

**Mississippi Cancer
Registry
Reporting Manual
Revised 2021**

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PROGRAM OVERVIEW

The Mississippi Cancer Registry (MCR) Reporting Manual has been created to assist hospital registries in reporting cancer cases to the MCR. This manual is being implemented due to the requirements from the National Program of Cancer Registries (NPCR), the North American Association for Central Cancer Registries (NAACCR) and the Commission on Cancer (CoC) Facility Oncology Required Data Standards (FORDS). There are also clarifications and rules that are in place in order to accurately complete abstraction of cancer cases.

In October 1992, Congress enacted the Cancer Registries Amendment Act (<http://www.cdc.gov/cancer/npcr/npcrpdfs/publaw.pdf>) that authorized the Centers for Disease Control and Prevention to establish a national program to support cancer registries. One of the goals of this program was to help establish statewide population-based cancer registries that would meet the minimum standards for quality and completeness set forth by the North American Association of Central Cancer Registries (NAACCR). In 1993, the Mississippi State Legislature authorized the Mississippi State Department of Health (MSDH) to establish a central population-based incidence cancer registry for the state of Mississippi (<http://mcr.umc.edu/documents/StateLaw.pdf>)

Cancer registration is an important and fundamental tool in assessing the true extent of cancer in Mississippi. The data collected through a statewide cancer registry can be used for epidemiological studies, medical research, and cancer control planning. This information is a valuable and essential tool in the identification of populations at high risk for cancer, the monitoring of cancer incidence trends, the facilitation of studies related to cancer prevention, the evaluation of cancer control initiatives, and the development of educational awareness programs.

The rules and regulations governing reporting of cancer cases to the Central Cancer Registry were presented to the State Department of Health for final adoption in April 1995. These rules and regulations which created a new Class IV level of reportable diseases became effective May 1995. Health Department regulations require that a cancer case be reported within six months of the first date of contact with the patient for the reportable condition.

The Mississippi Cancer Registry began collecting data on January 1, 1996. The MCR collects data that: 1) are compliant with required NPCR data elements; 2) are standard requirements designated by NAACCR for incidence reporting and endorsed by CDC; 3) assist in determining data quality; and 4) provide useful information, feedback and assistance to submitting facilities.

CONFIDENTIALITY

Per State Cancer Law (41-91-11) “Data obtained under this chapter directly from the medical records of a patient is for the confidential use of the department and the persons or public or private entities that the department determines are necessary to carry out the intent of this chapter. The data is privileged and may not be divulged or made public in a manner that discloses the identity of an individual whose medical records have been used for obtaining data

under this chapter.”

DISCLOSURE OF DATA

The Mississippi Cancer Registry (MCR) may exchange patient-specific information with the reporting facility or clinical facility for the purpose of completing a case record, provided these facilities comply with all MCR confidentiality policies.

To achieve complete case ascertainment, the MCR may exchange patient-specific information with other state cancer registries if reciprocal data sharing agreements and confidentiality provisions are in place. The MCR may not provide information on a patient of a hospital that was obtained from another hospital (i.e., follow-up information).

The Mississippi Cancer Registry Advisory Committee, formed in December 2006, was tasked with developing the protocol for release of information for the purpose of research. The Committee consists of medical representatives from the different disciplines related to cancer, cancer registry professionals and also experts on human subject research. This protocol is complete and available to researchers who are interested in using registry data. Since the MCR is charged with data collection and not authorized to use funds for research, there is a fee for researchers that want to use the MCR data. The protocol and fee schedule are located on the MCR web site (<http://mcr.umc.edu>). Data obtained from the VA hospitals, Keesler Medical Center and certain other states cannot be used for research.

GENERAL INSTRUCTIONS

The following information provides some basic rules regarding cancer reporting to the MCR.

- A. Healthcare providers including, but not limited to, hospitals, ambulatory surgery centers, laboratories, radiation therapy facilities, oncology facilities and physician offices are required to report cancer cases to the MCR. Hospitals need to abstract inpatient and outpatient cancer cases.
- B. All required data items should be collected and reported to the MCR. The list is based on the rules and regulations of NPCR and NAACCR.
- C. The ICD-O-3 coding scheme must be used for site and histology for cases diagnosed on or after January 1, 2001. The ICD-O-2 coding scheme must be used for cases diagnosed prior to January 1, 2001.
- D. The Collaborative Staging Manual is to be used for cases diagnosed between January 1, 2004 and December 31, 2015. Site Specific Factors will continued to be used for cases diagnosed January 1, 2016 to December 31, 2017. AJCC TNM Staging is to be used for cases diagnosed January 1, 2016 to December 31, 2017. The SEER Summary Staging Manual – 2000 is to be used for staging for cases diagnosed between January 1, 2001 and December 31, 2003 and cases diagnosed January 1, 2015 to December 31, 2017. The SEER Summary Staging Manual – 2018 is to be used for staging cases diagnosed January 1, 2018 and forward. The SEER Summary Staging Guide, 1986 reprint, is to be used for cases diagnosed prior to January 1, 2001.

- E. The 2007 Multiple Primary and Histology Coding Rules Manual is to be used for cases diagnosed January 1, 2007 to December 31, 2017. The 2018 Solid Tumor Rules should be used for cases diagnosed January 1, 2018 and forward. The Hematopoietic and Lymphoid Database should be used for all hematopoietic diseases diagnosed January 1, 2010 and later. Appendix A of the FORDS 2016 manual should be used to determine multiple primaries for hematopoietic diseases diagnosed before January 1, 2010.
- F. All cases diagnosed and/or treated for cancer in your facility on or after January 1, 1996, must be abstracted and reported to the MCR.
- G. Completed cases should be submitted to the MCR within six months of date of initial diagnosis for Class of Case 00 through 14 and within six months of the date of first contact for Class of Case 20 through 22.
- H. All pathology reports that are read by hospital pathology laboratories, but the cases are not the responsibility of the hospital registry to abstract, should be forwarded to MCR for further investigation. MCR will be responsible for contacting the physicians on the pathology reports to obtain the information needed to include the case in the registry database.
- I. It is important for all reporting facilities to submit data monthly. This will ensure that all data can be processed and submitted to the CDC by the established deadlines.

QUALITY CONTROL

Edits

All cases must pass edits when submitted to the MCR. Edits are built into Web Plus for data entry. No case can be released to the MCR from Web Plus if there are edits on that case. For file uploaders, edits will be run on your submission file. A copy of the state edit set has been provided to all hospital vendors. Since there is often a lag time in getting the edit set added to the hospital software, files with less than 10% edits will be accepted. Otherwise, the file will be rejected and require resubmission after the edits are corrected. The edit report will be provided to the facility.

Visual Review of Cases

All cases submitted to the MCR will be subject to a visual review of key data items. Items include age at diagnosis, sex, race fields, Spanish/Hispanic origin, date of diagnosis, primary site, laterality, histology, grade, staging fields, class of case, dates of first course treatment and treatment data fields. Other items may be included as time and resources permit. The items will be compared to the text submitted, so good text is important to this process. If problems are identified or codes are not justified in text, the facility will be contacted. This process serves to ensure the registry has high quality data.

Audits

The MCR Auditor conducts annual case finding and quality assurance (re-abstracting) audits as

required by NPCR. The purpose of these audits is to ensure that all reportable cases are being identified and reported to the cancer registry and that all information submitted to the registry is of good quality and accurately coded. The audits are scheduled in advance to enable the facilities to prepare for the arrival of the registry staff or to set up remote access for the auditor. Case finding audits require a medical record disease index that is usually reviewed before the audit begins. Reviewing of charts, pathology case finding and re-abstracting are performed on site or by remote access to the facility's electronic medical records. A report is provided to the medical record director and administrator which summarize the percentage of case ascertainment or completeness and any suggestions that would help to improve the reporting process.

Case finding audits are performed on inpatient and outpatient disease indices, pathology reports and other pertinent case finding documents such as: clinic sign-in logs, surgery logs, etc. These documents are reviewed and all reportable codes are compared with the MCR database for the facility being audited. All cases that are not identified in the database will have to be reconciled by the registrar at the audited facility. The registrar will have a minimum of 30 days to complete the reconciliation process and return an updated list to MCR with reasons why the identified cases were either not abstracted or not reportable. Cases that are reportable must be abstracted into their database and submitted to the MCR. All cases that were either diagnosed prior to January 1, 1996 or diagnosis/treatment was not performed at the reporting facility are removed from the reconciliation log and a percentage is calculated at that time. A report is sent to the medical record director and the administrator of the facility that summarizes the percentage of case ascertainment and provides suggestions to help improve the case ascertainment process.

Quality assurance or re-abstracting audits consist of the MCR Auditor re-abstracting specific fields selected by MCR and comparing with the original data that has been submitted. Any discrepancies are documented and sent to the audited facility in a summary report. Exceptions are taken into consideration if a case has been merged in the MCR database and the audited facility did not have this information. This could indicate that the other procedures were done elsewhere and not available to the audited facility at the time of abstraction.

REPORTING PROCEDURE

Reporting facilities with 25 or less confirmed cases per year may elect to report their cases on paper. Copies of the following from the medical record need to be sent in for each identified cancer case: Face Sheet, History and Physical, Operation Reports, Scans, X-Rays, Pathology, Chemotherapy, Radiation, and Name of Referring Physician. The paper reporting form should no longer be completed. These records need to be submitted on a monthly basis. It is recommended that the facility keep a copy of what is submitted to prevent duplicate case reporting. The submissions should be faxed to 601-815-5483. The fax machine is the property of the Mississippi Cancer Registry, so the data will not be received or viewed by anyone other than our staff. If you cannot fax the information, you may mail it to the following address:

Mississippi Cancer Registry
2500 N. State Street
Jackson, MS 39261-4583

Facilities with no cases for a given month need to send a letter to the MCR stating that there were no cases to report. For facilities that frequently have no cases, quarterly reporting is acceptable.

Facilities with more than 25 confirmed cases of cancer must report electronically monthly. Facilities with their own cancer registry software should submit a file of cases in the appropriate NAACCR layout to the secure website, <https://mscrrweb1.umc.edu/webplus/LogOnEn.aspx>. Facilities using Web Plus for direct data entry will abstract their cases and correct edit errors. Once the cases are complete and free of edits, the facility will release the abstracts to the MCR.

CASE FINDING

Cases can be identified via many sources. The pathology reports can provide cases diagnosed by histology, cytology, hematology, bone marrow, or autopsy. Other sources are clinic admission logs, daily discharges, disease indices, inpatient and outpatient surgery logs, radiotherapy consults, treatment reports and logs and oncology clinic treatment reports and logs. The pathology reports should never be the only source of case finding, due to the fact that cases not diagnosed, only treated at your facility, may not have a path report. Also, some cases are clinical diagnoses only. Oncology clinic logs will be a good source in locating these cases. Cases not diagnosed histologically will be either confirmed by the physician in the patient's medical record or on the medical record disease index. A system should be established that would enable you to review a copy of the disease index.

REPORTABLE NEOPLASMS

The Mississippi Cancer Registry produces lists of reportable conditions containing the ICD-9-CM codes and ICD-10-CM codes depending on year to aid hospitals in case finding. The reportable conditions do not change that often. However, the ICD-10-CM codes for reportable conditions may change as frequently as annually. We update the reportable list annually to reflect those changes, if necessary. The reportable list in its entirety will not be included in this manual since all of the hospitals are not abstracting the same month and year and may need different lists based on what month and year of cancer cases they are abstracting. The most current and all historical reportable lists can be found on our web site at the following link:

https://www.umc.edu/Administration/Outreach_Services/Mississippi_Cancer_Registry/Reportable_Diseases.aspx.

Not only do these reportable lists contain information on what is reportable and the ICD-9-CM or ICD-10-CM codes for those conditions, but they also include conditions that should be

excluded and conditions or procedures that should be screened for a reportable cancer case. Additionally, they provide a list of ambiguous terms that should be used to help in determining if a case is reportable.

Diagnosis Prior to Birth

Reportability requirements apply to diagnoses made in utero. Diagnoses made in utero are reportable only when the pregnancy results in a live birth. If you have no indication in the record of still birth, abortion or fetal death, assume that there was a live birth. When a reportable condition is confirmed prior to birth and disease is not evident at birth due to regression, report the case based on the pre-birth diagnosis.

Reportability Examples

1. Positive histology from a needle aspiration/biopsy followed by negative resection. This case is reportable based on the positive needle biopsy.
2. Ovarian mucinous borderline tumor with foci of intraepithelial carcinoma. This case is reportable because there are foci of intraepithelial carcinoma (carcinoma in situ)
3. “Squamous cell carcinoma of the anus, NOS.” Squamous cell carcinoma of the anus is reportable unless the primary site is confirmed to be the skin of the anus.

Not Reportable Examples

- Left thyroid lobectomy shows microfollicular neoplasm with evidence of minimal invasion. Micro portion of path report states “The capsular contour is focally distorted by a finger of the microfollicular nodule which appears to penetrate into the adjacent capsular and thyroid tissue.” Do not report this case based on the information provided. There is no definitive statement of malignancy. Search for additional information in the record. Contact the pathologist or the treating physician.

Cases Diagnosed Clinically are Reportable

In the absence of histologic or cytologic confirmation of a reportable cancer, accession a case based on the clinical diagnosis (when a recognized medical practitioner says the patient has cancer or carcinoma). A clinical diagnosis may be recorded in the final diagnosis, on the face sheet or other parts of the medical record.

A pathology report normally takes precedence over a clinical diagnosis. If the patient has a negative biopsy, the case would not be reported.

Exception 1: If the physician treats a patient for cancer in spite of the negative biopsy, report the case.

Exception 2: If enough time has passed that it is reasonable to assume that the physician has seen the negative pathology, but the clinician continues to call this a reportable disease, report

the case. A reasonable amount of time would be equal to or greater than 6 months.

Brain or CNS “Neoplasms”

A brain or CNS ‘neoplasm’ identified by diagnostic imaging is reportable even when no other information is available (from biopsy or resection, for example).

Ambiguous Terminology

Ambiguous terminology may originate in any source document, such as a pathology report, radiology report or clinical report. The terms listed below are reportable.

Ambiguous terms that are reportable:

- Apparent(ly)
- Appears
- Comparable with
- Compatible with
- Consistent with
- Favor(s)
- Malignant appearing
- Most likely
- Presumed
- Probable
- Suspect(ed)
- Suspicious (for)
- Typical (of)

Do not substitute synonyms such as “supposed” for “presumed” or “equal” for “comparable”. Do not substitute “likely” for “most likely.” There may be ambiguous terms preceded by a modifier, such as “mildly” suspicious. In general, ignore modifiers or other adjectives and accept the reportable ambiguous term.

This list of ambiguous terms needs to be used correctly. The first and foremost resource for the registrar for a questionable case is the physician who diagnosed the tumor. The ideal way to approach abstracting situations when the medical record is not clear is to follow up with the physician. If the physician is not available, the medical record, and any other pertinent reports (e.g, pathology, etc.) should be read closely for the required information. The purpose of the Ambiguous Terminology lists is so that in the case where wording in the patient record is ambiguous with respect to reportability and no further information is available from any resource, registrars will make consistent decisions. When there is a clear statement of malignancy, then registrars should not refer to the list of ambiguous terms. Registrars should only rely on this list when the situation is not clear and the case cannot be discussed with the appropriate physician/pathologist. **Please document any conversations you have about reportability with a physician in your text.**

Do **not** report a case based solely on **suspicious** cytology. Follow back on cytology diagnoses

using ambiguous terminology. Report the case when a reportable diagnosis is confirmed later. The date of diagnosis is the date of the later confirmation in this situation.

Note: "Suspicious cytology" means any cytology report diagnosis that uses an ambiguous term listed on the ambiguous terminology for reportable cancers list.

Cytology refers to the microscopic examination of cells in body fluids obtained from aspirations, washings, scrapings, and smears; usually a function of the pathology department.

Report cases with cytology diagnoses that are **positive** for malignant cells.

Urine cytology positive for malignancy is reportable. Code the primary site to C689 in the absence of any other information.

How to Use Ambiguous Terminology for Case Ascertainment

1. If any of the reportable ambiguous terms precede a word that is synonymous with an in situ or invasive tumor, report the case.

Example: The pathology report says: Breast biopsy with abnormal cells consistent with ductal carcinoma. Report the case.

Negative example: The final diagnosis reads: Rule out lung cancer. Do not report this case.

2. For benign and borderline primary intracranial and CNS tumors, report the case if any reportable ambiguous term precedes the either the word "tumor" or the word "neoplasm."

- a. "Mass" and "lesion" are not reportable terms for intracranial and CNS because they are not listed in ICD-O-3.2 with behavior codes of /0 or /1.

3. Discrepancies

- a. Report the case based on the reportable ambiguous term when there are reportable and non-reportable ambiguous terms in the medical record.
 - i. Do not report a case when subsequent documents refer to a history of cancer and the original source document used a non-reportable ambiguous term.

Example: Impression from a CT scan of the chest states probable malignant neoplasm of the lung. Discharge diagnosis states possible lung cancer. Report this case because probable lung cancer makes this case reportable

- b. When there is a single report, accept the reportable term and report the case when one section of a report uses a reportable term such as “apparently” and another section of the same report uses a term that is not on the reportable list.

Example: Abdominal CT reveals a 2cm liver lesion. “The lesion is consistent with hepatocellular carcinoma” appears in the discussion section of the report. The final diagnosis is “2 cm liver lesion, possibly hepatocellular carcinoma.” Report this case. “Consistent with” is reportable.

Exception: Do not accession a case based ONLY on suspicious cytology.

- c. Use these terms when screening diagnoses on pathology reports, operative reports, scans, mammograms, and other diagnostic testing other than tumor markers.
 - i. Do not report a case when resection, excision, biopsy, cytology, or physician’s statement proves the ambiguous diagnosis is not reportable.

Example 1: Stereotactic biopsy of the left breast is “focally suspicious for DCIS” and is followed by a negative needle localization excisional biopsy. Do not report the case. The needle localization excisional biopsy was performed to further evaluate the suspicious stereotactic biopsy finding. The suspicious diagnosis was proven to be false.

Example 2: CT report states “mass in the right kidney, highly suspicious for renal cell carcinoma.” CT-guided needle biopsy with final diagnosis “Neoplasm suggestive of oncocytoma. A malignant neoplasm could not be excluded.” Discharged back to the nursing home and no other information is available. Do not report the case. The suspicious CT finding was biopsied and not proven to be malignant. “Suggestive of” is not a reportable ambiguous term.

Hematopoietic and Lymphoid Neoplasms

For cases diagnosed January 1, 2010 or later, see the Reportability Instructions in the *Hematopoietic and Lymphoid Neoplasm Coding Manual and Database* (<http://seer.cancer.gov/tools/heme/index.html>).

CORRECTIONS, DELETIONS AND ADDITIONS

When reviewing cases submitted to the MCR, delete any duplicate records, such as a case that is found to have been abstracted more than once. Also, delete a previously reported case if subsequent evidence disproves the presence of cancer, or if what was thought to be a new primary cancer is later found to be a manifestation of an earlier primary cancer. All deletions must be reported to the MCR. If your facility reports a class 43 and then the patient comes to

your facility for treatment resulting in a class 20, 21, or 22, please resubmit that case to the MCR.

Example:

After a case of “probable lymphoma” had been reported, the patient was referred to a specialty center where additional workup and repeat biopsies were performed. The final diagnosis was changed to “atypical lymphocytic infiltrates,” and physicians decided to follow the patient closely but not treat the condition. Since the patient is now deemed not to have cancer, delete the case from the hospital’s registry and notify the MCR.

Changes to the Abstract

The information in an abstract should be changed for the following circumstances and should be reported to the MCR.

1. To correct abstracting errors
2. When clarifications or rule changes retroactively affect data item codes.
3. When better information is available later.

Example 1: Consults from specialty labs, pathology report addendums or comments or other information have been added to the chart after the registrar abstracted the case. Whenever these later reports give better information about the histology, grade of tumor, primary site, etc., change the codes to reflect the better information. Make sure these changes are reported to the MCR.

Example 2: The primary site was recorded as unknown at the time of diagnosis. At a later date, the physician determines that the cancer is primary to the testis. Change the primary site from unknown to testis. Update all other fields affected by the change in primary site. Make sure these changes are reported to the MCR.

Example 3: The original diagnosis was in situ. Metastases are diagnosed at a later date. Change the behavior code for the original diagnosis from in situ to invasive when no new primary has been diagnosed in the interim. Make sure these changes are reported to the MCR.

Example 4: Patient seen in Hospital A. The pathologic diagnosis was negative for malignancy. Patient goes to Hospital B and the slides from Hospital A are re-read. The diagnosis at Hospital B is reportable. Hospital B sends their slide report back to Hospital A. Hospital A reports the case based on the info from Hospital B. Make sure to enter supporting documentation in the text.

4. When the date of diagnosis is confirmed in retrospect to be earlier than the original date abstracted.

Example: Patient has surgery for a benign argentaffin carcinoid (8240/1) of the sigmoid colon in May 2009. In January 2010, the patient is admitted with widespread metastasis consistent with malignant argentaffin carcinoid. The registrar accessions the malignant argentaffin carcinoid as a 2010 diagnosis. Two months later, the pathologist reviews the slides from the May 2009 surgery and concludes that the carcinoid diagnosed in 2009 was malignant. Change the date of diagnosis to May 2009 and histology to 8241 and the behavior code to malignant (/3). Make sure these changes are reported to the MCR.

ADDITIONAL TREATMENT INFORMATION

If after a case is reported to the MCR, additional information on treatment is added to an abstract, the additional treatment should be reported to the MCR by resubmitting the abstract or by contacting us with the updated information. For cancer registries using their own abstracting software, please report these updated abstracts as modification records.

CODING RULES AND EXAMPLES

CLASS OF CASE

All cases are assigned a class of case based on the nature of involvement of the facility in the care of the patient.

Analytic Cases

Cases diagnosed at the reporting facility and/or administered any of the first course of treatment there are analytic. A network clinic or outpatient center belonging to the facility is considered part of the facility. These cases would be given a class of case 00 through 22.

Nonanalytic Cases

Nonanalytic cases include class of case 30 through 99. The reporting facility did not diagnosis and/or provide the first course of treatment with the exception of class 38 and class 43. The reporting facility may have performed the autopsy and diagnosed the cancer (Class 38). For a class 43, a specimen was read at the reporting facility. However, the patient never physically entered the facility.

The MCR requires that classes of case 00-22, 30, 31, 32, 34, 36, 38, 43 be reported by all facilities. Class of case 34 and 36 would be used for cases of VIN III, VAIN III and AIN III which are not reportable to the COC but must be reported to the MCR. Cancer registries must also report class of case 40-42 if they collect these cases. New cancer programs should submit class of case 35 and 37 for cases diagnosed January 1, 1996 or later. **DO NOT** use class of case 99. If you are not sure of what class of case to use, please contact the MCR for assistance.

MORPHOLOGY (HISTOLOGY, BEHAVIOR, AND GRADE):

The instructions for coding histology, behavior, and grade are found in the Morphology section of the ICD-O-3 “Coding Guidelines for Topography and Morphology” (ICD-O-3 pp. 27-34). Instructions for coding histology and behavior can be found under each data item in the “Data Item” section of this manual. The grade data is described below due to changes in the number of data items over time to collect grade.

Grade or Differentiation

This code occupies the 6th position of the morphology code. This number describes the grade or differentiation characteristics of the cancer. In most cases, the pathology report is the source for this description.

Grade is a measure of the aggressiveness of the tumor and an important prognostic indicator for many tumors. Prior to 2018, grade in cancer registries has been collected based on a generic classification with the following categories.

GRADE, DIFFERENTIATION OR CELL INDICATOR

Item Length: 1

NAACCR Item #: 440

NAACCR Name: Grade

Grade, Differentiation for solid tumors (Codes 1, 2, 3, 4, 9) and Cell Indicator for Lymphoid Neoplasms (Codes 5, 6, 7, 8, 9)

Code	Grade Description
1	Well differentiated
2	Moderately differentiated
3	Poorly differentiated
4	Undifferentiated or anaplastic
5	T-cell; T-precursor cell
6	B-cell; B-precursor cell
7	Null cell; Non-T-non-B
8	NK cell (natural killer cell)
9	Grade unknown, not stated, or not applicable

For cases diagnosed prior to January 1, 2014, use the grade codes in the FORDS 2012 manual. For cases diagnosed between January 1, 2014 and December 31, 2017, use the grade codes found at <https://seer.cancer.gov/tools/grade/>.

For solid tumors, diagnosed January 1, 2018 and forward, grade will be collected in three different data items, Grade Clinical, Grade Pathological, and Grade Post Therapy, and the coding instructions will depend on the type of cancer. The grade manual can be accessed at <https://www.naaccr.org/SSDI/Grade-Manual.pdf?v=1556221646>. Also, the NAACCR Site

Specific Data Items (SSDI)/Grade site version 2.0 (<https://apps.naaccr.org/ssdi/list/2.0>) can be used to code the grade data items.

Beginning in January 1, 2021, a new grade item has been added. Grade Post Therapy Clinical (yc). Grade Post Therapy has been renamed as Grade post Therapy Path (yp).

Prior to 2018, codes 6-8 for grade collected the cell lineage indicator for Hematopoietic and Lymphoid Neoplasms. For cases diagnosed prior to January 1, 2010, the cell lineage had be stated to be coded 6-8. If the lineage was not stated, then cases were coded to “9”. For cases diagnosed January 1, 2010 to December 31, 2017, the grade codes were included in the *Hematopoietic and Lymphoid Neoplasm Manual and Database*. Cell lineage indicator will no longer be collected for cases diagnosed January 1, 2018 and later.

MULTIPLE PRIMARIES:

For solid tumors diagnosed prior to January 1, 2007:

- For nonmalignant tumors of the central nervous system, use the instructions below under the heading **Determining Multiple Primaries for Nonmalignant CNS Tumors** to decide whether the tumor(s) is one site or multiple sites.
- Use the instructions below under the heading **Site Differences** to decide whether the tumor(s) is one site or multiple sites.
- Follow the instructions below under the heading **Histology Differences** to decide whether tumors other than lymphomas, leukemia, or benign or borderline CNS tumors represent a single histology or mixed/multiple histologies.
- Follow the instructions below under the heading **Timing** to decide if one or more primaries are involved.

For solid tumors diagnosed January 1, 2007 to December 31, 2017:

The SEER 2007 *Multiple Primary and Histology Coding Rules* contain site-specific rules for lung, breast, colon, melanoma of the skin, head and neck, kidney, renal pelvis/ureter/bladder, and brain. A separate set of rules addresses the specific and general rules for all other sites. The multiple primary rules guide and standardize the process of determining the number of primaries.

Apply the 2007 multiple primary rules to the tumor(s) diagnosed January 1, 2007 to December 31, 2017, when the patient had a previous tumor(s) diagnosed prior to 2007.

Example 1: Duct carcinoma of the right breast diagnosed in July 2006. In February 2007, duct carcinoma of the right breast is diagnosed in a separate tumor. Apply the 2007 rules to the tumor diagnosed in 2007. According to the 2007 rules, the 2007 tumor is not a new primary.

Example 2: Duct carcinoma of the right breast diagnosed in July 2006. In February 2007, duct carcinoma of the left breast is diagnosed. Apply the 2007 rules to the 2007 diagnosis. According to the 2007 rules, the 2007 diagnosis is a new primary.

For solid tumors diagnosed January 1, 2018 and later:

The SEER 2018 *Solid Tumor Rules* contain site specific rules for head and neck, colon, rectosigmoid, rectum, lung, cutaneous melanoma, breast, kidney, urinary sites, non-malignant and malignant CNS, and peripheral nerves. . A separate set of rules addresses the specific and general rules for all other sites.

Apply the 2018 *Solid Tumor Rules* to the tumor(s) diagnosed January 1, 2018 and later, when the patient had a previous tumor(s) diagnosed prior to 2018.

Example 1: Duct carcinoma of the right breast diagnosed in July 2016. In February 2018, duct carcinoma of the right breast is diagnosed in a separate tumor. Apply the 2018 rules to the tumor diagnosed in 2018. According to the 2018 rules, the 2018 tumor is not a new primary.

The SEER *Multiple Primary and Histology Coding Rules* do not apply to hematopoietic primaries (lymphoma and leukemia M9590-9989), or Kaposi Sarcoma (M9140) of any site. For hematopoietic and lymphoid neoplasms diagnosed prior to January 1, 2010, use the tables in Appendix D to decide whether differing histologies represent one or more primaries. Primary site and timing are not applicable for determining whether these malignancies represent one or more primaries. For hematopoietic and lymphoid neoplasms diagnosed January 1, 2010 and later, use the *Hematopoietic and Lymphoid Neoplasm Coding Manual and Database* (DB) to determine the correct number of primaries to report. Consider Kaposi sarcoma as one primary site.

Determining Multiple Primaries for Nonmalignant CNS Tumors (For cases diagnosed prior to January 1, 2007).

Source: *Data Collection of Primary Central Nervous System Tumors: National Program of Cancer Registries Training Materials*

Determining if Site is Same or Different

Each **subsite** (fourth-digit level) as delineated in *ICD-O-3* is considered a separate site.

- If separate tumors with the same histology occur **in the same subsite**, they are considered the same tumor and one abstract is completed. Therefore, if multiple tumors of the same histology occur in the cerebrum (C71.0), they are considered the same tumor regardless of when they occur and only one abstract is completed.
- If separate tumors with the same histology occur **in different subsites**, they are different tumors and separate abstracts are completed. Therefore, if a tumor occurs

in the cerebral meninges (C70.0), and a separate tumor occurs in the spinal meninges (C70.1), they are considered separate tumors. Likewise, if separate brain tumors of the same histology occur in the frontal lobe (C71.1) and in the occipital lobe (C71.4), they are also considered separate tumors.

- As with malignant tumors, if the first three digits are the same, and the fourth digit is a **9** or not otherwise specified (NOS) site, this is considered one site and should be coded to the **more specific site**. For example, if a tumor is identified as meninges, NOS (C70.9), and a separate tumor is identified as occurring in either the spinal (C70.1) or cerebral (C70.0) meninges, this is considered one tumor, and only one abstract should be completed using the more specific site code.
- Laterality is used to determine multiple primaries for nonmalignant CNS tumors for sites listed as being lateral. If multiple tumors of the same site and same histologic type are identified and **both sides** of a site (listed as lateral) are involved, the tumors should be considered to be separate tumors, and separate abstracts should be completed. For example, the right and left temporal lobes of the brain or the right and left acoustic nerves.

How to determine same vs. different histologies for benign and borderline primary intracranial and CNS tumors (C70.0-C72.9, C75.1-C75.3) (Based on histologic groupings)

Note: These rules do not apply to malignant primary intracranial and CNS tumors.

To determine if the histology in multiple nonmalignant CNS tumors is the same, follow these rules in priority order.

When multiple tumors are in the **same site**; the first three digits of the histology code are **the same**; and the codes are not found in the table below, then the histology is considered to be **the same**. Only one abstract should be completed

When multiple tumors are in the **same site**; the first three digits of the histology code are **different**, and the codes are not found in the table below, then the histology is considered to be **different**. Separate abstracts should be completed.

If all histologies are in the **same histologic group** in the table below, then the histology is considered to be the **same**, even though the first three digits are different. These two histologies represent a progression, differentiation, or subtype of a single histologic category. One abstract should be prepared.

Example: A patient has a diagnosis of a choroid plexus papilloma, NOS (9390/0), a nonspecific term, and subsequently has a diagnosis of atypical choroid plexus papilloma (9390/1), a more specific histology, in the same site. These terms are both in the grouping of choroids plexus neoplasms in the table below. The 9390/0 is the

first diagnosis (earliest diagnosis date), but 9390/1 is more specific. Therefore, 9390/1 should be used.

Example: A patient has subependymoma (9383/1), a specific histology, and is subsequently diagnosed with a choroid glioma (9444/1), also a specific histology. Both histologies are listed in the table below under the ependymomas grouping. In this instance, because both histologies are specific and in the same grouping, the first histology of subependymoma (9383/1) should be coded even though the second histology has a higher code.

If the first three digits are the **same** as the first three digits for any histologies in **one** of the groupings in the table below, then the histology is considered to be **the same**.

Example: A patient has a ganglioglioma (9505/1) listed in the table below in the grouping neuronal and neuronal-gliial neoplasm, as well as a separate Pacinian tumor (9507/0) which is not listed in the table below. These two tumors have the same first three digits. In this instance, the Pacinian tumor is considered the same as the ganglioglioma, and only one abstract should be completed. The first histology of ganglioglioma (9505/1) should be coded even though the second histology has a higher code.

If the first three digits are the **same** and the histologies are from two **different** groups in the histologic groupings table, the histologies are considered to be **different**.

Example: A patient has a choroid plexus papilloma (9390/0), listed in the table below in the choroid plexus neoplasm grouping, as well as a myxopapillary ependymoma (9394), which is listed in the ependymoma grouping. In this case, even though the first three digits are the same, the histologies are considered to be different for these tumors because they are listed in different groupings in the table below. Thus, two abstracts should be completed.

Tumor	ICD-O-3 Code Groupings
Choroid plexus neoplasms	9390/0, 9390/1
Ependymomas	9383, 9394, 9444
Neuronal and neuronal-gliial neoplasms	9384, 9412, 9413, 9442, 9505/1, 9506
Neurofibromas	9540/0, 9540/1, 9541, 9550, 9560/0
Neurinomatosis	9560/1
Neurothekeoma	9562
Neuroma	9570
Perineurioma, not otherwise specified	9571/0

Malignant Transformation

In rare cases, a diagnosed nonmalignant tumor transforms into a malignant tumor. In these cases, the behavior changes from code 0 or 1 to code 2 or 3.

When malignant transformation occurs in a **previously diagnosed nonmalignant tumor**, the tumors are considered **separate primaries**, and two abstracts should be completed because of the change from nonmalignant to malignant.

SITE DIFFERENCES (Cases diagnosed before January 1, 2007)

- The **third numeric digit** after the ‘C’ describes a subsite of the organ; it is **not used** to define individual (different) sites.

Exception: For the following sites, a difference in the third numeric digit designates a different primary site, **except** NOS (C_ _ .9) with a specific four-digit site code in the same site.

- Colon (C18.0-C18.9) except polyps involving multiple segments (see Colon and Rectum Polyps).
- Anus/anal canal (C21.0-C21.8)
- Pleura (visceral, parietal, NOS) (C38.4)
- Bone (C40.0-C41.9)
- Melanoma of the skin (C44.0-44.9)
- Peripheral nerves/autonomic nervous system (C47.0-C47.9)
- Connective tissue (C49.0-C49.9)
- Non-malignant meninges (C70.0–C70.9 with Behavior Code /0 or /1)
- Non-malignant brain (C71.0–C71.8 with Behavior Code /0 or /1)
- Non-malignant spinal cord, cranial nerves, and other parts of central nervous system (C72.0–C72.8 with Behavior Code /0 or /1)

Colon and Rectum Polyps

- Simultaneous lesions and polyps in the same segment of the colon are a single primary.
- Polyps may be present in more than one segment of the colon. If the diagnosis reads adenocarcinoma in multiple polyps, it is one primary, colon, NOS (C18.9).

Familial Polyposis

- This is a genetic disease characterized by polyps that increase in numbers and may cover the mucosal surface of the colon.
- If multiple segments of the colon, rectosigmoid and/or rectum are involved with adenocarcinoma in adenomatous polyposis coli or adenocarcinoma in multiple adenomatous polyps, it is a single primary. Code the primary site to colon, NOS (C18.9).

- If the **first two numeric digits** after the C are **identical**, it is the **same site**.

Possible Exception: Paired Organs

- It is one primary if a physician states the tumor in one organ is metastatic from the other. Code the laterality to the side in which the primary originated. If the side of origin is unknown, code '4' for laterality.
 - Code as separate primaries if the physician states these are independent primaries or there is no physician statement that one is metastatic from the other with the exception of the following:
 - Simultaneous bilateral involvement of the ovaries with the same histology is one primary. Laterality is coded '4' when the ovary of origin is unknown.
 - Bilateral retinoblastomas are a single primary with laterality of '4.'
 - Bilateral Wilms tumors are always a single primary with laterality of '4.'
 - Laterality should not be used to determine single or multiple primaries of malignant brain and central nervous system tumors. Laterality is used in the determination of single or multiple primaries for benign and borderline brain and central nervous system tumors.
- If there is any difference in the first two numeric digits after the C, it is a **different** site.

Exceptions: The following groups of three-character ICD-O-3 topography codes refer to single organs. Lesions within any combination of each group are considered to be the same primary site.

- C01 Base of tongue; C02 Other and unspecified parts of tongue
- C05 Palate; C06 Other and unspecified parts of mouth
- C07 Parotid gland; C08 Other and unspecified major salivary glands
- C09 Tonsil; C10 Oropharynx
- C12 Pyriform sinus; C13 Hypopharynx
- C23 Gallbladder; C24 Other and unspecified parts of biliary tract
- C30 Nasal cavity and middle ear; C31 Accessory sinuses
- C33 Trachea; C34 Bronchus and lung
- C37 Thymus; C38.0 Heart; C38.1-C38.3 Mediastinum; C38.8 Overlapping lesion of heart, mediastinum, and pleura.
- C51 Vulva; C52 Vagina; C57.7 Other specified female genital organs; C57.8-C57.9 Unspecified female genital organs
- C56 Ovary; C57.0 Fallopian tube
- C57.1 Broad ligament; C57.2 Round ligament
- C57.3 Parametrium; C57.4 Uterine adnexa
- C60 Penis; C63 Other and unspecified male genital organs
- C64 Kidney; C65 Renal pelvis; C66 Ureter; C68 Other and unspecified urinary organs
- C74 Adrenal gland; C75 Other endocrine glands and related structures

HISTOLOGY DIFFERENCES (Cases diagnosed before January 1, 2007)

Source: SEER Program and Coding and Staging Manual

The ICD-O-3 morphology code has five digits, for example 8500/3. The **fifth digit** of the ICD-O-3 morphology code is the behavior code. The first four characters are sometimes referred to as the histology code. Multiple terms may describe a single histology. Refer to the ICD-O-3 histology code to determine whether two or more lesions represent the same tumor histologically.

- If the first **three digits of the ICD-O-3 histology codes are the same**, it is the same histology.
- Lesion(s) with a single histology (the first three digits of the histology code are the same) containing invasive and in situ components are one primary. Code the behavior of the invasive component.
- A single lesion composed on one histologic type is a single primary, even if the lesion crosses site boundaries.
- A single lesion composed of multiple (different) histologic types is a single primary even if the lesion crosses site boundaries.
- If one lesion is invasive and another lesion of the same histologic type is in situ, or if two or more lesions have invasive and in situ components, this is a single primary.
- A difference in the first three digits of the ICD-O-3 histology code indicates a **different** histologic type.

Exception 1:

If one malignancy is stated to be carcinoma, NOS, adenocarcinoma, NOS, sarcoma, NOS, or melanoma, NOS and the second lesion is a more specific term, such as large cell carcinoma, mucinous adenocarcinoma, spindle cell sarcoma, or superficial spreading melanoma, consider this to be a **single** histology.

Exception 2:

For lymphatic and hematopoietic disease, refer to Appendix D to determine which histologies represent single or multiple primaries.

Exception 3:

Consider the following as a **single** histology, even though the first three digits of the ICD-O-3 morphology codes differ. Code the histology according to the rules for mixed histologies.

- Transitional cell carcinoma (8120-8131) of the bladder (C67._)

- Ductal (8500) and lobular (8520) adenocarcinoma of the breast (C50. _)

Exception 4:

See page 20 for determining single or multiple primaries for benign or borderline primary intracranial or CNS tumors.

TIMING RULES (Cases diagnosed prior to January 1, 2007)

- If two malignancies of the same histology occur in the same site simultaneously (within two months of each other), there is only **one** primary.

Exception 1:

Each occurrence of melanoma of the skin is a new or **separate** primary **unless** a physician states otherwise.

Exception 2:

Two primary intracranial and central nervous system tumors (C70.0–C72.9, C75.1–C75.3) in which one is malignant (behavior of /2 or /3) and one is non-malignant (behavior of /0 or /1) are always separate primaries regardless of timing. **Complete two abstracts.**

- If a tumor with the same histology is identified in the same site at least two months after the initial/original diagnosis, this is a separate primary.

Exception 1:

When there is an in situ followed by an invasive cancer at the same site more than 2 months apart, report the invasive cancer as a second primary even if stated by the physician to be recurrence. This is true for all sites including bladder.

- Multiple lesions with different histologies in a single site are **separate** primaries, whether they occur simultaneously or at different times.
- If two malignancies of the same histology (following the rules under *Histology Differences*) and in the same site (following the rules under *Site Differences*, including rules for laterality for paired sites) are identified **more** than two months apart, then there are **two** primaries. Complete a separate report for each one.

Exceptions:

The following are recurrences of the original disease without time limits.

- Invasive bladder primaries with morphology codes 8120-8130 (If there is an in situ followed by an invasive more than two months apart, report the invasive cancer as a second primary even if stated by the physician to be a recurrence)
- Invasive adenocarcinoma of the prostate, site code C61.9.
- Kaposi sarcoma (9140) of any site.

- Lymphoma and leukemia histologies that are determined from Appendix D to be the same primary.

EXAMPLES OF SINGLE OR MULTIPLE PRIMARY CODING (Cases diagnosed prior to January 1, 2007)

- A single lesion involving the tongue and floor of mouth is one primary.
- A single, large mucinous adenocarcinoma involving the sigmoid and descending colon segments is one primary.
- A single lesion containing both embryonal cell carcinoma and teratoma is a single primary and would be coded to 9081/3, mixed embryonal carcinoma and teratoma.
- A single lesion of the liver composed of neuroendocrine carcinoma (8246/3) and hepatocellular carcinoma (8170/3) is a single primary and would be coded to the more specific histology, neuroendocrine carcinoma 8246/3.
- At mastectomy for removal of a 2 cm invasive ductal carcinoma, an additional 5 cm area of intraductal carcinoma was noted. Abstract as one invasive primary.
- Adenocarcinoma in adenomatous polyp (8210) in sigmoid colon removed by polypectomy in December 2004. At segmental resection in January 2005, and adenocarcinoma in a tubular adenoma (8210) adjacent to the previous polypectomy site was removed. Count as one primary.
- Infiltrating duct carcinoma of the upper outer quadrant of the right breast diagnosed March 2004 and treated with lumpectomy. Previously unidentified mass in left inner quadrant right breast noted in July 2004 mammogram. This was removed and found to be infiltrating duct carcinoma. Abstract the case as two primaries.

DETERMINING MULTIPLE PRIMARIES AND HISTOLOGIES (Solid tumors diagnosed on or after January 1, 2007 and Hematopoietic and Lymphoid Neoplasms diagnosed January 1, 2001 and forward)

For solid tumors diagnosed January 1, 2007 to December 31, 2017, refer to the *Multiple Primary and Histology Coding Rules* manual for determining the number of primaries and the correct histology for each primary abstracted.

For cases diagnosed January 1, 2018 and later, refer to the *Solid Tumor Rules* manual for determining the number of primaries and the correct histology for each primary abstracted.

For hematopoietic and lymphoid neoplasms diagnosed January 1, 2001 and later, use the

Hematopoietic and Lymphoid Neoplasm Coding Manual and Database (DB) to determine the correct number of primaries, primary site, histology and grade to report.

STAGING

Collaborative Staging (CS)

(<https://cancerstaging.org/cstage/schema/Pages/version0205.aspx>) is to be used for cases diagnosed between January 1, 2004 and December 31, 2015. Site Specific Factors, Regional Nodes Positive and Examined, and Lymph-vascular Invasion from CS will continue to be used with cases diagnosed January 1, 2016 to December 31, 2017. SEER Summary Stage 2000 is used for cases diagnosed from January 1, 2001 to December 31, 2003 and cases diagnosed January 1, 2015 to December 31, 2017. AJCC TNM staging will be used with cases diagnosed January 1, 2016 to December 31, 2017. Summary Stage 2018 and the new Site Specific Data Items (SSDI) will be used for cases diagnosed January 1, 2018 and later.

Timing Rule for CS (Cases diagnosed between January 1, 2004 and December 31, 2015)

The timing rule for CS coding was designed to make use of the most complete information possible to yield the “best stage” information for the tumor at the time of diagnosis—“use all information gathered through completion of surgery(ies) in first course of treatment or all information available within four months of the date of diagnosis in the absence of disease progression, whichever is *longer*.” Disease progression is defined as further direct extension, regional lymph node involvement or distant metastasis known to have developed after the diagnosis was established. Information about tumor extension, lymph node involvement, or distant metastasis obtained after disease progression is documented should be excluded from the CS coding.

Ambiguous Terminology for CS (Do not use this list with AJCC TNM Staging)

If the wording in the patient record is ambiguous with respect to tumor spread, use the following guidelines.

Consider as Involvement/Extension		DO NOT Consider as Involvement
Adherent	Induration	Abuts
Apparent(ly)	Infringe/infringing	Approaching
Appears to	Into*	Approximates
Comparable with	Intrude	Attached
Compatible with	Invasion to, into, onto, out onto	Cannot be excluded/ruled out
Consistent with	Most likely	Efface/Effacing/Effacement
Contiguous/continuous with	Onto*	Encased/Encasing
Encroaching upon*	Overstep	Encompass(ed)
Extension to, into, onto, out onto	Presumed	Entrapped
Features of	Probable	Equivocal
Fixation to another structure**	Protruding into (unless encapsulated)	Extension to without invasion/involvement of
Fixed**	Suspected	Kiss/kissing
Impending perforation of	Suspicious	Matted (except for lymph nodes)
Impinging upon	To*	Possible
Impose/imposing on	Up to	Questionable
Incipient invasion		Reaching
		Rule out
		Suggests
		Very close to
		Worrisome

Coding CS Values

The complete instructions and site-histology defined codes are available in the *Collaborative Staging Manual and Coding Instructions (CS Manual)*. Part I, Section 1 provides general instructions and the instructions and codes for generic (non site-specific) items. Part I, Section 2 provides information on lab tests, tumor markers, and site specific factors. Part II contains the site-specific instructions and codes.

The MCR collects the following CS Items: CS Tumor Size, CS Extension, CS Tumor Size/Extension Eval, CS Lymph Nodes, CS Lymph Node Eval, Regional Nodes Examined, Regional Nodes Positive, CS Mets at Diagnosis, CS Mets Eval, Site Specific Factor 1 (Lung, Pleura, Retinoblastoma Brain and Central Nervous System and Breast cases only), Site Specific Factor 2 (Breast, corpus adenosarcoma, corpus carcinoma and corpus sarcoma cases only), Site Specific Factor 3 (Prostate Cases Only), Site Specific Factors 8-16 (Breast cases only), Site Specific 25 (Schema discriminator for various sites).

- Read the medical record carefully to identify the primary site and histology and

determine their ICD-O-3 codes. While you are reviewing the record, make mental notes about the tissues and lymph nodes that are involved by tumor.

- If the histology is melanoma (8720-8790), Kaposi sarcoma (9140), retinoblastoma (9510-9514), lymphoma (9590-9699 and 9702-9729), mycosis fungoides (9700-9701), or hematopoietic and reticuloendothelial system (9731-9989), use the histology-specific schema for the appropriate histology-site combination.
- Otherwise, use the correct site-specific schema in Part II of the *CS Manual*.
- Code your cases.

SEER Summary Stage 2000 (Cases diagnosed January 1, 2001 to December 31, 2003 or diagnosed January 1, 2015 to December 31, 2017)

Summary staging is a basic way to categorize spread of disease from the point of origin. Summary Stage 2000 applies to every anatomic site and to lymphomas and leukemias. Summary stage uses both the clinical and pathologic information in determining the extent of disease. Gross observations at surgery are important when all malignant tissue is not removed. Be sure to also review all clinical information to assure accurate staging. However, when the pathologic information disproves either the clinical information or the surgical observations, pathologic information takes precedent. The *SEER Summary Staging Manual - 2000* manual can be found at <https://seer.cancer.gov/tools/ssm/ssm2000/>. Summary stage should include all information available through the completion of surgery(ies) in the first course of treatment or within four months of diagnosis in the absence of disease progression, whichever is longer. Summary stage information obtained after radiation or systemic therapy has begun may be included unless it is beyond the timeframe stated in the previous sentence. Autopsy reports are used in summary stage just like pathology reports. If the only information available for stage is the values for T, N, and M from AJCC staging, record the summary stage that corresponds to that information. If there is a discrepancy between the documentation available and the recorded TNM stage, the documentation should be used to record the summary stage.

All schemas apply to all histologies, unless otherwise noted. Some exceptions to this rule include Kaposi's sarcoma and lymphomas which should be staged based on histology schemes and not the primary site scheme. Site specific guidelines take precedence over the general guidelines. Unlike with CS where site specific guidelines are prior to the coding rules, in Summary Stage 2000 manual, site specific guidelines are documented after the coding rules in each chapter.

The following is a list of ambiguous terms to determine whether or not tissue should be considered involved for summary stage.

Consider as Involvement/Extension		DO NOT Consider as Involvement
Adherent	Induration	Abuts
Apparent(ly)	Infringe/infringing	Approaching
Appears to	Into*	Approximates
Comparable with	Intrude	Attached
Compatible with	Invasion to, into, onto, out onto	Cannot be excluded/ruled out
Consistent with	Matted (for lymph nodes only)	Efface/Effacing/Effacement
Contiguous/continuous with	Most likely	Encased/Encasing
Encroaching upon*	Onto*	Encompass(ed)
Extension to, into, onto, out onto	Overstep	Entrapped
Features of	Presumed	Equivocal
Fixation to another structure**	Probable	Extension to without invasion/involvement of
Fixed**	Protruding into (unless encapsulated)	Kiss/kissing
Impending perforation of	Suspected	Matted (except for lymph nodes)
Impinging upon	Suspicious	Possible
Impose/imposing on	To*	Questionable
Incipient invasion	Up to	Reaching
		Rule out
		Suggests
* Interpreted as involvement whether the description is clinical or operative/pathological		Very close to
** Interpreted as involvement of other organ or tissue		Worrisome

AJCC TNM Staging (Cases diagnosed January 1, 2016 to December 31, 2017)

AJCC TNM Stage determines the anatomic extent of disease based on clinical, operative and pathologic findings. AJCC TNM Stage is used to make treatment decisions, assess prognosis, and measure outcomes. The rules in the current *AJCC Cancer Staging Manual* should be used to assign the clinical and pathologic T, N, M, and Stage Group. The following are general rules that apply to all sites.

- Clinical stage included any information obtained regarding the extent of disease before the initiation of definitive treatment (surgery, systemic or radiation therapy, active surveillance, or palliative care) or within four months after the date of diagnosis, whichever is shorter, as long as the cancer has not clearly progressed during that time.
- Pathologic stage includes any information obtained regarding the extent of disease through completion of definitive surgery as part of the first course of treatment or identified within four months after the date of diagnosis, whichever is longer, as long as there is no systemic or radiation therapy administered or the cancer has not clearly progressed during that time.
- When a patient has multiple primaries, stage each primary independently.

- If the stage group cannot be determined from the recorded components, then record it as unknown.
- When a patient with multiple simultaneous primaries has metastases, a biopsy may be able to distinguish the source of the distant disease. If the physician cannot determine which primary has metastasized, stage all primaries as having metastatic disease. If, at a later time, the physician identifies which primary has metastasized, update the stage(s) as appropriate.
- If a site/histology combination is not defined in the AJCC Manual, code 88 for clinical and pathologic T, N, and M as well as stage group.
- Even if complete AJCC TNM information is not available in the record, any piece of staging information should be collected and reported.
 - Example: If the T and N are available but not information is available on M, the T and N should be reported.

The AJCC items required to be completed are the following:

Clinical T
 Clinical N
 Clinical M
 Clinical Stage Group
 Clinical Stage (Prefix/Suffix) Descriptor
 Staged By (Clinical Stage)
 Pathologic T
 Pathologic N
 Pathologic M
 Pathologic Stage Group
 Pathologic Stage (Prefix/Suffix) Descriptor
 Staged By (Pathologic Stage)

SEER Summary Stage 2018 (Cases diagnosed January 1, 2018 and later)

Summary staging is a basic way to categorize spread of disease from the point of origin. Summary Stage 2018 applies to every anatomic site and to lymphomas and leukemias. Chapter-specific guidelines take precedence over general guidelines. Always read the information pertaining to a specific primary site or histology chapter. Summary stage uses both the clinical and pathologic information in determining the extent of disease. Gross observations at surgery are important when all malignant tissue is not removed. Be sure to also review all clinical information to assure accurate staging. However, when the pathologic information disproves either the clinical information or the surgical observations, pathologic information takes precedent. The *SEER Summary Staging Manual - 2018* manual can be found at <https://seer.cancer.gov/tools/ssm/>. Summary stage should include all information available through the completion of surgery(ies) in the first course of treatment or within four months of diagnosis in the absence of disease progression, whichever is longer Information for Summary Stage from a surgical resection after neoadjuvant treatment may be used, but ONLY if the extent of disease is greater than the

pre-treatment clinical findings. Autopsy reports are used in summary stage just like pathology reports. If the only information available for stage is the values for T, N, and M from AJCC staging, record the summary stage that corresponds to that information. If there is a discrepancy between the documentation available and the recorded TNM stage, the documentation should be used to record the summary stage.

All schemas apply to all histologies, unless otherwise noted. Some exceptions to this rule include Kaposi's sarcoma and lymphomas which should be staged based on histology schemes and not the primary site scheme. Site specific guidelines take precedence over the general guidelines.

The following is a list of ambiguous terms to determine whether or not tissue should be considered involved for summary stage.

Consider as Involvement/Extension		DO NOT Consider as Involvement
Adherent	Induration	Abuts
Apparent(ly)	Infringe/infringing	Approaching
Appears to	Into*	Approximates
Comparable with	Intrude	Attached
Compatible with	Invasion to, into, onto, out onto	Cannot be excluded/ruled out
Consistent with	Most likely	Efface/Effacing/Effacement
Contiguous/continuous with	Onto*	Encased/Encasing
Encroaching upon*	Overstep	Encompass(ed)
Extension to, into, onto, out onto	Presumed	Entrapped
Features of	Probable	Equivocal
Fixation to a structure other than primary**	Protruding into (unless encapsulated)	Extension to without invasion/involvement of
Fixed to another structure**	Suspected	Kiss/kissing
Impending perforation of	Suspicious	Matted (except for lymph nodes)
Impinging upon	To*	Possible
Impose/imposing on	Up to	Questionable
Incipient invasion		Reaching
		Rule out
		Suggests
		Very close to
		Worrisome
* Interpreted as involvement whether the description is clinical or operative/pathological		
** Interpreted as involvement of other organ or tissue		

Site Specific Data Items (Cases diagnosed January 1, 2018 and later)

In 2018, the Collaborative Staging Site Specific Factors (SSF) were discontinued and Site Specific Data Items (SSDI) were developed to replace them to capture prognostic information. SSDI's have unique names and are not limited to three characters like the SSF's were in Collaborative Staging. The SSDI manual can be accessed at

<https://apps.naaccr.org/ssdi/list/>. The MCR does not require all of the SSDI in the manual. The required SSDI are described in the data variable section of this manual.

First Course of Treatment

Treatment includes all types of therapy intended to modify, control, remove or destroy proliferating cancer cells. The first course of treatment includes all therapy planned and administered during the first diagnosis of cancer. “Active surveillance” is also a form of treatment administered to some patients and coded in *RX Summ-Treatment Status*. “No treatment” which is also coded in *RX Summ-Treatment Status* is an option that occurs if the patient or patient’s family refuses treatment, the physician recommends that no treatment be given, or the patient dies before starting treatment. For all malignancies except leukemias, any therapy administered after the discontinuation of first course of treatment is considered subsequent treatment. For leukemias, all therapy administered after the first relapse is secondary or subsequent treatment. Information on treatment can be found in the discharge plan, physician(s) treatment plan, or established protocol or accepted management guidelines in the absence of a treatment plan. If no plan, guideline or protocol is available and a physician cannot be consulted, use the following rule: “Initial treatment must begin within four months of the date of initial diagnosis.” If no treatment is given, record the date the decision was made not to treat or date of death. If the patient undergoes “Active surveillance,” record the date the decision for surveillance was made as the date of first course treatment.

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DATA ITEMS AND RECORD LAYOUT

The data items that are collected are those needed to produce age-, race-, sex-, and area-specific cancer incidence rates by cancer site and histologic type. Stage is included for cancer control studies. Administrative items are included that facilitate death clearance, patient matching, inter-field edit reviews, and hospital admission tracking. The fields are listed in the order that they will be entered into Web Plus for cases diagnosed January 1, 2021 and later. For abstracting cases diagnosed prior to January 1, 2021, make sure you use the Web Plus template that corresponds to the diagnosis year of the case.

The following data items are required:

REPORTING FACILITY

Item Length: 10
NAACCR Item #540
NAACCR Name: Reporting Facility
XML NAACCR ID: reportingFacility

Description

The Facility Identification Number is used to identify reporting facilities in the MCR database. This number is used to monitor data submissions and data accuracy.

Note: This number should be automatically coded by your software.

Examples:

Code	Reason
0006439999	6439999, General Hospital, Anywhere, MS
0010000099	10000099, Doctor's Medical Center, Anytown, MS

ABTRACTOR

Item Length: 3
NAACCR Item #570
NAACCR Name: Abstracted By
XML NAACCR ID: abstractedBy

Definition:

Records the initials or assigned code of the individual abstracting the case.

This item is used for quality control by the MCR.

Coding Instructions:

- Code the initials of the abstractor.

- This code will be automatically generated in Web Plus.

MEDICAL RECORD NUMBER

Item Length: 11

NAACCR Item #2300

NAACCR Name: Medical Record Number

XML NAACCR ID: medicalRecordNumber

Definition:

Records the medical record number used by the facility to identify the patient. Record the number assigned by the HIM Department, not a department specific number.

This number identifies the patient in a facility. It can be used by the MCR to reference a patient record and it helps to identify multiple reports on the same patient.

Coding Instructions:

- Record the medical record number.

Coding Examples:

Code (In addition to the medical record number)	Explanation
____NNNN	If the medical record number is fewer than 11 characters, use leading blanks and right justify the characters.
____SU	1-day surgery clinic without HIM number.
____RT	Radiation therapy department patient without HIM number
____UNK	The medical record number is unknown.

DATE OF 1ST CONTACT

Item Length: 8
 NAACCR Item #580
 NAACCR Name: Date of 1st Contact
 XML NAACCR ID: dateOf1stContact

Definition:

Date of first contact with the reporting facility as an inpatient or outpatient for diagnosis and/or treatment of this cancer.

Date of 1st Contact is one of several data items that can be used to measure timeliness of reporting by individual facilities to the MCR.

Coding Instructions:

- Date the patient first had contact with the facility as either an inpatient or outpatient for diagnosis and/or treatment of a reportable tumor.
- This may be the date of an outpatient visit for a biopsy, x-ray, or laboratory test, or the date a pathology specimen was collected at the hospital.
- For patients diagnosed in a staff physician office, the date of 1st contact should be the date the patient was seen at the reporting facility, not the date the patient was first seen in the staff physician office.
- When pathology-specimen-only tumors are collected (Class of Case 43), the date of specimen collection is the date of first contact.
- For autopsy only cases, the date of first contact is the same as the date of death.

Code	Definition
CCYYMMDD CCYYMM CCYY	The date the patient first had contact with the reporting facility for a diagnostic procedure; review or administration of treatment; palliative care; or, for pathology.
201104	Only description is “Spring” 2011
201107	Only description is “The middle of the year” 2011
201110	Only description is “Fall” 2011
CCYY1299 or CCYY0199	If the only description is “Winter,” try to determine if this is referring to the beginning or end of a year.
Blank	When it is unknown when the first patient contact occurred. This situation is rare. Set the appropriate date flag.

Coding Examples:

Code	Explanation
20110212	Patient has an outpatient mammography that is highly suspicious for malignancy on February 12, 2011, and subsequently undergoes an excisional biopsy on February 14, 2011. Record the date of the mammography (February 12, 2011) as the date of first contact/first admission to this facility.
20110414	Patient undergoes a biopsy in a physician's office on April 6, 2011. The pathology specimen was sent to the reporting facility and read as malignant melanoma. The patient enters that same reporting facility on April 14, 2011 for wide re-excision.
20111108	Patient has an MRI of the brain at your facility on November 8, 2011 for symptoms including severe headache and disorientation. The MRI findings are consistent with astrocytoma. Surgery on December 2 removes all gross tumor. The date of first contact is November 8, 2011.
20110604	Patient is admitted to your facility on June 1, 2011 for COPD. During a chest x-ray on June 4, 2011, lung cancer is discovered. The date of first contact is June 4, 2011, since the date of 1 st contact cannot be before the date of diagnosis.
201109	If the only information known is that the patient was admitted to the facility for cancer in September 2011 record the month and year.

DATE OF 1ST CONTACT FLAG

Item Length: 2

NAACCR Item #581

NAACCR Name: Date of 1st Contact Flag

XML NAACCR ID: dateOf1stContactFlag

Definition:

This flag explains why no appropriate value is in the field Date of 1st Contact. Prior to 2010, date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions:

- If a valid date is coded in Date of 1st Contact, then leave this field blank. A valid date may be just a partial date.
- If date of 1st contact is completely unknown, then leave date of 1st contact blank and record 12 in this field indicating that the date is unknown.

Code	Definition
12	A proper value is applicable but unknown (e.g., date of 1 st contact is unknown)
Blank	A valid date value is provided in item Date of 1 st Contact, or the date was not expected to have been transmitted.

Coding Examples:

Code	Explanation
Blank	Full date is known (CCYYMMDD) for Date of 1 st Contact
Blank	Partial date is known (CCYYMM or CCYY) for Date of 1 st Contact
12	Date is completely unknown for Date of 1 st Contact

CLASS OF CASE

Item Length: 2
NAACCR Item #610
NAACCR Name: Class of Case
XML NAACCR ID: classOfCase

Definition:

Classifies cases recorded in the database.

The MCR requires all analytic (class 00- 22) and non-analytic cases including class of case 30-32, 34, 36, 38, and 43 to be reported by all facilities. Class of case 34 and 36 would be used for cases of VIN III, VAIN III and AIN III which are not reportable to the COC but must be reported to the MCR. Cancer registries must also report class of case 40-42 if they collect these cases. New cancer programs should submit class of case 35 and 37 for cases diagnosed January 1, 1996 or later.

Do not use class of case 49 or 99. These are reserved for the central cancer registry only.

This data item divides case records into analytic and nonanalytic categories. This allows reporting facilities to select cases for use within their facility or to be reported to the MCR.

Coding Instructions:

- Analytic cases are coded 00–22.
- Nonanalytic cases are coded 30–99.
- Use the code that best describes the patient’s relationship to the reporting facility.
- Code 00 should be used only when it is known that the patient went somewhere else for treatment. If this information is not available, use code 10.
- Codes 34 and 36 should be used for VIN III, VAIN III and AIN III.
- A staff physician is a physician who is employed by the reporting facility, under contract with it, or a physician who has routine admitting privileges there.
- “In-transit” care is care given to a patient who is away from their primary source of care in order to prevent a break in their care. These cases would be coded as *Class of Case 31*. This would also include cases where the reporting facility is monitoring use of oral medications started somewhere else. If a patient begins infusion therapy or radiation therapy at another facility and then continues that care at the reporting facility, this is not a case of “in transit” care and should be coded as *Class of Case 21*.
-

Class of Case Definitions	
Case	Includes
Analytic Cases	
	Initial diagnosis at reporting facility or in a staff physician's office
Class 00	<ul style="list-style-type: none"> ▪ Initial diagnosis at the reporting facility AND all treatment or a decision not to treat was done elsewhere.
Class 10	<ul style="list-style-type: none"> ▪ Initial diagnosis at the reporting facility or in a physician's office with admitting privileges AND part or all of first course treatment or a decision not to treat was at the reporting facility, NOS.
Class 11	<ul style="list-style-type: none"> ▪ Initial diagnosis in a physician's office with admitting privileges AND part of first course treatment was done at the reporting facility.
Class 12	<ul style="list-style-type: none"> ▪ Initial diagnosis in physician's office with admitting privileges AND part of first course treatment was done at the reporting facility.
Class 13	<ul style="list-style-type: none"> ▪ Initial diagnosis at the reporting facility AND part of first course treatment was done at the reporting facility; part of first course treatment was done elsewhere.
Class 14	<ul style="list-style-type: none"> ▪ Initial diagnosis at the reporting facility AND all first course treatment or a decision not to treat was done at the reporting facility.
	Initial diagnosis elsewhere
Class 20	<ul style="list-style-type: none"> ▪ Initial diagnosis elsewhere AND all or part of first course treatment was done at the reporting facility, NOS.
Class 21	<ul style="list-style-type: none"> ▪ Initial diagnosis elsewhere AND part of first course treatment was done at the reporting facility; part of first course treatment was done elsewhere.
Class 22	<ul style="list-style-type: none"> ▪ Initial diagnosis elsewhere AND all first course treatment or a decision not to treat was done at the reporting facility.
Nonanalytic Cases	
	Patient appears in person at reporting facility
Class 30	<ul style="list-style-type: none"> ▪ Initial diagnosis and all first course treatment elsewhere AND reporting facility participated in diagnostic workup (for example, consult only, treatment plan only, staging workup after initial diagnosis elsewhere)
Class 31	<ul style="list-style-type: none"> ▪ Initial diagnosis and all first course treatment elsewhere AND reporting facility provided in-transit care; or hospital provided care that facilitated treatment elsewhere (for example, stent placement)
Class 32	<ul style="list-style-type: none"> ▪ Diagnosis AND all first course treatment provided elsewhere AND patient presents at reporting facility with disease recurrence or persistence (active disease)
Class 33	<ul style="list-style-type: none"> ▪ Diagnosis AND all first course treatment provided elsewhere AND patient presents at reporting facility with disease history only (disease not active)
Class 34	<ul style="list-style-type: none"> ▪ VIN III, VAIN III or AIN III AND initial diagnosis AND part or all of first course treatment by reporting facility.
Class 35	<ul style="list-style-type: none"> ▪ Case diagnosed before program's Reference Date AND initial diagnosis AND all or part of first course treatment by reporting facility.

Case	Includes
Nonanalytic Cases	
	Patient appears in person at reporting facility
Class 36	<ul style="list-style-type: none"> VIN III, VAIN III or AIN III AND initial diagnosis elsewhere AND all or part of first course treatment by reporting facility
Class 37	<ul style="list-style-type: none"> Cases diagnosed before program's Reference Data AND initial diagnosis AND all or part of first course treatment by reporting facility.
Class 38	<ul style="list-style-type: none"> Initial diagnosis established by autopsy at the reporting facility, cancer not suspected prior to death.
	Patient does not appear in person at reporting facility
Class 40	<ul style="list-style-type: none"> Diagnosis AND all first course treatment given at the same physician's office with admitting privileges.
Class 41	<ul style="list-style-type: none"> Diagnosis AND all first course treatment given in two or more different physician's offices with admitting privileges.
Class 42	<ul style="list-style-type: none"> Non-staff physician, clinic or other facility, not part of reporting facility, accessioned by reporting facility for diagnosis and/or treatment by that entity (for example, hospital abstracts cases from an independent radiation facility)
Class 43	<ul style="list-style-type: none"> Pathology or other lab specimen only
Class 49	<ul style="list-style-type: none"> Death certificate only. Used by central registry only.
Class 99	<ul style="list-style-type: none"> Nonanalytic case of unknown relationship to facility. Please contact the MCR before using this code.

Coding Examples:

Code	Explanation
00	Patient enters the reporting facility with dizziness and receives a clinical workup including CT and MRI of the brain. The results are positive for multiple metastatic deposits in both lobes of the brain. A CT of the lung is positive for hilar adenopathy. The patient is discharged to hospital B for treatment with a diagnosis of lung cancer.
14	Patient is admitted with hemoptysis. Workup reveals right upper lobe mass. A biopsy is positive for adenocarcinoma. The patient undergoes surgery followed by radiation therapy at the same facility.
21	Patient was diagnosed and had surgery at another facility for primary breast cancer. The patient then comes to your facility for radiation therapy.
32	Patient was diagnosed and treated for primary breast cancer four years prior to admission. Patient is admitted to your facility for recurrent breast cancer. She is treated with chemotherapy.
11	Patient diagnosed with breast cancer in Dr. Smith's office, who is a staff physician. The patient then comes to your hospital for surgery and radiation. She is then referred to an oncologist not associated with your facility and receives hormonal therapy.
43	Hospital pathology department received a tissue sample for evaluation which was positive for malignant melanoma. The patient never visited the hospital.

CASEFINDING SOURCE

Item Length: 2
 NAACCR Data Item #501
 NAACCR Name: Casefinding Source
 XML NAACCR ID: casefindingSource

Definition:

This variable codes the earliest source of identifying information. This data item will help reporting facilities, as well as, the MCR prioritize their casefinding activities. It will identify reportable tumors that were first identified through death clearance.

Coding Instructions:

Code the source that first identified the tumor.

Code	Definition
10	Reporting Hospital, NOS
20	Pathology Department Review (surgical pathology reports, autopsies, or cytology reports)
21	Daily Discharge Review (daily screening of charts of discharged patients in the medical records department)
22	Disease Index Review (review of disease index in the medical records department)
23	Radiation Therapy Department/Center
24	Laboratory Reports (other than pathology reports, code 20)
25	Outpatient Chemotherapy
26	Diagnostic Imaging/Radiology (other than radiation therapy, code 23; includes nuclear medicine)
27	Tumor Board
28	Hospital Rehabilitation Services or Clinic
29	Other Hospital Source (including clinic, NOS or outpatient department, NOS)
80	Death Certificate (case identified through death clearance)
99	Unknown

TYPE OF REPORTING SOURCE

Item Length: 1

NAACCR Data Item #500

NAACCR Name: Type of Reporting Source

XML NAACCR ID: typeOfReportingSource

Definition:

This variable codes the source documents used to abstract the majority of information on the tumor being reported. This may not be the source of the original case finding. For example, if a case is identified through a pathology laboratory report review and all source documents used to abstract the case are from a physician's office, code this item 4.

Coding Instructions:

- Code in the following priority order: 1, 2, 8, 4, 3, 5, 6, 7. This is a change to reflect the addition of codes 2 and 8 and to prioritize laboratory reports over nursing home reports. The source facilities included in the previous code 1 (hospital inpatient and outpatient) are split between codes 1, 2, and 8.
- This data item is intended to indicate the completeness of information available to the abstractor.
- Reports from health plans (e.g., Kaiser, Veterans Administration, military facilities) in which all diagnostic and treatment information is maintained centrally and is available to the abstractor are expected to be at least as complete as reports for hospital inpatients. This is the reason these sources are grouped with inpatients and given the code with the highest priority.
- Sources coded with '2' usually have complete information on the cancer diagnosis, staging, and treatment.
- Sources coded with '8' would include, but would not be limited to, outpatient surgery and nuclear medicine services. A physician's office that calls itself a surgery center should be coded as a physician's office. Surgery centers are equipped and staffed to perform surgical procedures under general anesthesia. If a physician's office calls itself a surgery center, but cannot perform surgical procedures under general anesthesia, code as a physician office.

Code	Definition
1	Hospital inpatient; Managed health plans with comprehensive, unified medical records.
2	Radiation Treatment Centers or Medical Oncology Centers (hospital-affiliated or independent).
3	Laboratory only (hospital-affiliated or independent).
4	Physician's office/private medical practitioner (LMD)
5	Nursing/convalescent home/hospice.
6	Autopsy only
7	Death Certificate Only (To be used by the MCR only)
8	Other hospital outpatient units/surgery centers (effective with diagnosis on or after 1/1/2006)

SEQUENCE NUMBER – HOSPITAL

Item Length: 2

NAACCR Item #560

NAACCR Name: Sequence Number—Hospital

XML NAACCR ID: sequenceNumberHospital

Definition:

Indicates the sequence of malignant neoplasms and benign and borderline brain and central nervous system tumors over the lifetime of the patient even if all were not diagnosed/treated at the reporting facility.

Coding Instructions:

- Codes 00-59 and 99 indicate neoplasms of in situ or malignant behavior (Behavior equals 2 or 3). Codes 60-88 indicate neoplasms of non-malignant behavior (Behavior equals 0 or 1).
- Code 00 only if the patient has a single malignant primary. If the patient develops a subsequent malignant or in situ primary tumor, change the code for the first tumor from 00 to 01, and number subsequent tumors sequentially.
- Code 60 only if the patient has a single non-malignant primary. If the patient develops a subsequent nonmalignant primary, change the code for the first tumor from 60 to 61, and assign codes to subsequent non-malignant primaries sequentially.
- If two or more malignant or in situ neoplasms are diagnosed at the same time, assign the lowest sequence number to the diagnosis with the worst prognosis. If no difference in prognosis is evident, the decision is arbitrary.
- If two or more non-malignant neoplasms are diagnosed at the same time, assign the lowest sequence number to the diagnosis with the worst prognosis. If no difference in prognosis is evident, the decision is arbitrary.
- Because the time period for Sequence Number is the patient's lifetime, reportable neoplasms not included in the hospital registry are allotted a sequence number. For example, a registry may contain a single record for a patient with a sequence number of 02 because the first reportable neoplasm occurred before the hospital's reference date or was diagnosed and treated at another facility. Document all previous primaries and the date or approximate date of diagnosis. Also document where the patient was diagnosed/treated if known.

Malignant, in situ or invasive:

Code	Definition
00	One malignant primary only in the patient's lifetime.
01	First of two or more independent malignant primaries.
02-59	Actual sequence of two or more malignant primary tumors
99	Unspecified malignant sequence number or unknown Always try to record the sequence rather than code 99.

Benign and Borderline Brain and Central Nervous System Tumors

Code	Definition
60	Only one non-malignant primary.
61	First of two or more independent non-malignant primaries.
62-87	Actual sequence of two or more <i>in situ</i> or malignant primary tumors.
88	Unspecified number of non-malignant tumors in this category. Always try to record the sequence rather than code 88.

Coding Examples:

Code	Explanation
00	A patient with no history of previous cancer is diagnosed with melanoma <i>in situ</i> on April 13, 2003.
01	The sequence number is changed when a patient with melanoma diagnosed on April 13, 2003, is diagnosed with a subsequent prostate cancer on August 30, 2004
02	The sequence number assigned to a prostate diagnosed on August 30, 2004, following a melanoma diagnosed on April 13, 2003.
03	A patient is admitted to the hospital for surgery as part of their first course of treatment for breast cancer. The patient had been diagnosed with two previous primary cancers. The other primaries were not diagnosed or treated at your facility. No abstracts are completed for primaries 01 or 02. Document the previous primaries in the text field.
60	The sequence number assigned to a benign brain tumor diagnosed on November 1, 2005, following a melanoma diagnosed on April 13, 2003 and a prostate cancer diagnosed on August 30, 2004.

DEMOGRAPHICS

NAME – LAST

Item Length: 40
NAACCR Item #2230
NAACCR Name: Name—Last
XML NAACCR ID: nameLast

Definition:

Last name of the patient, may also be called surname

Coding Instructions:

- Truncate name if more than 40 letters long.
- Blanks, spaces, hyphens, and apostrophes are allowed. Do not use other punctuation.
- If the patient’s last name is unknown, enter UNKNOWN.
- **If the name becomes available at a later admission, please send the correction to the MCR.**
- Record the most current name and update this data item if the last name changes. Enter previous names in the Alias data item (not included in this manual).

Examples:

Code	Reason
MC NAIR	Mc Nair
O’KEEFE	O’Keefe
JONES-VESSEY	Janet Jones marries Mike Vessey and changes her last name to Jones-Vessey.
UNKNOWN	If the patient’s last name is unknown, enter UNKNOWN

NAME - FIRST

Item Length: 40
NAACCR Item #2240
NAACCR Name: Name—First
XML NAACCR ID: nameFirst

Definition:

First name of the patient, may also be referred to as given name.

Coding Instructions:

- Truncate name if more than 40 letters long.
- Spaces, hyphens, and apostrophes are allowed. Do not use other punctuation.
- If the patient’s first name is unknown, enter UNKNOWN.
- **If the name becomes available at a later admission, please send the correction to the MCR.**
- Do not record nicknames in First Name.
 - Record nicknames in the Alias data item
 - **Example:** The patient’s nickname is Mike and the first name is Michael. Record Michael in the First Name field.

Examples:

Code	Reason
MICHAEL	Patient is admitted as Michael Hogan. Enter Hogan as the last name and Michael as the first name.
UNKNOWN	If patient’s first name is unknown, code UNKNOWN

NAME - MIDDLE

Item Length: 40
NAACCR Item #2250
NAACCR Name: Name—Middle
XML NAACCR ID: nameMiddle

Definition:

Middle name or, if middle name is unavailable, middle initial of the patient.

Coding Instructions:

- Truncate the name if more than 40 letters long.
- Blanks, spaces, hyphens, and apostrophes are allowed. Do not use other punctuation.
- Record the middle initial if the full middle name is not known.
- Leave blank if the middle name or middle initial is unknown. Do not use abbreviations such as NMI or NMN to denote that the middle initial/name is unknown.
- This field may be updated if the name changes.

Examples:

Code	Reason
DAVID	Patient is admitted as John David Smith. Enter Smith as the last name, John as the first name, and David as the middle name.
D	Patient is admitted as John D. Smith. Enter Smith as the last name, John as the first name, and D as the middle name.
(leave blank)	If patient does not have a middle name or initial, or if the middle name or initial are unknown, do not fill in the space.

NAME - BIRTH SURNAME

Item Length: 40

NAACCR Item #2332

NAACCR Name: Name—Birth Surname

XML NAACCR ID: nameBirthSurname

Description

Birth surname, effective 01/01/2021, is a gender-neutral replacement for the NAACCR data item Name – Maiden [2390]. Birth Surname reflects the last name of the patient at birth regardless of gender or marital status. Allowable values for Birth Surname are identical to those used for Name—Maiden.

Coding Instructions:

- Truncate the name if more than 40 letters long.
- Blanks, spaces, hyphens, and apostrophes are allowed. Do not use other punctuation.
- Record when known regardless of value in the Sex data item.
- Leave blank if the middle name or middle initial is unknown. Do not use abbreviations such as NMI or NMN to denote that the middle initial/name is unknown.

Examples:

Code	Reason
MC NAIR	Mc Nair
O'KEEFE	O'Keefe
JONES-VESSEY	Janet Jones marries Mike Vessey and changes her last name to Jones-Vessey.
UNKNOWN	If the patient's last name is unknown, enter UNKNOWN

NAME - ALIAS

Item Length: 40
NAACCR Item #2280
NAACCR Name: Name—Alias
XML NAACCR ID: nameAlias

Description

Records an alternate name or “AKA” (also known as) used by the patient, if known. Note that the birth surname (AKA maiden name) is entered in Name-Birth Surname.

NAME - SUFFIX

Item Length: 3
NAACCR Item #2270
NAACCR Name: Name—Suffix
XML NAACCR ID: nameSuffix

Description

Title that follows a patient's last name, such as a generation order or credential status (e.g., "MD," "Jr.")

SOCIAL SECURITY NUMBER

Item Length: 9
NAACCR Item #2320
NAACCR Name: Social Security Number
XML NAACCR ID: socialSecurityNumber

Definition:

Records the patient's Social Security number.

This data item can be used to identify patients with similar names.

Coding Instructions:

- Code the patient's Social Security number.
- A patient's Medicare claim number may not always be identical to the person's Social Security number.
- Code Social Security numbers that end with "B" or "D" as 999999999. This indicates the patient is using a spouse's social security number.
- **If the social security number becomes available at a later admission, update the number and send the correction to the MCR.**

Code	Definition
Fill spaces	Record the patient's Social Security number (SSN) without dashes or letter suffixes.
999999999	When the patient does not have a Social Security number or the information is not available.

BIRTH DATE

Item Length: 8
NAACCR Item #240
NAACCR Name: Date of Birth
XML NAACCR ID: dateOfBirth

Definition:

Identifies the patient's birth date.

This data item is useful for patient identification and analyzing tumors by age groups.

Coding Instructions:

- Record the patient's date of birth as indicated in the patient record.
- For *in utero* diagnosis and treatment, record the actual date of birth.
- If age at diagnosis and year of diagnosis are known, but year of birth is unknown, then year of birth should be calculated and so coded. Month and day would be left blank.
- Every effort must be made to collect the correct birth date.
- Estimate the birth date and note in the text field that the birth date is estimated. It is better to estimate the date than code as unknown.
- **Corrections to birth date should be sent to the MCR.**

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of birth is the year, month and day that the patient was born.
1937	A patient is admitted with unknown birth date. Note in the text field this is an <i>estimated birth date</i> . Record for example, a patient who is about 70 would have an estimated birth date of 1937.

BIRTH DATE FLAG

Item Length: 2
NAACCR Item #241
NAACCR Name: Date of Birth Flag
XML NAACCR ID: dateOfBirthFlag

Definition:

This flag explains why no appropriate value is in the field Birth date. Prior to 2010, date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions:

- If a valid date is coded in birth date, then leave this field blank. A valid date may be just a partial date.
- If birth date cannot be determined, then leave birth date blank and record 12 in this field indicating that the date is unknown.

Code	Definition
12	A proper value is applicable but unknown (e.g., birth date cannot be determined)
Blank	A valid date value is provided in item Birth date.

Coding Examples:

Code	Explanation
Blank	Full date is known (CCYYMMDD) for Birth date
Blank	Partial date is known (CCYYMM or CCYY) for Birth date
12	Date is completely unknown for Birth date and cannot be estimated

BIRTHPLACE—STATE

Item Length: 2

NAACCR Item #252

NAACCR Name: Birthplace—State

XML NAACCR ID: birthplaceState

Definition:

USPS abbreviation for state, commonwealth, U.S. possession in which the patient was born. CanadaPost abbreviations for the Canadian provinces can also be recorded if the patient was born in Canada. If the patient has multiple primaries, this data item should be coded the same for each primary.

Coding Instructions:

- Use the most specific code.

Code	Definition	Code	Definition
AL	Alabama	MI	Michigan
AK	Alaska	MN	Minnesota
AZ	Arizona	MS	Mississippi
AR	Arkansas	MO	Missouri
CA	California	MT	Montana
CO	Colorado	NE	Nebraska
CT	Connecticut	NV	Nevada
DE	Delaware	NH	New Hampshire
DC	District of Columbia	NJ	New Jersey
FL	Florida	NM	New Mexico
GA	Georgia	NY	New York
HI	Hawaii	NC	North Carolina
ID	Idaho	ND	North Dakota
IL	Illinois	OH	Ohio
IN	Indiana	OK	Oklahoma
IA	Iowa	OR	Oregon
KS	Kansas	PA	Pennsylvania
KY	Kentucky	RI	Rhode Island
LA	Louisiana	SC	South Carolina
ME	Maine	SD	South Dakota
MD	Maryland	TN	Tennessee
MA	Massachusetts	TX	Texas

Code	Definition	Code	Definition
UT	Utah	ZZ	U.S., NOS; Canada, NOS; Country Unknown
VT	Vermont	AB	Alberta
VA	Virginia	BC	British Columbia
WA	Washington	MB	Manitoba
WV	West Virginia	NB	New Brunswick
WI	Wisconsin	NL	Newfoundland and Labrador
WY	Wyoming	NS	Nova Scotia
AS	American Samoa	NT	Northwest Territories
GU	Guam	NU	Nunavut
MP	Northern Mariana Islands	ON	Ontario
PW	Palau	PE	Prince Edward Island
PR	Puerto Rico	QC	Quebec
UM	U.S. Outlying Islands	SK	Saskatchewan
VI	Virgin Islands of the United States	US	Resident of United States, NOS
FM	Federated States of Micronesia	AA	APO/FPO for Armed Services America
MH	Marshall Islands	AE	APO/FPO for Armed Services Europe
TT	Trust Territories	AP	APO/FPO for Armed Services Pacific
XX	Country Known, Not U.S., Not Canada		
YT	Yukon Territories		
YY	Country Unknown, Not U.S., Not Canada		

Coding Examples:

Code	Explanation
MS	If the state in which the patient was born is Mississippi, then uses the USPS code for the state of Mississippi.
XX	Born in a country other than the U.S. (including US territories, commonwealths, and possessions) or Canada and the country is known.
YY	Born in a country other than the U.S. (including US territories, commonwealths, and possessions) or Canada and the country is unknown.
US	Patient was born in the U.S. (including U.S. territories, commonwealths, or possessions) but the state is not given.
ZZ	Place of birth is unknown

BIRTHPLACE—COUNTRY

Item Length: 3

NAACCR Item #254

NAACCR Name: Birthplace—Country

XML NAACCR ID: birthplaceCountry

Definition:

Code for the country where the patient was born. If the patient has multiple tumors, all records should contain the same code. Place of birth is helpful in patient matching and quality review of race and ethnicity.

Coding Instructions:

- Use the most specific code.

Code	Definition	Code	Definition
ABW	Aruba	BGD	Bangladesh
AFG	Afghanistan	BGR	Bulgaria
AGO	Angola	BHR	Bahrain
AIA	Anguilla	BHS	Bahamas
ALA	Aland Islands	BIH	Bosnia and Herzogovina
ALB	Albania	BLM	St. Barthelemy
AND	Andorra	BLR	Belarus
ARE	United Arab Emirates	BLZ	Belize
ARG	Argentina	BMU	Bermuda
ARM	Armenia	BOL	Bolivia
ASM	American Samoa	BRA	Brazil
ATA	Antarctica	BRB	Barbados
ATF	French Southern Territories	BRN	Brunei
ATG	Antigua and Barbuda	BTN	Bhutan
AUS	Australia	BVT	Bouvet Island
AUT	Austria	BWA	Botswana
AZE	Azerbaijan	CAF	Central African Republic
BDI	Burundi	CAN	Canada
BEL	Belgium	CHE	Switzerland
BEN	Benin	CHL	Chile
BES	Bonaire, Saint Eustatius and Saba	CHN	China
BFA	Burkina Faso	CIV	Cote d'Ivoire

Code	Definition	Code	Definition
CMR	Cameroon	FSM	Micronesia
COD	Congo, Democratic Republic of	GAB	Gabon
COG	Congo	GBR	United Kingdom
COK	Cook Islands	GEO	Georgia
COL	Columbia	GGY	Guernsey
COM	Comoros	GHA	Ghana
CPV	Cape Verde	GIB	Gibraltar
CRI	Costa Rica	GIN	Guinea
CSK	Czechoslovakia	GLP	Guadelupe
CUB	Cuba	GMB	Gambia
CUW	Curacao	GNB	Guinea Bissau
CXR	Christmas Island	GNQ	Equatorial Guinea
CYM	Cayman Islands	GRC	Greece
CYP	Cyprus	GRD	Grenada
CZE	Czech Republic	GRL	Greenland
DEU	Germany	GTM	Guatemala
DJI	Djibouti	GUF	French Guiana
DMA	Dominica	GUM	Guam
DNK	Denmark	GUY	Guyana
DOM	Dominican Republic	HKG	Hong Kong
DZA	Algeria	HMD	Heard Island & McDonalds Islands
ECU	Ecuador	HND	Honduras
EGY	Egypt	HRV	Croatia
ENG	England	HTI	Haiti
ERI	Eritrea	HUN	Hungary
ESH	Western Sahara	IDN	Indonesia (Dutch East Indies)
ESP	Spain	IMN	Isle of Man
EST	Estonia	IND	India
ETH	Ethiopia	IOT	British Indian Ocean Territory
FIN	Finland	IRL	Ireland
FJI	Fiji	IRN	Iran
FLK	Falkland Islands	IRQ	Iraq
FRA	France	ISL	Iceland
FRO	Faroe Islands	ISR	Israel

Code	Definition	Code	Definition
ITA	Italy	MKD	Macedonia
JAM	Jamaica	MLI	Mali
JEY	Jersey	MLT	Malta
JOR	Jordan	MMR	Myanmar
JPN	Japan	MNE	Montenegro
KAZ	Kazakhstan	MNG	Mongolia
KEN	Kenya	MNP	Northern Mariana Islands
KGZ	Kyrgyzstan	MOZ	Mozambique
KHM	Cambodia	MRT	Mauritania
KIR	Kiribati	MSR	Montserrat
KNA	St. Kitts and Nevis	MTQ	Martinique
KOR	Korea, NOS	MUS	Mauritius
KOR	South Korea	MWI	Malawi
KWT	Kuwait	MYS	Malaysia
LAO	Laos	MYT	Mayotte
LBN	Lebanon	NAM	Namibia
LBR	Liberia	NCL	New Caledonia
LBY	Libya	NER	Niger
LCA	St. Lucia	NFK	Norfolk Island
LIE	Liechtenstein	NGA	Nigeria
LKA	Sri Lanka	NIC	Nicaragua
LSO	Lesotho	NIR	Northern Ireland (Ulster)
LTU	Lithuania	NIU	Niue
LUX	Luxembourg	NLD	Netherlands
LVA	Latvia	NOR	Norway
MAC	Macao	NPL	Nepal
MAF	Saint Martin (French part)	NRU	Nauru
MAR	Morocco	NZL	New Zealand
MCO	Monaco	OMN	Oman
MDA	Moldova	PAK	Pakistan
MDG	Madagascar	PAN	Panama
MDV	Maldives	PCN	Pitcairn Islands
MEX	Mexico	PER	Peru
MHL	Marshall Islands	PHL	Philippines

Code	Definition	Code	Definition
PLW	Palau(Trust Territory of Pacific Islands)	SVN	Slovenia
PNG	Papua New Guinea	SWE	Sweden
POL	Poland	SWZ	Swaziland
PRI	Puerto Rico	SXM	Sint-Maarten
PRK	North Korea	SYC	Seychelles
PRT	Portugal	SYR	Syria
PRY	Paraguay	TCA	Turks and Caicos
PSE	Palestine Territory, Occupied	TCD	Chad
PYF	French Polynesia	TGO	Togo
QAT	Qatar	THA	Thailand
REU	Réunion	TJK	Tajikistan
ROU	Romania	TKL	Tokelau Islands (New Zealand)
RUS	Russia	TKM	Turkmenistan
RWA	Rwanda	TLS	Timor-Leste
SAU	Saudi Arabia	TON	Tonga
SCT	Scotland	TTO	Trinidad and Tobago
SDN	Sudan	TUN	Tunisia
SEN	Senegal	TUR	Turkey
SGP	Singapore	TUV	Tuvalu
SGS	S Georgia & S Sandwich Islands	TWN	Taiwan
SHN	St Helena	TZA	Tanzania
SJM	Svalbard & Jan Mayen	UGA	Uganda
SLB	Solomon Islands	UKR	Ukraine
SLE	Sierra Leon	UMI	U.S. Minor Outlying Islands
SLV	El Salvador	URY	Uruguay
SMR	San Marino	USA	United States
SOM	Somalia	UZB	Uzbekistan
SPM	St Pierre and Miquelon	VAT	Vatican City
SRB	Serbia	VCT	St. Vincent & the Grenadines
SSD	South Sudan	VEN	Venezuela
STP	Sao Tome & Principe	VGB	British Virgin Islands
SUR	Suriname	VIR	U.S. Virgin Islands
SVK	Slovakia	VNM	Vietnam

Code	Definition	Code	Definition
VUT	Vanuatu		
WLF	Wallis and Fotuna		
WLS	Wales		
WSM	Samoa		
YEM	Yemen		
YUG	Yugoslavia		
ZAF	Republic of South Africa		
ZMB	Zambia		
ZWE	Zimbabwe		
ZZA	Asia, NOS		
ZZC	Central America, NOS		
ZZE	Europe, NOS		
ZZF	Africa, NOS		
ZZN	North America, NOS		
ZZP	Pacific, NOS		
ZZS	South America, NOS		
ZZU	Unknown		
ZZX	Non-US/Canada, NOS		

Coding Examples:

Code	Explanation
USA	United States
CAN	Canada
MEX	Mexico
ZZU	Place of Birth Unknown

ADDRESS AT DIAGNOSIS – NO & STREET

Item Length: 60

NAACCR Item #2330

NAACCR Name: Addr at Dx—No & Street

XML NAACCR ID: addrAtDxNoStreet

Definition:

The patient's address at diagnosis is the patient's place of residence at the time of the original diagnosis. It does not change if the patient moves. If the patient has more than one primary tumor, the address at diagnosis may be different for each primary.

Normally a residence is the home named by the patient. Legal status and citizenship are not factors in residency decisions. Rules of residency are identical to or comparable with the rules of the Census Bureau's definition, "the place where he or she lives and sleeps most of the time or the place the person considers to be his or her usual home." Vital statistics rules may differ from Census rules. **Do not record the residence from the death certificate.** Review each case carefully.

RULES FOR PERSONS WITH AMBIGUOUS RESIDENCE:

Persons with more than one residence (summer and winter homes): Use the address the patient specifies if a usual residence is not apparent.

Persons with no usual residence (transients, homeless): Use the address of the place the patient was staying when the cancer was diagnosed. This could be a shelter or a diagnosing facility. In the supplemental address field, add the word "homeless" or "transients", as applicable.

Persons away at school: College students are residents of the school area. Boarding school students below the college level are residents of their parents' homes.

Persons away in institutions: The Census Bureau states, "Persons under formally authorized, supervised care or custody," are residents of the institution. This includes the following:

- Incarcerated persons
- Persons in nursing, convalescent, and rest homes
- Persons in homes, schools, hospitals, or wards for the physically disabled, mentally retarded, or mentally ill
- Long-term residents of other hospitals, such as Veterans Affairs (VA) hospitals.

Temporary Residents

Code the place of usual residence rather than the temporary address for migrant workers, educators temporarily assigned to a university in the area, persons temporarily residing with family during cancer treatment, military personnel on temporary duty assignment (TDY), and boarding school students below college level (code the parent's residence).

Persons in the Armed Forces and on Maritime Ships: Members of the armed forces are residents of the installation area. Use the stated address for military personnel and their families. Military personnel may use the installation address or the surrounding community's address.

Coding Instructions:

- Record the number and street address or the rural mailing address of the patient's usual residence when the tumor was diagnosed.
- Record the physical number and street address of the patient at diagnosis. If the patient also has a Post Office (PO) Box address, record the PO Box address in Address at Diagnosis—Supplemental.
- The address should be fully spelled out with standardized use of abbreviations and punctuation per U.S. Postal Service postal addressing standards. The USPS Postal Addressing Standards, Pub 28, April 2010 can be found on the Internet at <http://pe.usps.gov/cpim/ftp/pubs/pub28/pub28.pdf>.
- Abbreviations should be limited to those recognized by the Postal Service standard abbreviations. They include, **but are not limited to:**

APT (apartment)	AVE (avenue)	BLDG (building)
BLVD (boulevard)	CIR (circle)	CT (court)
DEPT (department)	DR (drive)	E (east)
FL (floor)	N (north)	NE (northeast)
NW (northwest)	PKWY (parkway)	RD (road)
RM (room)	ST (street)	STE (suite)
S (south)	SE (southeast)	SW (southwest)
W (west)	CV (cove)	EXT (Extension)
HWY (highway)	PL (place)	RTE (route)
- Do not update this data item if the patient's address changes.
- See "Residency Rules" for further instructions.
- Punctuation marks should be avoided, except when punctuation is necessary to convey the meaning (a period in 39.2 RD, a slash in 100 ½ FIFTH ST, or a hyphen in 259-02 SMITH AVE). Use of the pound sign (#) to designate address units should be avoided whenever possible. If a pound sign is used, there must be a space between the pound sign and the secondary number.
- If the patient has multiple tumors, the address may be different for different primaries.
- Leave blanks between numbers and words.
- If the patient's address is unknown, enter UNKNOWN.

Coding Examples:

Code	Definition
100 EASY ST APT 302	Use recognized USPS standardized abbreviations; do not use punctuation unless absolutely necessary to clarify an address; leave blanks between numbers and words
UNKNOWN	If the patient's address is unknown, enter UNKNOWN

SUPPLEMENTAL PATIENT ADDRESS, NUMBER & STREET AT DIAGNOSIS

Item Length: 60

NAACCR Item #2335

NAACCR Name: Addr at Dx—Supplementl

XML NAACCR ID: addrAtDxSupplementl

Definition:

Provides the ability to store additional address information such as the name of a place or facility, a nursing home, or the name of an apartment complex. If the patient has multiple tumors, the address at diagnosis may be different for every tumor.

Coding Instructions:

- Record the place or facility (i.e., a nursing home or name of an apartment complex) of the patient's usual residence when the tumor was diagnosed.
- If the patient has a known street address and a post office box address, the post office box address should be placed in this field. The street address should be put in Address at Diagnosis-No & Street.
- Do not update this data item if the patient's address changes.
- Leave blank if this data item is not needed.
- See **ADDRESS AT DIAGNOSIS – NO & STREET** for further instructions.

Example:

SUNNY VIEW NURSING HOME

ADDRESS AT DIAGNOSIS - CITY

Item Length: 50
NAACCR Item #70
NAACCR Name: Addr at DX—City
XML NAACCR ID: addrAtDxCity

Definition:

Name of the city in which the patient resides at the time of diagnosis. If the patient has multiple primaries, the city of residence may be different for each primary.

Coding Instructions:

- Record the city of the physical number and street address of the patient at diagnosis. If the patient also has a Post Office (PO) Box address, record the PO Box address in Address at Diagnosis—Supplemental. Ensure the city recorded matches the street address and not the PO Box address if the two addresses have different cities.
- Do not update this data item if the patient’s city/town of residence changes.
- See **ADDRESS AT DIAGNOSIS – NO & STREET** for further instructions.
- Do not use punctuation, special characters, or numbers.
- Abbreviate where necessary.
- If the patient’s city or town is unknown type UNKNOWN.

Code	Definition
CITY NAME	Do not use punctuation, special characters, or numbers. Abbreviate when necessary.
UNKNOWN	If the patient’s city or town is unknown.

ADDRESS AT DIAGNOSIS - STATE

Item Length: 2
NAACCR Item #80
NAACCR Name: Addr at DX—State
XML NAACCR ID: addrAtDxState

Definition:

USPS abbreviation for the state, territory, commonwealth, or U.S. possession in which the patient resides at the time of diagnosis. If the patient has multiple primaries, the state of residence may be different for each primary.

Coding Instructions:

- If the patient is a foreign resident, then code either XX or YY depending on the circumstance.
- If the patient is a U.S. resident, but the state is unknown, code US.
- If the patient is a Canadian resident and the province is unknown, code CD.

Code	Definition
MS	If the state in which the patient resides at the time of diagnosis and treatment is Mississippi, then use the USPS code for the state of Mississippi
US	Resident of the United States, NOS (state/commonwealth/territory/possession unknown)
CD	Resident of Canada, NOS (province/territory unknown)
XX	Resident of a country other than the U.S. or Canada (including territories, commonwealths, or possessions) and the country is known .
YY	Resident of a country other than the U.S. or Canada (including its territories, commonwealths, or possessions) and the country is unknown .
ZZ	Residence unknown

ADDRESS AT DIAGNOSIS – POSTAL CODE

Item Length: 9

NAACCR Item #100

NAACCR Name: Addr at DX—Postal Code

XML NAACCR ID: addrPostalCode

Definition:

Postal code of the patient’s address at diagnosis. If the patient has multiple primaries, the postal code at diagnosis may be different for each primary.

Coding Instructions:

- For U.S. residents, use either the 5-digit or the extended 9-digit ZIP code. Blanks follow the 5-digit code.
- For Canadian residents, record the six-character alphanumeric postal code.
- When available, record the postal code for other countries.
- Do not update this data item if the patient’s postal code changes.
- See **ADDRESS AT DIAGNOSIS – NO & STREET** for further instructions.
- If the patient has both a street address and a Post Office Box address, record the street address in Address at Diagnosis-Number and Street and make sure the city and postal code match the street address and not the Post Office Box address.

Code	Definition:
(fill spaces)	The patient’s nine-digit U.S. extended postal code. Do not record hyphens.
38655_ _ _ _	When the nine-digit extended U.S. ZIP Code is not available, record the five-digit postal code, left justified, followed by four blanks.
888888888	Permanent address in a country other than Canada, United States, or U.S. possessions and postal code is unknown.
999999999	Resident of the U.S. (including its possessions, etc.) or Canada and the postal code is unknown

COUNTY AT DIAGNOSIS

Item Length: 3

NAACCR Item #90

NAACCR Name: County at DX Reported

XML NAACCR ID: countyAtDx

Definition:

Code the county of the patient's residence at the time the reportable tumor is diagnosed. If the patient has multiple primaries, the county at diagnosis may be different for each primary.

Coding Instructions:

- Refer to the table below for a list of Mississippi Counties and their FIPS codes.
- For residents of states other than Mississippi and residents of other countries at the time of diagnosis, code 998 for diagnosis county.
- If the county of the patient is unknown, code 999.
- Do not update this data item if the patient's county of residence changes.

FIPS Code	County	FIPS Code	County	FIPS Code	County	FIPS Code	County
001	Adams	043	Grenada	085	Lincoln	127	Simpson
003	Alcorn	045	Hancock	087	Lowndes	129	Smith
005	Amite	047	Harrison	089	Madison	131	Stone
007	Attala	049	Hinds	091	Marion	133	Sunflower
009	Benton	051	Holmes	093	Marshall	135	Tallahatchie
011	Bolivar	053	Humphreys	095	Monroe	137	Tate
013	Calhoun	055	Issaquena	097	Montgomery	139	Tippah
015	Carroll	057	Itawamba	099	Neshoba	141	Tishomingo
017	Chickasaw	059	Jackson	101	Newton	143	Tunica
019	Choctaw	061	Jasper	103	Noxubee	145	Union
021	Claiborne	063	Jefferson	105	Oktibbeha	147	Walthall
023	Clarke	065	Jefferson Davis	107	Panola	149	Warren
025	Clay	067	Jones	109	Pearl River	151	Washington
027	Coahoma	069	Kemper	111	Perry	153	Wayne
029	Copiah	071	Lafayette	113	Pike	155	Webster
031	Covington	073	Lamar	115	Pontotoc	157	Wilkinson
033	DeSoto	075	Lauderdale	117	Prentiss	159	Winston
035	Forrest	077	Lawrence	119	Quitman	161	Yalobusha
037	Franklin	079	Leake	121	Rankin	163	Yazoo
039	George	081	Lee	123	Scott	998	Out of State
041	Greene	083	Leflore	125	Sharkey	999	Unknown

SEX

Item Length: 1
Allowable Values: 1–6, 9
NAACCR Item #220
STORE 2018 pg. 90

Definition:

Identifies the sex of the patient.

Transsexual: A person who was assigned under one gender at birth based on physical characteristics but who self-identifies psychologically and emotionally as the other gender.

Transgender: See Transsexual

Transgendered person: A person who identifies with or expresses a gender identity that differs from the one which corresponds to the person's sex at birth.

Coding Instructions:

- Record the patient's sex as indicated in the medical record.
- If the medical record indicates unknown for sex contact the patient's primary physician to identify the sex of the patient.
- When gender is not known
 - Assign code 1 when the primary site is C600-C639
 - Assign code 2 when the primary site is C510-C589
 - Assign code 9 for primary sites not included above.
- Natality was added in 2015 for transsexuals. However, the codes can also be used when abstracting cases diagnosed prior to 2015
- Assign code 3 for Intersexed (persons with sex chromosome abnormalities)
- Assign code 5 for transsexuals who are natively male or transsexuals with primary site of C600-C639
- Assign code 6 for transsexuals who are natively female or transsexuals with primary site of C510-C589
- Assign code 4 for transsexuals with unknown natal sex and primary site is NOT C501-C589 or C600-C639

Codes	Definition
1	Male
2	Female
3	Other (intersex, disorders of sexual development/DSD)
4	Transsexual, NOS
5	Transsexual, natal male
6	Transsexual, natal female
9	Not Stated/Unknown

RACE 1, 2, 3, 4, 5

Item length: 2

NAACCR Item #160, 161, 162, 163, 164

NAACCR Name: Race 1, Race 2, Race 3, Race 4, Race 5

XML NAACCR ID: race1, race2, race3, race4, race5

Definition:

Race and ethnicity are defined by specific physical, hereditary and cultural traditions or origins, not necessarily by birthplace, place of residence, or citizenship. 'Origin' is defined by the U.S. Census Bureau as the heritage, nationality group, lineage, or in some cases, the country of birth of the person or the person's parents or ancestors before their arrival in the United States.

The five race data items (Race 1- Race 5) make it possible to code multiple races for one person, consistent with the 2000 Census. All resources in the facility, including the medical record, face sheet, physician, and nursing notes, photographs, and any other sources, must be used to determine race. If a facility does not print race in the medical record but does maintain it in an electronic form, the electronic data must also be reviewed.

Document how the race code(s) was (were) determined in a text field.

Coding Instructions:

- Do not use patient name as the basis for coding race.
 - a. See Coding Instruction 15, Exception, for the only situation in which name is taken into account when coding race.
- Code race using the highest priority source available according to the list below (a is the highest and c is the lowest) when race is reported differently by two or more sources.
 - a. The patient's self-declared identification
 - b. Documentation in the medical record
 - c. Death certificate
- Assign the same race code(s) for all tumors for one patient.
- Code the race(s) of the patient in data items Race 1, Race 2, Race 3, Race 4, and Race 5.
 - a. Code 88 for the remaining race data items (Race2-Race5) when at least one race, but fewer than five races, are reported.
- Priorities for coding multiple races:
 - a. Code 07 takes priority over all other codes.

- b. Codes 02-32, 96-98 take priority over code 01
 - c. Codes 04-17 take priority over code 96
 - d. Codes 16-17 take priority over code 15
 - e. Codes 20-32 take priority over code 97
 - f. Codes 02-32 and 96-97 take priority over code 98
 - g. Code 98 takes priority over code 99
- Code as 01 (White) when the race is described as Caucasian regardless of place of birth or there is a statement that the patient is Hispanic or Latino(a) with no further information. **DO NOT code 98 for Hispanic, NOS**
 - Code race as 02 (Black) when the stated race is African-American, Black or Negro.
 - Assign code 03 for any person stated to be Native American (western hemisphere) or Indian (whether from North, Central, South or Latin America)
 - Assign a specific code when a specific Asian race is stated. Code 96 is not applicable when a specific race is known.
 - Code a race based on birthplace information when the race is recorded as Oriental, Mongolian, or Asian and the place of birth is recorded as China, Japan, the Philippines, or another Asian nation.
 - Use the appropriate non-specific code 96 (Other Asian including Asian, NOS and Oriental, NOS), 97 (Pacific Islander, NOS) or 98 (Other) when there is no race code for a specific race.
 - When a face-sheet indicates “Race Other,” look for other descriptions of the patient’s race. **When no further race information is available, code race as 99 (unknown)** and document that the face-sheet indicates “Race Other,” and no further information is available.
 - Do not use code 96, 97, or 98 for a patient described as “multi-racial.”
 - If Race 1 is coded 99, then Race 2-5 must all be coded 99. Do not code Race 2-Race 5 as 99 if Race 1 is not coded 99.
 - Refer to Appendix D of the current *SEER Program Coding and Staging Manual* (<http://www.seer.cancer.gov/tools/codingmanuals/index.html>) to determine race when only nationality or place of birth is known.

Exception: Code Race 1 through 5 as 99 (Unknown) when patient’s name is incongruous with the race inferred on the basis of nationality.

- Example: Patient’s name is Siddhartha Rao and the birthplace listed is England. Code Race 1 through Race 5 as 99 (Unknown).

- Use the associated text field to document
 - a. Why a particular race code was chosen when there are discrepancies in race information.
 - b. That no race information is available.
- Patient photographs may be used with caution to determine race in the absence of documented race information.
 - a. Use caution when interpreting a patient photograph to assist in determining race. Review the patient record for a statement to verify race. The use of photographs alone to determine race may lead to misclassification of race.
- Code the race data items in the order stated when no other priority applies.
- The race of parents, when known, may be used with caution to determine patient’s race in the absence of other more specific information.

Code	Definition
01	White
02	Black
03	American Indian, Aleutian, Alaska Native or Eskimo (includes all indigenous populations of the Western hemisphere)
04	Chinese
05	Japanese
06	Filipino
07	Hawaiian
08	Korean
10	Vietnamese
11	Laotian
12	Hmong
13	Kampuchean, (Cambodian)
14	Thai
15	Asian Indian or Pakistani, NOS (Effective with 1/1/2010 dx)
16	Asian Indian (Effective with 1/1/2010 dx)
17	Pakistani (Effective with 1/1/2010 dx)
20	Micronesian, NOS
21	Chamorro/Chamoru
22	Guamanian, NOS
25	Polynesian, NOS

Code	Definition
26	Tahitian
27	Samoan
28	Tongan
30	Melanesian, NOS
31	Fiji Islander
32	New Guinean
88	No further race documented
96	Other Asian, including Asian, NOS and, Oriental, NOS
97	Pacific Islander, NOS
98	Other
99	Unknown

Coding Examples for Race 1:

Codes	Explanation
01,88,88,88,88	A patient is noted to be Hispanic NOS. Code also <i>Spanish/Hispanic Origin</i> .
08,88,88,88,88	Patient is described as Asian in a consult note and as second generation Korean-American in the history. Code Race 1 as 08 and Race 2 through Race 5 as 88. Do not code 96 in Race 2.
05,01,88,88,88	A patient has a Japanese father and a Caucasian mother. Code Race 1 to 05 Japanese, and Race 2 to white, 01. Always code the least common code to the first race field .
01,88,88,88,88	Patient is referred to as a native of Portugal
01,88,88,88,88	Patient is described as Arabian.
25,26,88,88,88	Patient states she has a Polynesian mother and Tahitian father. Code Race 1 as 25 (Polynesian), Race 2 as 26 (Tahitian) and Race 3 through Race 5 as 88.
04,02,88,88,88	Patient is stated to be Chinese and black. Code Race 1 as 04 (Chinese), code race 2 as 02 (Black). Code in the order stated when no other priority applies.
99,99,99,99,99	A patient's race is unknown. Race fields 2-5 must also be unknown.

SPANISH/HISPANIC ORIGIN

Item Length: 1

NAACCR Item #190

NAACCR Name: Spanish/Hispanic Origin

XML NAACCR ID: spanishHispanicOrigin

Definition:

Identifies persons of Spanish or Hispanic origin. Persons of Spanish or Hispanic origin may be of any race. If the patient has a Hispanic name, but there is reason to believe they are not Hispanic (e.g., the patient is a woman known to be non-Hispanic who has a Hispanic married name), then code the patient as non-Hispanic.

This code is used by hospital and central registries to identify whether or not the person should be classified as “Hispanic” for purposes of calculating cancer rates.

Coding Instructions:

- Coding Spanish Surname or Origin is not dependent on race. A person of Spanish
- Use all information to determine the Spanish/Hispanic Origin including
 - The ethnicity stated in the medical record
 - Self-reported information takes priority over other sources of information
 - Hispanic origin stated on the death certificate
 - Birthplace
 - Information about life history and/or language spoken found in the abstracting process
- Assign code 7 when the only evidence of the patient’s Hispanic origin is a surname or maiden name and there is no evidence that the patient is not Hispanic. Code 7 is ordinarily for central registry use only.
- Portuguese, Brazilians and Filipinos are not presumed to be Spanish or non-Spanish
 - Assign code 7 when the patient is Portuguese, Brazilian, or Filipino and their name appears on a Hispanic surname list.
 - Assign code 0 when the patient is Portuguese, Brazilian, or Filipino and their name does NOT appear on a Hispanic surname list.
- The following information should be referred to determine the Spanish/Hispanic Origin code: stated ethnicity in the medical record, stated Hispanic origin on the death

certificate, birthplace, information about the life history and/or language spoken found during the abstracting process, patient's last name or maiden name found on a list of Hispanic/Spanish names.

- A list of Hispanic surnames from the Census Bureau can be found at the following link:
<https://fcds.med.miami.edu/downloads/DataAcquisitionManual/dam2014/25%20Appendix%20E%20Census%20List%20of%20Spanish%20Surnames.pdf>
- If the patient has multiple tumors, all records should have the same code.

Codes	Definition:
0	Non-Spanish; non-Hispanic
1	Mexican (includes Chicano)
2	Puerto Rican
3	Cuban
4	South or Central American (except Brazil)
5	Other specified Spanish/Hispanic origin (includes European; excludes Dominican Republic)
6	Spanish, NOS, Hispanic, NOS, Latino, NOS There is evidence other than surname or surname at birth that the person is Hispanic, but he/she cannot be assigned to any category of 1–5.
7	Spanish surname only. The only evidence of the person's Hispanic origin is surname or surname at birth and there is no contrary evidence that the person is not Hispanic.
8	Dominican Republic. For use with patients who were diagnosed with cancer on January 1, 2005, or later.
9	Unknown whether Spanish or not stated in patient record.

TEXT--USUAL INDUSTRY

Item Length: 100
NAACCR Item #320
NAACCR Name: Text—Usual Industry
XML NAACCR ID: textUsualIndustry

Description

Source: “A Cancer Registrar’s Guide to Collecting Industry and Occupation”; DHHS (NIOSH) Publication No. 2011-173 (<http://www.cdc.gov/niosh/docs/2011-173/pdfs/2011-173.pdf>)

Text area for information about the patient’s usual industry.

Used to identify new work-related health hazards; serves as an additional measure of socioeconomic status; identifies industrial groups or worksite-related groups in which cancer screening or prevention activities may be beneficial.

The patient’s usual industry is the type of business or industry where the patient worked in his or her usual occupation.

Coding Instructions

- Be descriptive: Record the primary activity carried on by the industry at the location where the patient was employed.

Inadequate: “automobile industry”

Adequate: “automobile manufacturing”

- Be specific: In order to give a clear and exact description of the industry, the entry must indicate both a general and specific function for the employer.

General Industry (Inadequate)	Specific Industry (Adequate)
Mine	Copper mine
Manufacturer	Automobile manufacturer
Wholesale	Wholesale grocery
Retail	Retail bookstore
Construction	Road construction
Repair service	Shoe repair service

- Be complete: If the primary activity of the industry is unknown, record the name of the company (with city or town) in which the patient worked the most number of years before diagnosis.

Inadequate: “ABC, Inc.”

Adequate: “ABC, Inc., Los Angeles, CA”

- Instructions for reporting government agencies:
 - Record the level: federal, state, county.
Inadequate: “fire department”
Adequate: “city fire department”
 - Record the division of the agency, if available, to help clarify the specific activity of the patient.
 - Use full name of division/agency:
Inadequate: “Census”
Adequate: “U.S. Census Bureau”
- If a person worked only at home, then record industry as “own home.”
- If a patient worked at someone else’s home for pay, then record industry as “private home.”
- If patient ever worked outside the home, then report corresponding industry for longest-held job outside the home. Do not report “homemaker” in this case.
Note: This is an exception to the rule that the occupation with the greatest number of years should be recorded as “usual” occupation.
- If the patient is under 14 years of age, then record industry as “child.”
- If patient was a student at the time of diagnosis and had never held a job, then record industry as the type of school (“high school,” “college”)
- If patient was part of the military for most of his/her working life, then record industry as “military.”
- If patient was not a student or homemaker and had never worked, then record industry as “none.”
- “Unknown” should be entered only after you have tried your best to find job information in the medical record. It is better to enter “unknown” than to leave the field blank.
- A business at a person’s home should be reported in the same manner as regular business establishments. If work is in an office located in a private home, report the specific business. Do not report an individual’s name as the employer.
 Inadequate: “O’Keefe, Brown, & Smith”
 Adequate: “lawyer’s office”

- Here are some common incomplete entries to avoid:
 - “Retired”: If retired, enter the kind of work patient did during most of his or her working life if this can be determined. (Do not add “retired”.)
Inadequate: “retired plumber”
Adequate: “plumber”
 - “Institutionalized,” “Disabled,” or “Unemployed”: Do not record such a description if patient was ever employed. Record longest-held occupation and industry.
 - “Self-employed”: If self-employed, specify the kind of work performed.
Inadequate: “self-employed”
Adequate: “automobile manufacturing”

- Make sure that the recorded usual occupation matches the recorded industry.

Occupation	Business/Industry
Timber cutter	Logging
Shoe designer	Leather footwear factory
Tire tester	Tire manufacturing
Petroleum analyst	Petroleum refining
Carpenter	Building construction
Carpet installer	Retail carpet sales and installation company
Registered nurse	Hospital
Miner	Coal Mine
Mechanic, auto	Engine repair shop
Insurance agent	Life insurance company
Student	Junior college

TEXT USUAL OCCUPATION

Item Length: 100
NAACCR Item #310
NAACCR Name: Text—Usual Occupation
XML NAACCR ID: textUsualOccupation

Description

Source: “A Cancer Registrar’s Guide to Collecting Industry and Occupation”; DHHS (NIOSH) Publication No. 2011-173 (<http://www.cdc.gov/niosh/docs/2011-173/pdfs/2011-173.pdf>)

The patient’s usual occupation is the type of job the patient was engaged in for the longest time. It is not necessarily the highest paid job not the job considered the most prestigious, but the one that accounted for the greatest number of working years.

Coding Instructions

- Be descriptive: Record the word or words which most clearly describe the kind of work or type of duties performed by the patient.
Inadequate: “teacher”
Adequate: “preschool teacher,” “high school teacher”
- Be specific: General or vague terms are not adequate since they do not always provide enough information to code. You are allowed 100 characters.
Inadequate: “laborer”
Adequate: “residential bricklayer”

General Occupation (Inadequate)	Specific Occupation (Adequate)
Contractor	Building construction contractor
Consultant	Computer database consultant
Assembler	Aircraft engine assembler
Technician	Civil engineering technician
Laborer	Dairy farm laborer
Engineer	Chemical engineer, railroad engineer

- Be complete: Occupation entries that give only the department or a place of work are inadequate.
Inadequate: “worked in a warehouse”, “worked in a shipping department”
Adequate: “warehouse forklift operator”
- Here are some commonly confused occupations:
 - Contractor vs. skilled worker
 - A “contractor” mainly obtains contracts and supervises the work
 - A “skilled worker” works with his or her own tools as a carpenter, plasterer, plumber, or electrician.

- Machine operator vs. machinist vs. mechanic
 - A “machine operator” operates machines.
 - A “machinist” sets up and operates machines.
 - A “mechanic” repairs, installs, and adjusts machines
 - If a patient worked only at home, then record occupation as “homemaker.”
 - If a patient worked at someone else’s home for pay, then record occupation as “housekeeper” (or “nurse,” “babysitter,” etc.)
 - If patient ever worked outside the home, then report corresponding occupation for longest-held job outside the home. Do not report “homemaker” in such cases.
- Note:** This is an exception to the rule that the occupation with the greatest number of years should be recorded as “usual” occupation.
- If the patient is under 14 years of age, then record occupation as “child.”
 - If patient was a student at time of diagnosis and had never held a job, then record occupation as “student.”
 - If patient was part of the military for most of his/her working life, then record occupation as “military.”
 - If the patient was not a student or homemaker and had never worked, then record occupation as “never worked.”
 - “Unknown” should be entered only after you have tried your best to find job information in the medical record. It is better to enter “unknown” than to leave blank.
 - A business at a person’s home should be reported in the same manner as a regular business establishment. If work is in an office located in a private home, report the specific business. Do not report an individual’s name as the employer.
Inadequate: “works from home”
Adequate: “paralegal”
 - If retired, enter the kind of work patient did during most of his or her working life if this can be determined. (Do not add “retired.”) For example, record “plumber,” not “retired plumber.”
 - Do not record descriptions such as “institutionalized,” “disabled,” or “unemployed” if patient was ever employed. Record longest-held occupation.

- If self-employed, specify the kind of work performed.
Inadequate: “self-employed”
Adequate: “self-employed auto mechanic”

- Record “manager” only if patient worked most of time managing a business, but include specifics in these cases.
Inadequate: “manager”
Adequate: “operations manager”

PRIMARY PAYER AT DIAGNOSIS

Item Length: 2
NAACCR Item #630
NAACCR Name: Primary Payer at DX
XML NAACCR ID: primaryPayerAtDx

Description

Identifies the patient's primary payer/insurance carrier at the time of initial diagnosis and/or treatment. The Joint Commission on Accreditation of Healthcare Organizations requires the patient admission page document the type of insurance or payment structure that will cover the patient while being cared for at the hospital.

Coding Instructions

- Record the type of insurance reported on the patient's admission page.
- If the patient is diagnosed at the reporting facility, record the payer at the time of diagnosis.
- If the patient is diagnosed elsewhere or the payer at the time of diagnosis is not known, record the payer when the patient is initially admitted for treatment.
- Codes 21 and 65-68 are to be used for patients diagnosed on or after January 1, 2006.
- If more than one payer or insurance carrier is listed on the patient's admission page record the first.
- If the patient's payer or insurance carrier changes, do not change the code in the record.
- Use code 02 when the only information available is "self-pay."
- Use code 10 for prisoners when no further information is available.

Code	Label	Definition
01	Not insured	Patient has no insurance and is declared a charity write-off.
02	Not insured, self-pay	Patient has no insurance and is declared responsible for charges.
10	Insurance, NOS	Type of insurance unknown or other than the types listed in codes 20, 21, 31, 35, 60-68.
20	Private Insurance: Managed Care , HMO, or PPO	An organized system of prepaid care for a group of enrollees usually within a defined geographic area. Generally formed as one of four types: a group model, an independent physician association (IPA), a network, or a staff model. "Gate-keeper model" is another term for describing this type of insurance

Code	Label	Definition
21	Private Insurance: Fee-for-Service	An insurance plan that does not have a negotiated fee structure with the participating hospital. Type of insurance plan not coded as 20.
31	Medicaid	State government administered insurance for persons who are uninsured, below the poverty level, or covered under entitlement programs. Medicaid other than described in code 35.
35	Medicaid- Administered through a Managed Care plan	Patient is enrolled in Medicaid through a Managed Care program (e.g. HMO or PPO). The managed care plan pays for all incurred costs.
60	Medicare without supplement, Medicare, NOS	Federal government funded insurance for persons who are 62 years of age or older, or are chronically disabled (social security insurance eligible). Not described in codes 61, 62, 63
61	Medicare with supplement, NOS	Patient has Medicare and another type of unspecified insurance to pay costs not covered by Medicare.
62	Medicare- Administered through a Managed Care plan	Patient is enrolled in Medicare through a Managed Care plan (e.g. HMO or PPO). The Managed Care plan pays for all incurred costs.
63	Medicare with private supplement	Patient has Medicare and private insurance to pay costs not covered by Medicare.
64	Medicare with Medicaid eligibility	Federal government Medicare insurance with State Medicaid administered supplement.
65	TRICARE	Department of Defense program providing supplementary civilian-sector hospital and medical services beyond a military treatment facility to military dependents, retirees, and their dependents. Formally CHAMPUS (Civilian Health and Medical Program of the Uniformed Services).
66	Military	Military personnel or their dependents who are treated at a military facility.
67	Veterans Affairs	Veterans who are treated in Veterans Affairs facilities.
68	Indian/Public Health Service	Patient who receives care at an Indian Health Service facility or at another facility, and the medical costs are reimbursed by the Indian Health Service. Patient receives care at a Public Health Service facility or at another facility, and medical costs are reimbursed by the Public Health Service
99	Insurance status unknown	It is unknown from the patient's medical record whether or not the patient is insured.

Coding Examples:

Code	Reason
01	An indigent patient is admitted with no insurance coverage.
20	A patient is admitted for treatment and the patient admission page states the primary insurance carrier is an HMO
62	A 65-year old male patient is admitted for treatment and the patient admission page states the patient is covered by Medicare with additional insurance coverage from a PPO.

CANCER IDENTIFICATION

DATE OF DIAGNOSIS

Item Length: 8

NAACCR Item #390

NAACCR Name: Date of Diagnosis

XML NAACCR ID: dateOfDiagnosis

Definition:

Records the date of initial diagnosis by a recognized medical practitioner for the tumor being reported whether clinically or microscopically confirmed.

Coding Instructions:

- Code the month, day and year the tumor was first diagnosed, clinically or microscopically, by a recognized medical practitioner. When the first diagnosis includes reportable ambiguous terminology, record the date of that diagnosis.

Exception: Do not use the date of diagnosis from a cytology report using ambiguous terminology.

- When the only information available is a positive pathology or cytology report, code the date the biopsy was done, not the date the report was dictated or transcribed.
- The first diagnosis of cancer may be clinical (i.e., based on clinical findings or physician's documentation). Do not change the date of diagnosis when a clinical diagnosis is subsequently confirmed by positive histology or cytology.
- Positive tumor markers alone are not diagnostic of cancer. Use the date of clinical, histologic, or positive cytologic confirmation as the date of diagnosis.

Note: Positive tumor markers alone are never used for case ascertainment.

- Do not use cytology as a basis for diagnosis when ambiguous terms are used. Ambiguous cytology is not diagnostic of cancer. Use the date of clinical, histologic, or positive cytologic confirmation as the date of diagnosis.

Note 1: "Ambiguous" cytology means that the diagnosis is preceded by an ambiguous term such as apparently, appears, compatible with, etc.

Note 2: Do not use ambiguous cytology alone for case ascertainment.

- Code the earlier date as the date of diagnosis when
 1. A recognized medical practitioner says that, in retrospect, the patient had cancer at earlier date OR
 2. The original slides are reviewed and the pathologist documents that cancer was present. Code the date of the original procedure as the diagnosis date.

Note: Do not back-date the diagnosis when the information on the previous tumor is unclear and/or there is not review of previous slides and/or there is no physician’s statement that, in retrospect, the previous tumor was malignant.

- Use the date therapy was started as the date of diagnosis if the patient receives a first course of treatment before a definitive diagnosis.
- The date of death is the date of diagnosis for a Class of Case 38 or 49.
- Use the actual date of diagnosis as the Date of Initial Diagnosis for an *in utero* diagnosis, for cases diagnosed January 1, 2009 or later.
- If the year is unknown, it should be approximated. The month and day would be unknown.
- If the only information provided is “spring”, use April. Use July for “summer” or “mid-year” and October for “fall” or “autumn”. In winter, attempt to determine whether the diagnosis was “late in the year” (use December with the applicable year) or “early in year” (use January with the respective year).

Code	Definition
CCYYMMDD	The date of initial diagnosis is the month, day, and year that this primary cancer was first diagnosed by a recognized medical practitioner. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.

Coding Examples:

Code	Explanation
20050630	Key June 30, 2005
20110312	A mammogram on March 12, 2011 reveals a mass in the upper-outer quadrant of a patient’s right breast compatible with carcinoma. On March 20, 2011 the patient has an excisional breast biopsy that confirms infiltrating ductal carcinoma.
20110512	During a physical examination on May 12, 2011 the physician notes a prostate nodule that is suspicious for cancer. On June 15, 2011, an ultrasound guided needle biopsy of the prostate provides histologic confirmation of adenocarcinoma of the prostate.
201003	A patient has a total abdominal hysterectomy for endometriosis in March 2010. The patient is admitted to the hospital with abdominal pain and distention in November 2011. A laparoscopy with omental biopsy shows metastatic cystadenocarcinoma. Pathologists review the 2010 hysterectomy specimen. They identify an area of cystadenocarcinoma in the left ovary.

Code	Explanation
20210105	The patient has an elevated PSA and negative physical examination on 1/1/2021. The physician documents only that the patient is referred for a needle biopsy of the prostate. The biopsy on 1/5/2021 is positive for adenocarcinoma. The date of diagnosis is the date of the biopsy (do not code the date of the PSA or the date the procedure was dictated or transcribed).
20210101	The patient has an elevated PSA and negative physical examination on 1/1/2021. The physician documents that he/she suspects that the patient has prostatic cancer and is referring the patient for a needle biopsy. The biopsy on 1/5/2021 is positive for adenocarcinoma, confirming the physician's suspicion for cancer. The date of diagnosis is the date the physician documented that he/she suspects that the patient has prostatic cancer.
2010	Patient admitted to your facility June 7, 2011 for in-transit care for a lung cancer diagnosed in sometime in 2010.
20210130	Fetal intrahepatic mass consistent with hepatoblastoma diagnosed via ultrasound at 39 weeks gestation (01/30/2021). Live birth by C-section 02/04/2021. Code the date of diagnosis as 01/30/2021.
201110	If information is limited to the description "Fall," 2011

DATE OF DIAGNOSIS FLAG

Item Length: 2
NAACCR Item #391
NAACCR Name: Date of Diagnosis Flag
XML NAACCR ID: dateOfDiagnosisFlag

Definition:

This flag explains why no appropriate value is in the field Date of Diagnosis. Prior to 2010, date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions:

- If a valid date is coded in date of diagnosis, then leave this field blank. A valid date may be just a partial date.
- If date of diagnosis cannot be estimated or determined, then leave date of diagnosis blank and record 12 in this field indicating that the date is unknown.

Code	Definition
12	A proper value is applicable but cannot be determined or estimated (e.g., Date of diagnosis cannot be determined)
Blank	A valid date value is provided in item Date of Diagnosis.

Coding Examples:

Code	Explanation
Blank	Full date is known (CCYYMMDD) for Date of Diagnosis
Blank	Partial date is known (CCYYMM or CCYY) for Date of Diagnosis
12	Date is completely unknown for Date of Diagnosis

PRIMARY SITE

Item Length: 4
NAACCR Item #400
NAACCR Name: Primary Site
XML NAACCR ID: primarySite

Definition:

The primary site is defined as the organ or site in which the cancer originated or began. A metastatic site indicates that the primary (originating) tumor has spread from the original site to other areas in the body. Cancer registries only code the primary site in this field, using the ICD-O-2 (cases diagnosed prior to January 1, 2001) or ICD-O-3 (cases diagnosed on or after January 1, 2001) manual to determine the correct site code. The current Solid Tumor Rules (cases diagnosed January 1, 2018 and forward) contain additional coding instructions for some primary sites, including Head and Neck, Lung, and Urinary. Continue using ICD-O-3 beginning January 1, 2021 to assign topography codes. ICD-O-3.2 did not change any of the topography codes. Indications of metastatic sites are used in the registry for identifying the extent of the patient's disease and for staging purposes.

Coding Instructions:

- Record the ICD-O-3 topography code for the site of origin for cases diagnosed on or after January 1, 2001 and the ICD-O-2 topography code for cases diagnosed before January 1, 2001.
- Consult the attending physician to identify the primary site or the most definitive site code if the medical record does not contain that information.
- Topography codes are indicated by a “C” preceding the three-digit code number (do not record the decimal point).
- Unless otherwise instructed, use all available information, including pathology reports, scans, x-rays, MRIs, etc., to code the primary site.
- Unless otherwise instructed, use all available information, including pathology reports, scans, x-rays, MRIs, etc., to code the primary site.
- Use subcategory “.8” (overlapping lesion code) when a single tumor overlaps the boundaries of two or more categories or subcategories and its point of origin cannot be determined. Overlapping applies to sites that are contiguous (adjacent) to one another.
- Code the site of the invasive tumor when there is an invasive tumor and an in situ tumor in different subsites of the same anatomic.
- Code the last digit of the primary site to ‘9’ for single primaries, when multiple tumors arise in different subsites of the same anatomic site and the point of origin cannot be determined.

- Some histology/behavior terms in ICD-O-3.2 have a related site code in parenthesis; for example: Hepatoma (C22.0)
 1. Code the site as documented in the medical record and ignore the suggested ICD-O-3.2 code when a primary site is specified in the medical record.
 2. Use the site code suggested by ICD-O-3.2 when the primary site is the same as the site code suggested or the primary site is unknown.
- Code the primary site, not the metastatic site. If a tumor is metastatic and the primary site is unknown, code the primary site as unknown (C809).
- Code C42.2 (Spleen) as the primary site for angiosarcoma of spleen with mets to bone marrow.
- Code C50_ (breast) for angiosarcoma of breast. Although angiosarcoma actually originates in the lining of the blood vessels, an angiosarcoma originating in the breast has a poorer prognosis than many other breast tumors.
- Gastrointestinal Stromal Tumor (GIST): Code the primary site to the location where the malignant GIST originated.
- Code to Skin, NOS (C44.9) if a patient is diagnosed with metastatic melanoma and the primary site is unidentified.
- Transplanted organs or tissue may originate from organs or tissue from the patient's own body (called autograft) or another human donor (homograft or allograft). Report a new primary in the transplanted organ as you would any new primary, applying the current Solid Tumor Rules. Code the primary site to the location of the transplanted organ, i.e., code the malignancy where it resides/lies.
- In the absence of any additional information about the primary site, assign the codes listed for these primary sites/histologies.

Primary Site/Histology	Topography Code
Ampullary/peri-ampullary	C241
Anal margin	C445
Anal verge	C211
Angle of the stomach	C162
Angular incisura of stomach	C163
Book-leaf lesion (mouth)	C445
Colored/lipstick portion of upper lip	C000
Cutaneous leiomyosarcoma	C44
Distal conus	C720
Edge of tongue	C021

Primary Site/Histology	Topography Code
Frontoparietal (brain)	C718
Gastric angular notch (incisura)	C163
Gastrohepatic ligament	C481
Genu of pancreas	C250
Glossotonsillar sulcus	C109
Incisura, incisura angularis	C163
Infrahilar area of lung	C349
Interhemispheric fissure (cerebrum)	C710
Lateral tongue	C023
Leptomeninges	C709
Masticator space	C760
Melanoma, NOS	C449
Nail bed, thumb	C446
Pancreatobiliary	C269
Parapharyngeal space	C490
Perihilar bile duct	C240
Testis, descended post orchiopexy	C621
Uncinate of pancreas	C250

- Complete primary site coding rules are described in the ICD-O-2 manual under the heading “Topography,” pages xx-xxiii, and in the ICD-O-3 manual under *Coding Guidelines for Topography and Morphology*.

Note: Kaposi’s Sarcoma is coded to the site in which it originates. If the Kaposi’s sarcoma is present in the skin and another site simultaneously, code to the specified skin site, (C44._). If the primary site is unknown or cannot be determined, code skin, NOS (C44.9).

Note: The majority of sarcomas arise in mesenchymal or connective tissues that are located in the musculoskeletal system, which includes the fat, muscles, blood vessels, deep skin tissues, nerves, bones, and cartilage. The default code for sarcomas of unknown primary site is C49.9 rather than C80.9. Sarcomas may also arise in the walls of hollow organs and in the viscera covering an organ. Code the primary site to the organ of origin.

Specific Tissue with Ill-Defined Sites

If any of the following histologies appears only with an ill-defined site description (e.g., “abdominal” or “arm”), code it to the tissue in which such tumors arise rather than the ill-defined region (C76._) of the body, which contains multiple tissues.

Histology	Description	Code to This Site
8720-8790	Melanoma	C44._, Skin
8800-8811, 8813-8830, 8840-8921, 9040-9044	Sarcoma except periosteal fibrosarcoma and dermatofibrosarcoma	C49._, Connective, Subcutaneous and Other Soft Tissues
8990-8991	Mesenchymoma	C49._, Connective, Subcutaneous and Other Soft Tissues
9120-9170	Blood vessel tumors, lymphatic vessel tumors	C49._, Connective, Subcutaneous and Other Soft Tissues
9580-9582	Granular cell tumor and alveolar soft part sarcoma	C49._, Connective, Subcutaneous and Other Soft Tissues
9240-9252	Mesenchymal chondrosarcoma and giant cell tumors	C40._, C41._, for Bone and Cartilage C49._, Connective Subcutaneous and Other Soft Tissues
8940-8941	Mixed tumor, salivary gland type	C07._ for Parotid Gland C08._ for Other and Unspecified Major Salivary Glands

- When the medical record does not contain enough information to assign a primary site
 - a. Consult a physician advisor to assign the site code. Document your discussion in the text.
 - b. Use the NOS category for the organ system or the Ill-Defined Sites (C760-C768) if the physician advisor cannot identify a primary site.

Occult Tumors of the Head and Neck (Cases diagnosed January, 1, 2018 and later)

Assign primary site C119 (nasopharynx) for occult head and neck tumors with cervical lymph node metastasis in Levels I-VII, and other group lymph nodes positive for Epstein-Barr virus (EBV+) (regardless of p16 status) encoded small RNAs (EBER) identified by in situ hybridization.

Assign primary site C109 (oropharynx) for occult head and neck tumors with cervical lymph node metastasis in Levels I-VII, and other group lymph nodes, p16 positive with histology consistent with HPV-mediated oropharyngeal carcinoma (OPC).

Assign C760 for Occult Head and Neck primaries with positive cervical lymph nodes. Schema Discriminator 1: Occult Head and Neck Lymph Nodes is used to discriminate between these cases and other uses of C760.

- c. Assign the NOS code for the body system when there are two or more possible primary sites documented and all are within the same system.

Example: Two possible sites are documented in the GI system such as colon and small intestine; code to the GI tract, NOS (C269). Document the possible primary sites in a text field.

- d. Code unknown primary site when there is a physician statement of unknown primary site ONLY when none of the above instructions can be applied.
- e. Code Unknown Primary Site (C809) if there is not enough information to assign an NOS or Ill-Defined Site category.

Coding Instructions for Hematopoietic and Lymphoid Neoplasm (9590/3-9992/3)

For cases diagnosed January 1, 2010 and later, see the *Hematopoietic and Lymphoid Neoplasm Coding Manual and Database* (<http://seer.cancer.gov/tools/heme/>) for instructions on coding the primary site for hematopoietic and lymphoid neoplasms.

For hematopoietic and lymphoid neoplasms diagnosed prior to January 1, 2010, use the following rules for coding primary site:

Code all leukemias except myeloid sarcoma (9930/3) to the bone marrow C42.1. Myeloid sarcoma is coded to the site of origin.

Primary Site Coding Lymphoma (Cases diagnosed prior to January 1, 2010)

Use the following guidelines to determine the primary site(s) for malignant lymphomas:

- Code lymphomas arising in lymphatic tissue or nodes to the site of origin. The lymphatic sites are Lymph Node(s) C77.~, Tonsil C09.~, Spleen C42.2, Waldeyer's Ring C14.2, and Thymus C37.9.
- Code extralymphatic lymphomas (lymphatic cells in nonlymphatic organs such as intestine or stomach) to the organ of origin (Intestine C26.0, Stomach C16.0-C16.9).
- If extranodal/extralymphatic site is suspected but is unknown, code C80.9.
- Code mycosis fungoides and cutaneous lymphomas to Skin (C44.~).

- Code to Lymph Nodes, NOS (C77.9) when:
 1. the site of origin is not identified for a lymphoma
 2. a patient has diffuse lymphoma and a primary site is unknown or not specified
 3. a lymphoma mass is identified as “retroperitoneal,” “inguinal,” “mediastinal,” or “mesentery,” and no specific information is available to indicate what tissue is involved
 4. bone marrow metastases are present and the primary site of a lymphoma is unknown or not specified

- If origin of a lymphoma is unknown but is suggested by the histology code in ICD-O-3, code to the suggested site. Example: 9689/3 Splenic marginal zone B-cell lymphoma (C42.2)

- Code to Lymph Nodes, Multiple Regions (C77.8) when multiple lymph node chains are involved with disease.

Note: Carefully identify the origin of the tumor. Do not code the biopsy site or a metastatic site as the primary site. Lymphoma may be present in both an extranodal/extralymphatic organ and one or more lymph node chains. Code the primary site as the extranodal/extralymphatic organ or the lymph nodes as directed by the managing physician or physician advisor

Examples:

Code	Explanation
C508	Overlapping lesion of breast. Code overlapping lesion when a tumor involves both the upper inner quadrant and lower inner quadrant of the left breast.
C504	Patient has an invasive breast tumor in the upper-outer quadrant of the left breast and in situ tumors in multiple quadrants of the left breast. Code the primary site to C50.4 (upper outer quadrant of breast)
C503	The patient has a 2cm tumor in the right breast. The tumor originated in the lower inner quadrant and extends into the upper inner quadrant. Code the primary site to the lower inner quadrant of the breast (C50.3).
C679	Bladder, NOS. Use subcategory 9 when multiple lesions arise in both the bladder trigone (C67.0) and lateral wall (C67.2).
C700	Code C70.0 (Cerebral Meninges) when patient is diagnosed with a meningioma of the frontal lobe of the brain.
C269	Two possible sites are documented in the GI system such as colon and small intestine; code to the GI tract, NOS (C269). Document the possible primary sites in a text field.

Code	Explanation
C159	Diagnosis of malignancy in transplanted section of colon serving as esophagus. Code the primary site as esophagus. Document the situation in a text field.
C509	An excision of the right axillary nodes reveals metastatic infiltrating duct carcinoma. The right breast is negative. The ICD-O-3 shows infiltrating duct carcinoma (8500) with a suggested site of breast (C50.).
C250	The pathology report says “infiltrating duct carcinoma of the head of the pancreas.” The listing in ICD-O-3 is infiltrating duct carcinoma 8500/3 (C50.). Code the primary site to head of pancreas (C25.0), not to breast (C50.) as suggested by the ICD-O-3.

LATERALITY

Item Length: 1
NAACCR Item #410
NAACCR Name: Laterality
XML NAACCR ID: laterality

Definition:

Identifies the side of a paired organ or the side of the body on which the reportable tumor originated. This applies to the primary site only.

Coding Instructions:

- Always code unknown primary site (C80.9), and lymphoma to 0 (not paired).
- Laterality must be coded 1-5 or 9 for all sites listed in the table below. Laterality may be assigned for sites not listed in the table or tumors in those sites may be assigned a laterality of 0.
- Code the side where the primary tumor originated. If it is known the a primary tumor is confined to one side a paired organ, but that side is unknown, use code 3.
- Code 4 would rarely be used except when both ovaries are involved simultaneously with a single histology or epithelial histologies (8000-8799), there are diffuse bilateral lung nodules, there are bilateral retinoblastomas or bilateral Wilms tumors.
- Code 5, midline lesion, where the right and left sides of a paired site are contiguous and the tumor is at the intersection of the right and left side. The only sites for which code 5 would be used are C700, C710-C714, C722-C725, C443, C445.
- Code 9 should be used only when the laterality is unknown and there is no information that the tumor is confined to one side of a paired organ.

The following is a list of paired organs:

Site Code	Definition
C07.9	Parotid gland
C08.0	Submandibular gland
C08.1	Sublingual gland
C09.0	Tonsillar fossa
C09.1	Tonsillar pillar
C09.8	Tonsil, overlapping site
C09.9	Tonsil, NOS
C30.0	Nasal cavity (excluding nasal cartilage and nasal septum)
C30.1	Middle ear
C31.0	Maxillary sinus
C31.2	Frontal sinus
C34.0	Main bronchus (excluding carina)
C34.1—C34.9	Lung
C38.4	Pleura
C40.0	Long bones of upper limb and scapula
C40.1	Short bones of upper limb
C40.2	Long bones of lower limb
C40.3	Short bones of lower limb
C41.3	Rib and clavicle (excluding sternum)
C41.4	Pelvic bones (excluding sacrum, coccyx, and symphysis pubis)
C44.1	Skin of eyelid
C44.2	Skin of external ear
C44.3	Skin of face (midline code 9)
C44.5	Skin of trunk (midline code 9)
C44.6	Skin of upper limb and shoulder
C47.1	Peripheral nerves and autonomic nervous system of upper limb and shoulder
C47.2	Peripheral nerves and autonomic nervous system of lower limb and hip
C49.1	Connective, subcutaneous, and other soft tissues of upper limb and shoulder
C49.2	Connective, subcutaneous, and other soft tissues of lower limb and hip
C50.0- C50.9	Breast
C56.9	Ovary
C57.0	Fallopian tube
C62.0- C62.9	Testis
C63.0	Epididymis
C63.1	Spermatic cord
C64.9	Kidney, NOS

Site Code	Definition
C65.9	Renal pelvis
C66.9	Ureter
C69.0- C69.9	Eye and lacrimal gland
C70.0	Cerebral meninges, NOS (excluding diagnoses prior to 2004)
C71.0	Cerebrum (excluding diagnoses prior to 2004)
C71.1	Frontal Lobe (excluding diagnoses prior to 2004)
C71.2	Temporal Lobe (excluding diagnoses prior to 2004)
C71.3	Parietal Lobe (excluding diagnoses prior to 2004)
C71.4	Occipital Lobe(excluding diagnoses prior to 2004)
C72.2	Olfactory Nerve (excluding diagnoses prior to 2004)
C72.3	Optic Nerve (excluding diagnoses prior to 2004)
C72.4	Acoustic Nerve (excluding diagnoses prior to 2004)
C72.5	Cranial Nerve, NOS(excluding diagnoses prior to 2004)
C74.0- C74.9	Adrenal gland
C75.4	Carotid body

Code	Definition
0	Organ is not considered to be a paired site.
1	Origin of primary is right.
2	Origin of primary is left.
3	Only one side involved, right or left origin not specified.
4	Bilateral involvement, side of origin unknown, stated to be a single primary. This includes: <ul style="list-style-type: none"> ▪ Both ovaries simultaneously involved with a single histology ▪ Bilateral retinoblastoma ▪ Bilateral Wilms tumors
5	Paired site: midline tumor
9	Paired site, but lateral origin unknown

HISTOLOGIC TYPE ICD-O-3

Item Length: 4
NAACCR Item #522
NAACCR Name: Histologic Type ICD-O-3
XML NAACCR ID: histologicTypeIcdO3

Definition:

Histologic type refers to the *classification* of malignancy described in the pathology or cytology report.

Coding Instructions:

- Record histology using the ICD-O-3 codes (<https://seer.cancer.gov/icd-o-3/>) in the Numeric Lists/Morphology section (ICD-O-3, pp. 69–104) and in the Alphabetic Index (ICD-O-3, pp. 105–218).
- ICD-O-3 identifies the morphology codes with an “M” preceding the code number. Do not record the “M”.
- For cases diagnosed prior to January 1, 2007, follow the coding rules outlined on pages 20 through 40 of ICD-O-3. More specific rules are provided below.
- For cases diagnosed January 1, 2007 to December 31, 2017, use the 2007 *Multiple Primary and Histology Coding Rules* (<https://seer.cancer.gov/tools/mphrules/download.html>)
- For cases diagnosed January 1, 2018 and later, use the 2018 Solid Tumor Rules (<https://seer.cancer.gov/tools/solidtumor/>). Follow documentation in priority order
 1. Updated ICD-O histology codes and terms (https://www.naacr.org/wp-content/uploads/2020/11/Histology3_v21_FINAL_11_10_2020.xlsx)
 2. The current Solid Tumor Rules
 3. The ICD-O-3 manual

Site-specific rules cover the following

Primary Site	Topography Codes
Head and Neck	C000-C148, C300-C329, C410, C411, C442
Colon, Rectosigmoid, Rectum	C180-C189, C199, C209
Lung	C340-C349
Cutaneous Melanoma	C440-C449 with Histology 8720-8780
Breast	C500-C506, C508-C509
Kidney	C649
Urinary Sites	C659, C669, C670-C679, C680-C681, C688-C689
Non-malignant CNS	C700, C701, C709, C710-C719, C720-C725, C728, C729, C751-C753
Malignant CNS and Peripheral Nerves	C470-C479, C700, C701, C709, C710-C719, C720-C725, C728, C729, C751-C753
Other Sites	Excludes Head and Neck, Colon, Rectosigmoid, Rectum, Lung, Cutaneous Melanoma, Breast, Kidney, Urinary Sites, Peripheral Nerves, CNS

The General rules do not apply to hematopoietic primaries (lymphoma and leukemia) of any site or to the reportable benign or borderline intracranial or CNS tumors. The head and neck, colon, rectosigmoid and rectum, breast, kidney, urinary sites, and malignant CNS and peripheral nerves rules exclude lymphoma and leukemia (M9590-M9992) and Kaposi sarcoma (M9140). All other sites rules exclude lymphoma and leukemia (M9590-M9992).

- Carefully review all pathology reports.
- Code the final pathologic diagnosis for solid tumors.

Exception:

For cases diagnosed **prior to January 1, 2007**, if the final diagnosis is “Not Otherwise Specified” (carcinoma, NOS; melanoma, NOS; sarcoma, NOS; lymphoma, NOS; or malignant tumor, NOS), then code the histology from the microscopic description if it identifies a more specific histologic type (higher ICD-O-3 code) such as adenocarcinoma, amelanotic melanoma, spindle cell sarcoma.

- The codes for cancer, NOS (8000) and carcinoma, NOS (8010) **are not interchangeable**. If the physician says that the patient has carcinoma, then code carcinoma, NOS (8010).
- For leukemias, lymphomas and other hematopoietic diseases, follow the instructions in the *Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* and the Hematopoietic and Lymphoid Neoplasms Database (<https://seer.cancer.gov/tools/heme/>).

2021 ICD-O-3.2

For 2021, standard setters have agreed to implement new histology terms and codes for ICD-O-3 based on the current versions of the *World Health Organization Classification of Tumors*. The update, referred to as ICD-O-3.2 includes comprehensive tables listing histology codes and behavior codes in effect beginning with cases diagnosed 1/1/2021 and later. The new codes, new terms, and codes with changes to behavior are available at <https://www.naaccr.org/icdo3/>

2018 ICD-O-3 Update

There are new codes, changes in behavior codes, and new terms associated with current codes for all cases diagnosed January 1, 2018 and later. These changes reflect updates to the WHO Classifications for Tumors (Blue Books). The new codes, new terms, and codes with changes to behavior are available at: <https://www.naaccr.org/2018-implementation>.

ICD-O-3.1

The International Classification of Diseases for Oncology, Third Edition, First Revision has not been approved for use in the United States. This revision was published in 2013. It includes codes and terms which are not approved for use at this time.

2014 ICD-O-3 Update

See the *NAACCR Guidelines for ICD-O-3 Update Implementation* pages five and six <https://www.naaccr.org/wp-content/uploads/2019/12/ICD-O-3-Implementation-Guide-FINAL.pdf>. The terms and synonyms for existing ICD-O-3 histology codes listed in the document should be incorporated into your ICD-O-3 manual.

2015 and 2016 ICD-O-3 Update

Effective for 2015 diagnoses, code 8240/1 for Carcinoid tumor, NOS of appendix (C181) is obsolete. Code Carcinoid tumor, NOS of appendix to 8240/3 as this is now reportable (behavior code 3) in 2015.

Effective for 2015 diagnoses, one reportable histology codes is obsolete
8157/3 Enteroglucagonoma, malignant

Use histology code 8152/3 for Enteroglucagonoma, malignant as Enteroglucagonoma is now a related term for Glucagonoma.

New histology terms and codes have been introduced for ICD-O-3, but many cannot be used for 2015 and 2016 diagnoses because they are not included among the acceptable histology codes for the Collaborative Stage algorithms. See the *NAACCR Guidelines for ICD-O-3 Update Implementation* page 7 for new terms and codes. https://www.naacr.org/wp-content/uploads/2016/11/2015_Implementation_Guidelines_and_Recommendations_20150505.pdf

SOLID TUMORS DIAGNOSED BEFORE 2007; HEMATOPOIETIC DISEASES DIAGNOSED PRIOR TO 2010

Histology Coding Rules for a Single Tumor (Solid tumors diagnosed prior to January 1, 2007 and Hematopoietic and Lymphoid neoplasms diagnosed prior to January 1, 2010).

Source: *Seer Program and Coding and Staging Manual*

- The rules are in hierarchical order. Rule 1 has the highest priority.
 - Use the rules in priority order.
 - Use the first rule that applies to the case. (Do not apply any additional rules.)
1. Code the histology if only one type is mentioned in the pathology report.
 2. Code the **invasive** histology when both invasive and in situ tumors are present.

Example: Pathology report reads infiltrating ductal carcinoma and cribriform ductal carcinoma in situ. Code the invasive histology 8500/3.

Exception: If the histology of the invasive component is carcinoma, NOS, adenocarcinoma, NOS, melanoma, NOS, or sarcoma, NOS, then code the histology of the specific term associated with the in situ component and an invasive behavior code.

3. Use a **mixed** histology code if one exists.

Example: 9085 Mixed germ cell tumor

4. Use a **combination** histology code if one exists.

Example: 8255 Renal cell carcinoma, mixed clear cell and chromophobe types

5. Code the **more specific term** when one of the terms is ‘NOS’ and the other is a more specific description of the same histology.

Example 1: Pathology report reads poorly differentiated carcinoma, probably squamous in origin. Code the histology as squamous cell carcinoma rather than the non-specific term “carcinoma.”

Example 2: The pathology report from a nephrectomy reads renal cell carcinoma (8312) (renal cell identifies the affected organ system rather than the histologic cell type) in one portion of the report and clear cell carcinoma (8310) (a histologic cell type) in another section of the report. Code clear cell carcinoma (8310); renal cell carcinoma (8312) refers to the renal system rather than the cell type, so renal cell is the less specific code.

6. Code the **majority** of tumor.
 - a. Based on the pathology report description of the tumor.
 - b. Based on the use of majority terms.

Majority of Tumor:

Terms that mean the majority of tumor	Terms that DO NOT mean the majority of tumor
Predominantly	With foci of
With features of	Focus of/focal
Major	Areas of
Type (Effective 1/1/1999)	Elements of
With...Differentiation (Effective 1/1/1999)	Component (Effective 1/1/1999)

7. Code the **numerically higher** ICD-O-3 code. This is the rule with the lowest priority and should be used infrequently.

Histology Coding Rules for Multiple Tumors with Different Behaviors in the Same Organ Reported as a Single Primary (Cases diagnosed prior to January 1, 2007)

Source: *SEER Program Coding and Staging Manual*

1. Code the histology of the invasive tumor when one lesion is in situ (/2) and the other is invasive (/3).

Example: At mastectomy for removal of a 2 cm invasive ductal carcinoma, an additional 5 cm area of intraductal carcinoma was noted. Code histology and behavior as invasive ductal carcinoma (8500/3).

Histology Coding Rules for Multiple Tumors in Same Organ Reported as a Single Primary (Cases diagnosed prior to January 1, 2007)

Source: *SEER Program Coding and Staging Manual*

1. Code the histology when multiple tumors have the same histology.
2. Code the histology to adenocarcinoma (8140/_; in situ or invasive) when there is an adenocarcinoma and an adenocarcinoma in a polyp (8210/_, 8261/_, 8263/_) in the same segment of the colon or rectum.
3. Code the histology to carcinoma (8010/_; in situ or invasive) when there is a carcinoma and a carcinoma in a polyp (8210/_) in the same segment of the colon or rectum.
4. Use a **combination** code for the following:
 - a. Bladder: Papillary and urothelial (transitional cell) carcinoma (8130)
 - b. Breast: Paget Disease and duct carcinoma (8541)
 - c. Breast: Duct carcinoma and lobular carcinoma (8522)
 - d. Thyroid: Follicular and papillary carcinoma (8340)
5. Code the more specific term when one of the terms is 'NOS' and the other is a more specific description of the same histology.
6. Code all other multiple tumors with different histologies as multiple primaries.

BEHAVIOR CODE ICD-O-3

Item Length: 1
NAACCR Item #523
NAACCR Name: Behavior Code ICD-O-3
XML NAACCR ID: behaviorCodeIcdO3

Definition:

The behavior code occupies the 5th space (digit) of the morphology code. This component of the histologic code indicates the way in which the neoplasm will act or behave – invasive (3), non-invasive (2), borderline malignancy (1), benign (0).

Coding Instructions:

Intracranial and CNS tumors

Intracranial and CNS tumors with behavior codes 0 (benign) and 1 (borderline malignancy) are reportable beginning with January 1, 2004 diagnoses.

Code the behavior from CT scan Magnetic Resonance Imaging (MRI), or Positron Emission Tomography (PET) report when there is no tissue diagnosis (pathology or cytology report). Code the behavior listed on the scan. Do not use the WHO grade to code behavior.

Metastatic or Non-primary Sites

Cases reported to the MCR cannot have a metastatic (/6) behavior code. If the only pathologic specimen is from a metastatic site, code the appropriate histology code and the malignant behavior code (/3). The primary site and its metastatic site(s) have the same histology.

Code the behavior as malignant (/3) when malignant metastasis is present. Metastasis could be regional, nodal, or distant.

Exception: For in situ breast cancer; code as non-invasive (/2) in the presence of isolated tumor cells or if cells are artifactually displaced from previous procedure.

In Situ

Clinical evidence alone cannot identify the behavior as in situ; a behavior code of /2 (in situ) must be based on pathologic examination.

In Situ and Invasive

Code 3 if any invasion is present, no matter how limited, i.e., microinvasion.

Recode the behavior as malignant (Code 3) when metastases are attributed to a tumor originally thought to be in situ.

If an invasive behavior is ambiguous on a biopsy and the subsequent surgery shows only in situ cancer, code the behavior as in situ (/2).

ICD-O-3.2 Histology/Behavior Code Listing

Behavior is the fifth digit of the morphology code after the slash (/). The standard reference for coding behavior is the ICD-O-3.2. Pages 27 through 30 in ICD-O-3 discuss behavior. The following general rules are found on pages 29-30 in ICD-O-3.

- Usually a histologic term carries a clear indication of the likely behavior of the tumor, whether malignant or benign, and this is reflected in the behavior code assigned to it in the ICD-O.
- Although only a few histologic types of in situ neoplasms are actually listed in the ICD-O, the behavior code /2 could be attached to any histology code if an in situ form of the neoplasm is diagnosed.
- If the pathologist disagrees with the ICD-O behavior assignment in a particular case, code the behavior according to the pathologist's description of the behavior even if that histology/behavior combination is not listed in the ICD-O.

The pathologist has the final say on the behavior of the tumor. ICD-O-3 may have only one behavior code, in situ (/2) or malignant (/3), listed for a specific histology. If the pathology report describes the histology as in situ and the ICD-O-3 histology code is listed only with a malignant behavior code (/3), assign the in situ behavior code (/2). If the pathology report describes histology as malignant and the ICD-O-3 histology code is listed only with an in situ behavior code (/2), assign the malignant behavior code (/3). See the Morphology and Behavior Code Matrix discussion on page 29 in ICD-O-3.

- Example: The pathology report says large cell carcinoma in situ. ICD-O-3 lists large cell carcinoma only with a malignant behavior (8013/3). Code the histology and behavior as 8013/2 as stated by the pathologist.

Note: The ICD-O-3 behavior code for juvenile astrocytoma (9421/1) is coded as 3. Gastrointestinal stromal tumors (GIST) and thymomas are often non-malignant. However, they must be abstracted and coded with a behavior code 3 if they have multiple foci, positive lymph nodes, or metastasis.

Code	Label	Definition
0	Benign	Benign
1	Borderline	Borderline. Uncertain whether benign or malignant. Low malignant potential Uncertain malignant potential.
2	In situ and/or carcinoma in situ Synonyms with in situ	Behavior code '2' Bowen disease (not reportable for C440-C449) Clark level I for melanoma (limited to epithelium) Confined to epithelium Hutchinson melanotic freckle, NOS (C44_) Intracystic, noninfiltrating (carcinoma) Intraductal (carcinoma) Intraepidermal, NOS (carcinoma) Intraepithelial neoplasia, Grade III (e.g., AIN III, LIN III, SIN III, VAIN III, VIN III) Intraepithelial, NOS (carcinoma) Involvement up to, but not including the basement membrane Lentigo maligna (C44_) Lobular, noninfiltrating (C50_) (carcinoma) Noninfiltrating (carcinoma) Non-invasive (carcinoma) No stromal invasion/involvement Papillary, noninfiltrating or intraductal (carcinoma) Precancerous melanosis (C44_) Queyrat erythroplasia (C60_) Stage 0 (except Paget's disease (8540/3) of breast and colon or rectal tumors confined to the lamina propria)
3		Invasive or microinvasive.

Coding Examples:

Code	Explanation
3	The pathology report stated intraductal carcinoma (8500/2) with focal areas of invasion. Code the behavior code to the invasive component, infiltrating ductal carcinoma (8500/3).
3	The pathology report stated large in situ intraductal carcinoma (8500/2) with an area of microinvasion. Code the behavior to invasive even if only microinvasion.
2	The pathology report says large cell carcinoma in situ. ICD-O-3 lists large cell carcinoma only with a malignant behavior (8013/3). Code the histology and behavior as 8013/2 as stated by the pathologist.
3	Right colon biopsy reveals tubulovillous adenoma with microfocal carcinoma in situ; right hemicolectomy is negative for residual disease. Later core liver biopsy consistent with metastatic adenocarcinoma of gastrointestinal origin. Oncologist states most likely colon primary. Change the behavior code for the colon primary from /2 to /3. There were no other colon primaries in this case.

Code	Explanation
2	Needle biopsy of a breast tumor shows ductal carcinoma in situ with a focus suspicious for invasion. Subsequent lumpectomy showed ductal carcinoma in situ. Code the ductal carcinoma in situ (8500/2)

GRADE CLINICAL

Item Length: 1
NAACCR Item #3843
NAACCR Name: Grade Clinical
XML NAACCR ID: gradeClinical

Definition:

Grade Clinical is new in 2018. This data item records the grade of a solid primary tumor before any treatment (surgical resection or initiation of any treatment including neoadjuvant).

Coding Instructions:

For cases diagnosed January 1, 2018 and later, this data item, along with Grade Pathological and Grade Post-Therapy Path (yp), replaces the data item Grade [NAACCR Item #440] as well as site specific factors for cancer sites with alternative grading systems (e.g., breast [Bloom-Richardson], prostate [Gleason]). The data item Grade Post Therapy Clin (yc) was added in 2021 as another grade code.

This data item cannot be left blank for cases diagnosed January 1, 2018 and later.

Refer to the most recent version of the *Grade Coding Instructions and Tables* (<https://www.naacccr.org/SSDI/Grade-Manual.pdf>) for additional site-specific instructions.

GRADE PATHOLOGICAL

Item Length: 1
NAACCR Item #3844
NAACCR Name: Grade Pathological
XML NAACCR ID: gradePathological

Definition:

Grade Pathological is new in 2018. This data item records the grade of a solid primary tumor that has been resected and for which no neoadjuvant therapy was administered.

Coding Instructions:

For cases diagnosed January 1, 2018 and later, this data item, along with Grade Clinical and Grade Post-Therapy Path (yc), replaces the data item Grade [NAACCR Item #440] as well as site specific factors for cancer sites with alternative grading systems (e.g., breast [Bloom-Richardson], prostate [Gleason]). The data item Grade Post Therapy Clin (yc) was added in 2021 as another grade code.

If AJCC staging is being assigned, the tumor must have met the surgical resection requirements in the AJCC manual. This may include the grade from the clinical workup. Since all clinical information is used in pathological staging, record the highest grade documented from any microscopic specimen of the primary site whether from the clinical workup or the surgical resection.

This data item cannot be left blank for cases diagnosed January 1, 2018 and later.

Refer to the most recent version of the *Grade Coding Instructions and Tables* (<https://www.naacccr.org/SSDI/Grade-Manual.pdf>) for additional site-specific instructions.

GRADE POST THERAPY (yc)

Item Length: 1

NAACCR Item #1068

NAACCR Name: Grade Post Therapy Clin (yc)

XML NAACCR ID: gradePostTherapyClin

Grade, Post Therapy Clin (yc), effective 01/01/2021, records the grade of a solid primary tumor that has been microscopically sampled following neoadjuvant therapy or primary systemic/radiation.

Refer to the most recent version of the Grade Coding Instructions and Tables (<https://www.naacr.org/SSDI/Grade-Manual.pdf>) for additional site-specific instructions.

GRADE POST THERAPY (yp)

Item Length: 1

NAACCR Item #3845

NAACCR Name: Grade Post Therapy Path (yp)

XML NAACCR ID: gradePostTherapy

Definition:

Grade, Post Therapy Path (yp), effective 01/01/2018, records the grade of a solid primary tumor that has been resected following neoadjuvant therapy or primary systemic/radiation therapy. If AJCC staging is being assigned, the tumor must have met the surgical resection requirements in the AJCC manual. For some sites, grade is required to assign the post-neoadjuvant stage group. The name was updated from Grade Post Therapy to Grade Post Therapy Path (yp) in 2021.

Coding Instructions:

For cases diagnosed January 1, 2018 and later, this data item, along with Grade Clinical and Grade Pathological, replaces the data item Grade [NAACCR Item #440] as well as site specific factors for cancer sites with alternative grading systems (e.g., breast [Bloom-Richardson], prostate [Gleason]). The data item Grade Post Therapy Clin (yc) was added in 2021 as another grade code.

If AJCC staging is being assigned, the tumor must have met the surgical resection requirements in the AJCC manual. Record the highest grade documented from the surgical treatment resection specimen of the primary site following neoadjuvant therapy.

Refer to the most recent version of the *Grade Coding Instructions and Tables* (<https://www.naacr.org/SSDI/Grade-Manual.pdf>) for additional site-specific instructions.

AJCC ID

Item Length: 1
NAACCR Item #995
NAACCR Name: AJCC ID
XML NAACCR ID: acjId

Definition:

For 2018, AJCC ID is used to link AJCC staging eligible sites/histologies with the appropriate AJCC chapter and staging algorithm.

Coding Instructions:

AJCC ID will be derived by registry software based on the site and histology codes entered by the registrar.

Refer to the SSDI Manual Appendix A (<https://www.naacccr.org/wp-content/uploads/2020/08/SSDI-Manual.Appendix-A.pdf?v=1613490814>) for crosswalks for sites/histology, AJCC ID and Schema ID.

SCHEMA ID

Item Length: 1
NAACCR Item #3800
NAACCR Name: Schema ID
XML NAACCR ID: schemaId

Definition:

For 2018, Schema ID is used to link all combinations of sites and histologies with the appropriate stage data collection systems and site-specific data items.

Coding Instructions:

Schema ID will be derived by registry software based on the site and histology codes entered by the registrar.

Refer to the SSDI Manual Appendix A (<https://www.naacr.org/wp-content/uploads/2020/08/SSDI-Manual.Appendix-A.pdf?v=1613490814>) for crosswalks for sites/histology, AJCC ID and Schema ID.

SCHEMA DISCRIMINATOR 1

Item Length: 1

NAACCR Item #3926

NAACCR Name: Schema Discriminator 1

XML NAACCR ID: schemaDiscriminator1

Definition:

Introduced in CSV2, schema discriminators are used when the primary site and/or histology are not sufficient to identify the correct AJCC staging algorithm. Schema discriminators are used to define both AJCC ID and Schema ID.

Coding Instructions:

Refer to the SSDI Manual (<https://apps.naacr.org/ssdi/list/2.0>) for crosswalks for the schema discriminators and for codes and coding instructions.

SCHEMA DISCRIMINATOR 2

Item Length: 1

NAACCR Item #3927

NAACCR Name: Schema Discriminator 2

XML NAACCR ID: schemaDiscriminator2

Definition:

Introduced in CSv2, schema discriminators are used when the primary site and/or histology are not sufficient to identify the correct AJCC staging algorithm. Schema discriminators are used to define both AJCC ID and Schema ID.

Coding Instructions:

Refer to the SSDI Manual (<https://apps.naaccr.org/ssdi/list/2.0>) for crosswalks for the schema discriminators and for codes and coding instructions.

LYMPH-VASCULAR INVASION

Item Length: 1

NAACCR Item #1182

NAACCR Name: Lymphovascular Invasion

XML NAACCR ID: lymphVascularInvasion

Definition:

Lymph-vascular invasion is defined as the presence of tumor cells found inside small blood vessels or lymphatic channels within the tumor and surrounding tissues in the primary site. The tumor cells have broken free of the primary tumor and now have the capability to float throughout the body. Other names for lymph-vascular invasion are LVI, lymphovascular invasion, vascular invasion, blood vessel invasion, and lymphatic invasion. Vascular invasion is not the same as direct tumor extension from the primary tumor into adjacent blood vessels; LVI cells are not attached to or growing into the wall of the blood vessel. Lymphatic invasion is not the same as involvement of regional lymph nodes. Lymph-vascular invasion does not include perineural invasion.

Coding Instructions:

Revised CAP Protocols and 8th Edition chapters will indicate which chapters will use the new codes (2, 3, and 4) and which will only use the existing codes (0, 1, 8, 9), as there are some disease sites where distinguishing between L and V is not medically appropriate.

For cases diagnosed January 1, 2018 and later, new codes indicating lymphatic, small vessel, and/or large vessel invasion were added.

1. **Code from pathology report(s).** Code the absence or presence of lymph-vascular invasion as described in the medical record.
 - a. The primary sources of information about lymph-vascular invasion are the pathology check lists (synoptic reports) developed by the College of American Pathologists. If the case does not have a checklist or synoptic report, code from the pathology report or a physician's statement, in that order.
 - b. Do not code perineural invasion in this field.
 - c. Information to code this field can be taken from any specimen from the primary tumor (biopsy or resection.)
 - d. If lymph-vascular invasion is identified in any primary tumor specimen, it should be coded as present/identified.
 - e. Assign Code 8 Not applicable for benign/borderline brain and CNS tumors.
 - f. For cases treated with neoadjuvant therapy, refer to table below in order to code this field. However, if documentation in the medical record indicates information that conflicts with this table, code lymphovascular invasion with the documentation in the medical record.

LVI on pathology report PRIOR to neoadjuvant therapy	LVI on pathology report AFTER neoadjuvant therapy	Code LVI to:
0 - Not present/Not identified	0 - Not present/Not identified	<i>0 - Not present/Not identified</i>
0 - Not present/Not identified	1 - Present/Identified	<i>1 - Present/Identified</i>
0 - Not present/Not identified	9 - Unknown/Indeterminate	<i>9 - Unknown/Indeterminate</i>
1 - Present/Identified	0 - Not present/Not identified	<i>1 - Present/Identified</i>
1 - Present/Identified	1 - Present/Identified	<i>1 - Present/Identified</i>
1 - Present/Identified	9 - Unknown/Indeterminate	<i>1 - Present/Identified</i>
9 - Unknown/Indeterminate	0 - Not present/Not identified	<i>9 - Unknown/Indeterminate</i>
9 - Unknown/Indeterminate	1 - Present/Identified	<i>1 - Present/Identified</i>
9 - Unknown/Indeterminate	9 - Unknown/Indeterminate	<i>9 - Unknown/Indeterminate</i>

2. Use of codes.

- a. Use code 0 when the pathology report indicates that there is no lymphovascular invasion. Assign code 0 for in situ cases.
- b. Use code 1 when the pathology report or a physician's statement indicates that lymphovascular invasion (or one of its synonyms) is present in the specimen.

Synonyms include, but are not limited to

- i. Angiolymphatic invasion
 - ii. Blood vessel invasion
 - iii. Lymph vascular emboli
 - iv. Lymphatic invasion
 - v. Lymph-vascular invasion
 - vi. Vascular invasion
- c. Lymphovascular invasion must be coded 0, 1, 2, 3, 4, or 9 for the following Schemas/Schema IDs
 - Ampulla Vater 00270
 - Appendix 00190
 - Bile Ducts Distal 00260

Bile Ducts Intrahepatic 00230
Bile Ducts Perihilar 00250
Bladder 00620
Buccal Mucosa 00076
Cervix 00520
Colon and Rectum 00200
Corpus Adenosarcoma 00542
Corpus Carcinoma 00530
Corpus Sarcoma 00541
Esophagus (including GE Junction) (excluding Squamous) 00169
Esophagus (including GE Junction) Squamous 00161
Floor of Mouth 00074
Gum 00073
Hypopharynx 00112
Larynx Glottic 00132
Larynx Other 00130
Larynx Subglottic 00133
Larynx Supraglottic 00131
Lip 00071
Lung 00360
Major Salivary Glands 00080
Maxillary Sinus 00121
Melanoma Skin 00470
Merkel Cell Skin 00460
Mouth Other 00077
Nasal Cavity and Ethmoid Sinus 00122
NET Ampulla of Vater 00302
NET Appendix 00320
NET Colon and Rectum 00330
NET Duodenum 00301
NET Pancreas 00340
NET Stomach 00290
Oropharynx (p16-) 00111
Oropharynx (p16+) 00100
Palate Hard 00075
Pancreas 00280
Penis 00570
Placenta 00560
Small Intestine 00180
Stomach 00170
Testis 00590
Thymus 00350
Thyroid 00730
Thyroid Medullary 00740
Tongue Anterior 00072

Vagina 00510

Vulva 00500

- d. Lymphovascular invasion must be coded 0, 1, 2, 3, 4, 8, or 9 for the following Schemas/IDs

Adnexa Uterine Other 00558

Anus 00210

Biliary Other 00278

Breast (Invasive) 00480

Bone Appendicular Skeleton 00381

Bone Pelvis 00383

Bone Spine 00382

Brain 00721

Cervical Lymph Nodes, Occult Head and Neck 00060

Conjunctiva 00650

CNS Other 00722

Cutaneous Carcinoma Head and Neck 00150

Cystic Duct 00242

Digestive Other 00288

Endocrine Other 00778

Eye Other 00718

Fallopian Tube 00553

Gallbladder 00241

Genital Female Other 00559

Genital Male Other 00598

Heart, Mediastinum, and Pleura 00422

Ill-Defined Other 99999

Intracranial Gland 00723

Kaposi Sarcoma 00458

Kidney Parenchyma 00600

Kidney Renal Pelvis 00610

Lacrimal Gland 00690

Lacrimal Sac 00698

Liver 00220

Melanoma Choroid and Ciliary Body 00672

Melanoma Conjunctiva 00660

Melanoma Head and Neck 00140

Melanoma Iris 00671

Middle Ear 00119

NET Adrenal Gland 00770

Orbital Sarcoma 00700

Ovary 00551

Parathyroid 00750

Pharynx Other 00118

Pleural Mesothelioma 00370

Primary Peritoneal Carcinoma 00552

Prostate 00580
Respiratory Other 00378
Retinoblastoma 00680
Retroperitoneum 00440
Sinus Other 00128
Skin Eyelid 00640
Skin Other 00478
Soft Tissue Abdomen and Thorax 00421
Soft Tissue Head and Neck 00400
Soft Tissue Other 00450
Soft Tissue Trunk and Extremities 00410
Trachea 00358
Urethra 00631
Urethra-Prostatic 00632
Urinary Other 00638

e. Use code 8 for the following Schemas/IDs

GIST (2021+) 00430
HemeRetic 00830
Lymphoma 00790
Lymphoma (CLL/SLL) 00795
Lymphoma Ocular Adnexa 00710
Mycosis Fungoides (MF) 00811
Plasma Cell Disorder 00822
Plasma Cell Myeloma 00821
Primary Cutaneous Lymphoma (excluding MF and SS) 00812

d. Use code 9 when

- i. there is no microscopic examination of a primary tissue specimen
- ii. the primary site specimen is cytology only or a fine needle aspiration
- iii. the biopsy is only a very small tissue sample
- iv. it is not possible to determine whether lymph-vascular invasion is present
- v. the pathologist indicates the specimen is insufficient to determine lymph-vascular invasion
- vi. lymph-vascular invasion is not mentioned in the pathology report
- vii. primary site is unknown

e. Clarification between codes 8 and 9:

- i. Code 8 should only be used in the following situations: 1. Standard-setter does not require this item and you are not collecting it. 2. Those histologies noted above described in code 8 for which LVI is always not applicable.
- ii. For those cases where there is no information/documentation from the pathology report or other sources, use code 9

Code	Definition
0	Lymph-vascular Invasion stated as Not Present
1	Lymph-vascular Invasion Present/Identified
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
8	Not applicable
9	Unknown/Indeterminate/not mentioned in path report

DIAGNOSTIC CONFIRMATION

Item Length: 1

NAACCR Item #490

NAACCR Name: Diagnostic Confirmation

XML NAACCR ID: diagnosticConfirmation

Definition:

Records the best method of diagnostic confirmation of the cancer being reported **at any time** in the patient's history.

The codes and instructions for hematopoietic and lymphoid neoplasms are different from the codes for solid tumors. See below.

Codes for Solid Tumors

Code	Definition
Microscopically Confirmed	
1	Positive histology.
2	Positive cytology.
4	Positive microscopic confirmation, method not specified.
Not Microscopically Confirmed	
5	Positive laboratory test/marker study
6	Direct visualization without microscopic confirmation
7	Radiography and other imaging techniques without microscopic confirmation
8	Clinical diagnosis only (other than 5, 6, or 7).
Confirmation Unknown	
9	Unknown whether or not microscopically confirmed.

Coding Instructions for Solid Tumors (all tumors except histologies 9590-9992)

1. The codes are in priority order; code 1 has the highest priority. Always code the procedure with the lower numeric value when presence of cancer is confirmed with multiple diagnostic methods.
2. Change to a higher priority code, if at any time during the course of the disease, the patient has a diagnostic confirmation with a higher priority. Change to the higher-priority code even when diagnostic confirmation is based on the result of subsequent treatment.

Example: Benign brain tumor diagnosed on MRI. Assign diagnostic confirmation code 7. Patient later becomes symptomatic and the tumor is surgically removed. Change diagnostic confirmation code to 1.

3. Assign code 1 when the microscopic diagnosis is based on
 - a. Tissue specimen from biopsy, frozen section, surgery, autopsy or D&C
 - b. Bone marrow specimens (aspiration and biopsy)
4. Assign code 2 when the microscopic diagnosis is based on

- a. Examination of cells (rather than tissue) including but not limited to: sputum smears, bronchial brushings, bronchial washings, prostatic secretions, breast secretions, gastric fluid, spinal fluid, peritoneal fluid, pleural fluid, urinary sediment, cervical smears, or vaginal smears
 - b. Paraffin block specimens from concentrated spinal, pleural or peritoneal fluid.
5. Assign code 4 when there is information that the diagnosis of cancer was microscopically confirmed, but the type of confirmation is unknown.
6. Assign code 5 when the diagnosis of cancer is based on laboratory tests or marker studies that are clinically diagnostic for that specific cancer.

Example 1: The presence of alpha-fetoprotein for liver cancer

Example 2: If the workup for a prostate cancer patient is limited to a highly elevated PSA and the physician diagnoses and/or treats the patient based only on the PSA, code the diagnostic confirmation to 5.

Note: For tests and tumor markers that may be used to help diagnose cancer, see <https://www.cancer.gov/about-cancer/diagnosis-staging/diagnosis/tumor-markers-list>

- a. The surgeon's operative report from a surgical exploration or endoscopy such as colonoscopy, mediastinoscopy, or peritoneoscopy and no tissue was examined.
 - b. Gross autopsy findings (no tissue or cytologic confirmation)
7. Assign code 6 when the diagnosis is based only on
8. Assign code 7 when the only confirmation of malignancy was diagnostic imaging such as computerized axial tomography (CT scans), magnetic resonance imaging (MRI scans), or ultrasounds/sonography.
9. Assign code 8 when the case was diagnosed by any clinical method not mentioned in preceding codes. The diagnostic confirmation is coded 8 when the only confirmation of disease is a physician's clinical diagnosis.

Example: CT diagnosis is possible lung cancer. Patient returns to the nursing home with a DNR order. Physician enters a diagnosis of lung cancer in the medical record. Code the diagnostic confirmation to 8: there is a physician's clinical diagnosis-clinical diagnosis made by the physician using the information available for the case.

- a. When it is unknown if the diagnosis was confirmed microscopically
 - b. For death-certificate-only cases
10. Assign code 9

Codes for Hematopoietic and Lymphoid Neoplasms (9590/3-9993/3)

Code	Definition
Microscopically Confirmed	
1	Positive histology.
2	Positive cytology.
3	Positive histology PLUS: (ONLY for hematopoietic and lymphoid neoplasms (9590/3-9992/3). Effective for cases diagnosed 1/1/2010) <ul style="list-style-type: none">• Positive immunophenotyping AND/OR• Positive genetic studies
4	Positive microscopic confirmation, method not specified.
Not Microscopically Confirmed	
5	Positive laboratory test/marker study
6	Direct visualization without microscopic confirmation
7	Radiography and other imaging techniques without microscopic confirmation
8	Clinical diagnosis only (other than 5, 6, or 7).
Confirmation Unknown	
9	Unknown whether or not microscopically confirmed.

Coding Instructions for Hematopoietic and Lymphoid Neoplasms (9590/3-9993/3)

1. There is no priority order or hierarchy for coding the Diagnostic Confirmation for hematopoietic or lymphoid neoplasms. Most commonly the specific histology type is determined through immunophenotyping or genetic testing. See the Hematopoietic Database (DB) for information on the definitive diagnostic confirmation for specific types of tumors.
2. Assign code 1 when the microscopic diagnosis is based on:
 - a. Tissue specimens from biopsy, frozen section, surgery, or autopsy
 - b. Bone marrow specimens (aspiration and biopsy)
 - c. For leukemias only, complete blood count (CBC), white blood count (WBC), and peripheral blood smear

Note: Use code 1 when ONLY the tissue, bone marrow, or blood was used to diagnose the specific histology. Do not use code 1 if the diagnosis was based on immunophenotyping or genetic testing using tissue, bone marrow, or blood.

3. Code 2 would rarely be used for hematopoietic or lymphoid neoplasms. Use code 2 when the microscopic diagnosis is based on
 - a. Examination of cells (other than tissue) including but not limited to: spinal fluid, peritoneal fluid, or pleural fluid.
 - b. Paraffin block specimens from concentrated spinal, pleural or peritoneal fluid.
 - c. These methods are rarely used for hematopoietic or lymphoid tumors.

4. Assign code 3 when BOTH a histology positive for cancer AND also positive immunophenotyping and/or positive genetic testing are available. Do not use code 3 for neoplasms diagnosed prior to January 1, 2010.

Example 1: Bone marrow examination is positive for acute myeloid leukemia (9861/3). Genetic testing shows AML with inv(16)(p13.1q22) (9871/3). Code the Diagnostic Confirmation 3, positive histology and positive genetic testing.

Example 2: Skin biopsy positive for cutaneous T-cell lymphoma, NOS (9709/3). Immunophenotyping shows CD8 positive. Diagnosis is primary cutaneous CD8 positive aggressive epidermotropic T-cell lymphoma (9709/3). Code the Diagnostic Confirmation 3, positive histology and positive genetic testing.

5. Assign code 4 when there is information that the diagnosis of cancer was microscopically confirmed, but the type of confirmation is unknown.
6. Assign code 5 when the diagnosis of a hematopoietic or lymphoid neoplasm is based ONLY on laboratory tests or marker studies that are clinically diagnostic for that specific cancer, but no positive histologic confirmation.

Example: The only information available is that the patient had a positive JAK2 done on a blood sample and is diagnosed with polycythemia vera. Code 5 for diagnosis based on a marker study that is diagnostic of polycythemia vera.

7. Assign code 6 when the diagnosis is based only on
 - a. The surgeon's operative report from a surgical exploration or endoscopy and no tissue was examined.
 - b. Gross autopsy findings (no tissue or cytologic confirmation)
8. Assign code 7 when the only confirmation of malignancy was diagnostic imaging such as computerized axial tomography (CT), magnetic resonance imaging (MRI), or ultrasounds/sonography.
9. Assign code 8 when the case was diagnosed by any clinical method not mentioned in the preceding codes. A number of hematopoietic and lymphoid neoplasms are diagnosed clinically; these are called "diagnoses of exclusion" (the tests for the disease are equivocal and the physician does a clinical diagnosis based on the information from the equivocal tests and the patient's clinical presentation).

Note: The hematopoietic DB will identify clinical diagnosis as the definitive diagnostic method.

10. Assign code 9

- a. When it is unknown if the diagnosis was confirmed microscopically
- b. For death-certificate-only cases

AGE AT DIAGNOSIS

Item Length: 3
NAACCR Item #230
NAACCR Name: Age at Diagnosis
XML NAACCR ID: ageAtDiagnosis

Definition:

The item must be computer generated.

Records the age of the patient at diagnosis in complete years.

Coding Instructions:

If the patient has multiple primaries, then the age at diagnosis may be different for subsequent primaries.

STAGE/PROGNOSTIC FACTORS

TUMOR SIZE SUMMARY

Item Length: 3
NAACCR Item #756
NAACCR Name: Tumor Size Summary
XML NAACCR ID: tumorSizeSummary

Description

This data item records the most accurate measurement of a solid primary tumor, usually measured on the surgical resection specimen. Tumor size can indicate extent of disease and be used for quality assurance.

Coding Instructions

- **All measurements should be in millimeters(mm)**
- **Record size in the specified order:**
 1. Size measured on the surgical resection specimen, when **surgery is administered as the first definitive treatment. No pre-surgical treatment has been administered.**
 - If there is a discrepancy among tumor size measurements in the various sections of the pathology report, code the size from the synoptic report (also known as the CAP protocol or pathology checklist). If only a text report is available, use final diagnosis, microscopic, or gross examination, in that order.
 2. If neoadjuvant therapy followed by surgery, do not record the size of the pathologic specimen. Code the largest size of tumor prior to neoadjuvant treatment. If unknown, codes size as 999.
 3. If no surgical resection, then largest measurement of the tumor from physical exam, imaging, or other diagnostic procedures prior to any other form of treatment (See Coding Rules below).
 4. If 1, 2, and 3 do not apply, the largest size from all information available within four months of the date of diagnosis, in the absence of disease progression.
- **Coding Rules**
 1. Tumor size is the diameter of the tumor, not the depth or thickness of the tumor.

2. Recording less than/greater than Tumor Size

- If a tumor size is reported as less than x mm or less than x cm, the reported tumor size should be 1 mm less; for example if size is <10 mm, code size as 009. Often these are given in cm such as <1 cm which is coded as 009, <2 cm is coded as 019, <3 cm is coded as 029, <4 cm is coded as 039, <5 cm is coded as 049. If stated as less than 1 mm uses code 001.
- If tumor size is reported as more than x mm or more than x cm, code size as 1 mm more; for example if size is >10 mm, size should be coded as 011. Often these are given in cm such as >1 cm, which is coded as 011, >2 cm is coded as 021, >3 cm is coded as 031, >4 cm is coded as 041, >5 cm is coded as 051. If described as anything greater than 989 mm (98.9 cm) code as 989.
- If tumor size is reported to be between two sizes, record tumor size as the midpoint between the two: i.e., add the two sizes together and then divide by two (“between 2 and 3 cm” is coded as 025).

3. Rounding

- Round the tumor size only if it is described in fractions of millimeters. If the largest dimension of a tumor is less than 1 millimeter (between 0.1 and 0.9 mm), record size as 001 (do not round down to 000). If tumor size is greater than 1 millimeter, round tenths of millimeters in the 5-9 range up to the nearest whole millimeter. Do not round tumor size expressed in centimeters to the nearest whole centimeters (rather, move the decimal point one space to the right, converting the measurement to millimeters). For breast cancer, please follow the AJCC 8th Edition, Breast Chapter.

4. Priority of imaging/radiographic techniques

- Information on size from imaging/radiographic techniques can be used to code size when there is no more specific size information from a pathology or operative report, but it should be taken as low priority, over physical exam.

5. Tumor size discrepancies among imaging and radiographic reports

- If there is a difference in reported tumor size among imaging and radiographic techniques, unless the physician specifies which imaging is most accurate, record the largest size in the record, regardless of which imaging technique reports it.

6. **Always code the size of the primary tumor, not the size of the polyp, ulcer, cyst, or distant metastasis.**
 - If the tumor is described as a “cystic mass,” and only the size of the entire mass is given, code the size of the entire mass, since the cysts are part of the tumor itself.
7. **Record the size of the invasive component, if given.**
 - If both an in situ and invasive component are present and the invasive component is measured, record the size of the invasive component even if it is smaller.
 - If the size of the invasive component is not given, record the size of the entire tumor from the surgical report, pathology report, radiology report or clinical examination.
8. **Record the largest dimension or diameter of the tumor, whether it is from an excisional biopsy specimen or the complete resection of the primary tumor.**
9. **Record the size as stated for purely in situ lesions.**
10. **Disregard microscopic residual or positive surgical margins when coding tumor size.**
 - Microscopic residual tumor does not affect the overall tumor size. The status of primary tumor margins may be recorded in a separate data item.
11. **Do not add the size of pieces of chips together to create a whole.**
 - The pieces or chips may not be from the same location, or they may represent only a very small portion of a large tumor. However, if the pathologist states an aggregate or composite size (determined by fitting the tumor pieces together and measuring the total size), record that size. If the only measurement describes pieces or chips, record tumor size as 999.
12. **Multifocal/multicentric tumors**
 - If the tumor is multi-focal or if multiple tumors are reported as a single primary, code the size of the largest invasive tumor or if all of the tumors are in situ, code the size of the largest in situ tumor.

13. Tumor size code 999 is used when size is unknown or not applicable.

- Site/morphologies where tumor size is not applicable are listed here.
Hematopoietic, Reticuloendothelial, and Myeloproliferative neoplasms: histology codes 9590-9992
Kaposi Sarcoma
Melanoma Choroid
Melanoma Ciliary Body
Melanoma Iris

14. Tumor size code 000 is used for the following schema:

- Schema is Cervical Lymph Nodes and Unknown Primary 00060
- Occult Cervical Lymph Node (See STORE, Overview of Coding Principles, page 44).

15. Document the information to support coded tumor size in the appropriate text data item of the abstract.

Code	Definition
000	No mass/tumor found
001	1 mm or described as less than 1 mm
002-988	Exact size in millimeters (2 mm to 988 mm)
989	989 millimeters or larger
990	Microscopic focus or foci only and no size of focus is given
998	<p>SITE-SPECIFIC CODES: Alternate descriptions of tumor size for specific sites:</p> <p>Familial/multiple polyposis: Rectosigmoid and rectum (C19.9, C20.9) Colon (C18.0, C18.2-C18.9)</p> <p>If no size is documented:</p> <p>Circumferential: Esophagus (C15.0-C15.5, C15.8-C15.9)</p> <p>Diffuse; widespread: 3/4s or more; linitis plastica: Stomach and Esophagus Ge Junction (C16.0-C16.6, C16.8-C16.9)</p> <p>Diffuse, entire lung or NOS: Lung and main stem bronchus (C34.0-C34.3, C34.8-C34.9)</p> <p>Diffuse: Breast (C50.0-C50.6, C50.8-C50.9)</p>

Code	Definition
999	Unknown; size not stated Not documented in patient record Size of tumor cannot be assessed Not applicable

Coding Examples:

Code	Explanation
028	Chest x-ray shows 3.5 cm mass; the pathology report from the surgery states that the same mass is malignant and measures 2.8 cm (28 mm).
032	Pathology report states lung carcinoma is 2.1 cm x 3.2 cm x 1.4 cm. Record the largest dimension 032 (32 mm)
022	Patient has a 2.2 cm mass in the oropharynx; fine needle aspiration of mass confirms squamous cell ca. Patient receives a course of neoadjuvant combination chemotherapy. Pathologic size after total resection is 2.8. cm Record tumor size as 022 (22 mm) from the imaging prior to the neoadjuvant therapy.
007	Breast cancer described as 6.5 millimeters in size. Round up tumor size as 007.
002	Cancer in polyp described as 2.3 millimeters in size. Round down tumor size as 002.
001	Focus of cancer described as 1.4 mm in size. Round down as 001.
005	5.2 mm breast cancer. Round down to 5 mm and code as 005.
014	Tumor is mixed in situ and invasive adenocarcinoma, total 3.7 cm in size, of which 1.4 cm is invasive. Record tumor size as 014 (14 mm)
023	A breast tumor with infiltrating duct carcinoma with extensive in situ component, total size 2.3 cm. Record tumor size as 023 (23 mm)
019	Duct carcinoma in situ measuring 1.9 cm with an area of invasive ductal carcinoma. Record tumor size as 019 (19 mm)
051	Tumor is described as 2.4 x 5.1 x 1.8 cm in size. Record tumor size as 051 (51 mm)

SUMMARY STAGE 2018

Item Length: 1

NAACCR Item #764

NAACCR Name: Summary Stage 2018

XML NAACCR ID: summaryStage2018

Definition:

Summary Stage 2018 is effective for cases diagnosed 1/1/2018 or later. Summary stage groups cases into broad categories of in situ, local, regional, and distant. Summary stage can be used to evaluate disease spread at diagnosis, treatment patterns and outcomes over time.

Coding Instructions:

Code summary stage at the initial diagnosis or treatment of the reportable tumor. Summary stage should include all information available through completion of surgery(ies) in the first course of treatment or within 4 months of diagnosis in the absence of disease progression, whichever is longer.

- Refer to the site and histology-specific definitions of categories and coding instructions in the SEER Summary Staging Manual 2018 (<https://seer.cancer.gov/tools/ssm/>).
- Use Code 8 for benign and borderline brain/CNS cases.

Code	Definition
0	In situ
1	Localized only
2	Regional by direct extension only
3	Regional lymph nodes only
4	Regional by BOTH direct extension AND regional lymph nodes
7	Distant site(s)/node(s) involved
8	Benign, borderline*
9	Unknown if extension or metastasis (unstaged, unknown, or unspecified) Death certificate only case

*Applicable for the following Summary Stage 2018 chapters: Brain, CNS Other, Intracranial Gland.

BRAIN MOLECULAR MARKERS

Item Length: 2

NAACCR Item #3816

NAACCR Name: Brain Molecular Markers

XML NAACCR ID: brainMolecularMarkers

Definition:

This data item only applies to cancers with primary site C700, C710-C719.

Multiple brain molecular markers have become standard pathology components necessary for diagnosis. This data item captures clinically important brain cancer subtypes identified by molecular markers that are not distinguishable by ICD-O-3 codes.

Coding Instructions:

Note 1: This data item applies only to ICD-O-3 histology codes: 9400/3, 9401/3, 9440/3, 9450/3, 9451/3, 9471/3 and 9478/3. If a histology is not included in this list, assign, code 85.

Note 2: Physician statement of histologic subtype can be used to code this data item.

Note 3: Only one code is applicable for each tumor.

- IDH mutation status distinguishes between clinically important subtypes within ICD-O-3 9400/3, Diffuse astrocytoma and 9401/3, Anaplastic astrocytoma.
- IDH mutant and 1p/19q co-deletion distinguishes between clinically important subtypes within ICD-O-3 code 9450/3, Oligodendroglioma and 9451/3, Anaplastic Oligodendroglioma.
- IDH-wildtype distinguishes clinically important subtypes within ICD-O-3 9400/3, Diffuse astrocytoma, 9401/3, Anaplastic astrocytoma and 9440/3, Glioblastoma, Epithelioid glioblastoma and Glioblastoma, NOS (note that the new ICD-O-3 code 9445/3 applies to Glioblastoma, IDH-mutant; information regarding this subtype is not collected using this data item).
- SHH-activation and TP53-wildtype distinguishes between clinically important subtypes within ICD-O-3 histology code 9471/3, Medulloblastoma.
- C19MC alteration status distinguishes a clinically important highly aggressive subtype within ICD-O-3 9478/3, Embryonal tumor with multilayered rosettes.

Code	Definition
01	Diffuse astrocytoma, IDH-mutant (9400/3)
02	Diffuse astrocytoma, IDH-wildtype (9400/3)
03	Anaplastic astrocytoma, IDH-mutant (9401/3)
04	Anaplastic astrocytoma, IDH-wildtype (9401/3)
05	Glioblastoma, IDH-wildtype (9440/3)
06	Oligodendroglioma, IDH-mutant and 1 p/19q co-deleted (9450/3)
07	Anaplastic oligodendroglioma, IDH-mutant and 1p/19q co-deleted (9451/3)
08	Medulloblastoma, SHH-activated and TP53-wildtype (9471/3)
09	Embryonal tumor with multilayered rosettes, C19MC-altered (9478/3)
85	Not applicable: Histology not 9400/3, 9401/3, 9440/3, 9450/3, 9451/3, 9471/3, 9478/3
86	Benign or borderline tumor
87	Test ordered, results not in chart
88	Not applicable: Information not collected for this case (If this item is required by your standard setter, use of code 88 will result in an edit error.)
99	Not documented in patient record No microscopic confirmation Brain molecular markers not assessed or unknown if assessed

Coding Examples:

Code	Explanation
01	Biopsy of brain tumor, microscopic confirmation diagnosis: Diffuse Astrocytoma (9400/3). Additional testing done, and IDH-mutant is identified.
99	Biopsy of brain tumor, microscopic confirmation diagnosis: Anaplastic astrocytoma (9401/3). No further testing or results unknown.
99	MRI of brain tumor, clinical diagnosis: glioblastoma. No further workup.
85	Biopsy of brain tumor, microscopic confirmation diagnosis: Mixed glioma (9382/3).

LDH (LACTATE DEHYDROGENASE) LAB VALUE

Item Length: 7

NAACCR Item #3932

NAACCR Name: LDH Lab Value

XML NAACCR ID: ldhPretreatmentLabValue

Definition:

This data item only applies to cancers with primary site C000-C002, C440-C449, C500, C510-C512, C518-C519, C600-C602, C608-C609, C632 and histologies 8720-8790.

LDH (Lactate Dehydrogenase) Lab Value, measured in serum, is a predictor of treatment response, progression-free survival and overall survival for patients with Stage IV melanoma of the skin. It was previously collected as LDH Pretreatment Lab Value and Melanoma Skin, CS SSF# 5.

Coding Instructions:

Note 1: Physician statement of LDH Pretreatment Lab Value can be used to code this data item when no other information is available.

Note 2: LDH is only considered in melanoma staging in the setting of DISTANT metastasis. LDH level might only be ordered after re-excision/wide excision and/or nodal evaluation indicates a higher risk of distant metastasis. Imaging may then be performed and if distant metastasis are identified, LDH is ordered.

Note 3: Record the lab value of the highest serum LDH test results documented in the medical record either before or after surgical resection of the primary tumor with or without regional lymph node dissection. The LDH must be taken prior to systemic (chemo, immunotherapy, hormone), radiation therapy or surgery to a metastatic site. The lab value may be recorded in a lab report, history and physical, or clinical statement in the pathology report.

Code	Definition
0.0	0.0 (U/L)
0.1-99999.9	0.1 – 99,999.9 U/L
XXXXX.1	100,000 U/L or greater
XXXXX.7	Test ordered, results not in chart
XXXXX.8	Not applicable: Information not collected for this case (If this item is required by your standard setter, use of code XXXXX.8 will result in an edit error.)
XXXXX.9	Not documented in medical record LDH (Lactate Dehydrogenase) Pretreatment Lab Value not assessed or unknown if assessed.

BRESLOW TUMOR THICKNESS

Item Length: 4

NAACCR Item #3817

NAACCR Name: Breslow Tumor Thickness

XML NAACCR ID: breslowTumorThickness

Definition:

This data item only applies to cancers with primary site C000-C002, C440-C449, C500, C510-C512, C518-C519, C600-C602, C608-C609, C632 and histologies 8720-8790.

Breslow Tumor Thickness, the measurement of the thickness of a melanoma as defined by Dr. Alexander Breslow, is a prognostic factor for Melanoma of the Skin. Breslow Tumor Thickness is a Registry Data Collection Variable in AJCC. It was previously collected as Melanoma Skin, CS SSF# 1.

Coding Instructions:

Note 1: Physician statement of Breslow Tumor Thickness can be used to code this data item when no other information is available, or the available information is ambiguous.

Note 2: Code Breslow tumor thickness, not size. Record actual measurement in tenths of millimeters from the pathology report. Measurement given in hundredths of millimeters should be rounded to the nearest tenth.

Note 3: Code the greatest measured thickness from any procedure performed on the lesion, whether it is described as a biopsy or an excision.

Note 4: If there are multiple procedures and the pathologist adds the measurement together to get a final Breslow's depth, the register can use this.

- Do not add the measurements together, only the pathologist can do this.

Note 5: If the pathologist describes the thickness as "at least," use the appropriate A code. An exact measurement takes precedence over A codes.

- If the pathologist states "greater than" instead of "at least", code to XX.9, unless it is greater than 9.9 mm (Code AX.0).

Code	Definition
0.0	No mass/tumor found
0.1	Greater than 0.0 and less than or equal to 0.1
0.2-99.9	0.2 - 99.9 millimeters
XX.1	100 millimeters or larger
A0.1-A9.9	Stated as "at least" some measured value of 0.1 to 9.9
AX.0	Stated as greater than 9.9 mm

Code	Definition
XX.8	Not applicable: Information not collected for this schema (If this item is required by your standard setter, use of code XX.8 will result in an edit error)
XX.9	Not documented in medical record Microinvasion; microscopic focus or foci only and no depth given Cannot be determined by pathologist In situ melanoma Breslow Tumor Thickness not assessed or unknown if assessed

Coding Examples:

Code	Explanation
0.4	0.4 mm
1.0	1.0 mm
2.5	2.5 mm
2.6	2.56 mm
11.0	11 mm
12.4	12.35 mm
A2.0	Pathologist states the thickness is "at least 2.0 mm." Code A2.0
XX.9	Not applicable: Information not collected for this schema (If this item is required by your standard setter, use of code XX.8 will result in an edit error)
1.5	If a punch body with a thickness of 1.5 mm is followed by a re-excision with a thickness of residual tumor of 0.2 mm

ER (ESTROGEN RECEPTOR) SUMMARY

Item Length: 1

NAACCR Item #3827

NAACCR Name: Estrogen Receptor Summary

XML NAACCR ID: estrogenReceptorSummary

Definition:

This data item only applies to cancers with primary site C500-C506, C508-C509 with histologies 8000-8700, 8720-8790, 8982-8983, 9700-9701.

ER (Estrogen Receptor) Summary is a summary of results of the estrogen receptor (ER) assay. It was previously collected as Breast CS SSF # 1.

Coding Instructions:

Note 1: Physician statement of ER (Estrogen Receptor) Summary status can be used to code this data item when no other information is available.

Note 2: The result of the ER test performed on the primary breast tissue is to be recorded in this data item.

Note 3: Results from nodal or metastatic tissue may be used ONLY when there is no evidence of primary tumor.

Note 4: In cases where there are invasive and in situ components and ER is done on both, ignore the in situ results.

- If ER is positive on an in situ component and ER is negative on all tested invasive components, code ER as negative (code 0)
- If in situ and invasive components present and ER only done on the in situ component, code unknown (code 9)

Note 5: In cases where there is a single tumor with multiple biopsies and/or surgical resection with different ER results, use the highest (positive vs negative).

Note 6: In cases where there are multiple tumors with different ER results, code the results from the largest tumor size (determined either clinically or pathologically) when multiple tumors are present. Do not use specimen size to determine the largest tumor size.

Note 7: If neoadjuvant therapy is given, record the assay from tumor specimens prior to neoadjuvant therapy. If neoadjuvant therapy is given and there are no ER results from pre-treatment specimens, report the findings from post-treatment specimens.

Note 8: If the patient is ER positive and node negative, a multigene test such as Oncotype Dx may be performed, in which case another ER test will be performed. Do not record the results of that test in this field.

- Record only the results of the test which made the patient eligible to be given the multigene test.

Code	Definition
0	ER negative
1	ER positive
7	Test ordered, results not in chart
9	Not documented in medical record Cannot be determined (indeterminate) ER (Estrogen Receptor) Summary status not assessed or unknown if assessed

Coding Examples:

Code	Explanation
0	Patient had a biopsy 1/1/2018 which showed ER positive DCIS. Lumpectomy specimen from surgical resection on 2/5/18 showed a 1.1 cm invasive ductal carcinoma. No residual DCIS. Studies on the resected specimen were ER negative.
1	Patient diagnosed with ductal carcinoma on 3/5/2018. Studies on the biopsy are ER positive. Pt had neoadjuvant chemotherapy followed by a resection. The studies on the resected specimen are ER negative.

PR (PROGESTERONE RECEPTOR) SUMMARY

Item Length: 1

NAACCR Item #3915

NAACCR Name: Progesterone Receptor Summary

XML NAACCR ID: progesteroneRecepSummary

Definition:

This data item only applies to cancers with primary site C500-C506, C508-C509 with histologies 8000-8700, 8720-8790, 8982-8983, 9700-9701.

PR (Progesterone Receptor) Summary is a summary of results of the progesterone receptor (PR) assay. It was previously collected as Breast CS SSF # 2.

Coding Instructions:

Note 1: Physician statement of PR (Progesterone Receptor) Summary status can be used to code this data item when no other information is available.

Note 2: The result of the PR test performed on the primary breast tissue is to be recorded in this data item.

Note 3: Results from nodal or metastatic tissue may be used ONLY when there is no evidence of primary tumor.

Note 4: In cases where there are invasive and in situ components and PR is done on both, ignore the in situ results.

- If PR is positive on an in situ component and PR is negative on all tested invasive components, code PR as negative (code 0)
- If in situ and invasive components present and PR only done on the in situ component, code unknown (code 9)

Note 5: In cases where there is a single tumor with multiple biopsies and/or surgical resection with different PR results, use the highest (positive vs negative).

Note 6: In cases where there are multiple tumors with different PR results, code the results from the largest tumor size (determined either clinically or pathologically) when multiple tumors are present. Do not use specimen size to determine the largest tumor size.

Note 7: If neoadjuvant therapy is given, record the assay from tumor specimens prior to neoadjuvant therapy. If neoadjuvant therapy is given and there are no PR results from pre-treatment specimens, report the findings from post-treatment specimens.

Note 8: If the patient is PR positive and node negative, a multigene test such as Oncotype Dx may be performed, in which case another PR test will be performed. Do not record the results of that test in this field.

Code	Definition
0	PR negative
1	PR positive
7	Test ordered, results not in chart
9	Not documented in medical record Cannot be determined (indeterminate) PR (Progesterone Receptor) Summary status not assessed or unknown if assessed

Coding Examples:

Code	Explanation
0	Patient had a biopsy 1/1/2018 which showed PR positive DCIS. Lumpectomy specimen from surgical resection on 2/5/18 showed a 1.1 cm invasive ductal carcinoma. No residual DCIS. Studies on the resected specimen were PR negative.
1	Patient diagnosed with ductal carcinoma on 3/5/2018. Studies on the biopsy are PR positive. Pt had neoadjuvant chemotherapy followed by a resection. The studies on the resected specimen are PR negative.

HER2 OVERALL SUMMARY

Item Length: 1
NAACCR Item #3855
NAACCR Name: HER2 Overall Summary
XML NAACCR ID: her2OverallSummary

Definition:

This data item only applies to cancers with primary site C500-C506, C508-C509 with histologies 8000-8700, 8720-8790, 8982-8983, 9700-9701.

HER2 Overall Summary is a summary of results from HER2 testing. It was previously collected as Breast CS SSF # 15.

Coding Instructions:

Note 1: Physician statement of HER2 Overall Summary status can be used to code this data item when no other information is available.

Note 2: The result of the HER2 test performed on the primary breast tissue is to be recorded in this data item.

Note 3: Results from nodal or metastatic tissue may be used ONLY when there is no evidence of primary tumor.

Note 4: In cases where there are invasive and in situ components and HER2 is done on both, ignore the in situ results.

- If HER2 is positive on an in situ component and HER2 is negative on all tested invasive components, code HER2 as negative (code 0)
- If in situ and invasive components present and HER2 only done on the in situ component, code unknown (code 9)

Note 5: In cases where there is a single tumor with multiple biopsies and/or surgical resection with different HER2 results, use the highest (positive vs negative).

Note 6: In cases where there are multiple tumors with different HER2 results, code the results from the largest tumor size (determined either clinically or pathologically) when multiple tumors are present. Do not use specimen size to determine the largest tumor size.

Note 7: If neoadjuvant therapy is given, record the assay from tumor specimens prior to neoadjuvant therapy. If neoadjuvant therapy is given and there are no HER2 results from pre-treatment specimens, report the findings from post-treatment specimens.

Note 8: If the patient is HER2 positive and node negative, a multigene test such as Oncotype Dx may be performed, in which case another HER2 test will be performed. Do not record the results of that test in this field.

Code	Definition
0	HER2 negative; equivocal
1	HER2 positive
7	Test ordered, results not in chart
9	Not documented in medical record Cannot be determined (indeterminate) HER2 Overall Summary status not assessed or unknown if assessed

Coding Examples:

Code	Explanation
0	Patient had a biopsy 1/1/2018 which showed HER2 positive DCIS. Lumpectomy specimen from surgical resection on 2/5/18 showed a 1.1 cm invasive ductal carcinoma. No residual DCIS. Studies on the resected specimen were HER2 negative.
1	Patient diagnosed with ductal carcinoma on 3/5/2018. Studies on the biopsy are HER2 positive. Pt had neoadjuvant chemotherapy followed by a resection. The studies on the resected specimen are HER2 negative.

FIBROSIS SCORE

Item Length: 1
NAACCR Item #3835
NAACCR Name: Fibrosis Score
XML NAACCR ID: fibrosisScore

Definition:

This data item only applies to cancers with primary site C220-C221 with histologies 8000-8700, 8720-8790, 9700-9701.

Fibrosis Score, the degree of fibrosis of the liver based on pathological examination, is a prognostic factor for liver cancer. This data item was previously collected for Liver, CS SSF #2.

Coding Instructions:

Note 1: Physician statement of fibrosis score can be used to code this data item when no other information is available. However, code 7 when the physician statement of fibrosis score is not based on histologic examination of the liver.

Note 2: FIB-4 is NOT a pathological fibrosis score of 4. It is a scoring method using the patient's age and relevant lab values to calculate a score. The medical record may show something like "FIB-4 = 3.52." Do not code FIB-4 values in this data item.

Note 3: AJCC classifies Ishak fibrosis scores 0-4 (none to moderate fibrosis) as F0, and Ishak fibrosis scores 5-6 (cirrhosis/severe fibrosis) as F1. This is not the same as METAVIR score F0 or F1.

Note 4: Record the results based on information collected during the initial work-up. If multiple biopsies are taken and have conflicting scores, use the results from the biopsy closest to the start of treatment. Information collected after the start of treatment may not be used to code this data item.

Note 5: To use codes 0 and 1, you must have a histologic(microscopic) confirmation of fibrosis/cirrhosis. Code the absence (code 0) or presence (code 1) of fibrosis as documented in the pathology report.

Note 6: Use code 7 if there is a clinical diagnosis (no microscopic confirmation) of severe fibrosis or cirrhosis.

Note 7: If no score is mentioned, descriptive terms may be used to assign codes 0 and 1 - see specific terms in the table below.

Note 8: If a fibrosis score is stated but the scoring system is not recorded, consult with the physician. If no further information is available, code 9.

Code	Definition
0	Ishak fibrosis score 0-4 No to moderate fibrosis METAVIR score F0-F3 Batt-Ludwig score 0-3
1	Ishak fibrosis score 5-6 Advanced/severe fibrosis METAVIR score F4 Batt-Ludwig score 4 Developing cirrhosis Incomplete cirrhosis Transition to cirrhosis Cirrhosis, probable or definite Cirrhosis, NOS
7	Clinical statement of advanced/severe fibrosis or cirrhosis, AND Not histologically confirmed or unknown if histologically confirmed
8	Not applicable: Information not collected for this case (If this item is required by your standard setter, use of code 8 will result in an edit error.)
9	Not documented in medical record Stated in medical record that patient does not have advanced cirrhosis/advanced fibrosis, not histologically confirmed or unknown if histologically confirmed Fibrosis score stated but cannot be assigned to codes 0 or 1 Fibrosis score stated but scoring system not recorded Fibrosis Score not assessed or unknown if assessed

MICROSATELLITE INSTABILITY (MSI)

Item Length: 1

NAACCR Item #3890

NAACCR Name: Microsatellite Instability

XML NAACCR ID: microsatelliteInstability

Definition:

This data item only applies to cancers with primary site C180, C182-189, C199, C209 with histologies 8000-8149, 8154, 8157, 8160-8231, 8243-8248, 8250-8682, 8690-8700, 8720-8790, 9700-9701.

Microsatellite Instability (MSI) is a form of genetic instability manifested by changes in the length of repeated single- to six-nucleotide sequences (known as DNA microsatellite sequences). High MSI, found in about 15% of colorectal carcinomas, is an adverse prognostic factor for colorectal carcinomas and predicts poor response to 5-FU chemotherapy (although the addition of oxaliplatin in FOLFOX regimens negates the adverse effects [page 266 AJCC manual]). High MSI is a hallmark of hereditary nonpolyposis colorectal carcinoma, also known as Lynch syndrome. Microsatellite Instability (MSI) is a Registry Data Collection Variable in AJCC. It was previously collected as Colon and Rectum, CS SSF# 7.

Coding Instructions:

Note 1: Physician statement of MSI status can be used to code this data item when no other information is available.

Note 2: The microsatellite instability (MSI) test is a genetic test performed on tumor tissue to look for differences in length of certain non-functioning sections of DNA. The differences are caused by problems with the genes that encode proteins that normally repair certain types of DNA damage. A high proportion of colon cancers arising in patients with hereditary nonpolyposis colorectal cancer (HNPCC) (also known as Lynch syndrome) have high MSI and a smaller percentage of colon cancers not associated with Lynch syndrome have high MSI. Patients with colon cancers with high MSI may be further tested to determine if they have HNPCC. In addition, MSI is a useful prognostic marker in that patients with high MSI colon cancers have better response to surgery and survival.

Note 3: Testing for MSI may be done by immunology or genetic testing. Only genetic testing results will specify whether the MSI is low or high.

- MSI is looking at instability in informative markers.
- MSI results are recorded as
 - MSS (Code 0)

- Stable (Code 0)
- Negative (Code 0)
- Low probability of MSI-H (Code 0)
- MSS/MSI-L (Code 0)
- MSI-L (Code 1)
- Unstable, high (Code 2)
- Unstable, NOS (no designation of high or low) (Code 2)
- MSI-H (code 2)
- MSI-I (intermediate) (Code 9)

Note 4: Testing for Mismatch Repair (MMR) is usually done by immunohistochemistry (IHC).

- Most common markers are MLH1, MSH2, MSH6, PMS2
- MMR results are recorded as
 - No loss of nuclear expression (code 0)
 - Mismatch repair (MMR) intact (code 0)
 - MMR proficient (pMMR or MMR-P) (code 2)
 - MMR normal (code 0)
 - Loss of nuclear expression (code 2)
 - MMR deficient (pMMR or MMR-P) (code 2)
 - MMR abnormal (code 2)

Note 5: If both tests are done and one or both are positive, code 2.

Note 6: If all tests done are negative, code 0.

Code	Definition
0	Microsatellite instability (MSI) stable; microsatellite stable (MSS); negative, NOS AND/OR Mismatch repair (MMR) intact, no loss of nuclear expression of MMR proteins
1	MSI unstable low (MSI-L)
2	MSI unstable high (MSI-H) AND/OR MMR-D (loss of nuclear expression of one or more MMR proteins, MMR protein deficient)
8	Not applicable: Information not collected for this case (If this information is required by your standard setter, use of code 8 may result in an edit error.)
9	Not documented in medical record MSI-indeterminate Microsatellite instability not assessed or unknown if assessed

PSA (PROSTATIC SPECIFIC ANTIGEN) LAB VALUE

Item Length: 5
NAACCR Item #3920
NAACCR Name: PSA Lab Value
XML NAACCR ID: psaLabValue

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

PSA (Prostatic Specific Antigen) is a protein produced by cells of the prostate gland and is elevated in patients with prostate cancer. This data item pertains to PSA lab value. It was previously collected as Prostate, CS SSF# 1.

Coding Instructions:

Note 1: Physician statement of prostatic specific antigen (PSA) pre-diagnosis can be used to code this data item when no other information is available.

Note 2: PSA is a prognostic factor required for AJCC staging. It affects the stage group in most cases.

Note 3: Record to the nearest tenth in nanograms/milliliter (ng/ml) the last pre-diagnosis PSA lab value prior to diagnostic biopsy of prostate and treatment. The lab value may be recorded in the lab report, history and physical, or clinical statement in the pathology report, etc.

- A lab value expressed in micrograms per liter (ug/L) is equivalent to the same value expressed in nanograms per milliliter (ng/ml)
- Record 0.1 when the lab results are stated as less than 0.1 ng/ml with no exact value.

Note 4: A discrepancy between the PSA documented in the lab report and the PSA documented by the clinician may arise due to the clinician's adjusting the PSA value. Certain medications for benign prostatic hypertrophy (BPH) decrease the PSA.

- If there is documentation by a clinician within the medical record of an adjusted PSA value, record the adjusted value.
- The registrar does not adjust the PSA value based on BPH medication use.
- If there is no documentation by a clinician within the medical record of an adjusted PSA value, record the PSA value provided.

- The fact that an adjusted PSA value is being recorded should be documented in the Dx Proc - Lab Tests text field (NAACCR Item # 2550).

Code	Definition
0.1	0.1 or less nanograms/milliliter (ng/ml) (Exact value to nearest tenth of ng/ml)
0.2-999.9	0.2 - 999.9 ng/ml (Exact value to nearest tenth of ng/ml)
XXX.1	1,000 ng/ml or greater
XXX.7	Test ordered, results not in chart
XXX.9	Not documented in medical record PSA lab value not assessed or unknown if assessed

Coding Examples:

Code	Explanation
7.2	PSA of 7.2
10.0	PSA of 10.0
8.6	PSA of 8.56
110.4	PSA of 110.35

GLEASON SCORE CLINICAL

Item Length: 2

NAACCR Item #3840

NAACCR Name: Gleason Score Clinical

XML NAACCR ID: gleasonScoreClinical

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

This data item records the Gleason score based on adding the values for primary and secondary patterns in Needle Core Biopsy or TURP. Gleason Score Clinical is a Registry Data Collection Variable for AJCC. This data item was previously collected as Prostate, CS SSF# 8.

Coding Instructions

Note 1: Physician statement of Gleason Score Clinical can be used to code this data item when there is no other information available.

Note 2: Code the Gleason Score Clinical from needle core biopsy or transurethral resection of prostate (TURP) in this field. Gleason primary and secondary patterns provided for any prostate tissue identified from a transurethral resection of a bladder tumor (TURBT) specimen can also be used in this field.

Note 3: Code the Gleason Score prior to neoadjuvant treatment.

Note 4: Usually prostate cancers are graded using Gleason's score or pattern. Gleason's grading for prostate primaries is based on a 5-component system (5 histologic patterns). Prostatic cancer generally shows two main histologic patterns. The primary pattern, the pattern occupying greater than 50% of the cancer, is usually indicated by the first number of the Gleason grade, and the secondary pattern is usually indicated by the second number. These two numbers are added together to create a pattern score, ranging from 2 to 10.

- If there are two numbers, assume that they refer to two patterns (the first number being the primary pattern and the second number the secondary pattern), and sum them to obtain the score.
- If the only one number is given, and it is less than or equal to 5, code the total score to X9, unknown or no information.
- If only one number is given, and it is greater than 5, assume that it is a score and code as stated.

- If the pathology report specifies a specific number out a total of 10, the first number given is the score.

Note 5: If different scores are documented on multiple needle core biopsies, code the highest or most aggressive score.

Note 6: If needle core biopsy and TURP are both performed, code the highest score.

Note 7: Do not infer the Gleason Score from Grade Group (Code X9).

Note 8: Record the Gleason score based on the addition of the primary and secondary patterns coded in Gleason Patterns Clinical [NAACCR Data Item #3838].

Code	Definition
02	Gleason score 2
03	Gleason score 3
04	Gleason score 4
05	Gleason score 5
06	Gleason score 6
07	Gleason score 7
08	Gleason score 8
09	Gleason score 9
10	Gleason score 10
X7	No needle core biopsy/TURP performed
X8	Not applicable: Information not collected for this case (If this information is required by your standard setter, use of code X8 may result in an edit error.)
X9	Not documented in medical record Gleason Score Clinical not assessed or unknown if assessed.

Coding Examples:

Code	Explanation
03	The pathology report from the biopsy says Gleason's 3/10.
07	The pathology report from the biopsy says Gleason 4+3
X9	The pathology report from the biopsy says Gleason 4

GLEASON PATTERNS CLINICAL

Item Length: 2

NAACCR Item #3838

NAACCR Name: Gleason Patterns Clinical

XML NAACCR ID: gleasonPatternsClinical

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

Prostate cancers are graded using Gleason score or pattern. This data item represents the Gleason primary and secondary patterns from needle core biopsy or TURP. Gleason Patterns Clinical is a Registry Data Collection Variable for Clinical Stage for AJCC. This data item was previously collected as Prostate, CS SSF# 7.

Coding Instructions

Note 1: Physician statement of Gleason Patterns Clinical can be used to code this data item when there is no other information available.

Note 2: Code the Gleason primary and secondary patterns from needle core biopsy or transurethral resection of prostate (TURP) in this field. Gleason primary and secondary patterns provided for any prostate tissue identified from a transurethral resection of a bladder tumor (TURBT) specimen can also be used in this field.

Note 3: Code the Gleason primary and secondary patterns prior to neoadjuvant treatment.

Note 4: Usually prostate cancers are graded using Gleason score or pattern. Gleason grading for prostate primaries is based on a 5-component system (5 histologic patterns). Prostatic cancer generally shows two main histologic patterns. The primary pattern, the pattern occupying greater than 50% of the cancer, is usually indicated by the first number of the Gleason grade, and the secondary pattern is usually indicated by the second number. These two numbers are added together to create a pattern score.

- If there are two numbers, assume that they refer to two patterns (the first number being the primary pattern and the second number the secondary pattern), and sum them to obtain the score.
- If the only one number is given, and it is less than or equal to 5, assume that it describes a pattern (since scores of 5 or less would reflect Primary or Secondary Pattern Scores of 1 or 2). Code the number as the primary pattern and code the secondary pattern as Unknown.
- If only one number is given, and it is greater than 5, assume that it is a score.

- If the pathology report specifies a specific number out a total of 10, the first number given is the score.

Note 5: If different patterns are documented on multiple needle core biopsies, code the pattern that reflects the highest or most aggressive score regardless if the pathologist provides an overall pattern in a final summary. If different patterns equal the same high score, give priority to the highest primary pattern and then the highest secondary primary. Do not mix patterns from multiple specimens.

Note 6: If needle core biopsy and TURP are both performed, code the pattern that reflects the highest score.

Note 7: Do not infer Gleason Primary and Secondary Pattern from Grade Group (Code X9).

Note 8: The clinical score is recorded in Gleason Score Clinical [NAACCR Data Item #3840].

Code	Definition
11	Primary pattern 1, secondary pattern 1
12	Primary pattern 1, secondary pattern 2
13	Primary pattern 1, secondary pattern 3
14	Primary pattern 1, secondary pattern 4
15	Primary pattern 1, secondary pattern 5
19	Primary pattern 1, secondary pattern unknown
21	Primary pattern 2, secondary pattern 1
22	Primary pattern 2, secondary pattern 2
23	Primary pattern 2, secondary pattern 3
24	Primary pattern 2, secondary pattern 4
25	Primary pattern 2, secondary pattern 5
29	Primary pattern 2, secondary pattern 9
31	Primary pattern 3, secondary pattern 1
32	Primary pattern 3, secondary pattern 2
33	Primary pattern 3, secondary pattern 3
34	Primary pattern 3, secondary pattern 4
35	Primary pattern 3, secondary pattern 5
39	Primary pattern 3, secondary pattern unknown
41	Primary pattern 4, secondary pattern 1
42	Primary pattern 4, secondary pattern 2
43	Primary pattern 4, secondary pattern 3
44	Primary pattern 4, secondary pattern 4
45	Primary pattern 4, secondary pattern 5
49	Primary pattern 4, secondary pattern unknown
51	Primary pattern 5, secondary pattern 1
52	Primary pattern 5, secondary pattern 2

Code	Definition
53	Primary pattern 5, secondary pattern 3
54	Primary pattern 5, secondary pattern 4
55	Primary pattern 5, secondary pattern 5
59	Primary pattern 5, secondary pattern unknown
X6	TURP and/or Biopsy done, primary pattern unknown, secondary pattern unknown
X7	No needle core biopsy/TURP performed
X8	Not applicable; Information not collected for this case (If this information is required by your standard setter, use of code X8 may result in an edit error.)
X9	Not documented in medical record Gleason Patterns Clinical not assessed or unknown if assessed. Unknown whether TURP and/or Biopsy done

Coding Examples:

Code	Explanation
39	If only one number is given and it is a 3
X6	If only one number is given, and it is a 7.
X6	If the pathology report says Gleason 7/10
43	Two patterns 4+3=7 and 3+4=7 are found in the same specimen. Code the pattern that results in the highest score. If the patterns result in the same score code the highest primary pattern and the secondary pattern that matches.

GLEASON SCORE PATHOLOGICAL

Item Length: 2

NAACCR Item #3841

NAACCR Name: Gleason Score Pathological

XML NAACCR ID: gleasonScorePathological

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

This data item records the Gleason score based on adding the values for primary and secondary patterns prostatectomy or autopsy. Gleason Score Pathological is a Registry Data Collection Variable for AJCC. This data item was previously collected as Prostate, CS SSF# 10.

Coding Instructions

Note 1: Physician statement of Gleason Score Pathological can be used to code this data item when there is no other information available.

Note 2: Code the Gleason Score Pathological from prostatectomy or autopsy only in this field. Do not include patterns from tissues taken prior to prostatectomy.

Note 3: Usually prostate cancers are graded using Gleason's score or pattern. Gleason's grading for prostate primaries is based on a 5-component system (5 histologic patterns). Prostatic cancer generally shows two main histologic patterns. The primary pattern, the pattern occupying greater than 50% of the cancer, is usually indicated by the first number of the Gleason grade, and the secondary pattern is usually indicated by the second number. These two numbers are added together to create a pattern score, ranging from 2 to 10.

- If there are two numbers, assume that they refer to two patterns (the first number being the primary pattern and the second number the secondary pattern), and sum them to obtain the score.
- If the only one number is given, and it is less than or equal to 5, code the total score to X9, unknown or no information.
- If only one number is given, and it is greater than 5, assume that it is a score and code as stated.
- If the pathology report specifies a specific number out a total of 10, the first number given is the score.

Note 4: If neoadjuvant therapy was given, code Gleason pathological score as X9.

Note 5: Do not infer the Gleason Score from Grade Group (Code X9).

Note 6: Record the Gleason score based on the addition of the primary and secondary patterns coded in Gleason Patterns Pathological [NAACCR Data Item #3839].

Code	Definition
02	Gleason score 2
03	Gleason score 3
04	Gleason score 4
05	Gleason score 5
06	Gleason score 6
07	Gleason score 7
08	Gleason score 8
09	Gleason score 9
10	Gleason score 10
X7	No prostatectomy/autopsy performed
X8	Not applicable: Information not collected for this case (If this information is required by your standard setter, use of code X8 may result in an edit error.)
X9	Not documented in medical record Gleason Score Pathological not assessed or unknown if assessed.

Coding Examples:

Code	Explanation
03	The pathology report from the prostatectomy says Gleason's 3/10.
07	The pathology report from the prostatectomy says Gleason 4+3
X9	The pathology report from the prostatectomy says Gleason 4

GLEASON PATTERNS PATHOLOGICAL

Item Length: 2

NAACCR Item #3839

NAACCR Name: Gleason Patterns Pathological
XML NAACCR ID: gleasonPatternsPathological

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

Prostate cancers are graded using Gleason score or pattern. This data item represents the Gleason primary and secondary patterns from prostatectomy or autopsy. Gleason Patterns Pathological is a Registry Data Collection Variable for Clinical Stage for AJCC. This data item was previously collected as Prostate, CS SSF# 9.

Coding Instructions

Note 1: Physician statement of Gleason Patterns Pathological can be used to code this data item when there is no other information available.

Note 2: Code the Gleason primary and secondary patterns from prostatectomy or autopsy in this field. Do not include patterns from tissue taken prior to prostatectomy.

Note 3: Usually prostate cancers are graded using Gleason score or pattern. Gleason grading for prostate primaries is based on a 5-component system (5 histologic patterns). Prostatic cancer generally shows two main histologic patterns. The primary pattern, the pattern occupying greater than 50% of the cancer, is usually indicated by the first number of the Gleason grade, and the secondary pattern is usually indicated by the second number. These two numbers are added together to create a pattern score.

- If there are two numbers, assume that they refer to two patterns (the first number being the primary pattern and the second number the secondary pattern), and sum them to obtain the score.
- If the only one number is given, and it is less than or equal to 5, assume that it describes a pattern (since scores of 5 or less would reflect Primary or Secondary Pattern Scores of 1 or 2). Code the number as the primary pattern and code the secondary pattern as Unknown.
- If only one number is given, and it is greater than 5, assume that it is a score.
- If the pathology report specifies a specific number out a total of 10, the first number given is the score.

Note 4: If neoadjuvant therapy was given, code Gleason pathological patterns as X9.

Note 5: Do not infer Gleason Primary and Secondary Pattern from Grade Group (Code X9).

Note 6: If a tertiary pattern is documented on prostatectomy or autopsy, code in Gleason Tertiary Pattern [NAACCR Data Item #3842].

Note 7: The pathological score is recorded in Gleason Score Pathological [NAACCR Data Item #3841].

Code	Definition
11	Primary pattern 1, secondary pattern 1
12	Primary pattern 1, secondary pattern 2
13	Primary pattern 1, secondary pattern 3
14	Primary pattern 1, secondary pattern 4
15	Primary pattern 1, secondary pattern 5
19	Primary pattern 1, secondary pattern unknown
21	Primary pattern 2, secondary pattern 1
22	Primary pattern 2, secondary pattern 2
23	Primary pattern 2, secondary pattern 3
24	Primary pattern 2, secondary pattern 4
25	Primary pattern 2, secondary pattern 5
29	Primary pattern 2, secondary pattern 9
31	Primary pattern 3, secondary pattern 1
32	Primary pattern 3, secondary pattern 2
33	Primary pattern 3, secondary pattern 3
34	Primary pattern 3, secondary pattern 4
35	Primary pattern 3, secondary pattern 5
39	Primary pattern 3, secondary pattern unknown
41	Primary pattern 4, secondary pattern 1
42	Primary pattern 4, secondary pattern 2
43	Primary pattern 4, secondary pattern 3
44	Primary pattern 4, secondary pattern 4
45	Primary pattern 4, secondary pattern 5
49	Primary pattern 4, secondary pattern unknown
51	Primary pattern 5, secondary pattern 1
52	Primary pattern 5, secondary pattern 2
53	Primary pattern 5, secondary pattern 3
54	Primary pattern 5, secondary pattern 4
55	Primary pattern 5, secondary pattern 5
59	Primary pattern 5, secondary pattern unknown
X6	Prostatectomy done, primary pattern unknown, secondary pattern unknown
X7	No prostatectomy/autopsy performed

Code	Definition
X8	Not applicable; Information not collected for this case (If this information is required by your standard setter, use of code X8 may result in an edit error.)
X9	Not documented in medical record Gleason Patterns Pathological not assessed or unknown if assessed. Unknown if prostatectomy done

Coding Examples:

Code	Explanation
39	If only one number is given and it is a 3
X6	If only one number is given, and it is a 7.
X6	If the pathology report says Gleason 7/10

GLEASON TERTIARY PATTERN

Item Length: 2

NAACCR Item #3842

NAACCR Name: Gleason Tertiary Pattern

XML NAACCR ID: gleasonTertiaryPattern

Definition:

This data item only applies to cancers with primary site C619 with histologies 8000-8700, 8720-8790, 9700-9701.

Prostate cancers are graded using Gleason score or pattern. This data item represents the tertiary pattern value from prostatectomy or autopsy. Tertiary Gleason pattern on prostatectomy is a Registry Data Collection Variable for AJCC. This data item was previously collected as Prostate, CS SSF# 11.

Coding Instructions

Note 1: Physician statement of Gleason tertiary pattern can be used to code this data item when there is no other information available.

Note 2: If present, a high Gleason Tertiary Pattern appears to be an indication for a worse outcome.

Note 3: Record the tertiary pattern documented on prostatectomy or autopsy only. Record the tertiary pattern prior to neoadjuvant treatment.

- If a tertiary pattern is documented on needle core biopsy or transurethral resection of prostate (TURP), it should be disregarded.
- Do not code the tertiary pattern on prostatectomy or autopsy in Gleason Patterns Pathological.

Note 4: The CAP Prostate Protocol does not include Patterns 1 and 2 for Tertiary Pattern.

Note 5: If neoadjuvant therapy was given, code Gleason patterns as X9.

Code	Definition
10	Tertiary pattern 1
20	Tertiary pattern 2
30	Tertiary pattern 3
40	Tertiary pattern 4
50	Tertiary pattern 5
X7	No prostatectomy/autopsy performed
X8	Not applicable: Information not collected for this case (If this information is required by your standard setter, use of code X8 may result in an edit error.)
X9	Not documented in medical record Gleason Tertiary Pattern not assessed or unknown if assessed

TREATMENT – 1ST COURSE

DATE OF FIRST COURSE OF TREATMENT - COC

Item Length: 8

NAACCR Item #1270

NAACCR Name: Date 1st Crs Rx COC

XML NAACCR ID: date1stCrsRxCoc

Definition:

Records the date on which treatment (surgery, radiation, systemic, or other therapy) of the patient was initiated at any facility. This includes the date a decision was made not to treat or a decision was made for active surveillance.

Coding Instructions:

- Record the earliest of the following dates:
 - Date of First Surgical Procedure
 - Date Radiation Started
 - Date Systemic Therapy Started
 - Date Other Treatment Started

- In cases of non-treatment, in which a physician decides not to treat a patient or a patient’s family or guardian declines all treatment, the date of first course of treatment is the date this decision was made.

- If active surveillance (“watchful waiting”) was selected, record the date of that decision.

- Leave this item blank if the cancer was diagnosed at autopsy and not suspected prior to that.

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of first course of treatment is the year, month, day (YYYYMMDD) of the beginning of treatment (surgery, radiation, systemic, or other therapy) at any facility. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.

Coding Examples

Code	Explanation
20110301	A patient has an incisional, core, or fine needle biopsy on February 12, 2011 and subsequently undergoes an excisional biopsy or radical surgical procedure on March 1, 2011. Record the date of the excisional biopsy or radical surgery (March 1, 2011) as the date of first course of treatment. Do not record the date of the incisional, core, or fine needle biopsies as the date of first course of treatment.
201107	Admitting note stated the patient was diagnosed in June 2011, and treated in July 2011. Code as July 2011.
20110804	A patient has an excisional biopsy on August 11, 2011 followed by a radical surgical procedure on September 18, 2011. Record the date of the excisional biopsy (August 11, 2011) as the date of first course of treatment.
20110421	A patient begins receiving preoperative radiation therapy on April 21, 2011 and subsequent surgical therapy on June 2, 2011. Record the date of the preoperative radiation therapy (April 21, 2011) as the date of first course of treatment.
201101	A patient is diagnosed with cancer at your facility and receives radiation therapy in January 2011 at another facility before returning to your facility for surgery on February 2, 2011. Record the date of the radiation therapy (January 2011) as the date of first course of treatment, since the exact day of treatment is unknown.

DATE OF FIRST COURSE OF TREATMENT FLAG

Item Length: 2

NAACCR Item #1261

NAACCR Name: Date 1st Crs Rx COC Flag

XML NAACCR ID: date1stCrsRxCocFlag

Definition:

This flag explains why no appropriate value is in the field, Date of 1st Crs RX. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if Date of First Course Treatment has a full or partial date recorded.
- Assign code 10 when it is unknown whether any treatment was administered.
- Assign code 11 when the initial diagnosis was at autopsy.
- Assign code 12 if the Date of First Course Treatment cannot be determined, and the patient did receive first course treatment.
- Use code 12 if a decision not to treat was made or a decision to use active surveillance was made, but the date when the decision was made is totally unknown.

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any treatment was given)
11	No proper value is applicable in this context (for example, autopsy only)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, treatment was given but the date is unknown).
(blank)	A valid date value is provided in item Date of First Course of Treatment

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for Date of First Course of Treatment
Blank	Partial date is known (YYYYMM or YYYY) for Date of First Course of Treatment
10	Unknown if any treatment given
11	Diagnosed at autopsy
12	Date is completely unknown for Date of First Course of Treatment

RX SUMM – SURGERY PRIMARY SITE

Item Length: 2
NAACCR Item #1290

NAACCR Name: Rx Summ – Surg Prim Site
XML NAACCR ID: rxSummSurgPrimSite

Description

Site-specific codes for the type of surgery to the primary site performed as part of the first course of treatment. This includes treatment given at all facilities as part of the first course of treatment.

Coding Instructions

- Site-specific surgery codes for this data item are found in Appendix A of *STORE* 2018 (https://www.facs.org/-/media/files/quality-programs/cancer/ncdb/store_manual_2021.ashx) beginning on page 351 of the pdf.
- Document the most invasive surgical procedure for the primary site.
- For codes 00 through 79, the response positions are hierarchical. Last-listed responses take precedence over responses written above. Use codes 80 and 90 only if more precise information about the surgery is not available.
- Use code 98 for any cases codes to primary site C420, C421, C423, C424, C760-C768, C809
- Biopsies that remove the entire tumor and/or leave only microscopic margins are to be coded in this item.
- If a needle biopsy is done prior to an excisional biopsy or more extensive surgery, and no tumor remains in the specimen from the excisional biopsy or more extensive surgery, DO NOT consider the needle biopsy as an excisional biopsy.
- Surgery to remove regional tissue or organs is coded in item only if the tissue/organs are removed in continuity with the primary site, except where noted in Appendix A of *STORE*.
- If a previous surgical procedure to remove a portion of the primary site is followed by surgery to remove the remainder of the primary site, then code the total or final results.

Code	Definition
00	No surgical procedure of primary site. Diagnosed at autopsy.
10-19	Tumor destruction, no pathological specimen produced. Refer to Appendix A of <i>STORE</i> for the correct site-specific code for the procedure.
20-80	Refer to Appendix A of <i>STORE</i> for the correct site specific code for the procedure.
90	A surgical procedure to the primary site was done, but no information on the type of surgical procedure is provided.
98	Special code. Refer to Appendix A of <i>STORE</i> for the correct site-specific code for the procedure.
99	Patient record does not state whether a surgical procedure of the primary site was performed and no information is available. Death certificate only.

RX DATE OF FIRST SURGICAL PROCEDURE

Item Length: 8

NAACCR Item #1200

NAACCR Name: Rx Date Surgery

XML NAACCR ID: rxDateSurgery

Definition

Records the earliest date on which any first course surgical procedure was performed.

Coding Instructions

- Record the date of the first surgical procedure of the types coded as *Surgical Procedure of Primary Site* (NAACCR Item #1290), *Scope of Regional Lymph Node Surgery* (NAACCR Item #1292) (**excluding code 1**) or *Surgical Procedure/Other Site* (NAACCR Item #1294) performed at this or any facility.
- If a biopsy or aspirate of a regional node is the first surgical procedure or the only surgical procedure, do not code the date of that procedure in this field.
- The date recorded here may be the same as the date recorded in *Date of Most Definitive Surgical Resection of the Primary Site* if the patient only received one surgical procedure and it was a resection of the primary site.
- If surgery is the first or only treatment administered to the patient, then the date of surgery should be the same as the date entered into the item *Date of First Course of Treatment* (NAACCR Item #1270).

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of first surgical procedure is the year, month and day (YYYYMMDD) of the procedure at this or any facility. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.

Examples:

Code	Definition
20100402	Patient had a lumpectomy done April 2, 2010 followed by a MRM on April 27, 2010.
201105	Patient underwent an excisional biopsy in May 2011 for melanoma followed by a wide excision at your facility June 10, 2011
20100308	Patient had a needle aspiration of metastatic axillary lymph node March 8, 2010 followed by a MRM for breast cancer April 1, 2010.

RX DATE – SURGERY FLAG

Item Length: 2
NAACCR Item #1201
NAACCR Name: Rx Date Surgery Flag
XML NAACCR ID: rxDateSurgeryFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-Surgery. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Surgery has a full or partial date recorded.
- Assign code 10 when it is unknown whether any surgery was performed.
- Assign code 11 if no surgical procedure was performed.
- Assign code 12 if RX Date – Surgery cannot be determined, but it is known that surgery was performed as part of first course of treatment.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any surgery performed)
11	No proper value is applicable in this context (for example, no surgery performed)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, surgery was performed but the date is unknown).
(blank)	A valid date value is provided in item RX Date-Surgery.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Surgery
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Surgery
10	Unknown if any surgery was performed
11	No surgery performed
12	Surgery performed as first course treatment but date is completely unknown.

RX DATE – MOST DEFINITIVE SURGERY

Item Length: 8

NAACCR Item #3170

NAACCR Name: Rx Date Mst Defn Srg

XML NAACCR ID: rxDateMostDefinSurg

Definition

Records the date that the most definitive surgical procedure on the primary site was performed as part of first course treatment. This item is used to measure the lag time between diagnosis and the most definitive surgery performed on the primary site.

Coding Instructions

- Record the date that the procedure coded as *Surgical Procedure of Primary Site* (NAACCR Item #1290) was performed at this or any facility.
- The date recorded here may be the same as the date recorded in *RX-Date First Surgical Procedure* if the patient only received one surgical procedure and it was a resection of the primary site.

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of first surgical procedure is the year, month and day (YYYYMMDD) of the procedure at this or any facility. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.

Examples:

Code	Definition
20100427	Patient had a lumpectomy done April 2, 2010 followed by a MRM on April 27, 2010.
20110610	Patient underwent an excisional biopsy in May 2011 for melanoma followed by a wide excision at your facility June 10, 2011
2010401	Patient had a needle aspiration of metastatic axillary lymph node March 8, 2010 followed by a MRM for breast cancer April 1, 2010.

RX DATE - MOST DEFINITIVE SURGERY FLAG

Item Length: 2

NAACCR Item #3171

NAACCR Name: Rx Date Mst Defn Srg Flag

XML NAACCR ID: rxDateMostDefinSurgFlag

Definition

This flag explains why no appropriate value is in the field, *RX Date-Most Definitive Surgery*. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date – Most Definitive Surgery has a full or partial date recorded.
- Assign code 10 when it is unknown whether any surgery was performed.
- Assign code 11 if no surgical procedure was performed.
- Assign code 12 if RX Date – Most Definitive Surgery cannot be determined, but it is known that surgery was performed as part of first course of treatment.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any surgery performed)
11	No proper value is applicable in this context (for example, no surgery performed)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, surgery was performed but the date is unknown).
(blank)	A valid date value is provided in item RX Date-Most Definitive Surgery. Case diagnosed prior to January 1, 2003

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Surgery
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Surgery
10	Unknown if any surgery was performed
11	No surgery performed
12	Surgery performed as first course treatment but date is completely unknown.

Definition:

Scope of Regional Lymph Node Surgery describes the procedure of removal, biopsy, or aspiration of **regional** lymph nodes performed during the initial work-up or first course of therapy. Instructions for coding **sentinel lymph node biopsies** (SLNBx) have been clarified for 2012 and later, diagnoses. This data item can be used to evaluate the extent of surgery and should be collected even if a surgery of the primary site was not performed.

Coding Instructions:

- Use the operative report as the primary source document to determine whether the operative procedure was a SLNBx, or a more extensive dissection of regional lymph nodes, or a combination of both SLNBx and regional lymph node dissection. The operative report will designate the surgeon's planned procedure as well as a description of the procedure that was actually performed. The pathology report may be used to complement the information appearing in the operative report, but the operative report takes precedence when attempting to distinguish between SLNBx and regional lymph node dissection or a combination of these two procedures. Do not use the number of lymph nodes removed and pathologically examined as the sole means of distinguishing between a SLNBx and a regional lymph node dissection.
- Code regional lymph node procedures in this data item. Record distant lymph node removal in Surgical Procedure of Other Site.
 - Include lymph nodes that are regional in the current AJCC Staging Manual or EOD 2018
- Code regional lymph node procedures in this data item. Record distant lymph node removal in Surgical Procedure of Other Site.
- Code the procedure that is numerically higher. Codes 0-7 are hierarchical.
- Record all surgical procedures that remove, biopsy, or aspirate regional lymph node(s) whether or not there were any surgical procedures of the primary site. The regional lymph node surgical procedure(s) may be done to diagnose cancer, stage the disease, or as part of the initial treatment.
- Include lymph nodes obtained or biopsied during any procedure within the first course of treatment. A separate lymph node surgery is not required.
 - Code the removal of intra-organ lymph nodes in Scope Regional LN Surgery.

- Add the number of all of the lymph nodes removed during each surgical procedure performed as part of the first course of treatment. The Scope of Regional Lymph Node field is cumulative.
 - Lymph node aspirations
 - Do not double-count when a regional lymph node is aspirated and that node is in the resection field. Do not add the aspirated node to the total number.
 - Count as an additional node when a regional lymph node is aspirated and that node is NOT in the resection field. Add it to the total number.
- Code the removal of regional nodes for both primaries when the patient has two primaries with common regional lymph nodes.

Example: Patient has a cystoprostatectomy and pelvic lymph node dissection for bladder cancer. Pathology identifies prostate cancer as well as bladder cancer and 4/21/ nodes positive for metastatic adenocarcinoma. Code Scope of Regional Lymph Node Surgery to 5 (4 or more regional lymph nodes removed) for both primaries

- Assign code 0 when
 - Regional lymph node removal procedure was not performed. This excludes all sites and histologies that would be coded 9. OR
 - First course of treatment was active surveillance/watchful waiting. OR
 - The operative report lists a lymph node dissection, but no nodes were found by the pathologist.
- Assign code 2 when
 - The operative report states that a SLNBx was performed, OR
 - The operative report describes a procedure using injection of a dye, radio label, or combination to identify a lymph node (possibly more than one) for removal/examination.

Note: When a SLNBx is performed, additional non-sentinel nodes can be taken during the same operative procedure. These additional non-sentinel nodes may be discovered by the pathologist or selectively removed (or harvested) as part of the SLNBx procedure by the surgeon. Code this as a SLNBx (code 2). If review of the operative

report confirms that a regional lymph node dissection followed the SLNBx, code these cases as 6.

- Codes 3, 4, and 5: The operative report states that a regional lymph node dissection was performed (a SLNBx was not done during this procedure or in a prior procedure).
 - Code 3: Check the operative report to ensure this procedure is not a SLNBx only (code 2) or a SLNBx with a regional lymph node dissection (code 6 or 7).
 - Code 4 should be used infrequently. Review the operative report to ensure the procedure was not a SLNBx only.
 - Code 5: If a relatively small number of nodes was examined pathologically, review the operative report to confirm the procedure was not a SLNBx only (code 2). If a relatively large number of nodes was examined pathologically, review the operative report to confirm that there was not a SLNBx in addition to a more extensive regional lymph node dissection during the same, or separate, procedure (code 6 or 7).

Note: Infrequently, a SLNBx is attempted and the patient fails to map (i.e., no sentinel lymph nodes are identified by the dye and/or radio label injection). When mapping fails, surgeons usually perform a more extensive dissection of regional lymph nodes. Code these cases as 2 if no further dissection of regional lymph nodes was undertaken, or 6 when regional lymph nodes were dissected during the same operative event.

- Code 6: SLNBx and regional lymph node dissection (code 3, 4, or 5) during the same surgical event, or timing not known
 - Generally, SLNBx followed by a regional lymph node completion will yield a relatively large number of nodes. However it is possible for these procedures to harvest only a few nodes.
 - If relatively few nodes are pathologically examined, review the operative report to confirm whether the procedure was limited to a SLNBx only.
 - Infrequently, a SLNBx is attempted and the patient fails to map (i.e., no sentinel lymph nodes are identified by the dye and/or radio label injection.) When mapping fails, the surgeon usually performs a more extensive dissection of regional lymph nodes. Code these cases as 6.
- Code 7: SLNBx and regional lymph node dissection (code 3, 4, or 5) in separate surgical events.

- Generally, SLNBx followed by a regional lymph node completion will yield a relatively large number of nodes. However it is possible for these procedures to harvest only a few nodes.
- If relatively few nodes are pathologically examined, review the operative report to confirm whether the procedure was limited to a SLNBx only.
- Assign code 9 for
 - Primary sites
 - Brain (C700-C709) OR
 - Spinal cord (C710-C719) OR
 - Cranial nerves and other parts of the central nervous system (C720-C729, C751-C753) OR
 - Unknown or ill-defined sites (C760-C768, C809) (all histologies)
 - Lymphoma with primary site in lymph nodes (C770-C779) AND
 - 9590-9597 OR
 - 9650-9719 OR
 - 9724-9738
 - Hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative disease
 - Primary sites: C420, C421, C423 or C424 (all histologies)
 - Histologies: 9750, 9760-9764, 9800-9820, 9826, 9831-9920, 9931-9992 (all sites)

Coding Instructions – Sentinel lymph node biopsy (SLNBx), breast primary C500-C509

- Use the operative report as the primary source document to determine whether the operative procedure was a SLNBx, an axillary node dissection (ALND), or a combination of both SLNBx and ALND. The operative report will designate the surgeon's planned procedure as well as a description of the procedure that was actually performed. The pathology report may be used to complement the information appearing in the operative report, but the operative report takes precedence when

attempting to distinguish between SLNBx and ALND, or a combination of these two procedures. Do not use the number of lymph nodes removed and pathologically examined as the sole means of distinguishing between a SLNBx and an ALND.

- Code 1
 - Excisional biopsy or aspiration of regional lymph nodes for breast cancer is uncommon. Review the operative report to confirm whether an excisional biopsy or aspiration of regional lymph nodes was actually performed; it is highly possible that the procedure is a SLNBx (code 2) instead. If additional procedures were performed on the lymph nodes, such as axillary lymph node dissection, use the appropriate code 2-7.

- Code 2
 - If a relatively large number of lymph nodes, more than 5, are pathologically examined, review the operative report to confirm the procedure was limited to a SLNBx and did not include an axillary lymph node dissection (ALND).
 - Infrequently, a SLNBx is attempted and the patient fails to map (i.e., no sentinel lymph nodes are identified by the dye and/or radio label injection) and no sentinel nodes are removed. Review the operative report to confirm that an axillary incision was made and a node exploration was conducted. Patients undergoing SLNBx who fail to map will often undergo ALND. Use code 2 if no ALND was performed, or 6 when ALND was performed during the same operative event. Enter the appropriate number of nodes examined and positive in the data items Regional Lymph Nodes Examined (NAACCR Item #830) and Regional Lymph Nodes Positive (NAACCR Item #820).

- Code 3, 4, and 5: Generally, ALND removes at least 7-9 nodes. However, it is possible for these procedures to remove or harvest fewer nodes. Review the operative report to confirm that there was not a SLNBx in addition to a more extensive regional lymph node dissection during the same procedure (code 6 or 7).

- Code 6
 - Generally, SLNBx followed by ALND will yield a minimum of 7-9 nodes. However it is possible for these procedures to harvest fewer (or more) nodes.
 - If relatively few nodes are pathologically examined, review the operative report to confirm whether the procedure was limited to a SLNBx, or whether a SLNBx plus an ALND was performed.

- Code 7
 - Generally, SLNBx followed by ALND will yield a minimum of 7-9 nodes. However, it is possible for these procedures to harvest fewer (or more) nodes.
 - If relatively few nodes are pathologically examined, review the operative report to confirm whether the procedure was limited to a SLNBx only, or whether a SLNBx plus an ALND was performed.

Code	Definition
0	None. No regional lymph node surgery. No lymph nodes found in the pathologic specimen or diagnosed at autopsy.
1	Biopsy or aspiration of regional lymph nodes, NOS. Biopsy or aspiration of regional lymph node(s) regardless of the extent of involvement of disease.
2	Sentinel lymph node biopsy. Biopsy of the first lymph node or nodes that drain a defined area of tissue within the body. Sentinel node(s) are identified by the injection of a dye or radio label at the site of the primary tumor.
3	Number of regional nodes removed unknown or not stated; regional lymph nodes removed, NOS. Sampling or dissection of regional lymph node(s) and the number of nodes removed is unknown or not stated. The procedure is not specified as sentinel node biopsy.
4	1–3 regional lymph nodes removed. Sampling or dissection of regional lymph node(s) with fewer than four lymph nodes found in the specimen. The procedure is not specified as sentinel node biopsy.
5	4 or more regional lymph nodes removed. Sampling or dissection of regional lymph nodes with at least four lymph nodes found in the specimen. The procedure is not specified as a sentinel node biopsy.
6	Sentinel node biopsy and code 3, 4, or 5 at the same time, or timing not stated. Code 2 was performed in a single surgical event with code 3, 4, or 5. Or, code 2 and 3, 4, or 5 were performed, but timing was not stated in the patient record.
7	Sentinel node biopsy and code 3, 4, or 5 at different times. Code 2 was followed in a subsequent surgical event by procedures coded as 3, 4, or 5.
9	Unknown or not applicable. It is unknown whether regional lymph node surgery was performed; death certificate only; for lymphomas with a lymph node primary site; an unknown or ill-defined primary; or for hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative disease.

Coding Examples:

Code	Explanation
0	If an attempt was made to remove regional nodes but no nodes were found in the pathological specimen, code as not done.
1	Aspiration of regional lymph node to confirm histology of widely metastatic disease.
2	There was an attempt at a sentinel lymph node biopsy for a breast cancer, but no nodes were found in the pathological specimen.
3	Pelvic lymph node dissection for prostate cancer
6	Sentinel lymph node biopsy of right axilla, followed by right axillary lymph node dissection during a lumpectomy procedure.
9	If all you know is that the patient had surgery but do not code if lymph nodes were removed.

REGIONAL NODES EXAMINED

Item Length: 2

NAACCR Item #830

NAACCR Name: Regional Nodes Examined

XML NAACCR ID: regionalNodesExamined

Description

This field records the total number of regional lymph nodes that were removed and examined by the pathologist.

Instructions for Coding

- **Regional lymph nodes only.** Record information about only regional lymph nodes in this field.
- This field is based on pathologic information only. This field is to be recorded regardless of whether the patient received preoperative treatment.
- **Use of code 00.** Code 00 may be used in several situations.
 - When the assessment of lymph nodes is clinical.
 - When no lymph nodes are removed or examined.
 - When a “dissection” of a lymph node drainage area is found to contain no lymph nodes at the time of pathologic examination.
 - If Regional Nodes Examined is coded 00, Regional Nodes Positive is coded as 98.
- **Cumulative nodes removed and examined.** Record the total number of regional lymph nodes removed and examined by the pathologist.
 - The number of regional lymph nodes examined is cumulative from all procedures that removed lymph nodes through the completion of surgeries in the first course of treatment with the exception of aspiration or core biopsies coded to 95.
 - Do not count a positive aspiration or core biopsy of a lymph node in the same lymph node chain removed at surgery as an additional node in Regional Nodes Examined.
 - If the positive aspiration or core biopsy is from a node in a different node region, include the node in the count of Regional Nodes Examined.

- If the location of the lymph node that is aspirated or core-biopsied is not known, assume it is part of the lymph node chain surgically removed, and do not include it in the count of Regional Nodes Examined.
- When neither the type of lymph node removal procedure nor the number of lymph nodes examined is known, use code 98.
- **Priority of lymph node counts.** If there is a discrepancy regarding the number of regional lymph nodes examined, use information in the following priority:
 - Final diagnosis
 - Synoptic report (also known as CAP protocol or pathology report checklist)
 - Microscopic description
 - Gross description
- **Use of code 95.** Use code 95 when the only procedure for regional lymph nodes is a needle aspiration (cytology) or core biopsy (tissue).
- **Lymph node biopsy.** If a lymph node biopsy was performed, code the number of nodes removed, if known. If the number of nodes removed by biopsy is not known, use code 96.
- **Definition of “sampling” (code 96).** A lymph node “sampling” is removal of a limited number of lymph nodes. Other terms for removal of a limited number of nodes include lymph node biopsy, berry picking, sentinel lymph node procedure, sentinel node biopsy, selective dissection. Use code 96 when a limited number of nodes are removed but the number is unknown,
- **Definition of “dissection” (code 97).** A lymph node “dissection” is removal of most or all of the nodes in the lymph node chain(s) that drain the area around the primary tumor. Other terms include lymphadenectomy, radical node dissection, lymph node stripping. Use code 97 when more than a limited number of lymph nodes are removed and the number is unknown.
- **Multiple lymph node procedures.** If both a lymph node sampling and a lymph node dissection are performed and the total number of lymph nodes examined is unknown, use code 97.

▪ **Use of code 99.**

- Any case coded to primary site C420, C421, C423, C424, C589, C700-C709, C710-C729, C751-C753, C761-C768, C770-C779, or C809
- Lymphoma (excluding CLL/SLL) 00790
- Lymphoma (CLL/SLL) 00795
- Plasma Cell Disorders (excluding 9734/3) 00822
- Cases with no information about number of regional lymph nodes examined
- For the following schemas, the Regional Nodes Positive field is always coded as 99.

Placenta

Brain and Cerebral Meninges

Other Parts of Central Nervous System

Intracranial Gland

Hematopoietic, Reticuloendothelial, Immunoproliferative and Myeloproliferative Neoplasms

Hodgkin and non-Hodgkin Lymphoma

- Excludes cases collected in the following schemas: Lymphoma Ocular Adnexa, Primary Cutaneous Lymphomas and Mycosis Fungoides

Myeloma and Plasma Cell Disorders

- Excludes histology 9734

Other and Ill-Defined Primary Sites

- Excludes Spleen (C422)

Unknown Primary Site

Code	Explanation
00	No nodes examined
01-89	1 to 89 nodes examined (code the exact number of regional lymph nodes examined.)
90	90 or more nodes examined
95	No regional nodes removed, but aspiration or core biopsy of regional nodes performed.
96	Regional lymph node removal documented as a sampling, and the number of nodes unknown/not stated
97	Regional lymph node removal documented as dissection, and the number of nodes unknown/not stated
98	Regional lymph nodes surgically removed, but number of lymph nodes unknown/not stated and not documented as sampling or dissection; nodes examined, but the number unknown.

Code	Explanation
99	Unknown whether nodes were examined; not applicable or negative; not documented in patient record

Examples:

Code	Explanation
11	Lung cancer patient has a mediastinoscopy and positive core biopsy of a hilar lymph node. Patient then undergoes right upper lobectomy that yields 3 hilar and 2 mediastinal nodes positive out of 11 nodes dissected. Code Regional Nodes Positive as 05 and Regional Nodes Examined as 11 because the core biopsy was of a lymph node in the same chain as the nodes dissected.
06	Positive right cervical lymph node aspiration followed by right cervical lymph node dissection showing 1 of 6 nodes positive. Code Regional Nodes Positive as 01 and Regional Nodes Examined as 06.
09	Breast cancer patient has a positive core biopsy of a supraclavicular node and an axillary dissection showing 3 of 8 nodes positive. Code Regional Nodes Positive as 04 and Regional Nodes Examined as 09 because the supraclavicular lymph node is in a different, but still regional, lymph node chain.
14	Patient record states that core biopsy was performed at another facility and 7/14 regional lymph nodes were positive at the time of resection. Code Regional Nodes Positive as 07 and Regional Nodes Examined as 14.
11	A breast case is two separate primaries as determined by the SEER multiple primary rules. The pathology report states “3 of 11 lymph nodes positive for metastasis” with no further information available. Code Regional Nodes Positive as 03 and Regional Nodes Examined as 11 for both primaries.
95	Patient with esophageal cancer. Enlarged mid-esophageal node found on CT scan, which is aspirated and found to be positive. Patient undergoes radiation therapy and no surgery. Code Regional Nodes Positive as 95 and Regional Nodes Examined as 95.
06	Lung cancer patient has aspiration of suspicious hilar mass, which shows metastatic squamous carcinoma in lymph node tissue. Patient undergoes preoperative radiation therapy followed by lobectomy showing 6 negative hilar lymph nodes. Code Regional Nodes Positive as 95 and Regional Nodes Examined as the 06 nodes surgically resected.
10	Patient with carcinoma of the pyriform sinus has a mass in the mid neck. Fine needle aspiration (FNA) of one node is positive. The patient has neoadjuvant chemotherapy, then resection of the primary tumor and a radical neck dissection. In the radical neck dissection “several” of 10 nodes are positive; the remainder of the nodes show chemotherapy effect. Code Regional Nodes Positive as 97 because the total number of positive nodes biopsied and removed is unknown, and code Regional Nodes Examined as 10.

REGIONAL NODES POSITIVE

Item Length: 2
NAACCR Item #820
NAACCR Name: Regional Nodes Positive
XML NAACCR ID: regionalNodesPositive

Description

This field records the exact number of regional lymph nodes examined by the pathologist and found to contain metastases.

Instructions for Coding

- **Regional lymph nodes only.** Record information about only regional lymph nodes in this field. Distant lymph node information should be coded in the “CS Mets at Dx” field.
- This field is based on pathologic information only. This field is to be recorded regardless of whether the patient received preoperative treatment.
- True in situ cases cannot have positive lymph nodes, so the only allowable codes are 00 (negative) or 98 (not examined). Codes 01-97 and 99 are not allowed.
- **Nodes positive is cumulative.** Record the total number of regional lymph nodes removed and found to be positive by pathologic examination. Record lymph nodes removed and found to be positive during an autopsy for autopsy-only cases.
 - The number of regional lymph nodes positive is cumulative from all procedures that remove lymph nodes through the completion of surgeries in the first course of treatment.
 - Do not count a positive aspiration or core biopsy of a lymph node in the same lymph node chain removed at surgery as an additional node in Regional Nodes Positive when there are positive nodes in the resection. In other words, if there are positive regional lymph nodes in a lymph node dissection, do not count the core needle biopsy or the fine needle aspiration if it is in the same chain.
 - Include the nodes in the count of Regional Nodes Positive when the positive aspiration or core biopsy is from a node in a different node region..
 - Assume the lymph node that is core-biopsied or aspirated is part of the lymph node chain surgically removed and do not include it in the count of Regional Nodes Positive when its location is not know.
- **Priority of lymph node counts.** If there is a discrepancy regarding the number of positive lymph nodes, use information in the following priority:

- Final diagnosis
- Synoptic report (also known as CAP protocol or pathology report checklist)
- Microscopic description
- Gross description
- **Positive nodes in multiple primaries in same organ.**
 - Determine the histology of the metastases in the nodes and code the nodes as positive for the primary with that histology when there are multiple primary cancers with different histologic types in the same organ and the pathology report just states the number of nodes positive.
 - Code the nodes as positive for all primaries when no further information is available.
- **Isolated Tumor Cells (ITCs) in lymph nodes**
 - For all cases **except cutaneous melanoma and Merkel cell carcinoma of skin**
 - Count only lymph nodes that contain micrometastases or larger (metastases greater than 0.2 millimeters in size)
 - Assume the metastases are larger than 0.2 mm and count the lymph node(s) as positive when the path report indicates that nodes are positive but the size of metastasis is not stated.
 - Do not include in the count of lymph nodes positive any nodes that are identified as containing ITCs.
 - For cutaneous melanoma and Merkel cell carcinoma of skin
 - Count nodes with ITCs as positive lymph nodes
- **Use of code 95.** Use code 95 when the only procedure for regional lymph nodes is a needle aspiration (cytology) or core biopsy (tissue).
 - Use code 95 when a positive lymph node is aspirated and there are no surgically resected lymph nodes.

- Use code 95 when a positive lymph node is aspirated and surgically resected lymph nodes are negative.
- **Definition of code 97.** Use code 97 for any combination of positive aspirated, biopsied, sampled or dissected lymph nodes if the number of involved nodes cannot be determined on the basis of cytology or histology. Code 97 includes positive lymph nodes diagnosed by either cytology or histology.

Note: If the aspirated node is the only one that is microscopically positive, use code 95.

- **Use of code 98.** Code 98 may be used in several situations.
 - When the assessment of lymph nodes is clinical only.
 - When no lymph nodes are removed and examined.
 - When a “dissection” of a lymph node drainage area is found to contain no lymph nodes at the time of pathologic examination.
 - If Regional Nodes Positive is coded as 98, Regional Nodes Examined is usually coded 00.
- **Use of code 99.**
 - Any case coded to primary site C420, C421, C423, C424, C589, C700-C709, C710-C729, C751-C753, C761-C768, C770-C779, or C809
 - Lymphoma (excluding CLL/SLL) 00790
 - Lymphoma (CLL/SLL) 00795
 - Plasma Cell Disorders (excluding 9734/3) 00822
 - Cases with no information about positive regional lymph nodes
 - For the following schemas, the Regional Nodes Positive field is always coded as 99.

Placenta
 Brain and Cerebral Meninges
 Other Parts of Central Nervous System
 Intracranial Gland
 Hematopoietic, Reticuloendothelial, Immunoproliferative and
 Myeloproliferative Neoplasms

Hodgkin and non-Hodgkin Lymphoma

- Excludes cases collected in the following schemas: Lymphoma Ocular Adnexa, Primary Cutaneous Lymphomas and Mycosis Fungoides

Myeloma and Plasma Cell Disorders

- Excludes histology 9734

Other and Ill-Defined Primary Sites

- Excludes Spleen (C422)

Unknown Primary Site

Code	Explanation
00	All nodes examined are negative
01-89	1-89 nodes are positive (code exact number of nodes positive)
90	90 or more nodes are positive
95	Positive aspiration of lymph node(s) was performed
97	Positive nodes are documented, but the number is unspecified
98	No nodes were examined
99	It is unknown whether nodes are positive; not applicable; not stated in patient record

Examples:

Code	Explanation
05	Lung cancer patient has a mediastinoscopy and positive core biopsy of a hilar lymph node. Patient then undergoes right upper lobectomy that yields 3 hilar and 2 mediastinal nodes positive out of 11 nodes dissected. Code Regional Nodes Positive as 05 and Regional Nodes Examined as 11 because the core biopsy was of a lymph node in the same chain as the nodes dissected.
01	Positive right cervical lymph node aspiration followed by right cervical lymph node dissection showing 1 of 6 nodes positive. Code Regional Nodes Positive as 01 and Regional Nodes Examined as 06.
04	Breast cancer patient has a positive core biopsy of a supraclavicular node and an axillary dissection showing 3 of 8 nodes positive. Code Regional Nodes Positive as 04 and Regional Nodes Examined as 09 because the supraclavicular lymph node is in a different, but still regional, lymph node chain.
07	Patient record states that core biopsy was performed at another facility and 7/14 regional lymph nodes were positive at the time of resection. Code Regional Nodes Positive as 07 and Regional Nodes Examined as 14.
03	A breast case is two separate primaries as determined by the SEER multiple primary rules. The pathology report states “3 of 11 lymph nodes positive for metastasis” with no further information available. Code Regional Nodes Positive as 03 and Regional Nodes Examined as 11 for both primaries.
95	Patient with esophageal cancer. Enlarged mid-esophageal node found on CT scan, which is aspirated and found to be positive. Patient undergoes radiation therapy and no surgery. Code Regional Nodes Positive as 95 and Regional Nodes Examined as 95.

Code	Explanation
95	Lung cancer patient has aspiration of suspicious hilar mass, which shows metastatic squamous carcinoma in lymph node tissue. Patient undergoes preoperative radiation therapy followed by lobectomy showing 6 negative hilar lymph nodes. Code Regional Nodes Positive as 95 and Regional Nodes Examined as the 06 nodes surgically resected.
97	Patient with carcinoma of the pyriform sinus has a mass in the mid neck. Fine needle aspiration (FNA) of one node is positive. The patient has neoadjuvant chemotherapy, then resection of the primary tumor and a radical neck dissection. In the radical neck dissection “several” of 10 nodes are positive; the remainder of the nodes show chemotherapy effect. Code Regional Nodes Positive as 97 because the total number of positive nodes biopsied and removed is unknown, and code Regional Nodes Examined as 10.

RX SUMM—SURG OTH REG/DIS

Item Length: 1

NAACCR Item #1294

NAACCR Name: Rx Summ - Surg Other Reg/Dis

XML NAACCR ID: rxSummSurgOthRegDis

Definition:

Records the surgical removal of distant lymph nodes or other tissue(s)/organ(s) beyond the primary site.

Coding Instructions:

- Do not code tissue or organs such as an appendix that were removed incidentally, and the organ was not involved with cancer

Note: Incidental removal of organs means that tissue was removed for reasons other than removing cancer or preventing the spread of cancer. Examples of incidental removal of organ(s) would be removal appendix, gallbladder, etc., during abdominal surgery.

- Do not code removal of uninvolved contralateral breast in this data item. The removal of the uninvolved contralateral breast is included in the surgery of the primary site codes.
- Assign code 0 when
 - No surgical procedures were performed that removed distant lymph node(s) or other tissue(s) or organ(s) beyond the primary site, or
 - First course of treatment was active surveillance/watchful waiting.
- The codes are hierarchical
 - Codes 1-5 have priority over codes 0 and 9
- Assign code 1
 - When the involved contralateral breast is removed for a **single** primary breast cancer
 - When any surgery is performed to remove tumors for any case coded to primary site C420, C421, C423, C424, C760-C768, C770-C779, or C809
 - Excluding cases coded to the schema Cervical Lymph Nodes and Unknown Primary 00060
- Assign code 2 for sites that are regional. Include sites that are regional in the current AJCC Staging Manual or EOD 2018.

- Assign code 4 for sites that are distant. Include sites that are distant in the current AJCC Staging Manual or EOD 2018.
- Assign code 9 for death certificate only (DCO) cases.

Code	Definition
0	None. No surgical procedure of non-primary site was performed or diagnosed at autopsy.
1	Non-primary surgical procedure performed. Non-primary surgical resection to other site(s), unknown if whether the site(s) is regional or distant.
2	Non-primary surgical procedure to other regional sites. Resection of regional site.
3	Non-primary surgical procedure to distant lymph node(s). Resection of distant lymph node(s).
4	Non-primary surgical procedure to distant site. Resection of distant site.
5	Combination of codes. Any combination of surgical procedures 2, 3, or 4.
9	Unknown. It is unknown whether any surgical procedure of a non-primary site was performed or death certificate only.

Examples:

Code	Explanation
0	Removal of a small portion of the duodenum during a right hemicolectomy.
1	Surgical excision of a metastatic abdominal mass; unknown primary.
2	Surgical ablation of solitary liver metastasis, hepatic flexure primary.
4	Removal of liver mets for a lung cancer.
5	Excision of brain mets and an axillary lymph node for a lung primary.

REASON FOR NO SURGERY

Item Length: 1

NAACCR Item #1340

NAACCR Name: Reason for No Surgery

XML NAACCR ID: reasonForNoSurgery

Definition:

Records the reason that no surgery was performed on the primary site as part of the first course of treatment.

Coding Instructions:

- Assign code 0 when Surgery of Primary Site is coded in the range of 10-90 (surgery of the primary site was performed)
- Code 1 if Surgical Procedure of Primary Site is coded 98 (not applicable).
- If Surgical Procedure of Primary Site is coded 00, then assign a code in the range of 1-8.
 - Referral to a surgeon is equivalent to a recommendation for surgery.
 - Assign code 1 when
 - There is no information in the patient's medical record about surgery AND
 - It is known that surgery is not usually performed for this type and/or stage of cancer OR
 - There is no reason to suspect that the patient would have had surgery of primary site.
 - The treatment plan offered multiple options and the patient selected treatment that did not include surgery of the primary site
 - Surgery was part of the first course of treatment but was cancelled due to complete response to radiation and/or systemic therapy
 - The patient elected to pursue no treatment following the discussion of surgery. Discussion does not equal a recommendation. Patient's decision not to pursue surgery is not a refusal of surgery in this situation.
 - Watchful waiting/active surveillance is the first course (e.g., prostate)

- Assign code 6 when
 - It is known that surgery was recommended AND
 - It is known that surgery was not performed AND
 - There is no documentation explaining why surgery was not done.
- Assign code 7 when the patient
 - Refuses recommended surgery OR
 - Makes a blanket statement that he/she refused all treatment when surgery is a customary option according to NCCN guidelines and/or NCI PDQ for the primary site/histology.
 - Assign code 1 when surgery is not normally performed for the site/histology

Note: Coding Reason for No Surgery of Primary Site as “refused” does not affect the coding of the other treatment data items (e.g., Radiation, Chemotherapy, Hormone Therapy, etc.) Code 7 means surgery is exactly what was recommended by the physician and the patient refused. If two treatment alternatives were offered and surgery was not chosen, code Reason No Surgery of Primary Site as 1 [Surgery of the primary site was not performed because it was not part of the planned first-course treatment].

- Assign code 8 when surgery is recommended, but it is unknown if the patient actually had the surgery.
- Assign code 9
 - When there is no documentation that surgery was recommended or performed
 - Autopsy only cases

Code	Definition
0	Surgery of the primary site was performed.
1	Surgery of the primary site was not performed because it was not part of the planned first course treatment.
2	Surgery of the primary site was not recommended/performed because it was contraindicated due to patient risk factors (comorbid conditions, advanced age, etc).
5	Surgery of the primary site was not performed because the patient died prior to planned or recommended surgery.
6	Surgery of the primary site was not performed; it was recommended by the patient's physician, but was not performed as part of the first course of therapy. No reason was noted in the patient record.
7	Surgery of the primary site was not performed; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
8	Surgery of the primary site was recommended, but it is unknown if it was performed. Further follow-up is recommended.
9	It is unknown whether surgery of the primary site was recommended or performed. Diagnosed at autopsy or death certificate only.

Examples:

Code	Explanation
1	Prostate cancer patient is offered a choice of prostatectomy or radiation and chooses radiation.
6	The medical record has a recommendation that the patient have surgery. No further admissions or documentation of surgery found; the primary care physician replies that the patient did NOT have surgery. No further information is given; it is unknown if the patient refused surgery or if there were co-morbid conditions that prevented the surgical procedure.
8	There is documentation in the medical record that the primary care physician referred the patient to a surgical oncologist. Follow-back to the surgical oncologist and primary care physician yields no further information. Assign code 8, it is known that surgery was recommended but there is no information on whether or not the patient actually had the surgical procedure.

RX DATE – RADIATION

Item Length: 8
NAACCR Item #1210
NAACCR Name: RX Date Radiation
XML NAACCR ID: rxDateRadiation

Description

Records the date on which radiation therapy began at any facility that is part of the first course of treatment.

The date radiation started will typically be found in the radiation oncologist's summary letter for the first course of treatment. Determination of the date radiation started may require assistance from the radiation oncologist for consistent coding.

Coding Instructions

- Record the data of the first/earliest radiation treatment if radiation was given and recorded as part of the first course of therapy
 - Do not record the date of the initial radiation planning session.
- If radiation therapy is the first or only treatment administered to the patient, then the date radiation started should be the same as the date entered into the item *Date of First Course of Treatment* (NAACCR Item #1270).
- The date when treatment started will typically be found in the radiation oncologist's summary letter for the first course of treatment.

Code	Description
YYYYMMDD YYYYMM YYYY	The year, month and day (YYYYMMDD) that the first course of radiation therapy began at any facility. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.

Coding Examples:

Code	Reason
20100412	A patient has external beam radiation on April 12, 2010
201009	If the exact date of the beginning of treatment is not available, then record an approximate date. For example, September 2010.

RX DATE – RADIATION FLAG

Item Length: 2

NAACCR Item #1211

NAACCR Name: RX Date Radiation Flag

XML NAACCR ID: rxDateRadiationFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-Radiation. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Radiation has a full or partial date recorded.
- Assign code 10 when it is unknown whether any radiation was given.
- Assign code 11 if no radiation is planned or given.
- Assign code 12 if RX Date – Radiation cannot be determined, but it is known that radiation was given as part of first course of treatment.
- Assign code 15 if radiation is planned, but has not yet started and the start date is not yet available.

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any radiation was given)
11	No proper value is applicable in this context (for example, no radiation was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, radiation was given but the date is unknown).
15	Information is not available at this time, but it is expected that it will be available later (for example, radiation therapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up).
(blank)	A valid date value is provided in item RX Date-Radiation.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Radiation
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Radiation
10	Unknown if any radiation given
11	No radiation given
12	Radiation given as first course treatment but date is completely unknown.
15	Radiation not yet started but planned

REASON FOR NO RADIATION

Item Length: 1

NAACCR Item #1430

NAACCR Name: Reason for No Radiation

XML NAACCR ID: reasonForNoRadiation

Definition:

Records the reason the patient did not receive radiation treatment as part of first course of therapy.

Coding Instructions

- Assign code 0 if the patient received regional radiation as part of first course of therapy.
- If Regional Treatment Modality is coded 00, then assign a code in the range of 1-8.
 - Referral to a radiation oncologist is equivalent to a recommendation for radiation.
 - Assign code 1 when
 - There is no information in the patient's medical record about radiation AND
 - It is known that radiation is not usually given for this type and/or stage of cancer OR
 - There is no reason to suspect that the patient would have had radiation.
 - The treatment plan offered multiple options and the patient selected treatment that did not include radiation
 - The patient elected to pursue no treatment following the discussion of radiation. Discussion does not equal a recommendation.
 - Watchful waiting/active surveillance (e.g., prostate)
 - Assign code 6 when
 - It is known that radiation was recommended AND
 - It is known that radiation was not given AND
 - There is no documentation explaining why radiation was not given.

- Assign code 7 when the patient
 - Refuses recommended radiation OR
 - Makes a blanket statement that he/she refused all treatment when radiation is a customary option for the primary site/histology.
 - Assign code 1 when radiation is not normally performed for the site/histology
- Assign code 8 when radiation is recommended, but it is unknown if the patient actually was given the radiation.
- Assign code 9
 - When there is no documentation that radiation was recommended or given
 - Autopsy only

Code	Explanation
0	Radiation therapy was administered
1	Radiation therapy was not administered because it was not part of the planned first-course treatment.
2	Radiation therapy was not administered because it was contraindicated due to patient risk factors (comorbid conditions, advanced age, etc.)
5	Radiation therapy was not administered because the patient died prior to planned or recommended treatment.
6	Radiation therapy was not administered; it was recommended by the patient's physician, but was not administered as part of the first-course therapy. No reason was noted in the patient's record.
7	Radiation therapy was not administered; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
8	Radiation therapy was recommended, but it is unknown if it was administered.
9	It is unknown if radiation therapy was recommended or administered. Death-certificate-only and autopsy-only cases.

Coding Examples:

Code	Explanation
2	A pt has a new primary breast cancer 4/2010 in the same breast where three years earlier she had a lumpectomy and radiation for breast cancer. The patient has another lumpectomy for her 4/2010 breast cancer but cannot receive the recommended radiation because she has already had radiation to that breast.

RX SUMM—SURG/RAD SEQ

Item Length: 1

NAACCR Item #1380

NAACCR Name: RX Summ-Surg/Rad Seq

XML NAACCR ID: rxSummSurgRadSeq

Definition:

This field records the order in which surgery and radiation therapies were administered for those patients who had both surgery and radiation.

Coding Instructions:

For the purpose of coding this data item, ‘Surgery’ is defined as a Surgical Procedure to the Primary Site (codes 10-90) or Scope of Regional Lymph Node Surgery (codes 2-7) or Surgical Procedure of Other Site (codes 1-5)

- Assign code 0 when
 - The patient did not have either surgery or radiation
 - The patient had surgery but not radiation
 - The patient had radiation but not surgery
 - It is unknown whether or not the patient had surgery and/or radiation
- Assign codes 2-9 when first course of therapy consists of both cancer-directed surgery and radiation therapy.
 - Assign code 4 when there are at least two courses, episodes, or fractions of radiation therapy given before and at least two more after surgery to the primary site, scope of regional lymph node surgery (excluding code 1), surgery to other regional site(s), distant site(s), or distant node(s).
 - Assign code 7 when there are at least two surgeries; radiation was administered between one surgical procedure and a subsequent surgical procedure.

Code	Definition
0	No radiation and/or surgery as defined above; Unknown if surgery and/or radiation given; Diagnosed at autopsy
2	Radiation before surgery
3	Radiation after surgery
4	Radiation both before and after surgery
5	Intraoperative radiation therapy
6	Intraoperative radiation with other radiation given before or after surgery
7	Surgery both before and after radiation
9	Sequence unknown, but both surgery and radiation were given

Coding Examples:

Code	Explanation
7	Sentinel lymph node biopsy, followed by radiation therapy, which was then followed by surgery of primary site. Code Radiation Sequence with Surgery as 7 (surgery both before and after radiation)
7	Two regional lymph nodes removed, followed by radiation, followed by surgery of the primary site. Code Radiation Sequence with Surgery as 7 (surgery both before and after radiation) because regional lymph node removal is coded in Scope of Regional Lymph Node Surgery.
3	Lymph node aspiration, followed by radiation, which was then followed by surgery of primary site. Code Radiation Sequence with Surgery as 3 (Radiation after surgery) BECAUSE lymph node aspiration is coded in Scope of Regional Lymph Node Surgery as code 1.
4	Preoperative radiation therapy to primary site to shrink a bulky tumor, followed by resection, which was then followed by post-operative radiation therapy to area. Assign code 4 (Radiation both before and after surgery)

PHASE I RADIATION TREATMENT MODALITY

Item Length: 2

NAACCR Item #1506

NAACCR Name: Phase I Radiation Treatment Modality

XML NAACCR ID: phase1RadiationTreatmentModality

Definition:

This field identifies the radiation modality administered during the first phase of radiation treatment delivered during the first course of treatment. It is to be coded for cases diagnosed 1/1/2018 and later. For cases diagnosed prior to 1/1/2018, this information should be coded in Radiation Regional Modality.

Radiation modality reflects whether a treatment was external beam, brachytherapy, a radioisotope as well as their major subtypes, or a combination of modalities. This data item should be used to indicate the radiation modality administered during the first phase of radiation.

Coding Instructions:

- Radiation treatment modality will typically be found in the radiation oncologist's summary letter for the first course of treatment. Segregation of treatment components into Phases and determination of the respective treatment modality may require assistance from the radiation oncologist to ensure consistent coding.
- The first phase may be commonly referred to as an initial plan and a subsequent phase may be referred to as a boost or cone down, and would be recorded as Phase II, Phase III, etc. accordingly.
- A new phase begins when there is a clinically meaningful change in target volume, treatment fraction size (I.e. dose given during a session), modality or treatment technique. Any one of these changes will mean that a new radiation plan will be generated in the treatment planning system, and it should be coded as a new phase of radiation therapy.
- For purposes of this data item, photons, x-rays and gamma-rays are equivalent.
- Use code 13 - Radioisotopes, NOS for radioembolization procedures, e.g. intravascular Yttrium-90.
- This data item intentionally does not include reference to various MV energies because this is not a clinically important aspect of technique. A change in MV energy (e.g., 6MV to 12MV) is not clinically relevant and does not represent a change in treatment technique. It is rare for change in MV energy to occur during any phase of radiation therapy.

- Note: Do not confuse a radioiodine scan with treatment. Only treatment is recorded in this item.

Code	Definition
00	No radiation treatment
01	External beam, NOS
02	External beam, photons
03	External beam, protons
04	External beam, electrons
05	External beam, neutrons
06	External beam, carbon ions
07	Brachytherapy, NOS
08	Brachytherapy, intracavitary, LDR
09	Brachytherapy, intracavitary, HDR
10	Brachytherapy, Interstitial, LDR
11	Brachytherapy, Interstitial, HDR
12	Brachytherapy, electronic
13	Radioisotopes, NOS
14	Radioisotopes, Radium-223
15	Radioisotopes, Strontium-89
16	Radioisotopes, Strontium-90
99	Radiation treatment modality unknown; Unknown if radiation treatment administered

Examples:

Code	Explanation
10	69 y/o man with prostate cancer. Treated with iodine seed implant (2/21/2018) at a university hospital; Returned home for additional treatment; 4-field conformal pelvic radiation with 15 Mv photons (3/5/2018 to 4/6/2018, 4500cGy in 25 fractions) at your facility. Code modality to 10 (brachytherapy, interstitial, LDR)
02	46 y/o female with T2N1M0 breast cancer and conservation surgery, 3 of 5 nodes positive. Whole breast RT, 5040 cGy in 28 fractions given between 8/13/2018 and 9/19/2018 using 6 Mv photons, conformal. Axillary and supraclavicular (SC) nodes treated concurrently with 6 Mv photons, and anterior field covering both regions and a posterior field (PAB) added to the axilla. Code modality 02 (External beam photon)
02	76 y/o man with T3b prostate cancer. Treated initially 7/9/2018 to 8/10/2018 with whole pelvis RT to 4500 cGy in 25 fractions of 180 cGy using a 4-field approach, all fields shaped conformally to pelvic anatomy. 8/13/2018 to 9/7/2018: IMRT boost of 19 fractions in which the seminal vesicles receive an additional 3420 cGy while the prostate receives 3800 cGy. Code modality to 02 (External beam photons)

Code	Explanation
02	65 y/o male with Stage IV adenocarcinoma of the lung and multiple symptomatic sites of metastasis. Treatment to thoracic spine 11/10/2018 to 11/21/2018 with unblocked photon field, 3000 cGy in 10 fractions. 11/12/2018 to 11/23/2018 treatment to right femur, unblocked photon field, 3000 cGy in 10 fractions. 11/12/2018 to 11/16/2018: Left hip treated with conformal fields designed to spare adjacent bowel, bladder, and soft tissue 2000 cGy in 5 equal fractions. 11/12/2018 to 11/16/2018: Right humerus, open square field, 2000 cGy in 5 equal fractions. Code modality to 02 (External beam, photons)

RX DATE--SYSTEMIC

Item Length: 8

NAACCR Item #3230

NAACCR Name: RX Date Systemic

XML NAACCR ID: rxDateSystemic

Description

Records the date of initiation for systemic therapy that is part of the first course of treatment. Systemic therapy includes the administration of chemotherapy agents, hormonal agents, biological response modifiers, bone marrow transplants, stem cell harvests, and surgical and/or radiation endocrine therapy.

Coding Instructions

- Record the first or earliest date on which systemic therapy was administered. Systemic therapy includes *Chemotherapy* (NAACCR Item #1390), *Hormone Therapy* (NAACCR Item #1400), *Immunotherapy* (NAACCR Item #1410), *Hematologic Transplant and Endocrine Procedures* (NAACCR Item #3250).

Code	Definition
YYYYMMDD YYYYMM YYYY	The date systemic therapy started is the year, month, and day that systemic therapy was administered at this or any facility. The first four digits are the year, the next two the month and the last two digits the day.

Examples:

Code	Explanation
20110112	A patient with breast cancer begins her regimen of chemotherapy on January 12, 2011, and is subsequently given tamoxifen on February 20, 2011.
201006	A patient with prostate cancer had an orchiectomy in June 2010 and was started on a regimen of hormonal agents on July 20, 2010

RX DATE – SYSTEMIC FLAG

Item Length: 2

NAACCR Item #3231

NAACCR Name: RX Date Systemic Flag

XML NAACCR ID: rxDateSystemicFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-Systemic. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Systemic has a full or partial date recorded.
- Assign code 10 when it is unknown whether any systemic therapy was given.
- Assign code 11 if no systemic therapy is planned or given.
- Assign code 12 if RX Date – Systemic cannot be determined, but it is known that systemic therapy was given as part of first course of treatment.
- Assign code 15 if systemic therapy is planned, but has not yet started and the start date is not yet available.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any systemic therapy was given)
11	No proper value is applicable in this context (for example, no systemic therapy was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, systemic therapy was given but the date is unknown).
15	Information is not available at this time, but it is expected that it will be available later (for example, systemic therapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up).
(blank)	A valid date value is provided in item RX Date-Systemic.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Systemic
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Systemic
10	Unknown if any systemic therapy given
11	No systemic therapy given
12	Systemic therapy given as first course treatment but date is completely unknown.
15	Systemic therapy not yet started but planned

Definition:

Chemotherapy is cancer therapy that achieves its antitumor effect through the use of antineoplastic drugs that inhibit the reproduction of cancer cells by interfering with DNA synthesis and mitosis. This data item records the type of chemotherapy administered as first course treatment at this and all other facilities. If chemotherapy was not administered, then this item records the reason it was not administered to the patient.

Chemotherapeutic Agents

Chemotherapeutic agents are chemicals that affect cancer tissue by means other than hormonal manipulation. Chemotherapeutic agents can be divided into five groups

- Alkylating agents
- Antimetabolites
- Natural Products
- Targeted therapy
- Miscellaneous

Alkylating Agents

Alkylating agents are not cell-cycle-specific. Although they are toxic to all cells, they are most active in the resting phase of the cell. Alkylating agents directly damage DNA to prevent the cancer cell from reproducing. Alkylating agents are used to treat many different cancers including acute and chronic leukemia, lymphoma, Hodgkin disease, multiple myeloma, sarcoma, and cancers of the lung, breast and ovary. Because the drugs damage DNA they can cause long-term damage to the bone marrow and can, in rare cases, lead to acute leukemia. The risk of leukemia from alkylating agents is “dose-dependent.” Types of alkylating agents include:

- Mustard gas derivatives/nitrogen mustards: Mechlorethamine, Cyclophosphamide, Chlorambucil, Melphalan, and Isosfmid
- Ethylemines: Thiotepa and Hexamethylmelamine
- Alkylsulfonates: Busulfan

- Hydrazines and Trizines: Alkretamine, Procarbazine, Decarbazine, and Temozolomide
- Nitrosureas: Camustine, Lomustine and Streptozocin. Nitrosureas are unique because they can cross the blood-brain barrier and can be used in treating brain tumors
- Metal salts: Carboplatin, Cisplatin, and Oxaliplatin

Antimetabolites

Antimetabolites are cell-cycle specific. Antimetabolites are very similar to normal substances within the cell. When the cells incorporate these substances into the cellular metabolism, they are unable to divide. Antimetabolites are classified according to the substances with which they interfere.

- Folic acid antagonist: Methotrexate
- Pyrimidine antagonist: 5-Fluorouracil, Fluorouridine, Cytarabine, Capecitabine, and Gemcitabine
- Purine antagonist: 6-Mercaptopurine and 6-Thioguanine
- Adenosine deaminase inhibitor: Cladribine, Fludarabine, Nelarabine, and Pentostatin

Natural Products

1. Plant Alkaloids are cell-cycle specific which means they attack the cells during various phases of division. They block cell division by preventing microtubule function. Microtubules are vital for cell division. Without them, division cannot occur. Plant alkaloids, as the name implies, are derived from certain types of plants.
 - Vinca alkaloids: Vincristine, Vinblastine, and Vinorelbine
 - Taxanes: Paclitaxel and Docetaxel
 - Podophyllotoxins: Etoposide and Teniposide
 - Camptothecin analogs: Irinotecan and Topotecan
2. Antitumor antibiotics are also cell-cycle specific and act during multiple phases of the cell cycle. They are made from natural products and were first produced by the soil fungus *Streptomyces*. Antitumor antibiotics form free radicals that break DNA strands, stopping the multiplication of cancer cells.

- Anthracyclines: Doxorubicin, Danorubicin, Epirubicin, Mitoxantone, and Idabycin
 - Chromomycins: Dactinomycin and Plicamycin
 - Miscellaneous: Mitomycin and Bleomycin
2. Topoisomerase inhibitors interfere with the action of topoisomerase enzymes (topoisomerase I and II). They control the manipulation of the structure of DNA necessary for replication.
- Topoisomerase I inhibitors: Irinotecan, topotecan
 - Topoisomerase II inhibitors: Amasrine, etoposide, etoposide phosphate, teniposide

Targeted therapy

Targeted therapy agents are a group of newer cancer drugs that act directly against abnormal proteins in cancer cells

Miscellaneous

Miscellaneous Antineoplastics that are unique

- Ribonucleotide reductase inhibitor: Hydroxyurea
- Adrenocortical steroid inhibitor: Mitotane
- Enzymes: Asparaginase and Pegaspargase
- Antimicrotubule agent: Estramustine
- Retinoids: Bexatene, Isotretinoin, Tretinoin (ATRA)

Coding Instructions:

See SEER*Rx (<http://www.seer.cancer.gov/tools/seerrx/index.html>) for chemotherapy drug codes and for information on the drug's function.

Chemotherapy recommended: A consult recommended chemotherapy, or the attending physician documented that chemotherapy was recommended. A referral to a clinical oncologist is equivalent to a recommendation.

Multiple agent chemotherapy: Planned first course of therapy included two or more chemotherapeutic agents and those agents were administered. The planned first course of

therapy may or may not have included other agents such as hormone therapy, immunotherapy, or other treatments in addition to the chemotherapeutic agents.

Single agent chemotherapy: Only one chemotherapeutic agent was administered to destroy cancer tissue during the first course of therapy. The chemotherapeutic agent may or may not have been administered with other drugs classified as immunotherapy, hormone therapy, ancillary, or other treatment.

- Code the chemotherapeutic agents whose actions are chemotherapeutic only; do not code the method of administration
- When chemotherapeutic agents are used as radiosensitizers or radioprotectants, they are given at a much lower dosage and do not affect the cancer. Radiosensitizers and radioprotectants are classified as ancillary drugs. Do not code as chemotherapy. Note: Do not assume that a chemo agent given with radiation therapy is a radiosensitizer. Seek additional information. Compare the dose given to the dose normally given for treatment.
For additional information see
 - The National Cancer Institute’s Physician Data Query (PDQ), Health Professional Version <http://www.cancer.gov/cancertopics/pdq>

And/or

- The National Comprehensive Cancer Network (NCCN) Clinical Practice Guidelines in Oncology http://www.nccn.org/professionals/physician_gls/f_guidelines.asp
- If it is known that chemotherapy is usually administered for this type and stage of cancer, but was not administered to the patient, use code 82, 85, 86, or 87 to record the reason why it was not administered.
- The physician may change a drug during the first course of therapy because the patient cannot tolerate the original agent.
 - This is a continuation of the first course of therapy when the chemotherapeutic agent that is substituted belongs to the same group (alkylating, antimetabolites, natural products, or other miscellaneous)
 - Do not code the new agent as first course therapy when the original chemotherapeutic agent is changed to one that is NOT in the same group. Code only the original agent as first course.
- Code 87 if the patient refused recommended chemotherapy, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended.

- Code as treatment for both primaries when the patient receives chemotherapy. Chemotherapy would likely affect both primaries.
- Assign code 00 when
 - There is no information in the patient's medical record about chemotherapy AND
 - It is known that chemotherapy is not usually performed for this type and/or stage of cancer
 - OR
 - There is no reason to suspect that the patient would have had chemotherapy.
 - If the treatment plan offered multiple treatment options and the patient selected treatment that did not include chemotherapy.
 - Patient elects to pursue no treatment following discussion of chemotherapy. Discussion does not equal a recommendation
 - Watchful waiting/active surveillance (CLL)
 - Patient diagnosed at autopsy.
- Do not code combination of ancillary drugs administered with single agent chemotherapeutic agents as multiple chemotherapy. For example, the administration of 5-FU (antimetabolite) and Leucovorin (ancillary drug) is coded to single agent (Code 02).
- Assign code 82 when chemotherapy is a customary option for the primary site/histology but it was not administered due to patient risk factors, such as:
 - Advanced age
 - Comorbid condition(s) (heart disease, kidney failure, other cancer, etc.)
- Assign code 88 when the only information available is
 - The patient was referred to an oncologist
 - Insertion of port-a-cath

- Assign code 99 when there is no documentation that chemotherapy was recommended or administered.
- **Coding for Tumor Embolization.** Chemoembolization is a procedure in which the blood supply to the tumor is blocked surgically or mechanically and anticancer drugs are administered directly to the tumor. This permits a higher concentration of drug to be in contact with the tumor for a longer period of time.

Radioembolization is a tumor embolization combined with the injection of small radioactive beads or coils into an organ or tumor.

Tumor embolization is the intentional blockage of an artery or vein to stop the flow of blood through the desired vessel.

- Code as Chemotherapy when the embolizing agent(s) is a chemotherapeutic drug(s). Use SEER*Rx to determine whether the drugs used are classified as chemotherapeutic agents. Use codes 01, 02, 03 as specific information regarding the agent(s) is documented.
- Do not code pre-surgical embolization of hypervascular tumors with particles, coils or alcohol. These pre-surgical embolizations are typically performed to make the resection of the primary tumor easier. Examples where pre-surgical embolization is used include meningiomas, hemangioblastomas, paragangliomas, and renal cell metastases in the brain.
- **Important information that affects the classification of some systemic therapies.** There are six drugs in the table below that were classified as chemotherapy and will be classified as BRM/Immunotherapy beginning with cases diagnosed **January 1, 2013 and later**. Code these as chemotherapy for cases diagnosed prior to January 1, 2013. Notes about this change have been added to SEER*RX.

Drug Name(s)	Category Prior to 2013	Category 2013+
Alemtuzumab/Campath	Chemotherapy	BRM/Immunotherapy
Bevacizumab/Avastin	Chemotherapy	BRM/Immunotherapy
Rituximab	Chemotherapy	BRM/Immunotherapy
Trastuzumab/Herceptin	Chemotherapy	BRM/Immunotherapy
Pertuzumab/Perjeta	Chemotherapy	BRM/Immunotherapy
Cetuxumab/Erbitux	Chemotherapy	BRM/Immunotherapy

Code	Definition
00	None, chemotherapy was not part of the planned first course of therapy or diagnosed at autopsy.
01	Chemotherapy administered as first course therapy, but the type and number of agents is not documented in patient record.
02	Single-agent chemotherapy administered as first course therapy.
03	Multiagent chemotherapy administered as first course therapy.
82	Chemotherapy was not recommended/administered because it was contraindicated due to patient risk factors (i.e., comorbid conditions, advanced age).
85	Chemotherapy was not administered because the patient died prior to planned or recommended therapy.
86	Chemotherapy was not administered. It was recommended by the patient's physician, but was not administered as part of the first course of therapy. No reason was stated in patient record.
87	Chemotherapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, a patient's family member, or the patient's guardian. The refusal was noted in the patient record.
88	Chemotherapy was recommended, but it is unknown if it was administered.
99	It is unknown whether a chemotherapeutic agent(s) was recommended or administered because it is not stated in patient record or death certificate only.

Coding Examples:

Code	Explanation
01	A patient with primary lung cancer is known to have received chemotherapy; however, the name(s) of agent(s) administered is not stated in patient record.
00	Patient is diagnosed with multiple myeloma. There is no mention of treatment or treatment plans in the medical record. The patient died three months after diagnosis. There is no additional pertinent information available. Assign code 00 since there is no reason to suspect that the patient had been treated.
02	A patient with stage III colon cancer is treated with surgery and adjuvant therapy of 5-FU and leucovorin. Code the 5-FU as a single drug, chemotherapy and record the leucovorin only in the text as this is an ancillary drug.
03	A patient with breast cancer patient receives CMF, (Cyclophosphamide, Adriamycin, 5-FU) chemotherapy regimen.
86	Following surgical resection of a right colon cancer with positive nodes the physician recommends chemotherapy. The patient record states that chemotherapy was not subsequently administered to the patient, but the reason why chemotherapy was not administered is not given.

RX DATE – CHEMO

Item Length: 8
 NAACCR Item #1220
 NAACCR Name: RX Date Chemo
 XML NAACCR ID: rxDateChemo

Description

Date of initiation of chemotherapy that is part of the first course of treatment. The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

Coding Instructions

- Record the earliest date on which chemotherapy was administered at any facility.

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of initiation of chemotherapy is the year, month, and day that the patient received the first treatment with chemotherapy. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.
blank	No chemotherapy administered; autopsy-only case; Chemotherapy administered, date unknown; Unknown if chemotherapy administered.

RX DATE – CHEMO FLAG

Item Length: 2

Definition:

This flag explains why no appropriate value is in the field, RX Date-Chemo. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Chemo has a full or partial date recorded.
- Assign code 10 when it is unknown whether any chemotherapy was given.
- Assign code 11 if no chemotherapy is planned or given.
- Assign code 12 if RX Date – Chemo cannot be determined, but it is known that chemotherapy was given as part of first course of treatment.
- Assign code 15 if chemotherapy is planned, but has not yet started and the start date is not yet available.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any chemotherapy was given)
11	No proper value is applicable in this context (for example, no chemotherapy was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, chemotherapy was given but the date is unknown).
15	Information is not available at this time, but it is expected that it will be available later (for example, chemotherapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up).
(blank)	A valid date value is provided in item RX Date-Chemo.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Chemo
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Chemo
10	Unknown if any chemotherapy given
11	No chemotherapy given
12	Chemotherapy given as first course treatment but date is completely unknown.
15	Chemotherapy not yet started but planned

RX SUMM--HORMONE

Item Length: 2

NAACCR Item #1400

NAACCR Name: RX Summ—Hormone

XML NAACCR ID: rxSummHormone

Definition:

Hormone therapy is Cancer therapy that achieves its antitumor effect through changes in hormonal balance. This type of therapy includes the administration of hormones, agents acting via hormonal mechanisms, antihormones, and steroids. This data item records the type of hormone therapy administered as first course treatment at this and all other facilities. If hormone therapy was not administered, then this item records the reason it was not administered to the patient.

Hormone Categories

Hormones may be divided into several categories

- Androgens: Fluoxymesterone
- Anti-androgens: Bicalutamide (Casodex), flutamide (Eulexin), and nilutamide (Nilandron)
- Corticosteroids: Adrenocorticotrophic agents
- Estrogens
- Progestins
- Estrogen antagonists, Anti-estrogens: Fulvestrant (Faslodex), tamoxifen, and toremifene (Fareston)
- Aromatase inhibitors, Antiaromatase: Anastrozole (Arimidex), exemestane (Aromasin), and letrozole (Femara)
- GnRH or LH-RH: Lupron, Zoladex

Coding Instructions:

See SEER*Rx (<http://www.seer.cancer.gov/tools/seerrx/index.html>) for hormone therapy drug codes.

Note: Surgical removal of organs for hormone manipulation is not coded in this data item. Code these procedures in the data field Hematologic Transplant and Endocrine Procedures.

- Code the hormonal agent given as part of combination chemotherapy (e.g., R-CHOP), whether it affects the cancer cells or not
 - Check SEER*Rx to determine if a hormone agent is part of a combination chemotherapy regimen
- Do not code prednisone as hormone therapy when it is administered for reasons other than chemotherapeutic treatment.
- Assign code 00 when
 - The medical record states that hormone therapy was not given, was not recommended, or was not indicated.
 - There is no information on the patient's medical record about hormone therapy AND
 - It is known that hormone therapy is not usually performed for this type and/or stage of cancer

OR

 - There is no reason to suspect that the patient would have had hormone therapy.
 - If the treatment plan offered multiple treatment options and the patient selected treatment that did not include hormone therapy.
 - Patient elected to pursue no treatment following the discussion of hormone therapy treatment. Discussion does not equal a recommendation.
 - Watchful waiting/active surveillance (e.g., prostate)
 - Patient diagnosed at autopsy
 - Hormone treatment was given for a non-reportable condition or as a chemoprevention prior to diagnosis of a reportable condition.
- Assign code 87 when
 - The patient refused recommended hormone therapy
 - The patient made a blanket refusal of all recommended treatment and hormone therapy is a customary option for the primary site/histology

- The patient refused all treatment before any was recommended and hormone therapy is a customary option for the primary site/histology
- Assign code 88 when the only information available is that the patient was referred to an oncologist.
- Assign code 99 when there is no documentation that hormone therapy was recommend or performed

Code	Definition
00	None. Hormone therapy was not part of the planned first course of therapy, not usually administered for this type and/or stage of cancer or diagnosed at autopsy.
01	Hormone therapy administered as first course therapy.
82	Hormone therapy was not recommended/administered because it was contraindicated due to patient risk factors (i.e., comorbid conditions, advanced age).
85	Hormone therapy was not administered because the patient died prior to planned or recommended therapy.
86	Hormone therapy was not administered. It was recommended by the patient's physician, but was not administered as part of the first course of therapy. No reason was stated in patient record.
87	Hormone therapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, patient's family member, or the patient's guardian. The refusal was noted in the patient record.
88	Hormone therapy was recommended, but it is unknown if it was administered.
99	It is unknown whether a hormonal agent(s) was recommended or administered because it is not stated in patient record or death certificate only.

Coding Examples:

Code	Explanation
00	A patient has advanced lung cancer with multiple metastases to the brain. The physician orders Decadron to reduce the edema in the brain and relieve the neurological symptoms. Decadron is an ancillary agent which should not be coded.
00	A breast cancer patient is treated with aminoglutethimide (Cytadren, Elipten), which suppresses the production of glucocorticoids and mineralcorticoids. This patient is given hydrocortisone (glucocorticoid) and Florinef (mineralcorticoid) for replacement therapy.
01	A patient with prostate cancer in given Lupron injections.
87	A patient with hormone positive breast cancer refuses Tamoxifen which is noted in her record.

Code	Explanation
01	Endometrial cancer may be treated with progesterone. Code all administration of progesterone to patients with endometrial cancer in this data item. Even if the progesterone is given for menopausal symptoms, it has an effect on the growth or recurrence of endometrial cancer.
01	Follicular and papillary cancers of the thyroid are often treated with thyroid hormone to suppress serum thyroid-stimulating hormone (TSH). If a patient with papillary and/or follicular cancer of the thyroid is given a thyroid hormone, code the treatment in this data item.
01	Bromocriptine suppresses the production of prolactin, which causes growth in pituitary adenoma. Code bromocriptine as hormone treatment for pituitary adenoma.
01	Lupron is hormone therapy that has been approved as an ovarian suppressor for pre-menopausal breast cancer.

RX DATE – HORMONE

Item Length: 8
NAACCR Item #1230
NAACCR Name: RX Date Hormone
XML NAACCR ID: rxDateHormone

Description

Date of initiation of hormone that is part of the first course of treatment. The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

Coding Instructions

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of initiation of hormone therapy is the year, month and day that the patient received the first treatment with hormones. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.
blank	No hormone therapy administered; hormone therapy administered but date completely unknown; unknown if any hormone therapy administered; autopsy-only case

RX DATE – HORMONE FLAG

Item Length: 2

NAACCR Item #1231

NAACCR Name: RX Date Hormone Flag

XML NAACCR ID: rxDateHormoneFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-Hormone. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Hormone has a full or partial date recorded.
- Assign code 10 when it is unknown whether any hormone therapy was given.
- Assign code 11 if no hormone therapy is planned or given.
- Assign code 12 if RX Date – Hormone cannot be determined, but it is known that hormone therapy was given as part of first course of treatment.
- Assign code 15 if hormone therapy is planned, but has not yet started and the start date is not yet available.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any hormone therapy was given)
11	No proper value is applicable in this context (for example, no hormone therapy was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, hormone therapy was given but the date is unknown).
15	Information is not available at this time, but it is expected that it will be available later (for example, hormone therapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up).
(blank)	A valid date value is provided in item RX Date-Hormone.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Hormone
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Hormone
10	Unknown if any hormone therapy given
11	No hormone therapy given
12	Hormone therapy given as first course treatment but date is completely unknown.
15	Hormone therapy not yet started but planned

Definition:

Immunotherapy is cancer therapy that achieves its antitumor effect by altering the immune system or changing the host's response to the tumor cells. This data item records the type of immunotherapy administered as first course treatment at this and all other facilities. If immunotherapy was not administered, then this item records the reason it was not administered to the patient.

Immunotherapy uses the body's immune system, either directly or indirectly, to fight cancer or to lessen the side effects that may be caused by some cancer treatments. Record only those treatments administered to affect the cancer cells.

Immunotherapy is **designed** to:

1. Make **cancer cells** more **recognizable** and therefore more **susceptible** to destruction by the immune system.
2. **Boost** the killing power of **immune** system cells, such as T-cells, NK-cells, and macrophages.
3. **Alter** the **growth patterns** of cancer cells to promote behavior like that of healthy cells.
4. **Block** or **reverse** the process that **changes** a normal cell or a pre-cancerous cell into a cancerous cell.
5. **Enhance** the body's ability to **repair** or **replace** normal cells damaged or destroyed by other forms of cancer treatment, such as chemotherapy or radiation.
6. **Prevent** cancer cells from **spreading** to other parts of the body.

Types of immunotherapy

Cancer Vaccines: Cancer vaccines are still in the experimental phase and are not coded in this data item. They may be coded in the field Other Therapy. Currently clinical trials use cancer vaccines for brain, breast, colon, kidney, lung, melanoma, and ovary.

Interferons: Interferons belong to a group of proteins called cytokines. They are produced naturally by the white blood cells in the body. Interferon-alpha is able to slow tumor growth directly as well as activate the immune system. It is used for a number of cancers including

multiple myeloma, chronic myelogenous leukemia (CML), hairy cell leukemia, and malignant melanoma.

Interleukins (IL-2) are often used to treat kidney cancer and melanoma.

Monoclonal Antibodies: Monoclonal antibodies (Mab) are produced in a laboratory. The artificial antibodies are used in a variety of ways in systemic therapy and can be chemotherapy, immunotherapy, or ancillary drugs. Some are injected into the patient to seek out and disrupt cancer cell activities. When the monoclonal antibody disrupts tumor growth, it is coded as chemotherapy. Other Mabs are linked to radioisotopes (conjugated monoclonal antibodies). The Mab finds and attaches to the target tumor cells and brings with it the radioisotope that actually kills the tumor cell. The monoclonal antibody itself does nothing to enhance the immune system. Conjugated monoclonal antibodies such as tositumomab (Bexxar) or ibritumomab (Zevalin) are coded to the part of the drug that actually kills the cells, usually radioisotopes. A third function of Mabs is to enhance the immune response against the cancer, either by identifying tumor cells that are mimicking normal cells, or by boosting the body's natural defenses that destroy foreign cells. Consult SEER*Rx for the treatment category in which each monoclonal antibody should be coded.

Coding Instructions:

See SEER*Rx (<http://www.seer.cancer.gov/tools/seerrx/index.html>) for immunotherapy drug codes.

- Assign code 00
 - The medical record states that immunotherapy was not given, not recommended, or not indicated
 - When there is no information in the patient's medical record about immunotherapy AND
 - It is known that immunotherapy is not usually performed for this type and/or stage of cancer
 - OR
 - There is no reason to suspect that the patient would have had immunotherapy.
 - If the treatment plan offered multiple treatment options and the patient selected treatment that did not include immunotherapy.
 - Patient elects to pursue no treatment following the discussion of immunotherapy. Discussion does not equal a recommendation.
 - Watchful waiting, active surveillance (e.g., prostate)
 - Patient diagnosed at autopsy

- For anti-thymocyte globulin treatment. Anti-thymocyte globulin is used to treat transplant rejection. Do not code as immunotherapy.
- Assign code 87 when
 - The patient refused recommended immunotherapy
 - The patient made a blanket refusal of all recommended treatment and immunotherapy is a customary option for the primary site/histology
 - The patient refused all treatment before any was recommended and immunotherapy is a customary option for the primary site/histology
- Assign code 88 when the only information available is that the patient was referred to an oncologist.
- Assign code 99
 - When there is no documentation that immunotherapy was recommended or performed AND
 - Immunotherapy is usually given for this type and/or stage of cancer
- **Important information that affects the classification of some systemic therapies.** There are six drugs in the table below that were classified as chemotherapy and will be classified as BRM/Immunotherapy beginning with cases diagnosed **January 1, 2013 and later**. Code these as chemotherapy for cases diagnosed prior to January 1, 2013. Notes about this change have been added to SEER*RX.

Drug Name(s)	Category Prior to 2013	Category 2013+
Alemtuzumab/Campath	Chemotherapy	BRM/Immunotherapy
Bevacizumab/Avastin	Chemotherapy	BRM/Immunotherapy
Rituximab	Chemotherapy	BRM/Immunotherapy
Trastuzumab/Herceptin	Chemotherapy	BRM/Immunotherapy
Pertuzumab/Perjeta	Chemotherapy	BRM/Immunotherapy
Cetuxumab/Erbitux	Chemotherapy	BRM/Immunotherapy

Code	Definition
00	None, immunotherapy was not part of the planned first course of therapy, is not customary therapy for this cancer or diagnosed at autopsy.
01	Immunotherapy administered as first course therapy.
82	Immunotherapy was not recommended/administered because it was contraindicated due to patient risk factors (i.e., comorbid conditions, advanced age).
85	Immunotherapy was not administered because the patient died prior to planned or recommended therapy.
86	Immunotherapy was not administered. It was recommended by the patient's physician, but was not administered as part of the first course of therapy. No reason was stated in patient record.
87	Immunotherapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, a patient's family member, or the patient's guardian. The refusal was noted in patient record.
88	Immunotherapy was recommended, but it is unknown if it was administered.
99	It is unknown whether an immunotherapeutic agent(s) was recommended or administered because it is not stated in patient record. Death certificate only.

Examples:

Code	Explanation
01	A patient is treated with a TURBT followed by BCG for bladder cancer.
85	Immunotherapy is planned but patient died before receiving it.
01	Patient diagnosed with breast cancer January 5, 2021, and begins receiving Herceptin as part of first course therapy on January 30, 2021. Code the Herceptin in the BRM/Immunotherapy data item.

RX DATE – BRM

Item Length: 8
NAACCR Item #1240
NAACCR Name: RX Date BRM
XML NAACCR ID: rxDateBrm

Description

Date of initiation of immunotherapy that is part of the first course of treatment. The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

Coding Instructions

- Record the first date on which immunotherapy (BRM) was administered by any facility.
 - Code the date that the prescription was written if date administered unknown

Code	Definition
YYYYMMDD YYYYMM YYYY	The date of initiation of immunotherapy is the year, month, and day that the patient received the first treatment with immunotherapy. The first four digits are the year, the fifth and sixth digits are the month, and the last two digits are the day.
blank	No immunotherapy administered; immunotherapy administered but date completely unknown; unknown if immunotherapy administered; autopsy-only case

RX DATE – BRM FLAG

Item Length: 2
NAACCR Item #1241
NAACCR Name: RX Date BRM Flag
XML NAACCR ID: rxDateBrmFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-BRM. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - BRM has a full or partial date recorded.
- Assign code 10 when it is unknown whether any immunotherapy was given.
- Assign code 11 if no immunotherapy is planned or given.
- Assign code 12 if RX Date – BRM cannot be determined, but it is known that immunotherapy was given as part of first course of treatment.
- Assign code 15 if immunotherapy is planned, but has not yet started and the start date is not yet available.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any immunotherapy was given)
11	No proper value is applicable in this context (for example, no immunotherapy was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, immunotherapy was given but the date is unknown).
15	Information is not available at this time, but it is expected that it will be available later (for example, immunotherapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up).
(blank)	A valid date value is provided in item RX Date-BRM.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-BRM
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-BRM
10	Unknown if any immunotherapy given
11	No immunotherapy given
12	Immunotherapy given as first course treatment but date is completely unknown.
15	Immunotherapy not yet started but planned

Definition:

Endocrine therapy is cancer therapy that achieves its antitumor effect through the use of radiation or surgical procedures that suppress the naturally occurring hormonal activity of the patient (when the cancer occurs at another site) and, therefore, alter or affect the long-term control of the cancer's growth. Hematologic transplants are bone marrow or stem cell transplants performed to protect patients from myelosuppression or bone marrow ablation associated with the administration of high-dose chemotherapy or radiation therapy. This data item identifies systemic therapeutic procedures administered as part of the first course of treatment at this and all other facilities. If none of these procedures were administered, then this item records the reason they were not performed. These include bone marrow transplants, stem cell harvests, surgical and/or radiation endocrine therapy.

Bone marrow transplant (BMT): Procedure used to restore stem cells that were destroyed by chemotherapy and/or radiation. Replacing the stem cells allows the patient to undergo higher doses of chemotherapy.

BMT Allogeneic: Receives bone marrow or stem cells from a donor.

BMT Autologous: Uses the patient's own bone marrow and/or stem cells. The tumor cells are filtered out and the purified blood and stem cells are returned to the patient.

Note: Used for breast cancer, lymphoma, leukemia, aplastic anemia, myeloma, germ cell tumors, ovarian cancer, and small cell lung cancer.

Conditioning: High-dose chemotherapy with or without radiation administered prior to transplants such as BMT and stem cells to kill cancer cells. This conditioning also destroys normal bone marrow cells so the normal cells need to be replaced (rescue). The high dose chemotherapy is coded in the Chemotherapy field and the radiation is coded in the Radiation field.

Hematopoietic Growth Factors: A group of substances that support hematopoietic (blood cell) colony formation. The group includes erythropoietin, interleukin-3, and colony-stimulating factors (CSFs). The growth-stimulating substances are ancillary drugs and not coded.

Non-Myeloablative Therapy: Uses immunosuppressive drugs pre- and post-transplant to ablate (destroy) the bone marrow. These are not recorded as therapeutic agents.

Peripheral Blood Stem Cell Transplantation (PBSCT): Rescue that replaces stem cells after conditioning.

Rescue: Rescue is the actual BMT or stem cell transplant done after conditioning.

Stem Cells: Immature cells found in bone marrow, blood stream and umbilical cords. The stem cells mature into blood cells.

Stem cell transplant: Procedure to replenish supply of healthy blood-forming cells. Also known as bone marrow transplant or umbilical cord blood transplant, depending on the source of the stem cells.

Umbilical cord stem cell transplant: Treatment with stem cells harvested from umbilical cord blood.

Coding Instructions:

- Assign code 00
 - The medical record states that there was no hematologic transplant or endocrine therapy, or these were not recommended, or not indicated.
 - When there is no information in the patient’s medical record about transplant procedure or endocrine therapy AND
 - It is known that transplant procedure or endocrine therapy is not usually performed for this type and/or stage of cancer

OR

 - There is no reason to suspect that the patient would have had transplant procedure or endocrine therapy.
 - If the treatment plan offered multiple treatment options and the patient selected treatment that did not include transplant procedure or endocrine therapy.
 - Patient elects to pursue no treatment following the discussion of transplant procedure or endocrine therapy. Discussion does not equal a recommendation. Patient’s decision not to pursue transplant procedure or endocrine therapy is not a refusal or transplant procedure or endocrine therapy in this situation.
 - Watchful waiting/active surveillance (e.g., CLL)
 - Patient diagnosed at autopsy.
- Assign code 10 if the patient has a bone marrow transplant and it is unknown if autologous or allogeneic (BMT, NOS) or “mixed chimera transplant” (mini-transplant)

or non-myeloablative transplant). These transplants are a mixture of the patient's cells and donor cells.

- Codes 11 (Bone marrow transplant autologous) and 12 (Bone marrow transplant allogeneic) have priority over code 10 (BMT, NOS).
- Assign code 12 (allogeneic) for a syngeneic bone marrow transplant (from an identical twin) or for a transplant from any person other than the patient.
- Assign code 20 when
 - Allogeneic stem cell transplant
 - Peripheral blood stem cell transplant
 - Umbilical cord stem cell transplant (single or double)

Note: If the patient does not have a rescue, code the stem cell harvest as 88, (recommended, unknown if administered) or if harvested but unknown if infused.

- Assign code 30 for endocrine radiation and/or surgery. Endocrine organs are testes and ovaries. Endocrine radiation and/or surgical procedures must be bilateral, or must remove the remaining paired organ for hormonal effect.
- Assign code 87
 - If the patient refused recommended transplant or endocrine procedure.
 - If the patient made a blanket refusal of all recommended treatment and the treatment coded in this data item is a customary option for the primary site/histology.
 - If patient refused all treatment before any was recommended.
- Assign code 88 when
 - The only information available is that the patient was referred to an oncologist for consideration of hematologic transplant or endocrine procedure
 - A bone marrow or stem cell harvest was undertaken, but it was not followed by a rescue or reinfusion as part of the first course treatment

- Code 99 if it is not known whether a transplant or endocrine procedure is usually administered for this type and stage of cancer, and there is no mention in the patient record whether it was recommended or administered.

Code	Definition
00	No transplant procedure or endocrine therapy was administered as part of first course therapy; not customary therapy for this cancer; diagnosed at autopsy.
10	A bone marrow transplant procedure was administered, but the type was not specified.
11	Bone marrow transplant, autologous.
12	Bone marrow transplant, allogeneic.
20	Stem cell harvest (stem cell transplant) and infusion.
30	Endocrine surgery and/or endocrine radiation therapy as first course therapy.
40	Combination of endocrine surgery and/or radiation with a transplant procedure. (Combination of codes 30 and 10, 11, 12, or 20) as first course therapy.
82	Hematologic transplant and/or endocrine surgery/radiation was not recommended and/or administered because it was contraindicated due to patient risk factors (i.e., comorbid conditions, advanced age).
85	Hematologic transplant and/or endocrine surgery/radiation was not administered because the patient died prior to planned or recommended therapy.
86	Hematologic transplant and/or endocrine surgery/radiation was not administered. It was recommended by the patient's physician, but was not administered as part of the first course of therapy. No reason was stated in patient record.
87	Hematologic transplant and/or endocrine surgery/radiation was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, a patient's family member, or the patient's guardian. The refusal was noted in patient record.
88	Hematologic transplant and/or endocrine surgery/radiation was recommended, but it is unknown if it was administered.
99	It is unknown whether hematologic transplant and/or endocrine surgery/radiation was recommended or administered because it is not stated in patient record

RX SUMM—SYSTEMIC/SURGERY SEQUENCE

Item Length: 1

NAACCR Item #1639

NAACCR Name: RX Summ—Systemic/SurSeq

XML NAACCR ID: rxSummSystemicSurSeq

Description

Records the sequencing of systemic therapy (chemotherapy, hormone, BRM, and transplant/endocrine) and surgical procedures given as part of the first course of treatment.

Coding Instructions

- *Systemic/Surgery Sequence* is to be used for patients diagnosed on or after January 1, 2006.
- Code the administration of systemic therapy in sequence with the first surgery performed.
- If none of the following surgical procedures were performed: *Surgical Procedure of Primary Site* (codes 10-19), *Scope of Regional Lymph Node Surgery* (codes 2-7), *Surgical Procedure/Other Site* (codes 1-5), then code this item 0.
- If the patient received both systemic therapy and any one of the following surgical procedures: *Surgical Procedure of Primary Site* (codes 10-19), *Scope of Regional Lymph Node Surgery* (codes 2-7), *Surgical Procedure/Other Site* (codes 1-5), then code this item 2-9, as appropriate.
- Codes 4 and 7 are used for multiple episodes of therapy in first course treatment. Use the code that defines the first sequence that applies.

Code	Label	Definition
0	No systemic therapy and/or surgical procedures; Unknown if surgery and/or systemic therapy given	The patient did not have both systemic therapy and surgery. It is unknown whether or not the patient had surgery and/or systemic therapy
2	Systemic therapy before surgery	The patient had systemic therapy prior to surgery.
3	Systemic therapy after surgery	The patient had systemic therapy after surgery
4	Systemic therapy both before and after surgery	Systemic therapy was administered prior to surgery and also after surgery.
5	Intraoperative systemic therapy	The patient had intraoperative systemic therapy.
6	Intraoperative systemic therapy with other therapy administered before or after surgery	The patient had intraoperative systemic therapy and also had systemic therapy before and/or after surgery.
7	Surgery both before and after systemic therapy	Systemic therapy was administered between two separate surgical procedures to the primary site; regional lymph nodes; surgery to other regional site(s), distant site(s), or distant lymph node(s).
9	Sequence unknown	<ul style="list-style-type: none"> The patient had systemic therapy and also had surgery It is unknown whether the systemic therapy was administered prior to surgery, after surgery, or intraoperatively

Coding Examples:

Code	Reason
0	Patient receives chemo and radiation only for a lung cancer.
2	Patient with inflammatory breast cancer receives chemo prior to an MRM.
7	Patient has LN dissection, followed by chemo, followed by primary site surgery.
4	Patient with breast cancer receives pre-operative chemotherapy followed by post-operative Tamoxifen
9	A patient comes to your facility after having chemo and surgery for cancer. However, you do not know the sequence of the treatment modalities.

RX SUMM--OTHER

Item Length: 1
NAACCR Item #1420
NAACCR Name: RX Summ—Other
XML NAACCR ID: rxSummOther

Definition:

Identifies other treatment that cannot be defined as surgery, radiation, systemic therapy, or ancillary treatment.

A quote from the website for the National Cancer Institute (NCI), Office of Cancer Complementary and Alternative Medicine (OCCAM) defines Complementary and Alternative Medicine (CAM) as any medical system, practice, or product that is not thought of as “western medicine” or standard medical care.

- Complementary medicine means it is used along with standard medicine, also called conventional medicine.
- Alternative medicine is used in place of standard treatments.

CAM treatments may include dietary supplements, megadose vitamins, herbal preparations, acupuncture, massage therapy, magnet therapy, spiritual healing, and meditation.

The OCCAM was established to coordinate and enhance activities of the NCI in complementary and alternative medicine research as it relates to the prevention, diagnosis, and treatment of cancer, cancer-related symptoms and side effects of conventional cancer treatment.

See complete information on types of complementary and alternative medicine specific to cancer at <http://www.cancer.gov/cam/>. For additional information on cancer and other diseases, please visit <http://nccam.nih.gov/health/whatisacam/>.

Coding Instructions:

- Assign code 0 when
 - There is no information in the patient’s medical record about other therapy.
AND
 - There is no reason to suspect that the patient would have had other therapy.
 - If the treatment plan offered multiple treatment options and the patient selected treatment that did not include other therapy.
 - Patient elects to pursue no treatment following the discussion of other therapy. Discussion does not equal a recommendation. Patient’s decision not to pursue other therapy is not a refusal of other therapy in this situation.

- First course of treatment was active surveillance/watchful waiting
- Patient diagnosed at autopsy.
- Assign code 1 for
 - Hematopoietic treatments such as: phlebotomy or aspirin. Consult the **Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual** (<http://seer.cancer.gov/tools/heme/>) for instruction on coding care in this data item for specific conditions.

Note: Do not code blood transfusion as treatment

Rationale: Blood transfusions may be used for any medical condition that causes anemia. It would be virtually impossible for the registrar to differentiate between blood transfusions used for a co-morbidity (i.e., anemia) from those given as prophylactic treatment of a hematopoietic neoplasm.

- PUVA (Psoralen (P) and long-wave ultraviolet radiation (UVA)) in the RARE event that it is used as treatment for extremely thin melanomas or cutaneous T-cell lymphomas (e.g., mycosis fungoides)

Note: Code UVB phototherapy for mycosis fungoides as photodynamic therapy under Surgery of Primary Site for skin. Assign code 11 [Photodynamic therapy (PDT)] if there is no pathology specimen. Assign code 21 [Photodynamic therapy (PDT)] if there is a pathology specimen.

- Photophoresis. This treatment is used ONLY for thin melanoma or cutaneous T-cell lymphoma (mycosis fungoides).
- Peptide Receptor Radionuclide Therapy (PRRT).
- Cancer treatment that could not be assigned to the previous treatment fields (surgery, radiation, chemotherapy, immunotherapy, or systemic therapy)

- Assign code 2 for any experimental or newly developed treatment, such as a clinical trial, that differs greatly from proven types of cancer therapy

Note: Hyperbaric oxygen has been used to treat cancer in clinical trials, but it is also used to promote tissue healing following head and neck surgeries. Do not code the administration of hyperbaric oxygen to promote healing as an experimental treatment.

- Assign code 3 when the patient is enrolled in a double blind clinical trial. When the trial is complete and the code is broken, review and recode the therapy.

- Assign code 6 for
 - Cancer treatment administered by nonmedical personnel.
 - Unconventional methods whether they are the only therapy or are given in combination with conventional therapy.
 - Complementary and Alternative Medicine (CAM) as any medical system, practice, or product that is not thought of as “western medicine” or standard medical care. CAM treatments may include dietary supplements, megadose vitamins, herbal preparations, acupuncture, massage therapy, magnet therapy, spiritual healing, and meditation.
 - Alternate medicine is treatment that is used instead of standard medical treatments. Alternative therapy is when the patient receives no other type of standard treatment.
 - Complementary medicine. Treatments that are used along with standard medical treatments but are not standard treatments; also called conventional medicine. One example is using acupuncture to help lessen some side effects of cancer treatment in conjunction with standard treatment.
 - Integrative medicine. A total approach to medical care that combines standard medicine with the CAM practices that have shown to be safe and effective. They treat the patient’s mind, body, and spirit.

Note: See complete information on types of complementary and alternative medicine specific to cancer at NCI Office of Cancer Complementary and Alternative Medicine (<https://cam.cancer.gov/>). For additional information on cancer and other diseases, please visit NIH National Center for Complementary and Integrative Health (<https://www.nccih.nih.gov/health/complementary-alternative-or-integrative-health-whats-in-a-name>).

- Assign code 8 when other therapy was recommended by the physician but there is no information that the treatment was given.
- Assign code 9 when there is no documentation that other therapy was recommended or performed.
- **Coding Tumor Embolization:** Tumor embolization is the intentional blockage of an artery or vein to stop the flow of blood through the desired vessel.
 - Use code 1 when tumor embolization is performed using alcohol as the embolizing agent.

- Use code 1 for embolization to a site other than the liver where the embolizing agent is unknown.

Do not code pre-surgical embolization of hypervascular tumors with particles, coils or alcohol. These pre-surgical embolizations are typically performed to make the resection of the primary tumor easier. Examples where pre-surgical embolization is used include meningiomas, hemangioblastomas, paragangliomas, and renal cell metastases in the brain.

- A complete description of the treatment plan should be recorded in the text field for Other Treatment in the abstract.

Code	Definition
0	None. All cancer treatment was coded in other treatment fields (surgery, radiation, systemic therapy). Patient received no cancer treatment or diagnosed at autopsy.
1	Other cancer treatment that cannot be appropriately assigned to specified treatment data items (surgery, radiation, systemic). Use this code for treatment unique to hematopoietic diseases.
2	Experimental. This code is not defined. It may be used to record participation in institution based clinical trials.
3	Double-blind clinical trial. A patient is involved in a double-blind clinical trial. Code the treatment actually administered when the double-blind trial code is broken.
6	Unproven Cancer treatments administered by non-medical personnel.
7	Refused. Other treatment was not administered. It was recommended by the patient's physician, but this treatment (which would have been coded 1, 2, or 3) was refused by the patient, a patient's family member, or the patient's guardian. The refusal was noted in the patient record.
8	Recommended; unknown if administered. Other treatment was recommended, but it is unknown whether it was administered.
9	Unknown. It is unknown whether other treatment was recommended or administered, and there is no information in the medical record to confirm the recommendation or administration of other treatment or death certificate only.

Coding Examples:

Code	Explanation
1	For head and neck primaries: Ideally, an embolic agent is chosen that will block the very small vessels within the tumor but spare the adjacent normal tissue. Liquid embolic agents, such as ethanol or acrylic, and powdered particulate materials can penetrate into the smallest blood vessels of the tumor.
6	DC vax given for brain cancer. Assign code 6. DC vax is not an approved treatment for brain cancer and should not be coded in the immunotherapy or any of the other treatment data items.
6	Cannabis oil or medical marijuana that is used for treatment.

RX DATE—OTHER

Item Length: 8
 NAACCR Item #1250
 NAACCR Name: RX Date Other
 XML NAACCR ID: rxDateOther

Definition:

Records the date on which other treatment began at any facility.

Coding Instructions:

- Other treatment is that which cannot be defined as surgery, radiation, or systemic therapy according to the defined data items in this manual.
- If other treatment is the first or only treatment administered to the patient, then the date other treatment started should be the same as the Date of First Course of Treatment.

Code	Definition
YYYYMMDD YYYYMM YYYY	The month, day, and year other treatment began at any facility. The first two digits are the month, the third and fourth digits are the day, and the last four digits are the year.
blank	When no other treatment was administered; other treatment was administered but the date is completely unknown; unknown whether other treatment administered; diagnosed at autopsy.

Coding Examples:

Code	Explanation
20110404	A patient with metastatic disease was started on an experimental therapy on April 4, 2011.
201006	In June 2010, a patient started treatment which cannot be defined as surgery, radiation, or systemic therapy according to the defined data items in this manual.

RX DATE – OTHER FLAG

Item Length: 2
NAACCR Item #1251
NAACCR Name: Rx Date Other Flag
XML NAACCR ID: rxDateOtherFlag

Definition:

This flag explains why no appropriate value is in the field, RX Date-Other. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if RX Date - Other has a full or partial date recorded.
- Assign code 10 when it is unknown whether any other treatment was given.
- Assign code 11 if no other treatment is planned or given.
- Assign code 12 if RX Date – Other cannot be determined, but it is known that other treatment was given as part of first course of treatment.
- Assign code 15 if other treatment is planned, but has not yet started and the start date is not yet available.

Code	Explanation
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any other treatment was given)
11	No proper value is applicable in this context (for example, no other treatment was given)
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, other therapy was given but the date is unknown).
15	Other therapy is planned as part of the first course of treatment, but had not been started at the time of the most recent follow-up.
(blank)	A valid date value is provided in item RX Date-Other.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for RX Date-Other
Blank	Partial date is known (YYYYMM or YYYY) for RX Date-Other
10	Unknown if any other treatment given
11	No other treatment given
12	Other therapy given as first course treatment but date is completely unknown.

RX SUMM—RX STATUS

Item Length: 1

NAACCR Item #1285

NAACCR Name: RX Summ—Treatment Status

XML NAACCR ID: rxSummTreatmentStatus

Description

Treatment Status documents active surveillance (watchful waiting). Before this data item was implemented, active surveillance or watchful waiting was deduced from the codes in each of the treatment fields.

Coding Instructions

- This item may be left blank for cases diagnosed prior to 2010.
- Treatment administered after a period of active surveillance is considered subsequent treatment and is not coded in this item.
- Use code 0 when the patient does not receive any treatment.
 - Scope of Regional Lymph Node Surgery may be coded 0, 1-7, or 9.
- Assign code 1 when the patient receives treatment collected in any of the following data items
 - Surgery of Primary Site
 - Surgical Procedure of Other Site
 - Radiation Treatment Modality, Phase I
 - Chemotherapy
 - Hormone Therapy
 - Immunotherapy
 - Hematologic Transplant and Endocrine Procedures
 - Other Therapy
- Assign code 2 when there is documentation the patient is being monitored using active surveillance/watchful waiting/deferred therapy or other similar options.

Code	Explanation
0	No treatment given
1	Treatment given
2	Active surveillance (watchful waiting)
9	Unknown if treatment was given

Coding Examples:

Code	Explanation
0	Patient with metastatic lung cancer requested to be placed on Hospice.
9	Patient was diagnosed at your facility and no treatment is planned. No information is available on whether or not the patient went on to have treatment.
2	Treatment plan for a prostate cancer patient is active surveillance.

OVER-RIDES/CONVERSION SYSTEM ADMIN

EDITS AND OVER-RIDES:

Text must be included to justify overrides that have been set. Some of the edits identify rare, but possible, code combinations. For these edits, an override flag can be set if, upon review, the unusual combination is verified as being correct. Once set, the error message will not be repeated on subsequent EDITS passes.

- When no error message is generated by an edit that uses an override item, no action by the registrar is needed.
- If an error message is generated, the problem can often be resolved by checking the accuracy of the entry for each item that contributes to the edit and correcting any problems identified. If correction of data entry errors resolves the problem, no override entry is needed. If the codes reflect the information in the patient record, check for physician notes indicating the unusual combination of circumstances (for example, a colon adenocarcinoma in a child) has been confirmed.
- Enter the override code according to the instructions for the data item. If no comment regarding the unusual circumstances can be found in the record, it may be necessary to check with the managing physician or pathologist to determine whether it is appropriate to override the edit.

OVERRIDE AGE/SITE/MORPH

Item Length: 1

NAACCR Item #1990

NAACCR Name: Over-ride Age/Site/Morph

XML NAACCR ID: overRideAgeSiteMorph

Definition

This override is used with the following edits:

Age, Primary Site, Morphology ICD02 (SEER IF15)

Age, Primary Site, Morphology ICD03 (SEER IF15)

Age, Primary Site, Morph ICD03 – Adult (SEER)

Age, Primary Site, Morph ICD03 – Pediatric (NPCR)

Edits of the type, *Age, Primary Site, Morphology* differ in using ICD-O-2 or ICD-O-3 morphologies, and require review if a site-morphology combination occurs in an age group for which it is extremely rare:

If the edit generates an error or warning message, check that the primary site and histologic type are coded correctly and that the age, date of birth, and date of diagnosis are correct.

Coding Instructions

1. Leave blank if the program does not generate an error message for one of the edits listed above.
2. Leave blank and correct any errors for the case if an item is discovered to be incorrect.
3. Code 1 as indicated if review of items in the error or warning message confirms that all are correct.
4. Use code 2 if the case was diagnosed in utero.
5. If both codes 1 and 2 would apply, use code 3.

Codes

Code	Definition
1	Reviewed: An unusual occurrence of a particular age/site/histology combination for a given age group has been reviewed.
2	Reviewed; diagnosis in utero.
3	Reviewed; both conditions apply.
Blank	Not reviewed or reviewed and corrected.

OVERRIDE HISTOLOGY

Item Length: 1
NAACCR Item #2040
NAACCR Name: Over-ride Histology
XML NAACCR ID: overRideHistology

Definition

This override is used with the following edits:

Diagnostic Confirmation, Behavior ICDO2 (SEER IF31)
Diagnostic Confirmation, Behavior ICDO3 (SEER IF31)
Morphology -- Type/Behavior ICDO2 (SEER MORPH)
Morphology – Type/Behavior ICDO3 (SEER MORPH)

Edits of the type *Diagnostic Confirmation, Behavior Code* differ in the use of ICD-O-2 or ICD-O-3 and check that, for *in situ* cases (Behavior = 2), Diagnostic Confirmation specifies microscopic confirmation (1, 2, or 4). The distinction between *in situ* and invasive is very important to a registry, since prognosis is so different. Since the determination that a neoplasm has not invaded surrounding tissues, i.e., *in situ*, is made microscopically, cases coded *in situ* in behavior should have a microscopic confirmation code. However, very rarely, a physician will designate a case noninvasive or *in situ* without microscopic evidence.

Edits of the type, Morphology – Type/Behavior, perform the following check:

1. Codes listed in ICD-O-2 or ICD-O-3 with behavior codes of only 0 or 1 are considered valid, since the behavior matrix of ICD-O-2 and ICD-O-3 allows for the elevation of the behavior of such histologies when the tumor is *in situ* or malignant. This edit forces review of these rare cases to verify that they are indeed *in situ* or malignant.
2. The following histologies are generally not accepted as *in situ*: ICD-O-2 histologies 8000-8004, 8020, 8021, 8331, 8332, 8800-9054, 9062, 9082, 9083, 9110-9491, 9501-9989, ICD-O-3 histologies 8000-8005, 8020, 8021, 8331, 8332, 8800-9055, 9062, 9082, 9083, 9110-9493, 9501-9989. This edit forces review of these cases.
3. If a *Morphology – Type/Behavior* edit produces an error or warning message and the case is one in which the 4-digit morphology code is one that appears in ICD-O-2 or ICD-O-3 only with behavior codes of 0 or 1, verify the coding of morphology and that the behavior should be coded malignant or *in situ*. The registrar may need to consult a pathologist or medical advisor in problem cases.

Exceptions:

If year of *Date of Diagnosis* > 2000, then a behavior code of 1 is valid for the following ICD-O-2 histologies and no override flag is needed: 8931, 9393, 9538, 9950, 9960-9962, 9980-9984, and 9989. Similarly, the following ICD-O-3 histologies are valid with a behavior code of 1: 8442, 8451, 8462, 8472, and 8473.

If year of *Date of Diagnosis* > 2003, the following ICD-O-3 benign histologies will pass without review: 8146, 8271, 8861, 8897, 9121, 9122, 9131, 9161, 9350, 9351, 9352, 9360, 9361, 9383, 9384, 9394, 9412, 9413, 9444, 9492, 9493, 9506, 9531, 9532, 9533, 9534, 9537, 9541, 9550, 9562, and 9570.

Coding Instructions

- Leave blank if no edit is generated of either type.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1, 2, or 3 as indicated if review of all items in the error or warning message confirms that all are correct.

Codes	Explanation
1	Reviewed: The behavior code of the histology is designated as “benign” or “uncertain” in ICD-O-2 or ICD-O-3, and the pathologist states the primary to be “ <i>in situ</i> ” or “malignant” (flag for a “Morphology Type & Behavior” edit)
2	Reviewed: The behavior code is “ <i>in situ</i> ,” but the case is not microscopically confirmed (flag for a “Diagnostic Confirmation, Behavior Code”)
3	Reviewed: Conditions 1 and 2 above both apply
Blank	Not reviewed or reviewed and corrected

OVERRIDE LEUK, LYMPHOMA

Item Length: 1

NAACCR Item #2070

NAACCR Name: Over-Ride Leuk, Lymphoma

XML NAACCR ID: overRideLeukLymphoma

Definition

This override is used with the following edits:

Diagnostic Confirmation, Histology ICDO2 (SEER IF48)

Diagnostic Confirmation, Histol Typ ICDO3 (SEER IF48)

Edits of the type Diagnostic Confirmation, Histology differ in use of ICD-O-2 or ICD-O-3 and check the following:

1. Since lymphoma and leukemia are almost exclusively microscopic diagnoses, this edit forces review of any cases of lymphoma that have diagnostic confirmation of direct visualization or clinical, and any leukemia with a diagnostic confirmation of direct visualization.
2. If histology = 9590-9717 for ICD-O-2 or 9590-9729 for ICD-O-3 (lymphoma) then Diagnostic Confirmation cannot be 6 (direct visualization) or 8 (clinical).
3. If histology=9720-9941 for ICD-O-2 or 9731-9948 for ICD-O-3 (leukemia and other) then Diagnostic Confirmation cannot be 6 (direct visualization).

Coding Instructions

- Leave blank if no edit is generated.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- If the edit produces an error or warning message, verify that the Histologic Type and Diagnostic Confirmation are correctly coded. Remember that positive hematologic findings and bone marrow specimens are included as histologic confirmation (code 1 in Diagnostic Confirmation) for leukemia. Code 1 for the override indicates that a review has taken place and histologic type and diagnostic confirmation are correctly coded.

Code	Explanation
1	Reviewed
Blank	Not reviewed or reviewed and corrected.

OVERRIDE SITE/BEHAVIOR

Item Length: 1
NAACCR Item #2071
NAACCR Name: Over-Ride Site/Behavior
XML NAACCR ID: overRideSiteBehavior

Definition

This override is used with the following edits:

Primary Site, Behavior Code ICDO2 (SEER IF39)

Primary Site, Behavior Code ICDO3 (SEER IF39)

Edits of the type, Primary Site, Behavior Code, require review of the following primary sites with a behavior of *in situ* (ICD-O-2 or ICD-O-3 behavior = 2):

C269	Gastrointestinal Tract, NOS
C399	Ill-defined sites with respiratory system
C559	Uterus, NOS
C579	Female genital tract, NOS
C639	Male genital organs, NOS
C689	Urinary system, NOS
C729	Nervous system, NOS
C759	Endocrine gland, NOS
C760-C768	Ill-defined sites
C809	Unknown primary site

Since the designation of *in situ* is very specific and almost always requires microscopic confirmation, ordinarily specific information should also be available regarding the primary site. Conversely, if inadequate information is available to determine a specific primary site, it is unlikely that information about a cancer being *in situ* is reliable. If an *in situ* diagnosis is stated, try to obtain a more specific primary site. A primary site within an organ system site can sometimes be identified based on the diagnostic procedure or treatment given or on the histologic type. If no more specific site can be determined, it is usually preferable to code a behavior code of 3. In the exceedingly rare situation in which it is certain that the behavior is *in situ* and no more specific site code is applicable, set Over-ride Site/Behavior to 1.

Coding Instructions

- Leave blank if no edit is generated
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.

- Code 1 if review of site and behavior verifies that the patient has an *in situ* cancer of a nonspecific site and no further information about the primary site is available.

Code	Explanation
1	Reviewed
Blank	Not reviewed or reviewed and corrected.

OVERRIDE SITE/TYPE

Item Length: 1
NAACCR Item #2030
NAACCR Name: Over-Ride Site/Type
XML NAACCR ID: overRideSiteType

Definition

This override is used with the following edits:

- Primary Site, Morphology-Type ICDO2 (SEER IF25)
- Primary Site, Morphology-Type ICDO3 (SEER IF25)
- Primary Site, Morphology-Type, Behavior ICDO3 (SEER IF25)

There are multiple versions of edits of the type, *Primary Site, Morphology-Type*, which check for “usual” combinations of site and ICD-O-2 or ICD-O-3 histology. The SEER version of the edit is more restrictive than the CoC edit, and thus uses a different override flag. The CoC version of the edit will accept Override CoC-Site/Type or Override Site/Type as equivalent.

- The Site/Histology Validation List (available on the SEER website) contains those histologies commonly found in the specified primary site. Histologies that occur only rarely or never are not included. These edits require review of combinations **not** listed.

Review of these cases requires investigating whether the combination is biologically implausible or there are cancer registry coding conventions that would dictate different codes for the diagnosis. Review of these rare combinations often results in changes to the primary site and/or morphology, rather than a decision that the combination is correct.

Coding Instructions

1. Leave blank if the program does not generate an error message for the edits of the type Primary Site, Morphology-Type.
2. Leave blank and correct any errors for the case if an item is discovered to be incorrect.
3. Code 1 if the case has been reviewed and both the site and histology are correct.

Codes

Code	Definition
1	Reviewed: An unusual occurrence of a particular age/site/histology combination for a given age group has been reviewed.
Blank	Not reviewed or reviewed and corrected.

OVERRIDE SURG/DXCONF

Item Length: 1
NAACCR Item #2020
NAACCR Name: Over-Ride Surg/DxConf
XML NAACCR ID: overRideSurgDxconf

Definition

This override is used with the following edits:

RX Summ – Surg Prim Site, Diag Conf (SEER IF76)

This edit checks that cases with a primary site surgical procedure coded 20-90 are histologically confirmed. If the patient had a surgical procedure, most likely there was a microscopic examination of the cancer.

Coding Instructions

- Verify the surgery and diagnostic confirmation codes, and correct any errors.
- Sometimes there are valid reasons why no microscopic confirmation is achieved with the surgery, for example, the tissue removed may be inadequate for evaluation.
- Leave blank if the program does not generate an error message for the edit *RX Summ - Surg Prim Site, Diag Conf*.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of all items in the error or warning message confirms that all are correct.

Code	Definition
1	Reviewed
Blank	Not reviewed or reviewed and corrected

OVERRIDE HOSPSEQ/DXCONF

Item Length: 1

NAACCR Item #1986

NAACCR Name: Over-Ride HospSeq/DxConf

XML NAACCR ID: overRideHospseqDxconf

Definition

This override is used with the following edits:

Diagnostic Confirm, Seq Num – Hosp (CoC)

The edit, Diagnostic Confirm, Seq Num – Hosp (CoC), does the following:

1. If any case is one of multiple primaries and is not microscopically confirmed or lacks a positive lab test/marker study, i.e., Diagnostic Confirmation > 5 and Sequence Number – Hospital > 00 (more than one primary), review is required.
2. If Primary Site specifies an ill-defined or unknown primary (C760-C768, C809), no further checking is done.
3. If Sequence Number – Hospital is in the range of 60-88, this edit is skipped.

It is important to verify that the non-microscopically confirmed case is indeed a separate primary from any others that may have been reported. This edit forces review of multiple primary cancers when one of the primaries is coded to a site other than ill-defined or unknown and is not microscopically confirmed or confirmed by a positive lab test/marker study.

1. If the suspect cases are confirmed accurate as coded and if the number of primaries is correct, set the Over-ride HospSeq/DxConf to 1. Do not set the over-ride flag on the patient's other primary cancers.
2. If it turns out that the non-microscopically confirmed cancer is considered a manifestation of one of the patient's other cancers, delete the non-microscopically confirmed case. Check the sequence numbers of remaining cases, correcting them if necessary. Also check for other data items on the remaining cases that may need to be changed as a result of the corrections, such as stage and treatment.

Coding Instructions

- Leave blank if the program does not generate an error message for the edit Diagnostic Confirm, Seq Num – Hosp (CoC).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.

- Code 1 if review of all items in the error or warning message confirms that all are correct.

Code	Explanation
1	Reviewed
Blank	Not reviewed or reviewed and corrected.

OVERRIDE HOSPSEQ/SITE

Item Length: 1
NAACCR Item #1988
NAACCR Name: Over-Ride HospSeq/Site
XML NAACCR ID: overRideHospseqSite

Definition

This override is used with the following edits:

Seq Num – Hosp, Primary Site, Morph ICDO2 (CoC)

Seq Num – Hosp, Primary Site, Morph ICDO3 (CoC)

Edits of this type review of multiple primary cancers when one of the primaries is coded to a site/morphology combination that could indicate a metastatic site rather than a primary site.

1. If Sequence Number – Hospital indicates the person has had more than one primary, then any case with one of the following site/histology combinations requires review:
 - C760-C768 (ill-defined sites) or C809 (unknown primary) and IDC-O-2 or ICD-O-3 histology <9590. Look for evidence that the unknown or ill-defined primary is a secondary site from one of the patient’s other cancers. For example, a clinical discharge diagnosis of “abdominal carcinomatosis” may be attributable to the patient’s primary ovarian cystadenocarcinoma already in the registry, and should not be entered as a second primary.
 - C770-C779 (lymph nodes) and ICD-O-2 histology not in the range 9590-9717 or ICD-O-3 histology not in the range 9590-9729; or C420-C424 and ICD-O-2 histology not in the range 9590-9941 or ICD-O-3 not in the range 9590-9989. That combination is most likely a metastatic lesion. Check whether the lesion could be a manifestation of one of the patient’s other cancers.
 - Any site and ICD-O-2 histology in the range 9720-9723, 9740-9741, or ICD-O-3 histology in the range 9740-9758. Verify that these diagnoses are coded correctly and are indeed separate primaries from the others.
2. If it turns out that the suspect tumor is a manifestation of one of the patient’s other cancers, delete the metastatic or secondary case, re-sequence remaining cases, and correct the coding on the original case as necessary.

Coding Instructions

- Leave blank if the program does not generate an error message for an edit of the type Seq Num – Hosp, Primary Site, Morph.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.

- Code 1 if review of all items in the error or warning message confirms that hospital sequence number and site are both correct.

Code	Explanation
1	Reviewed
Blank	Not reviewed or reviewed and corrected.

OVERRIDE NAME/SEX

Item Length: 1
NAACCR Item #2078
NAACCR Name: Over-Ride Name/Sex
XML NAACCR ID: overRideNameSex

Definition

This over-ride is used with the following edit in the NAACCR Metafile of the EDITS software:

Sex, Name-First, Date of Birth (NAACCR)

This edit will be triggered by extremely rare or nonexistent combination of first name and sex, such as John/female.

Coding Instructions

- Verify that the sex is correct for the patient.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of all items in the error or warning message confirms that all are correct.
- Leave blank if the program does not generate an error message for the edit Sex, Name-First, Date of Birth (NAACCR).

Code	Definition
1	Reviewed
Blank	Not reviewed or reviewed and corrected

FOLLOW-UP/RECURRENCE/DEATH

DATE OF LAST CONTACT

Item Length: 8

NAACCR #1750

NAACCR Name: Date of Last Contact

XML NAACCR ID: dateOfLastContact

Definition:

Records the date of last contact with the patient or the date of death.

Coding Instructions:

- Record the last date on which the patient was known to be alive or the date of death.

- If a patient has multiple primaries, all records should have the same date of last contact.

Code	Definition
YYYYMMDD	The date of last contact is the year, month and day that last contact was made. The first four digits are the year, the fifth and sixth digits are the day, and the last two digits are the year.
YYYYMM	
YYYY	

Examples:

Code	Explanation
20040630	The patient's date of death was June 30, 2004.
2004	The medical record contains only the year of death (2003). Check the social security death index on line and correct the date if the patient expired.
20050114	A patient returns his follow-up inquiry with no date information, the Envelope is postmarked January 14, 2005.

DATE OF LAST CONTACT FLAG

Item Length: 2
NAACCR Item #1751

NAACCR Name: Date of Last Contact Flag
XML NAACCR ID: dateOfLastContactFlag

Definition:

This flag explains why no appropriate value is in the field, Date of Last Contact. Before Version 12 (through 2009 diagnosis), date fields included codes that provided information other than dates. As part of an initiative to standardize date fields, new fields were introduced to accommodate non-date information that had previously been transmitted in date fields.

Coding Instructions

- Leave this item blank if Date of Last Contact has a full or partial date recorded.
- Assign code 12 if Date of Last Contact cannot be determined.

Code	Explanation
12	A proper value is applicable but not known. This event occurred, but the date is unknown (for example, the date of last contact is unknown).
(blank)	A valid date value is provided in item Date of Last Contact.

Coding Examples:

Code	Explanation
Blank	Full date is known (YYYYMMDD) for Date of Last Contact
Blank	Partial date is known (YYYYMM or YYYY) for Date of Last Contact
12	Date of Last Contact is completely unknown.

VITAL STATUS

Item Length: 1
NAACCR Item #1760
NAACCR Name: Vital Status
XML NAACCR ID: vitalStatus

Definition:

Records the vital status of the patient as of the date entered in Date of Last Contact or Death.

Coding Instructions:

- This item is collected during the follow-up process with Date of Last Contact or Death.
- If a patient has multiple primaries, all records should have the same vital status.

Code	Definition
0	Dead
1	Alive

Coding Examples:

Code	Explanation
0	Death clearance information obtained from the CCR confirms the death of the patient within the past year.
1	In response to a follow-up letter to a patient's following physician, it is learned the patient is alive.

CAUSE OF DEATH

Item Length: 4
NAACCR Item #1910
NAACCR Name: Cause of Death
XML NAACCR ID: causeOfDeath

Description

Official cause of death as coded from the death certificate in valid ICD-7, ICD-8, ICD-9, and ICD-10 codes.

Coding Instructions

Beginning with 1999, Mississippi Death Certificates are coded using ICD-10 codes. Cause of death should be coded in ICD-10. If the ICD-10 cause of death **from the death certificate** is not known, please code cause of death as 7777.

Do not code what you think the cause of death is, only what is specifically stated as the underlying cause of death on the patient's death certificate.

Do not make the cause of death more specific. For example, the ICD-10 code for prostate cancer is C61. Do not code C619 because that is what we code for primary site in OCD-O-3.

If the cause of death is less specific than your primary site, code the cause of death as recorded on the death certificate. For example, cause of death is C349 but you have C341 as the primary site, the cause of death should be coded C349.

Special codes in addition to ICD-7, ICD-8, ICD-9, and ICD-10

Code	
0000	Patient alive at last contact
7777	State death certificate not available
7797	State death certificate available but underlying cause of death is not coded.

PLACE OF DEATH STATE

Item Length: 2

NAACCR Item #1942

NAACCR Name: Place of Death—State

XML NAACCR ID: placeOfDeathState

Description

USPS abbreviation for state, commonwealth, U.S. possession in which the patient died and the death certificate is filed. CanadaPost abbreviations for the Canadian provinces can also be recorded if the patient died in Canada. If the patient has multiple primaries, this data item should be coded the same for each primary.

Coding Instructions

This data item is left blank if the patient is still alive. Otherwise, use the most specific code from the table below.

Code	Definition	Code	Definition
AL	Alabama	MI	Michigan
AK	Alaska	MN	Minnesota
AZ	Arizona	MS	Mississippi
AR	Arkansas	MO	Missouri
CA	California	MT	Montana
CO	Colorado	NE	Nebraska
CT	Connecticut	NV	Nevada
DE	Delaware	NH	New Hampshire
DC	District of Columbia	NJ	New Jersey
FL	Florida	NM	New Mexico
GA	Georgia	NY	New York
HI	Hawaii	NC	North Carolina
ID	Idaho	ND	North Dakota
IL	Illinois	OH	Ohio
IN	Indiana	OK	Oklahoma
IA	Iowa	OR	Oregon
KS	Kansas	PA	Pennsylvania
KY	Kentucky	RI	Rhode Island
LA	Louisiana	SC	South Carolina
ME	Maine	SD	South Dakota
MD	Maryland	TN	Tennessee
MA	Massachusetts	TX	Texas

Code	Definition	Code	Definition
UT	Utah	ZZ	U.S., NOS; Canada, NOS; Country Unknown
VT	Vermont	AB	Alberta
VA	Virginia	BC	British Columbia
WA	Washington	MB	Manitoba
WV	West Virginia	NB	New Brunswick
WI	Wisconsin	NL	Newfoundland and Labrador
WY	Wyoming	NS	Nova Scotia
AS	American Samoa	NT	Northwest Territories
GU	Guam	NU	Nunavut
MP	Northern Mariana Islands	ON	Ontario
PW	Palau	PE	Prince Edward Island
PR	Puerto Rico	QC	Quebec
UM	U.S. Outlying Islands	SK	Saskatchewan
VI	Virgin Islands of the United States	US	Resident of United States, NOS
FM	Federated States of Micronesia	AA	APO/FPO for Armed Services America
MH	Marshall Islands	AE	APO/FPO for Armed Services Europe
TT	Trust Territories	AP	APO/FPO for Armed Services Pacific
XX	Country Known, Not U.S., Not Canada		
YT	Yukon Territories		
YY	Country Unknown, Not U.S., Not Canada		

PLACE OF DEATH COUNTRY

Item Length: 3

NAACCR Item #1944

NAACCR name: Place of Death—Country

XML NAACCR ID: placeOfDeathCountry

Description

Records the country in which the patient died and where the death certificate is filed.

Coding Instructions

Leave blank if the patient is alive. Otherwise, use the most specific code from the table below.

Code	Definition	Code	Definition
ABW	Aruba	BGD	Bangladesh
AFG	Afghanistan	BGR	Bulgaria
AGO	Angola	BHR	Bahrain
AIA	Anguilla	BHS	Bahamas
ALA	Aland Islands	BIH	Bosnia and Herzogovina
ALB	Albania	BLM	St. Barthelemy
AND	Andorra	BLR	Belarus
ARE	United Arab Emirates	BLZ	Belize
ARG	Argentina	BMU	Bermuda
ARM	Armenia	BOL	Bolivia
ASM	American Samoa	BRA	Brazil
ATA	Antarctica	BRB	Barbados
ATF	French Southern Territories	BRN	Brunei
ATG	Antigua and Barbuda	BTN	Bhutan
AUS	Australia	BVT	Bouvet Island
AUT	Austria	BWA	Botswana
AZE	Azerbaijan	CAF	Central African Republic
BDI	Burundi	CAN	Canada
BEL	Belgium	CHE	Switzerland
BEN	Benin	CHL	Chile
BES	Bonaire, Saint Eustatius and Saba	CHN	China
BFA	Burkina Faso	CIV	Cote d'Ivoire

Code	Definition	Code	Definition
CMR	Cameroon	FSM	Micronesia
COD	Congo, Democratic Republic of	GAB	Gabon
COG	Congo	GBR	United Kingdom
COK	Cook Islands	GEO	Georgia
COL	Columbia	GGY	Guernsey
COM	Comoros	GHA	Ghana
CPV	Cape Verde	GIB	Gibraltar
CRI	Costa Rica	GIN	Guinea
CSK	Czechoslovakia	GLP	Guadelupe
CUB	Cuba	GMB	Gambia
CUW	Curacao	GNB	Guinea Bissau
CXR	Christmas Island	GNQ	Equatorial Guinea
CYM	Cayman Islands	GRC	Greece
CYP	Cyprus	GRD	Grenada
CZE	Czech Republic	GRL	Greenland
DEU	Germany	GTM	Guatemala
DJI	Djibouti	GUF	French Guiana
DMA	Dominica	GUM	Guam
DNK	Denmark	GUY	Guyana
DOM	Dominican Republic	HKG	Hong Kong
DZA	Algeria	HMD	Heard Island & McDonalds Islands
ECU	Ecuador	HND	Honduras
EGY	Egypt	HRV	Croatia
ENG	England	HTI	Haiti
ERI	Eritrea	HUN	Hungary
ESH	Western Sahara	IDN	Indonesia (Dutch East Indies)
ESP	Spain	IMN	Isle of Man
EST	Estonia	IND	India
ETH	Ethiopia	IOT	British Indian Ocean Territory
FIN	Finland	IRL	Ireland
FJI	Fiji	IRN	Iran
FLK	Falkland Islands	IRQ	Iraq
FRA	France	ISL	Iceland
FRO	Faroe Islands	ISR	Israel

Code	Definition	Code	Definition
ITA	Italy	MKD	Macedonia
JAM	Jamaica	MLI	Mali
JEY	Jersey	MLT	Malta
JOR	Jordan	MMR	Myanmar
JPN	Japan	MNE	Montenegro
KAZ	Kazakhstan	MNG	Mongolia
KEN	Kenya	MNP	Northern Mariana Islands
KGZ	Kyrgyzstan	MOZ	Mozambique
KHM	Cambodia	MRT	Mauritania
KIR	Kiribati	MSR	Montserrat
KNA	St. Kitts and Nevis	MTQ	Martinique
KOR	Korea, NOS	MUS	Mauritius
KOR	South Korea	MWI	Malawi
KWT	Kuwait	MYS	Malaysia
LAO	Laos	MYT	Mayotte
LBN	Lebanon	NAM	Namibia
LBR	Liberia	NCL	New Caledonia
LBY	Libya	NER	Niger
LCA	St. Lucia	NFK	Norfolk Island
LIE	Liechtenstein	NGA	Nigeria
LKA	Sri Lanka	NIC	Nicaragua
LSO	Lesotho	NIR	Northern Ireland (Ulster)
LTU	Lithuania	NIU	Niue
LUX	Luxembourg	NLD	Netherlands
LVA	Latvia	NOR	Norway
MAC	Macao	NPL	Nepal
MAF	Saint Martin (French part)	NRU	Nauru
MAR	Morocco	NZL	New Zealand
MCO	Monaco	OMN	Oman
MDA	Moldova	PAK	Pakistan
MDG	Madagascar	PAN	Panama
MDV	Maldives	PCN	Pitcairn Islands
MEX	Mexico	PER	Peru
MHL	Marshall Islands	PHL	Philippines

Code	Definition	Code	Definition
PLW	Palau(Trust Territory of Pacific Islands)	SVN	Slovenia
PNG	Papua New Guinea	SWE	Sweden
POL	Poland	SWZ	Swaziland
PRI	Puerto Rico	SXM	Sint-Maarten
PRK	North Korea	SYC	Seychelles
PRT	Portugal	SYR	Syria
PRY	Paraguay	TCA	Turks and Caicos
PSE	Palestine Territory, Occupied	TCD	Chad
PYF	French Polynesia	TGO	Togo
QAT	Qatar	THA	Thailand
REU	Réunion	TJK	Tajikistan
ROU	Romania	TKL	Tokelau Islands (New Zealand)
RUS	Russia	TKM	Turkmenistan
RWA	Rwanda	TLS	Timor-Leste
SAU	Saudi Arabia	TON	Tonga
SCT	Scotland	TTO	Trinidad and Tobago
SDN	Sudan	TUN	Tunisia
SEN	Senegal	TUR	Turkey
SGP	Singapore	TUV	Tuvalu
SGS	S Georgia & S Sandwich Islands	TWN	Taiwan
SHN	St Helena	TZA	Tanzania
SJM	Svalbard & Jan Mayen	UGA	Uganda
SLB	Solomon Islands	UKR	Ukraine
SLE	Sierra Leon	UMI	U.S. Minor Outlying Islands
SLV	El Salvador	URY	Uruguay
SMR	San Marino	USA	United States
SOM	Somalia	UZB	Uzbekistan
SPM	St Pierre and Miquelon	VAT	Vatican City
SRB	Serbia	VCT	St. Vincent & the Grenadines
SSD	South Sudan	VEN	Venezuela
STP	Sao Tome & Principe	VGB	British Virgin Islands
SUR	Suriname	VIR	U.S. Virgin Islands
SVK	Slovakia	VNM	Vietnam

Code	Definition	Code	Definition
VUT	Vanuatu		
WLF	Wallis and Fotuna		
WLS	Wales		
WSM	Samoa		
YEM	Yemen		
YUG	Yugoslavia		
ZAF	Republic of South Africa		
ZMB	Zambia		
ZWE	Zimbabwe		
ZZA	Asia, NOS		
ZZC	Central America, NOS		
ZZE	Europe, NOS		
ZZF	Africa, NOS		
ZZN	North America, NOS		
ZZP	Pacific, NOS		
ZZS	South America, NOS		
ZZU	Unknown		
ZZX	Non-US/Canada, NOS		

ICD REVISION NUMBER

Item Length: 1
NAACCR Item #1920
NAACCR Name: ICD Revision Number
XML NAACCR ID: icdRevisionNumber

Description

Indicator for the coding scheme used to code the cause of death.

NOTE: If the patient is deceased, please use code 1 (ICD-10). See note under “Cause of Death” regarding the use of ICD-9 for cause of death.

Coding Instructions

Code	Definition
0	Patient alive at last follow-up
1	ICD-10
7	ICD-7
8	ICDA-8
9	ICD-9

AUTOPSY

Item Length: 1
NAACCR Item #1930
NAACCR Name: Autopsy
XML NAACCR ID: autopsy

Description

Code indicating whether or not an autopsy was performed.

Coding Instructions:

Code	Description
0	Not applicable; patient alive
1	Autopsy performed
2	No autopsy performed
9	Patient expired, unknown if autopsy performed

DEATH CERTIFICATE NUMBER

Item Length: 6
NAACCR Item #2380
NAACCR Name: DC State File Number
XML NAACCR ID: dcStateFileNumber

Description

Death certificate number as assigned by the vital statistics office in the place recorded in *Place of Death* (NAACCR Item #1940)

TEXT--DIAGNOSIS

TEXT--DX PROC – PE

Item Length: 1000
NAACCR Item #2520
NAACCR Name: Text—DX Proc—PE
XML NAACCR ID: textDxProcPe

Description:

Information relating to the diagnosis of this cancer discovered during physical examination at the time of admission, or documented in the admitting note or History and Physical. Also review consultative reports.

Suggestions for text:

History that relates to this cancer diagnosis
Age, sex, race/ethnicity
Date of physical exam in physician's office
Family History, Alcohol History, Tobacco History
Personal cancer history with dates.
Tumor location
Tumor Size
Palpable lymph nodes
Histology (if diagnosis prior to this admission)
Record positive and negative clinical findings. Record positive results first,
Impression (when stated and pertains to cancer diagnosis)
Treatment plan

Examples:

Elderly black female

- 95 YO BF

Multiple primaries

- 01 prostate 1993, 02 melanoma 2000, this is 3rd primary

Admitting/H & P

- MD noted suspicious lesion, L leg. No lymphadenopathy or skin ulceration, Bx in MD office pos for melanoma
- History of Rt breast cancer 1985, NED
- Large prostate nodule in Rt lobe felt during rectal exam at Dr Record's office.

TEXT--DX PROC - X-RAY/SCAN

Item Length: 1000

NAACCR Item #2530

NAACCR Name: Text—DX Proc—X-Ray/Scan

XML NAACCR ID: textDxProcXRayScan

Description

Record x-rays or scans that were performed.

Suggestions for Text

Date(s) of x-ray/scans

Type(s) of x-rays/scans

Location of tumor

Primary Site

Histology (if given)

Size of tumor

Lymph nodes

Metastatic disease

Record both positive and negative findings. Record positive results first.

Examples:

- 3/6/2004: Chest Xray, large mass, LUL
- 3/8/2004: lung CT, highly suspicious large mass LUL, extension to mediastinum
- 3/10/2004: Bone Scan-Neg

TEXT--DX PROC – SCOPES

Item Length: 1000
NAACCR Item #2540
NAACCR Name: Text—DX Proc—Scopes
XML NAACCR ID: textDxProcScopes

Description

Record endoscopic examinations that were performed.

Suggestions for text

Date(s) of endoscopic exam(s)

Primary Site

Histology (if given)

Tumor location

Tumor size

Lymph nodes

Record site and type of endoscopic biopsy

Record positive and negative clinical findings. Record positive first.

Examples:

- 6/1/2004: colonoscopy, obstructing lesion of sigmoid, pos. bx,

TEXT--DX PROC – LAB TESTS

Item Length: 1000

NAACCR Item #2550

NAACCR Name: Text—DX Proc—Lab Tests

XML NAACCR ID: textDxProcLabTests

Description

A record of the positive and negative laboratory tests that related to this cancer diagnosis.

Suggestions for text

Type of lab test/tissue specimen(s)

Record both positive and negative findings. Record positive results first.

Date(s) of laboratory test(s)

Information can include tumor markers, serum and urine electrophoresis, special studies, etc.

Tumor markers included, but are not limited to:

- Breast Cancer: Estrogen Receptor Assay (ERA), Progesterone Receptor Assay (PRA), Her2/neu (Record both the type(s) of tests, result(s) and interpretation)
- Prostate Cancer – Prostate Specific Antigen (PSA)

Example:

- 6/1/2004: PSA- 1000
- 6/1/2011: ER 85% positive, PR 5% positive, HER2/neu FISH 11.85 positive

TEXT--DX PROC—OP

Item Length: 1000

NAACCR Item #2560

NAACCR Name: Text—DX Proc—OP

XML NAACCR ID: textDxProcOp

Description

Dates and descriptions of biopsies and all other surgical procedures from which staging information was derived

Location of tumor

Number of lymph nodes removed

Size of tumor removed

Documentation of residual tumor

Evidence of invasion of surrounding areas

Both positive and negative findings during operation of tissues observed but not removed

Reason primary site surgery could not be completed.

Examples

- 5/13/07 Colonoscopy found tumor of the Ascending colon followed by a R/hemicolectomy in which the tumor appeared to be confined to the colon and was located in the ascending colon. 14 right colic nodes were removed. All other organs and structures appeared normal.

TEXT--DX PROC—PATH

Item Length: 1000

NAACCR Item #2570

NAACCR Name: Text—DX Proc—Path

XML NAACCR ID: textDxProcPath

Description

If the pathology report is a slide review or a second opinion from an outside source, such as AFIP, record any additional comments from the pathologist, including differential diagnoses and final ruling.

Suggestions for Text

Date(s) of procedure(s)

Anatomic source of specimen

Type of tissue specimen(s)

Tumor type and grade (include all modifying adjectives, i.e., predominantly, with features of, with foci of, elements of, etc.)

Gross tumor size

Extent of tumor spread

Involvement of resection margins

Number of lymph nodes involved and examined

Record both positive and negative findings. Record positive test results first.

Note if pathology report is a slide review or a second opinion from an outside source, i.e., AFIP, Mayo, etc.

Record any additional comments from the pathologist, including differential diagnoses considered and any ruled out or favored

Presence of lymph-vascular invasion

Examples:

- 5/13/07 Colonoscopy: tumor of ascending colon; positive for adenoca; grade 2
- 5/13/07: R/Hemicolectomy adenoca of cecum, 1.3 cm w/invasion into the submucosa; Margins clear; 0/5 LNS.

TEXT--PRIMARY SITE TITLE

Item Length: 1000
NAACCR Item #2580
NAACCR Name: Text—Primary Site Title
XML NAACCR ID: textPrimarySiteTitle

Description

Describe the location of the tumor with laterality, if applicable. If the tumor extends into multiple organs record the organ from which the tumor originated.

Suggestions for Text

State the specific location of the primary site, including subsite
Include available information on tumor laterality

Example:

- R UOQ Breast

- LLL lung

TEXT—HISTOLOGY TITLE

Item Length: 100
NAACCR Item #2590
NAACCR Name: Text—Histology Title
XML NAACCR ID: textHistologyTitle

Description

Record the histology with behavior and grade.

Suggestions for Text

Information on histologic type and behavior

Information on differentiation from scoring systems such as Gleason’s score, Bloom-Richardson Grade, etc.

Example:

- Poorly differentiated Adenocarcinoma

TEXT—STAGING

Item Length: 1000
NAACCR Item #2600
NAACCR Name: Text—Staging
XML NAACCR ID: textStaging

Description

Information used to establish the codes for Tumor Size Summary, TNM Staging and Summary stage, as well as, any site specific factors. Additionally, the TNM stage or summary stage should be recorded here.

Suggestions for Text

Date(s) of procedure(s), including clinical procedures that provided information for assigning stage

Organs involved by direct extension

Size of tumor

Status of margins

Number and sites of positive lymph nodes

Site of distant metastasis

Physician's comments

Example:

- Clinically 3.5 cm on imaging; Pathologically 3 cm tumor confined to the breast, negative margins; no nodal involvement clinically or pathologically; no distant mets per physical exam; Clinical: cT2cN0cM0 Stage Group IIA assigned by the managing physician; Pathologic: pT2pN0cM0 Stage group IIA assigned by the surgeon; Tumor Size Summary: 3 cm; Summary Stage: Local

TEXT—REMARKS

Item Length: 1000
NAACCR Item #2680
NAACCR Name: Text—Remarks
XML NAACCR ID: textRemarks

Description

Additional text not able to fit in the above text fields

Suggestions for Text

Smoking history

Family and personal history of cancer

Comorbidities

Information on sequence number if a person was diagnosed with another primary(s) prior to the one being reported. Include type of cancer and date.

Place of birth

Justification of over-ride flags

Information clarifying anything unusual such as a reason for reporting a case seemingly not reportable for that facility or reason for coding numerous fields as “unknown.”

Examples

- Pt. seen by family phys. On 4/20/05 concerning lump found by self- examination.

TEXT--PLACE OF DIAGNOSIS

Item Length: 50

NAACCR Item #2690

NAACCR Name: Text—Place of Diagnosis

XML NAACCR ID: textPlaceOfDiagnosis

Description

Record where the patient was diagnosed

Suggestions for Text

The complete name of the hospital or the physician office where diagnosis occurred. The initials of a hospital are not adequate.

For out-of-state residents and facilities, include the city and the state where the medical facility is located.

Example:

- Big Time hospital, 2/1/2004
- Dr. Bill Records office 3/1/2004

TEXT--TREATMENT

RX TEXT—SURGERY

Item Length: 1000
NAACCR Item #2610
NAACCR Name: RX Text—Surgery
XML NAACCR ID: rxTextSurgery

Description:

The surgical procedure(s) performed as part of first course of treatment.

Suggestions for Text

Date of each procedure

Type(s) of surgical procedure(s), including excisional biopsies and surgery to other and distant sites.

Lymph nodes removed (Both name of nodes and number of nodes)

Regional tissues removed

Metastatic sites

Facility where each procedure was performed

Record positive and negative findings. Record positive findings first.

Other treatment information, e.g., planned procedure aborted; unknown if surgery performed

If surgery not performed, then provide reason (i.e., not planned, patient/family refused, patient elected treatment that did not include surgery, comorbid conditions)

Examples:

- 4/2/05: MRM (8 axillary nodes removed, 3 internal mammary nodes removed)
- 5/1/05: TAH, BSO, pelvic lymphadenectomy (6 pelvic nodes removed)

RX TEXT – RADIATION (BEAM)

Item Length: 1000

NAACCR Item #2620

NAACCR Name: RX Text—Radiation (Beam)

XML NAACCR ID: rxTextRadiation

Description

Record regional radiation administered as first course of treatment.

When applicable this should include the site treated and if the radiation is pre, post or intraoperative.

Suggestions for Text

Date radiation treatment began

Where treatment was given, e.g., at this facility, at another facility

Type(s) of beam radiation, e.g., Orthovoltage, Cobalt 60, MV X-rays, Electrons, Mixed modalities

Other treatment information, e.g., patient discontinued after 5 treatments; unknown if radiation given; patient referred to radiation oncologist (provide name)

If radiation recommended but not given, record the reason radiation was not given.

Examples:

- 2/1/05 - 3/1/05; beam rad to prostate, 4500 cGy, 1500 boost
- 2/15/05: pt refused rad implants
- 6/1/05 – 7/1/05: pre op rad, 5000 cGy to large L parotid tumor

RX TEXT – RADIATION OTHER

Item Length: 1000

NAACCR Item #2630

NAACCR Name: RX Text—Radiation Other

XML NAACCR ID: rxTextRadiationOther

Description

Documents information regarding treatment of the tumor being reported with radiation other than beam radiation. This includes brachytherapy and systemic radiation therapy.

Suggestions for Text

Date radiation treatment began

Where treatment was given, e.g., at this facility, at another facility

Type(s) of nonbeam radiation, e.g., High Dose rate brachytherapy, seed plant, Radioisotopes (I-131)

Other treatment information, e.g., unknown if radiation was given

Examples

- 2/1/2007 Interstitial boost to breast using Ir-192

- 3/5/2006 Patient receives 2 Fletcher intracavitary implants for cervical cancer

RX TEXT—CHEMO

Item Length: 1000
NAACCR Item #2640
NAACCR Name: RX Text—Chemo
XML NAACCR ID: rxTextChemo

Description

Chemotherapeutic agents administered to the patient as first course of treatment.

Using standard abbreviations will save time and keying errors.

Ancillary drugs are not coded, but should be included in the text field.

Suggestions for Text

Date chemotherapy began

Where treatment was given, e.g., at this facility, at another facility

Type of chemotherapy, e.g., name of agent(s) or protocol

Other treatment information, e.g., treatment cycle incomplete, unknown if chemotherapy was given, patient referred to oncologist, patient/family refused chemotherapy, chemotherapy recommended but not administered (give reason why).

Examples

- 12/12/04: CAF
- 2/1/05: 5FU & Leucovorin
- 2/11/05: palliative 5FU
- 3/12/05: Epogen (ancillary drugs are not coded but recorded in the text)

RX TEXT-HORMONE

Item Length: 1000
NAACCR Item #2650
NAACCR Name: RX Text—Hormone
XML NAACCR ID: rxTextHormone

Description

Include information on hormone therapy including Hematologic Transplant and Endocrine Procedures.

This text field is still named Hormone.

Hormone therapy (systemic) administered to the patient as first course of treatment.

When Prednisone is given with chemotherapeutic agents, the Prednisone is coded to hormone.

Suggestions for Text

Date treatment was started

Where treatment was given, e.g., at this facility, at another facility

Type of hormone or antihormone, e.g., Tamoxifen

Type of endocrine surgery or radiation, e.g., orchiectomy

Other treatment information, e.g., treatment cycle incomplete; unknown if hormones were given

Examples:

- 12/12/04: Lupron Dr Record's office
- 2/19/04: Pt refused hormone rx
- 6/1/05: CHOPP

Transplant/Endocrine Procedure

- 4/1/04: Bilateral orchiectomy

RX TEXT—BRM

Item Length: 1000
NAACCR Item #2660
NAACCR Name: RX Text—BRM
XML NAACCR ID: rxTextBRM

Description

Immunotherapy (systemic) administered to the patient as first course of treatment.

Suggestions for Text

Date treatment was started

Where treatment was given, e.g., at this facility, at another facility

Type of BRM agent, e.g., Interferon, BCG

Other treatment information, e.g., treatment cycle incomplete; unknown if BRM was given

Examples:

- 5/1/04: Interferon at Main St hosp WV

RX TEXT—OTHER

Item Length: 1000
NAACCR Item #2670
NAACCR Name: RX Text—Other
XML NAACCIR ID: rxTextOther

Description

Other therapy includes treatment of reportable hematopoietic diseases and treatment that does not meet the usual definitions of modifies, controls, removes, or destroys cancer tissue.

Other therapy is not coded as systemic therapy.

Suggestions for Text

Date treatment was started

Where treatment was given, e.g., at this facility, at another facility

Type of other treatment, e.g., blinded clinical trial, hyperthermia

Other treatment information, e.g., treatment cycle incomplete; unknown if other treatment was given

Examples:

- Pt refused chemo, started acupuncture on 3/1/04