma (involvement of renal collecting tubules without stromal invasion):

7 Histologic Grade (G)

For squamous cell carcinoma and adenocarcinoma, the following grading schema is recommended.

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 60.1. Anatomic sites and subsites of the kidney.

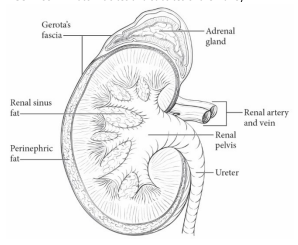


FIGURE 61.1. The regional lymph nodes of the renal pelvis.

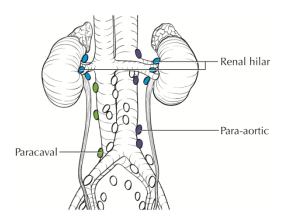
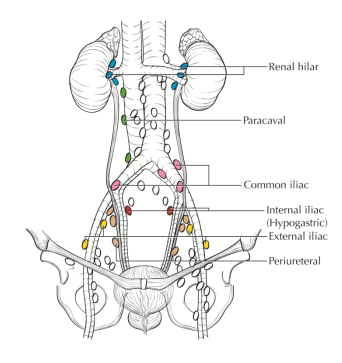


FIGURE 61.2. The regional lymph nodes of the ureter.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | ation Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Non-invasive papillary carcinoma |
| | Tis | Urothelial carcinoma in situ: "flat tumor" |
| | T1 | Tumor invades lamina propria (subepithelial connective tissue) |
| | T2 | Tumor invades muscularis propria |
| | pT2a | Tumor invades superficial muscularis propria (inner half) |
| | pT2b | Tumor invades deep muscularis propria (outer half) |
| | T3 | Tumor invades perivesical soft tissue |
| | pT3a | Tumor invades perivesical soft tissue microscopically |
| | pT3b | Tumor invades perivesical soft tissue macroscopically (extravesical mass) |
| | T4 | Extravesical tumor directly invades any of the following: prostatic stroma, seminal vesicles, uterus, vagina, pelvic |
| | | wall, abdominal wall |
| | T4a | Extravesical tumor invades directly into prostatic stroma, seminal vesicles, uterus, vagina |
| | T4b | Extravesical tumor invades pelvic wall, abdominal wall |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Ī | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Lymph nodes cannot be assessed |
| | N0 | No lymph node metastasis |
| | N1 | Single regional lymph node metastasis in the true pelvis (perivesical, obturator, internal and external iliac, or sacral lymph node) |
| | N2 | Multiple regional lymph node metastasis in the true pelvis (perivesical, obturator, internal and external iliac, or sacral lymph node metastasis) |
| | N3 | Lymph node metastasis to the common iliac lymph nodes |

| 1 | N Suffix | Definition |
|---|--|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Distant metastasis limited to lymph nodes beyond the common iliacs |
| | cM1b | Non-lymph-node distant metastases |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Distant metastasis limited to lymph nodes beyond the common iliacs, microscopically confirmed |
| | pM1b | Non-lymph-node distant metastases, microscopically confirmed |

${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|---------------|----------|----------|-------------------------|
| | Та | N0 | M0 | 0a |
| | Tis | NO | M0 | Ois |
| | T1 | N0 | M0 | 1 |
| | T2a | N0 | M0 | II |
| | T2b | NO | M0 | II |
| | T3a, T3b, T4a | N0 | M0 | IIIA |
| | T1 – T4a | N1 | M0 | IIIA |
| | T1 – T4a | N2, N3 | M0 | IIIB |
| | T4b | Any N | M0 | IVA |
| | Any T | Any N | M1a | IVA |
| | Any T | Any N | M1b | IVB |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Presence or absence of extranodal extension:
- 2. Total number of lymph nodes examined pathologically and total number positive:
- 3. Size of the largest tumor deposit in the lymph nodes:
- 4. World Health Organization/International Society of Urologic Pathology (WHO/ISUP) grade:
- 5. Presence of lymphovascular invasion:
- 6. Concurrent/associated noninvasive papillary (Ta) with carcinoma in situ (Tis):
- 7. Concurrent/associated noninvasive papillary (Ta) and/or carcinoma in situ (Tis) with invasive cancers:

7 Histologic Grade (G)

For urothelial histologies, a low- and high-grade designation is used to match the current WHO/ISUP recommended grading system

| | To distinct in the second and make a second in the second | |
|---|---|--------------|
| 1 | G | G Definition |
| | LG | Low-grade |
| | HG | High-grade |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description | |
|---|--------------|--|--|
| • | LVI Coding | | |
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 62.1. Extent of primary bladder cancer.

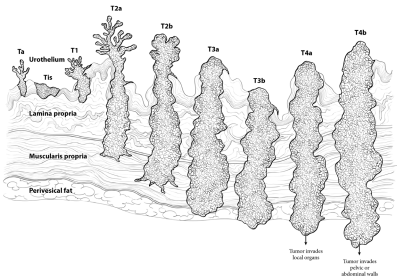
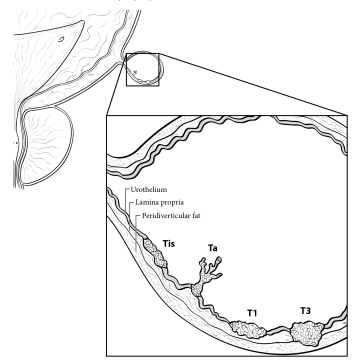


FIGURE 62.2. Extent of Tis, Ta, T1, and T3.



| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information | |
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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|----------------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Non-invasive papillary carcinoma |
| | Tis | Urothelial carcinoma in situ: "flat tumor" |
| | T1 | Tumor invades lamina propria (subepithelial connective tissue) |
| | T2 | Tumor invades muscularis propria |
| | pT2a | Tumor invades superficial muscularis propria (inner half) |
| | pT2b | Tumor invades deep muscularis propria (outer half) |
| | T3 | Tumor invades perivesical soft tissue |
| | pT3a | Tumor invades perivesical soft tissue microscopically |
| | pT3b | Tumor invades perivesical soft tissue macroscopically (extravesical mass) |
| | T4 | Extravesical tumor directly invades any of the following: prostatic stroma, seminal vesicles, uterus, vagina, pelvic |
| | wall, abdominal wall | |
| | T4a | Extravesical tumor invades directly into prostatic stroma, seminal vesicles, uterus, vagina |
| | T4b | Extravesical tumor invades pelvic wall, abdominal wall |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Lymph nodes cannot be assessed |
| | N0 | No lymph node metastasis |
| | N1 | Single regional lymph node metastasis in the true pelvis (perivesical, obturator, internal and external iliac, or sacral lymph node) |
| | N2 | Multiple regional lymph node metastasis in the true pelvis (perivesical, obturator, internal and external iliac, or sacral lymph node metastasis) |
| | N3 | Lymph node metastasis to the common iliac lymph nodes |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | cM1a | Distant metastasis limited to lymph nodes beyond the common iliacs |
| | cM1b | Non-lymph-node distant metastases |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Distant metastasis limited to lymph nodes beyond the common iliacs, microscopically confirmed |
| | pM1b | Non-lymph-node distant metastases, microscopically confirmed |

${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|---------------|----------|----------|-------------------------|
| | Та | N0 | M0 | 0a |
| | Tis | NO | M0 | Ois |
| | T1 | N0 | M0 | 1 |
| | T2a | N0 | M0 | II |
| | T2b | NO | M0 | II |
| | T3a, T3b, T4a | N0 | M0 | IIIA |
| | T1 – T4a | N1 | M0 | IIIA |
| | T1 – T4a | N2, N3 | M0 | IIIB |
| | T4b | Any N | M0 | IVA |
| | Any T | Any N | M1a | IVA |
| | Any T | Any N | M1b | IVB |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Presence or absence of extranodal extension:
- 2. Total number of lymph nodes examined pathologically and total number positive:
- 3. Size of the largest tumor deposit in the lymph nodes:
- 4. World Health Organization/International Society of Urologic Pathology (WHO/ISUP) grade:
- 5. Presence of lymphovascular invasion:
- 6. Concurrent/associated noninvasive papillary (Ta) with carcinoma in situ (Tis):
- 7. Concurrent/associated noninvasive papillary (Ta) and/or carcinoma in situ (Tis) with invasive cancers:

7 Histologic Grade (G)

For squamous cell carcinoma and adenocarcinoma, the following grading schema is recommended:

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 62.1. Extent of primary bladder cancer.

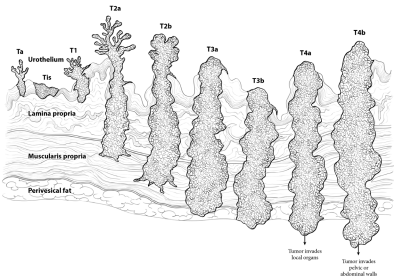
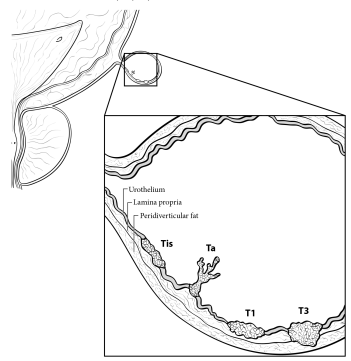


FIGURE 62.2. Extent of Tis, Ta, T1, and T3.



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|---------------------|-----------|
| Physician Signature | Date/Time |

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Non-invasive papillary carcinoma |
| | Tis | Carcinoma in situ |
| | T1 | Tumor invades subepithelial connective tissue |
| | T2 | Tumor invades any of the following: corpus spongiosum, periurethral muscle |
| | T3 | Tumor invades any of the following: corpus cavernosum, anterior vagina |
| | T4 | Tumor invades other adjacent organs (e.g., invasion of the bladder wall) |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Single regional lymph node metastasis in the inguinal region or true pelvis [perivesical, obturator, internal |
| | | (hypogastric) and external iliac], or presacral lymph node |
| | N2 | Multiple regional lymph node metastasis in the inguinal region or true pelvis (perivesical, hypogastric, obturator, |
| | | internal and external iliac, or presacral lymph node) |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| 1 | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | Ois |
| | Ta | N0 | M0 | 0a |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | N0 | M0 | III |
| | T3 | N1 | M0 | III |
| | T4 | NX | M0 | IV |
| | T4 | N0 | M0 | IV |
| | T4 | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. WHO/ISUP Grade:
- 2. Grade 1–3 for squamous cell carcinoma and adenocarcinoma:

7 Histologic Grade (G)

Grade is reported by the grade value. For urothelial histology, a low- and high-grade designation is used to match the current WHO/ISUP recommended grading system:

| ✓ | G | G Definition |
|---|----|--------------|
| | LG | Low grade |
| | HG | High grade |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

FIGURE 63.3. Female Urethra. Definition of primary tumor (T) for Ta, Tis, T1, and T2 with depth of invasion ranging from the epithelium to the urogenital diaphragm.

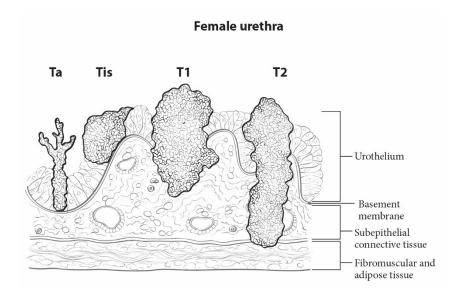
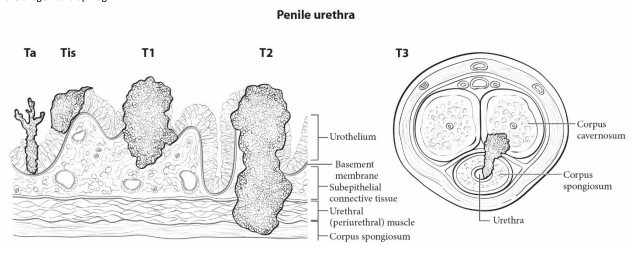
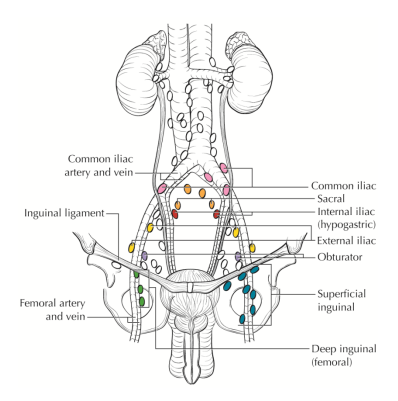


FIGURE 63.2. Penile Urethra. Definition of primary tumor (T) for Ta, Tis, T1, T2, and T3 with depth of invasion ranging from the epithelium to the urogenital diaphragm.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 63.1. Regional lymph nodes of the urethra.



| Physician Signature | Date/Time |
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| Patient Name/Information | |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Non-invasive papillary carcinoma |
| | Tis | Carcinoma in situ |
| | T1 | Tumor invades subepithelial connective tissue |
| | T2 | Tumor invades any of the following: corpus spongiosum, periurethral muscle |
| | T3 | Tumor invades any of the following: corpus cavernosum, anterior vagina |
| | T4 | Tumor invades other adjacent organs (e.g., invasion of the bladder wall) |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria |
|----------|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Single regional lymph node metastasis in the inguinal region or true pelvis [perivesical, obturator, internal |
| | | (hypogastric) and external iliac], or presacral lymph node |
| | N2 | Multiple regional lymph node metastasis in the inguinal region or true pelvis (perivesical, hypogastric, obturator, |
| | | internal and external iliac, or presacral lymph node) |

| | ✓ | N Suffix | Definition | |
|---|---|--|--|--|
| Г | | (sn) | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| Г | | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information | |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | Ois |
| | Ta | NO | M0 | 0a |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | NO | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | N0 | M0 | III |
| | T3 | N1 | M0 | III |
| | T4 | NX | M0 | IV |
| | T4 | N0 | M0 | IV |
| | T4 | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. WHO/ISUP Grade:
- 2. Grade 1–3 for squamous cell carcinoma and adenocarcinoma:

7 Histologic Grade (G)

For squamous cell carcinoma and adenocarcinoma, the following grading schema is recommended:

| 1 | G | G Definition | |
|---|----|---------------------------|--|
| | GX | Grade cannot be assessed | |
| | G1 | Well differentiated | |
| | G2 | Moderately differentiated | |
| | G3 | Poorly differentiated | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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9 Anatomy

FIGURE 63.3. Female Urethra. Definition of primary tumor (T) for Ta, Tis, T1, and T2 with depth of invasion ranging from the epithelium to the urogenital diaphragm.

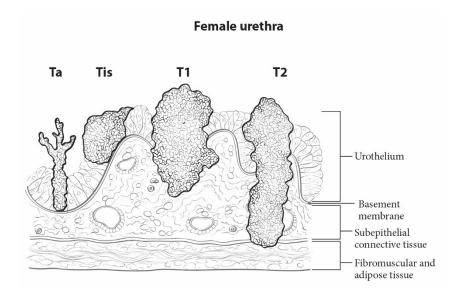
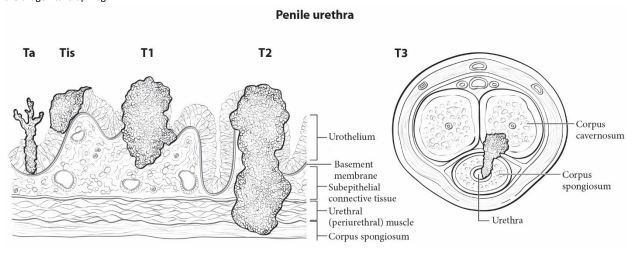
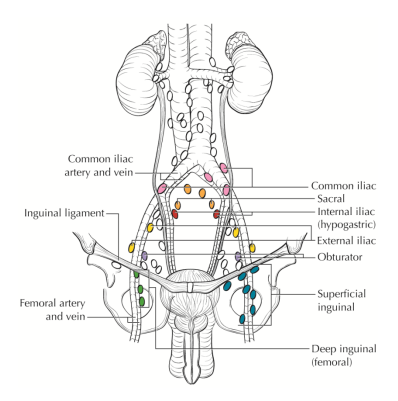


FIGURE 63.2. Penile Urethra. Definition of primary tumor (T) for Ta, Tis, T1, T2, and T3 with depth of invasion ranging from the epithelium to the urogenital diaphragm.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 63.1. Regional lymph nodes of the urethra.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification Definition | |
|---|---------------------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|-------------------------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Та | Non-invasive papillary carcinoma |
| | Tis | Carcinoma in situ involving the prostatic urethra or periurethral or prostatic ducts without stromal invasion |
| | T1 | Tumor invades urethral subepithelial connective tissue immediately underlying the urothelium |
| | T2 | Tumor invades the prostatic stroma surrounding ducts either by direct extension from the urothelial surface or by |
| | invasion from prostatic ducts | |
| | T3 | Tumor invades the periprostatic fat |
| | T4 | Tumor invades other adjacent organs (e.g., extraprostatic invasion of the bladder wall, rectal wall) |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|--|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | NO No regional lymph node metastasis | |
| | N1 | Single regional lymph node metastasis in the inguinal region or true pelvis [perivesical, obturator, internal | |
| | (hypogastric) and external iliac], or presacral lymph node | | |
| | N2 Multiple regional lymph node metastasis in the inguinal region or true pelvis (perivesical, hypogastric, obturator, | | |
| | | internal and external iliac, or presacral lymph node) | |

| ✓ | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | NO | M0 | Ois |
| | Ta | N0 | M0 | 0a |
| | T1 | NO | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | NO | M0 | III |
| | T3 | N1 | M0 | III |
| | T4 | NX | M0 | IV |
| | T4 | NO | M0 | IV |
| | T4 | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. WHO/ISUP Grade:
- 2. Grade 1–3 for squamous cell carcinoma and adenocarcinoma:

7 Histologic Grade (G)

Grade is reported by the grade value. For urothelial histology, a low- and high-grade designation is used to match the current WHO/ISUP recommended grading system:

| ✓ | G | G Definition |
|---|----|--------------|
| | LG | Low grade |
| | HG | High grade |

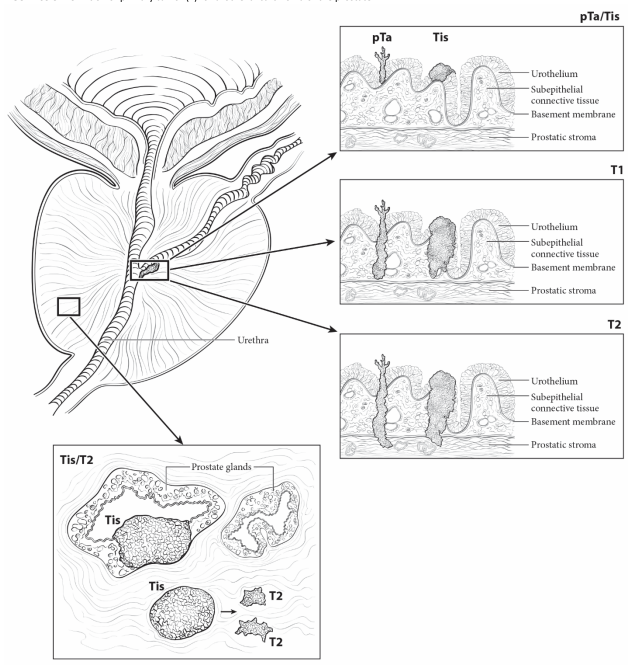
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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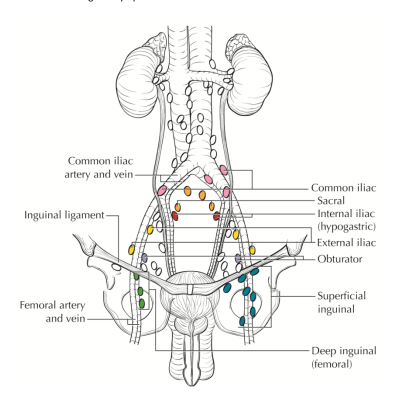
9 Anatomy

FIGURE 63.5. Definition of primary tumor (T) for urothelial carcinoma of the prostate.



| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

FIGURE 63.1. Regional lymph nodes of the urethra.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria | |
|---|------------|--|--|
| | TX | Primary tumor cannot be assessed | |
| | TO | No evidence of primary tumor | |
| | Та | Non-invasive papillary carcinoma | |
| | Tis | Carcinoma <i>in situ</i> involving the prostatic urethra or periurethral or prostatic ducts without stromal invasion | |
| | T1 | Tumor invades urethral subepithelial connective tissue immediately underlying the urothelium | |
| | T2 | Tumor invades the prostatic stroma surrounding ducts either by direct extension from the urothelial surface or by | |
| | | invasion from prostatic ducts | |
| | T3 | Tumor invades the periprostatic fat | |
| | T4 | Tumor invades other adjacent organs (e.g., extraprostatic invasion of the bladder wall, rectal wall) | |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node metastasis | |
| | N1 | Single regional lymph node metastasis in the inguinal region or true pelvis [perivesical, obturator, internal | |
| | | (hypogastric) and external iliac], or presacral lymph node | |
| | N2 | Multiple regional lymph node metastasis in the inguinal region or true pelvis (perivesical, hypogastric, obturator, | |
| | | internal and external iliac, or presacral lymph node) | |

| | ✓ | N Suffix | Definition | |
|---|---|--|---|--|
| Γ | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category M Criteria | |
|---|------------------------|---|
| | cM0 | No distant metastasis |
| | cM1 Distant metastasis | |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
| | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | Tis | N0 | M0 | Ois |
| | Ta | NO | M0 | 0a |
| | T1 | N0 | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | NO | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | N0 | M0 | III |
| | T3 | N1 | M0 | III |
| | T4 | NX | M0 | IV |
| | T4 | N0 | M0 | IV |
| | T4 | N1 | M0 | IV |
| | Any T | N2 | M0 | IV |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. WHO/ISUP Grade:
- 2. Grade 1–3 for squamous cell carcinoma and adenocarcinoma:

7 Histologic Grade (G)

For squamous cell carcinoma and adenocarcinoma, the following grading schema is recommended:

| 1 | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |

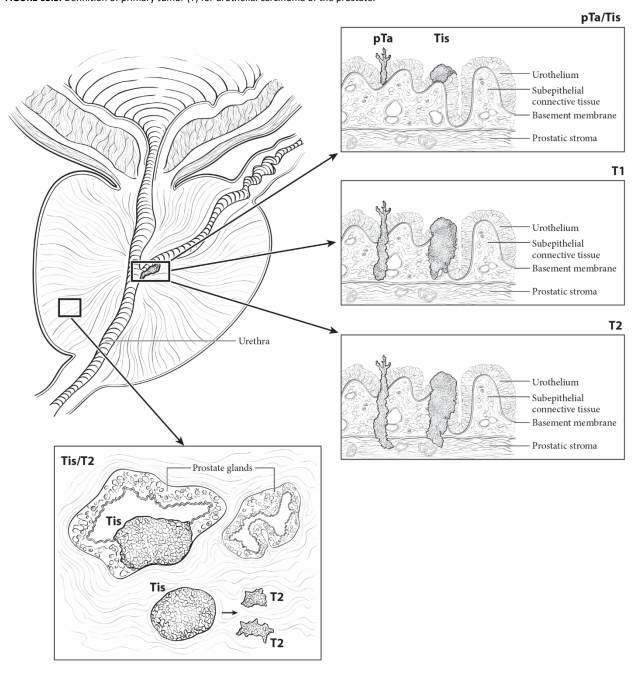
8 Lymphovascular Invasion (LVI)

| 1 | Component of LVI Coding | Description | |
|---|-------------------------|--|--|
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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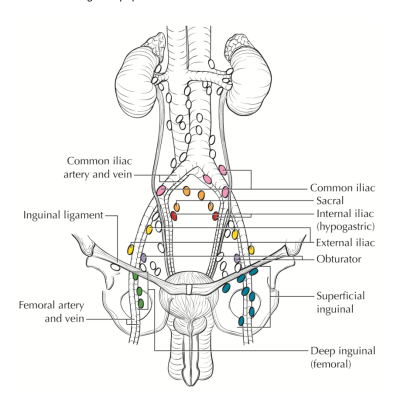
9 Anatomy

FIGURE 63.5. Definition of primary tumor (T) for urothelial carcinoma of the prostate.



| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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| | |

FIGURE 63.1. Regional lymph nodes of the urethra.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | Patient Name/Information | | |
|--------------------------|-----------------------------|--|--|
| | | | |
| | | | |
| | | | |
| | Patient Name/illionillation | | |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | | |
|-----------------------|--------------------------|--|--|
| | | | |
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| | | | |
| | ! | | |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---|--|--|
| | TX | Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | Tis | Carcinoma in situ | |
| | T1 | Tumor ≤10 mm in greatest dimension | |
| | T1a | Tumor does not invade the tarsal plate or eyelid margin | |
| | T1b | Tumor invades the tarsal plate or eyelid margin | |
| | T1c | Tumor involves full thickness of the eyelid | |
| | T2 | Tumor >10 mm but ≤20 mm in greatest dimension | |
| T2a Tumor does not invade the tarsal plate or eyelid margin | | Tumor does not invade the tarsal plate or eyelid margin | |
| T2b Tumor invades the tarsal plate or eyelid margin | | Tumor invades the tarsal plate or eyelid margin | |
| | T2c | Tumor involves full thickness of the eyelid | |
| | T3 | Tumor >20 mm but ≤30 mm in greatest dimension | |
| | T3a | Tumor does not invade the tarsal plate or eyelid margin | |
| | T3b | Tumor invades the tarsal plate or eyelid margin | |
| T3c Tumor involves full thickness of the eyelid | | Tumor involves full thickness of the eyelid | |
| T4 Any eyelid tumor that invades adjacent ocular, orbital, or facial structures | | Any eyelid tumor that invades adjacent ocular, orbital, or facial structures | |
| | T4a Tumor invades ocular or intraorbital structures | | |
| | T4b | Tumor invades (or erodes through) the bony walls of the orbit or extends to the paranasal sinuses or invades the | |
| lacrimal sac/nasolacrimal duct or brain | | | |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | | |
|---|---|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | | |
| | N0 | No evidence of lymph node involvement | | |
| | N1 | Metastasis in a single ipsilateral regional lymph node, ≤3 cm in greatest dimension | | |
| | N1a | Metastasis in a single ipsilateral lymph node based on clinical evaluation or imaging findings | | |
| | N1b | Metastasis in a single ipsilateral lymph node based on lymph node biopsy | | |
| | N2 Metastasis in a single ipsilateral lymph node, >3 cm in greatest dimension, or in bilateral or contralateral lym | | | |
| | nodes | | | |
| | N2a | Metastasis documented based on clinical evaluation or imaging findings | | |
| | N2b | Metastasis documented based on microscopic findings on lymph node biopsy | | |

| L | ✓ | N Suffix | Definition |
|---|---|----------|--|
| Ī | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| I | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | 1 Criteria | |
|----------|---|------------|--|
| | cM0 No distant metastasis | | |
| | cM1 Distant metastasis | | |
| | pM1 Distant metastasis, microscopically confirmed | | |

| Hospital Name/Address | Patient Name/Information | | |
|-----------------------|--------------------------|--|--|
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is And N is And M is | | And M is | Then the stage group is | | |
|---|-----------------------------|-------|----------|-------------------------|--|--|
| | Tis | NO | M0 | 0 | | |
| | T1 | N0 | M0 | IA | | |
| | T2a | N0 | M0 | IB | | |
| | T2b-c, T3 | N0 | M0 | IIA | | |
| | T4 | N0 | M0 | IIB | | |
| | Any T | N1 | M0 | IIIA | | |
| | Any T | N2 | M0 | IIIB | | |
| | Any T | Any N | M1 | IV | | |

6 Registry Data Collection Variables

See chapter for more details on these variables.

| 1. | Tumor size (greatest dimension in millimeters): |
|----|---|
| | (8 |

| 2. | Specific anatomic | location (e.g., | upper eyelid, | lower eyelid, bo | th eyelids, | medial canthus, | lateral canthus): |
|----|-------------------|-----------------|---------------|------------------|-------------|-----------------|-------------------|
|----|-------------------|-----------------|---------------|------------------|-------------|-----------------|-------------------|

| 3. Tu | umor thickness | (depth | of invasion) |
|-------|----------------|--------|--------------|
|-------|----------------|--------|--------------|

| 4 | D | / | _ £: | 1: |
|----|-----------|------------|--------------|------------|
| 4. | Presence. | /absence (| of perineura | i invasion |

| | 5. | Presence | /absence of I | lymphovascular i | nvasion |
|--|----|----------|---------------|------------------|---------|
|--|----|----------|---------------|------------------|---------|

| 6. | Mitotic | figures | ner | square | millimete | ٩r |
|----|---------|---------|-----|--------|-----------|----|
| | | | | | | |

| 8. | Sentinel | node hio | nsv status a | nd number | of sentinel | nodes (i | if applicable): |
|----|----------|----------|--------------|-----------|--------------|----------|--------------------------|
| ο. | Sentine | HOUE DIO | usv status a | na namber | OI SEIIUIIEI | HOUES H | ii abbiicabie <i>i</i> . |

| 9. | History | of HIV | infection |
|----|---------|--------|-----------|
| | | | |

| 10. | History | of solid | organ | transp | lant |
|-----|---------|----------|-------|--------|------|
|-----|---------|----------|-------|--------|------|

| 11 | History | of Mu | ir–Torre | syndron | no. |
|-----|---------|---------|----------|----------|-----|
| TT. | HISLOIV | OI IVIU | | Syllulul | пe. |

| 12. | History of xeroderm | a pigmentosum |
|-----|---------------------|---------------|
|-----|---------------------|---------------|

7 Histologic Grade (G)

A histologic grading system is used predominantly for SCCs and sebaceous carcinomas. It is not used for Merkel cell carcinoma or BCC.

| √ | G | G Definition |
|----------|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

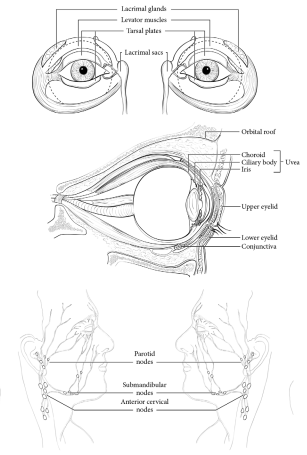
| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 64.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | Hospital Name/Address |
|--------------------------|-----------------------|
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
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| | |
| | ! |

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | Tis | Carcinoma in situ |
| | T1 | Tumor (≤5 mm in greatest dimension) invades through the conjunctival basement membrane without invasion of adjacent structures |
| | T2 | Tumor (>5 mm in greatest dimension) invades through the conjunctival basement membrane without invasion of adjacent structures |
| | T3 | Tumor invades adjacent structures (excluding the orbit) |
| | T4 | Tumor invades the orbit with or without further extension |
| | T4a | Tumor invades orbital soft tissues without bone invasion |
| | T4b | Tumor invades bone |
| | T4c | Tumor invades adjacent paranasal sinuses |
| | T4d | Tumor invades brain |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| | F | |
|---|------------|---|
| 1 | M Category | M Criteria |
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no proposal for anatomic stage and prognostic groups for conjunctival carcinoma. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| Hospital Name/Address | Patient Name/Information |
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6 Registry Data Collection Variables

See chapter for more details on these variables.

1. Ki-67 growth fraction, reported as percentage of positive tumor cells by immunohistochemistry:

7 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|---------------------------|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated |
| | G3 | Poorly differentiated |
| | G4 | Undifferentiated |

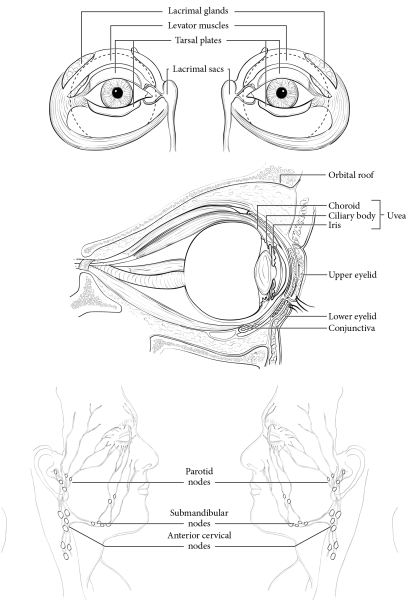
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description | |
|---|--------------|--|--|
| | LVI Coding | | |
| | 0 | LVI not present (absent)/not identified | |
| | 1 | LVI present/identified, NOS | |
| | 2 | Lymphatic and small vessel invasion only (L) | |
| | 3 | Venous (large vessel) invasion only (V) | |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion | |
| | 9 | Presence of LVI unknown/indeterminate | |

| Hospital Name/Address | Patient Name/Information |
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9 Anatomy

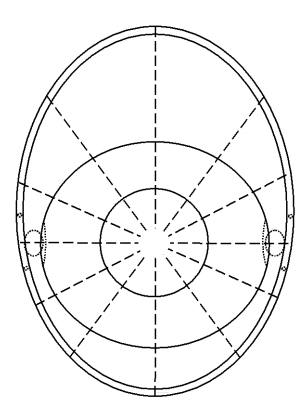
FIGURE 65.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



This form continues on the next page.

| Hospital Name/Address | Patient Name/Information | |
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FIGURE 65.2. Clinical mapping system for conjunctival carcinoma. The map displays the entire conjunctiva as a flat surface, with the central point located at the center of the cornea and concentric regions, such as the limbus, bulbar conjunctiva, fornix, palpebral conjunctiva, and eyelid, considered progressively more peripheral. Radial lines represent clock hours (Modified from Damato and Coupland).



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| | Patient Name/Information | |

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification Definition | |
|---|---------------------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

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| Hospital Name/Address | Patient Name/Information | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

4.1.1 Clinical T (cT)

| 1 | cT Category | cT Criteria |
|---|-------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor of the bulbar conjunctiva |
| | T1a | <1 quadrant |
| | T1b | ≥1 to <2 quadrants |
| | T1c | ≥2 to <3 quadrants |
| | T1d | ≥3 quadrants |
| | T2 | Tumor of the nonbulbar (forniceal, palpebral, tarsal) conjunctiva, and tumor involving the caruncle |
| | T2a | Noncaruncular, and ≤1 quadrant of the nonbulbar conjunctiva involved |
| | T2b | Noncaruncular, and >1 quadrant of the nonbulbar conjunctiva involved |
| | T2c | Caruncular, and ≤1 quadrant of the nonbulbar conjunctiva involved |
| | T2d | Caruncular, and >1 quadrant of the nonbulbar conjunctiva involved |
| | T3 | Tumor of any size with local invasion |
| | T3a | Globe |
| | T3b | Eyelid |
| | T3c | Orbit |
| | T3d | Nasolacrimal duct and/or lacrimal sac and/or paranasal sinuses |
| | T4 | Tumor of any size with invasion of the central nervous system |

| | ✓ | T Suffix | Suffix Definition | |
|---|---|----------|-------------------|--|
| Ī | (m) Select if synchronous primary tumors are found in single organ. | | | |

4.1.2 Pathological T (pT)

| ✓ | pT Category | pT Criteria |
|---|-------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | Tis | Melanoma confined to the conjunctival epithelium |
| | T1 | Tumor of the bulbar conjunctiva |
| | T1a | Tumor of the bulbar conjunctiva with invasion of the substantia propria, not more than 2.0 mm in thickness |
| | T1b | Tumor of the bulbar conjunctiva with invasion of the substantia propria, more than 2.0 mm in thickness |
| | T2 | Tumor of the nonbulbar (forniceal, palpebral, tarsal) conjunctiva, and tumor involving the caruncle |
| | T2a | Tumor of the nonbulbar conjunctiva with invasion of the substantia propria, not more than 2.0 mm in thickness |
| | T2b | Tumor of the nonbulbar conjunctiva with invasion of the substantia propria, more than 2.0 mm in thickness |
| | T3 | Tumor of any size with local invasion |
| | T3a | Globe |
| | T3b | Eyelid |
| | T3c | Orbit |
| | T3d | Nasolacrimal duct and/or lacrimal sac and/or paranasal sinuses |
| | T4 | Tumor of any size with invasion of the central nervous system |

| ✓ | T Suffix Definition | |
|---|---|--|
| | (m) Select if synchronous primary tumors are found in single organ. | |

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| Hospital Name/Address | Patient Name/Information | |
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4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition |
|---|--|---|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system.

Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no proposal for anatomic stage and prognostic groups for conjunctival melanoma.

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Tumor thickness: infiltration depth (measured in millimeters) into the substantia propria from the surface of the conjunctival epithelium:
- 2. Cytomorphology presence/absence of epithelioid cells:
- 3. Mitotic count number of mitosis per square millimiter:
- 4. Presence/absence of surface ulceration:
- 5. Presence/absence of growth regression:
- 6. Presence/absence of vessel invasion blood or lymphatic invasion:
- 7. Presence/absence of perineural invasion:
- 8. Status of all surgical margins (i.e., whether tumor extends to the lateral and deep margins):
- 9. Presence/absence of adjacent conjunctival melanoma in situ, including status within surgical margins:
- 10. Presence/absence of coexisting nevus:
- 11. Presence/absence of microsatellites:
- 12. The presence or absence of microscopic satellites/satellite in-transit metastases, which may be considered for future pathologic staging of pN level, as in the case of cutaneous melanoma*:

| Hospital Name/Address | Patient Name/Information |
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^{*}Satellite in-transit metastasis: discrete micronodule/nodule of melanoma <1 mm to several millimeters in diameter in subepithelial tissue close to but clearly separated from the primary melanoma by at least 1 to 2 mm or more of uninvolved connective tissue. Both these types of metastasis usually are angiotropic and may be solitary or often multiple.

7 Histologic Grade (G)

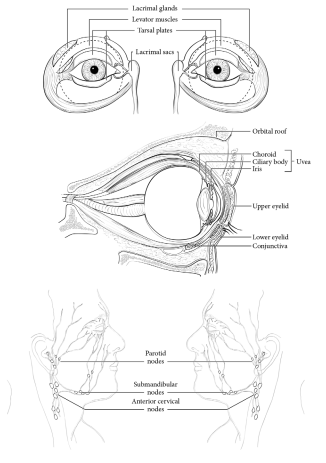
In accordance with melanomas at other anatomic sites, grading is not performed for conjunctival melanoma.

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

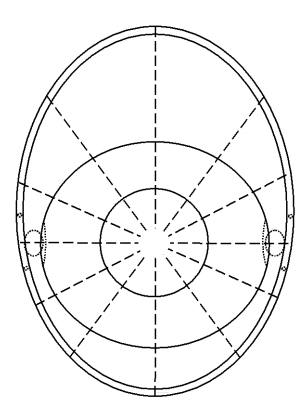
FIGURE 66.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



This form continues on the next page.

| Hospital Name/Address | Patient Name/Information | |
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FIGURE 66.2. Clinical mapping system for conjunctival melanoma. The map displays the entire conjunctiva as a flat surface, with the central point located at the center of the cornea and concentric regions such as the limbus, bulbar conjunctiva, fornix, palpebral conjunctiva, and eyelid considered progressively more peripheral. Radial lines represent clock hours (Modified from Damato and Coupland).



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| Physician Signature | Date/Time |
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| Hospital Name/Address | Patient Name/Information | |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|--|----------------|---|--|
| workup information, until first treatment, including clinical history and symptoms, physical examinati endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | M Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|------------|--|--|
| | TX | Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | T1 | Tumor limited to the iris | |
| | T1a | Tumor limited to the iris, not more than 3 clock hours in size | |
| | T1b | Tumor limited to the iris, more than 3 clock hours in size | |
| | T1c | Tumor limited to the iris with secondary glaucoma | |
| | T2 | Tumor confluent with or extending into the ciliary body, choroid, or both | |
| | T2a | Tumor confluent with or extending into the ciliary body, without secondary glaucoma | |
| | T2b | Tumor confluent with or extending into the ciliary body and choroid, without secondary glaucoma | |
| | T2c | Tumor confluent with or extending into the ciliary body, choroid, or both, with secondary glaucoma | |
| | T3 | Tumor confluent with or extending into the ciliary body, choroid, or both, with scleral extension | |
| | T4 | Tumor with extrascleral extension | |
| | T4a | Tumor with extrascleral extension ≤5 mm in largest diameter | |
| | T4b | Tumor with extrascleral extension >5 mm in largest diameter | |

Note: Iris melanomas originate from, and are predominantly located in, this region of the uvea. If less than half the tumor volume is located within the iris, the tumor may have originated in the ciliary body, and consideration should be given to classifying it accordingly.

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria | |
|----------|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node involvement | |
| | N1 | Regional lymph node metastases or discrete tumor deposits in the orbit | |
| | N1a | Metastasis in one or more regional lymph node(s) | |
| | N1b | No regional lymph nodes are positive, but there are discrete tumor deposits in the orbit that are not contiguous to | |
| | | the eye. (choroidal and ciliary body melanoma only) | |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|--|
| | cM0 | No distant metastasis by clinical classification |
| | cM1 | Distant metastasis |
| | cM1a | Largest diameter of the largest metastasis ≤3.0 cm |
| | cM1b | Largest diameter of the largest metastasis 3.1–8.0 cm |
| | cM1c | Largest diameter of the largest metastasis ≥8.1 cm |
| | pM1 | Distant metastasis, microscopically confirmed |
| | pM1a | Largest diameter of the largest metastasis ≤3.0 cm, microscopically confirmed |
| | pM1b | Largest diameter of the largest metastasis 3.1–8.0 cm, microscopically confirmed |
| | pM1c | Largest diameter of the largest metastasis ≥8.1 cm, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

There are no Prognostic Stage Groups for iris melanomas. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 Registry Data Collection Variables

| See cha | ee chapter for more details on these variables. | | | |
|---------|---|--|--|--|
| 1. | Tumor site (ICD code lacks specificity): iris (use this staging form) ciliary body (use Uveal Melanoma - Choroidal and | | | |
| | Ciliary Body Melanoma staging form) | | | |
| 2. | Largest basal diameter and thickness of tumor: | | | |
| 3. | Ciliary body involvement: | | | |
| 4. | Extraocular extension: | | | |
| 5. | Histologic type: | | | |
| 6. | Chromosome 3 and 8 loss or gain: | | | |
| 7. | Gene expression profile: | | | |
| 8. | Mitotic count (number of mitoses per 40 HPF, determined by using a 40x objective with a field area of 0.152 mm²): | | | |
| 9. | Extravascular matrix patterns (extracellular closed loops and networks, defined as at least three back-to-back closed loops, is | | | |
| | associated with death from metastatic disease): | | | |
| 10 | . Microvascular density: | | | |

7 Histologic Grade (G)

| ✓ | G | G Definition | |
|---|----|--|--|
| | GX | Grade cannot be assessed | |
| | G1 | Spindle cell melanoma (>90% spindle cells) | |
| | G2 | G2 Mixed cell melanoma (>10% epithelioid cells and <90% spindle cells) | |
| | G3 | Epithelioid cell melanoma (>90% epithelioid cells) | |
| | _ | | |

Note: Because of the lack of universal agreement regarding which proportion of epithelioid cells classifies a tumor as mixed or epithelioid, some ophthalmic pathologists currently combine grades 2 and 3 (nonspindle, i.e. epithelioid cells detected) and contrast them with grade 1 (spindle, i.e. no epithelioid cells detected).

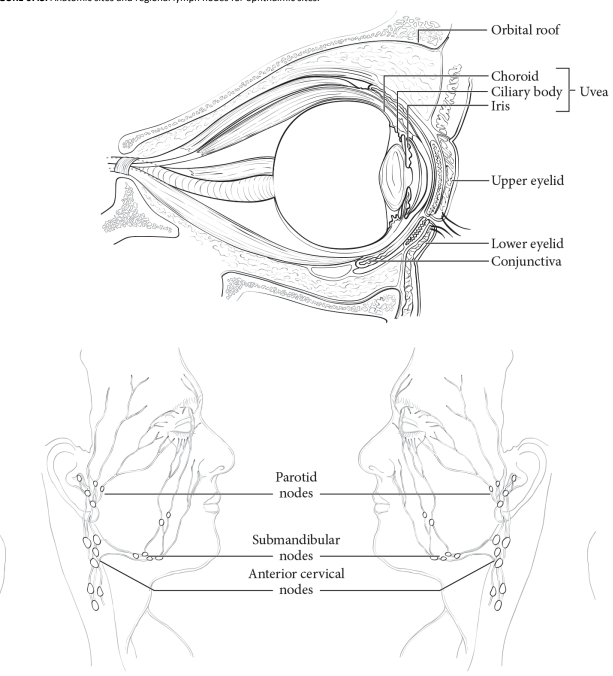
8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

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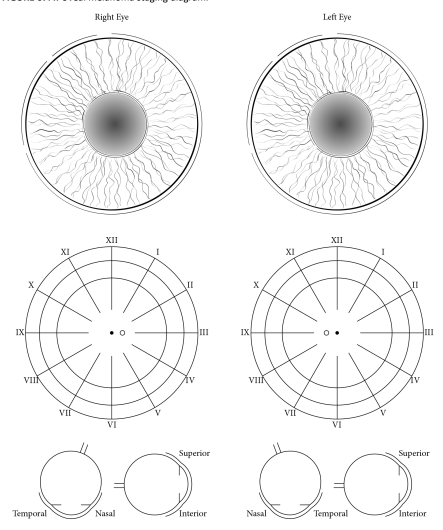
9 Anatomy

FIGURE 67.5. Anatomic sites and regional lymph nodes for ophthalmic sites.



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FIGURE 67.4. Uveal melanoma staging diagram.



| Physician Signature | Date/Time |
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| Patient Name/Information | |
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| | Patient Name/Information |

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
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| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

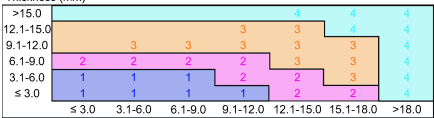
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

FIGURE 67.1. Classification of ciliary body and choroid uveal melanoma based on thickness and diameter

Thickness (mm)



Largest basal diameter (mm)

| ✓ | T Category | T Criteria | | |
|--|--|---|--|--|
| | TX | Primary tumor cannot be assessed | | |
| | T0 | No evidence of primary tumor | | |
| | T1 | Tumor size category 1 | | |
| | T1a | Tumor size category 1 without ciliary body involvement and extraocular extension | | |
| | T1b | Tumor size category 1 with ciliary body involvement | | |
| | T1c | Tumor size category 1 without ciliary body involvement but with extraocular extension ≤5 mm in largest diameter | | |
| | T1d | Tumor size category 1 with ciliary body involvement and extraocular extension ≤5 mm in largest diameter | | |
| | T2 | Tumor size category 2 | | |
| | T2a | Tumor size category 2 without ciliary body involvement and extraocular extension | | |
| | T2b | Tumor size category 2 with ciliary body involvement | | |
| T2c Tumor size category 2 without ciliary body involvement but with extraocular extension ≤5 mm in | | Tumor size category 2 without ciliary body involvement but with extraocular extension ≤5 mm in largest diameter | | |
| | | Tumor size category 2 with ciliary body involvement and extraocular extension ≤5 mm in largest diameter | | |
| T3 Tumor size category 3 | | Tumor size category 3 | | |
| | T3a | Tumor size category 3 without ciliary body involvement and extraocular extension | | |
| | T3b | Tumor size category 3 with ciliary body involvement | | |
| | T3c | Tumor size category 3 without ciliary body involvement but with extraocular extension ≤5 mm in largest diameter | | |
| | T3d | Tumor size category 3 with ciliary body involvement and extraocular extension ≤5 mm in largest diameter | | |
| | T4 | Tumor size category 4 | | |
| | T4a | Tumor size category 4 without ciliary body involvement and extraocular extension | | |
| T4b Tumor size category 4 with ciliary body involvement | | Tumor size category 4 with ciliary body involvement | | |
| | T4c | Tumor size category 4 without ciliary body involvement but with extraocular extension ≤5 mm in largest diameter | | |
| | T4d | Tumor size category 4 with ciliary body involvement and extraocular extension ≤5 mm in largest diameter | | |
| | T4e Any tumor size category with extraocular extension >5 mm in largest diameter | | | |

Notes:

- 1. Primary ciliary body and choroidal melanomas are classified according to the four tumor size categories defined in Figure 67.1
- 2. In clinical practice, the largest tumor basal diameter may be estimated in optic disc diameters (DD; average: 1 DD = 1.5 mm), and tumor thickness may be estimated in diopters (average: 2.5 diopters = 1 mm). Ultrasonography and fundus photography are used to provide more accurate measurements.
- 3. When histopathologic measurements are recorded after fixation, tumor diameter and thickness may be underestimated because of tissue shrinkage.

| ✓ | T Suffix | Suffix Definition | |
|---|----------|---|--|
| (m) Select if synchronous primary tumors are found in single organ. | | Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria | |
|----------|--|--|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No regional lymph node involvement | |
| | N1 | Regional lymph node metastases or discrete tumor deposits in the orbit | |
| | N1a | Metastasis in one or more regional lymph node(s) | |
| | N1b No regional lymph nodes are positive, but there are discrete tumor deposits in the orbit that are not contigue | | |
| | | the eye. (choroidal and ciliary body melanoma only) | |

| 1 | N Suffix | Definition | |
|---|--|------------|--|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | |
| | (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria | |
|---|---|--|--|
| | cM0 | No distant metastasis by clinical classification | |
| | cM1 | Distant metastasis | |
| | cM1a | Largest diameter of the largest metastasis ≤3.0 cm | |
| | cM1b | cM1b Largest diameter of the largest metastasis 3.1–8.0 cm | |
| | cM1c | Largest diameter of the largest metastasis ≥8.1 cm | |
| | pM1 Distant metastasis, microscopically confirmed | | |
| | pM1a Largest diameter of the largest metastasis ≤3.0 cm, microscopically confirmed | | |
| | pM1b Largest diameter of the largest metastasis 3.1–8.0 cm, microscopically confirmed | | |
| | pM1c Largest diameter of the largest metastasis ≥8.1 cm, microscopically confirmed | | |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

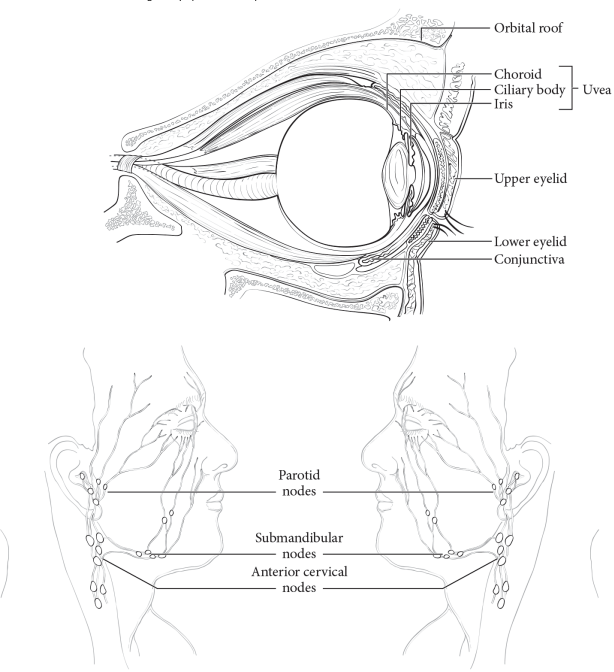
| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1a | N0 | M0 | I |
| | T1b-d | N0 | M0 | IIA |
| | T2a | N0 | M0 | IIA |
| | T2b | N0 | M0 | IIB |
| | T3a | N0 | M0 | IIB |
| | T2c-d | N0 | M0 | IIIA |
| | T3b-c | N0 | M0 | IIIA |
| | T4a | N0 | M0 | IIIA |
| | T3d | N0 | M0 | IIIB |
| | T4b-c | N0 | M0 | IIIB |
| | T4d-e | N0 | M0 | IIIC |
| | Any T | N1 | M0 | IV |
| | Any T | Any N | M1a-c | IV |

| Hospital Name/Address | Patient Name/Information | |
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67.2. Uveal Melanoma – Choroidal and Ciliary Body Melanoma

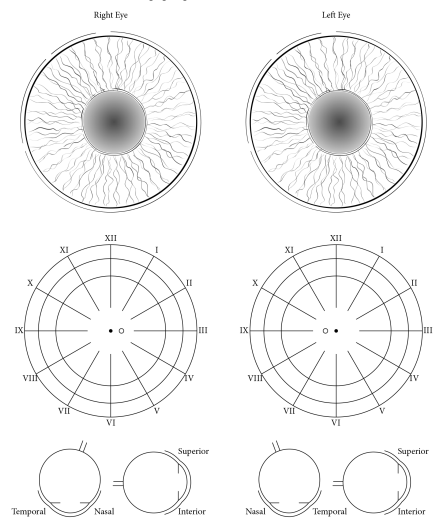
| 6 | | | | | | | |
|--|------------------------|--|--|--|-----------------------------|--|--|
| See chapter for more details on these variables. | | | | | | | |
| | 1. | Tumor site (ICD code lacks specificity): | | | use this staging form) | iris (use Uveal Melanoma – Iris | |
| | Melanoma staging form) | | ase this stagning form, | | | | |
| | - | | | | | | |
| | 2. | | al diameter and thickness of t | umor: | | | |
| | 3. | Ciliary body | involvement: | | | | |
| | 4. | Extraocular | extension: | | | | |
| | 5. | Histologic t | ype: | | | | |
| | 6. | Chromosor | ne 3 and 8 loss or gain: | | | | |
| | 7. | Gene expre | ssion profile: | | | | |
| | 8. | Mitotic cou | nt (number of mitoses per 40 | HPF, determined b | using a 40x objective wi | th a field area of 0.152 mm²): | |
| | 9. | Extravascul | ar matrix patterns (extracellu | lar closed loops and | networks, defined as at I | east three back-to-back closed loops, is | |
| | | associated | with death from metastatic di | sease): | | · | |
| | 10. | Microvascu | lar density: | | | | |
| | | | | | | | |
| | | | | | | | |
| 7 | Hi | stologic (| Grade (G) | | | | |
| _ | | | | | | | |
| / | G | | finition | | | | |
| | GX C1 | | cannot be assessed | lo colla) | | | |
| | G1 G2 | | le cell melanoma (>90% spind I cell melanoma (>10% epithe | | cnindle cells) | | |
| | G3 | | elioid cell melanoma (>90% ep | | spiriule celisj | | |
| Not | | • | | | artion of anithaliaid calls | classifies a tumor as mixed or epithelioid, some | |
| oph | thalm | | ts currently combine grades 2 | | | cted) and contrast them with grade 1 (spindle, | |
| | | | | | | | |
| 8 | Ly | mphovas | cular Invasion (LVI) | | | | |
| | | | | | | | |
| / | | nponent of | Description | | | | |
| | 0 | Coding | LVI not present (absent)/r | not identified | | | |
| | 1 | | LVI present/identified, NC | | | | |
| | 2 | | Lymphatic and small vesse | | | | |
| | 3 | | Venous (large vessel) inva | | | | |
| | 4 | | | BOTH lymphatic and small vessel AND venous (large vessel) invasion | | | |
| | 9 | | Presence of LVI unknown, | /indeterminate | | | |
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FIGURE 67.5. Anatomic sites and regional lymph nodes for ophthalmic sites.



| Hospital Name/Address | Patient Name/Information |
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FIGURE 67.4. Uveal melanoma staging diagram.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification Definition | | |
|---|---------------------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria : First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

4.1.1 Clinical T (cT)

| 4.1.1 Clinical T (cT) | | (01) |
|-----------------------|-------------|--|
| ✓ | cT Category | cT Criteria |
| | cTX | Unknown evidence of intraocular tumor |
| | cT0 | No evidence of intraocular tumor |
| | cT1 | Intraretinal tumor(s) with subretinal fluid |
| | | ≤5 mm from the base of any tumor |
| | cT1a | Tumors ≤3 mm and further than 1.5 mm |
| | | from disc and fovea |
| | cT1b | Tumors >3 mm or closer than 1.5 mm |
| | | from disc or fovea |
| | cT2 | Intraocular tumor(s) with retinal |
| | | detachment, vitreous seeding, or |
| | | subretinal seeding |
| | cT2a | Subretinal fluid >5 mm from the base of |
| | | any tumor |
| | cT2b | Vitreous seeding and/or subretinal |
| | | seeding |
| | cT3 | Advanced intraocular tumor(s) |
| | cT3a | Phthisis or pre-phthisis bulbi |
| | cT3b | Tumor invasion of choroid, pars plana, |
| | | ciliary body, lens, zonules, iris, or anterior |
| | | chamber |
| | cT3c | Raised intraocular pressure with |
| | | neovascularization and/or buphthalmos |
| | cT3d | Hyphema and/or massive vitreous |
| | | hemorrhage |
| | cT3e | Aseptic orbital cellulitis |
| | cT4 | Extraocular tumor(s) involving orbit, |
| | | including optic nerve |
| | cT4a | Radiologic evidence of retrobulbar optic |
| | | nerve involvement or thickening of optic |
| | | nerve or involvement of orbital tissues |
| | cT4b | Extraocular tumor clinically evident with |
| | | proptosis and/or an orbital mass |
| | | |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in |
| | | single organ. |

4.1.2 Pathological T (pT)

| ✓ | pT Category | pT Criteria |
|---|-------------|---|
| | pTX | Unknown evidence of intraocular tumor |
| | pT0 | No evidence of intraocular tumor |
| | pT1 | Intraocular tumor(s) without any local |
| | | invasion, focal choroidal invasion, or pre- |
| | | or intralaminar involvement of the optic |
| | | nerve head |
| | pT2 | Intraocular tumor(s) with local invasion |
| | pT2a | Concomitant focal choroidal invasion |
| | | and pre- or intralaminar involvement of |
| | | the optic nerve head |
| | pT2b | Tumor invasion of stroma of iris and/or |
| | | trabecular meshwork and/or Schlemm's |
| | | canal |
| | pT3 | Intraocular tumor(s) with significant |
| | | local invasion |
| | pT3a | Massive choroidal invasion (>3 mm in |
| | | largest diameter, or multiple foci of focal |
| | | choroidal involvement totalling >3 mm, |
| | | or any full-thickness choroidal |
| | | involvement) |
| | pT3b | Retrolaminar invasion of the optic nerve |
| | | head, not involving the transected end |
| | | of the optic nerve |
| | pT3c | Any partial-thickness involvement of the |
| | T2.1 | sclera within the inner two thirds |
| | pT3d | Full-thickness invasion into the outer |
| | | third of the sclera and/or invasion into |
| | .74 | or around emissary channels |
| | pT4 | Evidence of extraocular tumor: tumor at |
| | | the transected end of the optic nerve, |
| | | tumor in the meningeal spaces around |
| | | the optic nerve, full-thickness invasion of |
| | | the sclera with invasion of the episclera, |
| | | adjacent adipose tissue, extraocular |
| | | muscle, bone, conjunctiva, or eyelids |

| > | T Suffix | Definition | |
|---|----------|---|--|
| | (m) | Select if synchronous primary tumors are found in | |
| | | single organ. | |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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4.2 Definition of Regional Lymph Node (N)

4.2.1 Clinical N (cN)

| 1 | cN | cN Criteria |
|---|----------|--|
| • | Category | |
| | cNX | Regional lymph nodes cannot be assessed |
| | cN0 | No regional lymph node involvement |
| | cN1 | Evidence of preauricular, submandibular, and cervical lymph node involvement |

| ✓ | N Suffix | Definition |
|---|----------|---|
| | (sn) | Select if regional lymph node metastasis |
| | | identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis |
| | | identified by FNA or core needle biopsy only. |

4.2.2 Pathological N (pN)

| ✓ | pN Category | pN Criteria |
|---|----------------|--|
| | pNX | Regional lymph node involvement cannot be assessed |
| | pN0 | No lymph node involvement |
| | pN1 | Regional lymph node involvement |

| 1 | N Suffix | Definition | |
|---|----------|---|--|
| | (sn) | Select if regional lymph node metastasis | |
| | | identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis | |
| | | identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|------------|--|--|
| | cM0 | No signs or symptoms of intracranial or distant metastasis | |
| | cM1 | Distant metastasis without microscopic confirmation | |
| | cM1a | Tumor(s) involving any distant site (e.g., bone marrow, liver) on clinical or radiologic tests | |
| | cM1b | Tumor involving the CNS on radiologic imaging (not including trilateral retinoblastoma) | |
| | pM1 | Distant metastasis with histopathologic confirmation | |
| | pM1a | Histopathologic confirmation of tumor at any distant site (e.g., bone marrow, liver, or other) | |
| | pM1b | Histopathologic confirmation of tumor in the cerebrospinal fluid or CNS parenchyma | |

5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Heritable Trait (H)

| 1 | H Category | H Criteria | |
|---|------------|--|--|
| | HX | Unknown or insufficient evidence of a constitutional RB1 gene mutation. | |
| | Н0 | Normal RB1 alleles in blood tested with demonstrated high-sensitivity assays | |
| | H1 | Bilateral retinoblastoma, retinoblastoma with an intracranial primitive neuroectodermal tumor (i.e., trilateral | |
| | | retinoblastoma), patient with family history of retinoblastoma, or molecular definition of a constitutional RB1 gene | |
| | | mutation | |

| Patient Name/Information | |
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6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6.1 Clinical (cTNM)

| 1 | When T is | And N is | And M is | And H is | Then the stage |
|---|---------------|----------|------------|----------|----------------|
| * | | | | | group is |
| | cT1, cT2, cT3 | cN0 | cM0 | Any H | _ |
| | cT4a | cN0 | cM0 | Any H | Ш |
| | cT4b | cN0 | cM0 | Any H | Ш |
| | Any T | cN1 | cM0 | Any H | III |
| | Any T | Any N | cM1 or pM1 | Any H | IV |

6.2 Pathological (pTNM)

| 1 | When T is | And N is | And M is | And H is | Then the stage |
|---|---------------|----------|------------|----------|----------------|
| • | | | | | group is |
| | pT1, pT2, pT3 | pN0 | cM0 | Any H | 1 |
| | pT4 | pN0 | cM0 | Any H | II |
| | Any T | pN1 | cM0 | Any H | III |
| | Any T | Any N | cM1 or pM1 | Any H | IV |

7 Registry Data Collection Variables

Beyond the factors required to determine stage (T, N, M, and H), the authors have not noted any additional factors for registry data collection.

8 Histologic Grade (G)

| ✓ | G | G Definition |
|---|----|--|
| | GX | Grade cannot be assessed |
| | G1 | Tumor with areas of retinoma (fleurettes or neuronal differentiation) |
| | G2 | Tumor with many rosettes (Flexner–Wintersteiner or Homer Wright) |
| | G3 | Tumor with occasional rosettes (Flexner–Wintersteiner or Homer Wright) |
| | G4 | Tumor with poorly differentiated cells without rosettes and/or with extensive areas (more than half of tumor) of anaplasia |

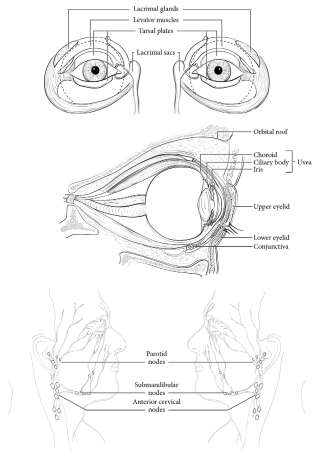
9 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information | |
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FIGURE 68.3. Anatomic sites and regional lymph nodes for ophthalmic sites.



This form continues on the next page.

| Hospital Name/Address | Patient Name/Information | |
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FIGURE 68.2. Staging diagram. Right Left Eye Eye IOP: IOP: Corneal Dia: Corneal Dia: XII XII ΧI Х VIII VIIÌ VI Superior Superior Temporal Interior Interior Nasal Clinical Extent: cT1b cT2a cT2b cT3a cT3b cT3c cT3d Right Eye Left Eye Imaging Orbital Optic Pre-Chiasmatic Chiasmatic Nerve Involved Optic Nerve Involvement Leptomengial Disease Extent: No Tumor Tumor Right Eye Left Eye Constitutional Hereditary Pinealoblastoma: Family History: RB1 Mutation: Trait: N Y N Y N Y Lymph Node: Metastasis: Systemic: N0 cN1 pN1 pM0 cM0 pM1a cM1a pM1b cM1b Pathology: pTX pT1 pT2a pT2b pT3a pT3b pT3c pT3d pT4 pT0 Right Eye Left Eye Physician Signature Date/Time

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤2 cm in greatest dimension with or without extraglandular extension into the orbital soft tissue |
| | T1a | No periosteal or bone involvement |
| | T1b | Periosteal involvement only |
| | T1c | Periosteal and bone involvement |
| | T2 | Tumor >2 cm and ≤4 cm in greatest dimension |
| | T2a | No periosteal or bone involvement |
| | T2b | Periosteal involvement only |
| | T2c | Periosteal and bone involvement |
| | T3 | Tumor >4 cm in greatest dimension |
| | T3a | No periosteal or bone involvement |
| | T3b | Periosteal involvement only |
| | T3c | Periosteal and bone involvement |
| | T4 | Involvement of adjacent structures, including sinuses, temporal fossa, pterygoid fossa, superior orbital fissure, |
| | | cavernous sinus, or brain |
| | T4a | Tumor ≤2 cm in greatest dimension |
| | T4b | Tumor >2 cm and ≤4 cm in greatest dimension |
| | T4c | Tumor >4 cm in greatest dimension |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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AJCC Prognostic Stage Groups

No stage groupings are currently recommended for lacrimal gland carcinomas. Always refer to the specific chapter for rules on clinical and

| | egistry Data Collection Variables | | | | |
|--|---|--|--|--|--|
| hap | er for more details on these variables. | | | | |
| | Freatment Related | | | | |
| 1. | Globe-sparing surgery performed: | | | | |
| 2. | Exenteration performed: | | | | |
| 3. | Orbital bone removed: | | | | |
| 4. | Postoperative radiotherapy: | | | | |
| 5. | Preoperative chemotherapy (intra-arterial vs. systemic): | | | | |
| 6. | Postoperative chemotherapy: | | | | |
| 7. | Concurrent chemoradiation: | | | | |
| 1. | Pathology Related Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) | | | | |
| | 0. | | | | |
| | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) | | | | |
| 2. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: | | | | |
| 2. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: | | | | |
| 2. 3. 4. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: | | | | |
| 2. 3. 4. 5. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): | | | | |
| 2. 3. 4. 5. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: | | | | |
| 2. 3. 4. 5. 6. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: For adenoid cystic carcinoma, approximate percentage of basaloid pattern present on pathological examination: | | | | |
| 2. 3. 4. 5. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: For adenoid cystic carcinoma, approximate percentage of basaloid pattern present on pathological examination: Tumor grade: | | | | |
| 2. 3. 4. 5. 6. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: For adenoid cystic carcinoma, approximate percentage of basaloid pattern present on pathological examination: Tumor grade: Presence of high-grade transformation in any tumor type: | | | | |
| 2. 3. 4. 5. 6. 7. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: For adenoid cystic carcinoma, approximate percentage of basaloid pattern present on pathological examination: Tumor grade: | | | | |
| 2. 3. 4. 5. 6. 7. 8. | Tumor location (ICD code lacks specificity): Lacrimal Gland Lacrimal Sac (not staged using AJCC TNM) Greatest diameter of the tumor: Histopathologic type: Perineural invasion present on pathological examination: Ki-67 growth fraction (percentage of tumor cells positive for Ki-67 on immunohistochemistry): For carcinoma ex pleomorphic adenoma, extent of invasion beyond capsule of pleomorphic adenoma: For adenoid cystic carcinoma, approximate percentage of basaloid pattern present on pathological examination: Tumor grade: Presence of high-grade transformation in any tumor type: | | | | |

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

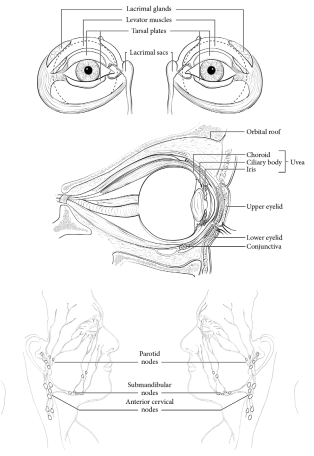
| ✓ | G | G Definition |
|---|----|---|
| | GX | Grade cannot be assessed |
| | G1 | Well differentiated |
| | G2 | Moderately differentiated: includes adenoid cystic carcinoma without basaloid (solid) pattern |
| | G3 | Poorly differentiated: includes adenoid cystic carcinoma with basaloid (solid) pattern |
| | G4 | Undifferentiated |

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 69.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|--|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤2 cm in greatest dimension |
| | T2 | Tumor >2 cm in greatest diameter without invasion of bony walls or globe |
| | T3 | Tumor of any size with invasion of bony walls |
| | T4 | Tumor of any size with invasion of globe or periorbital structures, including eyelid, conjunctiva, temporal fossa, |
| | | nasal cavity, paranasal sinuses, and/or central nervous system |

| | ✓ | T Suffix | Definition |
|---|---|---|------------|
| Ī | | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |

| ✓ | N Suffix | Definition | |
|--|----------|--|--|
| (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | | | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There is no proposal for prognostic stage groupings at this time. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 Registry Data Collection Variables

Beyond T, N, and M, there are no additional variables recommended for collection at this time. See chapter for more details on these variables.

| Hospital Name/Address | Patient Name/Information |
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7 Histologic Grade (G)

Currently, the preferred system for grading of sarcomas is the one proposed by the French Federation of Cancer Centers Sarcoma Group (FNCLCC), otherwise known as the French grading system. It uses three independent prognostic factors to determine the grade: mitotic activity, necrosis, and degree of differentiation of the primary tumor. Each feature is scored separately, and the three scores are added to obtain the grade. Grade 1 is defined as a total score of 2 or 3, grade 2 as a total score of 4 or 5, and grade 3 as a total score of 6 to 8. To enhance the reproducibility of the system, the parameters are defined as precisely as possible. The main value of the grading is to determine risk of distant metastases and overall survival, rather than local recurrence, which depends more on adequate surgical margins.

| ✓ | G | G Definition | |
|--|--|---|--|
| | GX Grade cannot be assessed | | |
| | G1 Total differentiation, mitotic count and necrosis score of 2 or 3 | | |
| | G2 Total differentiation, mitotic count and necrosis score of 4 or 5 | | |
| G3 Total differentiation, mitotic count and necrosis score of 6, 7, or 8 | | Total differentiation, mitotic count and necrosis score of 6, 7, or 8 | |

7.1 Mitotic Count

In the most mitotically active area of the sarcoma, 10 successive high-power fields (HPF; one HPF at 400× magnification = 0.1734 mm²) are assessed using a 40× objective.

| 1 | Mitotic Count Score | Definition |
|---|---------------------|--------------------------|
| | 1 | 0–9 mitoses per 10 HPF |
| | 2 | 10–19 mitoses per 10 HPF |
| | 3 | ≥20 mitoses per 10 HPF |

7.2 Tumor Necrosis

Tumor necrosis is evaluated on gross examination and validated with histologic sections. Necrosis related to previous surgery or to ulceration is not be taken into account, nor is hemorrhage or hyalinization.

| | Necrosis | Definition |
|---|----------|---------------------|
| • | Score | |
| | 0 | No necrosis |
| | 1 | <50% tumor necrosis |
| | 2 | ≥50% tumor necrosis |

7.3 Tumor Differentiation

Tumor differentiation is histology specific and is a mixture of histologic type and subtype and/or true differentiation.

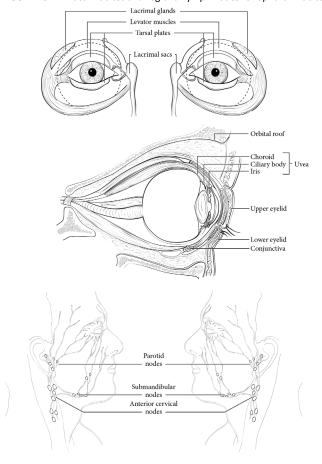
| √ | Differentiation Score | Definition |
|----------|-----------------------|--|
| | 1 | Sarcomas closely resembling normal adult mesenchymal tissue (e.g., low-grade leiomyosarcoma) |
| | 2 | Sarcomas for which histologic typing is certain (e.g., myxoid/round cell liposarcoma) |
| | 3 | Embryonal and undifferentiated sarcomas, sarcomas of doubtful type, synovial sarcomas, |
| | | soft tissue osteosarcoma, Ewing sarcoma /primitive neuroectodermal tumor (PNET) of soft tissue |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 70.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification Definition | |
|---|--|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information diagnostic workup from clinical staging combined with operative findings, and pathology review of resected support specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|--|--|--|
| | TX | Lymphoma extent not specified | |
| | T0 | No evidence of lymphoma | |
| | T1 | Lymphoma involving the conjunctiva alone without eyelid or orbital involvement | |
| | T2 | Lymphoma with orbital involvement with or without conjunctival involvement | |
| | T3 | Lymphoma with preseptal eyelid involvement with or without orbital involvement and with or without | |
| | | conjunctival involvement | |
| | T4 Orbital adnexal lymphoma and extraorbital lymphoma extending beyond the orbit to adjacent structures, such a: | | |
| | | bone, maxillofacial sinuses, and brain. | |

| ✓ | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Involvement of lymph nodes not assessed |
| | N0 | No evidence of lymph node involvement |
| | N1 | Involvement of lymph node region or regions draining the ocular adnexal structures and superior to the |
| | | mediastinum (preauricular, parotid, submandibular, and cervical nodes) |
| | N1a | Involvement of a single lymph node region, superior to the mediastinum |
| | N1b | Involvement of two or more lymph node regions, superior to the mediastinum |
| | N2 | Involvement of lymph node regions of the mediastinum |
| | N3 | Diffuse or disseminated involvement of peripheral and central lymph node regions |

| ✓ | N Suffix | Definition |
|----------|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No evidence of involvement of other extranodal sites |
| | cM1 | Evidence of involvement of other extranodal sites |
| | cM1a | Noncontiguous involvement of tissues or organs external to the ocular adnexa (e.g., parotid glands, submandibular |
| | | gland, lung, liver, spleen, kidney, breast) |
| | cM1b | Lymphomatous involvement of the bone marrow |
| | cM1c | Both M1a and M1b involvement |
| | pM1 | Evidence of involvement of other extranodal sites, microscopically confirmed |
| | pM1a | Noncontiguous involvement of tissues or organs external to the ocular adnexa (e.g., parotid glands, submandibular |
| | | gland, lung, liver, spleen, kidney, breast) |
| | pM1b | Lymphomatous involvement of the bone marrow |
| | pM1c | Both M1a and M1b involvement |

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 AJCC Prognostic Stage Groups

There is no prognostic stage grouping for ocular adnexal lymphoma. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. History of rheumatoid arthritis:
- 2. History of Sjögren's syndrome:
- 3. History of connective tissue disease:
- 4. History of recurrent dry eye syndrome (sicca syndrome):
- 5. History of IgG4 ocular adnexal disease:
- 6. Any evidence of previous or current infection with hepatitis B, hepatitis C, or HIV:
- 7. Any evidence of *Helicobacter pylori* infection:
- 8. Any evidence of an infection caused by *Chlamydia psittaci*:
- 9. Presence or absence of an A20 deletion:
- 10. IGH-locus translocation or somatic mutation pattern (EMZL):
- 11. Concordant/discordant bone marrow involvement (DLBCL):
- 12. Centroblastic/immunoblastic (DLBCL):

7 Histologic Grade (G)

Grade is assigned only to follicular lymphomas, as described by the 2008 WHO classification^{10,18} for malignant lymphomas as follows. For data collection purposes, WHO grade 3a is collected as G3 and WHO grade 3b as G4.

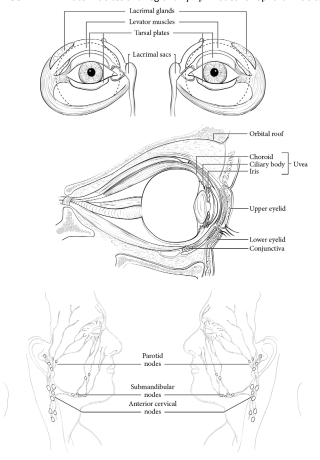
| ✓ | G | G Definition | |
|---|----|---|--|
| | GX | Grade cannot be assessed | |
| | G1 | 1–5 centroblasts per 10 high-power fields (HPF) | |
| | G2 | Between 5 and 15 centroblasts per 10 HPF | |
| | G3 | More than 15 centroblasts per 10 HPF but with admixed centrocytes | |
| | G4 | More than 15 centroblasts per 10 HPF but without centrocytes | |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| * | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 71.1. Anatomic sites and regional lymph nodes for ophthalmic sites.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
|-------------|---|
| pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Not applicable to tumors of the central nervous system. Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

5 AJCC Prognostic Stage Groups

Not applicable to tumors of the central nervous system. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 Registry Data Collection Variables

The variables in this section apply to gliomas. See chapter for more details on these variables.

- 1. IDH mutation:
- 2. WHO grade classification:
- 3. Ki-67/MIB1 labeling index (LI): brain
- 4. Functional neurologic status—e.g., Karnofsky performance scale (KPS):
- 5. Methylation of MGMT
- 6. Chromosome 1p: loss of heterozygosity (LOH)
- 7. Chromosome 19q: LOH
- 8. Extent of surgical resection
- 9. Unifocal versus multifocal tumor

7 Histologic Grade (G)

CNS WHO tumor grades are used in histologic grading. This provides uniformity of classification and categorization of CNS tumors (72.2).

| G | G Definition | |
|----|--|--|
| ı | Circumscribed tumors of low proliferative potential associated with the possibility of cure following resection | |
| II | Infiltrative tumors with low proliferative potential with increased risk of recurrence | |
| Ш | Tumors with histologic evidence of malignancy, including nuclear atypia and mitotic activity, associated with an aggressive clinical | |
| | course | |
| IV | Tumors that are cytologically malignant, mitotically active, and associated with rapid clinical progression and potential for | |
| | dissemination | |

8 Lymphovascular Invasion (LVI)

| ./ | Component of | Description |
|----|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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| 72. Brain and Spinal Cord | |
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2 Instructions

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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|-----|-----------------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤2 cm in greatest dimension limited to the thyroid |
| | T1a | Tumor ≤1 cm in greatest dimension limited to the thyroid |
| | T1b | Tumor >1 cm but ≤2 cm in greatest dimension limited to the thyroid |
| | T2 | Tumor >2 cm but ≤4 cm in greatest dimension limited to the thyroid |
| | T3 | Tumor >4 cm limited to the thyroid, or gross extrathyroidal extension invading only strap muscles |
| | T3a | Tumor >4 cm limited to the thyroid |
| | T3b | Gross extrathyroidal extension invading only strap muscles (sternohyoid, sternothyroid, thyrohyoid, or omohyoid |
| | | muscles) from a tumor of any size |
| | T4 | Includes gross extrathyroidal extension beyond the strap muscles |
| | T4a | Gross extrathyroidal extension invading subcutaneous soft tissues, larynx, trachea, esophagus, or recurrent |
| | | laryngeal nerve from a tumor of any size |
| | T4b | Gross extrathyroidal extension invading prevertebral fascia or encasing the carotid artery or mediastinal vessels |
| | | from a tumor of any size |
| Not | e: All categories may | be subdivided: (s) solitary tumor and (m) multifocal tumor (the largest tumor determines the classification). |

✓ T Suffix Definition

| | ✓ | T Suffix | Definition |
|---|---|----------|-----------------------------|
| Ī | | (s) | Select if solitary tumor. |
| | | (m) | Select if multifocal tumor. |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria |
|----------|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No evidence of locoregional lymph node metastasis |
| | N0a | One or more cytologically or histologically confirmed benign lymph nodes |
| | N0b | No radiologic or clinical evidence of locoregional lymph node metastasis |
| | N1 | Metastasis to regional nodes |
| | N1a | Metastasis to level VI or VII (pretracheal, paratracheal, or prelaryngeal/Delphian, or upper mediastinal) lymph |
| | | nodes. This can be unilateral or bilateral disease. |
| | N1b | Metastasis to unilateral, bilateral, or contralateral lateral neck lymph nodes (levels I, II, III, IV, or V) or |
| | | retropharyngeal lymph nodes |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 Prognostic Factors Required for Stage Grouping

5.1 Definition of Age at Diagnosis

| 1 | / | Age at Diagnosis |
|---|---|------------------|
| | | < 55 years |
| | | ≥ 55 years |

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When age at diagnosis is | When T is | And N is | And M is | Then the stage group is |
|---|--------------------------|-----------|----------|----------|-------------------------|
| | <55 years | Any T | Any N | M0 | 1 |
| | <55 years | Any T | Any N | M1 | II . |
| | ≥55 years | T1 | NO/NX | M0 | 1 |
| | ≥55 years | T1 | N1 | M0 | II |
| | ≥55 years | T2 | NO/NX | M0 | I |
| | ≥55 years | T2 | N1 | M0 | II . |
| | ≥55 years | T3a/T3b | Any N | M0 | II . |
| | ≥55 years | T4a | Any N | M0 | III |
| | ≥55 years | T4b | Any N | M0 | IVA |
| | ≥55 years | Any T | Any N | M1 | IVB |

7 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Histology:
- 2. Age at diagnosis:
- 3. Number of involved lymph nodes:
- 4. Maximum diameter of involved lymph nodes:
- 5. Size of largest metastatic foci within an involved lymph node:

8 Histologic Grade (G)

There is no formal grading system for thyroid cancers.

9 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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| 73.1. Thyroid – Differentiated | | |
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| Hospital Name/Address | Patient Name/Information | |
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FIGURE 73.1. Anatomy of the thyroid gland.

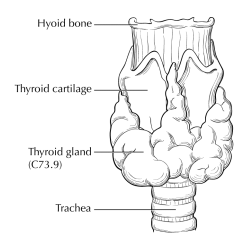
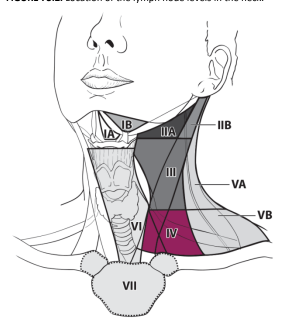


FIGURE 73.2. Location of the lymph node levels in the neck.



| Physician Signature | Date/Time |
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| Patient Name/Information | |
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| | Patient Name/Information |

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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information | |
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| T Category | T Criteria | |
|---|--|--|
| TX | Primary tumor cannot be assessed | |
| T0 | No evidence of primary tumor | |
| T1 | Tumor ≤2 cm in greatest dimension limited to the thyroid | |
| T1a | Tumor ≤1 cm in greatest dimension limited to the thyroid | |
| T1b | Tumor >1 cm but ≤2 cm in greatest dimension limited to the thyroid | |
| T2 | Tumor >2 cm but ≤4 cm in greatest dimension limited to the thyroid | |
| T3 | Tumor >4 cm limited to the thyroid, or gross extrathyroidal extension invading only strap muscles | |
| T3a | Tumor >4 cm limited to the thyroid | |
| T3b Gross extrathyroidal extension invading only strap muscles (sternohyoid, sternothyroid, thyrohyoid, or omoh muscles) from a tumor of any size | | |
| T4 | Includes gross extrathyroidal extension beyond the strap muscles | |
| T4a | Gross extrathyroidal extension invading subcutaneous soft tissues, larynx, trachea, esophagus, or recurrent laryngeal nerve from a tumor of any size | |
| T4b | Gross extrathyroidal extension invading prevertebral fascia or encasing the carotid artery or mediastinal vessels from a tumor of any size | |
| | T0 T1 T1a T1b T2 T3 T3a T3b T4 T4 | |

| ✓ | T Suffix | Definition |
|----------|----------|-----------------------------|
| | (s) | Select if solitary tumor. |
| | (m) | Select if multifocal tumor. |

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria | |
|----------|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No evidence of locoregional lymph node metastasis | |
| | N0a | One or more cytologically or histologically confirmed benign lymph nodes | |
| | N0b | No radiologic or clinical evidence of locoregional lymph node metastasis | |
| | N1 | Metastasis to regional nodes | |
| | N1a | Metastasis to level VI or VII (pretracheal, paratracheal, or prelaryngeal/Delphian, or upper mediastinal) lymph | |
| | | nodes. This can be unilateral or bilateral disease. | |
| | N1b | Metastasis to unilateral, bilateral, or contralateral lateral neck lymph nodes (levels I, II, III, IV, or V) or | |
| | | retropharyngeal lymph nodes | |

| 1 | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|----------|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

| Hospital Name/Address | Patient Name/Information |
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5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1–T3a | NO/NX | M0 | IVA |
| | T1-T3a | N1 | M0 | IVB |
| | T3b | Any N | M0 | IVB |
| | T4 | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Histology:
- 2. Age at diagnosis:
- 3. Number of involved lymph nodes:
- 4. Maximum diameter of involved lymph nodes:
- 5. Size of largest metastatic foci within an involved lymph node:

7 Histologic Grade (G)

There is no formal grading system for thyroid cancers.

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 73.1. Anatomy of the thyroid gland.

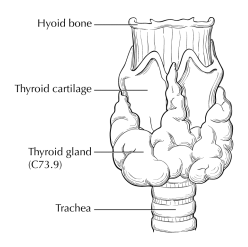
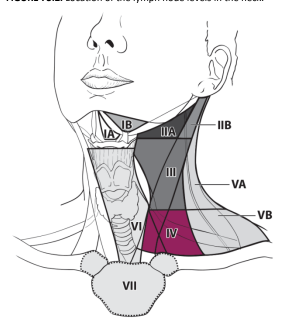


FIGURE 73.2. Location of the lymph node levels in the neck.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
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| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
|-----------------------|--------------------------|--|
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Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria | |
|---|---|---|--|
| | TX | Primary tumor cannot be assessed | |
| | T0 | No evidence of primary tumor | |
| | T1 | Tumor ≤2 cm in greatest dimension limited to the thyroid | |
| | T1a | Tumor ≤1 cm in greatest dimension limited to the thyroid | |
| | T1b | Tumor >1 cm but ≤2 cm in greatest dimension limited to the thyroid | |
| | T2 Tumor >2 cm but ≤4 cm in greatest dimension limited to the thyroid | | |
| | T3 | Tumor >4 cm or with extrathyroidal extension | |
| | T3a | Tumor >4 cm in greatest dimension limited to the thyroid | |
| | T3b | Tumor of any size with gross extrathyroidal extension invading only strap muscles (sternohyoid, sternothyroid, thyrohyoid or omohyoid muscles) | |
| | T4 Advanced disease | | |
| | T4a | Moderately advanced disease; tumor of any size with gross extrathyroidal extension into the nearby tissues of the neck, including subcutaneous soft tissue, larynx, trachea, esophagus, or recurrent laryngeal nerve | |
| | T4b | Very advanced disease; tumor of any size with extension toward the spine or into nearby large blood vessels, gross extrathyroidal extension invading the prevertebral fascia, or encasing the carotid artery or mediastinal vessels | |

| / | T Suffix | Definition |
|---|---|------------|
| | (m) Select if synchronous primary tumors are found in single organ. | |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|---|------------|---|--|
| | NX | Regional lymph nodes cannot be assessed | |
| | N0 | No evidence of locoregional lymph node metastasis | |
| | N0a | One or more cytologically or histologically confirmed benign lymph nodes | |
| | N0b | No radiologic or clinical evidence of locoregional lymph node metastasis | |
| | N1 | Metastasis to regional nodes | |
| | N1a | Metastasis to level VI or VII (pretracheal, paratracheal, or prelaryngeal/Delphian, or upper mediastinal) lymph | |
| | | nodes. This can be unilateral or bilateral disease. | |
| | N1b | Metastasis to unilateral, bilateral, or contralateral lateral neck lymph nodes (levels I, II, III, IV, or V) or | |
| | | retropharyngeal lymph nodes | |

| ✓ | N Suffix | Definition | |
|---|----------|--|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. | |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

 ${\it This form\ continues\ on\ the\ next\ page}.$

| Hospital Name/Address | Patient Name/Information | |
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AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | NO | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T3 | N0 | M0 | II |
| | T1-3 | N1a | M0 | III |
| | T4a | Any N | M0 | IVA |
| | T1-3 | N1b | M0 | IVA |
| | T4b | Any N | M0 | IVB |
| | Any T | Any N | M1 | IVC |

Registry Data Collection Variables

See

| 1. | Age at diagnosis: |
|-----|--|
| 2. | Gender: |
| 3. | Race: |
| 4. | Histology: |
| 5. | Size of primary tumor: |
| 6. | Number of involved lymph nodes: |
| 7. | Presence of extranodal extension: |
| 8. | Size of the involved lymph nodes: |
| 9. | Size of the metastatic focus in the involved lymph nodes: |
| 10. | Completeness of resection: |
| 11. | Preoperative calcitonin: |
| 12. | Preoperative CEA: |
| 13. | Genetic mutations, including specific codon information for mutations in the <i>RET</i> protooncogene, including the method of measurement, if available. Other mutations to be documented are in the <i>RAS</i> (<i>HRAS</i> , <i>KRAS</i> , or <i>NRAS</i>) group. |
| 14. | Whether the patient has medullary thyroid carcinoma that is sporadic or hereditary, if known: |

Histologic Grade (G)

Grade is not used in the staging for medullary thyroid carcinoma.

| Hospital Name/Address | Patient Name/Information |
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8 Lymphovascular Invasion (LVI)

| ✓ | Component of | Description |
|---|--------------|--|
| | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

9 Anatomy

FIGURE 73.1. Anatomy of the thyroid gland.

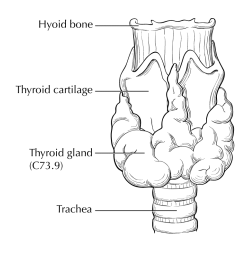
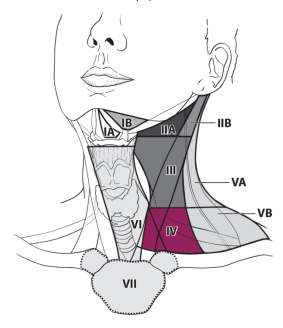


FIGURE 74.2. Location of the lymph node levels in the neck.



| | - |
|---------------------|---------------|
| Physician Signature | Date/Time |

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria : First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category | T Criteria |
|---|------------|---|
| | TX | Primary tumor cannot be assessed |
| | TO | No evidence of primary tumor |
| | Tis | Atypical parathyroid neoplasm (neoplasm of uncertain malignant potential)* |
| | T1 | Localized to the parathyroid gland with extension limited to soft tissue |
| | T2 | Direct invasion into the thyroid gland |
| | T3 | Direct invasion into recurrent laryngeal nerve, esophagus, trachea, skeletal muscle, adjacent lymph nodes, or |
| | | thymus |
| | T4 | Direct invasion into major blood vessel or spine |

*Defined as tumors that are histologically or clinically worrisome but do not fulfill the more robust criteria (i.e., invasion, metastasis) for carcinoma. They generally include tumors that have two or more concerning features, such as fibrous bands, mitotic figures, necrosis, trabecular growth, or adherence to surrounding tissues intraoperatively. Atypical parathyroid neoplasms usually have a smaller dimension, weight, and volume than carcinomas and are less likely to have coagulative tumor necrosis.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Regional lymph node metastasis |
| | N1a | Metastasis to level VI (pretracheal, paratracheal, and prelaryngeal/Delphian lymph nodes) or superior mediastinal lymph nodes (level VII) |
| | N1b | Metastasis to unilateral, bilateral, or contralateral cervical (level I, II, III, IV, or V) or retropharyngeal nodes |

| | ✓ | N Suffix | Definition |
|---|---|----------|--|
| ſ | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| Ī | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| ✓ | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

There are not enough data to propose prognostic stage groups for parathyroid carcinoma. Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

 ${\it This form\ continues\ on\ the\ next\ page.}$

| Hospital Name/Address | Patient Name/Information |
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| 6 | | | a Collection Variables tails on these variables. |
|------|---------|----------------------|--|
| see | спарі | er for more de | tails off these variables. |
| | 1. | Age at diagno | osis: |
| | 2. | Gender: | |
| | 3. | Race: | |
| | 4. | Size of prima | ry tumor in millimeters: |
| | 5. | Location of p | rimary tumor: left or right and superior (upper) or inferior (lower): |
| | 6. | | surrounding tissue: |
| | 7. | Distant meta | |
| | 8. | | mph nodes removed (by level): |
| | 9. | • | mph nodes positive (by level): |
| | 10. | | perative calcium (number in tenths in milligrams per deciliter [e.g., 11.5 mg/dL]): |
| | 11. | | perative PTH (whole number in picograms per milliliter [e.g., 350 pg/mL]): |
| | 12. | Lymphovascu | |
| | 13. | | Low Grade High Grade |
| | 14. | | mary tumor (in milligrams): |
| | 15. | Mitotic rate: | |
| | 16. | Time to recur | rrence (months): |
| 7 | Hi | stologic Gr | rade (G) |
| | | | rined as low grade or high grade. |
| Cytt | onacic | ar grade is der | med us low grade of high grade. |
| / | G | G Defi | nition |
| | LG | _ | ide: round monomorphic nuclei with only mild to moderate nuclear size variation, indistinct nucleoli, and chromatin |
| | HG | | eristics resembling those of normal parathyroid or of adenoma ade: more pleomorphism, with a nuclear size variation greater than 4:1; prominent nuclear membrane irregularities; |
| | | | tin alterations, including hyperchromasia or margination of chromatin; and prominent nucleoli. High-grade tumors |
| | | snow se | everal discrete confluent areas with nuclear changes. |
| _ | | _ | |
| 8 | Ly | mphovasc | ular Invasion (LVI) |
| | | | |
| / | | nponent of Coding | Description |
| | 0 | county | LVI not present (absent)/not identified |
| | 1 | | LVI present/identified, NOS |
| | 2 | | Lymphatic and small vessel invasion only (L) |
| | 3 | | Venous (large vessel) invasion only (V) |
| | 9 | | BOTH lymphatic and small vessel AND venous (large vessel) invasion Presence of LVI unknown/indeterminate |
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FIGURE 75.1. Anatomy of the parathyroid gland.

FIGURE 75.2. Lymph node levels in the neck.

Superior parathyroid glands lighands larger parathyroid glands larger parathyroid

| Hospital Name/Address | Patient Name/Information | |
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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information from diagnostic workup from clinical staging combined with operative findings, and pathology review of resected surgical specimens |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| ✓ | T Category T Criteria | |
|---|-----------------------|---|
| | TX | Primary tumor cannot be assessed |
| | T0 | No evidence of primary tumor |
| | T1 | Tumor ≤5 cm in greatest dimension, no extra-adrenal invasion |
| | T2 | Tumor >5 cm, no extra-adrenal invasion |
| | T3 | Tumor of any size with local invasion but not invading adjacent organs |
| | T4 | Tumor of any size that invades adjacent organs (kidney, diaphragm, pancreas, spleen, or liver) or large blood |
| | 14 | vessels (renal vein or vena cava) |

| | ✓ | T Suffix | Definition |
|---|---|----------|---|
| Ī | | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria |
|---|------------|---|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No regional lymph node metastasis |
| | N1 | Metastasis in regional lymph node(s) |

| | ✓ | N Suffix | Definition |
|---|---|----------|--|
| Г | | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| Г | | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|---|
| | cM0 | No distant metastasis |
| | cM1 | Distant metastasis |
| | pM1 | Distant metastasis, microscopically confirmed |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|---|-----------|----------|----------|-------------------------|
| | T1 | NO | M0 | 1 |
| | T1 | N1 | M0 | III |
| | T2 | N0 | M0 | II |
| | T2 | N1 | M0 | III |
| | T3 | Any N | M0 | III |
| | T4 | Any N | M0 | III |
| | Any T | Any N | M1 | IV |

| Hospital Name/Address | Patient Name/Information |
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6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Tumor weight in grams:
- 2. Vascular invasion:
- 3. Mitotic count:
- 4. Ki-67 proliferative index:
- 5. Weiss score:

7 Histologic Grade (G)

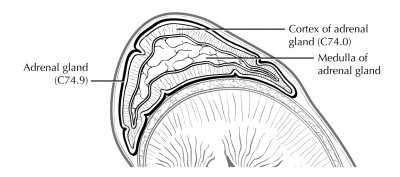
| ✓ | G | G Definition |
|---|----|---|
| | LG | Low grade (≤20 mitoses per 50 HPF) |
| | HG | High grade (>20 mitosis per 50 HPF); TP53 or CTNNB mutation |

8 Lymphovascular Invasion (LVI)

| 1 | Component of | Description |
|---|--------------|--|
| • | LVI Coding | |
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information | |
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FIGURE 76.1. Anatomy of the adrenal gland.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Hospital Name/Address | Patient Name/Information |
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3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|---|--|--|--|--|
| | cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | | |
| | pTNM Pathological Classification: Used for patients if surgery is the first definitive therapy. Composed of information of diagnostic workup from clinical staging combined with operative findings, and pathology review of resected sur specimens | | | |
| | ycTNM | Posttherapy Clinical Classification: after primary systemic and/or radiation therapy, or after neoadjuvant therapy and before planned surgery. Criteria: First therapy is systemic and/or radiation therapy | | |
| | ypTNM | Posttherapy Pathological Classification: Used for staging after neoadjuvant therapy and planned post neoadjuvant therapy surgery. Criteria: First therapy is systemic and/or radiation therapy and is followed by surgery. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

| Hospital Name/Address | Patient Name/Information | |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria | | |
|--|---|------------|--|--|
| | TX Primary tumor cannot be assessed | | | |
| | T1 PH <5 cm in greatest dimension, no extra-adrenal invasion | | | |
| | T2 PH ≥ 5 cm or PG-sympathetic of any size, no extra-adrenal invasion | | | |
| T3 Tumor of any size with invasion into surrounding tissues (e.g., liver, pancreas, spleen, kidneys) | | | | |

PH: within adrenal gland

PG Sympathetic: functional

PG Parasympathetic: nonfunctional, usually in the head and neck region

Note: Parasympathetic Paragaglioma are not staged because they are largely benign.

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

4.2 Definition of Regional Lymph Node (N)

| ✓ | N Category | N Criteria | |
|-----------------------------|--|--------------------------------|--|
| | NX Regional lymph nodes cannot be assessed | | |
| NO No lymph node metastasis | | No lymph node metastasis | |
| | N1 | Regional lymph node metastasis | |

| ✓ | N Suffix | Definition |
|--|--|------------|
| | (sn) Select if regional lymph node metastasis identified by SLN biopsy only. | |
| (f) Select if regional lymph node metastasis identified by FNA or core needle biopsy only. | | |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria | |
|---|--|-----------------------|--|
| | cM0 | No distant metastasis | |
| | cM1 | Distant metastasis | |
| | cM1a Distant metastasis to only bone | | |
| | cM1b Distant metastasis to only distant lymph nodes/liver or lung | | |
| | cM1c Distant metastasis to bone plus multiple other sites | | |
| | pM1 Distant metastasis, microscopically confirmed | | |
| | pM1a Distant metastasis to only bone, microscopically confirmed | | |
| | pM1b Distant metastasis to only distant lymph nodes/liver or lung, microscopically confirmed | | |
| | pM1c Distant metastasis to bone plus multiple other sites, microscopically confirmed | | |

| Hospital Name/Address | Patient Name/Information | |
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| | | |

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

| ✓ | When T is | And N is | And M is | Then the stage group is |
|----------|-----------|----------|----------|-------------------------|
| | T1 | N0 | M0 | 1 |
| | T2 | N0 | M0 | II |
| | T1 | N1 | M0 | III |
| | T2 | N1 | M0 | III |
| | T3 | Any N | M0 | III |
| | Any T | Any N | M1 | IV |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Primary tumor size (measured in centimeters):
- 2. Primary tumor location: PH, PG (specific location: e.g., aortic bifurcation, mediastinum):
- 3. Regional lymph node metastases:
- 4. Location of distant metastases:
- 5. Hormonal function: 24-hour urinary fractionated metanephrines/plasma metanephrines:
- 6. Chromogranin A:
- 7. Mitotic count:
- 8. Germline mutation status:
- 9. Plasma methoxytyramine:

7 Histologic Grade (G)

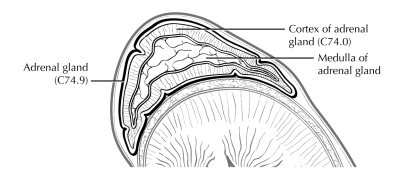
There is no recommended histologic grading system at this time.

8 Lymphovascular Invasion (LVI)

| ✓ | Component of LVI Coding | Description |
|---|-------------------------|--|
| | 0 | LVI not present (absent)/not identified |
| | 1 | LVI present/identified, NOS |
| | 2 | Lymphatic and small vessel invasion only (L) |
| | 3 | Venous (large vessel) invasion only (V) |
| | 4 | BOTH lymphatic and small vessel AND venous (large vessel) invasion |
| | 9 | Presence of LVI unknown/indeterminate |

| Hospital Name/Address | Patient Name/Information |
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FIGURE 76.1. Anatomy of the adrenal gland.



| Physician Signature | Date/Time |
|---------------------|-----------|

| Patient Name/Information | |
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| | Patient Name/Information |

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3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
|--|----------------|--|--|--|
| cTNM or TNM Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imagin endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and othe relevant examinations | | | | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage | | Stage description |
|--|---|---|
| ✓ | Limited stage | |
| | 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other lymph node regions on the same side of the diaphragm |
| | II bulky* | Stage II with disease bulk** |
| | Advanced stage | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm with spleen involvement |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without associated lymph node involvement |
| or <i>noncontiguous</i> extralymphatic organ involvement in conjunction with nodal Start or <i>any</i> extralymphatic organ involvement in nodal Stage III disease | | or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease or any extralymphatic organ involvement in nodal Stage III disease |
| | | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other than by direct extension in IIE disease). |
| | age II bulky may be considered either early HL prognostic factors). | r- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion |
| ** | The definition of disease bulk varies accord | ing to the lymphoma histology. In the Lugano classification, bulk in HL is defined as a mass |
| gre | ater than one third of the thoracic diameter | er on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by |

lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation.^{2,3} In DLBCL, cutoffs ranging from 5

6 Registry Data Collection Variables

to 10 cm have been used, although 10 cm is recommended. 4

See chapter for more details on these variables.

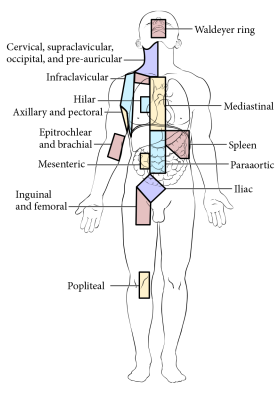
| 1. | Size of the larg | est mass in m | illimeters for al | ll stages: essentia | I for Stages Land | ł II |
|----|------------------|---------------|-------------------|---------------------|-------------------|------|

 ${\it This form\ continues\ on\ the\ next\ page.}$

Note: A/B is no longer used in NHL.

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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- 1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.
- 2. Arcaini L, Rattotti S, Gotti M, Luminari S. Prognostic assessment in patients with indolent B-cell lymphomas. *ScientificWorldJournal*. 2012;2012:107892.
- 3. Federico M, Bellei M, Marcheselli L, et al. Follicular lymphoma international prognostic index 2: a new prognostic index for follicular lymphoma developed by the international follicular lymphoma prognostic factor project. *J Clin Oncol.* 2009;27(27):4555-4562.
- 4. Pfreundschuh M, Ho AD, Cavallin-Stahl E, et al. Prognostic significance of maximum tumour (bulk) diameter in young patients with good-prognosis diffuse large-B-cell lymphoma treated with CHOP-like chemotherapy with or without rituximab: an exploratory analysis of the MabThera International Trial Group (MInT) study. *The lancet oncology*. 2008;9(5):435-444.

| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Classification Definition | | |
|--|----------------|---|--|--|
| workup information, until first treatment, including clinical history and symptoms, physical examination endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or s | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage | | Stage description | |
|--|--|--|--|
| ✓ Limited stage | | | |
| | 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) | |
| IE Single extralymphatic site in the absence of nodal involvement (rare in HL) II Involvement of two or more lymph node regions on the same side of the diaphra | | Single extralymphatic site in the absence of nodal involvement (rare in HL) | |
| | | Involvement of two or more lymph node regions on the same side of the diaphragm | |
| | | Contiguous extralymphatic extension from a nodal site with or without involvement of other | |
| | | lymph node regions on the same side of the diaphragm | |
| | II bulky* | Stage II with disease bulk** | |
| | Advanced stage | | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm | |
| | | with spleen involvement | |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without | |
| | | associated lymph node involvement | |
| | | or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease | |
| | | or <i>any</i> extralymphatic organ involvement in nodal Stage III disease | |
| | | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other | |
| | | | |
| | than by direct extension in IIE disease). | | |
| | | y- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion | |
| of | HL prognostic factors). | | |
| ** | The definition of disease bulk varies accord | ling to the lymphoma histology. In the Lugano classification, ¹ bulk in HL is defined as a mass | |
| gre | ater than one third of the thoracic diamet | er on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by | |

lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation. ^{2,3} In DLBCL, cutoffs ranging from 5 to 10 cm have been used, although 10 cm is recommended.4

Note: A/B is no longer used in NHL.

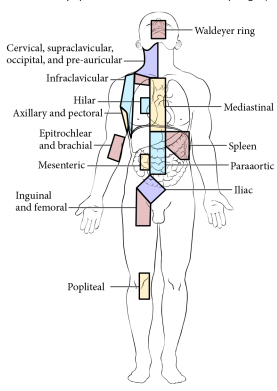
Registry Data Collection Variables

See chapter for more details on these variables.

- Size of the largest mass in millimeters for all stages; essential for Stages I and II:
- NCCN IPI points (0-8):
- 3. IHC-determined COO:

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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- 4. Pfreundschuh M, Ho AD, Cavallin-Stahl E, et al. Prognostic significance of maximum tumour (bulk) diameter in young patients with good-prognosis diffuse large-B-cell lymphoma treated with CHOP-like chemotherapy with or without rituximab: an exploratory analysis of the MabThera International Trial Group (MInT) study. *The lancet oncology.* 2008;9(5):435-444.

| Patient Name/Information | |
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| | Patient Name/Information |

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2 Instructions

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage | | Stage description |
|----------|---|---|
| ✓ | Limited stage | • |
| | 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other |
| | | lymph node regions on the same side of the diaphragm |
| | II bulky* | Stage II with disease bulk** |
| | Advanced stage | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm with spleen involvement |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without associated lymph node involvement or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease or any extralymphatic organ involvement in nodal Stage III disease |
| | | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other than by direct extension in IIE disease). |
| | age II bulky may be consider HL prognostic factors). | ed either early- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion |
| ** | The definition of disease bulk | varies according to the lymphoma histology. In the Lugano classification,¹ bulk in HL is defined as a mass |
| gre | ater than one third of the th | oracic diameter on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by |

 $lymphoma\ histology.\ In\ follicular\ lymphoma,\ 6\ cm\ has\ been\ suggested\ based\ on\ the\ FLIPI-2\ and\ its\ validation.^{2,3}\ In\ DLBCL,\ cutoffs\ ranging\ from\ 5$

6 Registry Data Collection Variables

to 10 cm have been used, although 10 cm is recommended.4

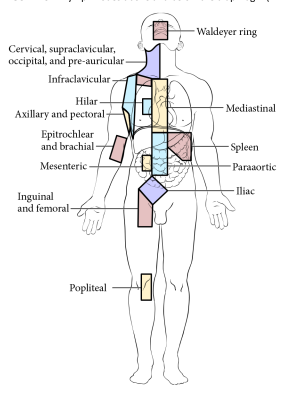
See chapter for more details on these variables.

Note: A/B is no longer used in NHL.

- 1. Size of the largest mass in millimeters for all stages; essential for Stages I and II:
- 2. Proliferation index (% of positivity with either the Ki-67 or MIB1 monoclonal antibodies):

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
|---------------------|-----------|

- 1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.
- 2. Arcaini L, Rattotti S, Gotti M, Luminari S. Prognostic assessment in patients with indolent B-cell lymphomas. *ScientificWorldJournal*. 2012;2012:107892.
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- 4. Pfreundschuh M, Ho AD, Cavallin-Stahl E, et al. Prognostic significance of maximum tumour (bulk) diameter in young patients with good-prognosis diffuse large-B-cell lymphoma treated with CHOP-like chemotherapy with or without rituximab: an exploratory analysis of the MabThera International Trial Group (MInT) study. *The lancet oncology*. 2008;9(5):435-444.

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3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stag | je | Stage description |
|--------|--|---|
| ✓ | Limited stage | |
| | I | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other lymph node regions on the same side of the diaphragm |
| | II bulky* | Stage II with disease bulk** |
| | Advanced stage | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm with spleen involvement |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without associated lymph node involvement or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease |
| | | or <i>any</i> extralymphatic organ involvement in nodal Stage III disease Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other than by direct extension in IIE disease). |
| *\$+20 | To II hulky may be considered either o | early- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion |
| - | ge if bulky may be considered either e prognostic factors). | rarry- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion |
| **Th | e definition of disease bulk varies acc | ording to the lymphoma histology. In the Lugano classification, bulk in HL is defined as a mass |

^{**}The definition of disease bulk varies according to the lymphoma histology. In the Lugano classification, bulk in HL is defined as a mass greater than one third of the thoracic diameter on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation. DLBCL, cutoffs ranging from 5 to 10 cm have been used, although 10 cm is recommended.

Note: A/B is no longer used in NHL.

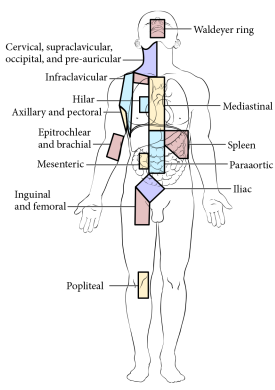
6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Size of the largest mass in millimeters for all stages; essential for Stages I and II:
- 2. Tumor disease burden (high [one or more factors] vs. low [0 factors]) based on the presence or absence of GELF criteria:
- 3. FLIPI (as FLIPI-1 or FLIPI-2):

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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3 Time of Classification (select one):

| ✓ | Classification | Definition | | |
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| workup information, until first treatment, including clinical history and symptoms, physical examination, image endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or samp | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | | |
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| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | | |

4 Definitions of AJCC TNM

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| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Sto | Stage Stage description | | | |
|----------|---|---|--|--|
| √ | Limited stage | | | |
| | 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) | | |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) | | |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm | | |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other lymph node regions on the same side of the diaphragm | | |
| | II bulky* | Stage II with disease bulk** | | |
| | Advanced stage | | | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm with spleen involvement | | |
| | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without associated lymph node involvement or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease or any extralymphatic organ involvement in nodal Stage III disease | | | |
| | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other than by direct extension in IIE disease). | | | |
| | age II bulky may be considered either earling. HL prognostic factors). | y- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion | | |
| ** | he definition of disease bulk varies accord | ling to the lymphoma histology. In the Lugano classification, bulk in HL is defined as a mass | | |
| gre | ater than one third of the thoracic diamet | er on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by | | |

lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation.^{2,3} In DLBCL, cutoffs ranging from 5

6 Registry Data Collection Variables

to 10 cm have been used, although 10 cm is recommended. 4

See chapter for more details on these variables.

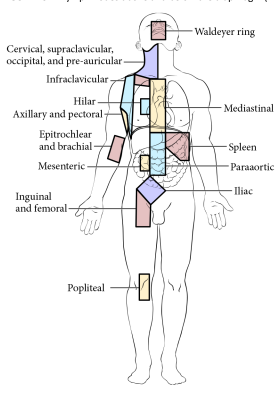
| 1. | Size of the | largest mass | in millimeter | s for all stages: | essential for Stages | I and II: |
|----|-------------|--------------|---------------|-------------------|----------------------|-----------|

 ${\it This form\ continues\ on\ the\ next\ page.}$

Note: A/B is no longer used in NHL.

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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- 1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.
- 2. Arcaini L, Rattotti S, Gotti M, Luminari S. Prognostic assessment in patients with indolent B-cell lymphomas. *ScientificWorldJournal*. 2012;2012:107892.
- 3. Federico M, Bellei M, Marcheselli L, et al. Follicular lymphoma international prognostic index 2: a new prognostic index for follicular lymphoma developed by the international follicular lymphoma prognostic factor project. *J Clin Oncol.* 2009;27(27):4555-4562.
- 4. Pfreundschuh M, Ho AD, Cavallin-Stahl E, et al. Prognostic significance of maximum tumour (bulk) diameter in young patients with good-prognosis diffuse large-B-cell lymphoma treated with CHOP-like chemotherapy with or without rituximab: an exploratory analysis of the MabThera International Trial Group (MInT) study. *The lancet oncology.* 2008;9(5):435-444.

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79.5. Non-Hodgkin Lymphomas: Chronic Lymphocytic Leukemia and Small Lymphocytic Lymphoma

1 Terms of Use

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2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

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The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

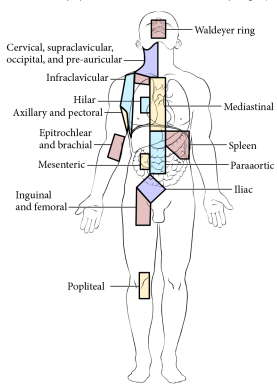
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| | Lymphoma | | |
|------------|---|---|--|
| <u></u> | A DESCRIPTION OF A LOCATION | | |
| 4 | Definitions of AJCC TNM | | |
| TNI | M does not apply to this disease. Always r | efer to the specific chapter | r for explicit instructions on classification for this disease. |
| 5 | AJCC Prognostic Stage Grou | ps | |
| Alw | vays refer to the specific chapter for rules | on clinical and pathological | I classification of this disease. |
| 5.1 | L Lugano Classification for Hoo | dgkin and Non-Hodgk | kin Lymphoma¹ |
| Sto | age | Stage description | |
| √ | Limited stage | _ stage acstription | |
| | I | Involvement of a single | lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| | IE | _ ' ' | te in the absence of nodal involvement (rare in HL) |
| | II | | nore lymph node regions on the same side of the diaphragm |
| | IIE | , , | itic extension from a nodal site with or without involvement of other the same side of the diaphragm |
| | II bulky* | Stage II with disease bul | |
| | Advanced stage | _ stage if with disease bar | 0 |
| | III | Involvement of lymph no with spleen involvement | ode regions on both sides of the diaphragm; nodes above the diaphragm |
| | IV | | involvement of one or more extralymphatic organs, with or without |
| | | associated lymph node i | |
| | | | ymphatic organ involvement in conjunction with nodal Stage II disease |
| | | or <i>any</i> extralymphatic of | rgan involvement in nodal Stage III disease |
| | | Stage IV includes any inv | volvement of the CSF, bone marrow, liver, or multiple lung lesions (other in IIF disease). |
| *St | age II bulky may be considered either earl | • | se based on lymphoma histology and prognostic factors (see discussion |
| | HL prognostic factors). | | |
| gre lym | ater than one third of the thoracic diamet | er on CT of the chest or a r 6 cm has been suggested b | ology. In the Lugano classification, bulk in HL is defined as a mass mass >10 cm. For NHL, the recommended definitions of bulk vary by based on the FLIPI-2 and its validation. In DLBCL, cutoffs ranging from 5 |
| | te: A/B is no longer used in NHL. | econinienaea. | |
| | | | |
| 6 | Registry Data Collection Va | riables | |
| | <u> </u> | | more details on these variables |
| CLL | CLL and SLL should always be abstracted as lymphoma. See chapter for more details on these variables. | | |
| | | | |
| | 1. Size of the largest mass in millimet | ters for all stages; essentia | l for Stages I and II: |
| | 2. ALC >5,000 cells/μL: yes | □no | |
| | 3. Adenopathy (presence of lymph no | <u> </u> | es no |
| | | | |
| | 4. Organomegaly (enlarged liver and/or spleen on PE): yes no | | |
| | 5. Anemia (Hgb <11.0 g/dL): | | |
| | 6. Thrombocytopenia (Plt <100,000/ | μL): ges nc | |
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79.5. Non-Hodgkin Lymphomas: Chronic Lymphocytic Leukemia and Small Lymphocytic Lymphoma

7 Anatomy

FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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- 1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.
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| Hospital Name/Address | Patient Name/Information |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|--|--|---|--|
| workup information, until first treatment, including clinical history and symptoms, physical examination endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or | | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | |
| | rTNM Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

| Hospital Name/Address | Patient Name/Information |
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Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage | | Stage description | |
|--|--|--|--|
| ✓ | Limited stage | | |
| | 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) | |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) | |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm | |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other | |
| | | lymph node regions on the same side of the diaphragm | |
| | II bulky* | Stage II with disease bulk** | |
| | Advanced stage | | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm | |
| | | with spleen involvement | |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without | |
| | | associated lymph node involvement | |
| | | or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease | |
| | | or any extralymphatic organ involvement in nodal Stage III disease | |
| Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung le | | | |
| | | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other | |
| | than by direct extension in IIE disease). | | |
| *St | age II bulky may be considered either ear | ly- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion | |
| | HL prognostic factors). | | |
| ** | The definition of disease bulk varies accor | ding to the lymphoma histology. In the Lugano classification, ¹ bulk in HL is defined as a mass | |
| _ | | ter on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by | |
| lyn | lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation. ^{2,3} In DLBCL, cutoffs ranging from 5 | | |

⁶ Registry Data Collection Variables

to 10 cm have been used, although 10 cm is recommended.4

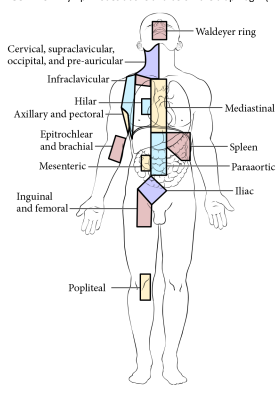
See chapter for more details on these variables.

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Note: A/B is no longer used in NHL.

| Hospital Name/Address | Patient Name/Information |
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FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

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4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage | | Stage description |
|-------|----------------|--|
| 1 | Limited stage | · |
| | I | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| | IE | Single extralymphatic site in the absence of nodal involvement (rare in HL) |
| | II | Involvement of two or more lymph node regions on the same side of the diaphragm |
| | IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other |
| | | lymph node regions on the same side of the diaphragm |
| | II bulky* | Stage II with disease bulk** |
| | Advanced stage | |
| | III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm |
| | | with spleen involvement |
| | IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without |
| | | associated lymph node involvement |
| | | or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease |
| | | or any extralymphatic organ involvement in nodal Stage III disease |
| | | Stage IV includes <i>any</i> involvement of the CSF, bone marrow, liver, or multiple lung lesions (other |
| | | than by direct extension in IIE disease). |
| *C+ | | , |

^{*}Stage II bulky may be considered either early- or advanced-stage disease based on lymphoma histology and prognostic factors (see discussion of HL prognostic factors).

Note: HL uses an A or B designation with stage group. A/B is no longer used in NHL

Select one:

| ✓ | Designation | Definition | |
|---|-------------|--|--|
| | Α | Asymptomatic (No B symptoms) | |
| | | | |
| | В | Any B symptom(s): | |
| | | Fevers. Unexplained fever with temperature above 38°C | |
| | | Night sweats. Drenching sweats (e.g., those that require change of bedclothes) | |
| | | 3. Weight loss. Unexplained weight loss of more than 10% of the usual body weight in the 6 months prior to | |
| | | diagnosis | |

6 Registry Data Collection Variables

See chapter for more details on these variables.

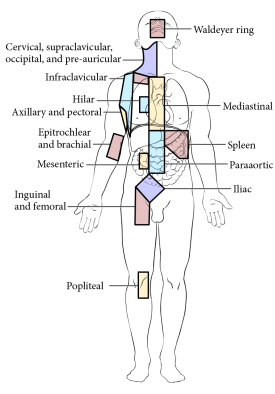
- 1. Size of the largest mass in millimeters for all stages; essential for Stages I and II:
- 2. A or B designation for symptoms must be part of the stage:
- 3. IPS:

| Hospital Name/Address | Patient Name/Information |
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^{**}The definition of disease bulk varies according to the lymphoma histology. In the Lugano classification, bulk in HL is defined as a mass greater than one third of the thoracic diameter on CT of the chest or a mass >10 cm. For NHL, the recommended definitions of bulk vary by lymphoma histology. In follicular lymphoma, 6 cm has been suggested based on the FLIPI-2 and its validation. DLBCL, cutoffs ranging from 5 to 10 cm have been used, although 10 cm is recommended.

7 Anatomy

FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
|---------------------|-----------|

8 Bibliography

- 1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.
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3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Lugano Classification for Hodgkin and Non-Hodgkin Lymphoma¹

| Stage Stage description | |
|--|---|
| ✓ Limited stage | |
| 1 | Involvement of a single lymphatic site (i.e., nodal region, Waldeyer's ring, thymus, or spleen) |
| IE | Single extralymphatic site in the absence of nodal involvement (rare in Hodgkin lymphoma) |
| II | Involvement of two or more lymph node regions on the same side of the diaphragm |
| IIE | Contiguous extralymphatic extension from a nodal site with or without involvement of other |
| | lymph node regions on the same side of the diaphragm |
| II bulky* | Stage II with disease bulk** |
| Advanced stage | |
| III | Involvement of lymph node regions on both sides of the diaphragm; nodes above the diaphragm |
| | with spleen involvement |
| IV | Diffuse or disseminated involvement of one or more extralymphatic organs, with or without |
| | associated lymph node involvement; |
| | or noncontiguous extralymphatic organ involvement in conjunction with nodal Stage II disease |
| | or any extralymphatic organ involvement in nodal Stage III disease |
| | Stage IV includes any involvement of the CSF, bone marrow, liver, or multiple lung lesions (other |
| | than by direct extension in IIE disease). |
| *Stage II bulky may be considered either early or advanced stage based on lymphoma histology and prognostic factors (see discussion of | |

^{*}Stage II bulky may be considered either early or advanced stage based on lymphoma histology and prognostic factors (see discussion of Hodgkin lymphoma prognostic factors).

Select one:

| ✓ | Designation | Definition | |
|---|-------------|--|--|
| | Α | Asymptomatic (No B symptoms) | |
| | | | |
| | В | Any B symptom(s): | |
| | | 1. Fevers. Unexplained fever with temperature above 38°C | |
| | | 2. Night sweats. Drenching sweats (e.g., those that require change of bedclothes) | |
| | | 3. Weight loss. Unexplained weight loss of more than 10% of the usual body weight in the 6 months prior to | |
| | | diagnosis | |

6 Registry Data Collection Variables

See chapter for more details on these variables.

- 1. Size of the largest mass in millimeters for all stages; essential for Stages I and II:
- 2. A or B designation for symptoms must be part of the stage:

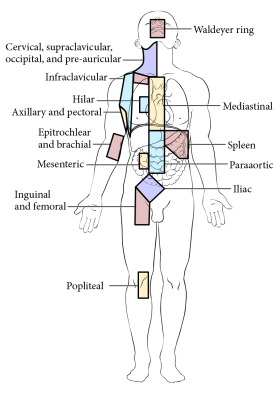
| Hospital Name/Address | Patient Name/Information |
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^{**}The definition of *disease bulk* varies according to lymphoma histology. In the Lugano classification, bulk in Hodgkin lymphoma is defined as a mass greater than one third of the thoracic diameter on CT of the chest or a mass >10 cm.

Note: Hodgkin lymphoma uses A or B designation with stage group.

7 Anatomy

FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
|---------------------|-----------|

8 Bibliography

1. Cheson BD, Fisher RI, Barrington SF, et al. Recommendations for initial evaluation, staging, and response assessment of Hodgkin and non-Hodgkin lymphoma: the Lugano classification. *J Clin Oncol.* 2014;32(27):3059-3068.

| ormation |
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2 Instructions

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| ✓ | Classification | Definition | |
|---|----------------|---|--|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations | |
| | pTNM | Pathological Classification: The use of the term <i>pathological staging</i> is reserved for patients who undergo staging laparotomy with an explicit intent to assess the presence of abdominal disease or to define histologic microscopic disease extent in the abdomen. As a result of improved diagnostic imaging, staging laparotomy and pathological staging generally are no longer performed. | |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. | |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). | |

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4 Definitions of AJCC TNM

TNM does not apply to this disease. Always refer to the specific chapter for explicit instructions on classification for this disease.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6 AJCC Prognostic Stage Groups

6.1 St. Jude Children's Research Hospital Staging System for Non-Hodgkin Lymphoma

| ✓ | Stage | Stage description |
|---|-------|---|
| | 1 | A single tumor (extranodal) or single anatomic area (nodal), with the exclusion of the |
| | | mediastinum or abdomen |
| | II | A single tumor (extranodal) with regional node involvement |
| | | Two or more nodal areas on the same side of the diaphragm |
| | | Two single (extranodal) tumors with or without regional node involvement on the same side of the diaphragm |
| | | A primary gastrointestinal tract tumor, usually in the ileocecal area, with or without involvement of associated mesenteric nodes only* |
| | III | Two single tumors (extranodal) on opposite sides of the diaphragm |
| | | Two or more nodal areas above and below the diaphragm |
| | | All the primary intrathoracic tumors (mediastinal, pleural, and thymic) |
| | | All extensive primary intra-abdominal disease* |
| | | All paraspinal or epidural tumors, regardless of other tumor site(s) |
| | IV | Any of the above with initial CNS and/or bone marrow involvement** |

^{*}A distinction is made between apparently localized gastrointestinal tract lymphoma versus more extensive intra-abdominal disease because of their quite different patterns of survival after appropriate therapy. Stage II disease typically is limited to a segment of the gut plus or minus the associated mesenteric nodes only, and the primary tumor can be completely removed grossly by segmental excision. Stage III disease typically exhibits spread to para-aortic and retroperitoneal areas by implants and plaques in mesentery or peritoneum, or by direct infiltration of structures adjacent to the primary tumor. Ascites may be present, and complete resection of all gross tumor is not possible.

Modified from Murphy SB.1

7 Registry Data Collection Variables

See chapter for more details on these variables.

1. Size of the largest mass in millimeters for all stages; essential for Stages I and II:

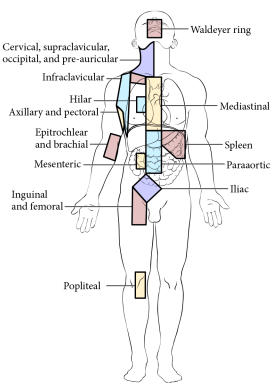
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^{**}If marrow involvement is present initially, the number of abnormal cells must be ≤25% in an otherwise normal marrow aspirate with a normal peripheral blood picture.

8 Anatomy

FIGURE 79.1. Lymph nodes above and below the diaphragm (Ann Arbor/Lugano classification).



Lymph nodes above the diaphragm

- 1. Waldeyer's ring
- 2. Cervical, supraclavicular, occipital, and pre-auricular
- 3. Infraclavicular
- 4. Axillary and pectoral
- 5. Mediastinal
- 6. Hilar
- 7. Epitrochlear and brachial

Lymph nodes below the diaphragm

- 8. Spleen
- 9. Mesenteric
- 10. Paraaortic
- 11. Iliac
- 12. Inguinal and femoral
- 13. Popliteal

| Physician Signature | Date/Time |
|---------------------|-----------|

9 Bibliography

1. Murphy SB. Classification, staging and end results of treatment of childhood non-Hodgkin's lymphomas: dissimilarities from lymphomas in adults. *Semin Oncol.* 1980;7(3):332-339.

| Hospital Name/Address | Patient Name/Information |
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| ✓ | Classification | Definition |
|---|----------------|---|
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| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |

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| Hospital Name/Address | Patient Name/Information |
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4 Definitions of AJCC TNM

Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T) (Skin)

ISCL/EORTC revision to the classification of mycosis fungoides and Sézary Syndrome

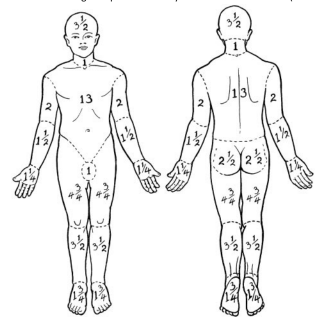
| ✓ | T Category | T Criteria |
|---|------------|---|
| | T1 | Limited patches,* papules, and/or plaques** covering <10% of the skin surface |
| | T1a | T1a (patch only) |
| | T1b | T1b (plaque ± patch) |
| | T2 | Patches, papules, or plaques covering ≥10% of the skin surface |
| | T2a | T2a (patch only) |
| | T2b | T2b (plaque ± patch) |
| | T3 | One or more tumors*** (≥1 cm in diameter) |
| | T4 | Confluence of erythema covering ≥80% of body surface area |

^{*}For skin, patch indicates any size skin lesion without significant elevation or induration. Presence/absence of hypo- or hyperpigmentation, scale, crusting, and/or poikiloderma should be noted.

^{***}For skin, *tumor* indicates at least one 1-cm diameter solid or nodular lesion with evidence of depth and/or vertical growth. Note the total number of lesions, total volume of lesions, largest size lesion, and region of body involved. Also note whether there is histologic evidence of large cell transformation. Phenotyping for CD30 is encouraged.

| 1 | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

FIGURE 81.1. Regional percent of body surface area in the adult (From Olsen et al., with permission).



| Patient Name/Information | |
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| | Patient Name/Information |

^{**}For skin, plaque indicates any size skin lesion that is elevated or indurated. Presence/absence of scale, crusting, and/or poikiloderma should be noted. Histologic features such as folliculotropism, large cell transformation (>25% large cells), and CD30 positivity or negativity, as well as clinical features such as ulceration, are important to document.

81.1. Primary Cutaneous Lymphoma: Mycosis Fungoides and Sézary Syndrome

4.2 Definition of Regional Lymph Node (N) (Node)

| ✓ | N Category | N Criteria |
|---|------------|--|
| | NX | Clinically abnormal peripheral lymph nodes; no histologic confirmation |
| | N0 | No clinically abnormal peripheral lymph nodes*; biopsy not required |
| | N1 | Clinically abnormal peripheral lymph nodes; |
| | | histopathology Dutch grade 1 or National Cancer Institute (NCI) LNO-2 |
| | N1a | Clone negative** |
| | N1b | Clone positive** |
| | N2 | Clinically abnormal peripheral lymph nodes; |
| | | histopathology Dutch grade 2 or NCI LN3 |
| | N2a | Clone negative** |
| | N2b | Clone positive** |
| | N3 | Clinically abnormal peripheral lymph nodes; |
| | | Histopathology Dutch grades 3–4 or NCI LN4; |
| | | clone positive or negative |

^{*}For node, abnormal peripheral lymph node(s) indicates any palpable peripheral node that on physical examination is firm, irregular, clustered, fixed or ≥1.5 cm in diameter. Node groups examined on physical examination include cervical, supraclavicular, epitrochlear, axillary, and inguinal. Central nodes, which generally are not amenable to pathological assessment, currently are not considered in the nodal classification unless used to establish N3 histopathologically.

^{**}A T-cell clone is defined by polymerase chain reaction (PCR) or Southern blot analysis of the TCR gene.

| ٧ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M) (Visceral)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| 1 | M Category | M Criteria |
|---|------------|--|
| | сМ0 | No visceral organ involvement |
| | cM1 | Visceral involvement (spleen and liver may be diagnosed by imaging criteria, and organ involved should be specified) |
| | pM1 | Visceral involvement (must have pathology confirmation, and organ involved should be specified) |

| Hospital Name/Address | Patient Name/Information |
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5 Prognostic Factors Required for Stage Grouping

5.1 Peripheral Blood Involvement (B)

| 1 | B Category | B Criteria |
|---|------------|--|
| | В0 | Absence of significant blood involvement: ≤5% of peripheral blood lymphocytes are |
| | | atypical (Sézary) cells* |
| | BOa | Clone negative** |
| | B0b | Clone positive** |
| | B1 | Low blood tumor burden: >5% of peripheral blood lymphocytes are atypical (Sézary) cells, |
| | | but does not meet the criteria of B2 |
| | B1a | Clone negative** |
| | B1b | Clone positive** |
| | B2 | High blood tumor burden: ≥1,000/μL Sézary cells* with positive clone** |

^{*}For blood, Sézary cells are defined as lymphocytes with hyperconvoluted cerebriform nuclei. If Sézary cells cannot be used to determine tumor burden for B2, then one of the following modified ISCL criteria, along with a positive clonal rearrangement of the TCR, may be used instead: (1) expanded CD4+ or CD3+ cells with a CD4/CD8 ratio of ≥10, or (2) expanded CD4+ cells with abnormal immunophenotype, including loss of CD7 or CD26

From Olsen et al., with permission from the American Society of Hematology. 1

6 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

6.1 ISCL/EORTC Revision to the Staging of Mycosis Fungoides and Sézary Syndrome

| 1 | When T is | And N is | And M is | And B is | Then the stage |
|---|-----------|----------|----------|----------|----------------|
| | | | | | group is |
| | T1 | NO | M0 | B0,1 | IA |
| | T2 | N0 | M0 | B0,1 | IB |
| | T1,2 | N1,2 | M0 | B0,1 | IIA |
| | T3 | N0-2 | M0 | B0,1 | IIB |
| | T4 | N0-2 | M0 | B0,1 | III |
| | T4 | N0-2 | M0 | ВО | IIIA |
| | T4 | N0-2 | M0 | B1 | IIIB |
| | T1-4 | N0-2 | M0 | B2 | IVA1 |
| | T1-4 | N3 | M0 | B0-2 | IVA2 |
| | T1-4 | N0-3 | M1 | B0-2 | IVB |

From Olsen et al., 1 with permission from the American Society of Hematology.

| Hospital Name/Address | Patient Name/Information |
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^{**}A T-cell clone is defined by PCR or Southern blot analysis of the TCR gene.

81.1. Primary Cutaneous Lymphoma: Mycosis Fungoides and Sézary Syndrome **Registry Data Collection Variables** See chapter for more details on these variables. Peripheral blood involvement: Physician Signature Date/Time 8 **Bibliography** Olsen E, Vonderheid E, Pimpinelli N, et al. Revisions to the staging and classification of mycosis fungoides and Sezary syndrome: a proposal of the International Society for Cutaneous Lymphomas (ISCL) and the cutaneous lymphoma task force of the European Organization of Research and Treatment of Cancer (EORTC). Blood. 2007;110(6):1713-1722.

| Hospital Name/Address | Patient Name/Information | |
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81.2. Primary Cutaneous Lymphoma: Primary Cutaneous B-Cell/T-Cell (non-MF/SS) Lymphoma

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This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Time of Classification (select one):

| 1 | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |

| Hospital Name/Address | Patient Name/Information |
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81.2. Primary Cutaneous Lymphoma: Primary Cutaneous B-Cell/T-Cell (non-MF/SS) Lymphoma

4 Definitions of AJCC TNM

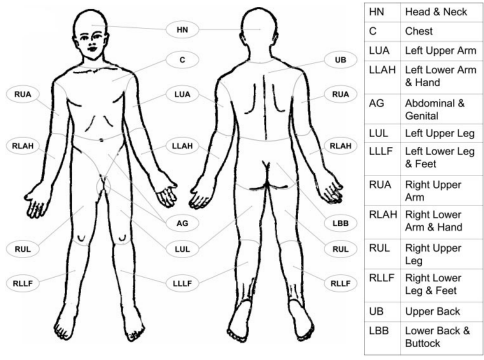
Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

4.1 Definition of Primary Tumor (T)

| 1 | T Category | T Criteria |
|---|------------|---|
| | T1 | Solitary skin involvement |
| | T1a | Solitary lesion <5 cm |
| | T1b | Solitary lesion ≥5 cm |
| | T2 | Regional skin involvement: multiple lesions limited to one body region or two contiguous body regions |
| | T2a | All disease encompassing in a <15-cm circular area |
| | T2b | All disease encompassing in a ≥15-cm and <30-cm circular area |
| | T2c | All disease encompassing in a ≥30-cm circular area |
| | T3 | Generalized skin involvement |
| | T3a | Multiple lesions involving 2 noncontiguous body regions |
| | T3b | Multiple lesions involving ≥3 body regions |

| ✓ | T Suffix | Definition |
|---|----------|---|
| | (m) | Select if synchronous primary tumors are found in single organ. |

FIGURE 81.2. Body regions as defined in the proposed TNM system for designating T (skin involvement) category. Left and right extremities are assessed as separate body regions. The designation of these body regions are based on regional lymph node drainage patterns (From Kim et al., with permission).



| Hospital Name/Address | Patient Name/Information |
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81.2. Primary Cutaneous Lymphoma: Primary Cutaneous B-Cell/T-Cell (non-MF/SS) Lymphoma

4.2 Definition of Regional Lymph Node (N)

| √ | N Category | N Criteria |
|----------|------------|--|
| | NX | Regional lymph nodes cannot be assessed |
| | N0 | No clinical or pathological lymph node involvement |
| | N1 | Involvement of one peripheral node region that drains an area of current or prior skin involvement |
| | N2 | Involvement of two or more peripheral node regions or involvement of any lymph node region that does not drain |
| | | an area of current or prior skin involvement |
| | N3 | Involvement of central nodes |

| ✓ | N Suffix | Definition |
|---|----------|--|
| | (sn) | Select if regional lymph node metastasis identified by SLN biopsy only. |
| | (f) | Select if regional lymph node metastasis identified by FNA or core needle biopsy only. |

4.3 Definition of Distant Metastasis (M)

The terms pM0 and MX are NOT valid categories in the TNM system. Assignment of the M category for clinical classification may be cM0, cM1, or pM1. Any of the M categories (cM0, cM1, or pM1) may be used with pathological stage grouping.

| √ | M Category | M Criteria |
|----------|------------|---|
| | сМ0 | No evidence of extracutaneous non–lymph node disease |
| | cM1 | Extracutaneous non–lymph node disease present |
| | pM1 | Extracutaneous non-lymph node disease present, microscopically proven |

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

There is no stage group for other primary cutaneous lymphomas – including cutaneous T-cell, B-cell, NK cell and non-MF/SS lymphoma – at this time

| Physician Signature | Date/Time |
|------------------------|-----------|
| riiysiciaii sigilature | Date/Time |

6 Bibliography

 Kim YH, Willemze R, Pimpinelli N, et al. TNM classification system for primary cutaneous lymphomas other than mycosis fungoides and Sezary syndrome: a proposal of the International Society for Cutaneous Lymphomas (ISCL) and the Cutaneous Lymphoma Task Force of the European Organization of Research and Treatment of Cancer (EORTC). Blood. 2007;110(2):479-484.

| Hospital Name/Address | Patient Name/Information |
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3 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
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| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
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82. Plasma Cell Myeloma and Plasma Cell Disorders

4 Definitions of AJCC TNM

TNM does not apply to this classification. Always refer to the specific chapter for explicit instructions on clinical and pathological classification for this disease.

5 AJCC Prognostic Stage Groups

Always refer to the specific chapter for rules on clinical and pathological classification of this disease.

5.1 Revised International Staging System (RISS) Adopted by the International Myeloma Working Group

| ✓ | RISS stage group | Factors | |
|---|------------------|---|--|
| | Stage I | Serum β ₂ -microglobulin <3.5 mg/L | |
| | | and | |
| | | serum albumin ≥3.5 g/dL | |
| | | and | |
| | | no high-risk cytogenetics* | |
| | | and | |
| | | Normal LDH | |
| | Stage II | Not stage I or III | |
| | | | |
| | Stage III | Serum β₂-microglobulin ≥5.5 mg/L | |
| | | and | |
| | | high-risk cytogenetics* | |
| | | and/or | |
| | | high LDH | |

^{*}High-risk cytogenetics consist of one or more of the following: del17p, t(4;14), or t(14;16).

Note: The following variables must be collected at the time of diagnosis for staging of multiple myeloma according to the RISS: serum β_2 -microglobulin, serum albumin, serum LDH, and FISH results from the bone marrow specimen for t(4;14), t(14;16), and del17p.

Adapted from Palumbo et al.1

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
| | |
| | |
| | |
| | |

| 6 | Re | gistry Data Co | llection Variab | les | | | |
|------|---------|--|--------------------------|---|---------------------------|---|--------------------------|
| See | chapt | er for more details o | n these variables. | | | | |
| | 1. | ISS stage group (if documented): | | | | | |
| | 2. | Imaging elements: | bone disease demon | strated on imaging, p | lain film (skeletal surve | y), CT, MR imaging, PE | т/ст: |
| | 3. | Number of bone le | sions identified on in | naging: no | ne one mo | re than one | |
| | 4. | Hemoglobin; all me | easurements are pret | reatment: | | | |
| | 5. | Serum β ₂ -microglo | bulin in milligrams pe | er liter, xx.x; all measu | rements are pretreatm | ent: | |
| | 6. | Serum albumin in g | grams per deciliter, x | x; all measurements | are pretreatment: | | |
| | 7. | Serum calcium in n | nilligrams per decilite | er, xx.x; all measureme | ents are pretreatment: | | |
| | 8. | Serum creatinine in | n milligrams per decil | iter, x.x; all measuren | nents are pretreatment | : | |
| | 9. | LDH, normal or abo | ove normal, xx,xxx un | its per liter; all measu | rements are pretreatm | ent: | |
| | 10. | IgG in milligrams po | er deciliter, xx,xxx; al | I measurements are p | retreatment: | | |
| | 11. | IgA in milligrams pe | er deciliter, xx,xxx; al | measurements are p | retreatment: | | |
| | 12. | IgM in milligrams p | er deciliter, xx,xxx; a | II measurements are p | oretreatment: | | |
| | 13. | Monoclonal protein measurements are | | urine (M spike): gran | ns per deciliter for seru | m, xx.x; grams for 24-h | our urine, xx.x; all |
| | 14. | Serum free kappa I measurements are | | ams per liter, xx,xxx (| milligrams per deciliter | × 10 to convert to gra | ms per liter); all |
| | 15. | Serum free lambda measurements are | • | grams per liter, xx,xxx | (milligrams per decilite | $\mathrm{er} 	imes 10$ to convert to gr | ams per liter); all |
| | 16. | Cytogenetics: | t(4;14) | □t(14;16) | t(14;20) | t(11;14) | t(6;14) |
| | | | add1q | del1p | del17p | trisomy 3 | trisomy 5 |
| | | | trisomy 7 | trisomy 9 | trisomy 11 | trisomy 15 | trisomy 19 |
| | | | trisomy 21 | <u></u> , | ш , | _ , | |
| | | | | | | | |
| Phys | ician | Signature | | | | Date/Time | |
| 7 | | bliography | | | | · | |
| 1. | | Palumbo A, Avet-Lo | | ıl. Revised Internation 2015;33(26):2863-286 | | Iultiple Myeloma: A Re | eport From International |
| Hosi | oital I | Name/Address | | | Patient Name/Inform | ation | |
| | | | | | · | | |

The cancer staging form is a specific document in the patient record; it is not a substitute for documentation of history, physical examination, and staging evaluation, or for documenting treatment plans or follow-up. The staging forms available in conjunction with the *AJCC Cancer Staging Manual, Eighth Edition* may be used by individuals without permission from the ACS or the publisher. They cannot be sold, distributed, published, or incorporated into any software (including any electronic record systems), product, or publication without a written license agreement with ACS. The forms cannot be modified, changed, or updated without the express written permission of ACS.

2 Instructions

See Principles of Cancer Staging (Chapter 1) of the AJCC Cancer Staging Manual, Eighth Edition for complete staging rules. Always refer to the respective chapter in the Manual for disease-specific rules for classification, as this form is not representative of all rules, exceptions and instructions for this disease.

This form may be used by physicians to record data on T, N, and M categories; prognostic stage groups; additional prognostic factors; cancer grade; and other important information. This form may be useful for recording information in the medical record and for communicating information from physicians to the cancer registrar.

The staging form may be used to document cancer stage at <u>different points in the patient's care</u> and during the course of therapy, including before therapy begins, after surgery and completion of all staging evaluations, or at the time of recurrence. It is best to use a separate form for each time point staged along the continuum for an individual cancer patient. However, if all time points are recorded on a single form, the staging basis for each element should be identified clearly.

This form may provide more data elements than required for collection by standard setters such as NCI SEER, CDC NPCR, and CoC NCDB.

3 Select Diagnosis

This form may be used for the following diagnoses discussed in the AJCC Cancer Staging Manual, Eighth Edition.

| ✓ | Diagnosis |
|---|---|
| | 83.1 Acute Myeloid Leukemia |
| | 83.2 Acute lymphoblastic leukemia in children |
| | 83.3 Acute Lymphocytic Leukemia in Adults |
| | 83.4 Chronic Myeloid Leukemia |
| | 83.0 Unspecified or Other Type of Leukemia |

4 Time of Classification (select one):

| ✓ | Classification | Definition |
|---|----------------|---|
| | cTNM or TNM | Clinical Classification: Used for all patients with cancer identified before treatment. It is composed of diagnostic workup information, until first treatment, including clinical history and symptoms, physical examination, imaging, endoscopy, biopsy of the primary site, biopsy or excision of a single regional node or sentinel nodes, or sampling of regional nodes, with clinical T, biopsy of distant metastatic site, surgical exploration without resection, and other relevant examinations |
| | rTNM | Recurrence or Retreatment Classification: Used for assigning stage at time of recurrence or progression until treatment is initiated. |
| | aTNM | Autopsy Classification: Used for cancers not previously recognized that are found as an incidental finding at autopsy, and not suspected before death (i.e., this classification does not apply if an autopsy is performed in a patient with a previously diagnosed cancer). |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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5 Prognostic Factors Required for Clinical Care

5.1 Acute Myeloid Leukemia

5.1.1 Age:

5.1.2 Zubrod performance status (PS)

| ✓ | PS | Definition |
|---|--------|------------------------|
| | 0 or 1 | Minimal symptoms |
| | 2 | Between 1 & 3 |
| | 3 | In bed 50-100% of time |
| | 4 | Bed ridden |

5.1.3 Hematopoietic cell transplantation comorbidity index (HCT-CI):

5.1.4 Cytogenetics (20 metaphase):

| ✓ | Description |
|---|--------------|
| | Favorable |
| | Intermediate |
| | Adverse |

5.1.5 Status of NPM, FLT3 and CEBPA genes:

| ✓ | Status |
|---|---|
| | NPM1 mutation in absence FLT3 internal tandem duplication |
| | Bi allelic CEBPA mutation |
| | FLT3 internal tandem duplication |

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

5.2 Acute Lymphoblastic Leukemia in Children

| 5.2.1 | n | 0: |
|-------|--------|----|
| | ١g | С. |

| ✓ | Age |
|---|--------------|
| | 1- <10 years |
| | ≥10 years |

5.2.2 WBC count at diagnosis ($<50,000 \text{ to } \ge 50,000 \text{ }\mu\text{L}$):

5.2.3 Timmunophenotype:

| ✓ | T Immunophenotype |
|---|-------------------|
| | CD5 |
| | CD7 |
| | CD8 |
| | CD4 |
| | CD2 |
| | CD1a |

| | 03101 | The second second second | 2 T T | |
|-------|--------|--------------------------|-------------|------------|
| 5.2.4 | CNS in | volvement | (hlacte on | cvtospin): |
| | | | | |

- 5.2.5 Hyperdiploidy (>50-67 chromosomes or specific trisomies (e.g. 4 and 10):
- 5.2.6 t(12;21) (p13:q22) EVT6/RUNX1 (Cryptic translocation detected by FISH, RT-PCR):
- 5.2.7 Hypodiploidy (<44 chromosomes by karyotype):
- 5.2.8 *MLL* rearrangements (Karotype or FISH (>100 fusion partners defined)):
- 5.2.9 iAMP21 (Three or more extra copies of *RUNX1* on an abnormal chromosome 21):
- 5.2.10 t(9;22)(q24;q11) Ph+ (FISH or karyotype):
- 5.2.11 MRD (flow cytometry or antigen receptor/fusion gene PCR):

| Hospital Name/Address | Patient Name/Information |
|-----------------------|--------------------------|
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| | |

| 5.3 | Acute L | ymphoc | ytic Leukemia | in Adults |
|-----|---------|--------|---------------|-----------|
|-----|---------|--------|---------------|-----------|

| 5.3.1 CNS involvement (presence of bla | asts in cerebrospinal fluid): |
|--|-------------------------------|
|--|-------------------------------|

| _ | 3.2 | Testicular involvement | st (tooti oulou moo | a of blacks on biomerr | A - |
|---|-----|------------------------|---------------------|----------------------------|-----|
| | | | | | |
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|-----------------------|--------------------------|
| Hospital Name/Address | Patient Name/Information |
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| 5.4 | Chronic Myeloid Leukemia |
|----------|--|
| 5.4.1 | Bone marrow (blast count): |
| 5.4.2 | Cytogenetics, Ph chromosome: |
| 5.4.3 | Cytogenetics, additional clonal changes: |
| | |
| | |
| | |
| Physicia | n Signature Date/Time |
| | |

| Hospital Name/Address | Patient Name/Information |
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| | |

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