



pennsylvania
DEPARTMENT OF HEALTH

**PENNSYLVANIA
CANCER
REGISTRY**

*Prevention, Control and Research
Commonwealth of Pennsylvania
Department of Health*

Tom Wolf, Governor

The Department of Health is an equal opportunity provider of grants, contracts, service and employment.

January 2010, revised January 2011, January 2012, January 2013, January 2014, January 2015, January 2016

PENNSYLVANIA CANCER REGISTRY MANUAL
January 2010, revised January 2011, January 2012, January 2013,
January 2014, January 2015, January 2016

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PREFACE

ABOUT THE PENNSYLVANIA CANCER REGISTRY

The rate of new cancer cases in Pennsylvania is among the highest in the nation. More than 70,000 Pennsylvania residents are diagnosed with cancer each year. Without information on these new cases of cancer, it is difficult to plan prevention, education, screening, early detection, treatment, and rehabilitation programs. The Pennsylvania Cancer Registry (PCR) records the incidence of cancer for the Commonwealth of Pennsylvania and provides data to help physicians, researchers, and other health professionals plan and evaluate cancer programs.

Probably not surprising to learn is the impetus for the PCR was public concern about excess radiation exposure to persons living near a nuclear-powered utility plant. What might be surprising, however, is this plant was not the Three Mile Island nuclear plant; site of the 1979 accident, but rather a plant located at Shippingport, Pennsylvania, and the year was 1973.

Then Governor Milton Shapp appointed a committee of epidemiologists and radiation physicists to examine the question of possible increased risk of cancer for individuals residing near the nuclear-powered plant. That committee reported in 1974 no conclusions could be reached due to the absence of reliable data. As a result, a task force of physicians and laypersons was appointed to examine the Commonwealth's role in helping to combat cancer, specifically emphasizing consideration of a statewide cancer registry. In 1976, the task force published a report with several recommendations for cancer control, including mandating a statewide population-based cancer incidence registry and requiring cancer be made a reportable disease.

In 1977, legislation was introduced which proposed funding for cancer research and control programs in the Commonwealth. The Pennsylvania Cancer Control, Prevention, and Research Act, Act 224 (see *Appendix A, Pennsylvania Cancer Control, Prevention and Research Act*) was signed into law on December 18, 1980. Passage of the bill into law resulted from the work of the Governor's Task Force on Cancer, the Pennsylvania and Philadelphia Divisions of the American Cancer Society, the Pennsylvania Cancer Coordination Committee, and many other concerned Pennsylvania organizations and individuals.

The Act provided enabling legislation for the Pennsylvania Department of Health to establish nine cancer control priorities, which are as follows:

- 1) Cancer Registry
- 2) Cancer screening, detection, and prevention
- 3) Cancer epidemiology and biostatistical studies
- 4) Cancer community outreach programs
- 5) Cancer rehabilitation
- 6) Communication and planning among cancer institutions
- 7) Cancer education and information
- 8) Cancer training
- 9) Cancer clinical research

The Pennsylvania Cancer Control, Prevention, and Research Act identified the cancer registry as the number one priority for cancer control, mandated reporting of cancer cases, and clearly outlined reporting responsibility. In addition, the law established a Cancer Control, Prevention, and Research Advisory Board to advise the Secretary of Health with respect to cancer control, prevention and research in Pennsylvania and to approve and implement the *Pennsylvania Cancer Plan*. Act 224 was, and continues to be, a key element in the success of the PCR and statewide cancer control activities.

The Act mandated all hospitals and laboratories to report cancer cases to the PCR. The implementation of reporting by acute care hospitals took place in four geographic regions over a period from July 1982 to September 1984 with 1985 being the first full year of statewide reporting. To assure complete statewide incidence statistics, efforts to increase reporting by freestanding laboratories and other non hospital sources began in 1995 and continue to be an ongoing challenge as more patients are diagnosed and treated in these settings.

As a population-based cancer incidence registry, the PCR collects demographic, diagnostic, and first course treatment information on all Pennsylvania residents diagnosed with cancer. All information collected and maintained in the PCR database is strictly confidential. Only summary statistical information is published for general distribution and public knowledge. The Department of Health may permit use of in-depth information for research, subject to careful screening, strict supervision, and only to accomplish approved program objectives.

PCR data are used for cancer research and surveillance activities, as well as epidemiologic and other special studies. State-specific incidence and mortality data are published annually in *Pennsylvania Cancer Incidence and Mortality* that includes the most recent five years of data. Data from this and other statistical reports and publications are available on the Department of Health website at <http://www.health.pa.gov/MyRecords/Health-Statistics>

With a strong and stable history, the PCR is recognized as a state-of-the-art cancer reporting system, an important component in the Department of Health's Cancer Program, and a valuable resource for cancer data. The PCR uses current technology and national data collection standards to continually enhance the completeness, accuracy and timeliness of cancer data. As the volume of PCR incidence data increases over time, the utility of these data for program planning, evaluation, and epidemiologic studies increases as well. The PCR depends on the support, cooperation, and accurate reporting by each facility for the ongoing operation of the statewide cancer registry. Working together, sufficient and reliable cancer incidence data will continue to be available to provide answers to our questions, to reduce the burden of cancer in Pennsylvania, and to improve the lives of both present and future patients with cancer.

**PART ONE:
REPORTING REQUIREMENTS**

PCR MANUAL, JANUARY 2010 EDITION

Cancer registration in the year 2010 experienced changes nationally. The January 2010 edition of the *Pennsylvania Cancer Registry (PCR) Manual* was written to convey these changes and their effect on PCR reporting requirements.

This updated manual must be used to abstract and submit reportable cases with a Date of Diagnosis of January 1, 2010 and after to the PCR. Cases diagnosed prior to January 1, 2010 maybe abstracted using this manual after the conversion has been completed.

Note: As revisions are made to the *PCR Manual, January 2010 Edition* the changes will be documented in *Appendix N*. In addition, the revised pages will have the date the revision is effective in the footer.

WHAT IS THE PCR

The Pennsylvania Cancer Registry (PCR) is a population-based cancer incidence registry responsible for the collection of demographic, diagnostic, and treatment information on all patients diagnosed and treated at hospitals, laboratories, and other health care facilities in Pennsylvania.

The PCR is also defined as an incidence only cancer registry. Incidence only registries gather only the information necessary to determine the incidence of cancer by geographic areas, by demographic characteristics and by stage at diagnosis for each type of cancer. Treatment information and other prognostic factors have also been added to the information collected.

The term *central cancer registry* is also used in referring to the PCR. Although a central registry does not have to be population-based, this term is frequently used to mean a statewide cancer registry. A central registry is designed to aggregate data from various sources. The contributing sources required to report to the PCR provide statewide coverage of the population.

WHY REPORT TO THE PCR

The mission of the PCR is to collect and provide complete, accurate, and timely statewide cancer incidence data for determination of cancer rates, trends in the population, use in cancer control and research. To fulfill this mission, the PCR depends on complete ascertainment of cases and use of the data.

1. **The Law and Regulations**- Statewide collection and dissemination of data on cancer by the Pennsylvania Department of Health is mandated in two state laws and Pennsylvania Department of Health disease-reporting regulations. The state laws include the Pennsylvania Cancer Control, Prevention and Research Act, 35 P.S. §5631 *et seq.*, and the Disease Prevention and Control Law of 1955, 35 P.S. §521.1 *et seq.* (*Appendix A*) According to these statutes, each designated hospital and laboratory in the Commonwealth shall report all cases of cancer, which are diagnosed and/or treated at the hospital or laboratory. These cases shall be submitted in the format prescribed by the Pennsylvania Cancer Registry. Regulations mandating reporting cancer cases by hospitals, clinical laboratories, other health care facilities and health care practitioners appear in Section 27.31(b) of 28 Pa. Code Chapter 27 (Communicable and Noncommunicable Diseases). (*Appendix B*)
2. **Cancer Control**- The ultimate value of the registry lies not in collection of the data but in the degree to which the data are used for cancer control. The basis for any successful cancer control program is a comprehensive registry system. Registry data provide answers to questions, the means to target limited cancer control resources, and the mechanism to evaluate cancer control activities.

HEALTH INSURANCE PORTABILITY AND ACCOUNTABILITY ACT (HIPAA)

HIPAA allows for the reporting of identifiable cancer data to public health entities. Because the PCR falls under the definition of a public health entity, HIPAA allows facilities to report data to the PCR in compliance with Pennsylvania state laws and regulations. Written informed consent from each cancer patient reported to public health entities is not required under HIPAA.

The PCR depends on reporting facilities to submit quality data. Through the dedicated efforts of these facilities, the PCR is able to provide accurate information used to establish and enhance cancer control programs, and thus improve the lives of present and future patients with cancer.

PCR REFERENCE DATE

Reference date refers to the start date after which all eligible records must be included in the registry. The PCR reference date is January 1, 1985. This means complete statewide cancer incidence data are available from the PCR for 1985 to the present.

PCR REPORTING SOURCES

Act 224 mandates each designated hospital and laboratory in the Commonwealth shall report all cases of cancer, which are diagnosed and/or treated at the hospital or laboratory. In addition, the PCR has agreements with other states to exchange data.

Hospitals

1. Registry Hospitals - The term *registry hospital* refers to hospitals with a cancer registry functioning as an integral component of the hospital cancer program. They may or may not be accredited by the American College of Surgeons Commission on Cancer. Generally, the cancer registrar or cancer program manager at a registry hospital is delegated the responsibility of reporting to the PCR.
2. Non Registry Hospitals - The term *non registry hospital* refers to hospitals that do not have cancer registries functioning as an integral component of a hospital cancer program. Personnel in the Health Information Management (HIM) Department are delegated the responsibility of reporting to the PCR.

Laboratories

The addition of these cases provides the PCR data on cases never seen in the hospital setting, thereby increasing the overall completeness of PCR data.

1. Hospital Laboratories - Required reporting of cases by hospital laboratories is performed by cancer registry or HIM personnel as described above.
2. Free-Standing Pathology Laboratories - Reporting of cases by designated free-standing laboratories is performed through electronic submission of pathology reports.

Non-Hospital Sources

The Department of Health's regulations concerning the Reporting of Communicable and Non Communicable Disease were revised in January 2002 to expand cancer reporting requirements to include additional non-hospital sources. Two types of non-hospital sources were added to Chapter 27, Section 27.31:

- Other Health Care Facilities – The requirement for reporting by other health care facilities that provide screening, diagnostic or therapeutic service for cancer patients was added to Section 27.31(a). Other health care facilities include facilities such as Radiation Centers and Ambulatory Surgery Centers.
- Health Care Practitioners – The requirement for reporting by health care practitioners who provide screening, diagnostic, or therapeutic services to cancer patients for cancer was added as Section 27.31(b).

Data Exchange

The PCR has written agreements to exchange data with other cancer registries including all contiguous states. This ensures a resident of Pennsylvania who was diagnosed and/or treated out-of-state will be included in the PCR database.

HOSPITAL REPORTING METHODS

Reporting hospitals are required to abstract all cases electronically using commercial or Abstract Plus software. Electronic files of reportable cases are submitted to the PCR via a secure website using Web Plus. Web Plus is an internet-based application developed by the Centers for Disease Control and Prevention (CDC), National Program of Cancer Registries (NPCR). Web Plus has been designed as a highly secure application that can be used to transmit data between reporting facilities and the PCR safely over the public internet. See *Appendix C, Web Plus File Upload Instructions*.

1. Commercial Software - Registry hospitals are required to report cases included in the hospital cancer registry electronically using commercial software.
2. Abstract Plus Software - Non registry hospitals report electronically using Abstract Plus Software provided by the PCR at no cost. Abstract Plus is developed and maintained by the Centers for Disease Control and Prevention (CDC).

REPORTABLE DIAGNOSES

ICD-9-CM and ICD-10-CM Codes

Use the list of ICD-9-CM and ICD-10-CM codes documented in *Appendix D* to identify reportable conditions.

Solid Tumors

1. ICD-O Behavior /2 or /3- All histologic diagnoses with a behavior code of /2 (in situ) or /3 (malignant) in the *International Classification of Diseases for Oncology, Second Edition (ICD-O-2)* or *Third Edition (ICD-O-3)* are reportable.

Exception 1: Cervical intraepithelial neoplasia, grade III, also called CIN III (code 8077/2 with primary site C53.X in ICD-O-3) is not reportable.

Exception 2: Prostatic intraepithelial neoplasia, grade III, also called PIN III (code 8148/2 in ICD-O-3) is not reportable.

Exception 3: Pilocytic/Juvenile astrocytoma (code 9421/3 in ICD-O-2 and 9421/1 in ICD-O-3) is reportable and must be coded with a behavior of /3 (malignant).

Note 1: If a pathologist verifies a /0 (benign) or /1 (uncertain whether benign or malignant) behavior code term in ICD-O as /2 (in situ) or /3 (malignant), these records are reportable.

2. ICD-O Behavior /0 or /1 – All histologic diagnoses with a behavior code of /0 (benign) or /1 (borderline or uncertain) when **primary to the intracranial and central nervous system (CNS) sites listed below** are reportable when diagnosed on or after **January 1, 2004**.

Reportable Intracranial and Central Nervous System (CNS) Primary Sites Effective January 1, 2004	
Meninges (C70.0 - C70.9)	Other CNS (C72.8, C72.9)
Brain (C71.0 - C71.9)	Pituitary gland (C75.1)
Spinal Cord (C72.0)	Craniopharyngeal duct (C75.2)
Cauda equina (C72.1)	Pineal gland (C75.3)
Cranial nerves (C72.2 - C72.5)	

3. Basal and Squamous Cell Carcinomas- Basal and squamous cell carcinomas are reportable except when primary to the skin, C44.0-C44.9 (see *Part One, Exclusions*). Carcinomas originating in **mucoepidermoid sites** are reportable. These sites include: lip (C00.0-C00.9), anus (C21.0), vulva (C51.0-C51.9), vagina (C52.9), penis (C60.0-C60.9), and scrotum (C63.2). Basal and squamous cell carcinomas originating in the nasal cavity (C30.0) and middle ear (C30.1) are also reportable.

4. Class IV and Class V Cytologies- Cytology results of Class IV or Class V are reportable to the PCR.

Exception: If the terminology on the cytology report further defines the Class IV and Class V as *suspicious* then the record is not reportable. Report this record only if a positive biopsy or a physician's clinical impression of cancer supports the cytology findings. Suspicious cytology only is not diagnostic of cancer.

Note: See *Part Three, Data Item Instructions, Diagnostic Confirmation* for clarification of histology and cytology using cell block and smear preparation of specimens.

5. Low Malignant Potential/Borderline Malignancy of Ovary or Peritoneum- Cystadenomas or tumors primary to the ovary or peritoneum qualified by the phrases *borderline malignancy* or *low malignant potential* are reportable only if **diagnosed prior to January 1, 2001**.

6. Intraepithelial Neoplasia- Patients with the following diagnoses of intraepithelial neoplasia are reportable. See also *Part One, Exclusions, Intraepithelial Neoplasia*.

- Vaginal intraepithelial neoplasia 3 (VAIN III)
- Vulvar intraepithelial neoplasia 3 (VIN III)
- Anal intraepithelial neoplasia 3 (AIN III)

7. Carcinoid of the Appendix – Effective with cases diagnosed on or after January 1 2015 and after all *Carcinoids of the appendix* **are reportable** and should be coded to 8240/3.

8. Severe/High Grade Dysplasia of the Colon and/or Esophagus- If your facility considers the terminology of severe dysplasia or high grade dysplasia of the colon and/or esophagus as synonymous with carcinoma in-situ; use the following guidelines for reporting these cases to the PCR:

- a. Obtain a statement from your pathologists outlining the terminology policy of their department. The statements should be separate, one for colon and one for esophagus, because different physicians may be involved in reviewing the statement for each site.
- b. Submit the statement(s) to the appropriate medical staff committee for approval. Registry hospitals will normally submit the statement(s) to the Cancer Committee.
- c. Document a policy that states the sites diagnosed with severe dysplasia and/or high grade dysplasia that will be abstracted as carcinoma in-situ based on the pathologists approved statements.
- d. Add the policy to your Operations Manual attaching the approved statement(s) from your pathologists.
- e. Forward a copy of the policy and statement(s) to the PCR to keep on permanent file.
- f. Abstract all colon and/or esophagus cases diagnosed with severe dysplasia and/or high grade dysplasia as carcinoma in-situ according to the statement(s) and policy. In the text for each abstract, document the final pathologic diagnosis along with the statement "in-situ per pathologist".

Hematopoietic and Lymphoid Neoplasms

1. Reportability Instructions- For all hematopoietic or lymphoid neoplasms, refer to the *Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual, Case Reportability Instructions*. <http://seer.cancer.gov/seertools/hemelymph/>

Note: The *Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* **must be used** because not all reportable hematopoietic or lymphoid terms are indexed in ICD-O-3.

Example: Essential Thrombocytosis is coded to ICD-9-CM code 238.71. It is not indexed in ICD-O-3. However when checked in the *Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual, Case Reportability Instructions* it is considered reportable; therefore the case is reportable and must be abstracted and submitted to the PCR

2. Behavior Code Change from /1 to /3 – The behavior code for hematopoietic neoplasms listed below changed from /1 (borderline or uncertain) to /3 (malignant) and are reportable when diagnosed on or after **January 1, 2010**.

Histologic Terms and Codes with Changes in Case Reportability Effective January 1, 2010	
Name	ICD-O-3 Code
Chronic lymphoproliferative disorder of NK-cells	9831/3
T-cell large granular lymphocytic leukemia	9831/3
Langerhans cell histiocytosis, NOS (9751/1)	9751/3
Langerhans cell histiocytosis, unifocal (9752/1)	9751/3
Langerhans cell histiocytosis, multifocal (9753/1)	9751/3
Myelodysplastic/Myeloproliferative neoplasm, unclassifiable	9975/3
Myeloproliferative disease, NOS	9975/3
Myeloproliferative neoplasm, unclassifiable	9975/3

AMBIGUOUS TERMINOLOGY FOR SOLID TUMORS

A patient has a reportable condition if a *recognized medical practitioner* says so. In most cases the patient’s record clearly presents the diagnosis by use of specific terms, which are synonymous with the diagnosis. However, the physician may not always be certain or the recorded language definitive. PCR rules concerning the usage of ambiguous terminology are as follows:

1. Terms That Constitute a Diagnosis- Interpret the following terms as a reportable diagnosis:

<i>apparent(ly)</i>	<i>consistent with</i>	<i>neoplasm**</i>	<i>suspicious (for)</i>
<i>appears</i>	<i>favor(s)</i>	<i>presumed</i>	<i>tumor**</i>
<i>comparable with</i>	<i>malignant appearing</i>	<i>probable</i>	<i>typical (of)</i>
<i>compatible with</i>	<i>most likely</i>	<i>suspect(ed)</i>	

**Beginning with 2004 diagnoses and only for meninges, brain, central nervous system, pituitary gland, craniopharyngeal duct, and pineal gland (C70.0-C72.9, C75.1-C75.3).

Note: Do not substitute synonyms such as “supposed” for presumed or “equal” for comparable. Do not substitute “likely” for “most likely.”

Exception: Do not report a case based only on suspicious cytology. The case is reported only if proven by positive cytology or other diagnostic methods including a physician’s clinical diagnosis.

2. Terms That Do Not Constitute a Diagnosis- Do not interpret the following terms as a diagnosis. Do not report patients who have a final diagnosis consisting only of these terms without additional information to support reportability:

<i>cannot be ruled out</i>	<i>potentially malignant</i>	<i>suggests</i>
<i>equivocal</i>	<i>questionable</i>	<i>worrisome</i>
<i>possible</i>	<i>rule(d) out</i>	

3. How To Use Ambiguous Terminology For Case Ascertainment

- a. In Situ and Invasive (Behavior codes /2 and /3)

1. **If any of the reportable ambiguous terms precede** a word that is **synonymous** with an in situ or invasive tumor (e.g., cancer, carcinoma, malignant neoplasm, etc.), the case is reportable.

Example 1: The pathology report says: “Prostate biopsy with markedly abnormal cells that are typical of adenocarcinoma.” Report the case.

Example 2: The final diagnosis on the outpatient report reads: “Rule out leukemia.” Do not report the case.

2. Discrepancies: If one section of the medical record(s) uses a reportable term such as “apparently” and another section of the medical record(s) uses a non-reportable term such as “cannot be ruled out”, accept the reportable term and report the case.

Exception: Do not report a case based only on suspicious cytology. The case is reported only if proven by positive cytology or other diagnostic methods including a physician’s clinical diagnosis.

3. Use these terms when screening diagnoses on pathology reports, operative reports, scans, mammograms, and other diagnostic testing other than tumor markers.

Note: If the ambiguous diagnosis is **proven to be not reportable** by biopsy, cytology, or physician’s statement, **do not report** the case.

Example: Mammogram shows calcifications suspicious for intraductal carcinoma. The biopsy of the area surrounding the calcifications is negative for malignancy. Do not report the case.

- b. Benign and borderline primary intracranial and CNS tumors

1. Use the “Ambiguous Terms that are Reportable” list to identify benign and borderline primary intracranial and CNS tumors that are reportable.

2. If any of the reportable **ambiguous terms precede** either the word “**tumor**” or the word “**neoplasm**,” the case is reportable. Report the case.

Example: The mass on the CT scan is consistent with pituitary tumor. Report the case.

3. Discrepancies: If one section of the medical record(s) uses a reportable term such as "apparently" and another section of the medical record(s) uses a non-reportable term such as “cannot be ruled out”, accept the reportable term and accession the case.

Exception: Do not report a case based only on suspicious cytology. The case is reported only if proven by positive cytology or other diagnostic methods including a physician’s clinical diagnosis.

4. Use these terms when **screening** diagnoses on pathology reports, scans, ultrasounds, and other diagnostic testing other than tumor markers.

Note: If the **ambiguous** diagnosis is proven to be **not reportable** by biopsy, cytology, or physician’s statement, **do not report** the case.

- c. Confirmation of an Ambiguous Diagnosis- Subsequent admissions for patients whose initial diagnosis contained ambiguous terminology must be reviewed. It is established practice to accept the information at the time of the latest admission, or the most complete or detailed information.

AMBIGUOUS TERMINOLOGY FOR HEMATOPOIETIC AND LYMPHOID NEOPLASMS

Reportability Instructions- For all hematopoietic or lymphoid neoplasms, refer to the *Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual, Case Reportability Instructions*.
<http://seer.cancer.gov/seertools/hemelymph/>

DATE OF DIAGNOSIS REPORTABILITY

All reportable cases included under *Part One, Reportable Diagnoses* diagnosed or treated at the facility are required to be reported to the PCR only if the date of diagnosis is **known to be on January 1, 1995 or after**. Known diagnosis date means the **year or the month and year, or the month, day and year** are known or can be estimated. Patients with an unknown date of diagnosis are **not** reportable.

Exception 1: Conditions only reportable if diagnosed on January 1, 2001 and after are **not** reportable if the date of diagnosis is between January 1, 1995 and December 31, 2000 or unknown.

Exception 2: Benign and borderline tumors of the intracranial and central nervous system are only reportable if diagnosed on or after January 1, 2004 and are **not** reportable if the date of diagnosis is prior to January 1, 2004 or unknown.

Example 1: If a patient is admitted on January 3, 2010 and is diagnosed with lung cancer on January 7, 2010, the case is reportable.

- Example 2:* If a patient is admitted on January 3, 2010 and receives palliative care for bone metastasis from a breast primary diagnosed in 1999, the case is reportable.
- Example 3:* If a patient is admitted on January 3, 2010 and receives a blood transfusion for polycythemia vera, originally diagnosed in November 1999, the case is not reportable per *Exception 1* above.
- Example 5:* If a patient is admitted on January 3, 2010 and receives treatment for hemangioblastoma, originally diagnosed in November 2003, the case is not reportable per *Exception 2* above.

REPORTABLE SITUATIONS

A case is reportable to the PCR if it is a condition included under *Part One, Reportable Diagnoses* and meets the following criteria:

1. Patients diagnosed or treated in your inpatient or outpatient departments, emergency room, ambulatory care center, or other units included under your hospital license.
 - a. Patients Diagnosed At Your Hospital - The reportable diagnosis has been made at your hospital. This diagnosis can be made on the basis of histology (including autopsy), hematology, cytology, endoscopy or other direct visualization, diagnostic radiology or clinical findings.

Clinical Diagnosis Only - A “clinical diagnosis only” is a diagnosis based solely on clinical judgment; diagnostic procedures were not performed or did not confirm the diagnosis. Patients diagnosed clinically are reportable to the PCR.
 - b. Patients Treated at Your Hospital - The PCR requires patients receiving treatment, cancer-directed or non cancer-directed, to be reported provided they have not been previously reported by your hospital.

The PCR recognizes the following definitions of treatment:

1. Cancer-Directed Treatment - Cancer-directed treatment is tumor directed, and its purpose is to modify, control, remove or destroy primary or metastatic cancer tissue. Physicians administer the therapy (ies) to remove or minimize the size of tumor or to delay the spread of disease.
2. Non Cancer-Directed Treatment - Non cancer-directed treatments prolong the patient’s life, alleviate pain, make the patient comfortable, or prepare the patient for cancer-directed therapy. They are not meant to destroy or control the tumor or delay the spread of disease. Non cancer-directed procedures include diagnostic tests and supportive care (treatments designed to relieve symptoms and minimize the effects of the disease).

Examples of non cancer-directed treatment include - Incisional biopsy, exploratory procedures with or without biopsies, port-a-catheter insertion, IV access for chemotherapy, pain medications, oxygen, antibiotics for an associated infection, transfusions, intravenous fluids to maintain fluid or nutritional balance, and laser therapy for relieving symptoms.

Note for registry hospitals: Because some patients receiving non cancer-directed treatment are nonanalytic, registry hospitals following ACOS Commission on Cancer guidelines may choose not to include these records in their hospital-based registries. These records, however, must be abstracted and reported to the PCR.

2. Patients Diagnosed at Autopsy- Final autopsy reports containing reportable diagnoses or incidental findings of reportable conditions must be reported to the PCR. Although autopsies performed as coroner's cases are not required to be reported, hospitals are encouraged to submit them to the PCR.
3. Patients Diagnosed Elsewhere- Patients diagnosed elsewhere and newly admitted to your hospital for further diagnostic workup or treatment, cancer-directed or non cancer-directed are to be reported.

Although this may result in multiple records on one patient, it enables the PCR to assure complete statewide casefinding and to have the most comprehensive information on each patient. Because the PCR is a population-based registry, every attempt must be made to receive all cases diagnosed within Pennsylvania to provide accurate statistical reports.

4. Brain and CNS Tumors Identified by Diagnostic Imaging Only- Patients with a brain or CNS tumor identified by diagnostic imaging (CT scans, MRI scans, or ultrasounds/sonography) are reportable even when no other information is available (from biopsy or resection, for example). The behavior for the tumor is coded as /1 (uncertain whether benign or malignant).
5. Residual Tumor-The PCR requires all records in which the pathology report states "no residual tumor" to be reported. The re-excision is considered cancer-directed treatment.

Example: Outside the hospital setting, a patient has a biopsy and is diagnosed with a malignant melanoma. The patient is seen at your hospital for a wide excision. The tissue report from the excision states no residual tumor. This record is reportable to the PCR. Even though the cancer was diagnosed elsewhere, the patient's hospital visit was for cancer related treatment.

6. Private Outpatient Specimens (POP) (Path Only)- Private outpatient specimens (POP) are specimens submitted from a physician's office to be read by the hospital pathologist as part of the Pathology Department's regular course of business. The patient is not registered as an inpatient or outpatient at the hospital. POPs are reportable to the PCR.

Example: A physician performs a biopsy in the office and sends the specimen to your Pathology Department where a reportable diagnosis is made.

- a. POP reports should be held for two to three months because many of these patients may return for treatment and more information can be obtained from these records.
- b. If the patient does not return as an inpatient or hospital outpatient, abstract the record using all available information. Every effort must be made to obtain accurate information. This information can be found through hospital billing systems, clinical history, or if needed by contacting physician offices.
- c. Data items should be completed as *unknown* only after further investigation does not provide more specific information.
- d. The specimen date should not be considered the date of diagnosis without further confirmation

from the clinical history or physician.

7. Ownership of the Medical Record- When the distinction between a hospital department and a freestanding facility cannot readily be made, such as a radiation therapy group practice versus a hospital unit; the ownership of the medical record is used to determine whether or not a record must be reported. If the medical record is the property of the institution, the record **must** be reported. If the hospital is part of a corporation, ownership of the record refers to the facility, not the corporation.

EXCLUSIONS

Non-Reportable Diagnosis

The following diagnoses are **not** reportable to the PCR.

1. Skin Cancers

- a. The following site/histology combinations for skin cancers are not reportable:

8000-8005	Neoplasms malignant, NOS of the skin (C44.0-C44.9)
8010-8046	Epithelial carcinomas of the skin (C44.0-C44.9)
8050-8084	Papillary and squamous cell carcinomas of the skin (C44.0-C44.9)
8090-8110	Basal cell carcinomas of the skin (C44.0-C44.9)

- b. ICD-O codes C44.0-C44.9 include skin of the lip, eyelid, external ear, face, nose, scalp, neck, trunk, perineum, (peri) anus, umbilicus, upper and lower limbs, shoulders, hips, and skin around ostomy sites.

Note: The above lesions are reportable when the primary tumor originates in a mucoepidermoid site (See *Part One, Reportable Diagnoses*).

- c. Skin of nose - Basal and squamous cell carcinomas originating in the external nose (C44.3) are not reportable; however, those primary to the nasal cavity (C30.0) such as nostril, nasal septum, and nares are reportable.
- d. Metastasis from non-reportable sites - If the primary site is not reportable but the cancer has metastasized to other sites, the record is still not reportable.

2. Carcinoma-In-Situ of the Cervix (CIS) - The diagnosis carcinoma in situ of the cervix (CIS) is not reportable. Terms indicating in situ include: *noninvasive*, *preinvasive*, *intraepithelial*, and *FIGO Stage 0*. A diagnosis of carcinoma in situ with endocervical gland involvement is still considered in situ and is not reportable.

Note: Diagnoses of **invasive** carcinoma of the cervix **are** reportable. A diagnosis of carcinoma in situ of the cervix with microinvasion is considered invasive and is therefore reportable.

3. Intraepithelial Neoplasia- Patients with the following diagnoses of intraepithelial neoplasia are not

reportable:

- Cervical intraepithelial neoplasia (CIN)
- Prostatic intraepithelial neoplasia (PIN)

See also *Part One, Reportable Diagnoses, Intraepithelial Neoplasia*.

4. Other Precancerous Conditions and Benign Tumors- Patients with precancerous conditions or benign tumors are not reportable. An example of such a diagnosis includes atypical adenoma. Registry hospitals may elect to collect these cases; however, they are not reportable to the PCR.

Exception 1: Ovary and Peritoneum- Cystadenomas or tumors primary to the ovary or peritoneum qualified by the phrases *borderline malignancy* or *low malignant potential* are reportable if **diagnosed prior to January 1, 2001**.

Exception 2: Brain and Central Nervous System- All primary intracranial and central nervous system (CNS) tumors are reportable beginning with cases diagnosed **on or after January 1, 2004**. This includes benign and borderline tumors for the following sites:

Reportable Intracranial and Central Nervous System (CNS) Primary Sites Effective January 1, 2004	
Meninges (C70.0 - C70.9)	Other CNS (C72.8, C72.9)
Brain (C71.0 - C71.9)	Pituitary gland (C75.1)
Spinal Cord (C72.0)	Craniopharyngeal duct (C75.2)
Cauda equina (C72.1)	Pineal gland (C75.3)
Cranial nerves (C72.2 - C72.5)	

Non-Reportable Situations

A case is **not** reportable to the PCR if it meets any of the following criteria:

1. Consult Only Records- Patients seen in consultation to provide a second opinion to confirm an established diagnosis or treatment plan are not reportable. Also, if the reporting institution provides services not available at the diagnosing or treatment facility, such as Computerized Tomography (CT) scans or Magnetic Resonance Imaging (MRI) scans, the case is not reportable.

Exception: Consults are reportable if they are done to establish a diagnosis or to develop and document a treatment plan. See *Part Three, Guidelines for Recording First Course of Treatment* for a definition of a treatment plan.

Example: The primary care physician (PCP) sends the patient to a consulting physician for his opinion and the consulting physician subsequently develops a treatment plan, the case is reportable

2. Slide Reviews- Records in which slides are sent to your hospital’s pathologist for a second opinion are encouraged to be reported, but are not required. Since the slide was already read by another

pathologist, the facility requesting the slide review is required to report the final diagnosis as determined after the slide review.

3. History of- Patients with a history of a reportable condition who are clinically free of disease are not reportable. If, however, the patient has actually received treatment during this admission the record must be reported. For example: if a patient is admitted for an unrelated condition, has a history of breast cancer and the hospital administers Tamoxifen during their admission, the case is reportable.
4. Positive Imaging Study- Facilities are not expected to report non-brain/CNS cases on the basis of a positive imaging study only. However, if the patient meets reporting requirements at a later time, the case must be reported.

Exception: A brain or a CNS 'neoplasm' identified by diagnostic imaging is reportable even when no other information is available.

5. Suspicious cytology only is not diagnostic of cancer. Report this record only if a positive biopsy or a physician's clinical impression of cancer supports the cytology findings.
6. Transient Care- Patients receiving transient care at the reporting institution to prevent interruption of the first course of treatment are not reportable. This only applies to patients vacationing or visiting in the area, or equipment failure at the primary treating institution which requires the patient to temporarily receive treatment elsewhere.

Exception: Cancer patients evacuated to other states due to natural disasters may receive diagnostic/treatment services in facilities in that state. If this occurs at your facility, consider these cases reportable to the Pennsylvania Cancer Registry (PCR). They should not be excluded as transient care or consult only cases.

When abstracting these cases, please record the **patient's usual residence when the tumor was diagnosed** in the Address at Diagnosis fields. Do not enter the patient's current address if the patient was diagnosed prior to relocating permanently or temporarily to Pennsylvania or other nearby state.

7. Readmitted Patients- If a patient is readmitted and new or additional metastatic sites are diagnosed or documented, the record is not reportable provided it has already been reported for the original primary site. Records of readmitted patients must be reviewed to determine if a new primary site has been diagnosed. Each new primary must be reported separately.
8. Metastatic Sites- Do not report the metastatic or secondary sites of a malignant neoplasm; however, check to make sure the primary site was previously reported. A diagnosis of metastatic cancer with an unknown primary site not previously reported should be submitted with the primary site documented or coded as **unknown**.
9. Special Units- Patients admitted to a skilled nursing unit or other separately licensed units are encouraged to be reported but are not required. These patients are either discharged from an acute care hospital unit and readmitted to a separately licensed unit or are admitted directly to the separately licensed unit.

10. Diagnosed Prior to 1995- Patients with a date of diagnosis prior to January 1, 1995 are not reportable.
11. Unknown Date of Diagnosis- Cases with an unknown month, day and year of diagnosis are not reportable.

Note: Estimating Diagnosis Date - If information is available to estimate at a minimum the year of diagnosis, the case is reportable. Instructions provided in the *PCR Manual Part Three: Data Item Instructions, General Information-Dates* should be used to approximate the diagnosis date whenever possible.

MULTIPLE PRIMARY DETERMINATION

Solid Malignant Tumors and Benign and Borderline Central Nervous System Tumors

1. Cases diagnosed on or after January 1, 2007- Effective with cases diagnosed on or after January 1, 2007 the PCR requires the use of *The Multiple Primary and Histology Coding Rules (MP/H Rules)* for multiple primary determination and histology coding of solid malignant tumors and benign and borderline Central Nervous System tumors. The *MP/H Rules Manual* can be downloaded from the following site:
2. Cases diagnosed prior to January 1, 2007- Pages 7-18 of the *SEER Program Coding and Staging Manual 2004* should be used for solid malignant tumors and benign and borderline Central Nervous System tumors diagnosed prior to January 1, 2007. The *SEER Program Coding and Staging Manual 2004* can be downloaded from the following site:

<http://seer.cancer.gov/tools/mphrules/download.html>

http://seer.cancer.gov/archive/manuals/2004Revision1/SPM_2004_maindoc.r1.pdf

Hematopoietic and Lymphoid neoplasms

1. Cases diagnosed on or after January 1, 2010- Effective with cases diagnosed on or after January 1, 2010 the PCR requires the use of *The Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* and *The Hematopoietic Database* for multiple primary determination, primary site and histology coding of hematopoietic and lymphoid neoplasms. The manual and database can be downloaded from the following site:
2. Cases diagnosed prior to January 1, 2010- The SEER table *Definitions of Single and Subsequent Primaries for Hematologic Malignancies* should be used to determine multiple primaries for hematopoietic and lymphoid neoplasms diagnosed prior to January 1, 2010. The SEER table *Definitions of Single and Subsequent Primaries for Hematologic Malignancies* can be downloaded from the following site: http://seer.cancer.gov/archive/icd-o-3/hematopoietic_primaries.d03152001.pdf

CONFLICTING STANDARDS

When standards of regulatory agencies differ, hospitals must implement procedures to comply with PCR standards.

WHEN IN DOUBT

When in doubt about submitting records to the PCR, ask the following question:

1. Did your facility diagnose and/or treat the patient for a condition included under *Part One, Reportable Diagnoses*?
2. Is the Date of Diagnosis known and is it on or after January 1, 1995?

If the answer to these questions is yes and the record was not previously submitted by your hospital, report the record. If you are in doubt about a particular record, call your PCR Field Representative at 1-800-272-1850.

PCR REQUIRED DATA ITEMS

The PCR requires specific data items to be completed for each reportable case. These data items include demographic, cancer identification, treatment, hospital-specific and text information. A listing of the PCR Required Data Set is included in *Appendix F*. Instructions on completing each data item are provided in *Part Three, Data Item Instructions*.

All data items required for participation in the National Program of Cancer Registries (NPCR) are included in the PCR data set. PCR-required codes and definitions comply with national standards established by the North American Association of Central Cancer Registries (NAACCR) and American College of Surgeons Commission on Cancer (ACOS COC).

CHANGING INFORMATION

A change includes updating or correcting previously submitted information.

Importance of Change/Deletion Procedure

The change procedure ensures the most accurate information is available to users of PCR data by enabling reporting facilities to provide updated or corrected information after a record has been accessioned by the PCR.

Changes are submitted electronically as an M record. An M record is any record Modified since previous submission to central registry (identical in format to the “A” record type).

Example: At the time a record was reported to the PCR, the primary site was unknown. On a subsequent admission, the primary site was documented as upper lobe of left lung. A modification record or change sheet must be submitted to update the primary site, laterality, and stage (as was known during first course of treatment). The PCR will update this information on the patient’s record on the PCR data file.

What to Change

1. Change any required data item when incorrect or unknown information was initially reported or when more specific/correct information is later available.
2. Change Collaborative Stage data items and SEER Summary Stage only if additional information is available through 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.
3. Submit a change for name when incorrectly spelled on record and when name is changed due to marital status or other reason.
4. **Do not** submit changes to update address changes or admission/discharge dates when the patient is readmitted.

How to Change Information Electronically

1. Make the change to the case in your database. Follow the procedures provided by your vendor to generate an M record. Abstract Plus users refer to the *Abstract Plus Users Guide* for instructions.
2. When creating your normal file for submission (A Records) each month, a separate file of M records should also be created. The A and M records cannot be combined into one file, a separate file of each is needed.
3. Upload both your A record file and your M record file in Web Plus each month. For your M record file, write “### M Records” in the comment box. Indicating the number of M records being submitted.

How to Notify PCR of a Deletion

1. If it is later determined a previously reported case is NOT reportable, contact your PCR Field Representative and provide the following information:
 - a. Patient Name
 - b. Patient Date of Birth
 - c. Patient Social Security Number
 - d. Reason for deleting the case

Example: At the time a record was reported to the PCR, the patient's initial diagnosis was *probable carcinoma*. After further review, it was determined the patient does not have cancer. Such cases must be deleted. Contact your PCR Field Representative with the above information. The case will then be deleted from the PCR database.

2. Delete the case from your database.

PCR TRANSMITTAL FORM

A *PCR Transmittal Form* is only used for the following reason:

- **No Records to Report**- If a facility has no cancer records to report, an electronic *PCR Transmittal Form* with zero (0) entered for the number of new records must be completed and submitted via Web Plus by the 15th of the month. In addition, the reason for not submitting any records must be documented on the transmittal form in the space provided.

An electronic version of the *PCR Transmittal Form* to use when there are no records to report can be downloaded from the following link:

<http://www.health.pa.gov/MyRecords/Registries/Cancer/Pages/Hospital%20Reporting.aspx>

Instructions for Completing a *PCR Transmittal Form*

1. **Facility Name**- Enter full facility name
2. **PCR Identification Number**- Enter four-digit PCR identification number
3. **Date Submitted** - Enter date shipment was sent
4. **Number of New Records*** - Enter “0” if there are no records to report for the month and document the reason on the lines provided.

HOW TO REPORT

Records containing all required data items must be uploaded to the PCR using Web Plus. Detailed instructions for completing the required data items can be found in the *Part Three, Data Item Instructions*. See *Appendix C* for instructions to upload files via Web Plus.

WHEN TO REPORT

Submission Dates

Reporting facilities must upload abstract files on the 15th of every month. If the 15th falls on a weekend or holiday, files must be uploaded on the last working day **before** the 15th. Abstract files may be uploaded more than once per month but **no more frequently than once a week**. Special submissions such as reconciliations, death clearance cases, etc, are not considered regular reporting and therefore may be transmitted the same week as a routine transmission.

Timeliness of Reporting

1. **180 Days**- The PCR requires 90% of abstracts submitted by reporting facilities to be received by the PCR within 180 days from *Date of Inpatient Disch* if an inpatient or *Date of 1st Contact* if an outpatient. The months of May and June should be used to perform quality assurance procedures to ensure all cases have been identified and reported. These cases may fall into the 10% over 180 days. This is expected and acceptable. The timeliness requirement was established at 90% to provide a cushion of 10% to encourage late reporting of missed cases to assure reporting completeness.
2. **Year End Deadline**- The first working day in July is the deadline for submitting all reportable cases from the previous year.

FACILITY CONTACT PEOPLE FOR PCR

One person at each reporting facility is designated as the PCR contact person. This person will receive all PCR correspondence regarding PCR requirements and quality control results. For example: changes in reporting requirements and timeliness reports. Each facility must designate the Managing PCR contact person such as the supervisor or manager that oversees the staff responsible for reporting to the PCR.

The reporting facility can inform the PCR of additional staff they would like included in all email correspondence. To maintain proper communication, inform the PCR of any changes in the contact people at your facility by calling 1-800-272-1850 or emailing your PCR Field Representative.

TRAININGS

The PCR conducts trainings throughout the year to provide specific information on PCR reporting requirements and data collection. Trainings are free of charge.

Announcements listing dates and locations of trainings are sent to PCR contacts periodically. If interested in attending a training, refer to the announcement or call the PCR. See *Part Four, Quality Control: PCR, Trainings*.

CONTACTING THE PCR

If you have any questions regarding the PCR, contact us at: 1-800-272-1850.

All faxes for PCR staff should be sent to 1-866-531-8238.

A list of current PCR Field Representatives can be found on the *Contact Information* page of the PCR website.

<http://www.health.pa.gov/MyRecords/Registries/Cancer/Pages/Contact%20Information.aspx>

**PART TWO:
CASEFINDING**

CASEFINDING

Casefinding Procedures

Casefinding is a system for identifying patients with a reportable diagnosis. Because cancer incidence can be most accurately reflected only when every reportable diagnosis is identified and submitted to the central registry, effective casefinding procedures are essential.

Although casefinding procedures will vary among reporting facilities, the key to effective casefinding is the identification of reportable conditions in all areas where patients are diagnosed or treated in a routine and systematic manner. The following concepts should be considered when developing procedures to insure complete identification of cases reportable to the Pennsylvania Cancer Registry (PCR).

Reportable Conditions

The first step in establishing effective casefinding procedures is to know what conditions are reportable. These conditions are defined in the following references:

- ICD-9-CM AND ICD-10-CM Codes – Appendix D, List of Reportable ICD-9-CM and ICD-10-CM Codes provides a list of ICD-9-CM and ICD-10-CM codes used to identify reportable diagnoses.

Casefinding Sources

The second step in establishing effective casefinding procedures is to identify all areas in the facility where these reportable conditions are either diagnosed or treated and the sources for casefinding in each area. The Health Information Management (HIM) Department and Pathology Department must be included as casefinding sources by all facilities; the remaining sources listed below should be included as applicable. Copies of reports forwarded for review to the person responsible for reporting to the PCR serve as a pending or tickler file to cross-reference with medical records flagged in the HIM Department.

The term “records” as used in the descriptions below refers to all patient records, i.e., inpatient, outpatient, Emergency Room, ambulatory care, short stay procedures, radiation therapy, chemotherapy. For each source, review all of the following reports and records.

1. Health Information Management Department (HIM) - Disease Index - Records assigned an ICD-9-CM or ICD-10-CM code included in *Appendix D* should be reviewed to identify reportable cases. In addition to casefinding, the disease index should also be used as a quality control measure to make sure all reportable diagnoses have been submitted. See also *Part Four, Quality Control: Reporting Facilities*.
2. Pathology Department/Laboratory Medicine- Casefinding from Pathology Department/Laboratory Medicine must include identification of reportable diagnoses made on inpatient, outpatient, and private outpatient (POP) specimens.
 - a. Histology - Surgical pathology reports should be reviewed for a reportable diagnosis. If your Pathology Department screens the reports and forwards copies of those reports to the person responsible for PCR reporting, they must be provided with a copy of *Appendix D*. Surgical pathology reports showing “no residual malignancy (or tumor)” and reports resulting from orchiectomy or oophorectomy performed for prostate or breast malignancies should be included in what is copied and forwarded to the person responsible for PCR reporting.

- b. Cytology - All cytology reports should be reviewed for a malignant diagnosis and, when identified, a copy forwarded to the person responsible for PCR reporting. An alternative would be to review a log of positive or abnormal cytologies.
 - c. Hematology - Peripheral blood reports should be reviewed for a diagnosis of malignancy and, when identified, a copy forwarded to the person responsible for PCR reporting.
 - d. Bone Marrow - All bone marrow reports should be reviewed for a diagnosis of malignancy and, when identified, a copy forwarded to the person responsible for PCR reporting.
 - e. Autopsy - All final autopsy reports should be reviewed for reportable diagnoses including incidental findings and, when identified, a copy forwarded to the person responsible for PCR reporting. Reportable diagnoses on autopsy reports from coroner's cases should also be identified. See *Part One, Patients Diagnosed at Autopsy*.
3. Outpatient Departments
 - a. Short Procedure/Same Day Surgery/Ambulatory Care Unit - A system must be implemented to routinely review all outpatient records maintained within or separate from the HIM Department for diagnoses. If reporting criteria are met, cases must be submitted to the PCR.
 - b. Emergency Room (ER) - Pathology and cytology reports from procedures performed in the ER should be screened and reported if a reportable diagnosis is made.
 4. Oncology Services
 - a. Radiation Therapy - Radiation therapy records, appointment logs, or patient rosters must be reviewed. If reporting criteria are met, cases must be submitted to the PCR. Patients diagnosed elsewhere but treated at your facility must be reported.
 - b. Medical Oncology/Chemotherapy - Chemotherapy records, appointment logs, or patient rosters must be reviewed. If reporting criteria are met, cases must be submitted to the PCR. Patients diagnosed elsewhere but treated at your facility must be reported.
 5. Other Areas- Records from other areas of the hospital where reportable conditions are either diagnosed or treated must be reviewed and submitted if a reportable diagnosis is made.
 6. Records Maintained Separately from the HIM Department- Reporting cases when records are maintained separately from the HIM Department is based on ownership of the medical record. The determination must be made as to who owns the medical record - either the hospital or the medical practitioner who uses space/equipment within the hospital to perform procedures and/or treat private patients. If the hospital owns the record, the case must be reported. If the hospital does not own the record, the PCR requests but does not require the case to be reported.

Completeness of Casefinding

After all reportable diagnoses have been identified through routine casefinding procedures; the final step to effective casefinding is quality control. Procedures should be in place to verify all cases were identified and reported to the PCR. *PCR Manual Part Four, Quality Control* describes various quality control strategies to assure complete casefinding and reporting.

Most Effective Casefinding Procedure

The most effective approach to identifying all reportable diagnoses for reporting to the PCR should include the following:

1. Review the ICD-9-CM or ICD-10-CM disease index daily, weekly or monthly of all inpatient and outpatient medical records with an ICD-9-CM or ICD-10-CM diagnosis code listed in *Appendix D*.
2. Review reports from all inpatient, outpatient, and private outpatient (POP) pathology, cytology, bone marrow, hematology, and autopsy specimens analyzed at your facility.
3. Review records, appointment logs, or rosters of patients seen in the chemotherapy, radiation therapy, and any other area where reportable conditions are diagnosed or treated.
4. Perform quality control procedures to assure all reportable cases were identified and reported to the PCR.

**PART THREE:
DATA ITEM INSTRUCTIONS**

GENERAL INFORMATION

Data Item Completion

Each case reported to the PCR must include all data items identified in *Appendix F, Required Data Set for Reporting Facilities*. These data items must be completed according to codes, definitions, and instructions specified for each item in this section. The codes and definitions for each required data item conform to national cancer registration standards as defined by NAACCR (North American Association of Central Cancer Registries) and ACOS COC (American College of Surgeons Commission on Cancer).

Every effort must be made to obtain specific, complete, and accurate information for each required data item. Inpatient and outpatient health records, clinical history on pathology reports, hospital billing records, and contact with physician offices should be used as sources of information in completing data items.

Recording Unknown or Not Applicable Information

Data items should be recorded as *unknown* only after **all** efforts to obtain specific information prove unsuccessful.

1. Unknown, Text - When specific information is not available for any data item requiring an alphabetic entry, record the word *unknown* in the field as specified in the data item instructions in this section.
2. Unknown, Code 9 - When specific information is not available for any data item requiring a numeric entry, record the code for unknown, *9*, in the field as specified in the data item instructions in this section.

Recording Dates

1. Date Format- Beginning in 2010, the way dates are transmitted was changed to improve the interoperability or communication of cancer registry data with other electronic record systems. Registry software may display dates in the traditional manner or in the interoperable format. Traditional dates are displayed in MMDDCCYY form, with 99 representing unknown day or month portions, and 99999999 representing a completely unknown date. In the traditional form, some dates also permit 88888888 or 00000000 for special meaning.

Interoperable dates are displayed in CCYYMMDD form, with the unknown portions of the date filled with blank spaces. If a date is entirely blank, an associated date flag is used to explain the missing date. If full or partial date is entered, the date flag fields are left blank.

Note: In Abstract Plus, interoperable dates will be entered; therefore the *PCR Manual* will provide details on entering interoperable dates. Registry hospitals should obtain instructions from their vendor on how dates should be entered.

2. Date Flag Fields- Only actual known dates are entered in interoperable date items, date flags explain the reason when there is no value in the corresponding date item. Date flag fields are left blank when a full or partial date is entered. See next page for date flag codes and definitions.

Date Flag Codes and Definitions		
Flag Codes	Description	Definition
bb (blank)	Date provided	Full or partial date was provided; therefore Date Flag field is left blank.
10	No Information	No information whatsoever can be inferred from this exceptional value. It is unknown if this event occurred. (e.g. surgery)
11	Not Applicable	No proper value is applicable in this context. (e.g. no surgery performed)
12	Unknown	A proper value is applicable but not known. The event occurred but the date is unknown. (e.g. birth date)
13	Not Asked	This information has not been sought. (i.e. the patient was not asked)
14	Asked but unknown	Information was sought but not found. (i.e. the patient was asked but did not know)
15	Temporarily unavailable	Information is not available at this time, but it is expected that it will be available later.

The table below illustrates an example of the relationship among these items for *RX Date- Surgery*.

Note: Lower case ‘b’ will be used throughout the *PCR Manual* to indicate blanks in the date fields.

RX Date Surgery Example			
Description	Traditional <i>RX Date-Surgery</i> . Date entered in MMDDCCYY; 99 is entered for unknown portions	Interoperable <i>RX Date-Surgery</i> . Date entered in CCYYMMDD; leaving unknown portions blank. Omit the date if the date is completely unknown.	<i>RX Date-Surgery</i> Flag
Full date known	MMDDCCYY (example: 01102010)	CCYYMMDD (example: 20100110)	bb*
Month and year known	MM99CCYY (example: 01992010)	CCYYMMbb (example: 201001bb)	bb
Year only known	9999CCYY (example: 99992010)	CCYYbbbb (example: 2010bbbb)	bb
Unknown if any surgery performed	99999999 (example: 99999999)	bbbbbbbb (example: bbbbbbbb)	10
No Surgery Performed	00000000 (example: 00000000)	bbbbbbbb (example: bbbbbbbb)	11
<i>RX Date- Surg Primary Site</i> is unknown, but surgery performed	99999999 (example: 99999999)	bbbbbbbb (example: bbbbbbbb)	12

*b= blank

3. Allowable Values:

Year	Month		Day
Use four-digit year	01 January	08 August	01
Blanks* Year Unknown	02 February	09 September	02
9999** Year Unknown	03 March	10 October	03
	04 April	11 November	...
	05 May	12 December	31
	06 June	Blanks*	Blanks* Day Unknown
	07 July	99** Month unknown	99 Day Unknown

*Blanks are only used if entering dates in the interoperable date format CCYYMMDD

**Unknown (99 or 9999) is only used if entering dates in the traditional date format MMDDCCYY

4. Approximating Dates - If the exact date is unknown, use guidelines below to estimate dates from descriptive terms.

a. Approximating Year: Use the following guidelines to estimate year from descriptive terms:

1. Code 'a couple of years' to two years earlier
2. Code 'a few years' to three years earlier.
3. Use whatever information is available to calculate the year, such as 7 years ago.
4. If a descriptive term is not included in this guideline or if there are no descriptive terms available, **do not enter fictitious dates or default values**. Apply the following rules:

- Interoperable Date entry- leave blank
- Traditional Date entry- code as unknown (9999).

b. Approximating Month: Use the following guidelines to estimate month from descriptive terms:

1. Code 'spring of' to April (04)
2. Code 'summer' or 'middle of year' to July (07)
3. Code 'fall' or 'autumn' to October (10)
4. For 'winter of', try to determine whether the physician means the first of the year or the end of the year and code January (01) or December (12) as appropriate.
5. Code 'early in year' to January (01)
6. Code 'late in year' to December (12)
7. Use whatever information is available to calculate the month, such as 7 months ago.
8. If a descriptive term is not included in this guideline or if there are no descriptive terms available, **do not enter fictitious dates or default values**. Apply the following rules:

- Interoperable Date entry- leave blank
 - Traditional Date entry- code as unknown (99)
- c. **Approximating Day:** No approximation of day is acceptable. **Do not enter fictitious days or default values**
- Interoperable Date Entry- leave blank
 - Traditional Date Entry- If day is unknown; enter 99 in the appropriate position in the date field for dates where this is acceptable.
5. **Unknown Dates** - If the month, day, or year is unknown with no descriptions or information to calculate:
- Interoperable Date Entry- leave the appropriate date field position blank. If the date is completely unknown, assign appropriate Date Flag.
 - Traditional Date Entry- enter 99 in the appropriate position in the date field. If the date is completely unknown, assign appropriate Date Flag.
6. **Fictitious Dates** - If any part of a date is unknown and there is no description or guideline to approximate a date for fields where this is acceptable, leave blank for interoperable date entry or enter 99 for traditional date entry. Do not enter fictitious dates or default values such as 15 for unknown day or 0101 when month and day are unknown. Because fictitious dates or default values cannot be differentiated from exact dates when comparing dates reported by different facilities, incorrect dates may be chosen over exact dates during the record consolidation process.

Ill-defined Sites

Throughout the *PCR Manual* "ill-defined sites" is referenced and often has special rules. Below is a listing of what is considered an ill-defined site.

Ill-defined Sites	
C76.0	Head, face and neck, NOS
C76.1	Thorax, NOS
C76.2	Abdomen, NOS
C76.3	Pelvis, NOS
C76.4	Upper limb, NOS
C76.5	Lower limb, NOS
C76.7	Other ill-defined sites
C76.8	Overlapping ill-defined sites

Hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative diseases

Throughout the *PCR Manual* "hematopoietic, reticuloendothelial, immunoproliferative, and myeloproliferative diseases" are referenced and often have special rules. Below is a listing of the primary sites and histologies included in these diseases.

Hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative diseases	
C42.0	Blood
C42.1	Bone marrow
C42.3	Reticuloendothelial system
C42.4	Hematopoietic system
9731/9734	Plasmacytoma
9732	Multiple myeloma
9733	Plasma cell leukemia
9734	Plasmacytoma, extramedullary
9740-9742	Mast cell tumors
9750-9754	Malignant histiocytosis
9755	Histiocytic sarcoma
9756	Langerhans cell sarcoma
9758	Follicular dendritic cell sarcoma
9760	Immunoproliferative disease, NOS
9761	Waldenstrom macroglobulinemia
9762	Heavy chain disease
9764	Immunoproliferative small intestinal disease (Mediterranean lymphoma)
9800-9827, 9831-9920, 9940-9948	Leukemias
9930	Myeloid sarcoma
9931	Acute panmyelosis with fibrosis
9950	Polycythemia vera
9960	Chronic myeloproliferative disorder
9961	Myelosclerosis with myeloid metaplasia
9962	Essential thrombocythemia
9963	Chronic neutrophilic leukemia
9964	Hypereosinophilic syndrome
9965-9967	Myeloid and lymphoid neoplasms
9971	Polymorphic PTLD
9975	Myelodysplastic/Myeloproliferative neoplasm, unclassifiable, Myeloproliferative disease, NOS, Myeloproliferative neoplasm, unclassifiable
9980-9985	Refractory anemias
9986	Myelodysplastic syndrome with 5q deletion syndrome
9987	Therapy-related myelodysplastic syndrome, NOS
9989	Myelodysplastic syndrome, NOS
9991	Refractory neutropenia
9992	Refractory thrombocytopenia

Questions

If you have any questions regarding completion of PCR required data items:

1. Refer to the appropriate section of the *PCR Manual* for detailed instructional information regarding completion of the data item in question.
2. Call your PCR Field Representative at 1-800-272-1850.

NAME - LAST

Record the patient's full last name. Do not leave blank.

Recording Name-Last

1. Blanks, Spaces, Hyphens and Apostrophes are allowed. No other special characters are allowed.
2. Change to Name- This data item should be updated on the hospital abstract if the last name changes and the change must be submitted to the PCR. See *Part One, Changing Information*.

Example: Janet White marries and becomes Janet Black. Change the last name to Black and record White in the maiden name field; forward the change to the PCR.

3. Suffixes and Prefixes- Name suffixes when available must be entered in the field *Name - Suffix* and not included in the *Name - Last* field. Do **not** include name prefixes (e.g., Sister, Reverend, Brother, Dr) as part of the patient last name. Name prefixes are not collected by the PCR and must not be included in any of the required name fields.

NAME - SUFFIX

Record the patient’s name suffix.

Name suffix is a title that follows a patient’s last name. The suffix can identify the generation order in families and provide credential status.

Recording Name-Suffix

1. Blanks, Spaces, Hyphens and Apostrophes are allowed. No other special characters are allowed.
2. No Suffix- Leave this data item blank if the patient does not have a name suffix.
3. Suggested Abbreviation-

<u>Title</u>	<u>Abbreviation</u>
Doctor	MD, PhD
Junior	Jr
Senior	Sr
Third	III
Fourth	IV

4. Multiple Suffixes- If multiple suffixes are used, the generation specific suffix is to be recorded.

Example: The patient’s name is John C. Smith III, MD. Record the III.

5. Prefixes- Do **not** include name prefixes (e.g., Sister, Reverend, Brother, Dr) as part of the patient name suffix. Name prefixes are not collected by the PCR and must not be included in any of the required name fields.

NAME – FIRST

Record the patient's full first name. Do not leave blank.

Recording Name-First

1. Blanks, Spaces, Hyphens and Apostrophes are allowed. No other special characters are allowed.
2. First Initial Only- If the patient uses the initial of their first name and their full middle name, enter the patient's first initial in the *Name - First* field. Record the middle name in the *Name - Middle* field.

Example: Patient's name is M. Jane
(*Name - First*) = M
(*Name - Middle*) = Jane

3. Prefixes- Do **not** include name prefixes (e.g., Sister, Reverend, Brother, Dr) as part of the patient first name. Name prefixes are not collected by the PCR and must not be included in any of the required name fields.

NAME - MIDDLE

Record the patient's middle name.

Recording Name-Middle

1. Middle Initial - Record the middle initial if full name is unknown.
2. No Middle Name or Unknown- Leave this item blank if the patient does not have a middle name or initial, or if the middle name or initial is unknown. Do **not** record *not applicable*, *N/A* or *unknown*.
3. Punctuation- Do not use any punctuation.

NAME - MAIDEN

Record the maiden name of female patients who are or have been married. This item is useful for matching multiple records on the same patient.

Recording Name-Maiden

1. Hyphens are allowed.
2. Unknown or Not Applicable- Leave this data item blank if the patient does not have a maiden name, information is not available, or it is not applicable to the patient as in the case of a male. Do **not** record *not applicable*, *N/A* or *unknown*.

NAME - ALIAS

Record any alternate name or "AKA" (also known as) used by the patient, if known. This item is useful for matching multiple records on the same patient.

Recording Name-Alias

1. Unknown or Not Applicable- Leave this data item blank if the patient does not have an alias or if the information is not available. Do **not** record *not applicable*, *N/A* or *unknown*.
2. Maiden Name- Do not record maiden name in this field. It should be recorded in the *Name-Maiden* field.

GUIDELINES FOR RECORDING PATIENT ADDRESS

The address is the home or residence named by the patient at the time he/she was diagnosed. Legal status and citizenship are not factors in residency decisions. Rules of residency are identical to, or comparable with, the rules of the United States Census Bureau whenever possible. Resolve residency questions by using 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 Statistic rules may differ from census rules. Do not record residence from the death certificate. Review each record carefully to determine correct residence. If address at diagnosis is unavailable, use current address.

Coding Priority

1. Code the Street Address of usual residence as stated by the patient.
2. A post office box is not a reliable source to identify the residency at diagnosis. Post office box addresses do not provide accurate geographical information for analyzing cancer incidence. Use the post office box address only if no street address information is available.

Rules for Persons without Apparent Residences:

1. Persons with More Than One Residence (summer and winter homes): Record the address where the patients spends the majority of time (usual residence). If the usual residence is not known or the information is not available, code the residence the patient specifies at the time of diagnosis.
2. Persons with No Usual Residence (transients, homeless): Use the address of the place they were staying when the cancer was diagnosed. This could be a shelter or the diagnosing institution.
3. Persons Away at School: College students are residents of the school area. Boarding school children below college level are residents of their parents' home.
4. Persons 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 Administration (VA) hospitals
5. 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 family. Military personnel may use the installation address or the surrounding community's address. The Census Bureau has detailed residency rules for Navy personnel, Coast Guard, and maritime ships. Refer to the Census Bureau publications for these detailed rules. <http://www.census.gov>

ADDR AT DX - NO & STREET

Record the number and street address of the patient's usual residence at the time the tumor was initially diagnosed.

Patient address is used to provide census tract and other geocodes for incidence statistics and epidemiologic research. The PCR uses geocoding software for automated assignment of geocodes. To increase the rate of automated geocoding, improve the quality of residence data, and enhance the specificity of residence information available for research, addresses must conform to the following format rules. The USPS Postal Addressing Standards, Pub 28, November 2000 can be found on the Internet at <http://pe.usps.gov/cpim/ftp/pubs/pub28/pub28.pdf>.

Recording Addr at Dx - No & Street

1. Blanks - Leave a blank between numbers and words if space permits.
2. Capital Letters - The use of capital letters is preferred.
Example: 103 First Avenue should be recorded as 103 FIRST AVENUE
3. Multiple Tumors- If the patient has multiple tumors; the address may be different for each primary.
4. No Address at Diagnosis - If no information is available on address at diagnosis, assume the current address was also address at time of original diagnosis.
5. Unknown - If the patient's current address is not known, record UNKNOWN only after **all** efforts to obtain this information prove unsuccessful.
6. Do Not Update this data item if the patient's address changes over time. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.
7. Punctuation: Punctuation marks should be avoided, except when punctuation is necessary to convey the meaning.
 - a. Punctuation normally is limited to periods when the period carries meaning (e.g., 39.2 RD), slashes for fractional addresses (e.g., 101 ½ MAIN ST) and hyphens when the hyphen carries meaning (e.g., 289-01 MONTGOMERY AVE).
 - b. Pound signs- The use of pound signs (#) to designate address units should be avoided whenever possible. The preferred notation is as follows:
Example: Address: 1234 Main St., Apartment #12
 Record as: 1234 MAIN ST APT 12
If a pound sign is used, there must be a space between the pound sign and secondary number (e.g., 425 FLOWER BLVD # 72).
 - c. Do not use commas, semicolons, colons, dashes, question marks, exclamation points, apostrophes, parentheses, brackets, braces, quotation marks or asterisks (*) when recording address.

8. Abbreviations: Enter complete street names without abbreviation. Abbreviate only directional prefixes, directional suffixes and street type suffixes as included on the following PCR list, *Standardized Abbreviations for Street Address*. Use of abbreviations for these terms will enable the entire street address to be recorded.

Examples: 101 W PINE ST HOMESTEAD PA 15120 is in Whitaker Borough
 101 W PINE WAY HOMESTEAD PA 15120 is in Homestead Borough

9. PO Box: Avoid using PO Box numbers in place of street address. Use of street address is necessary for more accurate geocoding.

Example: Address: P.O. Box 20, 221 Springfield Rd
 Record as: 221 SPRINGFIELD RD

10. Postal Route Numbers: Avoid using postal route numbers in place of street address. Confirm the house number is not part of the postal route. Use of street address is necessary for more accurate geocoding.

Example: Address: RD2 35 Sycamore St
 Record as: 35 SYCAMORE ST when it is known that 35 is the street number

11. Apartment Numbers or Letters: Enter apartment numbers or letters in *Address at DX- Supplemental* field.

Example: Address: Apartment F at 321 Knollwood Dr.
 Record *Addr at DX-No and Street* as: 321 KNOLLWOOD DR
 Record *Address at DX- Supplemental* as: APT F

12. Intersections: Use one of the following formats when an intersection is used in place of a street number:

SMITH AND JONES ST (not Sts or Streets)
 SMITH ST AND JONES ST
 SMITH AT JONES

13. Nursing Home or Other Institution: If residence is a nursing home or other institution, enter the street address given in this field. The name of the institution should be entered in the *Address at DX- Supplemental* field.

Example: Address: Oak Nursing Home, 1530 Elm Ave
 Record *Addr at DX-No and Street* as: 1530 ELM AVE
 Record *Address at DX- Supplemental* as: OAK NURSING HOME

PCR STANDARD ABBREVIATIONS FOR STREET ADDRESS

Directional Prefix or Suffix Abbreviations

Prefix/Suffix	Abb
North	N
South	S

Prefix/Suffix	Abb
East	E
West	W

Prefix/Suffix	Abb
Northeast	NE
Northwest	NW

Prefix/Suffix	Abb
Southeast	SE
Southwest	SW

Street Prefix Abbreviations

Prefix	Abb
Avenue	AV, AVE
Boulevard	BLVD
Calle	CLL
Caminito	CMT

Prefix	Abb
Camino	CMN
Circulo	CIR
Corte	CT
Drive	DR

Prefix	Abb
Paseo	PAS
Place/Placita	PL
Plaza	PLZ
Rue	RUE

Prefix	Abb
Via	VIA
Vista	VISTA

Street Suffix Abbreviations

Suffix	Abb
Alley	AL
Alley	ALY
Arcade	ARC
Avenue	AV, AVE
Boulevard	BLVD
Bypass	BYP
Calle	CLL
Causeway	CSWY
Center	CTR
Circle	CIR
Concourse	CONC
Court	CT
Crescent	CRES

Suffix	Abb
Crossing	CRSG
Drive	DR
Expressway	EXWY
Expressway	EXY
Freeway	FRWY
Freeway	FWY
Gardens	GDNS
Highway	HWY
Lane	LA
Loop	LOOP
Mews	MEWS
Motorway	MTWY
Oval	OVAL

Suffix	Abb
Overpass	OVPS
Park	PARK
Parkway	PKWY
Parkway	PKY
Pass	PASS
Path	PATH
Pike	PKE
Place	PL
Plaza	PLZ
Road	RD
Row	ROW
Rue	RUE
Skyway	SKWY

Suffix	Abb
Square	SQ
Street	ST
Terrace	TER
Trafficway	FWY
Throughway	THWY
Trail	TRL
Turnpike	TPKE
Underpass	UNP
Walk	WALK
Way	WY

ADDR AT DX - SUPPLEMENTL

Record additional address information such as the name of a place or facility (e.g., a nursing home or name of an apartment complex) at the time of diagnosis.

Recording Addr at Dx - Supplementl

1. Not Applicable- If additional address space is not needed, leave blank.
2. Do Not Update this data item if the patient's address changes over time. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.

ADDR AT DX - CITY

Record the city or town of the patient's usual residence when the tumor was initially diagnosed. The address is a part of the patient's demographic data and has multiple uses. It will provide a referral pattern report and allow analysis of cancer clusters or environmental studies.

Recording Addr at DX-City

1. Do Not Update this data item if the patient's address changes over time. Changing this data item would destroy its usefulness. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.
2. Rural area- If the patient resides in a rural area, record the name of the city or town used in his or her mailing address.
3. Punctuation- Do not use punctuation, special characters, or abbreviations.
4. Capital Letters- The use of capital letters is preferred.
5. Multiple Tumors- If the patient has multiple tumors; the address may be different for each primary.
6. Unknown- If the city is not known, record UNKNOWN only after **all** efforts to obtain this information prove unsuccessful.
7. No Information- If no information is available on address at time of diagnosis, use current address.

ADDR AT DX – STATE

Record the US postal service abbreviation for the state or Canadian province of the patient’s usual residence when the tumor was diagnosed.

Recording Addr at DX-State

1. Multiple Tumors- If the patient has multiple tumors; the address may be different for each primary.
2. Do Not Update this data item if the patient’s address changes over time. Changing this data item would destroy its usefulness. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.
3. Abbreviations- Only abbreviations on the following three tables are acceptable.

Abbreviations - US States & Possessions

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

Abbreviations - Other US Possessions

US Possession		US Possession	
American Samoa	AS	Marshall Islands	MH
Guam	GU	Outlying Islands	UM
Puerto Rico	PR	APO/FPO Armed Services America	AA
Virgin Islands	VI	APO/FPO Armed Services Europe	AE
Palau	PW	APO/FPO Armed Services Pacific	AP
Micronesia	FM		

Abbreviations - Canadian Provinces

Province		Province	
Alberta	AB	Nunavut	NU
British Columbia	BC	Ontario	ON
Manitoba	MB	Prince Edward Island	PE
New Brunswick	NB	Quebec	QC
Newfoundland/Labrador	NL	Saskatchewan	SK
Northwest Territories	NT	Yukon	YT
Nova Scotia	NS		

Abbreviations - Other

Other Country or Unknown	
Resident of a country other than the U.S. (including its territories, commonwealths, or possessions) or Canada and the country is known.	XX
Resident of a country other than the U.S. (including its territories, commonwealths, or possessions) or Canada and the country is unknown.	YY
Resident of U.S., NOS (including its territories, commonwealths, or possessions); Canada, NOS; residence unknown.	ZZ

The address is a part of the patient’s demographic data and has multiple uses. It will provide a referral pattern report and allow analysis of cancer clusters or environmental studies. Do not update this data item if the patient’s address changes over time. Changing this data item would destroy its usefulness. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.

ADDR AT DX - POSTAL CODE

For US residents, record the patient’s nine-digit extended postal (ZIP) code when the tumor was diagnosed. The address is a part of the patient’s demographic data and has multiple uses. It will provide a referral pattern report and allow analysis of cancer clusters or environmental studies.

Example: The extended postal code 60611-2797 is recorded as 606112797.

Recording Addr at DX- Postal Code

1. Only Five-digit Available- When the nine-digit extended code is unavailable, record the five-digit postal code.

Example: When only five digits, 60611, are available, record 60611_ _ _ _.

2. Canadian Residents- for Canadian residents, record the six-character postal code.
3. Hyphens- Do not record hyphens.
4. Do Not Update this data item if patient’s address changes over time. Changing this data item would destroy its usefulness. See *Part Three, Guidelines for Recording Patient Address* for detailed residency rules.
5. Multiple Tumors- If the patient has multiple tumors; the postal code may be different for each primary.
6. Other countries- When available, record the postal code for other countries.
7. Unknown Postal Code- If the street address, city and state are known, but the postal code is unknown, the following US Postal Service's web site may be used to determine the correct postal code:
<http://www.usps.com/>
8. Unknown Address- If street address, city, state and postal code are unknown and the information cannot be obtained from any other sources, use the following codes:

Codes and Definitions

Code	Definition
888888888	Permanent address in a country other than Canada, United States or US possessions and postal code is unknown.
999999999	Permanent address in Canada, United States, or US possession, and postal code is unknown. Permanent address (street, city and state) is totally unknown.

COUNTY AT DX

Record the county of the patient’s usual residence when the tumor was diagnosed using county codes issued by the Bureau of Standards in the Federal Information Processing Standards (FIPS). The FIPS codes for Pennsylvania counties are listed in *Appendix H* and are generally incorporated into abstracting software.

Recording County at Dx

1. **Multiple Tumors-** If the patient has multiple tumors; the county may be different for each primary.
2. **Do Not Leave Blank-** This data item must contain either the specific code for county at diagnosis or the appropriate unknown code. If the city and state are known, but the county is unknown, the following web site may be used to determine the correct county:
<http://www.melissadata.com/Lookups/addressverify.asp>.
3. **Pennsylvania Resident-** If the patient is a Pennsylvania resident, the specific county must be recorded when known.
4. **Outside of Pennsylvania-** If the patient resides in a state other than Pennsylvania, in Canada, or in a US possession, the specific county is not required but encouraged to be reported if known.
5. **Known Other Countries-** If the patient resides outside the US, Canada, or a US possession and the country is known. (XX is entered for *Addr at Dx - State*); the country of residence at diagnosis is recorded in this data item.
 - **SEER Geo-Codes-** Record the three-digit country code in which the patient resided at diagnosis. These codes are generally incorporated into abstracting software but they are also listed in *Appendix H, SEER Geo-Codes for Country*.
6. **Unknown Other Country-** If the patient resides outside the US, Canada, or a US possession but the country is unknown. (YY is entered for *Addr at Dx - State*); Record 999.

Additional Codes and Definitions

In addition to the FIPS and Geo-codes, the following codes are acceptable:

Code	Definition
998	Patient resides outside of Pennsylvania and exact county code cannot be found.
999	Unknown county/country

AGE AT DIAGNOSIS

Record the patient's age at his/her last birthday before diagnosis

000	Less than one year old
001	One year old, but less than two years old
002	Two years old
...	(Actual age in years)
101	One hundred one years old
...	
999	Unknown age

Recording Age at Diagnosis

1. Calculating Age- If age at diagnosis is unavailable, but the year of diagnosis and year of birth are known, calculate approximate age at diagnosis.
2. Date of Birth Unknown- Use 999 if the date of birth is unknown.
3. Date of Diagnosis Unknown- Use 999 if the date of diagnosis is unknown.

DATE OF BIRTH

Record the patient's date of birth.

Note: In the examples below the lowercase letter "b" is used to represent each blank space.

Recording Date of Birth

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 1906 as 19060630.

2. Date Unavailable, but Age Known- When age is known, estimate year of birth when further information is not available. It is better to estimate than to record as an unknown year.

Example 1: The patient is 60 years old when diagnosed on June 15, 2010. The medical record does not have a date of birth. Estimate the year as 1950 and leave month and day blank (1950bbbb).

Example 2: Record the patient's date of birth as 1927 when the medical record contains only the year of birth (1927bbbb).

3. Calculating Date of Birth- If date of birth is unavailable, but the year of diagnosis and age are known, calculate approximate date of birth. Leave this field blank only if the age at diagnosis is also unknown.
4. Unknown Month, Day and/or Year - If date is not known, leave blank. If only part of the date is known, record what is known and enter approximation for month and/or year if descriptions are available or leave blank for what is unknown. No approximation of day is acceptable. Refer to *Part Three: Data Item Instructions, General Information, Dates* for instructions regarding Approximating Dates and Unknown Dates. Fictitious dates or default values are not acceptable to be entered for month, day, or year.

Special Instruction for Registry Hospitals

Some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.

DATE OF BIRTH FLAG

Record the date flag in the event a complete *Date of Birth* was not entered to explain why.

Recording Date of Birth Flag

1. Full or Partial Date- Leave this field blank if *Date of Birth* has a full or partial date recorded.
2. Unknown Date- Code 12 if the *Date of Birth* cannot be determined at all.
3. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
12	A proper value is applicable but not known (for example, date of birth is unknown)
(blank)	A valid date value is provided in item <i>Date of Birth</i>

The following table illustrates the use of the date flag and the traditional and interoperable date formats for coding *Date of Birth* and *Date of Birth Flag*. In the table below, the lowercase letter “b” is used to represent each blank space.

Description	Traditional Date of Birth (MMDDCCYY)	Interoperable Date of Birth (CCYYMMDD)	Date of Birth Flag
Full date known	MMDDCCYY Example:04031968	CCYYMMDD Example: 19680403	bb
Month and year known	MM99CCYY Example: 04991968	CCYYMMbb Example: 196804bb	bb
Year only known	9999CCYY Example: 99991968	CCYYbbbb Example: 1968bbbb	bb
Unknown date	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12

b=blank

Special Instruction for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

BIRTHPLACE- STATE

Record the state, commonwealth, United States possession or Canadian Province in which the patient was born. This item corresponds to *Birthplace-Country*.

Recording Birthplace-State

Born in the United States/Canada

1. United States- Record the patient’s state of birth using the standard United States Postal Service Abbreviations. See *Appendix I* for list state abbreviations and their respective country abbreviations.
2. Canada- Record the patient’s province/territory of birth using the standard CanadaPost Abbreviations. See *Appendix I* for list province abbreviations and their respective country abbreviations.
3. Unknown State- If it is known the patient was born in the United States but the specific state is unknown, record *US*.
4. Unknown Province/territory- If it is known the patient was born in Canada but the specific province/territory is unknown, record *CD*.

Born Outside the United States

1. Country of Birth- If patient was born in a country other than the United States (including territories, commonwealths or possessions) or Canada and the country is known, enter *XX* and code the country of birth in *Birthplace-Country*.
2. Unknown Country- If patient was born in a country other than the United States (including territories, commonwealths or possessions) or Canada and the country is unknown, enter *YY*.

Birth Place Unavailable

- Information Unavailable- Record *ZZ* when place of birth is unavailable.

Non-Standard Codes and Definitions

Code	Definition
XX	Born in a country other than the United States (including territories, commonwealths or possessions) or Canada and the country is <u>known</u>
YY	Born in a country other than the United States (including territories, commonwealths or possessions) or Canada and the country is <u>unknown</u>
US	Born in the United States (including territories, commonwealths or possessions) and the state is unknown
CD	Born in Canada and the province/territory is unknown
ZZ	Place of birth is unknown, not mentioned in patient record

BIRTHPLACE- COUNTRY

Record the country in which the patient was born. This item corresponds to *Birthplace-Country*.

Recording Birthplace-Country

1. United States- If it is known the patient was born in the United States record *USA*. The corresponding state is recorded in *Birthplace-State*
2. Canada- If it is known the patient was born in Canada record *CAN*. The corresponding province/territory is recorded in *Birthplace-State*
3. Other Countries- If it known the patient was born in a country other than the United States or Canada, record the patient's country of birth using the country codes provided in *Appendix I*. The codes are based on International Organization for Standardization (ISO) with some custom codes.
4. Unknown Place of Birth- If it is completely unknown where the patient was born, record *ZZU*.

SOCIAL SECURITY NUMBER

Record the patient's Social Security Number (SSN) without dashes.

Recording Social Security Number

1. No Social Security Number- When a patient does not have a Social Security Number, or the information is not available, record 999999999.
2. Correct Social Security Number- It is important to enter the correct Social Security Number since this data item is used for record linkage to match patients at the PCR as well as to match PCR information with the Social Security Number on the hospital's Disease Index. Verify entries for missing values and transpositions. Do not record Social Security Numbers that end with B or D. These are the spouse's Social Security Number.
3. Invalid Entry- According to how a Social Security Number is assigned by the Social Security Administration, the following are invalid entries:
 - a. First three digits cannot = 000 or 666
 - b. Fourth and Fifth digits cannot = 00
 - c. Last four digits cannot = 0000
 - d. First digit cannot = 8 or 9 unless entire SSN is unknown (999999999)
4. Correction- If a correction is made to the Social Security Number, a change sheet must be submitted to the PCR. See *Part One, Changing Information*.

SEX

Record the patient’s sex.

Codes and Definitions

Code	Definition
1	Male
2	Female
3	Other (Hermaphrodite, Intersexed, persons with sex chromosome abnormalities)
4	Transsexual, NOS
5	Transsexual, natal male
6	Transsexual, natal female
9	Not stated/Unknown

Guidelines

1. **Transsexual/Transgender Definition:** A person who was assigned to one gender at birth based on physical characteristics but who self-identifies psychologically and emotionally as the other gender.
2. **Transgendered person Definition:** A person who identifies with or expresses a gender identity that differs from the one which corresponds to the person’s sex at birth.
3. **Intersexed:** Assign code **3** for Intersexed (persons with sex chromosome abnormalities)
4. **Natally Male/ Male Primary Sites:** Assign code **5** for transsexuals who are natally male or transsexuals with primary site of C600-C639
5. **Natally Female/Female Primary Sites:** Assign code **6** for transsexuals who are natally female or transsexuals with primary site of C510-C589
6. **Unknown Natal Sex/Non gender specific primary site:** Assign code **4** for transsexuals with unknown natal sex and primary site is not C510-C589 or C600-C639
7. **Unknown Gender:** When gender is not known:
 - a. Assign code **1** when the primary site is C600-C639
 - b. Assign code **2** when the primary site is C510-C589
 - c. Assign code **9** for primary sites not included above

SPANISH/HISPANIC ORIGIN

Record the Spanish/Hispanic origin. This item identifies persons of Spanish or Hispanic ethnicity.

Codes and Definitions

Code	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 maiden name 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 maiden name and there is no contrary evidence that the person is not Hispanic)
8	Dominican Republic
9	Unknown whether Spanish or not

Recording Spanish/Hispanic Origin

1. Any race- A person of Spanish/Hispanic origin may be any race, but these categories are generally not used for Native Americans, Filipinos, or others who may have Spanish names.
2. Portuguese and Brazilian- Code 0 (Non-Spanish; non-Hispanic) for Portuguese and Brazilian persons.
3. Multiple Tumors- If a patient has multiple tumors all records should have the same code.
4. Hispanic Surname or Maiden Name- Code 7 (Spanish surname only) 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.
5. Unknown- Code 9 (Unknown whether Spanish or not) if the Spanish/Hispanic information is unknown.

6. Use All Information- All information should be used to determine the Spanish/Hispanic Origin including:
- The stated ethnicity in the medical record
 - Stated Hispanic origin on the death certificate
 - Birthplace
 - Information about life history and/or language spoken found in the abstracting process
 - A last name or maiden name found on a list of Hispanic/Spanish names

RACE

RACE 1, RACE 2, RACE 3, RACE 4, RACE 5

Record the appropriate codes for the patient’s race(s) in *Race 1, Race 2, Race 3, Race 4, and Race 5*. Race is coded separately from Spanish/Hispanic Origin.

Race (and ethnicity) is defined by specific physical, hereditary and cultural traditions or origins, not necessarily by birthplace, place of residence, or citizenship. ‘Origin’ is defined by the US 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.

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 electronic form, the electronic data must also be reviewed.

Codes and Definitions

Code	Definition	Code	Definition
01	White	20	Micronesian
02	Black	21	Chamorroan
03	American Indian, Aleutian, Eskimo	22	Guamanian, NOS
04	Chinese	25	Polynesian, NOS
05	Japanese	26	Tahitian
06	Filipino	27	Samoan
07	Hawaiian	28	Tongan
08	Korean	30	Melanesian, NOS
10	Vietnamese	31	Fiji Islander
11	Laotian	32	New Guinean
12	Hmong	88	No further race documented (Do Not use in Race 1)
13	Kampuchean, includes Khmer & Cambodian	96	Other Asian, includes Asian NOS & Oriental NOS
14	Thai	97	Pacific Islander, NOS
15	Asian Indian, Pakistani, NOS (formerly code 09)	98	Other
16	Asian Indian	99	Unknown
17	Pakistani, NOS		

Recording Race

Single Race

1. **One Race-** If only one race is reported for the patient, in *Race 1* enter the race code and in *Race 2* through *Race 5*, enter 88.
2. **A specific race code** (other than 88 or 99) must not occur more than once.

Example 1: If the patient's race is listed as white, in *Race 1* enter 01 and in *Race 2* through *Race 5* enter 88. Do not code 01 in *Race 1* signifying one parent and 01 again in *Race 2* for other parent.

Example 2: A patient was born in Mexico of Mexican parentage. Code *Race 1* as 01 and *Race 2* through *Race 5* as 88

Multiple Races

1. **Primary Race(s)** - Code primary race(s) of the patient in fields *Race 1*, *Race 2*, *Race 3*, *Race 4*, and *Race 5*. The five race fields allow for the coding of multiple races consistent with the Census 2000. Rules 2-6 further specify how to code *Race 1* through *Race 5*.

2. **Less Than Five Specific Races-** If less than five specific race codes apply for a patient, code 88 in the remaining race fields.

Example: A patient has a Hawaiian father, black mother, Japanese grandfather, and Korean grandmother. Code *Race 1* as 07 Hawaiian, *Race 2* as 02 Black, *Race 3* as 05 Japanese, *Race 4* as 08 Korean, and *Race 5* as 88

3. **White and Other Races-** If a person's race is a combination of white and any other race(s), code the appropriate other race(s) first and code white in the next race field.
4. **Hawaiian and Other Races-** If a person's race is a combination of Hawaiian and any other race(s), code *Race 1* as 07 Hawaiian and code the other races in *Race 2*, *Race 3*, *Race 4*, and *Race 5* as appropriate.

Example: Patient is described as Japanese and Hawaiian. Code *Race 1* as 07, Hawaiian, *Race 2* as 05 Japanese, and *Race 3* through *Race 5* as 88

5. **Combination without Hawaiian-** If the person is not Hawaiian, code *Race 1* to the first stated non-white race (02-98).

Example: Patient is stated to be Vietnamese and Black. Code *Race 1* as 10 Vietnamese, *Race 2* as 02 Black, and *Race 3* through *Race 5* as 88

6. **Specific Race and Non-Specific Race-** Code only the specific race when both a specific race code and a non-specific race code apply.
 - a. Codes 04-17 take priority over code 96
 - b. Codes 16-17 take priority over code 15
 - c. Codes 20-32 take priority over code 97
 - d. Codes 02-32 and 96-97 take priority over code 98
 - e. Code 98 takes priority over code 99

7. Stated Race- Code the patient's stated race. Refer to Appendix D "Race and Nationality Descriptions from the 2000 Census and Bureau of Vital Statistics" of the SEER Program and Staging Manual for guidance. <http://seer.cancer.gov/tools/codingmanuals/index.html>

Example 1: Patient is stated to be Japanese. Code *Race 1* as 05 Japanese and *Race 2* through *Race 5* as 88

Example 2: Patient is stated to be German-Irish. Code *Race 1* as 01 White and *Race 2* through *Race 5* as 88

Example 3: Patient is described as Arabian. Code *Race 1* as 01 White and *Race 2* through *Race 5* as 88

Exception: When the race is recorded as Oriental, Mongolian, or Asian (coded to 96 Other Asian) and the place of birth is recorded as China, Japan, the Philippines, or another Asian nation, code the race based on birthplace information.

Example 4: The person's race is recorded as Asian and the place of birth is recorded as Japan. Code *Race 1* as 05 Japanese because it is more specific than 96 Asian, NOS and *Race 2* through *Race 5* as 88

Example 5: The person describes himself as an Asian-American born in Laos. Code *Race 1* as 11 Laotian because it is more specific than 96 Asian, NOS and *Race 2* through *Race 5* as 88.

8. Based on Race of Relatives- If the patient's race is determined on the basis of the races of relatives, there is no priority to coding race, other than to list the non-white race(s) first.

Example: The patient is described as Asian-American with Korean parents. Code *Race 1* as 08 Korean because it is more specific than 96 Asian, NOS and code *Race 2* through *Race 5* as 88.

No Race Stated

1. Race Category- If no race is stated in the medical record, or if the stated race cannot be coded, review the documentation for a statement of race category.

Example 1: Patient described as a black female in the physical exam, consultation or nursing notes, Code *Race 1* as 02 Black and *Race 2* through *Race 5* as 88.

Example 2: Patient describes herself as multi-racial (nothing more specific) and nursing notes say 'African-American.' Code *Race 1* as 02 Black and *Race 2* through *Race 5* as 88

Example 3: 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

2. If race is unknown, not stated in the medical record, or not stated specifically, refer to the race-specific guidelines on the next page. If none apply, code *Race 1* through *Race 5* as unknown (99). Do not use patient name in determining race.

Race-Specific Guidelines

1. White (01) includes Mexican, Central American, South American, Puerto Rican, Cuban, and all other Caucasians.
2. Black (02) includes the designations Negro or African-Americans.
3. Native American (03) should be used for any person stated to be Native American or [western hemisphere] Indian, whether from North, Central, South, or Latin America.
4. Birthplace Information- Race is based on birthplace information when place of birth is given as China, Japan, or the Philippines and race is reported only as Asian, Oriental, or Mongolian.

Example: If the patient's race is recorded as Asian and the place of birth is recorded as Japan, code *Race 1* as 05 Japanese and *Race 2* through *Race 5* as 88.

5. Asian- Do not code Asian in a subsequent race field if a specific Asian race has already been coded.

Use of Code 88 (No further race documented)

1. Race 1- Code 88 is valid for *Race 2* through *Race 5*; it is not valid for *Race 1*.
2. Race 2-5- If *Race 2* is coded to 88, then *Race 3* through *Race 5* must be coded to 88.

Use of Code 99 (Unknown)

1. If the patient's race is unknown, enter 99 in *Race 1* through *Race 5*.
2. If any race equals 99 then all race codes (*Race 1*, *2*, *3*, *4*, and *5*) must equal 99.

PRIMARY PAYER AT DIAGNOSIS

Record the patient's primary payer/insurance carrier at the time of initial diagnosis and/or treatment.

This item is used in financial analysis and as an indicator for quality and outcome analyses. Joint Commission on Accreditation of Healthcare Organizations (JCAHO) requires the patient admission page to document the type of insurance or payment structure that will cover the patient while being cared for at the facility.

Codes and Definitions

Code	Definition
01	<i>Not Insured-</i> Patient has no insurance and is declared a charity write-off.
02	<i>Not Insured, Self Pay-</i> Patient has no insurance and is declared responsible for charges.
10	<i>Insurance, NOS-</i> Type of insurance is unknown or other than types listed in codes 20, 21, 31, 35, 60-68.
20	<i>Private Insurance: Managed Care, HMO, or PPO-</i> 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.
21	<i>Private Insurance: Fee-for-Service-</i> An insurance plan that does not have a negotiated fee structure with the participating facility. Type of insurance plan not coded as 20.
31	<i>Medicaid-</i> State government administered insurance for persons who are uninsured, below poverty level, or covered under entitlement programs. Medicaid other than described in code 35.
35	<i>Medicaid-Administered through a Managed Care plan-</i> Patient is enrolled in Medicaid through a Managed Care program (e.g. HMO or PPO). The managed care plan pays for incurred costs.
60	<i>Medicare without supplement, Medicare, NOS-</i> Federal government funded insurance for persons who are 62 years of age and older, or are chronically disabled (social security insurance eligible). Not described in codes 61, 62, or 63.
61	<i>Medicare with supplement, NOS-</i> Patient has Medicare and another type of unspecified insurance to pay costs not covered by Medicare.
62	<i>Medicare-Administered through a Managed Care Plan-</i> 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	<i>Medicare with private supplement-</i> Patient has Medicare and private insurance to pay costs not covered by Medicare.

Code	Definition
64	<i>Medicare with Medicaid eligibility</i> - Federal government Medicare with State Medicaid administered supplement.
65	<i>TRICARE</i> - Department of Defense program providing supplementary civilian-sector hospital and medical services beyond a military treatment facility to military dependents, retirees, and their dependents. Formerly CHAMPUS (Civilian Health and Medical Program of the Uniformed Services)
66	<i>Military</i> - Military personnel or their dependents that are treated at a military facility.
67	<i>Veterans Affairs</i> - Veterans who are treated in Veterans Affairs facilities.
68	<i>Indian/Public Health Service</i> - Patient who receives care at an Indian Health Service facility or another facility, and the 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	<i>Insurance Status Unknown</i> - It is unknown from the patient's medical record whether or not the patient is insured.

Recording Primary Payer at Diagnosis

1. Type of Insurance- Record the type of insurance reported on the patient's admission page.
2. More than One Payer- If more than one payer or insurance carrier is listed on the patient's admission page, record the first.
3. Changes in Payer- If the patient's payer or insurance carrier changes do not change the initially recorded code.

TEXT - USUAL OCCUPATION

Record the patient's usual occupation, the kind of work performed during most of the patient's working life before diagnosis of this tumor.

This data item is used to identify new work-related health hazards, serves as an additional measure of socioeconomic status, and identifies occupational groups in which cancer screening or prevention activities may be beneficial.

Usual occupation is defined identically as on death certificates and conforms to the 1989 revision of the US Standard Certificate of Death.

Recording Text-Usual Occupation

1. Retired- Do not record "retired".
2. If Not Available or Unknown- If *usual* occupation is not available or is unknown, record the patient's current or most recent occupation or any known occupation.
3. Update this data item if better information is obtained as to the usual occupation of the patient. However, it is not the responsibility of facility abstractors to update abstracts with information provided on death certificates. Comparison with death certificate information is the function of the PCR.
4. Housewife/househusband- If the patient was a housewife/househusband and also worked outside the home most of her/his adult life, record the usual occupation outside the home. If the patient was a housewife/ househusband and did not work outside the home for most of her/his adult life, record *housewife* or *househusband*.
5. Never Worked- If the patient was not a student or housewife and never worked, record *never worked* as the usual occupation.
6. No Information- If no information is available, record *unknown*.
7. This data item cannot be blank unless the patient is under 14 years old. It applies only to patients who are 14 years or older at the time of diagnosis. For patients under the age of 14, leave blank.
8. Finding the Information- The patient's occupation may be found on the face sheet, nursing assessment, history and physical or consult reports in the medical record.

TEXT - USUAL INDUSTRY

Record the primary type of activity carried on by the business/industry where the patient was employed for the most number of years before diagnosis of this tumor.

Both occupation and business/industry are required to accurately describe an individual's occupation. These data items are used to identify new work-related health hazards, serve as an additional measure of socioeconomic status, and identify occupational groups in which cancer screening or prevention activities may be beneficial.

Usual industry (also known as "kind of business/industry") is defined identically as on death certificates and conforms to the 1989 revision of the US Standard Certificate of Death.

Recording Text-Usual Industry

1. Distinguish the Component- Be sure to distinguish among *manufacturing, wholesale, retail,* and *service* components of an industry that performs more than one of these components.
2. Primary Activity Unknown- If the primary activity carried on at the location where the patient worked is unknown, it may be sufficient to record the name of the company (with city or town) for which the patient performed his/her usual occupation. In these situations, if resources permit, the PCR may be able to use the employer name and city/town to determine the type of activity conducted at that location.
3. If Most Recent Occupation was Recorded- If current or most recent occupation, rather than usual occupation was recorded, record the patient's current or most recent business/industry.
4. Update this data item if better information is obtained as to the usual industry of the patient. However, it is not the responsibility of facility abstractors to update abstracts with industry information provided on death certificates. Comparison with death certificate information is the function of the PCR.
5. No Information Available- There must be an entry for usual industry when any occupation is reported. If no information is available regarding the industry in which the reported occupation was carried out or the occupation is unknown, record *unknown*.
6. This data item cannot be blank unless the patient is under 14 years old. It applies only to patients who are 14 years or older at the time of diagnosis. For patients under the age of 14, leave blank.

MEDICAL RECORD NUMBER

Record the patient's medical record number. The medical record number is a patient identification number usually assigned by the reporting facility.

Recording Medical Record Number

1. Match to Disease Index- This item is used to locate the medical record. It may also be used to link records and should be recorded **exactly** as it is recorded on your Disease Index.
2. Fewer Than Eleven Characters- If the medical record number is fewer than 11 characters, right justify the characters and allow leading blanks.

Example: Medical record number 811234 would be recorded bbbbb811234 (b=blank)

3. Departments without Medical Record Numbers- Record standard abbreviations for departments that do not use medical record numbers.

Examples: Radiation Therapy record bbbbbbRT (b=blank)

One-day surgery clinic record bbbbbbSU (b=blank)

4. Unknown- If the medical record number is unknown, record bbbbbbUNK (b=blank)

ACCESSION NUMBER - HOSP

Record the registry accession number assigned by your facility.

Recording Accession Number-Hosp

1. First Four Numbers- The first four numbers of the registry accession number specify the century and year in which the patient was first seen at the reporting institution for diagnosis and/or treatment.
2. Last Five Numbers- The last five numbers are the numeric order in which the registry entered the record into the database.

Example: A patient is diagnosed at the reporting institution in 2010. The first four digits of the registry accession number are 2010. This is the 33rd patient accessioned in 2010, making the last five digits of the registry accession number 00033. The full registry accession number is 201000033.

3. Unique Number for Each Patient- Assign a unique accession number to each patient. The accession number identifies the patient even if multiple primaries exist. Use the same accession number for all subsequent primaries. A patient's accession number should not be reassigned.
4. Options- Hospitals may choose one of the following three options for completing this data item:
 - a. Complete as described above (For more detailed instructions on assigning accession number, refer to *ACOS COC Facility Oncology Registry Data Standards (FORDS)*).
 - b. Use to sequentially number your records forwarded to the PCR.
 - c. Leave this item blank.

SEQUENCE NUMBER – HOSPITAL

Record the sequence number representing the order of this primary. Sequence number counts the occurrence of *independent, malignant and non-malignant neoplasms* except basal and squamous cell cancer of the skin **during the patient's lifetime**. Each neoplasm is assigned a different number. This number may change over the lifetime of the patient.

Codes 00-35 and 99 indicate neoplasms of in situ or malignant behavior (2 or 3). Codes 60-88 indicate neoplasms of non-malignant behavior (0, benign or 1, borderline).

Sequence Numbers for Malignant or In Situ Primaries

00	One malignant or in situ primary only in the patient's lifetime
01	First of two or more independent malignant or in situ primaries
02	Second of two or more independent malignant or in situ primaries
...	(Actual sequence of this malignant or in situ primary)
35	Thirty-fifth of thirty-five independent malignant or in-situ primaries
99	Unspecified malignant or in situ sequence number or unknown

Sequence Numbers for Non-Malignant Tumors

60	Only one non-malignant primary in the patient's lifetime
61	First of two or more independent non-malignant primaries
62	Second of two or more independent non-malignant primaries
...	(Actual number of this primary)
87	Twenty-seventh of twenty-seven independent non-malignant primaries
88	Unspecified number of neoplasms in this category

Recording Sequence Number

1. Single Malignant Primary Tumor- Code 00 only if the patient has a single malignant primary.
2. Subsequent Malignant or In Situ Primary Tumor- If the patient develops a subsequent malignant primary or in situ primary tumor, change the sequence number for the first tumor from 00 to 01, and number subsequent tumors sequentially.

Example: In January 2009, the registry assigns sequence number 00 to a patient with malignant melanoma. The patient develops a second primary cancer of the lung in July 2010. Assign sequence number 02 to the second cancer (lung). Change the sequence number of the first cancer (malignant melanoma) to 01.

Note: Reporting institutions are not required to forward a change sheet to the PCR when changing sequence number from 00 to 01.

3. Single Non-Malignant Primary Tumor- Code 60 only if the patient has a single non-malignant primary.

4. Subsequent Non-Malignant Primary Tumor- If the patient develops a subsequent non-malignant primary, change the sequence number of the first tumor from 60 to 61, and number subsequent non-malignant tumors sequentially.

Note: Reporting institutions are not required to forward a change sheet to the PCR when changing sequence number from 60 to 61.

5. Two or More Malignant Neoplasms Diagnosed at the Same Time- 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.

Example 1: A patient enters the reporting institution with simultaneous carcinoma in situ of the breast and invasive adenocarcinoma of the colon. Assign sequence number 01 to the colon primary and sequence number 02 to the breast primary.

Example 2: A patient has simultaneous adenocarcinoma in situ in a colon polyp and squamous cell carcinoma in situ in a vocal cord polyp. Assign sequence numbers in any order, since both primaries have similar prognoses.

6. Two or More Non-Malignant Neoplasms Diagnosed at the Same Time- 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.

7. In Situ Tumor Followed by an Invasive Cancer- If an in situ tumor is followed by an invasive cancer in the same site more than two months apart, report as two primaries even if stated to be a recurrence. The invasive primary should be reported with the date of the invasive diagnosis. Assign sequence numbers to both primaries with the in situ cancer being the first of the two.

8. Location and Date of Diagnosis- The sequence number counts the patient's independent, primary tumors regardless of the location(s) or institution(s) where those primaries were diagnosed and treated or the date of diagnosis.

Example: The reporting institution diagnosed colon cancer. The patient has a history of kidney cancer diagnosed in 1980. The colon cancer is the second of this patient's primary cancers. Assign a sequence number 02 to colon cancer.

9. Newly Reportable Conditions- If the patient has a condition that was diagnosed prior to the condition being reportable do not count that condition when assigning sequence number.

Example: A patient was diagnosed with refractory anemia on June 25, 1999 (not reportable until 2001) and then was later diagnosed with acute myelogenous leukemia on March 21, 2010 at your facility. Abstract only the acute myelogenous leukemia and assign a Sequence Number of 00.

10. Unaccessioned Tumor- Sequence numbers should be reassigned if the facility learns later of an unaccessioned tumor that affects the sequence.

11. Unknown- Use the sequence number 99 when it is impossible to estimate whether the patient has been diagnosed with an earlier malignancy (primary). If more information becomes available, change the sequence number(s).

Example: A patient is diagnosed in the reporting facility with cancer of the colon. The medical

record contains the statement “The patient recently had a salivary gland tumor removed. The patient does not know if the lesion was malignant.” Assign a 99 sequence number to the colon primary. The patient returns to the reporting facility a year later for treatment of prostate cancer. The medical record says “The patient has a history of a malignant salivary gland tumor.” Change the sequence number of the colon cancer from 99 to 02. Assign the sequence number 03 to the prostate cancer.

CLASS OF CASE

Record the Class of Case code to reflect the facilities role in the managing the cancer, whether the cancer is required to be reported and whether the case was diagnosed after a program’s Reference Date. *Class of Case* divides cases into two groups. Analytic cases (codes 00–22) which are grouped according to the location of diagnosis and first course of treatment. Nonanalytic cases (codes 30–49 and 99) which are grouped according to the reason a patient who received care at the facility is nonanalytic, or the reason a patient who never received care at the facility may have been abstracted.

Analytic and Nonanalytic Cases

Class of Case shows the role the reporting institution played in the patient's diagnosis or treatment. *Class of Case* codes are further categorized into analytic and nonanalytic categories to reflect where the initial diagnosis or treatment occurred. Both analytic and nonanalytic cases are reportable to the PCR.

1. Analytic Cases (Class of Case 00-22) - Cases diagnosed and/or administered any of the first course of treatment at the accessioning facility after the registry’s reference date are analytic (Class of Case 00-22). A network clinic or outpatient center belonging to the facility is considered part of the facility.
2. Nonanalytic Cases (Class of Case 30-49 and 99) – are to be abstracted by the facility to meet PCR reporting requirements.

Codes and Definitions

Analytic Classes of Case	
Initial Diagnosis At Reporting Facility	
00	Initial diagnosis at the reporting facility AND all treatment or a decision not to treat was done elsewhere <i>Note:</i> Code 00 applies only when it is known the patient went elsewhere for treatment. If that information is not available, code <i>Class of Case</i> 10.
10	Initial diagnosis at the reporting facility or in a staff physician’s office AND part or all of first course treatment or a decision not to treat was at the reporting facility, NOS
11	Initial diagnosis in staff physician’s office AND part of first course treatment was done at the reporting facility <i>Example:</i> Patient was diagnosed by a staff physician, received radiation at another facility, then underwent surgical resection at the reporting facility
12	Initial diagnosis in staff physician’s office AND all first course treatment or a decision not to treat was done at the reporting facility
13	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.
14	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	
20	Initial diagnosis elsewhere AND all or part of first course treatment was done at the reporting facility, NOS
21	Initial diagnosis elsewhere AND part of first course treatment was done at the reporting facility; part of first course treatment was done elsewhere.
22	Initial diagnosis elsewhere AND all first course treatment was done at the reporting facility or a decision not to treat was done at the reporting facility

Non-Analytic Classes of Case	
Patient Appears In Person At Reporting Facility	
30	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)
31	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)
32	Diagnosis AND all first course treatment provided elsewhere AND patient presents at reporting facility with disease recurrence or persistence (active disease) <i>Example:</i> After treatment failure, the patient was admitted to the facility for supportive care
33	Diagnosis AND all first course treatment provided elsewhere AND patient presents at reporting facility with disease history only (disease not active)
34	Type of case not required to be accessioned (for example, a benign colon tumor) AND initial diagnosis AND part or all of first course treatment by reporting facility <i>Note:</i> Registry hospitals will assign this Class of Case to patients diagnosed with VAIN III, VIN III or AIN III. Non-registry hospitals will assign these cases to the appropriate class of case other than 34 or 36. Class of Case 34 and 36 will not be used by non-registry hospitals.
35	Case diagnosed before program's Reference Date AND initial diagnosis AND all or part of first course treatment by reporting facility
36	Type of case not required to be accessioned (for example, a benign colon tumor) AND initial diagnosis elsewhere AND all or part of first course treatment by reporting facility <i>Note:</i> Registry hospitals will assign this Class of Case to patients diagnosed with VAIN III, VIN III or AIN III. Non-registry hospitals will assign these cases to the appropriate class of case other than 34 or 36. Class of Case 34 and 36 will not be used by non-registry hospitals.
37	Case diagnosed before program's Reference Date AND initial diagnosis elsewhere AND all or part of first course treatment by facility
38	Initial diagnosis established by autopsy at the reporting facility, cancer not suspected prior to

	death
Patient Does Not Appear In Person At Reporting Institution	
40	Diagnosis AND all first course treatment given at the same staff physician’s office
41	Diagnosis and all first course treatment given in two or more different staff physician offices
42	Nonstaff physician or 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) <i>Example:</i> Patients from an unaffiliated, free-standing clinic across the street that hospital abstracts with its cases because many physicians work both at the clinic and the hospital.
43	Pathology or other lab specimens only
49	Death certificate only
99	Nonanalytic case of unknown relationship to facility (not for use for analytic cases).

Recording Class of Case:

1. A Staff Physician (codes 10-12, 41) is a physician who is employed by the reporting facility, under contract with it, or a physician who has routine practice privileges there.
2. Ownership of Physician Practice- If the hospital has purchased a physician practice, it will be necessary to determine whether the practice is now legally considered part of the hospital (their activity is coded as the hospital’s) or not. If the practice is not legally part of the hospital, it will be necessary to determine whether the physicians involved are staff physicians or not, as with any other physician.

Special Instructions

Non-registry hospitals use January 1, 1985 as the reference date. (See *Part One, Reference Date*)

TYPE OF REPORTING SOURCE

Record the code identifying the source documents used to abstract the majority of information on the condition being reported. This may be different than the source used for the original casefinding.

Codes and Definitions

Code	Definition	Source Document	Priority
1	Hospital inpatient , Managed health plans with comprehensive, unified medical records	Hospital inpatient Offices/facilities with unit record <ul style="list-style-type: none"> • HMO physician office or group • HMO affiliated free-standing laboratory, surgery, radiation or oncology clinic Includes outpatient services of HMOs and large multi-specialty physician group practices with unit records.	1
2	Radiation Treatment Centers or Medical Oncology Centers hospital-affiliated or independent	Facilities with serial record (not a unit record) <ul style="list-style-type: none"> • Radiation treatment centers • Medical oncology centers (hospital affiliated or independent) There were no source documents from code 1	2
3	Laboratory only hospital or private, POP	Laboratory with serial record (not a unit record) There were no source documents from codes 1, 2, 8, or 4.	5
4	Physician office/ private medical practitioner	Physician’s office that is NOT an HMO or large multi-specialty physician group practice. There were no source documents from codes 1, 2 or 8.	4
5	Nursing home , convalescent home, convalescent hospital, hospice	Nursing or convalescent home or a hospice There were no source documents from codes 1, 2, 8, 4, or 3.	6
6	Autopsy Only	Autopsy The cancer was first diagnosed on autopsy. There are no source documents from codes 1, 2, 8, 4, 3, or 5.	7
7	Death certificate only (PCR use only)	Death certificate Death certificate is the only source of information; follow-back activities did not identify source documents from codes 1, 2, 8, 4, 3, 5 or 6. If another source document is subsequently identified, the Type of Reporting Source code must be changed to the appropriate code in the range of 1, 2, 8, 4, 3 or 6.	8
8	Other hospital outpatient units/ surgery centers	Other hospital outpatient units/surgery centers. Includes, but not limited to, outpatient surgery and nuclear medicine services. There are no source documents from codes 1 or 2.	3

Definitions

1. Comprehensive, unified medical record- A hospital or managed health care system that maintains a single record for each patient. That record includes all encounters in affiliated locations.
2. Stand-alone medical record- An independent facility; a facility that is not part of a hospital or managed care system. An independent medical record containing only information from encounters with that specific facility.
3. Managed health plan- Any facility where all of the diagnostic and treatment information is maintained in one unit record. The abstractor is able to use the unit record when abstracting the case.

Example: HMOs or other health plan such as Kaiser, Veterans Administration, and military facilities

4. Physician office- A physician office performs examinations and tests. Some physician offices may perform limited surgical procedures.

Note: The category “physician’s office” also includes facilities called surgery centers when those facilities cannot perform surgical procedures under general anesthesia.

5. Serial record- The office or facility stores information separately for each patient encounter (has a separate record for each encounter).
6. Surgery center- Surgery centers are equipped and staffed to perform surgical procedures under general anesthesia. The patient does not stay overnight.

Note If the facility cannot perform surgical procedures under general anesthesia, code as physician’s office.

5. Unit record: The office or facility stores information for all of a patient’s encounters in one record with one record number.

Recording Type of Reporting Source

Priority Order- Assign codes in the following priority: 1, 2, 8, 4, 3, 5, 6, 7

Example: The patient was first found through your pathology department as a private outpatient specimen (Code 3). The patient was admitted as an inpatient to your hospital a month later for surgery. The inpatient record is used for abstracting (Code 1). Code this data item to 1.

DATE OF 1ST CONTACT

Record the date of the first contact with your facility for diagnosis and/or first course treatment of this reportable condition.

Recording Date of 1st Contact

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630

2. **Outpatient Visit- Date of 1st Contact** may be an outpatient visit for a biopsy, x-ray or laboratory test or the date of a pathology specimen was collected at the facility.
3. **Report Actual Date Only. Blank or approximation of month, day, century, or year is not acceptable when reporting this data item to the PCR. Fictitious dates or default values are also not acceptable.**
4. **Never Leave Blank**- This data item can never be blank.
5. **Analytic Cases**- The *Date of 1st Contact* is the date the patient qualifies as an analytic case Class of Case 00-22. Usually, the *Date of 1st Contact* is the date of admission for diagnosis or for treatment. If the patient was initially diagnosed at the facility and went elsewhere for treatment (Class of Case 00), but then returned for treatment that was initially expected to occur elsewhere, the Class of Case is updated to 13 or 14 but the *Date of 1st Contact* is not changed because it still represents the date the patient became analytic.
6. **Nonanalytic Cases**- The *Date of 1st Contact* is the date the patient's nonanalytic status begins with respect to the cancer. For example, for a patient receiving supportive care (Class of Case 32), the date the patient received the supportive care is the *Date of 1st Contact*.
7. **Non-Analytic Case Changes to Analytic**- If the Class of Case changes from nonanalytic (for example, staging workup after initial diagnosis, Class of Case 30) to analytic (for example, part of first course treatment administered at the facility, Class of Case 21), the *Date of 1st Contact* is updated to the date the case became analytic (the date the patient was admitted for treatment).
8. **Autopsy Only**- If an autopsy-only case, use date of death.
9. **Readmission for a Newly Reportable Condition**- When a patient is readmitted on or after January 1, 2001 for a condition not previously required to be reported, the *Date of 1st Contact* is the earliest date on or after January 1, 2001 the patient was seen at the reporting institution. (Registry hospitals see also *Special Instructions*.)

Example: The patient was diagnosed with a colon primary in 1980. The patient came to the reporting facility in 1999 for treatment. The case was not reportable in 1999 due to the date of diagnosis being prior to January 1, 1990. The patient returns to the reporting facility for cancer treatment on March 23, 2003. The case is now reportable with a *Date of 1st Contact* of 20030323.

10. Earlier than Date of Inpatient Discharge- *Date of 1st Contact* must be earlier than the date of inpatient discharge.

Example 1: The patient is seen as an outpatient for a colonoscopy with a positive biopsy on October 6, 2010. He is admitted to the hospital from October 10, 2010 to October 15, 2010 for surgery. The *Date of 1st Contact* is 20101006.

Example 2: The patient is admitted from August 23, 2010 to August 27, 2010 for shortness of breath. On August 25, 2010 the patient has a lung biopsy, which is diagnostic of cancer. The *Date of 1st Contact* is 20100825.

11. Admission Unrelated to Cancer- If a patient is admitted for other reasons not related to cancer; use the diagnosis date as the *Date of 1st Contact*.

Example: Patient is admitted for a reason unrelated to cancer on 1/15/2010 and 1/17/2010 is incidentally diagnosed with cancer, the *Date of 1st Contact* is 20100117.

12. For Private Outpatient (POP) cases record the date the specimen was taken. If a patient was first identified as a POP and comes to your facility as an inpatient or outpatient during the three month holding period (See *Part One, Private Outpatient Specimens*) for further diagnosis or treatment, the *Date of 1st Contact* is the date of the patient's first in-person contact with your facility.

Example: Patient undergoes a biopsy in a physician's office on September 8, 2010. The pathology specimen was sent to your facility and was read as malignant melanoma. The patient enters your facility on September 14, 2010 for a wide excision. The *Date of 1st Contact* is 20100914.

13. Positive Imaging Study- Hospitals are not expected to report cases on the basis of a positive imaging study **only**. However, if the patient meets reporting requirements at a later time, the case must be reported using the date of the positive imaging study as the *Date of 1st Contact*.

Example: The patient has an outpatient mammogram on April 10, 2010 that is suspicious for cancer. The patient returns for a biopsy which is diagnostic of cancer on April 17, 2010. This case would be reportable at the time of the biopsy with a *Date of 1st Contact* of 20100410.

Special Instructions

Registry Hospitals - When a patient is readmitted on or after January 1, 2001 for a cancer not previously reportable to the PCR, but had been abstracted and included in your registry at the time of diagnosis, report the case to the PCR as it was originally abstracted. The *Date of 1st Contact* should not be updated to the current encounter date. These old cases will **not** be included in your hospital's timeliness calculations.

Completion of *Date of 1st Contact*, *Date of Inpatient Adm* and *Date of Inpatient Disch* and Date Flags

Patient Type	Date of 1st Contact	Date of Inpatient Admission	Date of Inpatient Discharge	Date of 1st Contact Flag	Date of Inpatient Admission Flag	Date of Inpatient Discharge Flag
Outpatient Only	Date of Outpatient Visit	bbbbbbbb	bbbbbbbb	bb	11	11
Inpatient Only	Date of Inpatient Adm or Date of Diagnosis	Date of Inpatient Adm	Date of Inpatient Disch	bb	bb	bb
Outpatient then Inpatient	Date of Outpatient Visit	Date of Inpatient Adm	Date of Inpatient Disch	bb	bb	bb
General Instructions	Must always contain date; Can never be blank.	Must contain date if patient was inpatient;	Must contain date if patient was inpatient;	Will always be left blank.	Must contain a flag if the patient was never an inpatient	Must contain a flag if the patient was never an inpatient

b= blank

DATE OF 1st CONTACT FLAG

The PCR requires *Date of 1st Contact* to be entered on all cases; therefore this field will always be left blank.

Codes and Definitions

Code	Definition
(blank)	A valid date value is provided in item <i>Date of 1st Contact</i> .

DATE OF INPATIENT ADM

Record the date of the inpatient admission to the facility for the most definitive surgery.

Recording Date of Inpatient Adm

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630

2. **No Surgery**- If the patient does not have surgery, use the Date of Inpatient Admission for any other cancer-directed therapy.
3. **No Cancer-Directed Therapy**- If the patient has no cancer-directed therapy, use the Date of Inpatient Admission for diagnostic evaluation.
4. **Never Admitted as an Inpatient**- If the patient was never admitted as an inpatient for this reportable condition, leave *Date of Inpatient Adm* blank.
5. **Most Definitive Treatment**- Use the Date of Inpatient Admission from the admission the patient was seen at the reporting institution for the most definitive treatment.

Example: The patient is admitted to the reporting institution from February 9, 2010 to February 11, 2010 for a modified radical mastectomy. Date of Inpatient Adm is 20100209.

6. **Report Actual Date Only.** Approximation of month, day, century, or year is not acceptable when reporting to the PCR. Fictitious dates or default values are also not acceptable.

DATE OF INPT ADM FLAG

Record the date flag in the event a complete Date of Inpatient Admission was not entered to explain why.

Recording Date of Inpt Adm Flag

1. Date Recorded- Leave this field blank if *Date of Inpt Adm* has a date recorded.
2. Not an Inpatient- Code 11 if the patient was never an inpatient.
3. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
11	No proper value is applicable in this context (e.g., patient was never an inpatient at the reporting facility)
(blank)	A valid date value is provided in item Date of Inpatient Admission

The following table illustrates the use of the date flag and the traditional and interoperable date formats for coding *Date of Inpatient Adm* and *Date of Inpt Adm Flag*. In the table below, the lowercase letter “b” is used to represent each blank space.

Description	Traditional Date of Inpatient Admission (MMDDCCYY)	Interoperable Date of Inpatient Admission (CCYYMMDD)	Date of Inpt Adm Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Never an inpatient	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11

b=blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

DATE OF INPATIENT DISCH

Record the date of the inpatient discharge from the facility for the most definitive surgery.

Recording Date of Inpatient Disch

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630

2. **No Surgery**- If the patient does not have surgery, use the Date of Inpatient Discharge for any other cancer-directed therapy.
3. **No Cancer-Directed Therapy**- If the patient has no cancer-directed therapy, use the Date of Inpatient Discharge for diagnostic evaluation.
4. **Never Admitted as an Inpatient**- If the patient was never admitted as an inpatient for this reportable condition, leave *Date of Inpatient Disch* blank.
5. **Most Definitive Treatment**- Use the Date of Inpatient Discharge from the admission the patient was seen at the reporting institution for the most definitive treatment.

Example: The patient is admitted to the reporting institution from Feb. 9, 2010 to Feb. 11, 2010 for a modified radical mastectomy. *Date of Inpatient Disch* is 20100211.

6. **Report Actual Date Only.** Approximation of month, day, century, or year is not acceptable when reporting to the PCR. Fictitious dates or default values are also not acceptable.

DATE OF INPT DISCH FLAG

Record the date flag in the event a complete Date of Inpatient Discharge was not entered to explain why.

Recording Date of Inpt Disch Flag

1. Date Recorded- Leave this field blank if *Date of Inpt Disch* has a date recorded.
2. Not an Inpatient- Code 11 if the patient was never an inpatient.
3. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
11	No proper value is applicable in this context (e.g., patient was never an inpatient at the reporting facility)
(blank)	A valid date value is provided in item Date of Inpatient Discharge

The following table illustrates the use of the date flag and the traditional and interoperable date formats for coding *Date of Inpatient Disch* and *Date of Inpt Disch Flag*. In the table below, the lowercase letter “b” is used to represent each blank space.

Description	Traditional Date of Disch (MMDDCCYY)	Interoperable Date of Disch (CCYYMMDD)	Date of Disch Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Never an inpatient	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11

b=blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

INSTITUTION REFERRED FROM

Record the facility that referred the patient to the reporting institution. The codes used to identify facilities in the PCR database are the Facility Identification Numbers (FIN) assigned by the American College of Surgeons Commission on Cancer (ACOS COC).

Recording Institution Referred From

1. Do not leave blank- This data item may not be blank.
2. Own Facility- Do not enter your own facility ID number.
3. Not Referred - Code 0000000000 if the patient was not referred to the reporting institution from another institution. *Institution Referred From* for Private Outpatient Specimens (POP) is coded to 0000000000.
4. Referring Facility - Use the codes in your current selection of facility ID numbers (FIN), i.e., Commission on Cancer (COC) codes, to identify the facility from which the patient was referred. This field is 10 characters. For facilities with seven-digit FINs assigned by the COC before January 1, 2001, the coded FIN will consist of three leading zeros followed by the full seven-digit number, e.g., 0006231234. For facilities with eight-digit FINs assigned by COC after January 1, 2001, the coded FIN will consist of two leading zeroes, followed by the full eight-digit number, e.g., 0010000000. *Note:* Most abstracting software including Abstract Plus will automatically zero-fill these remaining spaces when the selection is made from a table of facility codes.
5. Referred But Facility ID Number Unknown - Code 0099999999 if the patient was referred but the referring institution's ID number is unknown.
6. List of FINs- A complete list of FINs is available on the American College of Surgeons web site at <https://datalinks.facs.org/CPM/ACSHosp.cfm>

PHYSICIAN - FOLLOW-UP

Record the number used by your facility to uniquely identify the physician currently responsible for the patient's medical care.

Recording Physician-Follow-up

1. Do Not Leave Blank- This data item may not be blank.
2. Format- The identification number may include numbers, letters (8 characters or less) and embedded spaces, but may not include any punctuation.
3. If No Numbers are Assigned- When identification number is not assigned, enter up to eight characters of the physician's last name.
4. If Following Physician is Unknown- Enter 99999999 if the following physician is unknown.

DATE OF DIAGNOSIS

Record the date this reportable condition was first diagnosed by a recognized medical practitioner.

Note: In the examples below the lowercase letter “b” is used to represent each blank space.

Recording Date of Diagnosis

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630

2. **Date Reportable**- Use the earliest date of diagnosis whether clinically or histologically confirmed.

Example 1: The patient was diagnosed with stromal endometriosis August 24, 2010. The patient presents to the reporting institution for treatment of the stromal endometriosis on November 5, 2010. This case would be reportable with a *Date of Diagnosis* of 20100824.

Example 2: The patient has a history of breast cancer diagnosed September 10, 2000. The patient now presents to the reporting institution with metastasis from the breast. This case would be reportable with a *Date of Diagnosis* of 20000910.

3. **Exact Date Unavailable**- If the exact date is not available, refer to *Part Three: Data Item Instructions, General Information-Dates* for instructions regarding approximating dates and unknown dates. For analytic cases, at least the year of diagnosis should be known or be able to be approximated (month and day may be unknown).

Example: Patient is admitted on January 15, 2010 with severe flank pain with history of lung cancer diagnosed five years ago. The correct *Date of Diagnosis* is 2005bbbb. **Do not** leave entire date blank when descriptive information can be used to approximate the year.

4. **Partial Dates**- If only the month and/or year is known or can be approximated record what is available and leave the remaining parts of the date blank. No approximation of day is acceptable.

Example: Documentation in the patient's record from a June 2010 admission indicates the patient was diagnosed 'last year'. The correct *Date of Diagnosis* is 2009bbbb.

5. **Unknown Date**- If the date of diagnosis cannot be identified and is truly unknown, the case is not reportable. See *Part One, Date of Diagnosis Reportability*.

6. **Default Values**- Fictitious dates or default values are not acceptable to be entered for month, day, or year.

Example: Patient receives palliative treatment for breast cancer diagnosed in June 2005. The correct *Date of Diagnosis* is 200506bb. **Do not** record 20050615 where 15 is a default value for day.

7. Clinical Diagnosis- Use the date of diagnosis whether clinically or histologically confirmed. A clinical diagnosis often includes ambiguous terminology. See *Part One, Reportable Conditions* for a list of terms that constitute a diagnosis. Do not change the *Date of Diagnosis* when a later biopsy or cytology provides confirmation of a clinical diagnosis.

Example 1: A March 12, 2010 mammogram reveals a mass in the upper-outer quadrant of a patient's right breast compatible with carcinoma. On March 20, 2010, the patient has an excisional breast biopsy that confirms infiltrating ductal carcinoma. *Date of Diagnosis* is 20100312.

Example 2: A physician notes a prostate nodule possible for cancer during a May 12, 2010 physical exam. On June 15, 2010 a needle biopsy of the prostate histologically confirms adenocarcinoma. *Date of Diagnosis* is 20100615 because "possible for cancer" does not constitute a reportable diagnosis.

8. Earlier Date- If the physician states, in retrospect, the patient had a reportable condition at an earlier date, use the earlier date as the date of diagnosis.

Example: A patient has a total abdominal hysterectomy for endometriosis in January 2008. The patient is admitted to the hospital with abdominal pain in November 2010. An omental biopsy shows metastatic cystadenocarcinoma. Pathologists re-review the 2008 hysterectomy specimen. They identify an area of cystadenocarcinoma in the left ovary. *Date of Diagnosis* is 200801bb.

9. Non-reportable Conditions that Transform into a Reportable Condition- If a patient is diagnosed with a non-reportable condition that later transforms into a reportable condition, record the date the patient was diagnosed with the reportable condition.

Example: The patient was diagnosed with myelodysplastic syndrome on May 1, 2000 (not reportable until 2001) and it transforms into acute myelogenous leukemia on June 15, 2002. Abstract the case as acute myelogenous leukemia with a *Date of Diagnosis* of 20020615.

10. Diagnosed at Autopsy- The date of death is the date of diagnosis for a case diagnosed at autopsy.
11. Treatment Before a Definitive Diagnosis- Use the date therapy was started as the date of diagnosis if the patient receives cancer-directed treatment before a definitive diagnosis.
12. Suspicious Cytology Only is not diagnostic of cancer. Use the date of clinical, histologic, or positive cytologic confirmation as the date of diagnosis.
13. Diagnosed In Utero- Use the actual date of diagnosis for an *in utero* diagnosis, for cases diagnosed on or after January 1, 2009. For cases diagnosed prior to January 1, 2009 use the date of birth as the date of diagnosis.
14. Positive Tumor Markers alone are not diagnostic of cancer. Use the date of clinical, histologic, or positive cytologic confirmation as the date of diagnosis.

Example 1: The patient has an elevated PSA and the physical examination is negative. The physician documents only that the patient is referred for a needle biopsy of the prostate. The biopsy is positive for adenocarcinoma. The date of diagnosis is the date of the biopsy.

Example 2: The patient has an elevated PSA and the physical examination is negative. The physician documents that he/she suspects the patient has prostatic cancer and is referring the patient for a needle biopsy. The needle biopsy is positive. The date of diagnosis is the date the physician documented that he/she suspects the patient has prostatic cancer.

DATE OF DIAGNOSIS FLAG

The PCR on requires cases with a full or partial *Date of Diagnosis* to be reported on all cases; therefore this field will always be left blank.

Codes and Definitions

Code	Definition
(blank)	A valid date value is provided in item <i>Date of Diagnosis</i> .

PRIMARY SITE

This data item records the topography code for the primary site of the cancer/tumor condition being reported using ICD-O-3 or ICD-O-2 (*International Classification of Diseases for Oncology, Third or Second Edition* published by the World Health Organization).

1. Cases Diagnosed on or after January 1, 2001 - Code according to ICD-O-3.
2. Cases Diagnosed prior to January 1, 2001 - Code according to ICD-O-2.
3. ICD-O-3 Errata – All errata contained in *Appendix E* must be used when coding *Primary Site, Histology, Behavior and Grade*. It is recommended these errata be added to the ICD-O-3 coding book being used.

Coding Primary Site

1. Use all information available in the medical record for determining primary site. Operative reports, oncology consults and pathology reports will help in determining the correct primary site. If you cannot make this determination, consult a physician.
2. Be Specific- When determining the primary site, be as specific as possible. Many organs can be divided into specific segments or tissue types. It is important to code the exact segment or tissue involved.

Example 1: The wrist contains several tissue types; skin (C44.6), bone (C40.1), soft tissue (C49.1)

Example 2: The large intestine is divided as follows:

cecum (C18.0)	splenic flexure of colon (C18.5)
ascending colon (C18.2)	descending colon (C18.6)
hepatic flexure of colon (C18.3)	sigmoid colon (C18.7)
transverse colon (C18.4)	rectosigmoid colon (C19.9)

3. Primary Tumor Originated- Code the site in which the primary tumor originated, even if it extends onto/into an adjacent subsite.

Example 1: Final diagnosis is adenocarcinoma of the upper lobe of the right lung. Code the topography to lung, upper lobe (C341).

Example 2: Patient has a right brachial cleft cyst; the pathology report identifies an adenocarcinoma arising in an ectopic focus of thyroid tissue within the brachial cleft cyst. Thyroidectomy pathology is negative. Code the primary site to brachial cleft (C104).

Example 3: The patient had a total hysterectomy with a bilateral salpingo-oophorectomy ten years ago for non cancer reasons. She now has widespread cystadenocarcinoma in the peritoneum. Code the primary site to peritoneum, NOS (C482).

Example 4: The patient has a 4 cm tumor in the right breast. The tumor originated in the upper inner quadrant and extends into the lower inner quadrant. Code the primary site to upper inner quadrant of breast (C502).

4. Overlapping Tumor- Code the last digit of the primary site code to .8 when a **single tumor overlaps** an adjacent subsite(s) of an organ and the point of origin cannot be determined.

Example: The patient has a 5cm tumor that involves the dorsal surface and anterior 2/3 of tongue. Code the primary site to C028 (overlapping lesion of tongue).

5. Multiple Tumors/Different Subsites- Code the last digit of the primary site code 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.

Example 1: During a TURB, the physician describes multiple papillary tumors in the bladder neck (C675) and the lateral wall of the bladder (C672). Code the primary site as bladder, NOS (C679).

Example 2: Patient has an infiltrating duct tumor in the upper outer quadrant (C504) of the right breast and another infiltrating duct carcinoma in the lower inner (C503) quadrant of the right breast. Code the primary site as breast, NOS (C509).

6. Related Site Code- Some histology/behavior terms in ICD-O-3 have a related site code in parenthesis; for example: hepatoma (C220).

- a. Code the site as documented in the medical record and ignore the suggested ICD-O-3 code when a primary site is specified in the medical record.

Example: The pathology report says “ductal carcinoma of the head of the pancreas.” The listing in ICD-O-3 is ductal carcinoma 8500/3 (C50_). Code the primary site to head of pancreas (C250), NOT to breast (C50_) as suggested by the ICD-O-3.

- b. Use the site code suggested by ICD-O-3 when the primary site is the same as the site code suggested or the primary site is unknown

Example 1: The biopsy is positive for hepatoma, but there is no information available about the primary site. Code the primary site to liver (C220) as suggested by ICD-O-3.

Example 2: The patient has an excision of the right axillary nodes which reveals metastatic infiltrating duct carcinoma. The right breast is negative. The ICD-O-3 shows duct carcinoma (8500) with a suggested site of breast (C50_). Code the primary site as breast, NOS (C509).

7. Hematopoietic and Lymphoid Neoplasms-

- a. Cases diagnosed on or after January 1, 2010- Effective with cases diagnosed on or after January 1, 2010 the PCR requires the use of *The Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* and *The Hematopoietic Database* for primary site coding of hematopoietic and lymphoid neoplasms. The manual and database can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

- b. Cases diagnosed prior to January 1, 2010- Apply the following rules:

- i. Leukemia- Code leukemia primaries to bone marrow (C421); blood cells originate in the bone marrow.
- ii. Lymphoma Definitions-

- 1) Extra lymphatic: Originating in tissue or an organ that is not a part of the lymphatic system.
- 2) Extranodal lymphoma: Lymphoma originating in tissue or organ other than lymph nodes. Lymphatic system organs may be extranodal. (e.g.: Spleen is a lymphatic system organ and is also extranodal.)
- 3) Lymphatic system: An umbrella term that includes: lymph nodes, spleen, thymus, tonsils, Waldeyer's ring, and Peyer's patches.
- 4) Nodal lymphoma: A lymphoma originating in lymph nodes

iii. Lymphoma Coding Instructions

- 1) Single Lymph Node Chain- When a single lymph node chain is involved, code that chain as the primary site.
- 2) Multiple Lymph Node Chains- When multiple lymph node chains are involved at the time of diagnosis, do not simply code the lymph node chain that was biopsied.
 - a. If it is possible to determine where the disease originated, code the primary site to that lymph node chain.
 - b. If multiple lymph node chains are involved and all involved chains are located in the same lymph node region (i.e. the same primary site code) and it is not possible to determine the lymph node chain where the disease originated, code the primary site to lymph nodes of the specified nodal region (C77_).
 - c. If multiple lymph node chains are involved and the involved chains are in different lymph node regions, code C778 (lymph nodes of multiple regions).
- 3) Extranodal- When the lymphoma is extranodal and is
 - a. Confined to the organ of origin, code the organ of origin.

Example: Pathology from a stomach resection shows lymphoma. No other pathologic or clinical disease identified. Code the primary site as stomach, NOS (C169).
 - b. Present in an extranodal organ/site and in that organ/site's regional lymph nodes code the extranodal organ/site as the primary site. Lymphomas that are primary in an extranodal organ/site may metastasize to that organ/site's regional lymph nodes. In rare cases a lymphoma may spread from the lymph node to an extranodal site or extra lymphatic organ by direct extension.

Example 1: Lymphoma is present in the spleen and splenic lymph nodes. Code the primary site to spleen (C422).

Example 2: Lymphoma is present in the stomach and the gastric lymph nodes. Code the primary site to stomach, NOS (C169).

- c. Present in extranodal organ(s)/site and non-regional lymph nodes, consult the physician to determine the primary site. If a site cannot be determined, code primary site to Lymph Node, NOS (C779).

4) Unknown Primary- If the **primary site is unknown** or not given:

- a. Code retroperitoneal lymph nodes if described as retroperitoneal mass (772)
- b. Code inguinal lymph nodes if described as inguinal mass (C774)
- c. Code mediastinal lymph nodes if described as mediastinal mass (C771)
- d. Code mesenteric lymph nodes if described as mesenteric mass (C772)
- e. If the primary site is unknown code Lymph Nodes, NOS (C779)

Exception: Code unknown primary site (C809) only when there is no evidence of lymphoma in lymph nodes and/or the medical record documents that the physician suspects that it is an extranodal lymphoma

- 8. Melanoma - If a patient is diagnosed with metastatic melanoma and the primary site is not identified, the primary site is *skin NOS* (C44.9).
- 9. Kaposi Sarcoma - Kaposi sarcoma is a rare condition. When not AIDS-related, it usually presents as localized disease with an easily recognized primary site. AIDS-related Kaposi sarcoma usually presents as a disseminated disease with involvement of mucosal surfaces, visceral surfaces of organs, and skin. It is important to review consecutive records carefully to determine the extent of involvement at diagnosis. Review of a single record may reveal only the site being treated during that admission.
 - a. Code the Kaposi sarcoma to the primary site in which it arises.
 - b. If the Kaposi sarcoma is present in the skin and another site simultaneously, code to the specified skin site, (C44_).
 - c. If the primary site is unknown or cannot be determined, code skin, NOS (C449).
- 10. Sarcoma- The majority of sarcomas arise in mesenchymal or connective tissues that are located in the musculoskeletal system. The musculoskeletal system includes the fat, muscles, blood vessels, deep skin tissues, nerves, bones and cartilage. The default code for sarcomas of unknown primary site is C499 rather than C809.

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.

Example: The pathology identifies a mixed Mullerian tumor of the uterus. Code the primary site to uterus, NOS (C559)

11. Waldenstrom Macroglobulinemia - The primary site is *blood* (C42.0).
12. Unknown - When the primary site is not known, record as described below. Do not record a metastatic site when the primary is not known.
 - a. Osteosarcoma with unknown primary, record primary site as *bone NOS* (C41.9)
 - b. Other histologies with unknown primary, record primary site as *Unknown* (C80.9)

Text

Text to support this data item must be recorded in the specific text field. See *Part Three, Data Item Instructions, Text-Primary Site Title*. This text field is used by the PCR to validate the ICD-O topography and laterality codes reported.

LATERALITY

Record appropriate laterality code. Laterality describes the side of a paired organ or side of the body on which the reportable tumor originated. For each primary you need to determine whether laterality should be coded

Codes and Definitions

Code	Definition
0	Not a paired organ
1	Right: origin of primary
2	Left: origin of primary
3	Only one side involved, right or left origin unspecified
4	Bilateral involvement, side of origin unknown, stated to be a single primary
5	Paired site: midline tumor
9	Paired site, but no information concerning laterality

Recording Laterality

1. Unknown Primary Site- Record laterality for unknown primary site (C80.9) as 0 (not a paired site).
2. Metastatic Sites- Do not code laterality of metastatic sites.
3. Listing of Paired Sites-
 - a. Use codes 1-9 for the sites listed on the following page, except as noted.
 - b. Major categories- The listing includes major categories. Code laterality for all subheadings included in *ICD-O* under these headings, unless specifically excluded.
 - c. Exclusions should be coded to "0."
4. Single Side, Laterality Unknown- Assign code 3 if the laterality is not known but the tumor is confined to a single side of a paired organ.

Example: Pathology report: Patient has a 2 cm carcinoma in the upper pole of the kidney. Code laterality as 3 because there is documentation that the disease exists in only one kidney, but it is unknown if the disease originated in the right or left kidney.

5. Code 4 is seldom used **except** for the following diseases:

- Both ovaries involved simultaneously, single histology
- Bilateral retinoblastomas
- Bilateral Wilms tumors

6. Midline tumors- Where the right and left sides of paired sites are contiguous (come into contact) and the lesion is at the point of contact of the right and left sides, use code 5, midline.

Note: Most paired sites cannot develop midline tumors. Skin of the trunk can have a midline tumor, but breast cannot. “Midline of the right breast” is coded 1, right; midline in this usage indicates the primary site is C50.8 (overlapping sites).

7. Laterality Unknown- Assign code 9 when the disease originated in a paired site, but the laterality is unknown.

Example: Admitting history says patient was diagnosed with lung cancer based on positive sputum cytology. Patient is treated for painful bony metastases. There is no information about laterality in the diagnosis of this lung cancer, code to 9.

Paired Organ Sites

PAIRED SITE	ICD-O CODE
Acoustic nerve (excluding diagnoses prior to 2004)*	C72.4
Adrenal gland	C74.0-C74.9
Breast	C50.0-C50.9
Carotid body	C75.4
Cerebral meninges, NOS (excluding diagnoses prior to 2004)*	C70.0
Cerebrum (excluding diagnoses prior to 2004)*	C71.0
Connective, subcutaneous, and other soft tissues of lower limb and hip	C49.2
Connective, subcutaneous, and other soft tissues of upper limb and shoulder	C49.1
Cranial nerve, NOS (excluding diagnoses prior to 2004)*	C72.5
Epididymis	C63.0
Eye and lacrimal gland	C69.0-C69.9
Fallopian tube	C57.0
Frontal lobe (excluding diagnoses prior to 2004)*	C71.1
Frontal sinus	C31.2
Kidney, NOS	C64.9
Long bones of lower limb	C40.2
Long bones of upper limb and scapula	C40.0
Lung	C34.1-C34.9

PAIRED SITE	ICD-O CODE
Main bronchus (excluding carina, code to “0”)	C34.0
Maxillary sinus	C31.0
Middle ear	C30.1
Nasal cavity (excluding nasal cartilage and nasal septum, code to “0”)	C30.0
Occipital lobe (excluding diagnoses prior to 2004)*	C71.4
Olfactory nerve (excluding diagnoses prior to 2004)*	C72.2
Optic nerve (excluding diagnoses prior to 2004)*	C72.3
Ovary	C56.9
Parietal lobe (excluding diagnoses prior to 2004)*	C71.3
Parotid gland	C07.9
Pelvic bones (excluding sacrum, coccyx, and symphysis, code to “0”)	C41.4
Peripheral nerves and autonomic nervous system of lower limb and hip	C47.2
Peripheral nerves and autonomic nervous system of upper limb and shoulder	C47.1
Pleura	C38.4
Renal Pelvis	C65.9
Rib and clavicle (excluding sternum, code to “0”)	C41.3
Short bones of lower limb	C40.3
Short bones of upper limb	C40.1
Skin of eyelid	C44.1
Skin of external ear	C44.2
Skin of lower limb and hip	C44.7
Skin of other and unspecified parts of face (midline, code to “5”)	C44.3
Skin of trunk (midline, code to “5”)	C44.5
Skin of upper limb and shoulder	C44.6
Spermatic cord	C63.1
Sublingual gland	C08.1
Submandibular gland	C08.0
Temporal lobe (excluding diagnoses prior to 2004)*	C71.2
Testis	C62.0-C62.9
Tonsillar fossa	C09.0
Tonsillar pillar	C09.1
Tonsil, NOS	C09.9
Tonsil, Overlapping	C09.8
Ureter	C66.9

*For cases diagnosed prior to January 1, 2004 these sites are considered non-paired and should be coded to 0.

HISTOLOGIC TYPE

Record the code for histologic type of the cancer/tumor being reported using ICD-O-3 or ICD-O-2 (*International Classification of Diseases for Oncology, Third or Second Edition* published by the World Health Organization).

1. Cases Diagnosed on or after January 1, 2001 - Code according to ICD-O-3.
2. Cases Diagnosed prior to January 1, 2001 - Code according to ICD-O-2.
3. ICD-O-3 Errata – All errata contained in *Appendix E* must be used when coding *Primary Site, Histology, Behavior and Grade*. It is recommended these errata be added to the ICD-O-3 coding book being used.

Coding Histology

1. Solid tumors diagnosed on or after January 1, 2007- Effective with cases diagnosed on or after January 1, 2007 the PCR requires the use of *The Multiple Primary and Histology Coding Rules (MP/H Rules)* for coding histology coding of solid malignant tumors and benign and borderline Central Nervous System tumors. The *MP/H Rules Manual* can be downloaded from the following site:

<http://seer.cancer.gov/tools/mphrules/download.html>

2. Solid tumors diagnosed before January 1, 2007- Refer to *SEER Program Coding and Staging Manual 2004* at the following link for guidelines to code histology of solid malignant tumors and benign and borderline Central Nervous System tumors diagnosed prior to January 1, 2007.

http://seer.cancer.gov/archive/manuals/2004Revision1/SPM_2004_maindoc.r1.pdf

3. Hematopoietic and lymphoid neoplasms diagnosed on or after January 1, 2010- Effective with cases diagnosed on or after January 1, 2010 the PCR requires the use of *The Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* and *The Hematopoietic Database* for histology coding of hematopoietic and lymphoid neoplasms. The manual and database can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

4. Hematopoietic and lymphoid neoplasms diagnosed before January 1, 2010- Refer to *SEER Program Coding and Staging Manual 2007* at the following link for guidelines to code histology of hematopoietic and lymphoid neoplasms diagnosed prior to January 1, 2010.

http://seer.cancer.gov/archive/manuals/2007/SPCSM_2007_maindoc.pdf

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions, Text-DX Proc-Path* and *Text-Histology Title*. These text fields are used by the PCR to validate ICD-O histology codes reported.

BEHAVIOR

Record the code for the behavior of the cancer/tumor being reported using ICD-O-3 or ICD-O-2 (*International Classification of Diseases for Oncology, Third or Second Edition* published by the World Health Organization).

1. Cases Diagnosed on or after January 1, 2001 - Code according to ICD-O-3.
2. Cases Diagnosed prior to January 1, 2001- Code according to ICD-O-2.
3. ICD-O-3 Errata – All errata contained in *Appendix E* must be used when coding *Primary Site, Histology, Behavior and Grade*. It is recommended these errata be added to the ICD-O-3 coding book being used.

Coding Behavior

Behavior is part of the diagnosis. The behavior indicates whether a tumor is malignant, benign, in situ, or uncertain whether malignant or benign.

1. Reportable In Situ and Malignant Behaviors- The PCR requires the reporting of /2 (in situ) and /3 (malignant) tumors.
2. Behavior from Metastatic Site- If the only specimen is from a metastatic site, the behavior is malignant.
3. Reportable Benign and Borderline Behaviors- Effective with cases diagnosed on or after **January 1, 2004** primary intracranial and central nervous system tumors with a behavior code of /0 or /1 (benign and borderline or "non-malignant") are reportable regardless of histologic type for the sites listed below.

Reportable Intracranial and Central Nervous System (CNS) Primary Sites Effective January 1, 2004	
Meninges (C70.0 - C70.9)	Other CNS (C72.8, C72.9)
Brain (C71.0 - C71.9)	Pituitary gland (C75.1)
Spinal cord (C72.0)	Craniopharyngeal duct (C75.2)
Cauda equina (C72.1)	Pineal gland (C75.3)
Cranial nerves (C72.2 - C72.5)	

4. Hematopoietic Conditions with Change in Behavior Code- The following table lists hematopoietic conditions that are reportable beginning with cases diagnosed on or after **January 1, 2010**. Based on changes to the hematopoietic and lymphoid neoplasm rules, the behavior of the following diseases has changed from borderline or uncertain (/1) to malignant (/3). Adding the ICD-O-3 code and behavior in the alphabetic and tabular listing of the ICD-O-3 code book will assist in identifying them as reportable conditions and assigning the appropriate code.

Histologic Terms and Codes with Changes in Case Reportability Effective January 1, 2010	
Name	ICD-O-3 Code
Chronic lymphoproliferative disorder of NK-cells	9831/3
T-cell large granular lymphocytic leukemia	9831/3
Langerhans cell histiocytosis, NOS (9751/1)	9751/3
Langerhans cell histiocytosis, unifocal (9752/1)	9751/3
Langerhans cell histiocytosis, multifocal (9753/1)	9751/3
Myelodysplastic/Myeloproliferative neoplasm, unclassifiable	9975/3
Myeloproliferative disease, NOS	9975/3
Myeloproliferative neoplasm, unclassifiable	9975/3

5. In Situ Terminology- The following terms are synonymous with in situ (behavior code 2):

- *Adenocarcinoma in an adenomatous polyp with no invasion of stalk*
- *Bowen’s disease*
- *Clark’s level 1 for melanoma (limited to epithelium)*
- *Comedocarcinoma, noninfiltrating*
- *Confined to epithelium*
- *Hutchinson’s melanotic freckle, NOS*
- *Intracystic, noninfiltrating*
- *Intraductal*
- *Intraepidermal, NOS*
- *Intraepithelial, NOS*
- *Involvement up to but not including the basement membrane*
- *Lentigo maligna*
- *Lobular neoplasia, grade III (LN3)*
- *Lobular, noninfiltrating*
- *Noninfiltrating*
- *Noninvasive*
- *No stromal involvement*
- *Papillary, noninfiltrating or intraductal*
- *Precancerous melanosis*
- *Pre-invasive*
- *Queyrat’s erythroplasia*
- *Stage 0*
- *Vaginal epithelial neoplasia, grade 3 (VAIN III)*
- *Vulvar epithelial neoplasia, grade 3 (VIN III)*

6. Areas of Invasion- Record behavior as /3 (malignant) if any invasion is present, no matter how limited.

Example: The pathology report reads *intraductal carcinoma (8500/2) with focal areas of invasion*. The phrase *with focal areas of invasion* is an important component in determining behavior and impacts the proper ICD-O code assignment. The histologic type must include the invasive component, *intraductal carcinoma with focal areas of invasion (8500/3)*.

7. Severe/High Grade Dysplasia of the Colon and/or Esophagus- If your facility considers the terminology of severe dysplasia or high grade dysplasia of the colon and/or esophagus as synonymous with carcinoma in-situ; use the following guidelines for reporting these cases to the PCR:

- a. Obtain a statement from your pathologists outlining the terminology policy of their department. The statements should be separate, one for colon and one for esophagus, because different physicians may be involved in reviewing the statement for each site.
- b. Submit the statement(s) to the appropriate medical staff committee for approval. Registry hospitals will normally submit the statement(s) to the Cancer Committee.

- c. Document a policy that states the sites diagnosed with severe dysplasia and/or high grade dysplasia that will be abstracted as carcinoma in-situ based on the pathologists approved statements.
- d. Add the policy to your Operations Manual attaching the approved statement(s) from your pathologists.
- e. Forward a copy of the policy and statement(s) to the PCR to keep on permanent file.
- f. Abstract all colon and/or esophagus cases diagnosed with severe dysplasia and/or high grade dysplasia as carcinoma in-situ according to the statement(s) and policy. In the text for each abstract, document the final pathologic diagnosis along with the statement “in-situ per pathologist”.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions, Text-DX Proc-Path* and *Text-Histology Title*. These text fields are used by the PCR to validate ICD-O behavior codes reported.

GRADE

This data item records the code for grade or differentiation of the cancer/tumor being reported.

1. Cases Diagnosed on or after January 1, 2014 – Code grade according the guidelines outlined below. These guidelines are a consensus from COC, SEER and NPCR.
2. Cases Diagnosed prior to January 1, 2014- Code grade according the guidelines in the *SEER Program Coding and Staging Manual 2013*. <http://seer.cancer.gov/tools/codingmanuals/>

Hematopoietic and Lymphoid Neoplasms

Cell Indicator (Codes 5, 6, 7, 8) describes the lineage or phenotype of the cell. Codes 5, 6, 7, and 8 are used only for hematopoietic and lymphoid neoplasms. Code 9 indicates cell type not determined, not stated, or not applicable.

Coding Grade for Hematopoietic and Lymphoid Neoplasms

1. Determine the histology based on the current Hematopoietic and Lymphoid Neoplasm Manual
<http://seer.cancer.gov/seertools/hemelymph/>
2. Determine the Cell Indicator by applying the “Grade of Tumor Rules” within the current Hematopoietic and Lymphoid Neoplasm Manual to code the grade

<http://seer.cancer.gov/seertools/hemelymph/>

Grade codes for hematopoietic and lymphoid neoplasms

Terminology	Grade Code
T-cell; T-precursor	5
B-Cell; Pre-B; B-precursor	6
Null cell; Non T-non B	7
NK cell (natural killer cell)	8
Grade unknown, not stated, or not applicable	9

Solid Tumors

Pathologic examination determines the grade, or degree of differentiation, of the tumor. For these cancers, the grade is a measurement of how closely the tumor cells resemble the parent tissue (organ of origin). Well-differentiated tumor cells closely resemble the tissue from the organ of origin. Poorly differentiated and undifferentiated tumor cells are disorganized and abnormal looking; they bear little (poorly differentiated) or no (undifferentiated) resemblance to the tissue from the organ of origin. These similarities/differences may be based on pattern (architecture), cytology, nuclear (or nucleolar) features, or a combination of these elements, depending upon the grading system that is used. Some grading systems use only pattern, for example Gleason grading in prostate. Others use only a nuclear grade (usually size, amount of chromatin, degree of irregularity, and mitotic activity). Fuhrman’s grade for kidney is based only on nuclear features. Most systems use a combination of pattern and cytologic and

nuclear features; for example Nottingham's for breast combines numbers for pattern, nuclear size and shape, and mitotic activity. The information from this data item is useful for determining prognosis and treatment.

Pathologists describe the tumor grade using three systems or formats:

1. Two levels of similarity; also called a two-grade system
2. Three levels of similarity; also called a three-grade system (code according to "Coding for solid tumors."
 - a. Grade I, well
 - b. Grade II, moderately
 - c. Grade III, poorly (undifferentiated carcinoma is usually separated from this system, since "poorly" bears some, albeit little, similarity to the host tissue, while "undifferentiated" has none, e.g. Undifferentiated carcinoma).
3. Four levels of similarity; also called a four-grade system. The four-grade system describes the tumor as
 - a. Grade I; also called well-differentiated
 - b. Grade II; also called moderately differentiated
 - c. Grade III; also called poorly differentiated
 - d. Grade IV; also called undifferentiated or anaplastic

Breast and prostate grades may convert differently than other sites. These exceptions are noted in "Coding for Solid Tumors", #7-8 below.

Coding Grade for Solid Tumors

1. Systemic treatment and radiation can alter a tumor's grade. Therefore, it is important to code grade based on information prior to neoadjuvant therapy even if grade is unknown.
2. Code the grade from the primary tumor only.
 - a. Do NOT code grade based on metastatic tumor or recurrence. In the rare instance that tumor tissue extends contiguously to an adjacent site and tissue from the primary site is not available, code grade from the contiguous site.
 - b. If primary site is unknown, code grade to 9.
3. Code the grade shown below (6th digit) for specific histologic terms that imply a grade.
 - Carcinoma, undifferentiated (8020/34)
 - Carcinoma, anaplastic (8021/34)
 - Follicular adenocarcinoma, well differentiated (8331/31)
 - Thymic carcinoma, well differentiated (8585/31)
 - Sertoli-Leydig cell tumor, poorly differentiated (8631/33)
 - Sertoli-Leydig cell tumor, poorly differentiated with heterologous elements (8634/33)
 - Undifferentiated sarcoma (8805/34)
 - Liposarcoma, well differentiated (8851/31)
 - Seminoma, anaplastic (9062/34)
 - Malignant teratoma, undifferentiated (9082/34)
 - Malignant teratoma, intermediate type (9083/32)
 - Intraosseous osteosarcoma, well differentiated (9187/31)
 - Astrocytoma, anaplastic (9401/34)
 - Oligodendroglioma, anaplastic (9451/34)
 - Retinoblastoma, differentiated (9511/31)

Retinoblastoma, undifferentiated (9512/34)

4. In situ and/or combined in situ/invasive components:
 - a. If a grade is given for an in situ tumor, code it. Do NOT code grade for dysplasia such as high grade dysplasia.
 - b. If there are both in situ and invasive components, code only the grade for the invasive portion even if its grade is unknown.

5. If there is more than one grade, code the highest grade within the applicable system. Code the highest grade even if it is only a focus. Code grade in the following priority order using the first applicable system:
 - a. special grade systems for the sites listed in Coding for Solid Tumors #6
 - b. differentiation: use Coding for Solid Tumors #7: 2-, 3-, or 4- grade system
 - c. nuclear grade: use Coding for Solid Tumors #7: 2-, 3-, or 4- grade system
 - d. If it isn't clear whether it is a differentiation or nuclear grade and a 2-, 3-, or 4- grade system was used, code it.
 - e. Terminology (use Coding for Solid Tumors #8)

6. Use the information from the special grade systems first. If no special grade can be coded, continue with Coding for Solid Tumors #7-9.

Special grade systems for solid tumors

Grade information based on CS Site-specific factors for breast, prostate, heart, mediastinum, peritoneum, retroperitoneum, soft tissue, and kidney parenchyma is used to code grade. See **Special Grade System Rules** section below for details on how to use this information to code grade.

CS Schema	Special grade system
Breast	Nottingham or Bloom-Richardson (BR) Score/Grade (SSF7)
Prostate	Gleason's Score on Needle Core Biopsy/Transurethral Resection of Prostate (TURP) (SSF 8)
Prostate	Gleason's Score on Prostatectomy/Autopsy (SSF 10)
Heart, Mediastinum	Grade for Sarcomas (SSF 1)
Peritoneum	Grade for Sarcomas (SSF 1)
Retroperitoneum	Grade for Sarcomas (SSF 1)
Soft Tissue	Grade for Sarcomas (SSF 1)
Kidney Parenchyma	Fuhrman Nuclear Grade (SSF 6)

Do not use these tables to code grade for any other groups including WHO (CNS tumors), WHO/ISUP (bladder, renal pelvis), or FIGO (female gynecologic sites) grades.

7. Use the Two-, Three- or Four-grade system information
 - a. Two-grade system

Term	Description	Grade Code	Exception for Breast and Prostate Grade Code
1/2, I/II	Low grade	2	1
2/2, II/II	High grade	4	3

In transitional cell carcinoma for bladder, the terminology high grade TCC and low grade TCC

are coded in the two-grade system.

- b. Three-grade system

Term	Description	Grade Code	Exception for Breast and Prostate Grade Code
1/3	Low grade	2	1
2/3	Intermediate grade	3	2
3/3	High grade	4	3

- c. Four-grade system: Any four-grade system including Edmondson and Steiner grade for liver.

Term	Description	Grade Code
1/4	Grade I; Well differentiated	1
2/4	Grade II; Moderately differentiated	2
3/4	Grade III; Poorly differentiated	3
4/4	Grade IV; Undifferentiated	4

- 8. Terminology: use the ‘Description’ column or the ‘Grade’ column to code grade. Breast & Prostate use the same grade code with a few noted exceptions.

Description	Grade	Assign Grade Code	Exception for Breast and Prostate Grade Code
Differentiated, NOS	I	1	
Well differentiated	I	1	
Only stated as ‘Grade I’	I	1	
Fairly well differentiated	II	2	
Intermediate differentiation	II	2	
Low grade	I-II	2	1
Mid differentiated	II	2	
Moderately differentiated	II	2	
Moderately well differentiated	II	2	
Partially differentiated	II	2	
Partially well differentiated	I-II	2	1
Relatively or generally well differentiated	II	2	
Only stated as ‘Grade II’	II	2	
Medium grade, intermediate grade	II-III	3	2
Moderately poorly differentiated	III	3	
Moderately undifferentiated	III	3	
Poorly differentiated	III	3	
Relatively poorly differentiate	III	3	
Slightly differentiated	III	3	
Dedifferentiated	III	3	
Only stated as “Grade III”	III	3	
High Grade	III-IV	4	3
Undifferentiated, anaplastic, not differentiated	IV	4	
Only stated as “Grade IV”	IV	4	
Non-high grade		9	

9. If no description fits or grade is unknown prior to neoadjuvant therapy, code as a 9 (unknown).

SPECIAL GRADE SYSTEMS RULES

Breast (site: breast excluding lymphomas; CS schema: breast)

Use Bloom Richardson (BR) or Nottingham score/grade to code grade based on CSv2 site-specific factor 7 (SSF) as stated below. If your registry does not collect this SSF, use the description in the table below to determine grade. If you collect this SSF, codes 030-130 could be automatically converted into the grade field.

BR could also be referred to as: Bloom-Richardson, modified Bloom-Richardson, BR, BR grading, Scarff-Bloom-Richardson, SBR grading, Elston-Ellis modification of Bloom-Richardson score, Nottingham modification of Bloom-Richardson score, Nottingham modification of Scarff-Bloom-Richardson, Nottingham-Tenovus grade, or Nottingham grade.

Code the tumor grade using the following priority order

- a. BR scores 3-9
- b. BR grade (low, intermediate, high)

BR score may be expressed as a range, 3-9. The score is based on three morphologic features: degree of tubule formation/histologic grade, mitotic activity, nuclear pleomorphism/nuclear grade of tumor cells. If a report uses words such as low, intermediate, or high rather than numbers, use the table below to code grade.

If only a grade of 1 through 4 is given with no information on the score and it is unclear if it is a Nottingham or BR Grade, do not use the table below. Continue with the next priority according to “Coding for Solid Tumors” #7 above.

Code the highest score if multiple scores are reported (exclude scores from tests after neoadjuvant therapy began). Examples: different scores may be reported on multiple pathology reports for the same primary cancer; different scores may be reported for multiple tumors assigned to the same primary cancer.

CS Site-Specific Factor 7 Nottingham or Bloom-Richardson (BR) Score/Grade

Description	CS Code	Grade Code
Score of 3	030	1
Score of 4	040	1
Score of 5	050	1
Score of 6	060	2
Score of 7	070	2
Score of 8	080	3
Score of 9	090	3
Low Grade, Bloom-Richardson (BR) grade 1, score not given	110	1
Medium (Intermediate) Grade, BR grade 2, score not given	120	2
High Grade, BR grade 3, score not given	130	3

Kidney Parenchyma (Site: kidney parenchyma excluding lymphomas; CS schema: KidneyParenchyma):

The Fuhrman Nuclear Grade should be used to code grade for kidney parenchyma only based on CSv2 SSF 6 as stated below. Do not use for kidney renal pelvis. If your registry does not collect this SSF, use the description in the table to determine grade. If you collect this SSF, the information could be automatically converted into the grade field if it is coded 010-040. Fuhrman nuclear grade is a four-grade system based on nuclear diameter and shape, the prominence of nucleoli, and the presence of chromatin clumping in the highest grade.

Description	CS Code	Grade Code
Grade 1	010	1
Grade 2	020	2
Grade 3	030	3
Grade 4	040	4

SoftTissue (sites excluding lymphomas: soft tissue, heart, mediastinum, peritoneum, and retroperitoneum; for CS users: SoftTissue, HeartMediastinum, Peritoneum, Retroperitoneum schemas): Grade for Sarcomas

The Grade for Sarcomas should be used to code grade based on CSv2 SSF 1 as stated below. If your registry does not collect this SSF, use the description in the table to determine grade. If you collect this SSF, the information could be automatically converted into the grade field if it is coded 010-200. The grading system of the French Federation of Cancer Centers Sarcoma Group (FNCLCC) is the preferred system.

Record the grade from any three-grade sarcoma grading system the pathologist uses. For terms such as "well differentiated" or "poorly differentiated," go to Coding for Solid Tumors #8.

In some cases, especially for needle biopsies, grade may be specified only as "low grade" or "high grade." The numeric grade takes precedence over "low grade" or "high grade."

Description	CS Code	Grade Code
Specified as Grade 1 [of 3]	010	2
Specified as Grade 2 [of 3]	020	3
Specified as Grade 3 [of 3]	030	4
Grade stated as low grade, NOS	100	2
Grade stated as high grade, NOS	200	4

Prostate (site: prostate excluding lymphomas; CS schema: prostate)

Use the highest Gleason score from the biopsy/TURP or prostatectomy/autopsy. Use a known value over an unknown value. Exclude results from tests performed after neoadjuvant therapy began. This information is collected in CSv2 SSF 8 (Gleason score from biopsy/TURP) and SSF 10 (Gleason score from prostatectomy/autopsy) as stated below. Use the table below to determine grade even if your registry does not collect these SSFs. If you collect these SSFs, the information could be converted into the grade field automatically.

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, 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 only one number is given on a particular test and it is less than or equal to 5 and not specified as a score, do not use the information because it could refer to either a score or a grade. If only one number is given and it is greater than 5, assume that it is a score and use it. If the pathology report specifies a specific number out of a total of 10, the first number given is the score. Example: The pathology report says Gleason 3/10. The Gleason score would be 3.

Historic Perspective

Gleason Score	Description					
	CS Code	Grade Code	AJCC 7th	SEER 2003-2013	AJCC 6th	SEER prior 2003
2	002	1	G1	G1	G1	G1
3	003	1	G1	G1	G1	G1
4	004	1	G1	G1	G1	G1
5	005	1	G1	G2	G2	G2
6	006	1	G1	G2	G2	G2
7	007	2	G2	G3	G3	G2
8	008	3	G3	G3	G3	G3
9	009	3	G3	G3	G3	G3
10	010	3	G3	G3	G3	G3

Historical perspective on long term trends in prostate grade: The relationship of Gleason score to grade changed for 1/1/2014+ diagnoses in order to have the grade field in sync with AJCC 7th ed. Analysis of prostate grade before 2014 based solely on the grade field is not recommended. In Collaborative Stage (CS), Gleason score was originally coded in CSv1 in one field (SSF 6) and then it was split into two fields in CSv2 based on the tissue used for the test: needle biopsy/TURP (SSF 8) and prostatectomy/autopsy (SSF 10). For trends using data back to 2004, if one collected the various CS Gleason scores, one could design a recode to have the same criteria as the data collected 2014+. The original grade field would NOT be changed, but for analyses this recode could be based on the CS SSFs and the original grade code.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions, Text-DX Proc-Path* and *Text-Histology Title*. These text fields are used by the PCR to validate ICD-O grade codes reported.

DIAGNOSTIC CONFIRMATION

Record the diagnostic confirmation which specifies whether a diagnosis was confirmed microscopically at any time during the disease course.

Note: The codes and instructions for hematopoietic and lymphoid neoplasm are different than the codes for solid tumors. See below.

Codes and Definitions

Code	Definition
Microscopically Confirmed	
1	<i>Positive histology.</i> Microscopic diagnosis based on tissue specimens from biopsy, frozen section, surgery, autopsy, or dilatation and curettage. Bone marrow biopsy and bone marrow aspiration. Hematologic confirmation of leukemia (peripheral blood smear).
2	<i>Positive cytology, no positive histology.</i> Cytologic diagnosis based on microscopic examination of cells as contrasted with tissues. Fine-needle aspiration (FNA) is frequently used to obtain a cytologic specimen. Cells may be removed from exudates, secretions, or washings from tissue. (e.g., Sputum smears, bronchial brushings, bronchial washings, tracheal washings, prostatic secretions, breast secretions, gastric fluid, spinal fluid, peritoneal fluid, pleural fluid, and urinary sediment, cervical and vaginal smears.) Cytology also includes paraffin-block specimens from concentrated spinal, pleural, or peritoneal fluid.
3	<p><i>Positive histology PLUS</i></p> <ul style="list-style-type: none"> • <i>Positive immunophenotyping AND/OR</i> • <i>Positive genetic studies</i> <p>Histology is positive for cancer, and there are also positive immunophenotyping and/or genetic test results.</p> <p>Code 3 is used ONLY for hematopoietic and lymphoid neoplasms (9590/3-9992/3) Effective for cases diagnosed 1/1/2010 and later.</p>
4	<i>Positive microscopic confirmation, method not specified.</i> The record is reported as microscopically confirmed but no information is provided about the method (histology or cytology).
Not Microscopically Confirmed	
5	<i>Positive laboratory test/marker study.</i> A clinical diagnosis of cancer is based on laboratory tests/marker studies which are clinically diagnostic for cancer. This includes alpha-fetoprotein for liver cancer and abnormal electrophoretic spike for multiple myeloma. Elevated PSA is nondiagnostic of cancer. If the physician uses the PSA as a basis for diagnosing prostate cancer with no other workup, record as code 5.
6	<i>Direct visualization without microscopic confirmation.</i> Use this code only in the absence of positive histology or cytology. Diagnosis made at surgical exploration or by endoscopy. Autopsy only record (only information is from gross autopsy report).

Code	Definition
7	<i>Radiography and other imaging techniques without microscopic confirmation.</i> Use this code only in the absence of positive histology or cytology. Diagnosed by radiology, including ultrasound, computerized (axial) tomography (CT or CAT scans) and magnetic resonance imaging (MRI).
8	<i>Clinical diagnosis only (other than 5, 6, or 7).</i> Use this code only in the absence of positive histology or cytology. Records diagnosed by clinical methods not mentioned previously.
Confirmation Unknown	
9	<i>Unknown whether or not microscopically confirmed.</i> Death-certificate-only records (PCR use only). Method of confirmation is unknown.

Recording Diagnostic Confirmation for Solid Tumors

1. Solid Tumors- These guidelines should be used for solid tumors only.
2. Priority - This is a hierarchical coding scheme with code 1 taking precedence. **A lower number takes priority over all higher numbers.**
3. Changing Information- This data item is dynamic and must be changed to the lower code if a more definitive method confirms the diagnosis at any time during the course of the disease. See *Part One, Changing Information* for instructions on how to submit a change.

Example: A patient is admitted on 01/28/2009. A chest x-ray dated 02/01/2010 diagnoses a probable lung cancer. The patient refuses a diagnostic workup. The registry codes the diagnostic confirmation to radiography (7). The patient consents to a lymph node biopsy on 06/03/2010. The biopsy confirms small cell carcinoma. Change the diagnostic confirmation code to positive histology (1).

4. Assign code 1 when the microscopic diagnosis is based on:
 - a. Tissue specimens from biopsy, frozen section, surgery, autopsy or D&C
 - b. Bone marrow specimens (aspiration and biopsy)
5. 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 and vaginal smears.
 - b. Paraffin block specimens from concentrated spinal, pleural, or peritoneal fluid
6. Assign code 4 when there is information that the diagnosis of cancer was microscopically confirmed, but the type of confirmation is unknown.
7. Assign code 5 when the diagnosis of cancer is based on laboratory tests or marker studies which 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 that PSA, code the diagnostic confirmation to 5.

8. **Assign code 6** when the diagnosis is based only on:
 - 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).
9. **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.
10. **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 since there is a physician's clinical diagnosis.

11. **Assign code 9** if it is unknown if the diagnosis was confirmed microscopically and for Death certificate only cases.

Recording Diagnostic Confirmation for Hematopoietic and Lymphoid Neoplasms (9590/3-9992/3)

1. **Hematopoietic and Lymphoid Neoplasms**- These guidelines should be used for Hematopoietic and Lymphoid Neoplasms only.
2. **No Priority**- There is no priority order or hierarchy for coding the *Diagnostic Confirmation* for hematopoietic or lymphoid neoplasms. Most commonly the specific histologic type is diagnosed by immunophenotyping or genetic testing.

Note: See the glossary in the *2010 Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* for definitions of immunophenotyping and genetic testing. The manual can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

3. **Definitive Diagnostic Confirmation**- See the *Hematopoietic Database* for specific information on the definitive diagnostic confirmation for the neoplasm being abstracted. The database can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

4. **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.

5. 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
6. Assign code 3 when there is a histology positive for cancer and also positive immunophenotyping and/or positive genetic testing

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* to 3- positive histology and positive genetic testing.

Example 2: Skin biopsy positive for cutaneous T-cell lymphoma, NOS (9960/3). Immunophenotyping shows CD8 positive. Diagnosis is primary cutaneous CD 8 positive aggressive epidermitropic T-cell lymphoma (9709/3). Code the *Diagnostic Confirmation* to 3- positive histology and positive genetic testing.

7. Assign code 4 when there is information the diagnosis of cancer was microscopically confirmed, but the type of confirmation is unknown.
8. Assign code 5 when the diagnosis of a hematopoietic or lymphoid neoplasm is based ONLY on laboratory tests or marker studies which are diagnostic for that specific cancer.

Note: See the glossary in the *2010 Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual* for definitions of immunophenotyping and genetic testing which are marker studies for hematopoietic and lymphoid neoplasms. The manual can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

Example: The only information available is 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 for polycythemia vera.

9. 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).
10. **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.
11. **Assign code 8** when the case was diagnosed by any clinical method not mentioned in 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). Other than 6 or 7.

Note: The *Hematopoietic Database* will identify clinical diagnosis as the definitive diagnostic method. The database can be downloaded from the following site:

<http://seer.cancer.gov/seertools/hemelymph/>

12. **Assign code 9** when:
- a. It is unknown if the diagnosis was confirmed microscopically.
 - b. Death certificate only case.

Text

Record text to describe the findings of the diagnostic procedures in natural language

TUMOR SIZE SUMMARY

Record the most definitive size of a solid primary tumor.

Codes and Definitions

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)
990	Microscopic focus or foci only and no size of focus is given
998	Alternate descriptions of tumor size for the following specific sites: <ul style="list-style-type: none"> • Familial/multiple polyposis- Colon, Rectosigmoid and Rectum If no size is documented: <ul style="list-style-type: none"> • Circumferential: Esophagus • Diffuse; widespread: 3/4s or more; linitis plastica- Stomach and Esophagus GE Junction • Diffuse, entire lung or NOS- Lung and main stem bronchus • Diffuse- Breast
999	Unknown; size not stated Not documented in patient record Size of tumor cannot be assessed Not Applicable: <ul style="list-style-type: none"> • Kaposi Sarcoma • Melanoma Choroid • Melanoma Ciliary Body • Melanoma Iris • Hematopoietic, Reticuloendothelial, and Myeloproliferative neoplasms; histology codes 9590-9992

Recording Tumor Size Summary

1. Millimeters- Measurements recorded must be in millimeters.
2. Priority Order- Record size of the tumor in the following specified order:
 - a. Surgery with NO pre-surgical treatment- Record the size measured on the surgical resection specimen when surgery is administered as the first definitive treatment.
 - i. If there is a discrepancy among tumor size measurements in the various sections of the pathology report, code the size from the synoptic report (CAP protocol or checklist). If only a text report is available, use final diagnosis, microscopic, or gross examination, in this order.

- b. Neoadjuvant therapy before surgery- Code the largest size of tumor prior to neoadjuvant treatment; if unknown code size as 999
 - c. No Surgery- code the largest measurement of the tumor from physical exam, imaging, or other diagnostic procedures prior to any other form of treatment.
 - d. None of the above- code the largest size from all information available within four (4) months of the date of diagnosis, in the absence of disease progression.
3. Depth or Thickness- Do not record the depth or thickness for Tumor Size Summary; tumor size is the diameter of the tumor.
4. “Between, Less or Greater Than”- See below for specific instructions on how to record size of tumor when “between”, “less than” or “greater than” is stated only.
 - a. “Between”- If tumor size is reported to be between two sizes, record tumor size as the midpoint between the two.

Example: “The size of the tumor is between 2 and 3 cm.” Convert the size to mm and record the tumor size summary as 025, the midpoint.
 - b. “Less Than”- If the tumor size is stated as “less than” x mm, record the size as 1mm less than stated. If size is less than 1 mm, use code 001.

Example: “The size of the tumor is <10 mm in greatest diameter.” Code tumor size summary as 009.
 - c. “Greater Than”- If the tumor size is stated as “more than” x mm, record the size as 1mm higher than stated. If size is described as anything greater than 989 mm, code as 989.

Example: “The size of the tumor is >10 mm.” Code tumor size summary as 011.
5. Rounding- Round the size only if it is described in fractions of millimeters. Round tenths of millimeters in the 1-4 range down to the nearest whole millimeter and 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 centimeter; rather, move the decimal point one space to the right, converting the measurement to millimeters, then round if necessary.

Example #1: “Breast cancer described as 4.5 mm in size.” Round up the tumor size to 5 mm and record the tumor size summary as 005.

Example #2: “Focus of cancer described as 1.3 mm in size. Round down the size to 1 mm and record the tumor size summary as 001.
6. Discrepancies in Size among Imaging and Radiographic Reports- Unless the physician specifies which imaging is most accurate, record the largest size in the record, regardless of which imaging technique reports it.
7. Polyp, Cyst, Distant Mets- Do not record the size of a polyp, cyst, or distant metastasis.

Exception: 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 cysts are part of the tumor itself.

8. In Situ Only- Record the size as stated for purely in situ lesions.
9. In Situ and Invasive Components- Record the size of the invasive component, if given.
 - a. Both In Situ and Invasive size given- If both an in situ and an invasive component are present and the invasive component is measured, record the size of the invasive component even if it is smaller than the in situ component.

Example: Tumor is mixed in situ and invasive carcinoma, total 4.1 cm in size, of which 1.7 cm is invasive. Record the tumor size as 017 (17 mm).

- b. Invasive component NOT given- 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.

Example: Duct carcinoma in situ measuring 2.2 cm with an area of invasive ductal carcinoma. Record the tumor size as 022 (22 mm).

10. Largest Dimension or Diameter of Tumor- Record the largest dimension or diameter of the tumor whether it is from an excisional biopsy or the complete resection of the tumor.

Example: Tumor is described as 3.2 x 6.2 x 1.6 cm in size. Record tumor size as 062 (62 mm).

11. Residual or Positive Surgical Margins- Disregard microscopic residual or positive surgical margins when coding tumor size. Microscopic residual tumor does not affect overall tumor size.
12. Pieces or Chips of Tumor- Do not add the size of pieces or 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. If the only measurement describes pieces or chips, record the tumor size as 999.
13. Multifocal and Muticentric 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.
14. Unknown and Not Applicable- Use code 999 when size is unknown or not applicable. See table below for a list of not applicable sites and morphologies.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions*. These text fields are used by the PCR to validate the tumor size for each primary.

REGIONAL NODES EXAMINED

Record the total number of regional lymph nodes that were removed and examined by the pathologist.

Codes and Definitions

Code	Definition
00	No nodes examined
01 - 89	1 to 89 nodes examined (code the exact number of regional 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.
99	Unknown whether nodes were examined; not applicable or negative; not documented in patient record.

Recording Regional Nodes Examined

1. Regional Lymph Nodes- Record information about regional lymph nodes only in this field.
2. Pathologic Information Only- This field is based on pathologic information only.
3. Cumulative Nodes- 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.
 - a. Core Needle Biopsy or FNA of SAME LN Chain- 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.

Example: 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.*

- b. Core Needle Biopsy or FNA of DIFFERENT LN Chain- 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 Positive.

Example: 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.*

- c. Core Needle Biopsy or FNA of UNKNOWN LN Chain- If the location of the lymph node that is core-biopsied or aspirated 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 Positive.

Example: 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.*

4. No Nodes Examined- Record code 00 (no regional lymph nodes examined) when the assessment of lymph nodes is clinical or when no lymph nodes are removed and examined.
5. Priority- If there is a discrepancy regarding the number of lymph nodes examined, use information in the following priority order: final diagnosis, synoptic report (also known as CAP protocol or pathology report checklist), microscopic and gross.
6. Cytology/Core Biopsy- Use code 95 when the only procedure for regional lymph nodes is a needle aspiration (cytology) or core biopsy (tissue).
7. 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.
8. Sampling- 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.
9. Dissection- 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.
10. 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.
11. Unknown/Not Applicable- Code 99 if it is unknown whether nodes were removed or examined. Code 99 (Not Applicable) for the following sites: Placenta, Brain and Cerebral Meninges, Other Parts of CNS, Intracranial Gland, Hematopoietic, Reticuloendothelial, Immunoproliferative and Myeloproliferative Neoplasms, Hodgkin and non-Hodgkin Lymphoma, Myeloma and PlasmaCell Disorders, Other and Ill-Defined Primary Sites, and Unknown Primary Site.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions*. These text fields are used by the PCR to validate the numbers of regional lymph nodes examined.

REGIONAL NODES POSITIVE

Record the total number of positive regional lymph nodes that were examined by the pathologist.

Codes and Definitions

Code	Definition
00	All nodes examined negative
01 - 89	1 to 89 nodes positive (code exact number of nodes positive)
90	90 or more nodes positive
95	Positive aspiration or core biopsy of lymph node(s).
97	Positive nodes - number unspecified.
98	No nodes examined.
99	Unknown whether nodes are positive; not applicable; not documented in patient record.

Recording Regional Nodes Positive

1. Regional Lymph Nodes- Record information about regional lymph nodes only in this field.
2. **Pathologic Information Only**- This field is **based on pathologic information only**.
3. In-Situ Cases- True in situ cases cannot have positive lymph nodes, so the only allowable codes are 00 (negative) or 98 (not examined).
4. Cumulative Nodes- The number of regional lymph nodes positive is cumulative from all procedures that removed lymph nodes through the completion of surgeries in the first course of treatment.
 - a. Core Needle Biopsy or FNA of SAME LN Chain- 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.

Example: 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.*
 - b. Core Needle Biopsy or FNA of DIFFERENT LN Chain- 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 Positive.

Example: 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.*
 - c. Core Needle Biopsy or FNA of UNKNOWN LN Chain- If the location of the lymph node that is core-biopsied or aspirated 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 Positive.

Example: 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.*

5. Priority- If there is a discrepancy regarding the number of lymph nodes examined, use information in the following priority order: final diagnosis, synoptic report (also known as CAP protocol or pathology report checklist), microscopic and gross.
6. Positive Nodes in Multiple Primaries in Same Organ- If there are multiple primary cancers with different histologic types in the same organ and the pathology report just states the number of nodes positive, the registrar should first try to determine the histology of the metastases in the nodes and code the nodes as positive for the primary with that histology. If no further information is available, code the nodes as positive for all primaries.
7. Isolated Tumor Cells (ITCs)- For all primary sites 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). Do not include in the count of lymph nodes positive any nodes that are identified as containing isolated tumor cells (ITCs). If the path report indicates that nodes are positive but the size of metastasis is not stated, assume the metastases are larger than 0.2 mm and count the lymph node(s) as positive. For cutaneous melanoma and Merkel cell carcinoma, count nodes with ITCs as positive lymph nodes.
8. Cytology/Core Biopsy- Use code 95 when the only procedure for regional lymph nodes is a needle aspiration (cytology) or core biopsy (tissue).
9. Cannot Determine Number of Positive Nodes- 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.
10. No Nodes Examined- Use Code 98 when the assessment of lymph nodes is clinical only or when no lymph nodes are removed/examined. Code 98 is used often when regional lymph nodes examined is coded to 00.
11. Unknown/Not Applicable- Code 99 if it is unknown whether nodes were positive. Code 99 (Not Applicable) for the following sites: Placenta, Brain and Cerebral Meninges, Other Parts of CNS, Intracranial Gland, Hematopoietic, Reticuloendothelial, Immunoproliferative and Myeloproliferative Neoplasms, Hodgkin and non-Hodgkin Lymphoma, Myeloma and PlasmaCell Disorders, Other and Ill-Defined Primary Sites, and Unknown Primary Site.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions*. These text fields are used by the PCR to validate the numbers of regional lymph nodes positive.

LYMPH-VASCULAR INVASION (LVI)

Record for presence or absence of tumor cells in lymphatic channels (not lymph nodes) or blood vessels within the primary tumor as noted microscopically by the pathologist.

Codes and Definitions

Code	Definition
0	Lymph-vascular invasion not present, absent, not identified
1	Lymph-vascular invasion present
8	Not applicable
9	Unknown if lymph-vascular invasion present Indeterminate

Recording Lymph-Vascular Invasion

1. Pathology Report- The primary source of information about lymph-vascular invasion can be found within the pathology checklist or synoptic report. If the case does not have a checklist or synoptic report, code from the pathology report or a physician’s statement, in that order.
2. Perineural Invasion- Do not code perineural invasion in this field.
3. Specimen- Information to code lymph-vascular invasion can be taken from any specimen from the primary tumor (biopsy or resection). If lymph-vascular invasion is identified in any specimen, it should be coded as present/identified (1).
4. Benign/Borderline Tumor- For cases with benign or borderline behavior, code the lymph-vascular invasion documented (negative or positive) and, if not documented, code unknown.
5. Unknown- Use code 9 (Unknown) for the following situations:
 - a. there is no microscopic examination of a primary tissue specimen
 - b. the primary site specimen is cytology only or a fine needle aspiration
 - c. the biopsy is only a very small tissue sample
 - d. it is not possible to determine whether lymph-vascular invasion is present
 - e. the pathologist indicates the specimen is insufficient to determine lymph-vascular invasion
 - f. lymph-vascular invasion is not mentioned in the pathology report
 - g. primary site is unknown

- h. there is no documentation or information from the pathology report or any other sources
6. Not Applicable- Use code 8 for the following primary sites: Hodgkin and Non-Hodgkin lymphoma, Leukemias, Hematopoietic and reticuloendothelial disorders, Myelodysplastic syndromes including refractory anemias and refractory cytopenias, Myeloproliferative disorders.

Text

Text to support this data item must be recorded in the specific text fields. See *Part Three, Data Item Instructions*. These text fields are used by the PCR to validate the presence or absence of lymph vascular invasion.

METS AT DIAGNOSIS - BONE

Record for presence of BONE metastasis from the primary site at the time of diagnosis. Code information about bone metastases only (discontinuous or distant metastases to bone) identified at the time of diagnosis.

Codes and Definitions

Code	Definition
0	None; no bone metastases
1	Yes; distant bone metastases
8	Not applicable
9	Unknown whether bone is an involved metastatic site Not documented in patient record

Recording Mets at Diagnosis - Bone

1. Bone Marrow Involvement- This data item should not be coded for bone marrow involvement.
2. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
3. Clinical or Pathologic Information- Any clinical and/or pathologic information about bone involvement may be used to code this data item.
4. Preoperative Systemic Therapy- Code this data item for bone metastases even if the patient had any preoperative systemic therapy.
5. No Distant (discontinuous) Metastases- Assign code 0 when there is no bone metastases mentioned.
6. Distant (discontinuous) Metastases- Assign code 1 when bone is mentioned as an involved site.
7. Bone Primaries:
 - a. Assign code 1 if the medical record indicates that there are metastases in a different bone or bones than the primary site.
 - b. Do not assign code 1 for a bone primary with multifocal bone involvement of the same bone.
8. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

9. Unknown- Use code 9 when there are known distant metastases but it is not known whether the distant metastases includes bone.

METS AT DIAGNOSIS - BRAIN

Record for presence of BRAIN metastasis from the primary site at the time of diagnosis. Code information about brain metastases only (discontinuous or distant metastases to brain) identified at the time of diagnosis.

Codes and Definitions

Code	Definition
0	None; no brain metastases
1	Yes; distant brain metastases
8	Not applicable
9	Unknown whether brain is an involved metastatic site Not documented in patient record

Recording Mets at Diagnosis - Brain

1. Spinal Cord & CNS Involvement- This data item should not be coded for spinal cord or other parts of the central nervous system involvement.
2. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
3. Clinical or Pathologic Information- Any clinical and/or pathologic information about brain involvement may be used to code this data item.
4. Preoperative Systemic Therapy- Code this data item for brain metastases even if the patient had any preoperative systemic therapy.
5. No Distant (discontinuous) Metastases- Assign code 0 when there is no brain metastases mentioned.
6. Distant (discontinuous) Metastases- Assign code 1 when brain is mentioned as an involved site.
7. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

8. Unknown- Use code 9 when there are known distant metastases but it is not known whether the distant metastases includes brain.

METS AT DIAGNOSIS – DISTANT LYMPH NODES

Record for presence of DISTANT LYMPH NODE(S) metastasis from the primary site at the time of diagnosis. Code information about distant lymph node(s) metastases only identified at the time of diagnosis.

Codes and Definitions

Code	Definition
0	None; no distant lymph node metastases
1	Yes; distant lymph node metastases
8	Not applicable
9	Unknown whether distant lymph node(s) are involved metastatic site Not documented in patient record

Recording Mets at Diagnosis – Distant Lymph Nodes

1. Regional Lymph Nodes- This data item should not be coded for regional lymph node involvement with the exception of lymph nodes for placenta which are M1.
2. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
3. Clinical or Pathologic Information- Any clinical and/or pathologic information about distant lymph node involvement may be used to code this data item.
4. Preoperative Systemic Therapy- Code this data item for distant lymph node metastases even if the patient had any preoperative systemic therapy.
5. No Distant (discontinuous) Metastases- Assign code 0 when there is no distant lymph node metastases mentioned.
6. Distant (discontinuous) Metastases- Assign code 1 when distant lymph node is mentioned as an involved site.
7. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

8. Unknown- Use code 9 when there are known distant metastases but it is not known whether the distant metastases includes distant lymph node(s).

METS AT DIAGNOSIS – LIVER

Record for presence of LIVER metastasis from the primary site at the time of diagnosis. Code information about liver metastases only identified at the time of diagnosis.

Codes and Definitions

Code	Definition
0	None; no liver metastases
1	Yes; liver metastases
8	Not applicable
9	Unknown whether liver is involved metastatic site Not documented in patient record

Recording Mets at Diagnosis – Liver

1. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
2. Clinical or Pathologic Information- Any clinical and/or pathologic information about liver involvement may be used to code this data item.
3. Preoperative Systemic Therapy- Code this data item for liver metastases even if the patient had any preoperative systemic therapy.
4. No Distant (discontinuous) Metastases- Assign code 0 when there is no liver metastases mentioned.
5. Distant (discontinuous) Metastases- Assign code 1 when liver is mentioned as an involved site.
6. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2-C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

7. Unknown- Use code 9 when there are known distant metastases but it is not known whether the distant metastases includes liver.

METS AT DIAGNOSIS – LUNG

Record for presence of LUNG metastasis from the primary site at the time of diagnosis. Code information about distant lung metastases only identified at the time of diagnosis.

Codes and Definitions

Code	Definition
0	None; no lung metastases
1	Yes; distant lung metastases
8	Not applicable
9	Unknown whether lung is involved metastatic site Not documented in patient record

Recording Mets at Diagnosis – Lung

1. Pleural or Pleural Fluid- This data item should not be coded for pleural or pleural fluid involvement.
2. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
3. Clinical or Pathologic Information- Any clinical and/or pathologic information about lung involvement may be used to code this data item.
4. Preoperative Systemic Therapy- Code this data item for lung metastases even if the patient had any preoperative systemic therapy.
5. No Distant (discontinuous) Metastases- Assign code 0 when there is no lung metastases mentioned.
6. Distant (discontinuous) Metastases- Assign code 1 when lung is mentioned as an involved site.
7. Lung Primaries:
 - a. Assign code 1 if the medical record indicates that there are metastases in the contralateral lung
 - b. Do not assign code 1 for a lung primary with multifocal involvement of the same lung.
8. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

9. Unknown- Use code 9 when there are known distant metastases but it is not known whether the distant metastases includes lung.

METS AT DIAGNOSIS – OTHER

Record for presence of any OTHER metastasis (excluding bone, brain, liver, lung or distant lymph node) from the primary site at the time of diagnosis. Code information about other metastases only identified at the time of diagnosis.

Some examples of other sites include: adrenal gland, bone marrow, pleura, peritoneum and skin.

Codes and Definitions

Code	Definition
0	None; no other metastases
1	Yes; distant metastases in known site(s) other than bone, brain, liver, lung or distant lymph nodes
8	Not applicable
9	Unknown whether any other metastatic site Not documented in patient record

Recording Mets at Diagnosis – Other

1. Metastatic Sites- Do not code this data item for bone, brain, liver, lung or distant lymph node metastases.
2. Primary Site- This data item should be coded for all solid tumors, Kaposi sarcoma, Unknown Primary Site, and Other and Ill-Defined Primary Sites.
3. Clinical or Pathologic Information- Any clinical and/or pathologic information about lung involvement may be used to code this data item.
4. Preoperative Systemic Therapy- Code this data item for lung metastases even if the patient had any preoperative systemic therapy.
5. No Distant (discontinuous) Metastases- Assign code 0 when there is no other site(s) metastases mentioned.
6. Distant (discontinuous) Metastases- Assign code 1 when other site(s) is mentioned as an involved site.
7. Not Applicable- Use code 8 for the following site/histology combinations for which a code for distant metastasis is not clinically relevant:

ICD-O-3 Site	ICD-O-3 Histology	Description
C00.0-C80.9	9740-9809, 9840-9992	Mast cell, histiocytosis, immunoproliferative, leukemias coded to any site.
C42.0, C42.1, C42.4	9811-9818, 9823, 9827, 9837	Specific leukemia/lymphoma histologies coded to blood, bone marrow, hematopoietic
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9820, 9826, 9831-9834	Mostly lymphoid leukemias coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS
C00.0-C44.0, C44.2- C68.9, C69.1-C69.4, C69.8-C80.9	9731, 9732, 9734	Plasma cell tumors coded to any site except eyelid, conjunctiva, lacrimal gland, orbit, and eye overlapping and NOS

8. Unknown- Use code 9 when there are known distant metastases but it is not known specifically what they are.

THE COLLABORATIVE STAGE DATA COLLECTION SYSTEM

The Collaborative Stage Data Collection System (CS) is a carefully selected, medically relevant set of data items that describe how far a cancer has spread at the time of diagnosis. Most of the data items have traditionally been collected by cancer registries, including tumor size, extension, lymph node status, and metastatic status. New items were created to collect information necessary for the conversion algorithms, including the evaluation fields that describe how the collected data were determined, and site/histology-specific factors necessary to derive the final stage grouping for certain primary cancers. In addition to the items coded by the registrar, this unified data set also includes several data items derived from the computer algorithms that classify each case in multiple staging systems: the sixth and seventh editions of the AJCC TNM system (TNM), Summary Stage 1977 (SS77), and SEER Summary Stage 2000 (SS2000).

Effective Date

1. Date of Diagnosis 2016 and after- *Collaborative Stage Data Collection System Version 02.05 (CSv02.05) will be used for the collection of the Site-Specific Factors (SSFs) for cases diagnosed 1/1/2016 and forward.* In addition to the SSFs, Regional Nodes Positive and Examined and Lymphovascular Invasion will continue to be required. All other CS input data items are no longer required.
2. Date of Diagnosis 2004 to 2015-Stage cases diagnosed on or after January 1, 2014 using the *Collaborative Stage Data Collection System Version 02.05 (CSv02.05)*. Straggler cases diagnosed in 2004-2013 should be staged using CSv02.05 after software has been updated to NAACCR V14. CSv02.05 can be downloaded from the following link:

<https://cancerstaging.org/cstage/coding/Pages/Version-02.05.aspx>
3. Date of Diagnosis 2001 to 2003 - Stage cases diagnosed between January 1, 2001 and December 31, 2003 according to *SEER Summary Stage 2000* and leave CS items and *SEER Summary Stage 1977* blank.
4. Date of Diagnosis prior to 2001 - Stage cases diagnosed prior to January 1, 2001 according to *SEER Summary Stage 1977* and leave CS Items and SEER Summary 2000 blank.

How Collaborative Staging Works

CS is a site-specific coding system. CS codes are defined for every site and histology combination. Depending on the site or histology, the coding schema and instructions will vary. For each reportable case, the CS data items specific to the cancer are extracted from the medical record and coded in the Collaborative Stage Data Collection System fields. When CS data collection is complete, the registrar activates the computer algorithms to derive the values for the items in the TNM system (both 6th and 7th edition) and Summary Stage (both 1977 and 2000). The derived data items, assigned to specific "derived" fields, will be incorporated into the hospital registry database.

Collaborative Stage Data Collection System Manual (CS Manual)

The complete instructions and site-specific defined codes are documented in the *Collaborative Stage Data Collection System Manual*.

1. Versions- It is very important the correct version of the *CS Manual* is used. The version of the *CS Manual* to be used is based on the year of diagnosis in conjunction with the NAACCR Record Layout Version.
 - a. Date of Diagnosis in 2014 or after- Cases diagnosed on or after January 1, 2014 must be coded using CS V02.05.
 - b. Date of Diagnosis in 2012- Cases diagnosed on or after January 1, 2012 must be coded using CS V02.04. CS V02.04 can only be used with NAACCR Record Layout V12.2. This means you cannot abstract cases diagnosed in 2012 until V12.2 is installed.
 - c. Date of Diagnosis in 2011- Cases diagnosed on or after January 1, 2011 must be coded using CS V02.03 or later and must be reported in NAACCR Record Layout V12.1 or later. This means once your software has been converted to NAACCR V12.2, you may report any remaining cases from previous years using CS V02.04.
 - d. Date of Diagnosis in 2010- Cases diagnosed on or after January 1, 2010 must be coded using CS V02.02 or later and must be reported in NAACCR Record Layout V12 or later. This means once your software has been converted to NAACCR V12.1, you may report any remaining cases from previous years using CS V02.03.
2. CS Manual- the CS Manual consists of two parts and is provided in an electronic format.
 - a. Part I - provides general instructions and codes for generic (non site-specific) items. Part I must be reviewed in detail and the guidelines must be applied when coding all CS fields.
 - b. Part II - contains site-specific instructions and codes. Site-specific guidelines must also be followed when coding all CS fields. Site-specific guidelines take priority over the general instructions in Part I of the CS Manual.
1. Download- CSv2._ can be downloaded from the following link:
<https://cancerstaging.org/cstage/coding/Pages/Version-02.05.aspx>

The following table lists the CS data items required to be reported to the PCR by Diagnosis Year.

Diagnosed 2016 and After

Data Item	Description
Regional Lymph Nodes Positive	Number of positive regional lymph nodes from the pathology report
Regional Lymph Nodes Examined	Number of regional lymph nodes examined by the pathologist
Lymph-Vascular Invasion	Code to describe the absence or presence of tumor cells in lymphatic channels or blood vessels within the primary tumor
CS Site-Specific Factors (SSF) 1-25*	Codes to describe site-specific prognostic information.

Diagnosed Before 2016

Data Item	Description
CS Tumor Size	Code to describe tumor size
CS Extension	Code to describe how far the tumor has spread directly
CS Tumor Size/Ext Eval	Code to describe how the farthest tumor spread was determined
CS Lymph Nodes	Code to describe whether regional lymph nodes are involved
CS Lymph Nodes Eval	Code to describe how the farthest lymph node spread was determined
Regional Lymph Nodes Positive	Number of positive regional lymph nodes from the pathology report
Regional Lymph Nodes Examined	Number of regional lymph nodes examined by the pathologist
Lymph-Vascular Invasion	Code to describe the absence or presence of tumor cells in lymphatic channels or blood vessels within the primary tumor
CS Mets at DX	Code to describe the farthest distant metastasis (including distant lymph nodes)
CS Mets Eval	Code to describe how the distant metastasis was determined
CS Mets at DX- Bone	Code to indicate if bone is an involved metastatic site
CS Mets at DX- Brain	Code to indicate if brain is an involved metastatic site
CS Mets at DX- Liver	Code to indicate if liver is an involved metastatic site
CS Mets at DX- Lung	Code to indicate if lung is an involved metastatic site
CS Site-Specific Factors (SSF) 1-25*	Codes to describe site-specific prognostic information.

*Not all 25 SSF are required for all sites. See *Appendix J* for a list of the PCR required SSFs by schema.

Text

Text to support all Collaborative Stage data items must be recorded in the specific text fields. See *Part Three, Data Item Instructions, Text-Path, Text-DX Proc-X-ray/Scans, Text-DX Proc-OP, and Text-DX Proc-Scopes*.

SEER SUMMARY STAGE

Record the SEER Summary Stage at diagnosis in the appropriate Summary Stage field according to SEER Summary Stage 2000 (SS2000) or SEER Summary Staging 1977 (SS77) based on *Date of Diagnosis*.

Effective with cases diagnosed on or after January 1, 2015 and after, directly coded SEER Summary Stage 2000 is also required.

1. Date of Diagnosis 2015 and after- Stage cases diagnosed on or after January 1, 2015 according to *SEER Summary Stage 2000* and the *Collaborative Stage Data Collection System Version 2 (CSv2)*. Leave SEER Summary Stage 1977 blank. The *SEER Summary Stage Manual 2000* can be downloaded from:

<http://seer.cancer.gov/tools/ssm/>
2. Date of Diagnosis 2010-2014- Stage cases diagnosed from January 1, 2010 through December 31, 2014 using the *Collaborative Stage Data Collection System Version 2 (CSv2)*. SS2000 and SS1977 must be left blank.
3. Date of Diagnosis 2004-2009- Until software is updated to NAACCR V12, stage cases diagnosed between January 1, 2004 through December 31, 2009 using *Collaborative Staging Manual and Coding Instructions, version 1.04.00 (CSv1)*. SS2000 and SS1977 must be left blank
4. Date of Diagnosis 2001-2003- Stage cases diagnosed between January 1, 2001 and December 31, 2003 according to *SEER Summary Stage 2000* and leave CS items and SEER Summary Stage 1977 blank. *SEER Summary Stage Manual 2000* can be downloaded from:
5. Date of Diagnosis Prior to 2001 - Stage cases diagnosed prior to January 1, 2001 according to *SEER Summary Stage Guide 1977* and leave CS Items and SEER Summary 2000 blank. *SEER Summary Stage Guide 1977* can be downloaded from:

http://seer.cancer.gov/archive/manuals/historic/ssm_1977.pdf

Text

Text to support this data item must be recorded in the specific text fields. Positive and negative findings describing how far the cancer has spread from the organ of origin and the corresponding dates of the findings must be recorded in the appropriate *Text-DX* fields.

AJCC-TNM CLINICAL AND PATHOLOGIC STAGE

The American Joint Committee on Cancer (AJCC) established the way cancer is communicated. Clinicians and the surveillance community count on the AJCC for the most comprehensive anatomic staging data available, the Cancer Staging Manual and Cancer Staging Atlas. These AJCC publications are recognized as the authoritative guides for cancer staging information and are used by tens of thousands of medical professionals everyday

The TNM Staging System was developed and is maintained by the AJCC and the Union for International Cancer Control (UICC). It is the most commonly used staging system by medical professionals around the world. The TNM classification system was developed as a tool for doctors to stage different types of cancer based on certain, standardized criteria.

The TNM Staging System is based on the extent of the tumor (T), the extent of spread to the lymph nodes (N), and the presence of metastasis (M).

Because each cancer type has its own classification system, letters and numbers do not always mean the same thing for every kind of cancer. Once the T, N, and M are determined, they are combined, and an overall stage of 0, I, II, III, IV is assigned. Sometimes these stages are subdivided as well, using letters such as IIIA and IIIB.

In some cancer types, non-anatomic factors are required for assigning the anatomic stage/prognostic group. These are clearly defined in each chapter of the AJCC Cancer Staging Manual (e.g. Gleason Score in Prostate). These factors are collected separately from T, N, and M, which remain purely anatomic and are used to assign stage groups.

Where non-anatomic factors are used in groupings, there is a definition of the groupings provided for cases where the non-anatomic factor is not available (X) or where it is desired to assign a group ignoring the non-anatomic factor.

Additional information and further training opportunities for registrars can be found on the AJCC Website: <https://cancerstaging.org/CSE/Registrar/Pages/default.aspx>

Effective Date

Directly coded clinical and pathologic AJCC-TNM stage is required for all cases diagnosed on **1/1/2016** and after.

AJCC Cancer Staging Manual (AJCC Manual)

The complete instructions and site-specific defined codes are documented in the *AJCC Cancer Staging Manual*.

1. Edition- It is very important the correct version of the *AJCC Manual* is used.

- Date of Diagnosis of 2016 cases- Cases diagnosed in 2016 must be coded using the 7th Edition.

The following table lists the AJCC-TNM data items required to be reported to the PCR

Data Item	Description
TNM Path T	AJCC pathologic tumor*
TNM Path N	AJCC pathologic nodes*
TNM Path M	AJCC pathologic metastases*
TNM Path Stage Grp	AJCC pathologic stage group*
TNM Path Descriptor	Identifies the AJCC pathologic stage (prefix/suffix) descriptor*
TNM Clin T	AJCC clinical tumor*
TNM Clin N	AJCC clinical nodes*
TNM Clin M	AJCC clinical metastases*
TNM Clin Stage Grp	AJCC clinical stage group*
TNM Clin Descriptor	Identifies the AJCC clinical stage (prefix/suffix) descriptor*
TNM Edition Number	A code that indicates the edition of the AJCC manual used to stage the case*

* Detailed site-specific codes for the TNM fields are defined by AJCC.

Text

Text to support all AJCC-TNM data items must be recorded in the specific text fields. See *Part Three, Data Item Instructions, Text-Path, Text-DX Proc-X-ray/Scans, Text-DX Proc-OP, and Text-DX Proc-Scopes*.

GUIDELINES FOR RECORDING FIRST COURSE OF TREATMENT

First course of treatment includes all methods of cancer-directed therapy recorded in the treatment plan and administered to the patient before disease progression or recurrence. **Never** code treatment unless you know it has actually been administered at your facility or any other facility; record as None, 00 or 0.

No therapy is a treatment option (the patient refused therapy, the family/guardian refused therapy, the patient expired before therapy started, or the physician recommended no therapy). Therefore, first course of treatment may be no treatment. Use the date the decision was made not to treat as *Date of 1st Crs Rx*.

All modalities of treatment are included regardless of sequence or degree of completion of any component method.

Treatment Plan

A treatment plan describes the cancer-directed treatment intended to modify, control, remove or destroy proliferating cancer cells. The documentation confirming a treatment plan may be fragmented. It is frequently found in several different sources, e.g., medical or clinic records, consultation reports, and outpatient records. All cancer-directed therapies specified in the physician(s) treatment plan are a part of the first course of treatment. When a treatment plan is not available or unclear, consult a physician.

A discharge plan may contain part or the entire treatment plan.

A treatment plan may specify one or more modalities of therapy (surgery, radiation, chemotherapy, hormone therapy, immunotherapy, or other therapy). A treatment “regimen” may include combinations of concurrent or adjuvant therapies.

Example: A patient had a transurethral resection diagnostic of bladder cancer. Resection was followed by Cobalt-60 radiation, ileal loop diversion, and a complete cystectomy with node dissection. Code as follows:

Data Item	Treatment Code
Surgery to Primary Site	50 - Complete cystectomy
Radiation Regional RX Modality	22- Cobalt-60 radiation
Chemotherapy	00 - None
Hormone Therapy	00 - None
Immunotherapy	00 - None
Other Treatment	0 - No other cancer-directed therapy

Guidelines for Determining *First Course of Treatment*

First course of treatment includes all cancer-directed therapy planned and administered by the physician(s) during or after the first diagnosis of cancer. Planned treatment may include multiple modes of therapy and may encompass intervals of a year or more.

Time Period Rules for First Course of Treatment for Malignancies except Leukemias (in order of precedence)

1. If there is a documented, planned first course of treatment, first course ends at the completion of this treatment plan, regardless of the duration of the treatment plan.
2. If the patient is treated according to a facility's standards of practice (established protocol), first course ends at the completion of the treatment.
3. If there is no documented treatment plan, established protocol, or management guidelines, and consultation with a physician is not possible, use the principle: "initial treatment must begin within four months of the date of initial diagnosis."
4. If the patient refuses all treatment modalities, then changes his/her mind and the treatment is initiated, consult a physician to determine if this is part of first course of treatment.

Special Rules for Leukemias

The first course of definitive treatment is related to the first *remission* as follows:

1. If a remission, complete or partial, is achieved during the first course of therapy for the leukemic process, include:
 - All definitive therapy considered as *remission-inducing* for the first remission.
 - All definitive therapy considered as *remission-maintaining* for the first remission (maintenance chemotherapy or irradiation to the central nervous system).
 - Disregard all treatment administered to the patient after the relapse of the first remission.
2. If no remission is attained during the first course of therapy, record all treatment attempted to induce the remission. Disregard all treatment administered to the patient as a subsequent attempt to induce remission.

Watchful Waiting

If a treatment plan is given for symptoms/disease progression after period of *watchful waiting*, this treatment is not considered part of first course. For example, if physician and patient choose a *wait and watch* approach to prostate cancer and the patient becomes symptomatic, consider the symptoms to be an indication the disease has progressed and any further treatment is not part of first course.

Treatment Failure

Treatment failure or disease progression may prompt the physician to stop therapy before the full course has been completed. Any therapy administered after the discontinuation of first course must be considered as secondary or subsequent treatment.

Treatment for Recurrence or Progression

Treatment for recurrence or progression of disease includes all cancer-directed therapies administered after the first course of treatment is complete.

If the patient does not respond or if the disease progresses, a physician may stop the first course of treatment before it is complete. Therapy administered after the first course ends is not recorded as first course of treatment.

Non Cancer-Directed Treatment

Non cancer-directed treatments prolong the patient's life, alleviate pain, make the patient comfortable, or prepare the patient for cancer-directed therapy. They are not meant to destroy or control the tumor or delay the spread of disease. Non cancer-directed procedures include diagnostic tests and supportive care (treatments designed to relieve symptoms and minimize the effects of the cancer). Surgical procedures performed to diagnose/stage disease (exploratory) or for relief of symptoms (palliative) are non cancer-directed surgeries. **Non cancer-directed therapies should not be coded as treatment.**

Examples of non cancer-directed therapies include:

Diagnostic procedures:

- Incisional biopsies
- Exploratory procedures/surgery with or without biopsies, such as laparotomy, cystotomy, nephrotomy, gastrotomy, thoracotomy
- Brushings, washings, aspiration of cells, and hematologic findings (peripheral blood smears) are not surgical procedures

Palliative procedures:

- Colostomy
- Nephrostomy
- Esophagostomy
- Tracheostomy
- Gastrostomy

Supportive care/relieving symptoms:

- Pain medication
- Oxygen
- Antibiotics administered for an associated infection
- Intravenous therapy to maintain fluid or nutritional balance
- Laser therapy directed at relieving symptoms

Exception: Treatment for hematopoietic diseases can be supportive care, observation, or any treatment that does not meet the usual definition in which treatment "modifies, controls, removes, or destroys proliferating cancer tissue". See *Part Three, RX Summ-Other*.

Cancer-Directed Treatment

Cancer-directed treatment is tumor directed, and its purpose is to modify, control, remove, or destroy primary or metastatic cancer tissue. Physicians administer the therapy (-ies) to remove or minimize the size of tumor or to delay the spread of disease. Record all cancer-directed therapy administered to the patient. For complete treatment information, record therapies given in other institutions and failed treatments (the patient did not respond).

Example 1: A patient is diagnosed with stage IV small cell carcinoma of the lung. The treatment plan recommends radiation to shrink the metastatic tumor and alleviate the pain caused by rib metastases. The reporting institution delivers beam radiation. The data item *Rad--Reg RX Modality* is coded 22, beam radiation, NOS.

Example 2: A patient with breast cancer enters the reporting institution for a lumpectomy. The physician's treatment plan specifies radiation therapy to the breast following surgery. It is unknown if the patient had radiation. Code the data item *RX Summ - Surg Prim Site* to a partial or less than total mastectomy (22). Record the data item *Rad--Regional RX Modality* as (00), none. If additional follow-up information reveals the patient did receive radiation, change to the appropriate radiation code.

RX SUMM - SURG PRIM SITE

Record the most invasive, definitive cancer-directed procedure performed to the primary site as part of the first course of treatment at the reporting institution and other institutions. Cancer-directed surgery modifies, controls, removes, or destroys proliferating cancer tissue.

Recording Surgery to Primary Site

1. An excisional biopsy is cancer-directed surgery.

Example: The surgeon states the procedure is an excisional biopsy, but the pathology report shows microscopic involvement of the margins. Record the code for an excisional biopsy as *Rx Summ - Surg Prim Site*.

Note: Biopsies that remove all gross tumor or leave only microscopic margins should be coded to surgery of the primary site.

2. If no cancer-directed surgery was performed, code to 00.
3. If it is unknown if cancer-directed surgery was performed, code to 99.
4. Best Information- Use the best information in the operative/pathology reports to determine the operative procedure. Do **not** depend on the name of the procedure since it may be incomplete. If the operative report is unclear as to what was excised or if there is a discrepancy between the operative and pathology reports, use the pathology report, unless there is reason to doubt its accuracy.
5. Site-Specific Surgery Codes- Refer to *Appendix K* for surgical codes.
 - Codes 10 through 18 are site-specific descriptions of tumor-destruction procedures that do not produce a pathologic specimen.
 - Codes 20 through 79 are site-specific descriptions of resection procedures.
 - Codes 80 and 90 are NOS terms and should only be used if more precise information about the surgery is not available.
6. Hierarchy-For codes 00 through 79, the descriptions of the surgical procedures are hierarchical. Last-listed responses take precedence over earlier-listed responses. (Regardless of code or numeric value).

Example: A rectosigmoid primary surgically treated by polypectomy with electrocautery, which is listed after polypectomy alone, is coded 22.

20 Local tumor excision, NOS
 26 Polypectomy
 27 Excisional biopsy
 Combination of 20 or 26-27 WITH
 21 Photodynamic therapy (PDT)
 22 Electrocautery
 23 Cryosurgery

24 Laser ablation

7. Special Code 98 applies to specific tumors that cannot be clearly defined in terms of primary or nonprimary site. Code 98 takes precedence over all other codes. Surgical Procedure of Primary Site should be coded 98 for *Unknown and Ill-defined Primary Sites and Hematopoietic/Reticuloendothelial/Immunoproliferative/Myeloproliferative Disease* (See *Part Three, General Information* for a list of these sites and conditions). The item *RX Summ--Surg Oth Reg/Dis Site* is used to indicate whether surgery was performed for these tumors.
8. Total Resection- If a surgical procedure removes the remaining portion of an organ which had been partially resected previously for any condition, code as total removal of the organ. If none of the primary organ remains, the code should indicate this is the case.

Example 1: Resection of a stomach which had been partially excised previously is coded as total removal of stomach.

Example 2: Removal of a cervical stump is coded as total removal of uterus.

Example 3: Lobectomy of a lung with a previous wedge resection is coded as total removal of lobe.
9. Biopsies that remove the entire tumor and/or leave only microscopic margins are to be coded in this item.
10. Extranodal Lymphomas- Surgery for extranodal lymphomas should be recorded using the scheme for the extranodal site.

Example: Use the scheme for the stomach to record a gastrectomy for a primary lymphoma of the stomach.
11. Surgery for Multiple Primaries- If multiple primaries are treated by a single surgical event, code the appropriate surgical items for each primary.

Example 1: If a total abdominal hysterectomy was done for a patient with two primaries, one of the cervix and one of the endometrium, code each as having had a total abdominal hysterectomy.

Example 2: If a total colectomy was done for a patient with multiple primaries in several segments of the colon, code total colectomy for each of the primary segments.
12. Regional Tissue or Organs- Surgery to remove regional tissue or organs is coded in this item only if the tissue/organs are removed in continuity with the primary site, except where noted in *Appendix K*.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

Special Instructions

1. Registry Hospitals - If you can record multiple surgical procedures in your registry software, make sure the data item transmitted to the PCR as *RX Summ - Surg Prim Site* reflects the most extensive code.
2. Abstract Plus - You can only record one surgical procedure. If you have more than one procedure use the code for the most extensive.

RX DATE - SURGERY

Record the earliest date on which the patient had cancer-directed surgery for this primary or metastatic site. This includes *RX Summ-Surg Prim Site*, *RX Summ-Scope Reg LN Surg*, and *RX Summ-Surg Oth Reg/Dis*.

Recording RX Date-Surgery

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.

2. **No Surgery of Primary Site**- This data item may contain a date even when surgery to the primary site equals 00 (none).

Example: Patient has excision of a brain lesion on January 15, 2010; final pathology diagnosis is metastatic lung carcinoma. Patient refuses further work-up.

RX Summ - Surg Prim Site code = 00

RX Date - Surgery = 20100115

RX Summ - Surg Oth Reg/Dis = 4

3. **Collecting the dates** for each treatment modality allows sequencing of multiple treatments and aids evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
4. **Exact Date Unavailable**- If the exact date of cancer-directed surgery is not available, record an approximate date. Refer to *Part Three, General Information*.
5. **Leave RX Date- Surgery Blank** and assign the appropriate *RX Date-Surgery Flag* for the following reasons:
 - a. when no cancer-directed surgery is performed
 - b. when it is unknown if any cancer-directed surgery was performed
 - c. when the patient had surgery but the date is unknown
 - d. when the record was identified by death certificate only
 - e. when it is an autopsy-only case

Special Instructions

1. **Registry Hospitals** - If you can record multiple surgery dates, make sure the data item transmitted to the PCR as *RX Date-Surgery* reflects the earliest date of cancer-directed surgery. Also, some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.
2. **Abstract Plus** - You can only record one date and one surgical procedure. If you have more than one procedure, use the date of the first, but the code for the most invasive, extensive surgical procedure performed during the first course of treatment.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

RX DATE-SURGERY FLAG

Record the date flag in the event a complete *RX Date-Surgery* was not entered to explain why.

Recording RX Date-Surgery Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Surgery* has a full or partial date recorded.
2. Unknown if Surgery Performed- Code 10 if it is unknown whether any surgery was performed.
3. No Surgery- Code 11 if no surgical procedure was performed.
4. Unknown Date- Code 12 if the *RX Date-Surgery* cannot be determined at all, but the patient did receive first course surgery.
5. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
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 (for example, <i>RX Date-Surgery</i> is unknown)
(blank)	A valid date value is provided in item <i>RX Date-Surgery</i>

The following table illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Surgery* and *RX Date-Surgery Flag*. In the table below, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Surgery (MMDDCCYY)	Interoperable RX Date-Surgery (CCYYMMDD)	RX Date-Surgery Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any surgery done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No Surgery Performed	00000000 (example: 00000000)	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, surgery performed	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software

RX DATE--MOST DEFIN SURG

Records the date of the most definitive surgical procedure of the primary site performed as part of the first course of treatment. By definition, this would be the date on which the surgery coded in Surgical Procedure of Primary Site was performed at this or any facility.

Recording RX Date-Most Definitive Surgery

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1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.

2. Collecting this date enables the measurement of the lag time between diagnosis and the most definitive surgery of the primary site.
3. Exact Date Unavailable- If the exact date of cancer-directed surgery is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date- Most Defin Surg Blank and assign the appropriate *RX Date- Most Defin Surg Flag* for the following reasons:
 - a. when no surgery to the primary site is performed
 - b. when it is unknown if surgery to the primary site was performed
 - c. when the patient had surgery to the primary site but the date is unknown
 - d. when the record was identified by death certificate only
 - e. when it is an autopsy-only case

RX DATE-MOST DEFIN SURG FLAG

Record the date flag explain why a complete *RX Date-Most Definitive Surgery* was not entered.

Recording RX Date-Most Definitive Surgery Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Most Definitive Surg* has a full or partial date.
2. Unknown if Surgery Performed- Code 10 if it is unknown whether any surgery was performed.
3. No Surgery- Code 11 if no surgery to the primary site was performed.
4. Unknown Date- Code 12 if the *RX Date-Most Definitive Surg* cannot be determined at all, but the patient did receive surgery to the primary site.
5. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any surgery to the primary site was performed)
11	No proper value is applicable in this context (for example, no surgery to the primary site was performed)
12	A proper value is applicable but not known (for example, <i>RX Date-Most Definitive Surgery</i> is unknown)
(blank)	A valid date value is provided in item <i>RX Date-Most Definitive Surgery</i>

The following table illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Most Definitive Surg* and *RX Date-Most Definitive Surg Flag*. In the table below, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Most Definitive Surg (MMDDCCYY)	Interoperable RX Date-Most Definitive Surg (CCYYMMDD)	RX Date-Most Definitive Surg Flag
Full date known	MMDDCCYY Example: 04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any surgery done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No Surgery Performed	00000000 (example: 00000000)	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, surgery performed	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12

RX SUMM - SCOPE REG LN SURG

Record the code for the removal, biopsy or aspiration of regional lymph node(s) performed as first course of treatment. This includes treatment given at all facilities as part of the first course of treatment.

Codes and Definitions

The following instructions should be applied to all surgically treated cases for all types of cancers. The treatment of breast and skin cancer is where the distinction between sentinel lymph node biopsies (SLNBx) and more extensive dissection of regional lymph nodes is most frequently encountered. For all other sites, non-sentinel regional node dissections are typical, and codes 2, 6 and 7 are infrequently used.

Code	Label	General Instructions for All Sites	Additional Instructions for Breast Primaries
		Use the operative report as the primary source document to determine whether the operative procedure was a sentinel lymph node biopsy (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.	Use the operative report as the primary source document to determine whether the operative procedure was a sentinel lymph node biopsy (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.
0	<i>None</i>	No regional lymph node surgery. No lymph nodes found in pathologic specimen. Diagnosed at autopsy.	
1	<i>Biopsy or aspiration of regional lymph node, NOS</i>	Review the operative report of to confirm whether an excisional biopsy or aspiration of regional lymph nodes was actually performed. If additional procedures were performed on the lymph nodes, use the appropriate code 2-7.	Excisional biopsy or aspiration of regional lymph nodes for breast cancer is uncommon. Review the operative report of 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	Label	General Instructions for All Sites	Additional Instructions for Breast
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			Primaries
2	<i>Sentinel lymph node biopsy</i>	<ul style="list-style-type: none"> The operative report states that a SLNBx was performed. Assign Code 2 <i>Sentinel lymph node biopsy</i> when 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. When a SLNBx is performed, additional non-sentinel nodes can be taken during the same operative procedure. These additional nonsentinel 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 <i>Sentinel lymph node biopsy</i> (code 2). If review of the operative report confirms that a regional lymph node dissection followed the SLNBx, assign code 6, <i>Sentinel node biopsy and code 3, 4, or 5 at same time, or timing not stated</i> 	<ul style="list-style-type: none"> 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. Code these cases as 2, <i>Sentinel lymph node biopsy</i> if no ALND was performed, or 6, <i>Sentinel node biopsy and code 3, 4, or 5 at same time, or timing not stated</i> when ALND was performed during the same operative event.
3	<i>Number of regional nodes removed unknown or not stated; regional lymph nodes removed NOS</i>	<ul style="list-style-type: none"> 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 (<i>Number of regional lymph nodes removed unknown, not stated; regional lymph nodes removed, NOS</i>). 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). 	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).
4	<i>1-3 regional lymph nodes removed</i>	<ul style="list-style-type: none"> Code 4 (<i>1-3 regional lymph nodes removed</i>) should be used infrequently. Review the operative report to ensure the procedure was not a SLNBx only. 	
5	<i>4 or more regional lymph nodes removed</i>	<ul style="list-style-type: none"> Code 5 (<i>4 or more regional lymph nodes removed</i>). If a relatively small number of nodes were 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 (6 or 7). 	

Code	Label	General Instructions for All Sites	Additional Instructions for Breast
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			Primaries
6	<i>Sentinel node biopsy and code 3, 4, or 5 at same time, or timing not stated</i>	<ul style="list-style-type: none"> • SNLBx 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 SNLBx 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. 	<ul style="list-style-type: none"> • 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
7	<i>Sentinel node biopsy and code 3, 4, or 5 at different times</i>	<ul style="list-style-type: none"> • SNLBx and regional lymph node dissection (code 3, 4, or 5) in separate surgical events. • Generally, SLNBx followed by 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 SLNBx only. 	<ul style="list-style-type: none"> • 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 SLNBx only, or whether a SLNBx plus an ALND was performed.
9	<i>Unknown or not applicable</i>	<ul style="list-style-type: none"> • 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. • The status of regional lymph node evaluation should be known for surgically treated cases (cases coded 19-90 in the data item <i>Surgery of Primary Site</i>). Review surgically treated cases coded 9 in <i>Scope of Regional Lymph Node Surgery</i> to confirm the code. 	

Recording Scope of Regional Lymph Node Surgery

1. Regional Lymph Node List - Refer to all codes listed in the CS Lymph Node code tables in each site specific schema of *Collaborative Stage Data Collection System Version 2 (CSv2)* to identify site-specific regional lymph nodes. If a lymph node is not listed in CS Lymph Node code table, check an anatomy book or medical dictionary for a synonym. If the lymph node chain and its synonym are not listed in CS Lymph Nodes, code the lymph node surgery in *RX Summ - Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s)*.
2. Aspiration, biopsy or removal of lymph nodes- Record surgical procedures which aspirate, biopsy, or remove regional lymph nodes in an effort to diagnose or stage disease.
3. Minimum number- There is no minimum number of nodes that must be removed; code to the farthest regional lymph nodes removed regardless of involvement with disease (e.g., the biopsy of contralateral lung lymph nodes).
4. Hierarchy- Codes 0-7 are hierarchical. Code the procedure numerically higher.
5. Meninges, brain, spinal cord, cranial nerves and other parts of the central nervous system- For primaries of the meninges, brain, spinal cord, cranial nerves and other parts of the central nervous system (C70.0-C70.9, C71.0-C71.9, C72.0-C72.9, C75.1-C75.3), code to 9.
6. Lymphoma- For lymphomas with a lymph node primary site, code 9. For extranodal lymphomas, refer to the site-specific codes for the primary site.
7. Unknown or ill defined primary site or for hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative disease, code to 9. See *General Information* for a list of these sites and conditions.
8. Distant Lymph Nodes- Do not code *distant* lymph nodes removed during surgery to the primary site for this data item. Distant nodes are coded in the data field *Surgical Procedure/Other Site*.
9. No regional lymph nodes- This data item may not be blank. If no regional lymph nodes were removed or no surgery was performed, record 0.

Example 1: Aspiration of regional lymph node of a pharynx primary to confirm histology of widely metastatic disease is coded to 1.

Example 2: There was an attempt at sentinel lymph node dissection for a breast primary, but no lymph nodes were found in the pathological specimen, code to 2

Example 3: Patient has melanoma of the back. A sentinel lymph node dissection was done with the removal of one lymph node. This node was negative for disease, code to 2.

Example 4: Bilateral pelvic lymph node dissection for prostate cancer, code to 3.

Example 5: A patient with a breast primary has a sentinel lymph node biopsy of the right axilla, followed by right axillary lymph node dissection during the same surgical event, code to 6.

Example 6: Sentinel lymph node biopsy (SLNBx) of left axilla, followed in a second procedure 5 days later by a left axillary lymph node dissection (ALND) for a breast primary, code to 7.

Special Instructions

1. Registry Hospitals - If you can record multiple surgical procedures in your registry software, make sure the data item transmitted to the PCR as *RX Summ - Scope Reg LN Surg* reflects most extensive code.
2. Abstract Plus - You can only record one surgical procedure. If you have more than one procedure use the code for the most extensive (numerically highest).

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

RX SUMM - SURG OTH REG/DIS

Record the code for the type of surgery to sites other than the primary site and regional lymph nodes, performed as part of the first course of treatment. This includes treatment given at all facilities as part of the first course of treatment.

This data item describes the removal of tissues and organs other than the primary tumor or organ of origin. This data item is for all procedures that do not meet the definition of *RX Summ- Surgery of Primary Site* or *RX Summ - Scope of Regional Lymph Node Surgery*.

Codes and Definitions

Code	Definition
0	<i>None</i> - No surgical procedure of nonprimary site was performed. Diagnosed at autopsy.
1	<i>Nonprimary surgical procedure performed</i> - Nonprimary surgical resection to other site(s), unknown if the site(s) is regional or distant.
2	<i>Nonprimary surgical procedure to other regional sites</i> - Resection of regional site.
3	<i>Nonprimary surgical procedure to distant lymph node(s)</i> -Resection of distant lymph node(s).
4	<i>Nonprimary surgical procedure to distant site</i> - Resection of distant site.
5	<i>Combination of codes</i> - Any combination of surgical procedures 2, 3, or 4.
9	<i>Unknown</i> - It is unknown whether any surgical procedure of a nonprimary site was performed. Death certificate only.

Recording Surgery to Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s)

1. Hierarchy- The codes are hierarchical. Code the procedure numerically higher in this data item.
2. Suspicion of malignancy- Code the removal of non-primary tissue the surgeon suspected was involved with the malignancy even if the pathology is negative.
3. Unknown or ill-defined primary sites or for hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative disease- Assign code 1 if any surgical procedure is performed to treat these cases. See *Part Three, General Information* for a list of these sites and conditions.

Example: Surgical biopsy of metastatic lesion from liver; unknown primary is coded to 1.

4. Incidental removal- Do not code the incidental removal of tissue. Incidental is defined as tissue removed for reasons other than the malignancy.

Example: The incidental removal of the appendix during a surgical procedure to remove a primary malignancy in the right colon is not coded therefore; this case would be coded to 0.

5. No surgery- This data item may not be blank. If there was no surgery to other regional sites, distant sites or distant lymph nodes, record 0.

Special Instructions

1. Registry Hospitals - If you can record multiple surgical procedures in your registry software, make sure the data item transmitted to the PCR as *RX Summ - Surg Oth Reg/Dis* reflects the most extensive (numerically highest) code.
2. Abstract Plus - You can only record one surgical procedure. If you have more than one procedure use the code for the most extensive (numerically highest).

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

REASON FOR NO SURGERY

Record the reason for no Surgery of Primary Site. Codes 1-9 are valid only when *RX Summ - Surg Prim Site* is coded 00.

Codes and Definitions

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 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 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.

Recording Reason for No Surgery

1. No surgery in plan- Code 1 if the treatment plan offered multiple options and the patient selected treatment that did not include surgery of the primary site, or if the option of "no treatment" was accepted by the patient.
2. If Surgical Procedure of Primary Site is coded 98, code *Reason for No Surgery* to 1.
3. Patient refused- If the patient refused recommended surgical treatment, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 7.

4. Unknown treatment- If the treatment plan offered multiple choices, but it is unknown which treatment, if any was provided, code to 9.

Example 1: A patient with a primary tumor of the liver is not recommended for surgery due to advanced cirrhosis, code to 2.

Example 2: A patient is referred to another facility for recommended surgical resection of a gastric carcinoma, but further information from the facility to which the patient was referred is not available, code to 8.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

RAD- REGIONAL RX MODALITY

Record the dominant modality of radiation therapy used to deliver the most clinically significant regional dose to the primary volume of interest during the first course of treatment.

Codes and Definitions

Code	Definition
00	<i>No radiation treatment-</i> Radiation therapy was not administered to the patient. Diagnosis at autopsy
20	<i>External beam, NOS-</i> The treatment is known to be by external beam, but there is insufficient information to determine the specific modality.
21	<i>Orthovoltage-</i> External beam therapy administered using equipment with a maximum energy of less than one (1) million volts (MV). Orthovoltage energies are typically expressed in units of kilovolts (kV).
22	<i>Cobalt-60, Cesium-137-</i> External beam therapy using a machine containing either a Cobalt-60 or Cesium-137 source. Intracavitary use of these sources is coded either 50 or 51.
23	<i>Photons (2–5 MV) -</i> External beam therapy using a photon producing machine with a beam energy in the range of 2–5 MV.
24	<i>Photons (6–10 MV) -</i> External beam therapy using a photon producing machine with a beam energy in the range of 6–10 MV.
25	<i>Photons (11–19 MV) -</i> External beam therapy using a photon producing machine with a beam energy in the range of 11–19 MV.
26	<i>Photons (>19 MV) -</i> External beam therapy using a photon producing machine with a beam energy of more than 19 MV.
27	<i>Photons (mixed energies) -</i> External beam therapy using more than one energy over the course of treatment.
28	<i>Electrons-</i> Treatment delivered by electron beam.
29	<i>Photons and electrons mixed-</i> Treatment delivered using a combination of photon and electron beams.
30	<i>Neutrons, with or without photons/electrons-</i> Treatment delivered using neutron beam.
31	<i>IMRT -</i> Intensity modulated radiation therapy, an external beam technique that should be clearly stated in patient record.
32	<i>Conformal or 3-D therapy-</i> An external beam technique using multiple, fixed portals shaped to conform to a defined target volume. Should be clearly described as conformal or 3-D therapy in patient record.
40	<i>Protons-</i> Treatment delivered using proton therapy.
41	<i>Stereotactic radiosurgery, NOS-</i> Treatment delivered using stereotactic radiosurgery, type not specified in patient record.
42	<i>Linac radiosurgery -</i> Treatment categorized as using stereotactic technique delivered with a linear accelerator.
43	<i>Gamma Knife-</i> Treatment categorized as using stereotactic technique delivered using a Gamma Knife machine.

Code	Definition
50	<i>Brachytherapy, NOS</i> - Brachytherapy, interstitial implants, molds, seeds, needles, or intracavitary applicators of radioactive materials not otherwise specified.
51	<i>Brachytherapy, Intracavitary, LDR</i> - Intracavitary (no direct insertion into tissues) radioisotope treatment using low dose rate applicators and isotopes (Cesium-137, Fletcher applicator).
52	<i>Brachytherapy, Intracavitary, HDR</i> - Intracavitary (no direct insertion into tissues) radioisotope treatment using high dose rate after-loading applicators and isotopes.
53	<i>Brachytherapy, Interstitial, LDR</i> - Interstitial (direct insertion into tissues) radioisotope treatment using low dose rate sources.
54	<i>Brachytherapy, Interstitial, HDR</i> - Interstitial (direct insertion into tissues) radioisotope treatment using high dose rate sources.
55	<i>Radium</i> - Infrequently used for low dose rate (LDR) interstitial and intracavitary therapy
60	<i>Radioisotopes, NOS</i> - Iodine-131, Phosphorus-32, etc.
61	<i>Strontium-89</i> -Treatment primarily by intravenous routes for bone metastases.
62	<i>Strontium-90</i>
80*	<i>Combination modality, specified*</i> - Combination of external beam radiation and either radioactive implants or radioisotopes* This is a converted code and should not be coded for cases diagnosed on or after 1/1/2003.
85*	<i>Combination modality, NOS*</i> - Combination of radiation treatment modalities not specified in code 80.* This is a converted code and should not be coded for cases diagnosed on or after 1/1/2003.
98	<i>Other, NOS</i> -Radiation therapy administered, but the treatment modality is not specified or is unknown.
99	<i>Unknown</i> - It is unknown whether radiation therapy was administered. Death certificate only.

*For cases diagnosed prior to January 1, 2003, the codes reported in this data item describe any radiation administered to the patient as part or the entire first course of therapy. Codes 80 and 85 describe specific converted descriptions of radiation therapy and should not be used to record regional radiation for cases diagnosed on or after January 1, 2003.

Recording Radiation Regional Treatment Modality

1. Finding radiation information- 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 regional and boost and determination of the respective treatment modality may require assistance from the radiation oncologist to ensure consistent coding.
2. Regional vs. Boost- Radiation treatment is frequently delivered in two or more phases which can be summarized as "regional" and "boost" treatments.
 - a. Regional Radiation is directed at the cancer site and a larger area of surrounding tissue.

- b. Boost Radiation is a supplemental radiation dose targeted directly to the tumor site (or site of the original tumor). It is provided to a smaller area within the same volume as regional, in order to enhance the effect of the regional treatment.

The PCR only requires Regional Radiation to be reported.

3. Only one modality delivered- If only one radiation treatment modality is delivered to a patient and it is not specified as either regional or boost treatment, assume its regional treatment and code accordingly.
4. Multiple radiation modalities- In the event multiple radiation therapy modalities were employed in the treatment of the patient, record only the dominant modality.
5. Boost Treatment- In some circumstances, the boost treatment may precede the regional treatment.
6. Radioembolization- Code radioembolization as brachytherapy.
7. Brachytherapy with 125 seeds- Assign code 53 for brachytherapy with 125 seeds. Seeds are always low dose therapy because they are left in place and the radioactivity decays over time.
8. Terms- For purposes of this data item, photons and x-rays are equivalent.

Example 1: Patient receives 15 MV external pelvic treatments to 4,500 cGy for cervical carcinoma, and then receives two Fletcher intracavitary implants is coded to 25.

Example 2: A patient with carcinoma of the parotid receives daily treatments of which 60% are delivered by 15 MV photons and 40% of the dose is delivered by 16 MV electrons is coded to 29.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Radiation (Beam)* or *RX Text - Radiation Other*.

RX DATE - RADIATION

Record the date radiation started.

Recording RX Date- Radiation

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.
2. Collecting dates for each treatment modality allows sequencing of multiple treatments and evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
3. Exact Date Unavailable- If the exact date the radiation started is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date- Radiation Blank and assign the appropriate *RX Date-Radiation Flag* for the following reasons:
 - a. when no radiation is given
 - b. when radiation is planned, but has not yet started
 - c. when it is unknown if any radiation was given
 - d. when the patient received radiation but the date is unknown
 - e. when the record was identified by death certificate only
 - f. when it is an autopsy-only case

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Radiation (Beam)* or *RX Text - Radiation Other*.

Special Instructions for Registry Hospitals

Some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.

RX DATE-RADIATION FLAG

Record the date flag in the event a complete *RX Date-Radiation* was not entered to explain why.

Recording RX Date-Radiation Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Radiation* has a full or partial date recorded.
2. Unknown if Radiation Given- Code 10 if it is unknown whether any radiation was given.
3. No Radiation- Code 11 if no radiation is planned or given.
4. Unknown Date- Code 12 if *RX Date-Radiation* cannot be determined at all, but the patient did receive radiation.
5. Radiation Planned but Not Started- Code 15 if radiation is planned, but has not yet started and the start date is not yet available. If radiation is later started, update this item, *RX Date-Radiation*, and *Rad- Regional RX Modality* in your database and submit a change sheet to the PCR. See *Part One, Changing Information*.
6. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

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 given)
12	A proper value is applicable but not known (for example, <i>RX Date-Radiation</i> is unknown)
15	Information is not available at this time, but it is expected 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 the case was abstracted)
(blank)	A valid date value is provided in item <i>RX Date-Radiation</i>

The table on the next page illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Radiation* and *RX Date-Radiation Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Radiation (MMDDCCYY)	Interoperable RX Date-Radiation (CCYYMMDD)	RX Date-Radiation Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any radiation done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No radiation given	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, radiation given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12
Radiation not yet started	88888888 Example: 88888888	bbbbbbbb Example: bbbbbbbb	15

b=blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software

RX SUMM- SURG/RAD SEQ

Record the sequencing of radiation and surgical procedures given as part of first course of treatment.

The sequence of radiation and surgical procedures given as part of first course of treatment cannot always be determined using the date on which each modality was started or performed. This data item can be used to more precisely evaluate the timing of delivery of treatment to the patient.

Codes and Definitions

Code	Definition
0	<p><i>No radiation therapy and/or surgical procedures; unknown if surgery and/or radiation given-</i> No radiation therapy given; and/or no surgery of the primary site; no scope of regional lymph node surgery; no surgery to other regional site(s), distant site(s), or distant lymph node (s) or it is unknown whether any surgery and/or radiation was given. Diagnosed at autopsy.</p> <p><i>Example:</i> Due to other medical conditions surgery was not performed.</p>
2	<p><i>Radiation therapy before surgery-</i> Radiation therapy given before surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A patient has a large lung lesion and received radiation therapy prior to resection.</p>
3	<p><i>Radiation therapy after surgery-</i> Radiation therapy given after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A patient received a wedge resection of a right breast mass with axillary lymph node dissection followed by radiation to the right breast.</p>
4	<p><i>Radiation therapy both before and after surgery-</i> At least two courses of radiation therapy are given before and after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> Preoperative radiation was given to a large, bulky vulvar lesion and was followed by a lymph node dissection. This was then followed by radiation therapy to treat positive lymph nodes.</p>
5	<p><i>Intraoperative radiation therapy-</i> Intraoperative therapy given during surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A cone biopsy of the cervix is followed by intracavitary implant for IIIB cervical carcinoma.</p>

Code	Definition
6	<p><i>Intraoperative radiation therapy with other therapy administered before and/or after surgery-</i> Intraoperative radiation therapy given during surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node (s) with other radiation administered before and/or after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node (s).</p> <p><i>Example:</i> Stage IV vaginal carcinoma was treated with 5,000 cGy to the pelvis followed by a lymph node dissection and 2,500 cGy of intracavitary brachytherapy.</p>
7	<p><i>Surgery both before and after radiation-</i> Radiation therapy given before and after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p>
9	<p><i>Sequence unknown, but both surgery and radiation were given-</i> Administration of radiation therapy and surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s) were performed and the sequence of the treatment is not stated in the patient record.</p> <p><i>Example:</i> An unknown primary of the head and neck was treated with surgery and radiation prior to admission, but the sequence is unknown.</p>

Recording RX Summ-Surg/Rad Seq

1. Surgical procedures include *RX Summ- Surg Prim Site; RX Summ- Scope LN Surg; RX Summ- Surg Oth Reg/Dis*
2. No Surgery- If all surgery procedures listed above are coded to 0, then this item should be coded to 0.
3. Unknown if Surgery or Radiation Given- If it is unknown if surgery and/or radiation were given, this item should be coded to 0.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery, RX Text - Radiation (Beam) and RX Text - Radiation Other.*

REASON FOR NO RADIATION

Record the reason no regional radiation therapy was administered to the patient. Codes 1-9 are valid only when *Rad- Regional RX Modality* is coded 00.

Codes and Definitions

Code	Definition
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 recommended/administered because it was contraindicated due to other patient risk factors (comorbid conditions, advanced age, progression of tumor prior to planned radiation etc.).
5	Radiation therapy was not administered because the patient died prior to planned or recommended therapy.
6	Radiation therapy was not administered; it was recommended by the patient's physician, but was not administered as part of first course treatment. No reason was noted in patient 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 patient record.
8	Radiation therapy was recommended, but it is unknown whether it was administered.
9	It is unknown if radiation therapy was recommended or administered. Death certificate and autopsy cases only.

Recording Reason for No Radiation

1. No radiation therapy in plan- Code 1 if the treatment plan offered multiple options and the patient selected treatment that did not include radiation therapy, or if the option of "no treatment" was accepted by the patient.

Example: A patient with Stage I prostate cancer is offered either surgery or brachytherapy to treat his disease. The patient elects to be surgically treated, code to 1

2. Patient refused- If the patient refused recommended radiation therapy, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 7.

3. Recommended, but unknown if given- If it is known that a physician recommended radiation treatment, but no further documentation is available to confirm its administration, code to 8.
4. Unknown treatment- If the treatment plan offered multiple choices, but it is unknown which treatment, if any was provided, code to 9.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text – Radiation (Beam)* and *RX Text Radiation Other* for instructions.

RX SUMM - CHEMO

Record the type of chemotherapy administered as first course of treatment at your institution and at all other institutions. If chemotherapy was not administered, then this item records the reason it was not administered to the patient. Chemotherapy consists of a group of anticancer drugs that inhibit the reproduction of cancer cells by interfering with DNA synthesis and mitosis.

Codes and Definitions

Code	Definition
00	<i>None</i> - Chemotherapy was not part of the planned first course of therapy. Diagnosed at autopsy.
01	<i>Chemotherapy NOS</i> - 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 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. Death certificate only.

Recording Chemotherapy

1. Chemotherapy not usually given for this condition- If chemotherapy was not administered to the patient, and it is known it is not usually administered for this stage of cancer or type of condition, code to 00.
2. Patient did not select chemotherapy- If the treatment plan offered multiple options, and the patient selected treatment that did not include chemotherapy, code to 00.
3. Chemotherapy usually given for this condition- If it is known 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.
4. Chemoembolization- Code as chemotherapy when the embolizing agent(s) is a chemotherapy drug(s).
5. Patient refused- If the patient refused recommended chemotherapy, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 87.
6. Unknown- If it is not known whether chemotherapy 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 to 99.
7. Change to regimen- If the managing physician changes one of the agents in a combination regimen and the replacement agent belongs to a different group (See *Part Three, Chemotherapy Group Classifications*) than the original agent, the new regimen represents the start of subsequent therapy, and *only the original agent or regimen is recorded as first course therapy*.

Example: The physician documents a multimodality treatment plan that includes a combination regimen of chemotherapy. Velban is one of the drugs in the chemotherapy regimen. After two cycles of chemotherapy, the physician says the Velban will be replaced with Oncovin and the chemotherapy will continue as planned. This is a continuation of the planned first course of therapy since they are in the same group.

8. List of chemotherapeutic agents- Use *SEER RX* to determine if a drug is a chemotherapy agent. *SEER RX* is an interactive antineoplastic drug data base and it can be downloaded from this website:

<http://seer.cancer.gov/tools/seerrx/>

Methods of Administration

Method	Definition
Intravenous (IV) Infusion	A small plastic needle is inserted into a vein. Chemotherapy flows from the IV bag/bottle, through the needle and catheter into the bloodstream.
Orally	Medication taken in the form of either a pill or liquid taken by mouth.
Intrathecal	Administered directly into the cerebrospinal fluid through a lumbar puncture needle into an implanted access device (e.g., Ommaya reservoir).
Pleural/pericardial	Injected directly into pleural or pericardial space to control malignant effusions.
Intraperitoneal	Injected into the peritoneal cavity.
Hepatic artery	Injected into a catheter inserted into artery that supplies blood to liver.

Clarification of Terms

Term	Definition
Adjuvant chemotherapy	<p>Chemotherapy given after other methods have destroyed the clinically detectable cancer cells. Chemotherapy given to destroy micrometastases (undetectable cancer cells). The intent is to prevent or delay a recurrence.</p> <p><i>Example:</i> The patient has breast cancer with positive nodes. The patient is clinically free of disease after a modified radical mastectomy. The patient is treated with adjuvant chemotherapy to prevent or delay disease recurrence.</p>
Multimodality therapy Combined modality therapy Concurrent therapy	Chemotherapy given before, during, or after other treatment modalities (surgery, radiation) as a part of the treatment plan.
Neo-adjuvant therapy	<p>Given prior to surgical resection or radiation therapy to reduce the bulk of a locally advanced primary cancer.</p> <p><i>Example:</i> A patient with locally advanced breast cancer receives chemotherapy to reduce tumor size. Chemotherapy is followed by a modified radical mastectomy.</p>
Treatment cycles	Chemotherapy agents are administered in treatment cycles, either singly or in a combination regimen of two or more chemotherapy drugs. The interval of a treatment cycle varies and chemotherapy may be administered for several weeks or several years.

Chemotherapy Group Classifications

Group	Subgroup	Example
Alkylating agents	Nitrogen mustard	Mechlorethamine (Mustargen), phenylalanine mustard (Melphalan), chlorambucil (leukeran), cyclophosphamide (Cytosan)
	Ethylenimine derivatives	Triethylene-thiophosphoramide (Thio-TEPA)
	Alkyl sulfonates	Busulfan (Myleran)
	Nitrosoureas	Carmustine (Lomustine)
	Triazines	DTIC (Dacarbazine)
Antimetabolites	Folic acid analogues	Methotrexate (Amethopterin, MTX)
	Pyrimidine analogues	5-fluorouracil (5-FU)
	Purine analogues	6-mercaptopurine (6-MP)
Natural products	Anti-tumor	Dactinomycin (Actinomycin D), doxorubicin (Adriamycin), daunorubicin (Daunomycin), bleomycin (Blenoxane), mitomycin C (Mutamycin)
	Plant alkaloids	Vinblastine (Velban, VBL), vincristine (Oncovin, VCR)
	Enzymes	l-asparaginase (Elspar)
Miscellaneous		Cis-diammine dichloroplatinum II (Cisplatin), hydroxyurea (Hydrea), procarbazine (Matulane)

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Chemo*.

RX DATE-CHEMO

Record the date chemotherapy started.

Recording RX Date- Chemo

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.
2. Collecting dates for each treatment modality allows sequencing of multiple treatments and evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
3. Exact Date Unavailable- If the exact date the chemotherapy was started is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date-Chemo Blank and assign the appropriate *RX Date-Chemo Flag* for the following reasons:
 - a. when no chemotherapy is given
 - b. when chemotherapy is planned, but has not yet started
 - c. when it is unknown if any chemotherapy was given
 - d. when the patient received chemotherapy but the date is unknown
 - e. when the record was identified by death certificate only
 - f. when it is an autopsy-only case

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Chemotherapy*.

Special Instructions for Registry Hospitals

Some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.

RX DATE-CHEMO FLAG

Record the date flag in the event a complete *RX Date-Chemo* was not entered to explain why.

Recording RX Date-Chemo Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Chemo* has a full or partial date recorded.
2. Unknown if Chemotherapy Given- Code 10 if it is unknown whether any chemotherapy was given.
3. No Chemotherapy- Code 11 if no chemotherapy is planned or given.
4. Unknown Date- Code 12 if the *RX Date-Chemo* cannot be determined at all, but the patient did receive chemotherapy.
5. Chemotherapy Planned but Not Started- Code 15 if chemotherapy is planned, but has not yet started and the start date is not yet available. If chemotherapy is later started, update this item, *RX Date-Chemo*, and *RX Summ-Chemo* in your database and submit a change sheet to the PCR. See *Part One, Changing Information*.
6. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any chemotherapy given.)
11	No proper value is applicable in this context (for example, no chemotherapy given).
12	A proper value is applicable but not known (for example, <i>RX Date-Chemo</i> is unknown)
15	Information is not available at this time, but it is expected that it will be available later (for example, chemotherapy therapy is planned as part of the first course of therapy, but had not been started at the time the case was abstracted).
(blank)	A valid date value is provided in item <i>RX Date-Chemo</i>

The table on the following page illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Chemo* and *RX Date-Chemo Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Chemotherapy (MMDDCCYY)	Interoperable RX Date-Chemotherapy (CCYYMMDD)	RX Date-Chemotherapy Flag
Full date known	MMDDCCYY Example: 04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any chemotherapy done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No chemotherapy given	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, chemotherapy given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12
Chemotherapy not yet started	88888888 Example: 88888888	bbbbbbbb Example: bbbbbbbb	15

b= blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

RX SUMM - HORMONE

Record the type of hormone therapy the patient received as a part of first course of treatment at your institution and all other institutions. If hormone therapy was not administered, then this item records the reason it was not administered to the patient. Hormone therapy consists of a group of drugs that may affect the long-term control of a cancer’s growth. It is not usually used as a curative measure.

Hormone therapy achieves its effect on cancer tissue through change of the hormone balance. Included are the administration of hormones, agents acting via hormonal mechanisms, antihormones, and steroids.

Codes and Definitions

Code	Definition
00	None- hormone therapy was not part of the planned first course of therapy. 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, a patient’s family member, or the patient’s guardian. The refusal was noted in 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. Death certificate only.

Recording Hormone Therapy

1. All sites (primary and metastatic) - Hormones, agents acting via hormonal mechanisms, and antihormones (cancer-directed only) are to be coded for all sites (primary and metastatic).
2. Prednisone
 - a. Record prednisone as hormonal therapy when administered in combination with chemotherapy, such as MOPP (mechlorethamine, vincristine, procarbazine, prednisone) or COPP (cyclophosphamide, vincristine, procarbazine, prednisone).
 - b. Do not code prednisone as hormone therapy when it is administered for reasons other than cancer treatment.

Example 1: A patient has advanced lung cancer with metastases to the brain. The physician orders Decadron to reduce the edema in the brain and relieve the neurological symptoms. Decadron is not coded as hormone therapy.

Example 2: A patient with advanced disease is given prednisone to stimulate the appetite and improve nutritional status. Do not code the prednisone as hormone therapy.

3. Hormone replacement therapy- Tumor involvement or treatment may destroy hormone-producing tissue. Hormone replacement therapy will be given if the hormone is necessary to maintain normal metabolism and body function. Do not code hormone replacement therapy as part of first course therapy.

Example: Patients with breast cancer may be treated with aminoglutethimide (Cytadren, Elipten), which suppresses the production of glucocorticoids and mineralocorticoids. These patients must take glucocorticoid (hydrocortisone) and may also need a mineralocorticoid (Florinef) as a replacement therapy. Code *Rx Summ- Hormone* to 00, None.
4. Hormone therapy not usually given for this condition- If hormone therapy was not administered to the patient, and it is known it is not usually administered for this type and stage of cancer, code to 00.
5. Patient selected treatment option without hormone therapy- If the treatment plan offered multiple options, and the patient selected treatment that did not include hormone therapy, code to 00.
6. Thyroid replacement therapy- Code 01 for thyroid replacement therapy which inhibits TSH (thyroid-stimulating hormone). TSH is a product of the pituitary gland that can stimulate tumor growth.
7. Hormone therapy usually given for this condition- If it is known hormone therapy 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.
8. Patient refused- If the patient refused recommended hormone therapy, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 87.
9. Unknown- If it is not known whether hormone therapy 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 to 99.

10. List of hormonal agents- Use *SEER RX* to determine if a drug is a hormonal agent. *SEER RX* is an interactive antineoplastic drug data base and it can be downloaded from this website:

<http://seer.cancer.gov/tools/seerrx/>

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Hormone*.

RX DATE-HORMONE

Record the date hormone therapy started.

Recording RX Date- Hormone

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.
2. Collecting dates for each treatment modality allows sequencing of multiple treatments and evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
3. Exact Date Unavailable- If the exact date the hormone therapy started is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date-Hormone Blank and assign the appropriate *RX Date-Hormone Flag* for the following reasons:
 - a. when no hormone therapy is given
 - b. when hormone therapy is planned, but has not yet started
 - c. when it is unknown if any hormone therapy was given
 - d. when the patient received hormone therapy but the date is unknown
 - e. when the record was identified by death certificate only
 - f. when it is an autopsy-only case

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Hormone*.

Special Instructions for Registry Hospitals

Some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.

RX DATE-HORMONE FLAG

Record the date flag in the event a complete *RX Date-Hormone* was not entered to explain why.

Recording RX Date-Hormone Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Hormone* has a full or partial date recorded.
2. Unknown if Hormone Therapy Given- Code 10 if it is unknown whether any hormone therapy was given.
3. No Hormone Therapy- Code 11 if no hormone therapy is planned or given.
4. Unknown Date- Code 12 if the *RX Date-Hormone* cannot be determined at all, but the patient did receive hormone therapy.
5. Hormone Therapy Planned but Not Started- Code 15 if hormone therapy is planned, but has not yet started and the start date is not yet available. If hormone therapy is later started, update this item, *RX Date-Hormone*, and *RX Summ-Hormone* in your database and submit a change sheet to the PCR. See *Part One, Changing Information*.
6. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any hormone therapy given)
11	No proper value is applicable in this context (for example, no hormone therapy given)
12	A proper value is applicable but not known (for example, <i>RX Date-Hormone</i> 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 the case was abstracted)
(blank)	A valid date value is provided in item <i>RX Date-Hormone</i>

The table on the next page illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Hormone* and *RX Date-Hormone Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Hormone (MMDDCCYY)	Interoperable RX Date-Hormone (CCYYMMDD)	RX Date-Hormone Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any hormone therapy done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No hormone therapy given	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, hormone therapy given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12
Hormone therapy not yet started	88888888 Example: 88888888	bbbbbbbb Example: bbbbbbbb	15

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

RX SUMM - BRM

Record the immunotherapy (biological response modifier, BRM) the patient received as a part of first course of treatment at the reporting institution and all other institutions. If immunotherapy was not administered, then this item records the reason it was not administered to the patient. Immunotherapy consists of biological or chemical agents that alter the immune system or change the host’s response to the tumor cells.

Codes and Definitions

Code	Definition
00	None- Immunotherapy was not part of the planned first course of therapy. Diagnosed at autopsy.
01	Immunotherapy administered as first course therapy.
82	Immunotherapy was not recommended/administered because it was contra-indicated 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.

Recording Immunotherapy

1. Immunotherapy not usually given for this condition- If immunotherapy was not administered to the patient, and it is known that it is not usually administered for this type and stage of cancer, code to 00.
2. Patient selected treatment option without immunotherapy- If the treatment plan offered multiple options, and the patient selected treatment that did not include immunotherapy, code to 00.
3. Immunotherapy usually given for this condition- If it is known immunotherapy 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.

4. Patient refused- If the patient refused recommended immunotherapy, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 87.
5. Unknown- If it is not known whether immunotherapy 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 to 99.
6. List of Immunotherapy Agents- Use *SEER RX* to determine if a drug is an immunotherapy agent. *SEER RX* is an interactive antineoplastic drug data base and it can be downloaded from this website:

<http://seer.cancer.gov/tools/seerrx/>

7. Immunotherapy includes:

Allogeneic cells	Levamisole	Vaccine therapy
BCG	MVE - 2	Virus therapy
Interferon	Pyran copolymer	
LAK cells	Thymosin	

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - BRM*.

RX DATE-BRM

Record the date immunotherapy started.

Recording RX Date-BRM

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.
2. Collecting dates for each treatment modality allows sequencing of multiple treatments and evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
3. Exact Date Unavailable- If the exact date the immunotherapy started is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date-BRM Blank and assign the appropriate *RX Date-BRM Flag* for the following reasons:
 - a. when no immunotherapy is given
 - b. when immunotherapy is planned, but has not yet started
 - c. when it is unknown if any immunotherapy was given
 - d. when the patient received immunotherapy but the date is unknown
 - e. when the record was identified by death certificate only
 - f. when it is an autopsy-only case

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - BRM*.

Special Instructions for Registry Hospitals

Some vendor software may require dates to be entered in the traditional format (MMDDCCYY). Registry hospitals should obtain instructions from their vendor on how dates should be entered.

RX DATE-BRM FLAG

Record the date flag in the event a complete *RX Date-BRM* was not entered to explain why.

Recording RX Date-BRM Flag

1. Full or Partial Date- Leave this field blank if *RX Date-BRM* has a full or partial date recorded.
2. Unknown if Immunotherapy Given- Code 10 if it is unknown whether any immunotherapy was given.
3. No Immunotherapy- Code 11 if no immunotherapy is planned or given.
4. Unknown Date- Code 12 if the *RX Date-BRM* cannot be determined at all, but the patient did receive immunotherapy.
5. Immunotherapy Planned but Not Started- Code 15 if immunotherapy is planned, but has not yet started and the start date is not yet available. If immunotherapy is later started, update this item, *RX Date-BRM*, and *RX Summ-BRM* in your database and submit a change sheet to the PCR. See *Part One, Changing Information*.
6. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any immunotherapy given)
11	No proper value is applicable in this context (for example, no immunotherapy given)
12	A proper value is applicable but not known (for example, <i>RX Date-BRM</i> 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 the case was abstracted)
(blank)	A valid date value is provided in item <i>RX Date-BRM</i>

The table on the next page illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-BRM* and *RX Date-BRM Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-BRM (MMDDCCYY)	Interoperable RX Date-BRM (CCYYMMDD)	RX Date-BRM Flag
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any immunotherapy done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No immunotherapy given	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, immunotherapy given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12
Immunotherapy not yet started	88888888 Example: 88888888	bbbbbbbb Example: bbbbbbbb	15

b=blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

RX SUMM- TRANSPLNT/ENDROCR

Record the 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.

Codes and Definitions

Code	Definition
00	No transplant procedure or endocrine therapy was administered as part of first course therapy. 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 and infusion.
30	Endocrine surgery and/or endocrine radiation therapy.
40	Combination of endocrine surgery and/or radiation with a transplant procedure. (Combination of codes 30 and 10, 11, 12, or 20.)
82	Hematologic transplant and/or endocrine surgery/radiation was not recommended/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. Death certificate only.

Recording Hematologic Transplant and Endocrine Procedures

1. Bone marrow transplants should be coded as either autologous (bone marrow originally taken from the patient) or allogeneic (bone marrow donated by a person other than the patient). For cases in which the bone marrow transplant was syngeneic (transplanted marrow from an identical twin), the item is coded as allogeneic.
2. Stem cell harvests involve the collection of immature blood cells from the patient and the reintroduction by transfusion of the harvested cells following chemotherapy or radiation.
3. Endocrine irradiation and/or endocrine surgery
 - Procedures which suppress the naturally occurring hormonal activity of the patient and thus alter or affect the long-term control of the cancer's growth.
 - These procedures must be bilateral to qualify as endocrine surgery or endocrine radiation. If only one gland is intact at the start of treatment, surgery and/or radiation to that remaining gland qualifies as endocrine surgery or endocrine radiation.
4. These procedures are not usually administered for this condition- Code 00 if a transplant or endocrine procedure was not administered to the patient, and it is known these procedures are not usually administered for this type and stage of cancer.
5. Patient selected treatment option that did not include one of these procedures- Code 00 if the treatment plan offered multiple options, and the patient selected treatment that did not include a transplant or endocrine procedure.
6. These procedures are usually administered for this condition- If it is known a transplant or endocrine procedure is usually administered for this type and stage of cancer, but was not administered to patient, use code 82, 85, 86, or 87 to record reason why it was not.
7. Patient refused- If the patient refused a recommended transplant or endocrine procedure, made a blanket refusal of all recommended treatment, or refused all treatment before any was recommended, code to 87.
8. Unknown- 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 to 99.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery*.

RX SUMM - SYSTEMIC SUR SEQ

Record the sequencing of systemic therapy and surgical procedures given as part of first course of treatment.

The sequence of systemic therapy and surgical procedures given as part of first course of treatment cannot always be determined using the date on which each modality was started or performed. This data item can be used to more precisely evaluate the timing of delivery of treatment to the patient.

Codes and Definitions

Code	Definition
0	<p><i>No systemic therapy and/or surgical procedures; unknown if any surgery and/or systemic therapy given-</i> No systemic therapy given; and/or no surgery of the primary site; no scope of regional lymph node surgery; no surgery to other regional site(s), distant site(s), or distant lymph node(s) or it is unknown whether any surgery and/or systemic therapy was given.</p> <p>Diagnosed at autopsy.</p> <p><i>Example:</i> Due to other medical conditions surgery was not performed.</p>
2	<p><i>Systemic therapy before surgery-</i> Systemic therapy given before surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A patient with prostate cancer received hormone therapy prior to radical prostatectomy.</p>
3	<p><i>Systemic therapy after surgery-</i> Systemic therapy given after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A patient underwent a colon resection followed by a 5-FU based chemotherapy regimen.</p>
4	<p><i>Systemic therapy both before and after surgery-</i> At least two courses of systemic therapy were given before and after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> A patient with breast cancer receives pre-operative chemotherapy followed by post-operative Tamoxifen.</p>
5	<p><i>Intraoperative systemic therapy-</i> Intraoperative systemic therapy given during surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> Patient with an intracranial primary undergoes surgery at which time a glial wafer is implanted into the resected cavity.</p>

Code	Definition
6	<p><i>Intraoperative systemic therapy with other therapy administered before and/or after surgery-</i> Intraoperative systemic therapy given during surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node (s) with other systemic therapy administered before and/or after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p> <p><i>Example:</i> Patient with metastatic colon cancer receives intraoperative chemotherapy to the liver and postoperative 5-FU and leucovorin with irinotecan.</p>
7	<p><i>Surgery both before and after systemic therapy -</i> Systemic therapy given before and after surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s).</p>
9	<p><i>Sequence unknown-</i> Administration of systemic therapy and surgery of the primary site; scope of regional lymph node surgery; surgery to other regional site(s), distant site(s), or distant lymph node(s) were performed and the sequence of the treatment is not stated in the patient record.</p> <p><i>Example:</i> An unknown primary of the head and neck was treated with surgery and chemotherapy prior to admission, but the sequence is unknown.</p>

Recording RX Summ-Systemic Sur Seq

1. Surgical Procedures include RX Summ- Surg Prim Site; RX Summ- Scope LN Surg; RX Summ- Surg Oth Reg/Dis.
2. No Surgery- If all surgery procedures listed above are coded to 0, then this item should be coded to 0.
3. Unknown if Surgery or Systemic Therapy Given- If it is unknown if surgery and/or systemic therapy were given, this item should be coded to 0.

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Surgery, RX Text -Chemo, RX Text - BRM, and RX Text -Hormone.*

RX SUMM - OTHER

Record other cancer-directed therapy received by the patient as part of the first course of treatment at the reporting institution and all other institutions. Other treatment includes therapies designed to modify or control the cancer cells that are not defined in *Surgery*, *Radiation*, or *Systemic Therapy* fields.

Codes and Definitions

Code	Definition
0	None- All cancer treatment was coded in other treatment fields (surgery, radiation, systemic therapy). Patient received no cancer treatment. 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 (see next page).
2	Other-Experimental- This code is not defined. It may be used to record participation in institution based clinical trials.
3	Other-Double Blind- A patient is involved in a double-blind clinical trial. Code the treatment actually administered when the double-blind trial code is broken.
6	Other-Unproven Cancer treatments administered by nonmedical personnel.
7	Refusal- 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. Death certificate only.

Recording Other Treatment

1. Hematopoietic diseases- Treatment for reportable hematopoietic diseases can be supportive care, observation, or any treatment that does not meet the usual definition in which treatment “modifies, controls, removes, or destroys proliferating cancer tissue.” Such treatments include phlebotomy and blood thinners and anti-clotting agents, and should be coded 1.

Note: Transfusions are no longer coded as treatment for hematopoietic diseases effective for cases diagnosed on or after January 1, 2012.

- a. Blood thinners and/or anti-clotting agents can only be coded as treatment for the following conditions:

- 9741/3 Systemic mastocytosis
- 9742/3 Mast cell leukemia
- 9875/3 Chronic myelogenous leukemia BCR/ABL1 positive
- 9950/3 Polycythemia vera
- 9961/3 Primary myelofibrosis
- 9962/3 Essential thrombocythemia
- 9963/3 Chronic neutrophilic leukemia
- 9975/3 Myelodysplastic/myeloproliferative neoplasm, unclassifiable

Note: To determine whether aspirin is administered for pain, cardiovascular protection or thinning of platelets in the blood, use the following general guideline:

- Pain control is approximately 325–1000 mg every 3–4 hours.
- Cardiovascular protection starts at about 160 mg/day.
- Aspirin treatment for essential thrombocythemia is low dose, approximately 70–100 mg/day.

- b. Phlebotomy may be called blood removal, bloodletting, or venisection. Phlebotomies should only be coded for treatment of polycythemia vera. Do not code phlebotomies for other hematopoietic diseases.

2. Ancillary Drugs- Do not code ancillary drugs in this field. There is no coding scheme for ancillary drugs.

Examples: Aredia, Allopurinol, G-CSF (growth stimulating factors), Epopen, Nupogen/Neupogen, Leucovorin

Note: This is a partial list. See *SEER RX* to determine if a drug is an ancillary drug. *SEER RX* is an interactive antineoplastic drug data base and it can be downloaded from this website:

<http://seer.cancer.gov/tools/seerrx/>

3. Embolization- Assign code 1 for embolization using alcohol as an embolizing agent and for embolization to a site other than liver where the embolizing agent is unknown.

Note: Do not code pre-surgical embolization that is given for the purpose of shrinking the tumor.

4. PUVA- Assign code 1 in the rare event Psoralen (P) and long-wave ultraviolet radiation (UVA) it is used as treatment for extremely thin melanomas or cutaneous T-cell lymphomas (e.g. mycosis fungoides).

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text - Other*

RX DATE - OTHER

Record the date other treatment started.

Recording RX Date- Other

1. Date Format- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.

2. Collecting the dates for each treatment modality allows sequencing of multiple treatments and aids evaluation of time intervals (from diagnosis to treatment and from treatment to recurrence).
3. Exact Date Unavailable- If the exact date of other treatment is not available, record an approximate date. Refer to *Part Three, General Information*.
4. Leave RX Date- Other Blank and assign the appropriate *RX Date-Other Flag* for the following reasons:
 - a. when no other therapy is given
 - b. when other therapy is planned, but has not yet started
 - c. when it is unknown if any other therapy was given
 - d. when the patient received other therapy but the date is unknown
 - e. when the record was identified by death certificate only
 - f. when it is an autopsy-only case

Text

Text to support this data item must be recorded in specific text fields. See *Part Three, Data Item Instructions, RX Text-Other*

RX DATE-OTHER FLAG

Record the date flag in the event a complete *RX Date-Other* was not entered to explain why.

Recording RX Date-Other Flag

1. Full or Partial Date- Leave this field blank if *RX Date-Other* has a full or partial date recorded.
2. Unknown if other therapy Given- Code 10 if it is unknown whether other therapy was given.
3. No Other Therapy- Code 11 if no other therapy is planned or given.
4. Unknown Date- Code 12 if the *RX Date-Other* cannot be determined at all, but the patient did receive other therapy.
5. Other Therapy Planned but Not Started- Code 15 if other therapy is planned, but has not yet started and the start date is not yet available. If other therapy is later started, update this item, *RX Date-Other*, and *RX Summ-Other* in your database and submit a change sheet to the PCR. See *PCR Manual, Part One, Changing Information*.
6. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if other therapy given)
11	No proper value is applicable in this context (for example, no other therapy given)
12	A proper value is applicable but not known (for example, <i>RX Date-Other</i> is unknown)
15	Information is not available at this time, but it is expected that it will be available later (for example, other therapy is planned as part of the first course of therapy, but had not been started at the time the case was abstracted)
(blank)	A valid date value is provided in item <i>RX Date-Other</i>

The table on the next page illustrates the use of the date flag and the traditional and interoperable date formats for coding *RX Date-Other* and *RX Date-Other Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional RX Date-Other (MMDDCCYY)	Interoperable RX Date-Other (CCYYMMDD)	RX Date-Systemic Other
Full date known	MMDDCCYY Example:04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if other therapy done	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
No other therapy given	00000000 Example: 00000000	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, other therapy given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12
Other therapy not yet started	88888888 Example: 88888888	bbbbbbbb Example: bbbbbbbb	15

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software.

DATE OF 1ST CRS RX - COC

Record the date of first course treatment. This is the date of initiation of the first cancer-directed therapy for the cancer being reported

Recording Date of 1st CRS RX- COC

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.
2. **Earliest Date**- Record the earliest of the following dates: *RX Date- Surgery, RX Date- Radiation, RX Date- Chemo, RX Date-Hormone, RX Date-BRM, or RX Date- Other.*
3. **Physician Decides Not to Treat**- If the physician decides not to treat the patient, record the date of this decision as the *Date of 1st CRS RX-COC*. The physician may decide not to treat the patient because of co-morbid conditions, advanced disease, or because the accepted management of the cancer is to observe until the disease progresses or until the patient becomes symptomatic.

Example: On February 12, 2010 the physician says a low-stage prostate cancer patient will be observed until the Prostatic Specific Antigen (PSA) level starts to rise. Enter 20100212 as the date of first course treatment.
4. **Patient Refuses Treatment**- If the patient refuses treatment; record the date of this decision as the *Date of 1st CRS RX-COC*. If the patient is diagnosed at the reporting facility and no further information is available record the date the patient was last seen at the reporting institution.
5. **Incisional Biopsy**- Do not record the date of incisional, core or fine needle biopsy as the *Date of 1st CRS RX-COC*.
6. **Exact Date Unavailable**- If the exact date of first course of treatment is not available, record an approximate date. Refer to *Part Three, General Information*.
7. **Leave Date of 1st CRS RX-COC Blank** and assign the appropriate *Date of 1st CRS RX* Flag only for the following reasons:
 - a. when it is unknown if any cancer-directed treatment was given
 - b. when the patient had cancer-directed treatment but the date is unknown
 - c. when the record was identified by death certificate only
 - d. when it is an autopsy-only case

DATE OF 1ST CRS RX FLAG

Record the date flag in the event a complete date of first course treatment was not entered to explain why.

Recording Date of 1st CRS RX Flag

1. Full or Partial Date- Leave this field blank if *Date of 1st CRS RX-COC* has a full or partial date recorded.
2. Unknown if Cancer-Directed Treatment Given- Code 10 if it is unknown whether any cancer-directed treatment was given.
3. No Treatment- Code 11 if the initial diagnosis was at autopsy.
4. Unknown Date- Code 12 if the *Date of 1st CRS RX-COC* cannot be determined at all, but the patient did receive first course treatment.
5. Enter Directly- This field should be entered directly (when appropriate).

Codes and Definitions

Code	Definition
10	No information whatsoever can be inferred from this exceptional value (that is, unknown if any cancer-directed treatment was given.)
11	No proper value is applicable in this context (autopsy only)
12	A proper value is applicable but not known (for example, <i>Date of 1st CRS RX – COC</i> is unknown)
(blank)	A valid date value is provided in item <i>Date of 1st CRS RX – COC</i>

The table on the following page illustrates the use of the date flag and the traditional and interoperable date formats for coding *Date of 1st CRS RX – COC* and *Date of 1st CRS RX Flag*. In the table, the lowercase letter “b” is used to represent each blank space.

Description	Traditional Date of 1st CRS RX – COC (MMDDCCYY)	Interoperable Date of 1st CRS RX – COC (CCYYMMDD)	Date of 1st CRS RX Flag
Full date known	MMDDCCYY Example: 04032010	CCYYMMDD Example: 20100403	bb
Month and year known	MM99CCYY Example: 04992010	CCYYMMbb Example: 201004bb	bb
Year only known	9999CCYY Example: 99992010	CCYYbbbb Example: 2010bbbb	bb
Unknown if any treatment given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	10
Diagnosed at autopsy only	00000000 (example: 00000000)	bbbbbbbb Example: bbbbbbbb	11
Date is unknown, treatment given	99999999 Example: 99999999	bbbbbbbb Example: bbbbbbbb	12

b=blank

Special Note for Registry Hospitals

This field should be entered directly (when appropriate) even if the traditional form of date entry is used in the vendor software

RX SUMM-TREATMENT STATUS

Summarize whether the patient received any treatment or if the tumor is under active surveillance.

Codes and Definitions

Code	Definition
0	No treatment given
1	Treatment given
2	Active surveillance (watchful waiting)
9	Unknown if treatment was given

Recording Rx Summ-Treatment Status

1. Diagnosed Prior to January 1, 2010- This item may be left blank for cases diagnosed prior to January 1, 2010.
2. Code 0- Assign code 0 when:
 - a. No treatment is given.
 - b. Patient refused treatment
 - c. Physician decides not to treat for any reason, such as the presence of Comorbidities.

Example: An elderly patient with pancreatic cancer requested no treatment. Code *Rx Summ Treatment Status* to 0

3. Active Surveillance (watchful waiting) - Assign code 2 when the patient will be under active surveillance.

Example: Treatment plan for a lymphoma patient is active surveillance. Code *Rx Summ-Treatment Status* to 2

4. Subsequent Treatment- Treatment given after a period of active surveillance is considered subsequent treatment and is not coded in this item.

DATE OF LAST CONTACT

Record the date of last contact.

Recording Date of Last Contact

1. **Date Format**- Record date in year, month, day format (CCYYMMDD). Record the year in the first four spaces, the month in the fifth and sixth spaces, the day in the last two spaces. A zero must precede single-digit months and days. See *Part Three, General Instructions* for allowable values.

Example: Record June 30, 2010 as 20100630.

2. **Report Actual Date Only.** Blank or approximation of month, day, century, or year is not acceptable when reporting this data item to the PCR. Fictitious dates or default values are also not acceptable

Exception: If a patient is known to have expired after discharge from your facility, the month and/or day may be left blank if the exact month and/or day are not known.

3. **Inpatient Admission**- If the last contact with a patient is an inpatient admission, record the date of discharge.
4. **Outpatient Visit**- If the last contact with the patient was an outpatient visit, record the outpatient date.
5. **Treatment After Discharge**- If the patient receives treatment after discharge record the date of the treatment.

Example: The patient is admitted on November 1, 2010 and is discharged on November 3, 2010 and then starts his radiation treatment on December 1, 2010. The date of last contact is 20101201.

6. **Patient Deceased**- If the patient is deceased, record the date of death.

Note: *Date of Last Contact* does **not** have to be submitted as a change or update if the patient is readmitted or expires after the initial record was submitted.

DATE OF LAST CONTACT FLAG

The PCR requires a *Date of Last Contact* to be entered on all cases; therefore this field will always be left blank.

Codes and Definitions

Code	Definition
(blank)	A valid date value is provided in item <i>Date of Last Contact</i> .

VITAL STATUS

Record the appropriate code for the patient's vital status as of the date recorded in data item *Date of Last Contact*. Use the most accurate information available.

Codes and Definitions

Code	Definition
0	Dead
1	Alive

Note: Vital Status does **not** have to be submitted as a change or update if the patient expires after the initial record was submitted. The PCR periodically matches records on the PCR database against Pennsylvania death certificate files. As a result of this match, the PCR will send each hospital a list of its reported patients who have expired.

REPORTING HOSPITAL

Record the reporting facility identification (ID) number as described under special instructions below.

Special Instructions

1. Registry Hospitals - Record the ID number assigned to your facility by the American College of Surgeons, Commission on Cancer.
2. Abstract Plus - The ID number assigned to your facility by the American College of Surgeons, Commission on Cancer has been defaulted in your software.

ABSTRACTED BY

Record the initials of the individual completing the abstract.

Special Instructions

1. Registry Hospitals - Record the initials or assigned code of the individual who abstracted this record. Do not code the data entry person **unless** that person is also the abstractor.
2. Abstract Plus - Initials have been defaulted to the person logged into the software.

GUIDELINES FOR REPORTING TEXT

Text Requirements

The PCR requires all records to include text information to support specified fields. Text documentation is an essential component of a complete electronic abstract and is heavily utilized for quality control and special studies. Text is needed to justify coded values and to document supplemental information not transmitted within coded values. Text is used to validate data items, verify potential errors identified through standard edits, document clarifications, determine multiple primaries, and reconcile data item discrepancies when the same patient is submitted by several facilities.

The text fields must contain descriptions entered by the abstractor independently from the code(s). Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values. Cancer abstracting software must include specific fields designed to document text as defined by NAACCR fields. These fields must be transmitted to the PCR in addition to the other required data items when electronic shipments are prepared.

Completion of Text Fields

Text should be complete but concise. The text fields must summarize all cancer information recorded in the medical record. Text must be completed for primary site, laterality, histology, grade, and collaborative stage or summary stage on every record. Text should be completed for pathology and other diagnostic and treatment text fields as appropriate for studies performed and treatment provided. If information is missing from the record, state that it is missing. The text fields should be used to document information that will support the accuracy of data so anyone reviewing the abstract will be able to justify the coded information.

Amount of Text

Quality of text is more important than amount or quantity of text. The most useful text is brief, concise, and addresses pertinent issues. Often it is necessary to use abbreviations to provide adequate descriptions within the limited size of the text fields. Use standard medical abbreviations whenever possible. Refer to *Appendix L* for a list of PCR acceptable abbreviations. Include dates (month, day, and year) when appropriate.

Note the maximum field lengths for each text field. These lengths indicate how many characters will be transmitted to the PCR. Do not include irrelevant information. Do not repeat information from other text fields.

TEXT-DX PROC-PE**Maximum Field Length - 1000 characters**

Record text information from the history/physical examination that supports the diagnosis and history of the tumor as applicable.

Source Records:

The history/physical examination findings may be found in, but are not limited to, the following source records:

- History and Physical Report
- Consultation Reports
- Progress Notes

Suggestions for Text:

- Date of physical exam.
- Age, sex, race/ethnicity.
- History that relates to cancer diagnosis.
- Primary site.
- Histology (if diagnosis prior to this admission).
- Tumor location.
- Tumor size.
- Palpable lymph nodes.
- Record positive and negative clinical findings. Record positive results first.
- Impression (when stated and pertains to cancer diagnosis).
- Treatment plan.

Examples:

1. 5 cm mass palpated in UOQ rt breast
2. Abdominal pain, constipation
3. Enlarged lymph node in neck, fatigue
4. 35 year old white male with unexplained 30 lb weight loss.

TEXT-DX PROC-X-RAY/SCAN**Maximum Field Length - 1000 characters**

Record text information from diagnostic imaging reports as applicable. Document both positive and negative findings and the date(s) of the imaging result(s).

Source Records:

The diagnostic imaging findings may be found in, but are not limited to, the following source records:

- All Diagnostic X-ray reports including mammograms and CT scans
- History and Physical Report
- Consultation Reports
- Discharge Summary

Suggestions for Text:

- Date(s) of X-ray/Scan(s).
- Primary site.
- Histology (if given).
- Tumor location.
- Tumor size.
- Lymph nodes.
- Record positive and negative clinical findings. Record positive results first.
- Distant disease or metastasis.

Examples:

1. 01/15/2010 mammo-2 cm mass in UOQ rt breast; 01/16/2010 normal CXR
2. 01/15/2010 CT-abd-large mass in sigmoid colon with possible extension into pericolic fat; 01/16/2010 negative bone scan
3. 01/15/2010 CT-abd-diffuse adenopathy involving the retroperitoneal, periaortic, and inguinal LN
4. 1/15/2010 MRI glioblastoma multiforme, WHO grade III

TEXT-DX PROC-SCOPES

Maximum Field Length - 1000 characters

Record text information from endoscopic examinations as applicable. Document both positive and negative findings and the date(s) of the scope(s).

Source Records:

The endoscopic examination findings may be found in, but are not limited to, the following source records:

- Endoscopy Reports (i.e. Bronchoscopy, Colonoscopy, Laryngoscopy, Esophagoscopy)
- History and Physical Report
- Discharge Summary
- Consultation Reports

Suggestions for Text:

- Date(s) of endoscopic exam(s).
- Primary site.
- Histology (if given).
- Tumor location.
- Tumor size.
- Lymph nodes.
- Record positive and negative clinical findings. Record positive results first.

Examples:

1. Sigmoidoscopy 2/15/2010-Extrinsic compression of sigmoid most likely secondary to mass
2. Sigmoidoscopy 2/15/2010-Ulcerated lesion found in rectosigmoid with invasion through the serosa
3. EGD 2/1/2010-5 cm constricting mass in lower esophagus with metastasis to the cervical lymph nodes

TEXT-DX PROC-LAB TESTS

Maximum Field Length - 1000 characters

Record information from laboratory tests or marker studies other than cytology/histopathology that are clinically diagnostic of cancer as applicable. Document pertinent positive and negative findings with the date(s) of these test(s).

Source Records:

The laboratory examination findings may be found in, but are not limited to, the following source records:

- Laboratory Reports
- History and Physical Reports
- Consultation Reports

Suggestions for Text:

- Type of laboratory test/tissue specimen(s).
- Record both positive and negative findings. Record positive test results first.
- Information can include tumor markers, serum and urine electrophoresis, special studies, etc.
- Date(s) of laboratory test(s).
- Tumor markers included, but are not limited to:
 - Breast Cancer: Estrogen Receptor Assay (ERA), Progesterone Receptor Assay (PRA), Her2/neu.
 - Prostate Cancer: Prostatic Specific Antigen (PSA).
 - Testicular Cancer: Human Chorionic Gonadotropin (hCG), Alpha Fetoprotein (AFP), Lactate Dehydrogenase (LDH).
 - Polycythemia Vera- JAK2

Examples:

1. 1/15/2010 ER/PR studies positive
2. 1/15/2010 CEA = 10.4
3. 2/01/2010 Elevated WBC

TEXT-DX PROC-OP

Maximum Field Length - 1000 characters

Record text information from operative reports that support the diagnosis as applicable. Document both positive and negative findings and the date(s) of the procedure(s).

Source Records:

The operative findings may be found in, but are not limited to, the following source records:

- Operative Reports
- Consultation Reports

Suggestions for Text:

- Dates and descriptions of biopsies and all other surgical procedures from which staging information was derived.
- Number of lymph nodes removed.
- Size of tumor removed.
- Documentation of residual tumor.
- Evidence of invasion of surrounding areas.

Examples:

1. MRM 2/1/2010-Firm 3 cm mass excised from rt UOQ
2. Hemicolectomy 2/15/2010-No metastatic nodules noted in liver
3. Exp Lap 2/1/2010-Tumor arising from ileum and enlarged spleen consistent with lymphoma
4. Mastectomy 2/22/2010- 3 out of 12 lymph nodes found to have metastasis

TEXT-DX PROC-PATH

Maximum Field Length - 1000 characters

Record text from cytology and histopathology reports to support the final pathologic diagnosis. Include all descriptive terms from the histology or cytology **final diagnosis** to describe the specific diagnosis including nouns, adjectives, and phrases. Also include documentation to support unusual site/histology combinations, notes, comments, addenda, and results of consults and second opinions.

Either *Text-Histology Title* or *Text-Dx Proc-Path* must be completed on each record. Information to support the exact diagnosis has to appear in one of these two fields. *Text-Histology Title* is a 100 character field generally used to record clinical or other non pathologic diagnoses; *Text-Dx Proc-Path* is a 1000 character field generally used to record histologically and cytologically confirmed diagnoses from pathology reports.

This field should also include text to support multiple primaries diagnosed simultaneously and discrepancies between pathology reports. For example, if a definitive surgery pathology report has a more specific or differing diagnosis than the biopsy report, document the physician's final diagnosis. Include text to clarify site and/or histology information for cases discussed at Cancer Conference, especially if the site was unknown.

Terminology

If the reporting facility considers the terminology of severe dysplasia or high grade dysplasia of the colon as synonymous with carcinoma in-situ, follow the procedure described in *PCR Manual Part Three, Behavior*. Include text in this field to support the final pathologic diagnosis along with the statement "in-situ per pathologist". If any colon cases diagnosed with severe dysplasia and/or high grade dysplasia are submitted to the PCR without the text documentation "in-situ per pathologist", the cases will be deleted since the terminology alone is not reportable.

Mixed or multiple histologies may have documentation of various phrases describing the tumor. When documenting the description of the tissue, include the terminology type in the description. These terms are important because they impact the ICD-O code assignment.

Source Records:

The pathology findings may be found in, but are not limited to, the following source records:

- Pathology and Cytology Reports, Autopsy Reports
- Slide Consultation Reports

Suggestions for Text:

- Date(s) of procedure(s).
- 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 resected 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.).

Examples:

1. Poorly diff infiltrating duct and tubular carcinoma
2. Thyroidectomy specimen yields 8mm mass in RUL
3. Stains positive for mucin producing adenocarcinoma
4. Transitional cell Ca with foci of squamous differentiation
5. Negative bone marrow
6. Tumor directly extends into the trachea
7. 2 out of 10 peribronchial lymph nodes examined are positive on biopsy
8. Positive brain metastasis
9. Meningioma, WHO grade I

TEXT-PRIMARY SITE TITLE

Maximum Field Length - 100 characters

Record text describing the primary site including subsite information. Always document laterality when the site is paired. Refer to the listing of Paired Sites in *Part Three, Laterality. Text-Primary Site Title* must be completed on each record.

Source Records:

The primary site and laterality may be found in, but are not limited to, the following source records:

- Pathology Report
- Operative Report
- Xrays/Scans
- Discharge Summary
- Consultation Reports

Suggestions for Text:

- Include information on the location of the primary site of the tumor.
- Include available information on tumor laterality.

Examples:

1. Rt breast, UOQ
2. Sigmoid Colon
3. Cervical and Inguinal LNs
4. R Frontal Lobe of Brain

TEXT-HISTOLOGY TITLE

Maximum Field Length - 100 characters

Record text to support the patient's final diagnosis: clinical, other non pathologic diagnosis, or histologic diagnosis including cell type, behavior, and grade.

Either *Text-Histology Title* or *Text-Dx Proc-Path* must be completed on each record. Information to support the exact diagnosis has to appear in one of these two fields. *Text-Histology Title* is a 100 character field generally used to record clinical or other non pathologic diagnoses; *Text-Dx Proc-Path* is a 1000 character field generally used to record histologically and cytologically confirmed diagnoses from pathology reports.

Source Records:

The histologic diagnosis may be found in, but is not limited to, the following source records:

- Pathology and Cytology Reports
- History and Physical Report
- Discharge Summary
- Consultation Reports
- Slide Consultation Reports

Suggestions for Text:

- Information on histologic type and behavior.
- Information on differentiation from scoring systems such as Gleason's Score, Bloom-Richardson, Grade, etc.

Examples:

1. Ductal Ca w/ lobular features, well diff
2. Moderately to poorly diff Adenoca
3. Diffuse large B-cell lymphoma
4. Cancer

TEXT-STAGING

Maximum Field Length - 1000 characters

Record text to support any Collaborative Stage data items not already supported in other text fields (see *Part Three, Data Item Instructions, Collaborative Stage*). This field can also be used to continue Collaborative Stage text from another field.

Example: The only information available is the TNM stage, record *Physician stated this case is a TINIMO*.

Record text information to support the Summary Stage code assigned according to *SEER Summary Stage 2000 (SS2000)* or *SEER Summary Stage 1977 (SS77)* when applicable (see *Part Three, Data Item Instructions, SEER Summary Stage*). Document the extension of the disease that justifies the Summary Stage based on imaging studies, lab tests, scopes, and operative procedures performed. Also include both positive and negative findings and appropriate dates not already recorded in other *Text-DX* fields. If information is not sufficient to support a specific Summary Stage code, record *unknown* in this field.

Sources Records:

Information to determine Collaborative Stage data items and Summary Stage may be found in, but is not limited to, the following reports:

- Pathology Reports
- Operative procedures
- X-Rays/Scans
- Scopes
- Lab Tests
- Discharge Summary
- Consultations

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(s) of distant metastasis.
- Physician's comments.

Examples:

1. Work up and initial treatment for prostate primary included lung scan, bone scan, and CT/Pelvis. Based on these procedures, the Summary Stage is determined to be *Distant*, code 7. Document the following in the appropriate text fields:

Text-Dx Proc-X-ray/Scan: Bone Scan 1/15/2010-mets to pelvis; Lung scan 1/20/2010 no evidence of metastatic disease; CT/Pelvis-1/15/2010-positive iliac adenopathy

Text-Staging: Pelvic bone mets

2. Diagnosis of lymphoma and workup included CT scans and a bone marrow biopsy. Based on these procedures, the Summary Stage is determined to be *Regional NOS*, code 5. Document the following in the appropriate text fields:

Text-Dx Proc-X-ray/Scan: CT scans 1/15/2010-mediastinal and axillary LN suspicious for lymphoma, no pelvic or retroperitoneal adenopathy

Text-Dx Proc-Path: Bone marrow 2/01/2010 negative

Text-Staging: Multiple LN regions above diaphragm

3. If the only documentation is that the patient was diagnosed two years ago and now is admitted in January 2001 for treatment of recently discovered bone metastases, record:

Text-Staging: unknown at initial dx, bone mets 1/2010

RX TEXT-SURGERY

Maximum Field Length - 1000 characters

Record all surgical procedures, including dates, performed as first course of treatment as applicable. Surgical procedures used to treat regional lymph nodes and other regional and/or distant sites as first course of treatment should be documented. If applicable, text should also be included to describe the number of regional lymph nodes examined as part of the first course of treatment.

Source Records:

The surgical procedure information may be found in, but is not limited to, the following source records:

- Operative Reports
- Discharge Summary
- Consultation Reports
- History and Physical Report

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
- Regional tissues removed
- Metastatic sites
- Facility where each procedure was performed
- Record positive and negative findings; record positive findings first
- Reason surgery to primary site not performed

Examples:

1. 01/15/2010 Rt lumpectomy; 1/25/01 Rt MRM with LND (10 LN removed)
2. 01/15/2010 Rt hemicolectomy with LND (12 LN removed) & wedge resection of liver
3. 01/15/2010 Excision of cervical LN

RX TEXT-RADIATION (BEAM)

Maximum Field Length - 1000 characters

Record all beam radiation, including dates, given as first course of treatment as applicable.

Source Records:

The radiation information may be found in, but is not limited to, the following source records:

- Radiation Records or treatment letters
- Discharge Summary
- Consultation Reports

Suggestions for Text:

- Date when 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 five treatments; unknown if radiation was given)
- Reason no radiation therapy was administered

Examples:

1. Began 01/15/2010, 6000 cGy, 30fx
2. Radiation planned, patient refused
3. None

RX TEXT-RADIATION OTHER

Maximum Field Length - 1000 characters

Record all other radiation, including dates, given as first course of treatment as applicable.

Source Records:

The other radiation treatment may be found in, but is not limited to, the following source records:

- Radiation logbooks or treatment letters
- Discharge Summary
- Consultation Reports

Suggestions for Text:

- Date treatment was started
- Where treatment was given (e.g., at this facility, at another facility)
- Type(s) of nonbeam radiation (e.g., High Dose rate brachytherapy, seed implant, Radioisotopes [I-131])
- Other treatment information (e.g., unknown if radiation was given)
- Reason no radiation therapy was administered

Examples:

1. None
2. 01/15/2010 Brachytherapy
3. 01/15/2010 Iodine-131 at Cancer Radiation Center

RX TEXT-CHEMO

Maximum Field Length - 1000 characters

Record all chemotherapy, including dates, administered as first course of treatment as applicable.

Source Records:

The chemotherapy treatment information may be found in, but is not limited to, the following source records:

- Chemotherapy logbooks or treatment letters
- Discharge Summary
- Consultation Reports
- History and Physical Report

Suggestions for Text:

- Date when 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)

Examples:

1. 01/15/2010 Cisplatin & VP16 started, failed after 3 mths; Carboplatin & Taxol started
2. 01/15/2010 5 FU started
3. 01/15/2010 CHOP started at Cancer Treatment Center.

RX TEXT-HORMONE

Maximum Field Length - 1000 characters

Record all hormone therapy, including dates, administered as first course of treatment as applicable.

Source Records:

The hormone therapy information may be found in, but is not limited to, the following source records:

- Discharge Summary
- Consultation Reports
- History and Physical Report

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:

1. 01/15/2010 Tamoxifen prescription given to pt; unknown if filled
2. 01/15/2010 Prednisone started as part of CHOP regimen

RX TEXT-BRM

Maximum Field Length - 1000 characters

Record biological-response modifier treatment, including dates, administered as first course of therapy for cancer as applicable. This is also referred to as immunotherapy.

Source Records:

The biological-response modifier treatment information may be found in, but is not limited to, the following source records:

- Discharge Summary
- Consultation Reports
- History and Physical Report

Suggestions for Text:

- When treatment was given (e.g., at this facility; at another facility)
- Type of BRM agent (e.g., Interferon, BCG)
- BRM procedures (e.g., bone marrow transplant, stem cell transplant)
- Other treatment information (e.g., treatment cycle incomplete; unknown if BRM was given)

Examples:

1. 01/15/2010 Levamisole started
2. 01/15/2010 Stem cell transplant done at University Hospital.
3. 01/15/2010 Interferon started

RX TEXT-OTHER

Maximum Field Length - 1000 characters

Record all other treatment, including dates, performed as first course of treatment as applicable.

Source Records:

Other treatment may be found in, but is not limited to, the following source records:

- Discharge Summary
- Consultation Reports
- History and Physical Reports

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:

1. 01/15/2010 hyperbaric oxygen started at General Hospital
2. 01/15/2010 PUVA
3. None

TEXT-REMARKS

Maximum Field Length - 1000 characters

Record text information not elsewhere provided for or as an overflow from other text fields. The following information should be included in this field as applicable to the case:

- Document the site, laterality if applicable, histology, and date of diagnosis for all known previous primaries.
- Document text to explain any unusual or potentially questionable entry on the abstract. This will reduce the need to re-pull medical records at a later date.
- Document text to note particular issues or clarifications that were resolved prior to completion of the abstract. For example, clarifications made with a physician through quality assurance studies.

Source Records:

Information for this field may be found in, but is not limited to, the following source records:

- History and Physical Report
- Pathology Reports
- Discharge Summary
- Consultation Reports
- Cancer Conference Documentation

Suggestions for Text:

- Personal history of cancer.
- Comorbidities.
- Information on sequence numbers if a person was diagnosed with another cancer out-of-state or before the registry's reference date.
- Justification for unusual site/histology combinations.

Examples:

1. Simultaneous tumors: mucinous adenoca of sigmoid invading pericolic fat, 2/15 + LN and adenoca of cecum with lamina propria invasion
2. Primary site determined at Cancer Conference upon review of clinical presentation and diagnostic evaluation
3. Hx of thyroid cancer diagnosed in 1992 in New York.
4. Hx of benign meningioma 2003

TEXT-PLACE OF DIAGNOSIS

Maximum Field Length - 60 characters

Record text to indicate the facility, city, state or country where the diagnosis was made. This data item is required only when the patient was not diagnosed at the reporting facility.

Source Records:

Information for this field may be found in, but is not limited to, the following source records:

- Admission record
- History and Physical Report
- Consultation Reports

Suggestions for Text:

- Record the complete name of the hospital or the physician office where the diagnosis occurred. The initials of the hospital are not adequate.
- For out-of-state residents and facilities, include the city and state where the medical facility is located.

Examples:

1. University General Hospital, Anytown, NJ
2. Medical Center of London, England

SYSTEM CODES

System codes reflect types of coding systems used, record processing dates, and other information regarding how the data were collected. These codes are required to be transmitted on cases submitted electronically.

1. Registry Hospitals - Registry hospitals using commercial or hospital-developed software are responsible for making sure the correct system codes are submitted. Since most are computer generated, the registrar must communicate problems in complying with PCR code requirements to software vendors or facility Information Systems personnel.
2. Abstract Plus - Abstract Plus software will comply with all PCR code requirements. The only action the hospital abstracter must take is to make sure the correct version of Abstract Plus is installed at the hospital as directed by the PCR.

Required Codes and Definitions

PCR Required Data Item	NAACCR Item #	PCR Specific Instructions
Record Type	10	Must always contain "A" for <i>Full case abstract type, including text data item</i> ; length=22,824.
Registry Type	30	Allowable codes: "2" for central registry or hospital consortium (not population based); and "3" for single hospital/freestanding center.
NAACCR Record Version	50	Must always contain "140" for 2014 version (Version V15).
Site Coding Sys--Current	450	Cases diagnosed on or after 01/1/2001 must always contain "5" for ICD-O-3; cases diagnosed before 1/1/2001 must always contain "4" for ICD-O-2; cases with an unknown <i>Date of Diagnosis</i> and <i>Date of 1st Contact</i> on or after 01/01/2001 must always contain "5" for ICD-O-3; cases with an unknown <i>Date of Diagnosis</i> and <i>Date of 1st Contact</i> prior to 01/01/2001 must always contain "4" for ICD-O-2.
Morph Coding Sys--Current	470	Cases diagnosed on or after 1/1/2010 must always contain "8" for ICD-O-3 plus 2008 WHO terms. Cases diagnosed 01/1/2001 to 12/31/2009 must always contain "7" for ICD-O-3; cases diagnosed before 1/1/2001 must always contain "6" for ICD-O-2 plus REAL and FAB codes; cases with an unknown <i>Date of Diagnosis</i> and <i>Date of 1st Contact</i> on or after 1/1/2010 must always contain "8" for ICD-O-3 plus 2008 WHO terms. Cases with an unknown <i>Date of Diagnosis</i> and <i>Date of 1st Contact</i> 01/01/2001-12/31/2009 must always contain "7" for ICD-O-3; cases with an unknown <i>Date of Diagnosis</i> and <i>Date of 1st Contact</i> prior to 01/01/2001 must always contain "6" for ICD-O-2 plus REAL and FAB codes.
RX Coding System--Current	1460	Must always contain "06" for <i>Treatment data coded according to FORDS</i> .

PCR Required Data Item	NAACCR Item #	PCR Specific Instructions
ICD Revision Number	1920	Must always contain "1" for ICD-10.
Date Case Completed*	2090	Must contain the date abstract first passed all edits applied. Blank is not acceptable in any portion of the date.
Date Case Last Changed	2100	Contains the latest date the case was modified after completion at the reporting facility.
Date Case Report Exported	2110	Must contain the date the reporting facility exported the electronic abstract to a file for transmission to the central registry. Blank is not acceptable in any portion of the date.
ICD-O-3 Conversion Flag	2116	Cases diagnosed on or after 1/1/2001 must contain "0" for <i>Primary site and morphology originally coded in ICD-O-3</i> .
Vendor Name	2170	<i>Commercial Software</i> : name and version number must always be included; <i>Abstract Plus</i> : will contain name and version number as specified by the PCR.
CS Version Input Original	2935	Must contain the number of the version used to initially code the Collaborative Stage input fields.
CS Version Derived	2936	Must contain the number of the version of Collaborative Stage used most recently to derive the Collaborative Stage output fields.
CS Version Input Current	2937	Must contain the number of the version after Collaborative Stage input fields have been updated or recoded.

**Registry Hospital software only

**PART FOUR:
QUALITY CONTROL**

QUALITY CONTROL

The purpose of cancer data collection varies with the type and goals of the registry. The primary goal of hospital-based cancer registries is the improvement of patient care, and the primary goal of non-registry hospitals is to provide data to the central cancer registry. The primary objective of the central or population-based incidence registries is the determination of cancer rates and trends in the population. Whether data are reported to the Pennsylvania Cancer Registry (PCR) or reported by the PCR, there is a universal need for the data collected in any type of registry to be of the highest quality.

Quality can be defined as fitness for use. To assure data are of sufficient quality for use in meeting registry goals, quality control must be an integral component of the data collection system. Quality control involves the systematic execution of a carefully planned set of activities to monitor data quality and take appropriate action to positively affect future quality.

Activities and procedures to assure data quality should focus on three areas: completeness, accuracy and timeliness. Completeness refers to both case ascertainment and data collection. Accuracy refers to how well the abstracted data reflect the patient's diagnosis and treatment. Timeliness measures how the abstracting and reporting process are accomplished according to an expected schedule.

Evaluation of completeness, accuracy, and timeliness is the first step in quality control. To be effective, the registry's quality control plan must also involve a continuous loop of monitoring, communication, and feedback.

The following two sections describe various strategies used by reporting facilities and the PCR to assure data are as complete, accurate and timely as possible. The activities described for reporting facilities will enhance compliance to PCR reporting standards. Since communication and feedback are essential to the success of any quality control program, the major quality control procedures used by the PCR are described in order for hospital contacts to more fully understand the rationale for PCR requirements as well as verbal and written requests and questions made by the PCR.

QUALITY CONTROL: REPORTING FACILITIES

Reporting facilities must insure cancer data collected and submitted to the PCR are complete, accurate, and timely. Although some facilities may incorporate additional activities to assure quality, at a minimum, all facilities must include the following procedures to meet PCR reporting requirements and standards.

Completeness

1. **Casefinding Sources** - All areas where cancer patients are diagnosed or treated must be included in the casefinding system. This includes outpatient treatment areas, e.g., Radiation Therapy, Chemotherapy, Same Day Surgery Units, and Emergency Room. Review of pathology reports including private outpatient specimens and autopsy reports should also be included in casefinding.
2. **Disease Index** - Review of a Disease Index should be performed to verify all reportable cases are submitted to the PCR. If performed monthly, this review will simplify the annual reconciliation procedure (See *Part Four, Quality Control: PCR*) and aid in timeliness of reporting.
3. **Required Fields** - All data items required by the PCR must be submitted for each record. For a listing of these items, refer to *Appendix F*. Entries for each required data item must include specific demographic, diagnostic and treatment information that accurately reflects what is documented in the health record.

Accuracy

1. **Text Fields** - The *Required Data Set for Reporting Facilities* includes text fields (See *Appendix F*). The reason for requiring text is to enhance data accuracy. These fields give hospitals the ability to convey information to validate data items, document clarifications, reconcile data item discrepancies, support unusual site/histology combinations, provide history of previous cancers/reportable tumors, and explain any unusual or potentially questionable entry on the abstract. Required text information must be recorded in the designated text fields. (See also *Part Three, Data Item Instructions, Guidelines for Reporting Text*).
2. **Computer Edits** - Computer edits should be an integral component of any electronic abstracting system. These edits should check for completion of all required fields, allowable values and ranges, and interfield consistency. Edit checks should be performed on each completed abstract. Abstracts should be re-edited if any changes are made.

Abstract Plus includes the PCR required edits. A copy of the PCR edits is also provided to the cancer registry software vendors. All cases submitted to the PCR should be error free.

3. **Visual Editing** - The completed abstract should be visually reviewed to identify errors not detectable by the computer. Inconsistencies among data items could be identified when comparing text to coded items, e.g., stage coded to local with text indicating lymph node involvement.
4. **Physician Input** - Physicians should serve as resources to the abstractor. They should be consulted when questions arise during abstracting. Physician input may assist in identifying a primary site or provide clarification of conflicting statements or reports in the health record. Documentation of the physician input should be included in the text to support abstracted data.

Timeliness

1. 180 Days - 90% of the records must be received by the PCR within 180 days from *Date of Inpatient Disch* if an inpatient or *Date of 1st Contact* if an outpatient.
2. PCR Deadline - The first working day in July is the deadline for submitting all reportable cases seen at the reporting facility during the previous year.
3. PCR Reporting Schedule - This schedule should be followed to assure abstracts are received by the PCR within the required 180 days.

Cases with a Date of Inpatient Disch/ Date of 1st Contact in:	Upload on or before the 15th of:
January	June of same year
February	July of same year
March	August of same year
April	September of same year
May	October of same year
June	November of same year
July	December of same year
August	January of following year
September	February of following year
October	March of following year
November	April of following year
December	May** of following year

Example 1: All cases with a Date of Inpatient Disch/Date of 1st Contact on or between January 1 and January 31, 2010 must be uploaded by June 15, 2010.

Example 2: All cases with a Date of Inpatient Disch/Date of 1st Contact on or between December 1 and December 31, 2010 must be uploaded by May 15, 2011.

**The PCR deadline has not changed. The six weeks between May 15th and July 1st should be used to perform Quality Assurance procedures to ensure all cases for the year have been identified and reported. These cases may fall into the 10% over 180 days. This is expected and acceptable.

Note: This schedule should be used by reporting facilities as a guideline to assess timeliness of reporting but will not be used by the PCR to determine exact timeliness rates for reporting facilities. Reports provided by the PCR will show specific timeliness rates based on the number of days from *Date of Inpatient Disch* or *Date of 1st Contact* and the date the abstract was received by the PCR.

4. Incomplete and Suspense Cases - At a registry hospital, after identifying a potential case for the registry from a casefinding source, cases unable to be completely abstracted may be placed in an electronic suspense file. At a non-registry hospital using Abstract Plus software, incomplete abstracts may be saved as incomplete creating an electronic suspense file. A system should be in place to monitor these cases so they are completed and reported to the PCR in a timely manner. A case will not export out of Abstract Plus if it is incomplete.

Note: Incomplete/suspense cases should not be submitted to the PCR.

5. Method to Assure Timeliness - Review the Disease Index monthly using the reporting schedule as a guide to verify all reportable cases have been submitted within the 180-day timeframe.

QUALITY CONTROL: PCR

Quality control activities are conducted by the PCR to assure data in the central registry are complete, accurate, and timely. These activities fall into three categories: 1) internal procedures as data are processed, 2) on-site quality assessment reviews, and 3) trainings conducted by PCR staff or in conjunction with other organizations. These three major aspects of the PCR quality control program are described below.

Internal Quality Control Procedures

The quality control procedures described below are performed by the PCR routinely to enhance the quality of cancer data in the central cancer registry.

1. Completeness

- a. PCR Reporting Sources - The PCR establishes reporting from sources required to report and reporting through state data exchange agreements to assure all reportable cases are received. The PCR reporting sources (See *Part One, Reporting Requirements, PCR Reporting Sources*) include the following:
 - Acute Care Hospitals
 - Laboratories
 - Non Hospital Sources
 - States with Data Exchange Agreements
- b. Non-Reporting - All hospitals are required to submit on the 15th of every month or the last working day before the 15th if the 15th falls on a weekend or holiday. A listing of hospitals that have not submitted for two consecutive months is generated monthly at the PCR. A PCR Field Representative contacts hospitals appearing on this list and appropriate action is taken.
- c. Reconciliation - An annual comparison is made of each hospital's Disease Index with the PCR database to assure all cases have been reported. Each hospital receives a listing of cases identified as not being reported to the PCR with instructions to review each record to determine if the case is reportable. Cases missed, but now identified, must be reported. Cases that are not reportable must have justification documented on the listing explaining why the case is not reportable. Missed cases and listings must be returned to the PCR by the specified deadline.
- d. Death Clearance - The PCR conducts a Death Clearance procedure annually. This process involves identifying Pennsylvania Death Certificates with a reportable cause of death and matching them to the PCR files. Non-matched death certificates are potentially missed cases. Hospital contacts receive a listing of non-matched patients who expired at their hospital to determine if they were reportable. Missed cases must be reported. Cases that were not reportable must have justification documented on the listing. Missed cases and listings must be returned to the PCR by a specified deadline. At the conclusion of this process, the remaining non-matched cases are reviewed and may be abstracted at the PCR from the death certificates and defined as Death Certificate Only (DCO) cases. A DCO percentage (The number of DCO cases divided by the total number of incidence cases for that year) is computed. The PCR DCO percentage is

measured against the North American Association of Central Cancer Registries (NAACCR) DCO standard, which states a registry should have fewer than 5% DCO's in a given year.

2. Accuracy

- a. Computer Edits - Computer edits are performed on 100% of abstracts and consolidated records. The PCR utilizes a combination of NAACCR, SEER, and COC edits from the NAACCR metafile with PCR-developed edits added. These edits check for completion of all required fields, allowable ranges, allowable values, and interfield consistency. They check for invalid entries such as impossible site/histology combinations or flag unusual entries for review. PCR Field Representatives follow-up with hospital contacts and provide feedback on errors found.
- b. Visual Editing - Records are reviewed for consistency between coded data items and text documentation. This type of review is performed to detect discrepancies not detectable by the computer. PCR Field Representatives provide hospital contacts with feedback on these reviews.
- c. Electronic Reporting Approvals - An approval process is required for new contacts, hospital software changes, and updated NAACCR formats. Hospital shipments are monitored and PCR Field Representatives provide feedback to hospital contacts until acceptable accuracy is achieved.

The steps required for approval of electronic reporting are as follows:

- i. Trial Shipment - After all electronic reporting specifications have been met and records have been abstracted into the software, a trial shipment shall be prepared and submitted to the PCR for evaluation. The following must be included in the trial shipment:
 - a. Electronic File - The electronic file must be uploaded and contain 10-20 actual records not previously submitted to the PCR.
 - b. Paper Abstracts - An abstract from your software must be printed for each record on the trial shipment. All PCR required data items including text to support diagnostic findings, primary site, laterality, histology, behavior, grade, summary stage, and treatment must be printed on each abstract.
 - c. Copies of Supportive Documentation - Copies of supportive documentation from the medical record including pathology report, discharge summary, operative report, consultations, progress notes, radiology reports, and admission record must be included for each record on the electronic file. Paper documentation is used to verify the accuracy of required data fields reported in the electronic record.
- ii. Evaluation - The trial shipment is evaluated by the PCR for compliance to electronic reporting specifications, format, data quality, and completion of text fields. Hospitals may continue to abstract cases into their software system while the PCR is reviewing the trial shipment; however, no additional cases shall be submitted to the PCR until feedback is received on the current trial shipment.
- iii. Feedback - The PCR Field Representative will provide written feedback to the hospital contact to convey the results of the review. Errors and/or data items needing clarification will be identified and must be corrected or addressed in the trial shipment and for cases completed while the trial shipment was being reviewed. Additional trial shipments may be requested to resolve problems identified during evaluation(s).

- iv. Approval - When all aspects of the evaluation are acceptable, written approval for electronic reporting will be sent to the hospital contact. Approved hospitals do not have to send paper abstracts and supporting paper documentation with electronic files.
 - d. Unknown Values - The frequency of “*unknown*” or code for unknown in data items, such as age at diagnosis, sex, race, state, and county is monitored and follow-up is performed to eliminate as many unknowns as possible.
 - e. Resolution of Duplicates - To assure accuracy of incidence statistics, an incidence file containing all cases for a specified time period is created and a report is generated listing all cases alphabetically by last name. Cases with the same name are identified. Those determined to be the same person are then reviewed manually to determine whether they represent multiple primaries or duplications. While cases determined to be duplicates are deleted from the file, source records are retained and attached to the appropriate tumor in the PCR database.
3. Timeliness
- a. PCR Timeliness Standard - At least 90% of the records must be received by the PCR within 180 days from *Date of Inpatient Disch* if an inpatient or *Date of 1st Contact* if an outpatient.
 - b. Closeout Deadline -The first working in July is the deadline for submitting all reportable cases diagnosed/treated in the prior year.
 - c. Closeout Notification - Hospitals are notified annually of the closeout deadline and requested to notify the PCR when they anticipate closing out. Failure to meet the July deadline results in referral of the hospital to the Department of Health, Bureau of Facility Licensure and Certification.

On-Site Quality Assessment Review

Quality Assessment Reviews are routinely conducted at hospitals. Hospitals are scheduled for a review when certain criteria are met, such as unsatisfactory results from previous review, inability to perform annual reconciliation, reporting problems, and time lapse since last review. The reviews are designed to determine the quality of reporting to the PCR. During the review, casefinding completeness, data quality and timeliness of reporting are evaluated by PCR Field Representatives.

1. Notification of Quality Assessment Review- Hospitals receive a scheduling letter one month prior to the date of review. The scheduling letter includes:
 - a. Date and time of the review
 - b. *Reconciliation List*- sample of patients included on most recent reconciliation that were justified as not being reported.
 - c. *Reabstracting List*- list of randomly selected site specific cases reported to the PCR within the last twelve months that will be reabstracted by a PCR Field Representative
 - d. Request for private area with adequate work space for the PCR Field Representative
2. Hospital Preparation for a Quality Assessment Review- Hospitals must have the following available the day of the review:

- a. Health records for the patients on the *Reconciliation List*. The patient's complete health record must be provided including all inpatient and outpatient records.
 - b. Health records and copies of corresponding abstracts for all the cases on the *Reabstracting List*. All admissions used to abstract the case must be provided.
3. On Site Review Process- the PCR Field Representative will evaluate the following during their visit:
- a. Casefinding Completeness- The first component of the quality assessment review is the casefinding audit. The audit is a review and evaluation of the effectiveness of a facility's casefinding mechanisms used in submitting reportable cases to the PCR. The objective of the audit is to determine whether all reportable records are being identified and submitted to the PCR to insure PCR data accurately reflect cancer incidence in Pennsylvania.

The PCR Field Representative reviews the health records from the *Reconciliation List* to determine if these records are reportable and to identify any weaknesses or trends in a hospital's casefinding procedures.

The results of the casefinding audit are defined in terms of a completeness rate. The completeness rate indicates the percentage of reportable records submitted by the hospital to the PCR. The PCR acceptable completeness rate is 95 to 100%.

- b. Data Quality- The second component of the quality assessment review is a reabstracting study to evaluate data quality. Reabstracting compares the information in the health record to the previously abstracted data to determine the accuracy and completeness of the data. The PCR Field Representative reabstracts the cases on the *Reabstracting List* to identify any inaccurate information or misunderstandings of reporting guidelines.

The results of the reabstracting study are defined in terms of an accuracy rate. The accuracy rate indicates the percentage of data items reported correctly. The PCR standard for data quality is an accuracy rate of 95 to 100%.

- c. Timeliness- The third component of the quality assessment review is timeliness of reporting. For the PCR to provide timely statistics and reports, facilities must submit data in a timely manner. The timeliness standard established by the PCR to monitor hospital reporting requires at least 90% of the hospital's records be received by the PCR within 180 days from *Date of Inpatient Disch* if an inpatient or *Date of 1st Contact* if an outpatient. To evaluate timeliness, the PCR Field Representative uses reports generated by the PCR and assessment of cases currently being abstracted based on the reporting schedule (See *Quality Control, PCR Reporting Schedule*).
- d. Summation- At the conclusion of the review, the PCR Field Representative discusses findings and recommendations with appropriate hospital personnel during a summation conference. This provides the PCR Field Representative the opportunity to provide feedback relative to areas of compliance and concern. It also enables hospital personnel to be aware of the results of the review and ask questions regarding the findings and recommendations.
- e. Quality Assessment Review Report-The PCR sends a written report documenting findings, problems, recommendations, and rates to the hospital. A listing of missed records identified as reportable to the PCR and a listing of data items requiring correction are included in the report.

- f. Reporting Review Deficiencies- Hospital staff must submit the missed records and corrections to the PCR within 30 days of when they receive the report.
- g. Tracking Results- Upon completion of the Quality Assessment Review Report, completeness and accuracy rates by year review performed are entered into a tracking system at the PCR. This information provides a concise summary of review results for use in determining a hospital's performance over time and in identifying hospitals requiring more intense follow up.

Trainings

Education is an important part of quality control. The PCR offers trainings throughout the year. These trainings provide specific information on state reporting requirements and cancer data collection. For more information about training opportunities currently being offered, contact your PCR Field Representative.

**APPENDIX A:
PENNSYLVANIA CANCER CONTROL,
PREVENTION AND RESEARCH ACT
(P.L. 1241, No. 224)**

Official Advance Copy of Statute Enacted at 1980 Session

No. 1980-224

AN ACT

HB 230

Creating the Pennsylvania Cancer Control, Prevention and Research Advisory Board, providing authorization for the Secretary of Health, upon the recommendation of the Pennsylvania Cancer Control, Prevention and Research Advisory Board, to award grants and contracts for cancer control, prevention and research to associations organized in Pennsylvania and to governmental agencies in Pennsylvania.

The General Assembly of the Commonwealth of Pennsylvania hereby enacts as follows:

Section 1. Short title.

This act shall be known and may be cited as the "Pennsylvania Cancer Control, Prevention and Research Act."

Section 2. Definitions.

The following words and phrases when used in this act shall have, unless the context clearly indicates otherwise, the meanings given to them in this section:

"Board." The Pennsylvania Cancer Control, Prevention and Research Advisory Board established by this act.

"Cancer." All malignant neoplasms, regardless of the tissue of origin, including malignant lymphoma and leukemia.

"Secretary." The Secretary of Health of the Commonwealth of Pennsylvania.

Section 3. Pennsylvania Cancer Control, Prevention and Research Advisory Board.

(a) There is hereby created in the Department of Health the "Pennsylvania Cancer Control, Prevention and Research Advisory Board." The board shall consist of 11 members, all of whom shall be Pennsylvania residents, ten of whom the Governor shall appoint by and with the consent of a majority of the Senate. Of the ten appointed, three shall be distinguished scientists and physicians in the field of cancer, one shall be a qualified professional nurse engaged in the practice of oncological nursing, one shall be skilled in health care administration and two with substantial experience in the field of public health, one of whom shall be a professional nurse engaged in the practice of community health nursing, and three consumer members. The Secretary of Health shall be a member of the board.

(b) The terms of the members shall be four years from the respective date of their appointment except that the initial appointments shall be made in such a manner so that four members be appointed for terms of four years, three members be appointed for terms of three years, and three members be appointed for terms of two years.

(c) A chairman shall be appointed by the Governor for a term of four years.

(d) The board shall meet no less than four times annually at the call of the chairman or, in his absence or incapacity at the call of the Secretary of Health. Six members of the board shall constitute a quorum for the purpose of exercising all of the powers of the board. A vote of the majority of the members present shall be sufficient for all actions of the board.

(e) Each board member, except the secretary, shall receive actual travelling expenses and other necessary expenses.

(f) No member of the board shall participate in any discussions and decisions to recommend grants or contracts to any qualified association or to any agency of the Commonwealth or its political subdivisions with which the member is associated as a member of the governing body or as an employee, or with which the member has entered into any contractual arrangement.

Section 4. Responsibilities of the board.

(a) The board shall have the power to prescribe, amend and repeal bylaws governing the manner in which the business of the board is conducted.

(b) The board shall advise the secretary with respect to cancer control, prevention and research in Pennsylvania.

(c) The board shall approve each year a program for cancer control, prevention and research to be known as the "Pennsylvania Cancer Plan."

(d) In order to implement in whole or in part the Pennsylvania Cancer Plan, the board shall recommend to the secretary the awarding of grants and contracts to qualified associations or governmental agencies in order to plan, establish or conduct programs in cancer control or prevention, cancer education and training and cancer clinical research.

(e) Grants and contracts may be recommended for:

- (1) Cancer registry.
- (2) Cancer screening, detection and prevention.
- (3) Cancer epidemiology and biostatistical studies.
- (4) Cancer community outreach programs.
- (5) Cancer rehabilitation.
- (6) Communication and planning among cancer institutions.
- (7) Cancer education and information.
- (8) Cancer training.
- (9) Cancer clinical research.

(f) Consistent with the Pennsylvania Cancer Plan the board shall give its first priority to funding grants and contracts relating to subsection (e)(1), (2) and (3); second priority to funding grants and contracts relating to subsection (e)(4), (5) and (6); third priority to funding grants and contracts relating to subsection (e)(7), (8) and (9).

(g) The following criteria shall be given consideration for recommending grants and contracts for programs:

- (1) the relevancy of applicant's proposal to the Pennsylvania Cancer Plan; and
- (2) the feasibility of the applicant's proposal.

(h) The board shall recommend to the secretary rules and regulations consistent with law as it may deem necessary for the performance of its duties and the proper administration of this act.

(i) The board shall report annually to the Governor and the General Assembly. The report shall include, but not be limited to, a full description of the grants and contracts funded pursuant to this act, the amount of the grant or contract, an outline of the proposal on which the grant was based, and the results achieved as a result of the grant.

Section 5. Responsibilities of the secretary.

(a) The secretary shall award grants and contracts only from among those recommended by the board to qualified Pennsylvania associations and governmental agencies in order to plan, establish or conduct programs in cancer control and prevention, cancer education and training and cancer research. The secretary may request additional recommendations from the board.

(b) The secretary shall provide such staff, information and other assistance as the secretary may deem necessary for the completion of the board's responsibilities. Such staff shall be responsible to the secretary.

Section 6. Cancer registry.

(a) The Department of Health shall establish a system for the Statewide collection and dissemination of data on cases of cancer by anatomical site, medical and occupational history of patients, stage of disease and other data necessary to effectuate the provisions of this act as determined by the department.

(b) Persons in charge of hospitals and laboratories shall be required by the Department of Health, in accordance with its regulations adopted with the advice of the board to report cases of cancer on forms furnished by the department.

(c) The reports required pursuant to this act shall be confidential and not open to public inspection or dissemination. This shall not restrict the collection and analysis of data by the Department of Health or those with whom the department contracts, subject to strict supervision by the Department of Health to insure that the use of the reports is limited to specific research purposes.

Section 7. Sunset provisions.

With the exception of section 6, this act shall expire on June 30, 1984, unless otherwise extended by an act of the General Assembly.

Section 8. Effective date.

This act shall take effect January 1, 1981.

APPROVED—The 18th day of December, A. D. 1980.

DICK THORNBURGH

PRINTER'S NO. 2130

THE GENERAL ASSEMBLY OF PENNSYLVANIA

SENATE BILL

No. 1607 Session of 1996

INTRODUCED BY THOMPSON, ROBBINS, PETERSON, ULIANA, DELP, HART, WENGER AND MADIGAN, JUNE 18, 1996

REFERRED TO PUBLIC HEALTH AND WELFARE, JUNE 18, 1996

AN ACT

1 Amending the act of December 18, 1980 (P.L.1241, No.224),
2 entitled "An act creating the Pennsylvania Cancer Control,
3 Prevention and Research Advisory Board, providing
4 authorization for the Secretary of Health, upon the
5 recommendation of the Pennsylvania Cancer Control, Prevention
6 and Research Advisory Board, to award grants and contracts
7 for cancer control, prevention and research to associations
8 organized in Pennsylvania and to governmental agencies in
9 Pennsylvania," extending the expiration date.

10 The General Assembly of the Commonwealth of Pennsylvania
11 hereby enacts as follows:

12 Section 1. Section 7 of the act of December 18, 1980
13 (P.L.1241, No.224), known as the Pennsylvania Cancer Control,
14 Prevention and Research Act, reenacted and amended November 25,
15 1988 (P.L.1086, No.126) and amended June 30, 1992 (P.L.334,
16 No.67), is amended to read:

17 Section 7. Sunset provisions.

18 With the exception of section 6, this act shall expire on
19 June 30, [1996] 2006, unless otherwise extended by an act of the
20 General Assembly.

21 Section 2. This act shall take effect immediately.

E29L35JS/19960S1607B2130

APPENDIX B:
TITLE 28. HEALTH and SAFETY
PART III. PREVENTION of DISEASES
CHAPTER 27. COMMUNICABLE and
NONCOMMUNICABLE DISEASES

Annex A
TITLE 28. HEALTH AND SAFETY
PART III. PREVENTION OF DISEASES
CHAPTER 27. COMMUNICABLE AND NONCOMMUNICABLE DISEASES

Subchapter A. GENERAL PROVISIONS

§ 27.1. Definitions.

The following words and terms, when used in this chapter, have the following meanings, unless the context clearly indicates otherwise:

ACIP--The Advisory Committee on Immunization Practices of the Centers for Disease Control and Prevention, United States Department of Health and Human Services.

Caregiver--The entity or individual responsible for the safe and healthful care or education of a child in a child care group setting.

Case--A person or animal that is determined to have or suspected of having a disease, infection or condition.

Case report form--The form designated by the Department for reporting a case or a carrier.

Central office--Department headquarters located in Harrisburg.

Child--A person under 18 years of age.

Child care group setting--The premises in which care is provided at any one time to four or more children, unrelated to the operator.

Clinical laboratory--A laboratory for which a permit has been issued to operate as a clinical laboratory under the Clinical Laboratory Act (35 P. S. §§ 2151--2165).

Communicable disease--An illness which is capable of being spread to a susceptible host through the direct or indirect transmission of an infectious agent or its toxic product by an infected person, animal or arthropod, or through the inanimate environment.

Communicable period--The time during which an etiologic agent may be transferred directly or indirectly from an infected person to another person, or from an infected animal to a person.

Contact--A person or animal known to have had an association with an infected person or animal which presented an opportunity for acquiring the infection.

District office--One of the district headquarters of the Department located within this Commonwealth.

Health care facility--

- (i) A chronic disease, or other type of hospital, a home health care agency, a hospice, a long-term care nursing facility, a cancer treatment center using radiation therapy on an ambulatory basis, an ambulatory surgical facility, a birth center, and an inpatient drug and alcohol treatment facility, regardless of whether the health care facility is operated for profit, nonprofit or by an agency of the Commonwealth or local government.
- (ii) The term does not include:
 - a) An office used primarily for the private practice of a health care practitioner.
 - b) A facility providing treatment solely on the basis of prayer or spiritual means in accordance with the tenets of any church or religious denomination.

- c) A facility conducted by a religious organization for the purpose of providing health care services exclusively to clergy or other persons in a religious profession who are members of a religious denomination.

Health care practitioner--An individual who is authorized to practice some component of the healing arts by a license, permit, certificate or registration issued by a Commonwealth licensing agency or board.

Health care provider--An individual, a trust or estate, a partnership, a corporation (including associations, joint stock companies and insurance companies), the Commonwealth, or a political subdivision, or instrumentality (including a municipal corporation or authority) thereof, that operates a health care facility.

Household contact--A person living in the same residence as a case, including a spouse, child, parent, relation or other person, whether or not related to the case.

Infectious agent--Any organism, such as a virus, bacterium, fungus or parasite, that is capable of being communicated by invasion and multiplication in body tissues and capable of causing disease.

Isolation--The separation for the communicable period of an infected person or animal from other persons or animals, in such a manner as to prevent the direct or indirect transmission of the infectious agent from infected persons or animals to other persons or animals who are susceptible or who may spread the disease to others.

LMRO--Local morbidity reporting office--A district office of the Department or a local health department.

Local health authority--A county or municipal department of health, or board of health of a municipality that does not have a department of health. The term includes a sanitary board.

Local health department--Each county department of health under the Local Health Administration Law (16 P. S. §§ 12001--12028), and each department of health in a municipality approved for a Commonwealth grant to provide local health services under section 25 of the Local Health Administration Law (16 P. S. § 12025).

Local health officer--The person appointed by a local health authority to head the daily administration of duties imposed upon or permitted of local health authorities by State laws and regulations.

Medical record--An account compiled by physicians and other health professionals including a patient's medical history; present illness; findings on physical examination; details of treatment; reports of diagnostic tests; findings and conclusions from special examinations; findings and diagnoses of consultants; diagnoses of the responsible physician; notes on treatment, including medication, surgical operations, radiation, and physical therapy; and progress notes by physicians, nurses and other health professionals.

Modified quarantine--A selected, partial limitation of freedom of movement determined on the basis of differences in susceptibility or danger of disease transmission which is designated to meet particular situations. The term includes the exclusion of children from school and the prohibition, or the restriction, of those exposed to a communicable disease from engaging in particular activities.

Monitoring of contacts--The close supervision of persons and animals exposed to a communicable disease without restricting their movement.

Operator--The legal entity that operates a child care group setting or a person designated by the legal entity to serve as the primary staff person at a child care group setting.

Outbreak--An unusual increase in the number of cases of a disease, infection or condition, whether reportable or not as a single case, above the number of cases that a person required to report would expect to see in a particular geographic area or among a subset of persons (defined by a specific demographic or other features).

Physician--An individual licensed to practice medicine or osteopathic medicine within this Commonwealth.

Placarding--The posting on a home or other building of a sign or notice warning of the presence of communicable disease within the structure and the danger of infection there from.

Quarantine--

- (i) The limitation of freedom of movement of a person or an animal that has been exposed to a communicable disease, for a period of time equal to the longest usual incubation period of the disease, or until judged noninfectious by a physician, in a manner designed to prevent the direct or indirect transmission of the infectious agent from the infected person or animal to other persons or animals.
- (ii) The term does not exclude the movement of a person or animal from one location to another when approved by the Department or a local health authority under § 27.67 (relating to the movement of persons and animals subject to isolation or quarantine by action of a local health authority or the Department).

Reportable disease, infection, or condition--A disease, infection, or condition, made reportable by § 27.2 (relating to specific identified reportable diseases, infections and conditions)

SHC--State Health Center--The official headquarters of the Department in a county, other than a district office.

Segregation--The separation for special control or observation of one or more persons or animals from other persons or animals to facilitate the control of a communicable disease.

Sexually transmitted disease--A disease which, except when transmitted perinatally, is transmitted almost exclusively through sexual contact.

Surveillance of disease--The continuing scrutiny of all aspects of occurrence and spread of disease that are pertinent to effective control.

Volunteer--A person who provides services to a school or child care group setting without receiving remuneration.

§ 27.2. Specific identified reportable diseases, infections and conditions.

The diseases, infections and conditions in Subchapter B (relating to the reporting of diseases, infections and conditions) are reportable to the Department or the appropriate local health authority by the persons or entities in the manner and within the time frames set out in this chapter.

§ 27.3. Reporting outbreaks and unusual diseases, infections and conditions.

- a) A person required to report under this chapter shall report an outbreak within 24 hours, and in accordance with § 27.4 (relating to reporting cases)
- b) A person required to report under this chapter who suspects a public health emergency, shall report an unusual occurrence of a disease, infection or condition not listed as reportable in Subchapter B (relating to reporting of diseases, infections and conditions) or defined as an outbreak, within 24 hours, and in accordance with § 27.4.
- c) Any unusual or group expression of illness which the Department designates as a public health emergency shall be reported within 24 hours, and in accordance with § 27.4.

§ 27.4. Reporting cases.

- a) Except for reporting by a clinical laboratory, a case is to be reported to the LMRO serving the area in which a case is diagnosed or identified unless another provision of this chapter directs that a particular type of case is to be reported elsewhere. A clinical laboratory shall make reports to the appropriate office of the Department.
- b) Upon the Department's implementation of its electronic disease surveillance system for certain types of case reports, persons who make those reports shall do so electronically using an application and reporting format provided by the Department. At least 6 months in advance of requiring a type of case

report to be reported electronically, the Department will publish a notice in the *Pennsylvania Bulletin* announcing when electronic reporting is to begin.

- c) This section does not prohibit a reporter from making an initial report of a case to the Department or an LMRO by telephone. The reporter will be instructed on how to make a complete case report at the time of the telephone call.
- d) Department offices to which this chapter requires specified case reports to be filed are as follows:
 - 1) Cancer Registry, Division of Health Statistics, Bureau of Health Statistics and Research.
 - 2) Division of Infectious Disease Epidemiology, Bureau of Epidemiology.
 - 3) HIV/AIDS Epidemiology Section, Division of Infectious Disease Epidemiology, Bureau of Epidemiology
 - 4) Division of Maternal and Child Health, Bureau of Family Health.
- e) A case shall be reported using the appropriate case report format. Information solicited by the case report form shall be provided by the reporter, irrespective of whether the report is made by submitting the form directly in hard copy or by telecommunication or electronic submission. An appropriate case report form or format may be procured from the office to which the type of case is reportable.

§ 27.5a. Confidentiality of case reports.

Case reports submitted to the Department or to an LMRO are confidential. Neither the reports, nor any information contained in them which identifies or is perceived by the Department or the LMRO as capable of being used to identify a person named in a report, will be disclosed to any person who is not an authorized employe or agent of the Department or the LMRO, and who has a legitimate purpose to access case information, except for any of the following reasons:

- 1) When disclosure is necessary to carry out a purpose of the act, as determined by the Department or LMRO, and disclosure would not violate another act or regulation.
- 2) When disclosure is made for a research purpose for which access to the information has been granted by the Department or an LMRO. Access shall be granted only when disclosure would not violate another act or regulation. The research shall be subject to strict supervision by the LMRO to ensure that the use of information disclosed is limited to the specific research purpose and will not involve the further disclosure of information which identifies or is perceived as being able to be used to identify a person named in a report.

§ 27.6. Disciplinary consequences for violating reporting responsibilities

- a) Failure of a clinical laboratory to comply with the reporting provisions of this chapter may result in restrictions being placed upon or revocation of the laboratory's permit to operate as a clinical laboratory, as provided for in the Clinical Laboratory Act (35 P. S. §§ 2151--2165) unless failure to report is due to circumstances beyond the control of the clinical laboratory.
- b) Failure of a Department licensed health care facility to comply with the reporting provisions of this chapter may result in restrictions being placed upon or revocation of the health care facility's license, as provided for in the Health Care Facilities Act (35 P. S. §§ 448.101--448.904b)
- c) Failure of a health care practitioner to comply with the reporting provisions of this chapter may result in referral of that matter to the appropriate licensure board for disciplinary action.
- d) Failure of a child care group setting to comply with the reporting provisions of this chapter may result in referral of that matter to the appropriate licensing agency for appropriate action.

§ 27.7. Cooperation between clinical laboratories and persons who order laboratory tests.

To facilitate the reporting of cases by clinical laboratories, the following is required:

- 1) When a clinical laboratory is requested to conduct a test which, depending upon the results, would impose a reporting duty upon the clinical laboratory, the clinical laboratory shall provide to the person who orders the testing, a form that solicits all information which is required for completion of the applicable case report form.

- 2) A person who orders testing subject to paragraph (1) shall, at the time of ordering the test, provide the clinical laboratory with the information solicited by the form which that person either possesses or may readily obtain.

§27.8. Criminal penalties for violating the act or this chapter.

- a) A person who violates any provision of the act or this chapter shall, for each offense, upon conviction thereof in a summary proceeding before a district justice in the county wherein the offense was committed, be sentenced to pay a fine of not less than \$25 and not more than \$300, together with costs, and in default of payment of the fine and costs, shall be imprisoned in the county jail for a period not to exceed 30 days.
- b) A person afflicted with communicable tuberculosis, ordered to be quarantined or isolated in an institution, who leaves without consent of the medical director of the institution, is guilty of a misdemeanor, and upon conviction thereof, shall be sentenced to pay a fine of not less than \$100 nor more than \$500, or undergo imprisonment for not less than 30 days nor more than 6 months, or both.
- c) Prosecutions may be instituted by the Department, by a local health authority, or by any person having knowledge of a violation of the act or this chapter.

**Subchapter B. REPORTING OF DISEASES, INFECTIONS AND CONDITIONS
GENERAL**

§ 27.21. Reporting of AIDS cases by physicians and hospitals.

A physician or a hospital is required to report a case of AIDS within 5 workdays after it is identified to the local health department if the case resides within the jurisdiction of that local health department. In all other cases, the physician or hospital shall report the case to the HIV/AIDS Epidemiology Section, Division of Infectious Disease Epidemiology, Bureau of Epidemiology.

§ 27.21a. Reporting of cases by health care practitioners and health care facilities.

- a) Except as set forth in this section or as otherwise set forth in this chapter, a health care practitioner or health care facility is required to report a case of a disease, infection or condition in subsection (b) as specified in § 27.4 (relating to reporting cases), if the health care practitioner or health care facility treats or examines a person who is suffering from, or who the health care practitioner suspects, because of symptoms or the appearance of the individual, of having a reportable disease, infection or condition:
 - 1) A health care practitioner or health care facility is not required to report a case if that health care practitioner or health care facility has reported the case previously.
 - 2) A health care practitioner or health care facility is not required to report a case of influenza unless the disease is confirmed by laboratory evidence of the causative agent.
 - 3) A health care practitioner or health care facility is not required to report a case of chlamydia trachomatis infection unless the disease is confirmed by laboratory evidence of the infectious agent.
 - 4) A health care practitioner or health care facility is not required to report a case of cancer unless the health care practitioner or health care facility provides screening, therapy or diagnostic services to cancer patients.
 - 5) Only physicians and hospitals are required to report cases of AIDS.
- b) The following diseases, infections and conditions in humans are reportable by health care practitioners and health care facilities within the specified time periods and as otherwise required by this chapter:
 - 1) The following diseases, infections and conditions are reportable within 24 hours after being identified by symptoms, appearance or diagnosis:

Animal bite.
 Anthrax.
 Arboviruses.
 Botulism.
 Cholera.
 Diphtheria.
 Enterohemorrhagic E. coli.
 Food poisoning outbreak.
 Haemophilus influenzae invasive disease.
 Hantavirus pulmonary syndrome.
 Hemorrhagic fever.
 Lead poisoning.
 Legionellosis.
 Measles (rubeola).
 Meningococcal invasive disease.
 Plague.
 Poliomyelitis.
 Rabies.
 Smallpox
 Typhoid fever

- 2) The following diseases, infections and conditions are reportable within 5 work days after being identified by symptoms, appearance or diagnosis:

<p>AIDS Amebiasis. Brucellosis. Campylobacteriosis. Cancer. Chancroid. Chickenpox (varicella) Chlamydia trachomatis infections. Creutzfeldt-Jakob Disease. Cryptosporidiosis. Encephalitis. Giardiasis. Gonococcal infections. Granuloma inguinale. Guillain-Barre syndrome. Hepatitis, viral, acute & chronic cases. Histoplasmosis. Influenza. Leprosy (Hansen's disease). Leptospirosis. Listeriosis. Lyme disease. Lymphogranuloma venereum. Malaria. Maple syrup urine disease (MSUD) in children under 5 years of age. Meningitis (All types not caused by invasive Haemophilus influenza or Neisseria meningitis). Mumps.</p>	<p>Pertussis (whooping cough). Phenylketonuria (PKU) in children under 5 years of age. Primary congenital hypothyroidism in children under 5 years of age. Psittacosis (ornithosis). Rickettsial diseases. Rubella (German measles) and congenital rubella syndrome. Salmonellosis. Shigellosis. Sickle cell hemoglobinopathies in children under 5 years of age. Staphylococcus aureus, Vancomycin-resistant (or intermediate) invasive disease. Streptococcal invasive disease (group A). Streptococcus pneumoniae, drug-resistant invasive disease. Syphilis (all stages). Tetanus. Toxic shock syndrome. Toxoplasmosis. Trichinosis. Tuberculosis, suspected or confirmed active disease (all sites). Tularemia.</p>
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- c) A school nurse shall report to the LMRO any unusual increase in the number of absentees among school children. A caregiver at a child care group setting shall report to the LMRO any unusual increase in the number of absentees among children attending the child care group setting.

- d) A health care facility or health care practitioner providing screening, diagnostic or therapeutic services to patients with respect to cancer shall also report cases of cancer as specified in § 27.31 (relating to reporting cases of cancer).

§ 27.22. Reporting of cases by clinical laboratories.

- a) A person who is in charge of a clinical laboratory in which a laboratory examination of a specimen derived from a human body yields evidence significant from a public health standpoint of the presence of a disease, infection or condition listed in subsection (b) shall promptly report the findings, no later than the next work day after the close of business on the day on which the examination was completed, except as otherwise noted in this chapter.

- b) The diseases, infections and conditions to be reported include the following:

Amebiasis.	Phenylketonuria (PKU) in children under 5 years of age.
Anthrax.	Primary congenital hypothyroidism in children under 5 years of age.
An unusual cluster of isolates.	Plague.
Arboviruses	Poliomyelitis.
Botulism--all forms.	Psittacosis (ornithosis).
Brucellosis.	Rabies.
Campylobacteriosis.	Respiratory syncytial virus.
Cancer.	Rickettsial infections.
Chancroid.	Rubella.
Chickenpox (varicella).	Salmonella.
Chlamydia trachomatis infections.	Shigella.
Cholera.	Sickle cell hemoglobinopathies in children under 5 years of age.
Creutzfeldt-Jakob disease.	Staphylococcus Aureus Vancomycin-resistant (or intermediate) invasive disease.
Cryptosporidiosis.	Streptococcus pneumoniae, drug-resistant invasive disease.
Diphtheria infections.	Syphilis.
Enterohemorrhagic E. coli 0157 infections, or infections caused by other sub-types producing shiga-like toxin.	Tetanus.
Giardiasis.	Toxoplasmosis.
Gonococcal infections.	Trichinosis.
Granuloma inguinale.	Tuberculosis, confirmation of positive smears or cultures, including results of drug susceptibility testing.
Haemophilus influenzae infections--invasive from sterile sites.	Tularemia.
Hantavirus.	Typhoid.
Hepatitis, viral, acute and chronic cases.	
Histoplasmosis.	
Influenza.	
Lead poisoning.	
Legionellosis.	
Leprosy (Hansen's disease).	
Leptospirosis.	
Listeriosis.	
Lyme disease.	
Lymphogranuloma venereum.	
Malaria.	
Maple syrup urine disease (MSUD) in children under 5 years of age.	
Measles (rubeola).	
Meningococcal infections--invasive from sterile sites.	
Mumps.	
Pertussis.	

- c) The report shall include the following:
- 1) The name, age, address and telephone number of the person from whom the specimen was obtained.
 - 2) The date the specimen was collected.
 - 3) The source of the specimen (such as, serum, stool, CSF, wound).
 - 4) The name of the test or examination performed and the date it was performed.
 - 5) The results of the test.
 - 6) The range of normal values for the specific test performed.
 - 7) The name, address, and telephone number of the physician for whom the examination or test was performed.
 - 8) Other information requested in case reports or formats specified by the Department.
- d) The report shall be submitted by the person in charge of a laboratory, in either a hard copy format or an electronic transmission format specified by the Department.
- e) Reports made on paper shall be made to the LMRO where the case is diagnosed or identified. Reports made electronically shall be submitted to the Division of Infectious Disease Epidemiology, Bureau of Epidemiology. Reports of maple syrup urine disease, phenylketonuria, primary congenital hypothyroidism, sickle cell hemoglobinopathies, cancer and lead poisoning shall be reported to the location specifically designated in this subchapter. See §§ 27.30, 27.31 and 27.34 (relating to reporting cases of certain diseases in the newborn child; reporting cases of cancer; and reporting cases of lead poisoning).
- f) A clinical laboratory shall submit isolates of salmonella and shigella to the Department's Bureau of laboratories for serotyping within 5 work days of isolation.
- g) A clinical laboratory shall submit isolates of Neisseria meningitidis obtained from a normally sterile site to the Department's Bureau of Laboratories for serogrouping within 5 work days of isolation.
- h) A clinical laboratory shall send isolates of enterohemorrhagic E. coli to the Department's Bureau of Laboratories for appropriate further testing within 5 work days of isolation.
- i) A clinical laboratory shall send isolates of Haemophilus influenzae obtained from a normally sterile site to the Department's Bureau of Laboratories for serotyping within 5 work days of isolation.
- j) The Department, upon publication of a notice in the *Pennsylvania Bulletin*, may authorize changes in the requirements for submission of isolates based upon medical or public health developments when the departure is determined by the Department to be necessary to protect the health of the people of this Commonwealth. The change will not remain in effect for more than 90 days after publication unless the Board acts to affirm the change within that 90-day period.

§ 27.23. Reporting of cases by persons other than health care practitioners, health care facilities, veterinarians or laboratories.

Except with respect to reporting cancer, individuals in charge of the following types of group facilities identifying a disease, infection or condition listed in § 27.21a (relating to reporting of cases by health care practitioners and health care facilities) by symptom, appearance or diagnosis shall make a report within the time frames required in § 27.21a.

- 1) Institutions maintaining dormitories and living rooms.
- 2) Orphanages.
- 3) Child care group settings.

§ 27.24. (Reserved).

§ 27.24a. Reporting of cases by veterinarians.

A veterinarian is required to report a case, as specified in § 27.4 (relating to reporting cases), only if the veterinarian treats or examines an animal which the veterinarian suspects of having a disease set forth in § 27.35(a) (relating to reporting cases of disease in animals).

§§ 27.25--27.28. (Reserved).

§ 27.29. Reporting for special research projects.

A person in charge of a hospital or other institution for the treatment of disease shall, upon request of the Department, make reports of a disease or condition for which the Board has approved a specific study to enable the Department to determine and employ the most efficient and practical means to protect and to promote the health of the people by the prevention and control of the disease or condition. The reports shall be made on forms prescribed by the Department and shall be transmitted to the Department or to local health authorities as directed by the Department.

DISEASES AND CONDITIONS REQUIRING SPECIAL REPORTING**§ 27.30. Reporting cases of certain diseases in the newborn child.**

Reports of maple syrup urine disease, phenylketonuria, primary congenital hypothyroidism and sickle cell hemoglobinopathies shall be made to the Division of Maternal and Child Health, Bureau of Family Health, as specified in Chapter 28 (relating to metabolic diseases of the newborn) and those provisions of § 27.4 (relating to reporting cases) consistent with Chapter 28 and this section.

§ 27.31. Reporting cases of cancer.

- a) A hospital, clinical laboratory, or other health care facility providing screening, diagnostic or therapeutic services for cancer to cancer patients shall report each case of cancer to the Department in a format prescribed by the Cancer Registry, Bureau of Health Statistics and Research, within 180 days of the patient's discharge, if an inpatient or, if an outpatient, within 180 days following diagnosis or initiation of treatment.
- b) A health care practitioner providing screening, diagnostic or therapeutic services to cancer patients for cancer shall report each cancer case to the Department in a format prescribed by the Cancer Registry, Bureau of Health Statistics and Research, within 5 work days of diagnosis. Cases directly referred to or previously admitted to a hospital or other health care facility providing screening, diagnostic or therapeutic services to cancer patients in this Commonwealth, and reported by those facilities, are exceptions and do not need to be reported by the health care practitioner.
- c) The Department or its authorized representative shall be afforded physical access to all records of physicians and surgeons, hospitals, outpatient clinics, nursing homes and all other facilities, individuals or agencies providing services to patients which would identify cases of cancer or would establish characteristics of the cancer, treatment of the cancer or medical status of any identified cancer patient.
- d) Reports submitted under this section are confidential and may not be open to public inspection or dissemination. Information for specific research purposes may be released in accordance with procedures established by the Department with the advice of the Pennsylvania Cancer Control, Prevention and Research Advisory Board.
- e) Case reports of cancer shall be sent to the Cancer Registry, Division of Health Statistics, Bureau of Health Statistics and Research, unless otherwise directed by the Department.

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APPENDIX C:
WEB PLUS UPLOAD INSTRUCTIONS

**Procedure: Web Plus file upload**

Purpose: To send files via a secure website to the Pennsylvania Cancer Registry (PCR). This procedure is used when facilities have files ready to submit to the PCR (for example, monthly data submissions). Web Plus replaces the need for facilities to submit diskettes or CDs through the U.S. mail.

General Information:

1. **Security:** Web Plus is an Internet-based application developed by the Centers for Disease Control and Prevention (CDC), National Program of Cancer Registries (NPCR). Web Plus has been designed as a highly secure application that can be used to transmit data between reporting facilities and the PCR safely over the public internet.

Security is achieved by a combination of software features and network infrastructure. Web Plus is hosted on a secure Web server; the communication between the client and the server is encrypted with 128-bit encryption Secure Socket Layer (SSL) technology.

Security features of the application include:

- Web Plus keeps an extensive log of user logins, data accesses, and updates for auditing purposes.
 - User accounts can be locked out if invalid login attempts exceed a threshold value, configurable by the PCR Central Administrator.
 - Initial passwords are randomly generated by the system and the user will be forced to change it after their first successful login.
 - Current user activities are visible to the PCR Central Administrator through the Current user Activities page.
 - User passwords are stored in the database using a one-way hash encryption method.
 - The Web Plus configuration file will store the connection string to the SQL Server database in encrypted format.
 - The application times out after a specified time period.
 - Web Plus uses form-based authentication where users are required to enter their unique user ID and strong password to be authenticated by the application.
2. **Screen Resolution:** The resolution for Web Plus should be 1024 X 768. If the resolution set on your PC is different, you may still be able to use Web Plus, but Web Plus has been designed to be viewed best at 1024 X 768 or higher. You will receive a message on the Web Plus log-in screen if your resolution is not set correctly. Contact your IT department if you are unsure how to change your screen resolution.

3. Web Plus icon on desk top: It is recommended that your IT Department create an icon on your desk top from the Web Plus link for easy access to the application.
4. Password protected or encrypted files: Files uploaded to the PCR via Web Plus MUST NOT be password protected or encrypted. The security features of Web Plus replaces the need to password protect or encrypt files.
5. Password changes: You will be prompted to change your password the first time you log into Web Plus and then every 60 days after that.
6. Edits: When you upload your data file, edits will be run and an edit report will be made available for your review. PCR staff will monitor files submitted with errors and work individually with those facilities to determine the reason for the edit errors and the best way to correct errors before submission. **Note for Abstract Plus users**: Web Plus contains the same edit set that is in Abstract Plus. Since records cannot be exported until all errors are corrected, you should not receive any edit errors when uploading files through Web Plus.
7. What and when to submit: Reporting facilities must upload files by the 15th of every month as stated in the PCR Manual under “When To Report”. Files may be submitted more than once per month but no more frequently than once a week. Accession lists for the files uploaded are not required. A hard copy of the PCR Transmittal Form is no longer required unless you have zero (0) records to report (see #8).
8. No records to report: If a facility has no cancer records to report, an electronic PCR transmittal form with zero (0) entered for the number of new records must be completed and submitted via Web Plus by the 15th of the month. In addition, the reason for not submitting any records must be documented on the transmittal form in the space provided.
9. Changing Information: Updates or changes to previously submitted records should not be submitted using Web Plus. As corrections are made to records previously accessioned by the PCR, print a paper abstract from your software system and highlight the changed information. The PCR is investigating alternative methods for receiving change sheets. Until a new procedure is implemented, these modifications should continue to be mailed to the PCR in a PCR postage paid envelope along with a PCR Transmittal Form (see Changing Information on page 17 of the PCR Manual).
10. File size: There is no limit to the number of records in each upload file.
11. Duplicate files: Web Plus has restrictions on uploading files that are exact duplicates of a previously uploaded file. If you attempt to upload a duplicate file, you will receive the below message:

The screenshot shows the 'Web Plus' interface for the Pennsylvania Cancer Registry. At the top right, it says 'Pennsylvania Cancer Registry' and '1-800-272-1850'. Below this is a navigation bar with links: Home, New Upload, Previous Uploads, Download Files, Change Password, Help, and Log out. The main heading is 'Upload Abstract Bundle'. Below this is a paragraph of instructions: 'Select your upload type: NAACCR V13.x, Non-NAACCR, or NAACCR V12. If you have selected either NAACCR file upload option, the files must be in the correct NAACCR version record layout. NOTE: If you are uploading a NAACCR version 13.0 file, edits will be automatically run upon upload of the file and the edits error report will open in a separate window. For files uploaded in NAACCR version 12.x or NAACCR version 13.0 file formats, the file will be uploaded and submitted for edits processing by your central registry. You will be notified via e-mail when your error report becomes available for viewing.' Below the instructions are three radio buttons: 'NAACCR V13.x File' (selected), 'Non-NAACCR File', and 'NAACCR V12 File'. There is a text input field labeled 'Select a file to upload:' with a 'Browse...' button next to it. At the bottom, there is a 'Comment' field.

If it is necessary to re-submit a file already exported, contact the PCR for assistance.

Procedure:

Uploading files

1. Open Web Plus using the link in the e-mail provided by the PCR.
2. Type your User ID and Password. Note: Your User ID and Password were previously sent to you via e-mail.

Click Log in.

REGISTRY PLUS

NPCR NATIONAL PROGRAM OF CANCER REGISTRIES

Welcome to Web Plus
Application for Secure Cancer Reporting Over the WWW

Pennsylvania Cancer Registry

pennsylvania
DEPARTMENT OF HEALTH
Web Plus V3.3.0

Please log in

User ID

Password

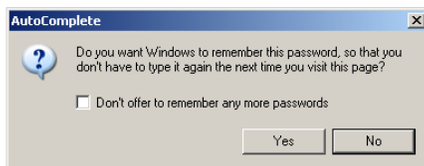
Log in

Notice to Users: Access to this system is restricted to authorized users. Unauthorized use of, or access to this resource may subject you to disciplinary action or criminal prosecution. If you are not authorized to access this resource, LOG OFF IMMEDIATELY.

HIPAA - WARNING
All users must comply with HIPAA PRIVACY RULE REQUIREMENTS while using this computer system, including -

- Log on only under your assigned user ID.
- Do not attempt to access health information that you are not authorized to use.
- Log off or lock up your workstation when it is unattended.

If the following message appears, click on No.



*The first time you log in, the screen below will appear, forcing you to change your password.

Enter a new password using the following criteria “Password must be between 8 to 20 characters, contain at least one digit and one alphabetic character, and must not contain any special characters”.

Change Password

You are required to change your password before proceeding further. Please enter your new password.

New password

Retype password

Click on Change.

If the password does not meet the criteria specified above or if the new password does not match the retype password line, you will receive a message ‘Password not changed’. The Change Password screen will remain until the password meets the criteria and the two password lines match.

3. The Web Plus home page for your facility opens.

Web Plus Pennsylvania Cancer Registry
1-800-272-1850

Web Plus Home Page for First Name Last Name

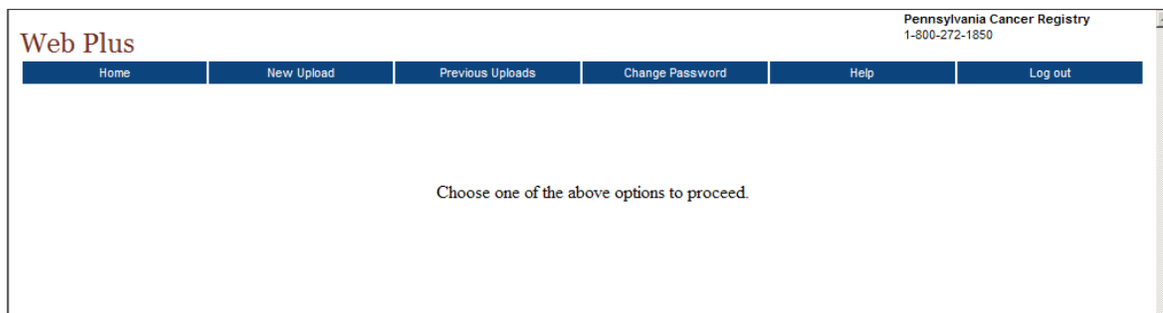
Please select a cancer reporting activity from those listed below the facility for which you would like to report.

Your facility name will display here

[File Upload](#)

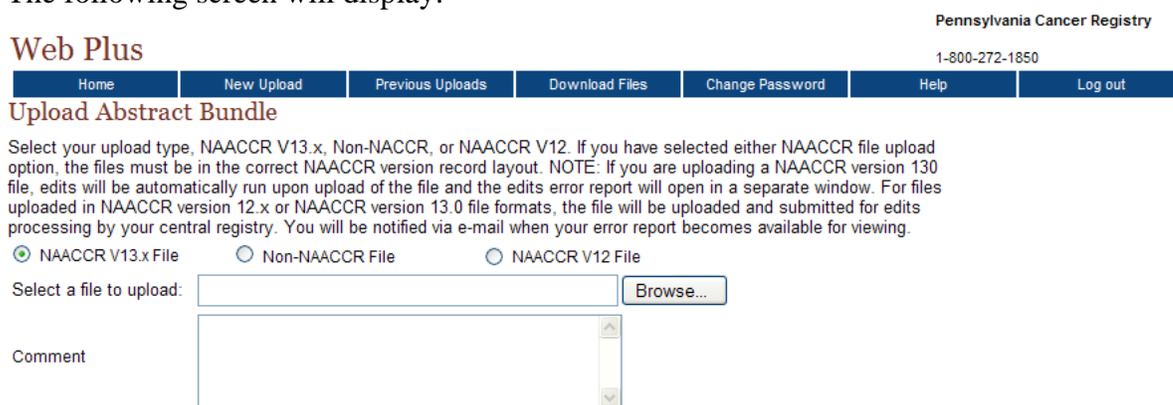
4. Click on File Upload.

The following screen will display:



5. Click on New Upload.

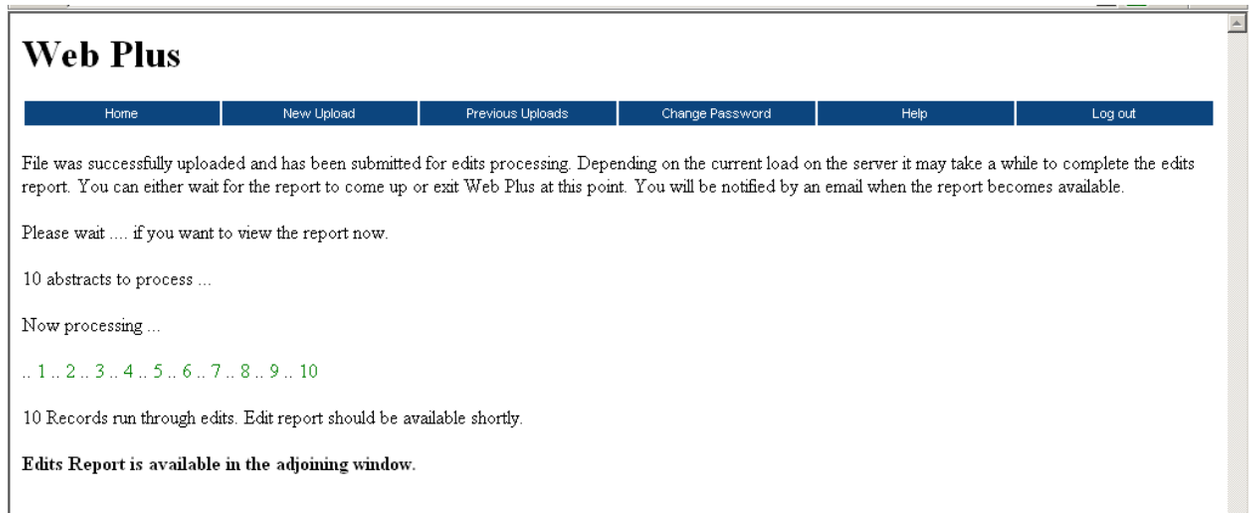
The following screen will display:



Uploading data files:

6. The button beside NAACCR v13.x File is defaulted, but make sure this button is selected prior to uploading the data file. **When uploading the actual data file, the button beside NAACCR file MUST be selected.** Select the data file to upload by clicking on the Browse button and navigating to the location of the file.
7. **In the comment section, you MUST enter the number of abstracts you are exporting.** The comment section can also be used for any special information. For example, if you are uploading a trial shipment, close-out or reconciliation file, this could be indicated here.
8. Click on Upload. The following screen will appear and as the records are processing, the case count will display under ‘Now processing’. Numbers in green indicate the record is error-free and numbers in red indicate the record has errors.

Note about edits: See #6 under General Information.



Web Plus

Home New Upload Previous Uploads Change Password Help Log out

File was successfully uploaded and has been submitted for edits processing. Depending on the current load on the server it may take a while to complete the edits report. You can either wait for the report to come up or exit Web Plus at this point. You will be notified by an email when the report becomes available.

Please wait if you want to view the report now.

10 abstracts to process ...

Now processing ...

.. 1 .. 2 .. 3 .. 4 .. 5 .. 6 .. 7 .. 8 .. 9 .. 10

10 Records run through edits. Edit report should be available shortly.

Edits Report is available in the adjoining window.

An edit report will display in an adjoining window. You can view and/or print the edit report now or close it by selecting File/Close or clicking on the X in the top right corner.

You will also receive the following e-mail when the edit report is available and it can be viewed at any time:

Dear First Name Last Name,

Edit report of the abstracts bundle, C:\yourdatafile.txt submitted on 12/5/2008 10:22:27 AM, is ready. Please log on to Web Plus and select "Previous Uploads" option from the menu. All your previous uploads will be listed on this page. Click on "View Edit Report" link to view the report of this bundle. The report will open in a separate window.

The bundle does not have any edit errors and has been accepted.

Thank you,
Web Plus System Administrator
Pennsylvania Cancer Registry

Uploading PCR Transmittal Form (Only necessary when zero records to report. See #8 under General Information):

1. Click on New Upload.

- Click on the button beside Non-NAACCR File. **When uploading the electronic accession list, the button beside Non-NAACCR file MUST be selected.** Select the PCR Transmittal Form to upload by clicking on the Browse button and navigating to the location of the file.

- Click on Upload. A message will appear at the bottom of the screen stating ‘The file has been uploaded as a Non-NAACCR file’.

You will receive the following e-mail:

Dear First Name Last Name,

Your non-NAACCR file: C:\PCR Transmittal Form.doc was successfully uploaded to Web Plus and received by Pennsylvania Cancer Registry on 12/5/2008 10:40:08 AM.

Web Plus System Administrator
 Pennsylvania Cancer Registry

- Click on Log out to close Web Plus.

View Cases Submitted to the Pennsylvania Cancer Registry via Web Plus

1. Log into Web Plus.
2. The Web Plus home page for your facility opens.



Web Plus Home Page for First Name Last Name

Please select a cancer reporting activity from those listed below the facility for which you would like to report.

Your facility name will display here

[File Upload](#)

3. Click on File Upload.
 - a. The following screen will display:



Choose one of the above options to proceed.

4. Click on Previous Uploads and Click Track File Uploads



Choose one of the above options to proceed.

- A List of previous Abstract Uploads will be displayed. Click the View Abstracts for any selected shipment you would like to review.

Pennsylvania Cancer Registry
 1-800-272-1850

Home
New Upload
Previous Uploads
Download Files
Change Password
Help
Log out

Web Plus

Previous Uploads

Abstract bundles previously uploaded from your facility are listed below. Click on View Edit Report link to view the report on a bundle. You can also view selected fields of the abstracts in a bundle by clicking on View Abstracts link. To view the files uploaded within a data range enter the date range below and click Search.

Date uploaded from: to:

Original File Name	Internal File Name	Date Uploaded	Status	Total Abstracts	Abstracts with Errors	Total Errors	Comment	Action
WEB PLUS TEST	F0028704.bun	1/22/2015 11:18:09 AM	Acceptable Errors	4	4	4	Test, Do Not Download	View Abstracts View Edit Report View Data Quality Report

- The list of Patients submitted for that shipment will be displayed.

Home
New Upload
Previous Uploads
Download Files
Change Password
Help
Log out

Web Plus

View Abstracts in Bundle

Abstract Bundle: WEB PLUS TEST, Bundle ID:F0028704.bun

Total Abstracts: 4

Abstracts with errors: 4

Total errors in bundle: 4

LastName	FirstName	BirthDate	MedRecNum	AccNumHosp	PSite
POTTER	HARRY	99/99/9999	100108988	201000713	C150
GRANGER	HERMIONE	99/99/9999	99832858	201101247	C341
WEASLEY	RONALD	99/99/9999	100566759	201201249	C029
WEASLEY	VIRGINIA	99/99/9999	100437996	201201250	C341

- If you would like to save a copy of this list for your files, this screen can be saved as an HTML document using the *save as* function in your browser. All browsers are different, so if you need assistance, please contact your IT staff.

Note: These lists will remain available until Web Plus is purged. Typically Web Plus is purged in July and anything >18 months old is removed. So in July of 2015 anything from 2013 and earlier will be removed.

**APPENDIX D:
ICD-9-CM and ICD-10-CM CASEFINDING LISTS
FOR REPORTABLE CONDITIONS**

CASEFINDING LIST FOR REPORTABLE CONDITIONS

ICD-9-CM Codes

Use the following ICD-9-CM list to identify potentially reportable. To determine if a diagnosis identified from the ICD-9-CM codes is reportable, refer to the reportability guidelines in Part One.

Some ICD-9-CM codes contain conditions that are not reportable. These records still need to be reviewed and assessed individually to verify whether or not they are reportable to the PCR.

ICD-9-CM Code	Description
140._ – 172._, 174._- 209.36, 209.7_	Malignant Neoplasms (excluding category 173) stated or presumed to be primary (of specified sites) and certain specified histologies
173.00, 173.09	Unspecified/other malignant neoplasm of skin of lip
173.10, 173.19	Unspecified/other malignant neoplasm of skin of eyelid, including canthus
173.20, 173.29	Unspecified/other malignant neoplasm of skin of ear and external auricular canal
173.30, 173.39	Unspecified/other malignant neoplasm of skin of other/unspecified parts of face
173.40, 173.49	Unspecified/other malignant neoplasm of scalp and skin of neck
173.50, 173.59	Unspecified/other malignant neoplasm of skin of trunk, except scrotum
173.60, 173.69	Unspecified/other malignant neoplasm of skin of upper limb, including shoulder
173.70, 173.79	Unspecified/other malignant neoplasm of skin of lower limb, including hip
173.80, 173.89	Unspecified/other malignant neoplasm of skin of other specified sites of skin
173.90, 173.99	Unspecified/other malignant neoplasm of skin, site unspecified
225.0- 225.9	Benign neoplasm of brain and spinal cord
227.3, 227.4	Benign neoplasm of pituitary gland, craniopharyngeal duct (pouch), pineal gland
228.02	Hemangioma of intracranial structures
228.1	Lymphangioma, any site (note: includes only Lymphangioma of the brain, other parts of nervous system and endocrine gland)
230.0 – 234.9	Carcinoma in situ
237.0 – 237.1	Neoplasm of uncertain behavior of endocrine glands and nervous system: pituitary gland, craniopharyngeal duct and pineal gland
237.5, 237.6, 237.9	Neoplasm of uncertain behavior of endocrine glands and nervous system: brain, spinal cord, meninges, endocrine glands and other and unspecified parts of nervous system.
238.4	Polycythemia Vera
238.7_	Other lymphatic and hematopoietic tissues

ICD-9-CM Code	Description
239.6, 239.7	Neoplasms of unspecified nature, brain, endocrine glands and other parts of nervous system
273.3	Macroglobulinemia (Waldenstrom's macroglobulinemia)
277.89	Other specified disorders of metabolism (reportable includes terms: Hand-Schuller Christian disease; histiocytosis (acute) (chronic); histiocytosis X (chronic))
V58.0	Admission for radiotherapy
V58.11	Admission for chemotherapy
V58.12	Encounter for immunotherapy for neoplastic condition.
V67.1	Radiation therapy follow-up
V67.2	Chemotherapy follow-up

ICD-10-CM Codes

The following ICD-10-CM list will be used to identify potentially reportable conditions once ICD-10-CM is implemented. To determine if a diagnosis identified from the ICD-10-CM codes is reportable, refer to the reportability guidelines in Part One.

Some ICD-10-CM codes contain conditions that are not reportable. These records still need to be reviewed and assessed individually to verify whether or not they are reportable to the PCR.

ICD-10-CM Code	Description
C00._ – C96._	Malignant neoplasms (excluding category C44), stated or presumed to be primary (of specified site) and certain specified histologies
C44.00, C44.09	Unspecified/other malignant neoplasm of skin of lip
C44.10_, C44.19_	Unspecified/other malignant neoplasm of skin of eyelid
C44.20_, C44.29_	Unspecified/other malignant neoplasm skin of ear and external auricular canal
C44.30_, C44.39_	Unspecified/other malignant neoplasm of skin of other/unspecified parts of face
C44.40, C44.49	Unspecified/other malignant neoplasm of skin of scalp & neck
C44.50_, C44.59_	Unspecified/other malignant neoplasm of skin of trunk
C44.60_, C44.69_	Unspecified/other malignant neoplasm of skin of upper limb, incl. shoulder
C44.70_, C44.79_	Unspecified/other malignant neoplasm of skin of lower limb, including hip
C44.80, C44.89	Unspecified/other malignant neoplasm of skin of overlapping sites of skin
C44.90, C44.99	Unspecified/other malignant neoplasm of skin of unspecified sites of skin
D00._ – D09._	In-situ neoplasms (<i>Note: Carcinoma in situ of the cervix (CIN III-8077/2) and Prostatic Intraepithelial Carcinoma (PIN III-8148/2) are not reportable.</i>)
D18.02	Hemangioma of intracranial structures and any site
D18.1	Lymphangioma, any site (<i>Note: Includes Lymphangiomas of Brain, Other parts of nervous system and endocrine glands, which are reportable.</i>)
D32._	Benign neoplasm of meninges (cerebral, spinal and unspecified)
D33._	Benign neoplasm of brain and other parts of central nervous system
D35.2 - D35.4	Benign neoplasm of pituitary gland, craniopharyngeal duct and pineal gland
D42._, D43._	Neoplasm of uncertain or unknown behavior of meninges, brain, CNS
D44.3 – D44.5	Neoplasm of uncertain or unknown behavior of pituitary gland, craniopharyngeal duct and pineal gland
D45	Polycythemia vera (9950/3)
D46._	Myelodysplastic syndromes (9980, 9982, 9983, 9985, 9986, 9989, 9991, 9992)

ICD-10-CM Code	Description
D47.1	Chronic myeloproliferative disease (9960/3, 9963/3)
D47.3	Essential (hemorrhagic) thrombocythemia (9962/3)
D47.4	Osteomyelofibrosis (9961/3)
D47.7	Other specified neoplasms of uncertain/unknown behavior of lymphoid, hematopoietic (9965/3, 9966/3, 9967/3, 9971/3, 9975/3, 9987/3)
D47.Z_	Other neoplasms of uncertain behavior of lymphoid, hematopoietic related tissue
D47.9	Neoplasm of uncertain behavior of lymphoid, hematopoietic and related tissue, unspecified (9960/3, 9970/1, 9931/3)
D49.6, D49.7	Neoplasm of unspecified behavior of brain, endocrine glands and other CNS
Z51.0	Encounter for antineoplastic radiation therapy
Z51.1	Encounter for antineoplastic chemotherapy and immunotherapy

Electronic Look up Lists

For an electronic list of each individual code, go to the PCR Website.

<http://www.health.pa.gov/MyRecords/Registries/Cancer/Pages/Hospital%20Reporting.aspx#.VQGqmnfD8qI>

Supplemental Casefinding List

Effective January 1, 2014, the PCR will no longer maintain a supplemental casefinding list. If time and resources permit, hospital registries may use the SEER supplemental list. The list is located on the SEER website:

<http://seer.cancer.gov/tools/casefinding/>

APPENDIX E:
**Errata to *International Classification of Diseases for
Oncology, Third Edition (ICD-O-3)*.**

Update ICD-O-3

All errata documented in this appendix must be added to all *International Classification of Diseases for Oncology, Third Edition (ICD-O-3)* coding books being used to ensure correct coding of primary site and histology. See also *Part Three, Primary Site* and *Part Three, Histology* for additional coding instructions.

Errata Effective January 1, 2016

These terms had not been included in any ICD-O-3 errata and therefore were not addressed throughout the cancer surveillance community. NPCR has reviewed the terms and determined that the following are reportable. Therefore, they are now reportable to the PCR. While there has not been an official errata to address these histology terms, the PCR recommends adding them your ICD-O-3 Manuals.

1. Non-invasive mucinous cystic neoplasm of the pancreas with high-grade dysplasia replaces mucinous cystadenocarcinoma, non-invasive (8470/2).
2. Solid pseudopapillary neoplasm of pancreas (8452/3) is synonymous with solid pseudopapillary carcinoma (C25._)
3. Based on pathologist consultation, metastases have been reported in some cystic pancreatic endocrine neoplasm (CPEN) cases. With all other pancreatic endocrine tumors now considered malignant, CPEN will also be considered malignant, until proven otherwise. Most CPEN cases are non-functioning and are REPORTABLE using histology code 8150/3, unless the tumor is specified as a neuroendocrine tumor, grade 1 (assign code 8240/3) or neuroendocrine tumor, grade 2 (assign code 8249/3)
4. Laryngeal intraepithelial neoplasia, grade III (LINIII) (8077/2), C320-C329)
5. Squamous intraepithelial neoplasia, grade III (SINIII) (8077/2), except Cervix and Skin
6. Mature teratoma of the testes in adults is malignant and REPORTABLE as 9080/3, but continues to be non-reportable in prepubescent children (9080/0). The following provides additional guidance:
 - Adult is defined as post puberty
 - Pubescence can take place over a number of years
 - Do not rely solely on age to indicate pre or post puberty status. Review all information (physical history, etc.) for documentation of pubertal status. When testicular teratomas occur in adult males, pubescent status is likely to be stated in the medical record because it is an important factor of the diagnosis.
 - Do not report if unknown whether patient is pre or post pubescence. When testicular teratoma occurs in a male and there is no mention of pubescence, it is likely that the patient is a child, or pre-pubescent, and the tumor is benign.

Errata Effective January 1, 2015

1. **Behavior code/Reportability changes-** Effective with cases diagnosed on or after 1/1/2015 Carcinoids, NOS of the appendix are now reportable and should be coded to 8240/3. Document this change in your ICD-O-3.
2. **Enteroglucagonoma, malignant-** Effective with cases diagnosed on or after 1/1/2015, Enteroglucagonoma, malignant is now coded to 8152/3. Instead of 8157/3. Document this change in your ICD-O-3.
3. **New codes and terms-** Many of the new codes slated to be effective in 2015 cannot be used because they are not among the acceptable histologies for the Collaborative Stage algorithms. The below table lists the new terms and what code to use for 2015. New instructions will be provided in 2016 for coding these terms, when Collaborative Stage is discontinued.

New Term	Code to use for 2015*
Pancreatobiliary-type carcinoma (C24.1) Adenocarcinoma, pancreatobiliary- type (C24.1)	8255/3
Serrated adenocarcinoma	8213/3
Micropapillary carcinoma, NOS (C18._, C19.9, C20.9)	8507/3
Mixed acinar ductal carcinoma	8523/3
Papillary tumor of the pineal region	9361/3
Pilomyxoid astrocytoma	9421/3
Angiocentric glioma	9380/1
Pituicytoma	9380/1
Papillary glioneuronal tumor	9505/1
Rosette-forming glioneuronal tumor	9505/1

*ICD-O-3 rule F applies (code the behavior stated by the pathologist).

Errata Effective January 1, 2014

Use the following new terms, synonyms, and related terms for existing ICD-O-3 codes for all cases diagnosed on or after January 1, 2014. Add these new terms/synonyms into your ICD-O-3 Coding Manual including the effective date.

Note: This listing does not change reportability requirements. Reportable histologies are still based on the following criteria:

- ICD-O Behavior /2 or /3- All histologic diagnoses with a behavior code of /2 (in situ) or /3 (malignant) in the *ICD, Second Edition* (ICD-O-2) or *Third Edition* (ICD-O-3) are reportable.
- ICD-O Behavior /0 or /1 – All histologic diagnoses with a behavior code of /0 (benign) or /1 (borderline or uncertain) when **primary to the intracranial and central nervous system (CNS) sites listed below** are reportable when diagnosed on or after **January 1, 2004**.

New preferred term	8150/0 Pancreatic endocrine tumor, benign (C25._)
Move former preferred term to synonym	8150/0 Islet cell adenoma (C25._)
New related term	8150/0 Pancreatic microadenoma (C25._)
New preferred term	8150/1 Pancreatic endocrine tumor, NOS (C25._)
Move former preferred term to synonym	8150/1 Islet cell tumor, NOS (C25._)
New preferred term	8150/3 Pancreatic endocrine tumor, malignant (C25._)
Move former preferred term to synonym	8150/3 Islet cell carcinoma (C25._)
New related term	8150/3 Pancreatic endocrine tumor, nonfunctioning (C25._)
New related term	8152/1 L-cell tumor
New related term	8152/1 Glucagon-like peptide-producing tumor (C25._)
New related term	8152/1 Pancreatic peptide and pancreatic peptide-like peptide within terminal tyrosine amide producing tumor
New synonym for related term	8152/1 PP/PYY producing tumor
New preferred term	8154/3 Mixed pancreatic endocrine and exocrine tumor, malignant (C25._)
New related term	8154/3 Mixed endocrine and exocrine adenocarcinoma (C25._)
New synonym for related term	8154/3 Mixed islet cell and exocrine adenocarcinoma (C25._)
New related term	8154/3 Mixed acinar-endocrine-ductal carcinoma
New related term	8201/3 Cribriform comedo-type carcinoma (C18._, C19.9, C20.9)
New synonym	8201/3 Adenocarcinoma, cribriform comedo-type (C18._, C19.9, C20.9)
New synonym to primary term	8213/0 Traditional serrated adenoma
New related term	8213/0 Sessile serrated adenoma
New related term	8213/0 Sessile serrated polyp
New related term	8213/0 Traditional sessile serrated adenoma

New related term	8240/3 Neuroendocrine tumor, grade 1
New related term	8240/3 Neuroendocrine carcinoma, low grade
New related term	8240/3 Neuroendocrine carcinoma, well-differentiated
New preferred term	8244/3 Mixed adenoneuroendocrine carcinoma
Move former preferred term to synonym	8244 Composite carcinoid
New synonym	8244/3 Combined/mixed carcinoid and adenocarcinoma
New synonym	8244/3 MANEC
New synonym	8249/3 Neuroendocrine tumor, grade 2
New related term	8249/3 Neuroendocrine carcinoma, moderately differentiated
New synonym	8263/0 Tubulo-papillary adenoma
New related term	8290/0 Spindle cell oncocytoma (C75.1)
New related term	8490/3 Poorly cohesive carcinoma
New related term	8811/0 Plexiform fibromyxoma
New related term	8970/3 Hepatoblastoma, epithelioid (C22.0)
New related term	8970/3 Hepatoblastoma, mixed epithelial-mesenchymal (C22.0)
New related term	9471/3 Medulloblastoma with extensive nodularity
New related term	9474/3 Anaplastic medulloblastoma
New related term	9506/1 Extraventricular neurocytoma

Errata Effective January 1, 2001

The ICD-O-3 Errata and Clarifications on the following pages were released by SEER in May 2003 and May 2001 to provide corrections to ICD-O-3 and further discussion of issues that were not well described in the printed editions of ICD-O-3.

**APPENDIX F:
REQUIRED DATA SET
FOR REPORTING FACILITIES**

Required Data Set for Reporting Facilities (Effective 1/1/2016)

*If the M Record column is checked, a Modification Record should be generated if the values in these fields change. If the column is not checked, DO NOT generate a Modification Record.

PCR Required Data Item	Field length	NAACCR Item #	M Record*
Name--Last	40	2230	✓
Name--Suffix	3	2270	
Name--First	40	2240	✓
Name--Middle	40	2250	✓
Name--Maiden	40	2390	✓
Name--Alias	40	2280	✓
Addr at DX--No & Street	60	2330	✓
Addr at DX--Supplementl	60	2335	✓
Addr at DX--City	50	70	✓
Addr at DX--State	2	80	✓
Addr at DX--Postal Code	9	100	✓
County at DX	3	90	
Addr Current--No & Street*	60	2350	✓
Addr Current—Supplementl*	60	2355	✓
Addr Current—City*	50	1810	✓
Addr Current—State*	2	1820	✓
Addr Current--Postal Code*	9	1830	✓
Age at Diagnosis	3	230	✓
Date of Birth	8	240	✓
Date of Birth Flag	2	241	✓
Birthplace-State	2	252	✓
Birthplace-Country	3	254	✓
Social Security Number	9	2320	✓
Sex	1	220	✓
Spanish/Hispanic Origin	1	190	✓
Race 1	2	160	✓
Race 2	2	161	✓
Race 3	2	162	✓
Race 4	2	163	✓
Race 5	2	164	✓
Primary Payer at DX	2	630	✓
Text--Usual Occupation	100	310	✓
Text--Usual Industry	100	320	✓
Medical Record Number	11	2300	
Sequence Number--Hospital	2	560	✓
Class of Case	2	610	✓
Type of Reporting Source	1	500	✓
Date of 1st Contact	8	580	✓
Date of 1st Contact Flag	2	581	✓

Required Data Set for Reporting Facilities (Effective 1/1/2016)

*If the M Record column is checked, a Modification Record should be generated if the values in these fields change. If the column is not checked, DO NOT generate a Modification Record.

PCR Required Data Item	Field length	NAACCR Item #	M Record*
Date of Inpatient Adm	8	590	✓
Date of Inpt Adm Flag	2	591	✓
Date of Inpatient Disch	8	600	✓
Date of Inpt Disch Flag	2	601	✓
Institution Referred From	10	2410	✓
Physician—Follow-up	8	2470	✓
Date of Diagnosis	8	390	✓
Date of Diagnosis Flag	2	391	✓
Primary Site	4	400	✓
Laterality	1	410	✓
Histology (92-00) ICD-O-2	4	420	✓
Behavior (92-00) ICD-O-2	1	430	✓
Histologic Type ICD-O-3	4	522	✓
Behavior Code ICD-O-3	1	523	✓
Grade	1	440	✓
ICD-O-3 Conversion Flag	1	2116	✓
Diagnostic Confirmation	1	490	✓
Tumor Size Summary	3	756	✓
Regional Nodes Examined	2	830	✓
Regional Nodes Positive	2	820	✓
Lymph-vascular Invasion	1	1182	✓
Mets at DX-Bone	1	1112	✓
Mets at DX-Brain	1	1113	✓
Mets at DX-Distant LN	1	1114	✓
Mets at DX-Liver	1	1115	✓
Mets at DX-Lung	1	1116	✓
Mets at DX-Other	1	1117	✓
CS Site-Specific Factor 1	3	2880	✓
CS Site-Specific Factor 2	3	2890	✓
CS Site-Specific Factor 3	3	2900	✓
CS Site-Specific Factor 4	3	2910	✓
CS Site-Specific Factor 5	3	2920	✓
CS Site-Specific Factor 6	3	2930	✓
CS Site-Specific Factor 7	3	2861	✓
CS Site-Specific Factor 8	3	2862	✓
CS Site-Specific Factor 9	3	2863	✓
CS Site-Specific Factor 10	3	2864	✓
CS Site-Specific Factor 11	3	2865	✓
CS Site-Specific Factor 12	3	2866	✓

Required Data Set for Reporting Facilities (Effective 1/1/2016)

*If the M Record column is checked, a Modification Record should be generated if the values in these fields change. If the column is not checked, DO NOT generate a Modification Record.

PCR Required Data Item	Field length	NAACCR Item #	M Record*
CS Site-Specific Factor 13	3	2867	✓
CS Site-Specific Factor 14	3	2868	✓
CS Site-Specific Factor 15	3	2869	✓
CS Site-Specific Factor 16	3	2870	✓
CS Site-Specific Factor 17	3	2871	✓
CS Site-Specific Factor 18	3	2872	✓
CS Site-Specific Factor 19	3	2873	✓
CS Site-Specific Factor 20	3	2874	✓
CS Site-Specific Factor 21	3	2875	✓
CS Site-Specific Factor 22	3	2876	✓
CS Site-Specific Factor 23	3	2877	✓
CS Site-Specific Factor 24	3	2878	✓
CS Site-Specific Factor 25	3	2879	✓
SEER Summary Stage 1977	1	760	✓
SEER Summary Stage 2000	1	759	✓
TNM Path T	4	880	✓
TNM Path N	4	890	✓
TNM Path M	4	900	✓
TNM Path Stage Group	4	910	✓
TNM Path Descriptor	1	920	✓
TNM Clin T	4	940	✓
TNM Clin N	4	950	✓
TNM Clin M	4	960	✓
TNM Clin Stage Group	4	970	✓
TNM Clin Descriptor	1	980	✓
TNM Edition Number	2	1060	✓
RX Summ--Surg Prim Site	2	1290	✓
RX Date--Surgery	8	1200	✓
RX Date--Surgery Flag	2	1201	✓
RX Date—Mst Defn Srg	8	3170	✓
RX Date—Mst Defn Srg	2	3171	✓
RX Summ--Scope Reg LN Sur	1	1292	✓
RX Summ--Surg Oth Reg/Dis	1	1294	✓
Reason for No Surgery	1	1340	✓
Rad--Regional RX Modality	2	1570	✓
RX Date--Radiation	8	1210	✓
RX Date--Radiation Flag	2	1211	✓
RX Summ-Surg/Rad Seq	1	1380	✓
Reason For No Radiation	1	1430	✓

Required Data Set for Reporting Facilities (Effective 1/1/2016)

*If the M Record column is checked, a Modification Record should be generated if the values in these fields change. If the column is not checked, DO NOT generate a Modification Record.

PCR Required Data Item	Field length	NAACCR Item #	M Record*
RX Date--Chemo	8	1220	✓
RX Date--Chemo Flag	2	1221	✓
RX Summ--Chemo	2	1390	✓
RX Date--Hormone	8	1230	✓
RX Date--Hormone Flag	2	1231	✓
RX Summ--Hormone	2	1400	✓
RX Date--BRM	8	1240	✓
RX Date--BRM Flag	2	1241	✓
RX Summ--BRM	2	1410	✓
RX Summ--Systemic Sur Seq	1	1639	✓
RX Summ--Transplnt/Endocr	2	3250	✓
RX Summ--Other	1	1420	✓
RX Date--Other	8	1250	✓
RX Date--Other Flag	2	1251	✓
Date of 1st Crs RX--COC	8	1270	✓
Date of 1st Crs RX Flag	2	1271	✓
RX Summ--Treatment Status	1	1285	✓
Date of Last Contact	8	1750	
Date of Last Contact Flag	2	1751	
Vital Status	1	1760	
Reporting Facility	10	540	
Abstracted By	3	570	
Text--DX Proc-PE	1000	2520	
Text--DX Proc-X-ray/Scan	1000	2530	
Text--DX Proc-Scopes	1000	2540	
Text--DX Proc-Lab Tests	1000	2550	
Text--DX Proc-Op	1000	2560	
Text--DX Proc-Path	1000	2570	
Text--Primary Site Title	100	2580	
Text--Histology Title	100	2590	
Text--Staging	1000	2600	
RX Text--Surgery	1000	2610	
RX Text--Radiation (Beam)	1000	2620	
RX Text--Radiation (Other)	1000	2630	
RX Text--Chemo	1000	2640	
RX Text--Hormone	1000	2650	
RX Text--BRM	1000	2660	
RX Text--Other	1000	2670	
Text--Remarks	1000	2680	

Required Data Set for Reporting Facilities (Effective 1/1/2016)

*If the M Record column is checked, a Modification Record should be generated if the values in these fields change. If the column is not checked, DO NOT generate a Modification Record.

PCR Required Data Item	Field length	NAACCR Item #	M Record*
Text--Place of Diagnosis	60	2690	

System Codes

PCR Required Data Item	Field length	NAACCR Item #	M Record*
Record Type	1	10	
Registry Type	1	30	
NAACCR Record Version	3	50	
Site Coding Sys--Current	1	450	
Morph Coding Sys--Current	1	470	
RX Coding System--Current	2	1460	
ICD Revision Number	1	1920	
Tumor Record Number	2	60	
Patient System ID-Hosp	8	21	
Date Case Completed	8	2090	
Date Case Last Changed	8	2100	
Date Case Report Exported	8	2110	
Vendor Name	10	2170	
CS Version Input Original	6	2935	
CS Version Derived	6	2936	
CS Version Input Current	6	2937	

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**APPENDIX G:
PCR Transmittal Form**



Date Received:

**PENNSYLVANIA CANCER REGISTRY
TRANSMITTAL FORM**

FACILITY NAME: _____

PCR 4-DIGIT IDENTIFICATION NUMBER: _____

DATE SUBMITTED: _____

NUMBER OF CHANGE RECORDS: _____

NUMBER OF NEW RECORDS: _____

Comments:

For Pennsylvania Cancer Registry Use Only

	Date	Initials
Web+:		
Log:		
Area:		

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**APPENDIX H:
FEDERAL INFORMATION PROCESSING
STANDARDS (FIPS) COUNTY CODES
FOR PENNSYLVANIA**

Federal Information Processing Standards Publication, Counties and Equivalent Entities of the United States, Its Possessions, and Associated Areas. U.S. Department of Commerce, National Institute of Standards and Technology, Gaithersburg, MD, August 31, 1990, pp. 21-22

AND

**SEER GEOCODES
FOR COUNTRY AT DIAGNOSIS**

Note: All codes are used in County at DX field

FIPS COUNTY CODES FOR PENNSYLVANIA

001	Adams	093	Montour
003	Allegheny	095	Northampton
005	Armstrong	097	Northumberland
007	Beaver	099	Perry
009	Bedford	101	Philadelphia
011	Berks	103	Pike
013	Blair	105	Potter
015	Bradford	107	Schuylkill
017	Bucks	109	Snyder
019	Butler	111	Somerset
021	Cambria	113	Sullivan
023	Cameron	115	Susquehanna
025	Carbon	117	Tioga
027	Centre	119	Union
029	Chester	121	Venango
031	Clarion	123	Warren
033	Clearfield	125	Washington
035	Clinton	127	Wayne
037	Columbia	129	Westmoreland
039	Crawford	131	Wyoming
041	Cumberland	133	York
043	Dauphin		
045	Delaware		
047	Elk		
049	Erie		
051	Fayette		
053	Forest		
055	Franklin		
057	Fulton		
059	Greene		
061	Huntington		
063	Indiana		
065	Jefferson		
067	Juniata		
069	Lackawanna		
071	Lancaster		
073	Lawrence		
075	Lebanon		
077	Lehigh		
079	Luzerne		
081	Lycoming		
083	McKean		
085	Mercer		
087	Mifflin		
089	Monroe		
091	Montgomery		

SEER GEOCODES

A		680	Asia, East
585	Abyssinia	640	Asia, Mid-East
629	Aden	610	Asia Minor, NOS
583	Afars and Issas	610	Asia, Near-East
638	Afghanistan	650	Asia, Southeast
500	Africa	620	Asian Arab countries
570	Africa, East	634	Asian Republics of the former U.S.S.R.
510	Africa, North	109	Atlantic/Caribbean area, other U.S. possessions
540	Africa, South	100	Atlantic/Caribbean area, U.S. possessions
545	Africa, South West	711	Australia
530	Africa, West	711	Australian New Guinea
580	African Coastal Islands (previously included in 540)	436	Austria
037	Alabama	633	Azerbaijan
091	Alaska	633	Azerbaijan S.S.R.
481	Albania	445	Azores
224	Alberta		
513	Algeria	B	
250	America, Central	247	Bahamas
260	America, North (use more specific term if possible)	629	Bahrain
300	America, South	443	Balearic Islands
121	American Samoa	463	Baltic Republic, NOS
611	Anatolia	463	Baltic States, NOS
641	Andaman Islands	645	Bangladesh
443	Andorra	245	Barbados
543	Angola	245	Barbuda
245	Anguilla	545	Basutoland
665	Annam	431	Bavaria
750	Antartica	545	Bechuanaland
245	Antigua	457	Belarus
245	Antilles, NOS	541	Belgian Congo
245	Antilles, Netherlands	433	Belgium
625	Arab Palestine	252	Belize
629	Arabia, Saudi	539	Benin
629	Arabian Peninsula	246	Bermuda
365	Argentina	456	Bessarabia
087	Arizona	643	Bhutan
071	Arkansas	539	Bioko (Fernando Poo)
633	Armenia (U.S.S.R.)	452	Bohemia
611	Armenia (Turkey)	355	Bolivia
245	Aruba	545	Bophuthatswana
600	Asia, NOS		

SEER GEOCODES

673	Borneo	361	Chile
453	Bosnia-Herzegovina	681	China, NOS
545	Botswana	665	China, Cochin
341	Brazil	682	China, People's Republic of
226	British Columbia	684	China, Republic of
331	British Guiana	723	Christmas Island
252	British Honduras	545	Ciskel
245	British Virgin Islands	665	Cochin China
245	British West Indies, NOS	711	Cocos (Keeling) Islands
671	Brunei	311	Columbia
454	Bulgaria	083	Colorado
520	Burkina Faso (Upper Volta)	580	Comoros
649	Burma (see Myanmar)	226	Columbia, British
579	Burundi	022	Columbia, District of
457	Byelorussian S.S.R.	539	Congo-Brazzaville
	C	541	Congo-Leopoldville
		541	Congo, Belgian
		539	Congo, French
543	Cabinda	541	Congo Kinshasa
245	Caicos Islands	007	Connecticut
097	California	124	Cook Islands
663	Cambodia	441	Corsica
539	Cameroon	256	Costa Rica
220	Canada	539	Cote d'Ivoire (Ivory Coast)
110	Canal Zone	471	Crete
443	Canary Islands	453	Croatia
122	Canton Islands	241	Cuba
545	Cape Colony	245	Curacao
445	Cape Verde Islands	495	Cyprus
245	Caribbean, NOS	517	Cryonic
245	Caribbean Islands, other	452	Czechoslovakia
123	Caroline Islands	452	Czech Republic
711	Cartier Islands		D
633	Caucasian Republics of the former U.S.S.R.	539	Dahomey
245	Cayman Islands	453	Dalmatia
500	Central Africa, NOS	017	Delaware
539	Central African Republic	425	Denmark
250	Central America	022	District of Columbia
499	Central Europe, NOS	583	Djibouti
060	Central Midwest States	449	Dobruja
647	Ceylon	245	Dominica
520	Chad	243	Dominican Republic
401	Channel Islands (British)	673	Dutch East Indies

SEER GEOCODES

332 Dutch Guiana

530 French West Africa, NOS

245 French West Indies

E

570 East Africa

680 East Asia

431 East Germany

673 East Indies, Dutch

645 East Pakistan

499 Eastern Europe, NOS

345 Ecuador

519 Egypt

410 Eire

254 El Salvador

125 Ellice Islands

122 Enderbury Islands

401 England

500 Equatorial Africa, NOS

539 Equatorial Guinea (Spanish
Guinea)

585 Eritrea

458 Estonia

458 Estonian S.S.R. (Estonia)

585 Ethiopia

499 Europe, NOS

470 Europe, other mainland

F

420 Faroe (Faeroe) Islands

381 Falkland Islands

431 Federal Republic of Germany

539 Fernando Poo

721 Fiji

429 Finland

035 Florida

684 Formosa

721 Fortuna

441 France

545 Free State (Orange Free State)

539 French Congo

333 French Guiana

725 French Polynesia

583 French Somaliland

G

539 Gabon

345 Galapagos Islands

539 Gambia

631 Gaza Strip

033 Georgia (U.S.A.)

633 Georgia (U.S.S.R.)

430 Germanic countries

431 German Democratic Republic

431 Germany

431 Germany, East

431 Germany, Federal Republic of

431 Germany, West

539 Ghana

485 Gibraltar

122 Gilbert Islands

471 Greece

210 Greenland

245 Grenada

245 Grenadines, The

245 Guadeloupe

126 Guam

251 Guatemala

401 Guernsey

331 Guiana, British

332 Guiana, Dutch

333 Guiana, French

539 Guinea

539 Guinea-Bissau (Portuguese
Guinea)

539 Guinea, Equatorial

--- Guinea, New (See New Guinea)

539 Guinea, Portuguese

331 Guyana

H

242 Haiti

099 Hawaii

432 Holland

SEER GEOCODES

253	Honduras	663	Kampuchea
252	Honduras, British	065	Kansas
683	Hong Kong	634	Kazakh S.S.R.
475	Hungary	634	Kazakhstan
	I	047	Kentucky
		575	Kenya
421	Iceland	634	Kirghiz S.S.R.
081	Idaho	122	Kiribati
061	Illinois	695	Korea
641	India	695	Korea, North
045	Indiana	695	Korea, South
673	Indies, Dutch East	629	Kuwait
660	Indochina	634	Kyrgystan
673	Indonesia	634	Kyrgyz
053	Iowa		L
637	Iran		
627	Iraq	221	Labrador
620	Iraq-Saudi Arabian Neutral Zone	661	Laos
410	Ireland (Eire)	420	Lapland, NOS
404	Ireland, Northern	265	Latin America, NOS
410	Ireland, NOS	459	Latvia
410	Ireland, Republic of	459	Latvian S.S.R. (Latvia)
401	Isle of Man	623	Lebanon
631	Israel	245	Leeward Island, NOS
583	Issas	545	Lesotho
447	Italy	539	Liberia
539	Ivory Coast	517	Libya
	J	437	Liechtenstein
		122	Line Islands, Southern
244	Jamaica	461	Lithuania
423	Jan Mayen	461	Lithuanian S.S.R. (Lithuania)
693	Japan	073	Louisiana
673	Java	434	Luxembourg
401	Jersey		M
631	Jewish Palestine		
127	Johnston Atoll	686	Macao
625	Jordan	686	Macau
453	Jugoslavia	453	Macedonia
	K	555	Madagascar
		445	Madeira Islands
539	Kameroon	002	Maine
		555	Malagasy Republic

SEER GEOCODES

551	Malawi	553	Mozambique
671	Malay Peninsula	629	Muscat
671	Malaysia	649	Myanmar (see Burma)
640	Maldives		
520	Mali		N
491	Malta		
224	Manitoba	545	Namibia
129	Mariana Islands	133	Nampo-shoto, Southern
221	Maritime Provinces, Canada	545	Natal
131	Marshall Islands	723	Nauru
245	Martinique	610	Near-East Asia
021	Maryland	067	Nebraska
005	Massachusetts	643	Nepal
520	Mauritania	432	Netherlands
580	Mauritius	245	Netherlands Antilles
580	Mayotte	332	Netherlands Guiana
490	Mediterranean Islands, other	085	Nevada
721	Melanesian Islands	245	Nevis
610	Mesopotamia, NOS	221	New Brunswick
230	Mexico	724	New Caledonia
041	Michigan	001	New England
123	Micronesia Islands [Federated States of] (Caroline Islands, Trust Territory of Pacific Islands)	673	New Guinea, except Australian and North East
723	Micronesia Islands (except possessions of the U.S.A.)	711	New Guinea, Australian
640	Mid-East Asia	711	New Guinea, North East
132	Midway Islands	003	New Hampshire
052	Minnesota	721	New Hebrides
249	Miquelon	008	New Jersey
039	Mississippi	086	New Mexico
063	Missouri	011	New York
456	Moldavia	715	New Zealand
456	Moldavian S.S.R.	221	Newfoundland
456	Moldova	255	Nicaragua
441	Monaco	520	Niger
691	Mongolia	531	Nigeria
056	Montana	715	Niue
453	Montenegro	711	Norfolk Island
245	Montserrat	510	North Africa, NOS
452	Moravia	260	North America, NOS (use more specific term if possible)
511	Morocco	240	North American Islands
080	Mountain States	671	North Borneo (Malaysia)
		025	North Carolina
		040	North Central States

SEER GEOCODES

054	North Dakota	257	Panama
711	North East New Guinea	711	Papua New Guinea
695	North Korea	371	Paraguay
010	North Mid-Atlantic States	014	Pennsylvania
499	Northern Europe, NOS	629	People's Democratic Republic of Yemen
404	Northern Ireland	682	People's Republic of China
129	Northern Mariana Islands	637	Persia
050	Northern Midwest States	629	Persian Gulf States, NOS
549	Northern Rhodesia	351	Peru
225	Northwest Territories (Canada)	675	Philippine Islands
423	Norway	675	Philippines
998	Not United States, NOS	725	Pitcairn
221	Nova Scotia	451	Poland
227	Nunavut	725	Polynesian Islands
551	Nyasaland	445	Portugal
	O	539	Portuguese Guinea
043	Ohio	224	Prairie Provinces, Canada
075	Oklahoma	221	Prince Edward Island
629	Oman	543	Principe
223	Ontario	101	Puerto Rico
545	Orange Free State		Q
095	Oregon		
403	Orkney	629	Qatar
	P	222	Quebec
			R
120	Pacific area, U.S. possessions	684	Republic of China
090	Pacific Coast States	545	Republic of South Africa
720	Pacific Islands	580	Reunion
123	Pacific Islands, Trust Territory of the (code to specific islands if possible)	006	Rhode Island
639	Pakistan	547	Rhodesia
645	Pakistan, East	549	Rhodesia, Northern
639	Pakistan, West	547	Rhodesia, Southern
139	Palau (Trust Territory of the Pacific Islands)	539	Rio Muni
625	Palestine, Arab	440	Romance-language countries
631	Palestine, Jewish	449	Romania
631	Palestine, NOS	449	Roumania
631	Palestinian National Authority-- PNA	577	Ruanda
		449	Rumania
		455	Russia, NOS
		455	Russia S.F.S.R.

SEER GEOCODES

457	Russian, White	581	Somaliland
455	Russian Federation (former U.S.S.R.)	583	Somaliland, French
577	Rwanda	540	South Africa
134	Ryukyu Islands	545	South Africa, Republic of
		545	South Africa, Union of
		300	South America
		380	South American Islands
		026	South Carolina
		055	South Dakota
		695	South Korea
		020	South Mid-Atlantic States
		545	South West Africa
		650	Southeast Asia
		030	Southeastern States
		499	Southern Europe, NOS
		122	Southern Line Islands
		070	Southern Midwest States
		133	Southern Nampo-shoto
		547	Southern Rhodesia
		629	Southern Yemen
		---	Soviet Union (see individual republics)
		443	Spain
		520	Spanish Sahara
		647	Sri Lanka
		520	Sudan (Anglo-Egyptian Sudan)
		520	Sudanese countries
		673	Sumatra
		332	Suriname
		423	Svalbard
		135	Swan Islands
		545	Swaziland
		427	Sweden
		435	Switzerland
		621	Syria
			T
		634	Tadzhik S.S.R.
		684	Taiwan
		634	Tajikistan
		571	Tanzania
		571	Tanganyika
		571	Tanzanyika
520	Sahara, Western		
121	Samoa, American		
725	Samoa, Western		
245	St. Christopher-Nevis		
580	St. Helena		
245	St. Kitts (see St. Christopher-Nevis)		
245	St. Lucia		
249	St. Pierre		
245	St. Vincent		
447	San Marino		
543	Sao Tome		
447	Sardinia		
224	Saskatchewan		
629	Saudi Arabia		
420	Scandinavia		
403	Scotland		
539	Senegal		
453	Serbia		
580	Seychelles		
403	Shetland Islands		
651	Siam		
447	Sicily		
539	Sierra Leone		
643	Sikkim		
671	Singapore		
450	Slavic countries		
453	Slavonia		
452	Slovak Republic		
452	Slovakia		
453	Slovenia		
721	Solomon Islands		
581	Somali Republic		
581	Somalia		

SEER GEOCODES

031	Tennessee	084	Utah
077	Texas	634	Uzbekistan
651	Thailand (Siam)	634	Uzbek, S.S.R.
685	Tibet		
245	Tobago		V
539	Togo		
136	Tokelau Islands	721	Vanuatu
725	Tonga	447	Vatican City
665	Tonkin	545	Venda
625	Trans-Jordan	321	Venezuela
545	Transkei	004	Vermont
545	Transvaal	665	Vietnam
449	Transylvania	245	Virgin Islands (British)
245	Trinidad	102	Virgin Islands (U.S.)
517	Tripoli	023	Virginia
517	Tripolitania		
629	Trucial States		W
515	Tunisia		
611	Turkey	137	Wake Island
634	Turkmen S.S.R.	402	Wales
634	Turkmenistan	449	Wallachia
245	Turks Islands	721	Wallis
125	Tuvalu	093	Washington (state)
		022	Washington D.C.
	U	530	West Africa, NOS
		539	West African countries, other
573	Uganda	631	West Bank
456	Ukraine	431	West Germany
456	Ukrainian S.S.R.	245	West Indies, NOS (see also individual islands)
404	Ulster	639	West Pakistan
545	Union of South Africa	024	West Virginia
---	Union of Soviet Socialist Republics	499	Western Europe, NOS
	(U.S.S.R.) (see individual republics)	520	Western Sahara
629	United Arab Emirates	725	Western Samoa
519	United Arab Republic	457	White Russia
400	United Kingdom	245	Windward Islands
000	United States	051	Wisconsin
102	U.S. Virgin Islands	082	Wyoming
999	Unknown		
520	Upper Volta		Y
375	Uruguay		
579	Urundi	629	Yemen

SEER GEOCODES

- 629 Yemen, People's Democratic
Republic of
- 453 Yugoslavia (former Yugoslavia
region)
- 225 Yukon Territory

Z

- 541 Zaire
- 549 Zambia
- 571 Zanzibar
- 547 Zimbabwe

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APPENDIX I:
International Organization for Standardization (ISO)
For Birthplace-State and Birthplace-Country

ISO CODES

United States Country Code = USA	
State	Code
Alabama	AL
Alaska	AK
Arizona	AZ
Arkansas	AR
Armed Forces Americas	AA
Armed Forces Canada, Europe, Middle East, Africa	AE
Armed Forces Pacific	AP
California	CA
Colorado	CO
Connecticut	CT
Delaware	DE
District of Columbia	DC
Florida	FL
Georgia	GA
Hawaii	HI
Idaho	ID
Illinois	IL
Indiana	IN
Iowa	IA
Kansas	KS
Kentucky	KY
Louisiana	LA
Maine	ME
Maryland	MD
Massachusetts	MA
Michigan	MI
Minnesota	MN
Mississippi	MS
Missouri	MO
Montana	MT
Nebraska	NE
Nevada	NV
New Hampshire	NH
New Jersey	NJ
New Mexico	NM
New York	NY
North Carolina	NC
North Dakota	ND
Ohio	OH
Oklahoma	OK
Oregon	OR
Pennsylvania	PA
Rhode Island	RI
South Carolina	SC

ISO CODES

United States Country Code = USA	
State	Code
South Dakota	SD
Tennessee	TN
Texas	TX
Utah	UT
Vermont	VT
Virginia	VA
Washington	WA
West Virginia	WV
Wisconsin	WI
Wyoming	WY
United States, NOS (specific state unknown)	US

Canada (Province and Territory) Country Code = CAN	
Province/Territory	Code
Alberta	AB
British Columbia	BC
Manitoba	MB
New Brunswick	NB
Newfoundland and Labrador	NL
Northwest Territories	NT
Northwest Territories, Yukon Territory	YN
Nova Scotia	NS
Nunavut	NU
Ontario	ON
Prince Edward Island	PE
Quebec	QC
Saskatchewan	SK
Yukon Territory	YT
Canada, NOS (specific Province/Territory unknown)	CD

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
Afghanistan	AFG	XX
African Coastal Islands (previously in South Africa, NOS)	XIF	YY
Aland Islands	ALA	XX
Albania	ALB	XX
Algeria	DZA	XX
American Samoa	ASM	AS
Andorra	AND	XX
Angola (Sao Tome, Principe, Cabinda)	AGO	XX
Anguilla	AIA	XX
Antarctica	ATA	XX
Antigua and Barbuda	ATG	XX
Arabian Peninsula	XAP	YY
Argentina	ARG	XX
Armenia	ARM	XX
Aruba	ABW	XX
Australia	AUS	XX
Australia and Australian New Guinea	AUS	XX
Austria	AUT	XX
Azerbaijan	AZE	XX
Bahamas	BHS	XX
Bahrain	BHR	XX
Bangladesh (East Pakistan)	BGD	XX
Barbados	BRB	XX
Belgium	BEL	XX
Belize (British Honduras)	BLZ	XX
Benin	BEN	XX
Bermuda	BMU	XX
Bhutan	BTN	XX
Bolivia	BOL	XX
Bonaire, Saint Eustatius and Saba	BES	XX
Bosnia and Herzegovina	BIH	XX
Botswana	BWA	XX
Bouvet Island	BVT	XX
Brazil	BRA	XX
British Indian Ocean Territory	IOT	XX
British Virgin Islands	VGB	XX
Brunei	BRN	XX
Bulgaria	BGR	XX
Burkina Faso	BFA	XX
Burma	MMR	XX
Burundi (Urundi)	BDI	XX
Byelorussia (Byelorussian SSR, White Russia)	BLR	XX
Cambodia	KHM	XX
Cameroon	CMR	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
Canal Zone	PAN	XX
Cape Verde	CPV	XX
Caucasian Republics of the USSR	XCR	YY
Cayman Islands	CYM	XX
Central African Republic	CAF	XX
Ceylon (Sri Lanka)	LKA	XX
Chad	TCD	XX
Chile	CHL	XX
China (Peoples Republic of China)	CHN	XX
China, NOS	XCH	YY
Christmas Island	CXR	XX
Cocos (Keeling) Islands	CCK	XX
Colombia	COL	XX
Comoros	COM	XX
Congo	COG	XX
Cook Islands (New Zealand)	COK	XX
Costa Rica	CRI	XX
Cote d'Ivoire	CIV	XX
Croatia	HRV	XX
Cuba	CUB	XX
Curacao	CUW	XX
Cyprus	CYP	XX
Czech Republic	CZE	XX
Czechoslovakia (former)	CSK	YY
Denmark, Faroe Islands	DNK	XX
Djibouti	DJI	XX
Dominica	DMA	XX
Dominican Republic	DOM	XX
East Africa	XEY	YY
Ecuador	ECU	XX
Egypt (United Arab Republic)	EGY	XX
El Salvador	SLV	XX
England	ENG	XX
England, Channel Islands, Isle of Man	XEN	XX
Equatorial Guinea	GNQ	XX
Eritrea	ERI	XX
Estonian SSR (Estonia)	EST	XX
Ethiopia	ETH	XX
Ethiopia (Abyssinia), Eritrea	XET	YY
Falkland Islands	FLK	XX
Faroe Islands	FRO	XX
Fiji	FJI	XX
Finland	FIN	XX
France, Corsica, Monaco	FRA	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
French Guiana	GUF	XX
French Polynesia	PYF	XX
French Southern Territories	ATF	XX
Gabon	GAB	XX
Gambia	GMB	XX
Georgia	GEO	XX
Germanic Countries	XGR	YY
Germany (East and West)	DEU	XX
Ghana	GHA	XX
Gibraltar	GIB	XX
Greece	GRC	XX
Greenland	GRL	XX
Grenada	GRD	XX
Guadeloupe	GLP	XX
Guam	GUM	GU
Guatemala	GTM	XX
Guernsey	GGY	XX
Guinea	GIN	XX
Guinea Bissau	GNB	XX
Guyana (British Guiana)	GUY	XX
Haiti	HTI	XX
Heard Island and McDonald Islands	HMD	XX
Honduras	HND	XX
Hong Kong	HKG	XX
Hungary	HUN	XX
Iceland	ISL	XX
India	IND	XX
Indochina	XSE	YY
Indonesia (Dutch East Indies)	IDN	XX
Iran (Persia)	IRN	XX
Iraq	IRQ	XX
Ireland (Eire) (Ireland NOS, Republic of Ireland)	IRL	XX
Isle of Man	IMN	XX
Israel	ISR	XX
Israel and former Jewish Palestine	XIS	YY
Italy (Sardinia, Sicily), San Marino, Vatican City	ITA	XX
Jamaica	JAM	XX
Japan	JPN	XX
Jersey	JEY	XX
Johnston Atoll	UMI	UM
Jordan (Transjordan) and former Arab Palestine	JOR	XX
Kazakhstan	KAZ	XX
Kenya	KEN	XX
Kiribati (Canton, Enderbury, Gilbert, S Lines, Phoenix)	KIR	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
Korea (North and South)	KOR	XX
Kuwait	KWT	XX
Kyrgyzstan	KGZ	XX
Laos	LAO	XX
Latvian SSR (Latvia)	LVA	XX
Lebanon	LBN	XX
Lesotho	LSO	XX
Liberia	LBR	XX
Libya (Tripoli, Tripolitania, Cyrenaica)	LBY	XX
Liechtenstein	LIE	XX
Lithuania (Lithuanian SSR)	LTU	XX
Luxembourg	LUX	XX
Macao (Macau)	MAC	XX
Macedonia	MKD	XX
Madagascar (Malagasy Republic)	MDG	XX
Malawi (Nyasaland)	MWI	XX
Malaysia	MYS	XX
Malaysia, Singapore, Brunei	XMS	YY
Mali	MLI	XX
Malta	MLT	XX
Mariana Islands (Trust Territory of Pacific Islands)	MNP	MP
Marshall Islands (Trust Territory Pacific Islands)	MHL	MH
Martinique	MTQ	XX
Mauritania	MRT	XX
Mauritius	MUS	XX
Mayotte	MYT	XX
Melanesian Islands, Solomon Islands	XML	YY
Mexico	MEX	XX
Micronesia (Fed States of) (Caroline, Trust Terr of Pacific)	FSM	FM
Micronesian Islands	XMC	YY
Mid-East Asia NOS, Maldives	MDV	XX
Moldova	MDA	XX
Monaco	MCO	XX
Mongolia	MNG	XX
Montenegro	MNE	XX
Montserrat	MSR	XX
Morocco	MAR	XX
Mozambique	MOZ	XX
Namibia	NAM	XX
Nampo-Shoto, Southern	JPN	XX
Nauru	NRU	XX
Nepal, Bhutan, Sikkim	NPL	XX
Netherlands	NLD	XX
New Caledonia	NCL	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
New Zealand	NZL	XX
Nicaragua	NIC	XX
Niger	NER	XX
Nigeria	NGA	XX
Niue	NIU	XX
Norfolk Island	NFK	XX
North Africa	XNF	YY
North American Islands	XNI	YY
North Korea	PRK	XX
Northern Ireland (Ulster)	NIR	XX
Norway (Svalbard, Jan Mayen)	NOR	XX
Oman	OMN	XX
Other Asian Republics of the USSR	XOR	YY
Other Caribbean Islands	XCB	YY
Other West African Countries	XWF	YY
Pakistan (West Pakistan)	PAK	XX
Palau (Trust Territory of Pacific Islands)	PLW	PW
Palestine Territory, Occupied	PSE	XX
Panama	PAN	XX
Papua New Guinea	PNG	XX
Paraguay	PRY	XX
Peru	PER	XX
Philippines (Philippine Islands)	PHL	XX
Pitcairn Islands	PCN	XX
Poland	POL	XX
Polynesian Islands	XPL	YY
Portugal (Madeira Islands, Azores, Cape Verde Islands)	PRT	XX
Puerto Rico	PRI	PR
Qatar	QAT	XX
Republic of South Africa	ZAF	XX
Republic of South Africa, Botswana, Lesotho, Namibia, Swaziland	XSF	YY
Réunion	REU	XX
Romania	ROU	XX
Russian SFSR (Russia)	RUS	XX
Rwanda (Ruanda)	RWA	XX
Ryukyu Islands (Japan)	JPN	XX
Samoa	WSM	XX
San Marino	SMR	XX
Sao Tome & Principe	STP	XX
Saudi Arabia	SAU	XX
Scandinavia	XSC	YY
Scotland	SCT	XX
Senegal	SEN	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
Serbia	SRB	XX
Seychelles	SYC	XX
Sierra Leone	SLE	XX
Singapore	SGP	XX
Sint-Maarten	SXM	XX
Slavic Countries	XSL	YY
Slovakia	SVK	XX
Slovenia	SVN	XX
Solomon Islands	SLB	XX
Somalia (Somali Republic, Somaliland)	SOM	XX
South Africa, NOS	XSF	YY
South Georgia and the South Sandwich Islands	SGS	XX
South Sudan	SSD	XX
Southeast Asia	XSE	YY
Spain (Canary Islands, Balearic Islands), Andorra	ESP	XX
St Pierre and Miquelon	SPM	XX
St. Barthelemy	BLM	XX
St. Helena	SHN	XX
St. Kitts and Nevis	KNA	XX
St. Lucia	LCA	XX
St. Martin (French part)	MAF	XX
St. Vincent and the Grenadines	VCT	XX
Sudan	SDN	XX
Sudanese Countries	XSD	YY
Suriname (Dutch Guiana)	SUR	XX
Svalbard and Jan Mayen	SJM	XX
Swan Islands	UMI	UM
Swaziland	SWZ	XX
Sweden	SWE	XX
Switzerland	CHE	XX
Syria	SYR	XX
Taiwan (Formosa) (Republic of China)	TWN	XX
Tajikistan	TJK	XX
Tanzania (Tanganyika, Zanzibar)	TZA	XX
Thailand (Siam)	THA	XX
Tibet	CHN	XX
Timor-Leste	TLS	XX
Togo	TGO	XX
Tokelau Islands (New Zealand)	TKL	XX
Tonga	TON	XX
Trinidad and Tobago	TTO	XX
Tunisia	TUN	XX
Turkey	TUR	XX
Turkmenistan	TKM	XX

ISO CODES

Countries outside the United States and Canada		
Country	Country Code	State Code
Turks and Caicos	TCA	XX
Tuvalu (Ellice Islands)	TUV	XX
U.S. Virgin Islands	VIR	VI
Uganda	UGA	XX
Ukraine	UKR	XX
Ukraine and Moldavia	XUM	YY
United Arab Emirates	ARE	XX
United Kingdom	GBR	XX
Uruguay	URY	XX
Uzbekistan	UZB	XX
Vanuatu	VUT	XX
Vatican City	VAT	XX
Venezuela	VEN	XX
Vietnam (Tonkin, Annam, Cochin China)	VNM	XX
Wake Island	UMI	UM
Wales	WLS	XX
Wallis and Fotuna	WLF	XX
West Africa, NOS (French Africa, NOS)	XWF	YY
Western Sahara	ESH	XX
Yemen	YEM	XX
Yugoslavia (former)	YUG	YY
Zaire (Congo-Leopoldville, Belgian Congo, Congo/Kinshasa)	COD	XX
Zambia (Northern Rhodesia)	ZMB	XX
Zimbabwe (Rhodesia, Southern Rhodesia)	ZWE	XX

ISO CODES

General Codes		
Only use in the absence of more specific information		
Country	Country Code	State Code
Africa, NOS (Central, Equatorial)	ZZF	YY
Asia, NOS	ZZA	YY
Asian and Arab Countries	ZZA	YY
Atlantic/Caribbean Area	ZZN	YY
Baltic Republic(s), NOS (Baltic States, NOS)	ZZE	YY
Central America	ZZC	YY
East Asia	ZZA	YY
Europe, NOS (Central, Eastern, Northern, Southern, Western)	ZZE	YY
Latin America, NOS	ZZU	YY
Near East	ZZA	YY
North America, NOS	ZZN	YY
Not U.S., but no other information	ZZX	YY
Other Atlantic/Caribbean Area	ZZN	YY
Other Mainland Europe	ZZE	YY
Other Mediterranean Isles	ZZE	YY
Other Pacific Area	ZZP	YY
Pacific Area	ZZP	YY
Pacific Islands	ZZP	YY
Romance-Language Countries	ZZE	YY
South America, NOS	ZZS	YY
South American Islands	ZZS	YY
Trust Territories	ZZP	TT
Unknown	ZZU	ZZ

APPENDIX J:

Pennsylvania Cancer Registry (PCR)

Required Site Specific Factors (SSFs)

PCR REQUIRED SITE SPECIFIC FACTORS

Below is a list of the Site Specific Factors (SSF) required to be reported to the PCR for all cases diagnosed on or after January 1, 2010 or any case originally coded in CSv2 (Collaborative Stage Version 2). If a schema is not listed, the PCR does not require any SSF for that schema.

Note to Registry Hospitals: All SSFs required by the PCR are also required by the American College of Surgeons (ACOS) Commission on Cancer (COC).

CS Schema Name	SSF Number	SSF Description
Appendix	2	Clinical Assessment of Regional Lymph Nodes
	11	Histopathological Grading
BileDuctsDistal	25	Schema Discriminator: Subsite of Extrahepatic Bile Ducts
BileDuctsIntraHepatic	10	Tumor Growth Pattern
BileDuctsPerihilar	25	Schema Discriminator: Subsite of Extrahepatic Bile Ducts
Bladder	1	WHO/ISUP Grade
	2	Size of Metastasis in Lymph Nodes
Brain	1	WHO Grade Classification
Breast	1	Estrogen Receptor Assay (ERA)
	2	Progesterone Receptor Assay (PRA)
	3	Number of Positive Ipsilateral Level I-II Axillary Lymph Nodes
	4	Immunohistochemistry (IHC) of Regional Lymph Nodes
	5	Molecular Studies of Regional Lymph Nodes
	7	Nottingham or Bloom-Richardson (BR) Score/Grade
	8	HER2: IHC Test Lab Value
	9	HER2: IHC Test Interpretation
	11	HER2: FISH Test Interpretation
	13	HER2: CISH Test Interpretation
	14	HER2: Result of other or unknown test
15	HER2: Summary Result of Testing	
BuccalMucosa	1	Size of Lymph Nodes
CarcinoidAppendix	2	Clinical Assessment of Regional Lymph Nodes
CNSOther	1	WHO Grade Classification
Colon	2	Clinical Assessment of Regional Lymph Nodes
Conjunctiva	1	Tumor Size
CorpusAdenosarcoma	2	Peritoneal Cytology

CS Schema Name	SSF Number	SSF Description
CorpusCarcinoma	2	Peritoneal Cytology
CorpusSarcoma	2	Peritoneal Cytology
CysticDuct	25	Schema Discriminator: Subsite of Extrahepatic Bile Ducts
EpiglottisAnterior	1	Size of Lymph Nodes
Esophagus	1	Clinical Assessment of Regional Lymph Nodes
EsophagusGEJunction	1	Clinical Assessment of Regional Lymph Nodes
	25	Schema Discriminator: Involvement of Cardia and Distance from Esophagogastric Junction (EGJ)
FloorMouth	1	Size of Lymph Nodes
GISTAppendix	11	Mitotic Count
GISTColon	11	Mitotic Count
GISTEsophagus	6	Mitotic Count
GISTPeritoneum	5	Mitotic Count
	10	Location of Primary Tumor
GISTRectum	11	Mitotic Count
GISTSmallIntestine	6	Mitotic Count
GISTStomach	6	Mitotic Count
GumLower	1	Size of Lymph Nodes
GumOther	1	Size of Lymph Nodes
GumUpper	1	Size of Lymph Nodes
HeartMediastinum	1	Grade for Sarcomas
HemeRetic	1	JAK-2
Hypopharynx	1	Size of Lymph Nodes
IntracranialGland	1	WHO Grade Classification
KidneyParenchyma	6	Fuhrman Nuclear Grade
KidneyRenalPelvis	1	WHO/ISUP Grade
LacrimalGland	25	Schema Discriminator: Lacrimal Gland/Lacrimal Sac
LacrimalSac	25	Schema Discriminator: Lacrimal Gland/Lacrimal Sac
LarynxGlottic	1	Size of Lymph Nodes
LarynxOther	1	Size of Lymph Nodes
LarynxSubglottic	1	Size of Lymph Nodes
LarynxSupraglottic	1	Size of Lymph Nodes

CS Schema Name	SSF Number	SSF Description
LipLower	1	Size of Lymph Nodes
LipOther	1	Size of Lymph Nodes
LipUpper	1	Size of Lymph Nodes
Lung	1	Separate Tumor Nodules/Ipsilateral Lung
Lymphoma	2	Systemic Symptoms at Diagnosis
LymphomaOcularAdnexa	2	Systemic Symptoms at Diagnosis
MelanomaChoroid	2	Measured Basal Diameter
	3	Measured Thickness (Depth)
	4	Size of Largest Metastasis
MelanomaCiliaryBody	2	Measured Basal Diameter
	3	Measured Thickness (Depth)
	4	Size of Largest Metastasis
	25	Schema Discriminator: Melanoma Ciliary Body/Melanoma Iris
MelanomaConjunctiva	1	Measured Thickness (Depth)
	2	Quadrants
MelanomaIris	4	Size of Largest Metastasis
	25	Schema Discriminator: Melanoma Ciliary Body/Melanoma Iris
MelanomaSkin	1	Measured Thickness (Depth), Breslow's Measurement
	2	Ulceration
	3	Clinical Status of Lymph Node Mets
	4	LDH
	7	Primary Tumor Mitotic Count/Rate
MerkelCellPenis	3	Clinical Status of Lymph Node Mets
MerkelCellScrotum	3	Clinical Status of Lymph Node Mets
MerkelCellSkin	3	Clinical Status of Lymph Node Mets
MerkelCellVulva	3	Clinical Status of Lymph Node Mets
	11	Regional Lymph Node - Laterality
MouthOther	1	Size of Lymph Nodes
MycosisFungoides	1	Peripheral Blood Involvement
NasalCavity	1	Size of Lymph Nodes
Nasopharynx	1	Size of Lymph Nodes

CS Schema Name	SSF Number	SSF Description
	25	Schema Discriminator
NETColon	2	Clinical Assessment of Regional Lymph Nodes
NETRectum	2	Clinical Assessment of Regional Lymph Nodes
NETStomach	1	Clinical Assessment of Regional Lymph Nodes
Oropharynx	1	Size of Lymph Nodes
PalateHard	1	Size of Lymph Nodes
PalateSoft	1	Size of Lymph Nodes
ParotidGland	1	Size of Lymph Nodes
Penis	17	Extranodal Extension of Regional Lymph Nodes
Peritoneum	1	Grade for Sarcomas
	25	Schema Discriminator
PeritoneumFemaleGen	25	Schema Discriminator
PharyngealTonsil	1	Size of Lymph Nodes
	25	Schema Discriminator
Placenta	1	Prognostic Scoring Index Table 1
Pleura	1	Pleural Effusion
Prostate	1	Prostatic Specific Antigen (PSA) Lab Value
	3	CS Extension – Pathologic Extension
	7	Gleason’s Primary Pattern and Secondary Pattern Value on Needle Core Biopsy/TURP
	8	Gleason’s Score on Needle Core Biopsy/TURP
	9	Gleason’s Primary Pattern and Secondary Pattern Value on Prostatectomy/Autopsy
	10	Gleason’s Score on Prostatectomy /Autopsy
Rectum	2	Clinical Assessment of Regional Lymph Nodes
Retinoblastoma	1	Extension Evaluated at Enucleation
Retroperitoneum	1	Grade for Sarcomas
SalivaryGlandOther	1	Size of Lymph Nodes
Scrotum	12	High Risk Features
	16	Size of Lymph Nodes
SinusEthmoid	1	Size of Lymph Nodes
SinusMaxillary	1	Size of Lymph Nodes
Skin	12	High Risk Features

CS Schema Name	SSF Number	SSF Description
	16	Size of Lymph Nodes
SkinEyelid	6	Perineural Invasion
SmallIntestine	2	Clinical Assessment of Regional Lymph Nodes
SoftTissue	1	Grade for Sarcomas
Stomach	1	Clinical Assessment of Regional Lymph Nodes
	25	Involvement of Cardia and Distance from Esophagogastric Junction (EGJ)
SubmandibularGland	1	Size of Lymph Nodes
Testis	4	Radical Orchiectomy Performed
	5	Size of Metastasis in Lymph Nodes
	7	Preorchiectomy Alpha Fetoprotein (AFP) Interpretation
	9	Preorchiectomy Human chorionic gonadotropin (hCG) Interpretation
	10	Preorchiectomy LDH Interpretation
	13	Post-Orchiectomy Alpha Fetoprotein (AFP) Range
	15	Post-Orchiectomy Human Chorionic Gonadotropin (hCG) Range
	16	Post-Orchiectomy Lactate Dehydrogenase (LDH) Range
TongueAnterior	1	Size of Lymph Nodes
TongueBase	1	Size of Lymph Nodes
Urethra	1	WHO/ISUP Grade
Vulva	11	Regional Lymph Node - Laterality

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**APPENDIX K:
SURGERY CODES**

Alphabetical Index by Primary Site for Surgery Codes

Primary Site	ICD-O Code(s)	Appendix K Page #
Accessory sinuses	C31.0-C31.9	44
Adnexa of eye	C69.0-C69.9	44
Adrenal gland	C74.0-C74.9	44
Anal canal and anus	C21.0-C21.8	17
Anus and anal canal	C21.0-C21.8	17
Articular cartilage, bones, & joints of limbs & other & unspecified sites	C40.0-C41.9	24
Autonomic nervous system and peripheral nerves	C47.0-C47.9	24
Biliary tract, other, and unspecified parts of	C24.0-C24.9	44
Bladder	C67.0-C67.9	39
Blood	C42.0	22
Bone marrow	C42.1	22
Bones, joints, and articular cartilage of limbs and other and unspecified sites	C40.0-C41.9	24
Brain	C71.0-C71.9	41
Breast	C50.0-C50.9	27
Bronchus and lung	C34.0-C34.9	21
Cervix uteri	C53.0-C53.9	29
Central Nervous System (CNS) parts, other, spinal cord, and cranial nerves	C72.0-C72.9	41
Colon	C18.0-C18.9	11
Connective, subcutaneous, and other soft tissues	C49.0-C49.9	24
Corpus uteri	C54.0-C55.9	31
Cranial nerves, spinal cord, and other CNS parts	C72.0-C72.9	41
Digestive organs, other, and ill-defined digestive organs	C26.0-C26.9	44
Ear, middle	C30.1	44
Endocrine glands, other, and related structures	C75.0-C75.9	44
Esophagus	C15.0-C15.9	8
Eye and adnexa of eye	C69.0-C69.9	44
Gallbladder	C23.9	44
Gastrointestinal tract, NOS	C26.9	44
Genital organs, female, other, and unspecified organs	C57.0-C57.9	44
Genital organs, male, other, and unspecified organs	C63.0-C63.9	44
Gum	C03.0-C03.9	4
Heart	C38.0	44
Heart, mediastinum, and pleura, overlapping lesion of	C38.8	44
Hematopoietic Diseases	C42.0-C42.4	22
Hypopharynx	C13.0-C13.9	7
Ill-defined primary	C76.0-C76.8	45
Immunoproliferative Diseases	C42.0-C42.4	22
Intestinal tract, NOS	C26.0	44
Intrahepatic bile ducts	C22.1	18
Intrathoracic organs and respiratory system, other, and ill-defined sites	C39.0-C39.9	44
Joints, bones, & articular cartilage of limbs & other & unspecified	C40.0-C41.9	24

Primary Site	ICD-O Code(s)	Appendix K Page #
Kidney	C64.9	38
Larynx	C32.0-C32.9	20
Lip	C00.0-C00.9	4
Oral cavity, and pharynx, other, and ill-defined sites in	C14.2-C14.8	44
Liver	C22.0	18
Lung and bronchus	C34.0-C34.9	21
Lymph nodes	C77.0-C77.9	43
Mediastinum	C38.1-C38.3	44
Mediastinum, heart, and pleura, overlapping lesion of	C38.8	44
Meninges	C70.0-C70.9	41
Mouth, floor of	C04.0-C04.9	4
Mouth, other, and unspecified parts of	C06.0-C06.9	4
Myeloproliferative Disorders	C42.1	22
Nasal cavity	C30.0	44
Nasopharynx	C11.0-C11.9	7
Oral cavity	C00.0-C06.9	4
Oral cavity and pharynx, other and ill-defined sites of	C14.2-C14.8	44
Oropharynx	C10.0-C10.9	7
Ovary	C56.9	33
Palate	C05.0-C05.9	4
Pancreas	C25.0-C25.9	19
Parotid gland	C07.9	5
Penis	C60.0-C60.9	44
Peripheral nerves and autonomic nervous system	C47.0-C47.9	24
Peritoneum	C48.1-C48.2	44
Peritoneum and retroperitoneum, overlapping lesion of	C48.8	44
Pharynx, oral cavity, other and ill-defined sites of	C14.2-C14.8	44
Pharynx, NOS	C14.0	7
Placenta	C58.9	44
Pleura, heart, mediastinum, overlapping lesion of	C38.8	44
Pleura, NOS	C38.4	44
Prostate gland	C61.9	35
Pyiform sinus	C12.9	7
Rectosigmoid junction	C19.9	13
Rectum	C20.9	15
Renal pelvis	C65.9	38
Respiratory system and intrathoracic organs, other, and ill-defined sites within	C39.0-C39.9	44
Reticuloendothelial System	C42.3	22
Retroperitoneum	C48.0	44
Retroperitoneum and peritoneum, overlapping lesion of	C48.8	44
Salivary glands, major; other, and unspecified glands	C08.0-C08.9	5
Skin	C44.0-C44.9	26
Small intestine	C17.0-C17.9	44
Soft tissues, other, and connective and subcutaneous tissues	C49.0-C49.9	24
Spinal cord, cranial nerves, and other CNS parts	C72.0-C72.9	41
Spleen	C42.2	25

Primary Site	ICD-O Code(s)	Appendix K Page #
Stomach	C16.0-C16.9	9
Subcutaneous, connective, and other soft tissues	C49.0-C49.9	24
Testis	C62.0-C62.9	37
Thymus	C37.9	44
Thyroid gland	C73.9	42
Tongue, base of	C01.9	4
Tongue, other, and unspecified parts of	C02.0-C02.9	4
Tonsil	C09.0-C09.9	7
Trachea	C33.9	44
Unknown primary site	C80.9	45
Ureter	C66.9	38
Urinary bladder	C67.0-C67.9	39
Urinary organs, other and unspecified organs	C68.0-C68.9	44
Uterus, NOS	C55.9	31
Vagina	C52.9	44
Vulva	C51.0-C51.9	44

ORAL CAVITY

**Lip C00.0-C00.9, Base of Tongue C01.9, Other Parts of Tongue
C02.0-C02.9, Gum C03.0-C03.9, Floor of Mouth C04.0-C04.9, Palate
C05.0-C05.9, Other Parts of Mouth C06.0-C06.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

00 None; no surgery of primary site; autopsy ONLY

10 Local tumor destruction, NOS

11 Photodynamic therapy (PDT)

12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)

13 Cryosurgery

14 Laser

No specimen sent to pathology from surgical events 10–14.

20 Local tumor excision, NOS

26 Polypectomy

27 Excisional biopsy

Any combination of 20 or 26–27 WITH

21 Photodynamic therapy (PDT)

22 Electrocautery

23 Cryosurgery

24 Laser ablation

25 Laser excision

30 Wide excision, NOS

Code 30 includes:

Hemiglossectomy

Partial glossectomy

40 Radical excision of tumor, NOS

41 Radical excision of tumor ONLY

42 Combination of 41 WITH resection in continuity with mandible (marginal, segmental, hemi-, or total resection)

43 Combination of 41 WITH resection in continuity with maxilla (partial, subtotal, or total resection)

Codes 40–43 include:

Total glossectomy

Radical glossectomy

Specimen sent to pathology from surgical events 20–43.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

PAROTID AND OTHER UNSPECIFIED GLANDS
Parotid Gland C07.9, Major Salivary Glands C08.0-C08.9
(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY

- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser

No specimen sent to pathology from surgical events 10–14.

- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy
 - Any combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision

- 30 Less than total parotidectomy, NOS; less than total removal of major salivary gland, NOS
 - 31 Facial nerve spared
 - 32 Facial nerve sacrificed
 - 33 Superficial lobe ONLY
 - 34 Facial nerve spared
 - 35 Facial nerve sacrificed
 - 36 Deep lobe (Total)
 - 37 Facial nerve spared
 - 38 Facial nerve sacrificed

- 40 Total parotidectomy, NOS; total removal of major salivary gland, NOS
 - 41 Facial nerve spared
 - 42 Facial nerve sacrificed

- 50 Radical parotidectomy, NOS; radical removal of major salivary gland, NOS
 - 51 WITHOUT removal of temporal bone
 - 52 WITH removal of temporal bone
 - 53 WITH removal of overlying skin (requires graft or flap coverage)

- 80 Parotidectomy, NOS

Specimen sent to pathology from surgical events 20–80.

- 90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

PHARYNX

Tonsil C09.0-C09.9, Oropharynx C10.0-C10.9, Nasopharynx C11.0-C11.9, Pyriform Sinus C12.9, Hypopharynx C13.0-C13.9, Pharynx C14.0

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Stripping

No specimen sent to pathology from surgical events 10–15.
- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy
 - Any combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision
 - 28 Stripping
- 30 Pharyngectomy, NOS
 - 31 Limited/partial pharyngectomy; tonsillectomy, bilateral tonsillectomy
 - 32 Total pharyngectomy
- 40 Pharyngectomy WITH laryngectomy OR removal of contiguous bone tissue, NOS (does NOT include total mandibular resection)
 - 41 WITH Laryngectomy (laryngopharyngectomy)
 - 42 WITH bone
 - 43 WITH both 41 and 42
- 50 Radical pharyngectomy (includes total mandibular resection), NOS
 - 51 WITHOUT laryngectomy
 - 52 WITH laryngectomy

Specimens sent to pathology from surgical events 20–52.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**ESOPHAGUS
C15.0-C15.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE**Codes**

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser**No specimen sent to pathology from surgical events 10–14.**
- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy
 - Any combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision
- 30 Partial esophagectomy
- 40 Total esophagectomy, NOS
- 50 Esophagectomy, NOS WITH laryngectomy and/or gastrectomy, NOS
 - 51 WITH laryngectomy
 - 52 WITH gastrectomy, NOS
 - 53 Partial gastrectomy
 - 54 Total gastrectomy
 - 55 Combination of 51 WITH any of 52–54
- 80 Esophagectomy, NOS

Specimen sent to pathology from surgical events 20–80.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

STOMACH C16.0-C16.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
- No specimen sent to pathology from surgical events 10–14.**

- 20 Local tumor excision, NOS
- 26 Polypectomy
 - 27 Excisional biopsy
- Any combination of 20 or 26–27 WITH
- 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
- 25 Laser excision
- 30 Gastrectomy, NOS (partial, subtotal, hemi-)
- 31 Antrectomy, lower (distal-less than 40% of stomach)***
 - 32 Lower (distal) gastrectomy (partial, subtotal, hemi-)
 - 33 Upper (proximal) gastrectomy (partial, subtotal, hemi-)

Code 30 includes:

Partial gastrectomy, including a sleeve resection of the stomach
Billroth I: anastomosis to duodenum (duodenostomy)
Billroth II: anastomosis to jejunum (jejunostomy)

- 40 Near-total or total gastrectomy, NOS
- 41 Near-total gastrectomy
 - 42 Total gastrectomy
- A total gastrectomy may follow a previous partial resection of the stomach.**
- 50 Gastrectomy, NOS WITH removal of a portion of esophagus
- 51 Partial or subtotal gastrectomy
 - 52 Near total or total gastrectomy
- Codes 50–52 are used for gastrectomy resection when only portions of esophagus are included in procedure.**

- 60 Gastrectomy with a resection in continuity with the resection of other organs, NOS***

61 Partial or subtotal gastrectomy, in continuity with the resection of other organs***

62 Near total or total gastrectomy, in continuity with the resection of other organs***

63 Radical gastrectomy, in continuity with the resection of other organs***

Codes 60–63 are used for gastrectomy resections with organs other than esophagus.

Portions of esophagus may or may not be included in the resection.

80 Gastrectomy, NOS

Specimen sent to pathology from surgical events 20–80

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

*** Incidental splenectomy NOT included

COLON

Code 70 includes: Any colectomy (partial, hemicolectomy, or total) WITH a resection of any other organs in continuity with the primary site. Other organs may be partially or totally removed. Other organs may include, but are not limited to, oophorectomy, partial proctectomy, rectal mucosectomy, or pelvic exenteration.

80 Colectomy, NOS

Specimen sent to pathology from surgical events 20–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

**RECTOSIGMOID
C19.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Code removal/surgical ablation of single or multiple liver metastases under the data item *Surgical Procedure/Other Site*.

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser ablation

No specimen sent to pathology from surgical events 10–14.

- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy

Combination of 20 or 26–27 WITH

 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation- 25 Laser excision
- 30 Wedge or segmental resection; partial proctosigmoidectomy, NOS
 - 31 Plus resection of contiguous organs; example: small bowel, bladder

Procedures coded 30 include, but are not limited to:

- Anterior resection
 - Hartmann operation
 - Low anterior resection (LAR)
 - Partial colectomy, NOS
 - Rectosigmoidectomy, NOS
 - Sigmoidectomy
- 40 Pull through WITH sphincter preservation (colo-anal anastomosis)
 - 50 Total proctectomy
 - 51 Total colectomy
 - 55 Total colectomy WITH ileostomy, NOS
 - 56 Ileorectal reconstruction
 - 57 Total colectomy WITH other pouch; example: Koch pouch
 - 60 Total proctocolectomy, NOS
 - 65 Total proctocolectomy WITH ileostomy, NOS
 - 66 Total proctocolectomy WITH ileostomy and pouch

Removal of the colon from cecum to the rectosigmoid or a portion of the rectum.

70 Colectomy or proctocolectomy resection in continuity with other organs; pelvic exenteration

80 Colectomy, NOS; Proctectomy, NOS

Specimen sent to pathology from surgical events 20–27.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

RECTUM C20.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Code removal/surgical ablation of single or multiple liver metastases under the data item *Surgical Procedure/Other Site*.

Codes

- 00 None; no surgery of primary site; autopsy ONLY

- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser

No specimen sent to pathology from surgical events 10-14.

- 20 Local tumor excision, NOS
 - 27 Excisional biopsy
 - 26 Polypectomy
 - Any combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision
 - 28 Curette and fulguration

- 30 Wedge or segmental resection; partial proctectomy, NOS

Procedures coded 30 include, but are not limited to:

 - Anterior resection
 - Hartmann's operation
 - Low anterior resection (LAR)
 - Transsacral rectosigmoidectomy

- 40 Pull through WITH sphincter preservation (coloanal anastomosis)

- 50 Total proctectomy- **Procedure coded 50 includes, but is not limited to:** Abdominoperineal resection (Miles Procedure)

- 60 Total proctocolectomy, NOS

- 70 Proctectomy or proctocolectomy with resection in continuity with other organs; pelvic exenteration

- 80 Proctectomy, NOS

Specimen sent to pathology from surgical events 20–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

**ANUS
C21.0-C21.8**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE**Codes**

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Thermal Ablation
- No specimen sent to pathology from surgical events 10–15.**
- 20 Local tumor excision, NOS
- 26 Polypectomy
 - 27 Excisional biopsy
- Any combination of 20 or 26–27 WITH
- 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
- 25 Laser excision
- 60 Abdominal perineal resection, NOS (APR; Miles procedure)
- 61 APR and sentinel node excision
 - 62 APR and unilateral inguinal lymph node dissection
 - 63 APR and bilateral inguinal lymph node dissection
- The lymph node dissection should also be coded under *Scope of Regional Lymph Node Surgery*.**

Specimen sent to pathology from surgical events 20–63.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

LIVER AND INTRAHEPATIC BILE DUCTS

C22.0-C22.1

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Alcohol (Percutaneous Ethanol Injection-PEI)
 - 16 Heat-Radio-frequency ablation (RFA)
 - 17 Other (ultrasound, acetic acid)
- No specimen sent to pathology from surgical events 10–17.**
- 20 Wedge or segmental resection, NOS
 - 21 Wedge resection
 - 22 Segmental resection, NOS
 - 23 One
 - 24 Two
 - 25 Three
 - 26 Segmental resection AND local tumor destruction
- 30 Lobectomy, NOS
 - 36 Right lobectomy
 - 37 Left lobectomy
 - 38 Lobectomy AND local tumor destruction
- 50 Extended lobectomy, NOS (extended: resection of a single lobe plus a segment of another lobe)
 - 51 Right lobectomy
 - 52 Left lobectomy
 - 59 Extended lobectomy AND local tumor destruction
- 60 Hepatectomy, NOS
 - 61 Total hepatectomy and transplant
- 65 Excision of a bile duct (for an intra-hepatic bile duct primary only)
 - 66 Excision of an intrahepatic bile duct PLUS partial hepatectomy
- 75 Extrahepatic bile duct and hepatectomy WITH transplant

Specimen sent to pathology from surgical events 20–75.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**PANCREAS
C25.0-C25.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 25 Local excision of tumor, NOS
- 30 Partial pancreatectomy, NOS; example: distal
- 35 Local or partial pancreatectomy and duodenectomy
 - 36 WITHOUT distal/partial gastrectomy
 - 37 WITH partial gastrectomy (Whipple)
- 40 Total pancreatectomy
- 60 Total pancreatectomy and subtotal gastrectomy or duodenectomy
- 70 Extended pancreatoduodenectomy
- 80 Pancreatectomy, NOS
- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

LARYNX

C32.0-C32.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE**Codes**

- 00 None; no surgery of primary site; autopsy ONLY

- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Stripping**No specimen sent to pathology from surgical events 10–15.**

- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy
 - Any combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision
 - 28 Stripping

- 30 Partial excision of the primary site, NOS; subtotal/partial laryngectomy NOS; hemilaryngectomy, NOS
 - 31 Vertical laryngectomy
 - 32 Anterior commissure laryngectomy
 - 33 Supraglottic laryngectomy

- 40 Total or radical laryngectomy, NOS
 - 41 Total laryngectomy ONLY
 - 42 Radical laryngectomy ONLY

50 Pharyngolaryngectomy

80 Laryngectomy, NOS

Specimen sent to pathology from surgical events 20–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

LUNG C34.0-C34.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 19 Local tumor destruction or excision, NOS
Unknown whether a specimen was sent to pathology for surgical events coded 19
- 15 Local tumor destruction, NOS
 - 12 Laser ablation or cryosurgery
 - 13 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
No specimen sent to pathology from surgical events 12–13 and 15.
- 20 Excision or resection of less than one lobe, NOS
 - 23 Excision, NOS
 - 24 Laser excision
 - 25 Bronchial sleeve resection ONLY
 - 21 Wedge resection
 - 22 Segmental resection, including lingulectomy
- 30 Resection of lobe or bilobectomy, but less than the whole lung (partial pneumonectomy, NOS)
 - 33 Lobectomy WITH mediastinal lymph node dissection
The lymph node dissection should be coded under *Scope of Regional Lymph Node Surgery*.
- 45 Lobe or bilobectomy extended, NOS
 - 46 WITH chest wall
 - 47 WITH pericardium
 - 48 WITH diaphragm
- 55 Pneumonectomy, NOS
 - 56 WITH mediastinal lymph node dissection (radical pneumonectomy)
The mediastinal lymph node dissection should also be coded under *Scope of Regional Lymph Node Surgery*.
- 65 Extended pneumonectomy
 - 66 Extended pneumonectomy plus pleura or diaphragm
- 70 Extended radical pneumonectomy
The mediastinal lymph node dissection should also be coded under *Scope of Regional Lymph Node Surgery*.
- 80 Resection of lung, NOS
- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

HEMATOPOIETIC/RETICULOENDOTHELIAL/IMMUNOPROLIFERATIVE

**/MYELOPROLIFERATIVE DISEASE
C42.0, C42.1, C42.3, C42.4 (with any histology) or
9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-
9967, and 9975-9992 (with any site)**

Hematopoietic, reticuloendothelial, immunoproliferative, or myeloproliferative diseases	
C42.0	Blood
C42.1	Bone marrow
C42.3	Reticuloendothelial system
C42.4	Hematopoietic system
9731/9734	Plasmacytoma
9732	Multiple myeloma
9733	Plasma cell leukemia
9734	Plasmacytoma, extramedullary
9740-9742	Mast cell tumors
9750-9754	Malignant histiocytosis
9755	Histiocytic sarcoma
9756	Langerhans cell sarcoma
9758	Follicular dendritic cell sarcoma
9760	Immunoproliferative disease, NOS
9761	Waldenstrom macroglobulinemia
9762	Heavy chain disease
9764	Immunoproliferative small intestinal disease (Mediterranean lymphoma)
9800-9827, 9831-9920, 9940-9948	Leukemias
9930	Myeloid sarcoma
9931	Acute panmyelosis with fibrosis
9950	Polycythemia vera
9960	Chronic myeloproliferative disorder
9961	Myelosclerosis with myeloid metaplasia
9962	Essential thrombocythemia
9963	Chronic neutrophilic leukemia
9964	Hypereosinophilic syndrome
9965-9967	Myeloid and lymphoid neoplasms
9971	Polymorphic PTLD
9975	Myelodysplastic/Myeloproliferative neoplasm, unclassifiable, Myeloproliferative disease, NOS, Myeloproliferative neoplasm, unclassifiable
9980-9985	Refractory anemias
9986	Myelodysplastic syndrome with 5q deletion syndrome
9945	Therapy-related myelodysplastic syndrome, NOS
9989	Myelodysplastic syndrome, NOS
9991	Refractory neutropenia
9992	Refractory thrombocytopenia

SURGERY OF PRIMARY SITE

Code

- 98 All hematopoietic/reticuloendothelial/immunoproliferative/myeloproliferative disease sites and/or histologies, WITH or WITHOUT surgical treatment.

Surgical procedures for hematopoietic/reticuloendothelial/immunoproliferative/myeloproliferative primaries are to be recorded using the data item *Surgical Procedure/Other Site*.

**BONES, JOINTS, AND ARTICULAR CARTILAGE C40.0-C41.9
PERIPHERAL NERVES AND AUTONOMIC NERVOUS SYSTEM**

C47.0-C47.9
CONNECTIVE, SUBCUTANEOUS, AND OTHER SOFT TISSUES
C49.0-C49.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 19 Local tumor destruction or excision, NOS
Unknown whether a specimen was sent to pathology for surgical events coded 19 (principally for cases diagnosed prior to January 1, 2003).
- 15 Local tumor destruction
No specimen sent to pathology from surgical event 15.
- 25 Local excision
- 26 Partial resection
- 30 Radical excision or resection of lesion WITH limb salvage
- 40 Amputation of limb
 - 41 Partial amputation of limb
 - 42 Total amputation of limb
- 50 Major amputation, NOS
 - 51 Forequarter, including scapula
 - 52 Hindquarter, including ilium/hip bone
 - 53 Hemipelvectomy, NOS
 - 53 Internal hemipelvectomy

Specimen sent to pathology from surgical events 25–54.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

SPLEEN
C42.2

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

00 None; no surgery of primary site; autopsy ONLY

19 Local tumor destruction or excision, NOS

Unknown whether a specimen was sent to pathology for surgical events coded 19 principally for cases diagnosed prior to January 1, 2003).

21 Partial splenectomy

22 Total splenectomy

80 Splenectomy, NOS

Specimen sent to pathology from surgical events 21–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

**SKIN
C44.0-C44.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser ablation
- No specimen sent to pathology from surgical events 10–14.**
- 20 Local tumor excision, NOS
- 26 Polypectomy
 - 27 Excisional biopsy
- Any combination of 20 or 26–27 WITH
- 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
- 25 Laser excision
- 30 Biopsy of primary tumor followed by a gross excision of the lesion (does not have to be done under the same anesthesia)
- 31 Shave biopsy followed by a gross excision of the lesion
 - 32 Punch biopsy followed by a gross excision of the lesion
 - 33 Incisional biopsy followed by a gross excision of the lesion
 - 34 Mohs surgery, NOS
 - 35 Mohs with 1-cm margin or less
 - 36 Mohs with more than 1-cm margin
- 45 Wide excision or reexcision of lesion or minor (local) amputation with margins more than 1 cm, NOS. Margins **MUST** be microscopically negative.
- 46 WITH margins more than 1 cm and less than or equal to 2 cm
 - 47 WITH margins greater than 2 cm
- If the excision does not have microscopically negative margins greater than 1 cm, use the appropriate code, 20-36.**
- 60 Major amputation
- Specimen sent to pathology from surgical events 20–60.**
- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

BREAST C50.0-C50.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

00 None; no surgery of primary site; autopsy ONLY

19 Local tumor destruction, NOS

No specimen was sent to pathology for surgical events coded 19 (principally for cases diagnosed prior to January 1, 2003).

20 Partial mastectomy, NOS; less than total mastectomy, NOS

21 Partial mastectomy WITH nipple resection

22 Lumpectomy or excisional biopsy

23 Reexcision of the biopsy site for gross or microscopic residual disease

24 Segmental mastectomy (including wedge resection, quadrantectomy, tylectomy)

Procedures coded 20–24 remove the gross primary tumor and some of the breast tissue (breast-conserving or preserving). There may be microscopic residual tumor.

30 Subcutaneous mastectomy

A subcutaneous mastectomy, also called a nipple sparing mastectomy, is the removal of breast tissue without the nipple and areolar complex or overlying skin. It is performed to facilitate immediate breast reconstruction. Cases coded 30 may be considered to have undergone breast reconstruction.

40 Total (simple) mastectomy

41 WITHOUT removal of uninvolved contralateral breast

43 Reconstruction NOS

44 Tissue

45 Implant

46 Combined (Tissue and Implant)

42 WITH removal of uninvolved contralateral breast

47 Reconstruction NOS

48 Tissue

49 Implant

75 Combined (Tissue and Implant)

A total (simple) mastectomy removes all breast tissue, the nipple, and areolar complex. An axillary dissection is not done, but sentinel lymph nodes may be removed.

For single primaries only, code removal of involved contralateral breast under the data item *Surgical Procedure/Other Site*.

If contralateral breast reveals a second primary, each breast is abstracted separately. The surgical procedure is coded 41 for the first primary. The surgical code for the contralateral breast is coded to the procedure performed on that site.

Reconstruction that is planned as part of first course treatment is coded 43-49 or 75, whether it is done at the time of mastectomy or later

- 76 Bilateral mastectomy for a single tumor involving both breasts, as for bilateral inflammatory carcinoma.
- 50 Modified radical mastectomy
- 51 WITHOUT removal of uninvolved contralateral breast
 - 53 Reconstruction, NOS
 - 54 Tissue
 - 55 Implant
 - 56 Combined (Tissue and Implant)
 - 52 WITH removal of uninvolved contralateral breast
 - 57 Reconstruction, NOS
 - 58 Tissue
 - 59 Implant
 - 63 Combined (Tissue and Implant)

Removal of all breast tissue, nipple, areolar complex, and variable amounts of breast skin in continuity with axilla. Specimen may or may not include portion of pectoralis major muscle.

If contralateral breast reveals a second primary, it is abstracted separately. The surgical procedure is coded 41 or 51 for the first primary. The surgical code for the contralateral breast is coded to the procedure performed on that site.

For single primaries only, code removal of involved contralateral breast under the data item *Surgical Procedure/Other Site*.

- 60 Radical mastectomy, NOS
- 61 WITHOUT removal of uninvolved contralateral breast
 - 64 Reconstruction, NOS
 - 65 Tissue
 - 66 Implant
 - 67 Combined (Tissue and Implant)
 - 62 WITH removal of uninvolved contralateral breast
 - 68 Reconstruction, NOS
 - 69 Tissue
 - 73 Implant
 - 74 Combined (Tissue and Implant)
- 70 Extended radical mastectomy
- 71 WITHOUT removal of uninvolved contralateral breast
 - 72 WITH removal of uninvolved contralateral breast

- 80 Mastectomy, NOS

Specimen sent to pathology from surgical events 20–80.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**CERVIX UTERI
C53.0-C53.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE**Codes**

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Loop Electrocautery Excision Procedure (LEEP)
 - 16 Laser ablation
 - 17 Thermal ablation
- No specimen sent to pathology from surgical events 10–17.**
- 20 Local tumor excision, NOS
- 26 Excisional biopsy, NOS
 - 27 Cone biopsy
 - 24 Cone biopsy WITH gross excision of lesion
 - 29 Trachelectomy; removal of cervical stump; cervicectomy
- Any combination of 20, 24, 26, 27 or 29 WITH
- 21 Electrocautery
 - 22 Cryosurgery
 - 23 Laser ablation or excision
- 25 Dilatation and curettage; endocervical curettage (for in situ only)
- 28 Loop electrocautery excision procedure (LEEP)
- 30 Total hysterectomy (simple, pan-) WITHOUT removal of tubes and ovaries
Total hysterectomy removes both the corpus and cervix uteri and may also include a portion of vaginal cuff.
- 40 Total hysterectomy (simple, pan-) WITH removal of tubes and/or ovary
Total hysterectomy removes both the corpus and cervix uteri and may also include a portion of vaginal cuff.
- 50 Modified radical or extended hysterectomy; radical hysterectomy; extended radical hysterectomy
- 51 Modified radical hysterectomy
 - 52 Extended hysterectomy
 - 53 Radical hysterectomy; Wertheim procedure
 - 54 Extended radical hysterectomy
- 60 Hysterectomy, NOS, WITH or WITHOUT removal of tubes and ovaries
- 61 WITHOUT removal of tubes and ovaries
 - 62 WITH removal of tubes and ovaries
- 70 Pelvic exenteration
- 71 Anterior exenteration

Includes bladder, distal ureters, and genital organs WITH their ligamentous attachments and pelvic lymph nodes.

- 72 Posterior exenteration
Includes rectum and rectosigmoid WITH ligamentous attachments and pelvic lymph nodes.
- 73 Total exenteration
Includes removal of all pelvic contents and pelvic lymph nodes.
- 74 Extended exenteration
Includes pelvic blood vessels or bony pelvis.

Specimen sent to pathology from surgical events 20–29.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**CORPUS UTERI
C54.0-C55.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 19 Local tumor destruction or excision, NOS
Unknown whether a specimen was sent to pathology for surgical events coded 19 (principally for cases diagnosed prior to January 1, 2003).
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Loop Electrocautery Excision Procedure (LEEP)
 - 16 Thermal ablation
- No specimen sent to pathology from surgical events 10–16.**
- 20 Local tumor excision, NOS; simple excision, NOS
- 24 Excisional biopsy
 - 25 Polypectomy
 - 26 Myomectomy
- Any combination of 20 or 24–26 WITH
- 21 Electrocautery
 - 22 Cryosurgery
 - 23 Laser ablation or excision
- 30 Subtotal hysterectomy/supracervical hysterectomy/fundectomy WITH or WITHOUT removal of tube(s) and ovary(ies).
- 31 WITHOUT tube(s) and ovary(ies)
 - 32 WITH tube(s) and ovary(ies)
- 40 Total hysterectomy (simple, pan-) WITHOUT removal of tube(s) and ovary(ies)
Removes both the corpus and cervix uteri. It may also include a portion of the vaginal cuff.
- 50 Total hysterectomy (simple, pan-) WITH removal of tube(s) and/or ovary(ies)
Removes both the corpus and cervix uteri. It may also include a portion of the vaginal cuff.
- 60 Modified radical or extended hysterectomy; radical hysterectomy; extended radical hysterectomy
- 61 Modified radical hysterectomy
 - 62 Extended hysterectomy
 - 63 Radical hysterectomy; Wertheim procedure
 - 64 Extended radical hysterectomy
- 65 Hysterectomy, NOS, WITH or WITHOUT removal of tube(s) and ovary(ies)
- 66 WITHOUT removal of tube(s) and ovary(ies)
 - 67 WITH removal of tube(s) and ovary(ies)

- 75 Pelvic exenteration
 - 76 Anterior exenteration
Includes bladder, distal ureters, and genital organs WITH their ligamentous attachments and pelvic lymph nodes.
 - 77 Posterior exenteration
Includes rectum and rectosigmoid WITH ligamentous attachments and pelvic lymph nodes.
 - 78 Total exenteration
Includes removal of all pelvic contents and pelvic lymph nodes.
 - 79 Extended exenteration
Includes pelvic blood vessels or bony pelvis.

Specimen sent to pathology from surgical events 20–79.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**OVARY
C56.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 17 Local tumor destruction, NOS
No specimen sent to pathology from surgical event 17.
- 25 Total removal of tumor or (single) ovary, NOS
26 Resection of ovary (wedge, subtotal, or partial) ONLY, NOS; unknown if hysterectomy done
27 WITHOUT hysterectomy
28 WITH hysterectomy
- 35 Unilateral (salpingo-)oophorectomy; unknown if hysterectomy done
36 WITHOUT hysterectomy
37 WITH hysterectomy
- 50 Bilateral (salpingo-)oophorectomy; unknown if hysterectomy done
51 WITHOUT hysterectomy
52 WITH hysterectomy
- 55 Unilateral or bilateral (salpingo-)oophorectomy WITH OMENTECTOMY, NOS; partial or total; unknown if hysterectomy done
56 WITHOUT hysterectomy
57 WITH hysterectomy
- 60 Debulking; cytoreductive surgery, NOS
61 WITH colon (including appendix) and/or small intestine resection (not incidental)
62 WITH partial resection of urinary tract (not incidental)
63 Combination of 61 and 62
Debulking is a partial or total removal of the tumor mass and can involve the removal of multiple organ sites. It may include removal of ovaries and/or the uterus (a hysterectomy). The pathology report may or may not identify ovarian tissue. A debulking is usually followed by another treatment modality such as chemotherapy.
- 70 Pelvic exenteration, NOS
71 Anterior exenteration
Includes bladder, distal ureters, and genital organs WITH their ligamentous attachments and pelvic lymph nodes.

72 Posterior exenteration
Includes rectum and rectosigmoid WITH ligamentous attachments and pelvic lymph nodes.

73 Total exenteration
Includes removal of all pelvic contents and pelvic lymph nodes.

74 Extended exenteration
Includes pelvic blood vessels or bony pelvis.
- 80 (Salpingo-)oophorectomy, NOS

Specimen sent to pathology from surgical events 25–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

PROSTATE C61.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Do not code an orchiectomy in this field. For prostate primaries, orchiectomies are coded in the data item *Hematologic Transplant and Endocrine Procedures*.

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 18 Local tumor destruction or excision, NOS
- 19 Transurethral resection (TURP), NOS and no specimen sent to pathology or unknown if sent
Unknown whether a specimen was sent to pathology for surgical events coded 18 or 19 (principally for cases diagnosed prior to January 1, 2003).
- 10 Local tumor destruction, NOS
- 14 Cryoprostatectomy
 - 15 Laser ablation
 - 16 Hyperthermia
 - 17 Other method of local tumor destruction
- No specimen sent to pathology from surgical events 10–17.**
- 20 Local tumor excision, NOS
- 21 Transurethral resection (TURP), NOS, with specimen sent to pathology
 - 22 TURP—cancer is incidental finding during surgery for benign disease
 - 23 TURP—patient has suspected/known cancer
- Any combination of 20–23 WITH
- 24 Cryosurgery
 - 25 Laser
 - 26 Hyperthermia
- 30 Subtotal, segmental, or simple prostatectomy, which may leave all or part of the capsule intact
- 50 Radical prostatectomy, NOS; total prostatectomy, NOS
Excised prostate, prostatic capsule, ejaculatory ducts, seminal vesicle(s) and may include a narrow cuff of bladder neck.
- 70 Prostatectomy WITH resection in continuity with other organs; pelvic exenteration
Surgeries coded 70 are any prostatectomy WITH resection in continuity with any other organs. The other organs may be partially or totally removed. Procedures may include, but are not limited to, cystoprostatectomy, radical cystectomy, and prostatectomy.
- 80 Prostatectomy, NOS

Specimen sent to pathology from surgical events 20–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

TESTIS
C62.0-C62.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 12 Local tumor destruction, NOS
No specimen sent to pathology from surgical event 12.
- 20 Local or partial excision of testicle
- 30 Excision of testicle WITHOUT cord
- 40 Excision of testicle WITH cord/or cord not mentioned (radical orchiectomy)
- 80 Orchiectomy, NOS (unspecified whether partial or total testicle removed)

Specimen sent to pathology from surgical event 20-80.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

KIDNEY, RENAL PELVIS, AND URETER

Kidney C64.9, Renal Pelvis C65.9, Ureter C66.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
 - 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Thermal ablation

No specimen sent to pathology from this surgical event 10–15.
 - 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy

Any combination of 20 or 26–27 WITH

 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation - 25 Laser excision
- 30 Partial or subtotal nephrectomy (kidney or renal pelvis) or partial ureterectomy (ureter)

Procedures coded 30 include, but are not limited to:

Segmental resection; Wedge resection
- 40 Complete/total/simple nephrectomy—for kidney parenchyma
Nephroureterectomy

Includes bladder cuff for renal pelvis or ureter.
- 50 Radical nephrectomy

May include removal of a portion of vena cava, adrenal gland(s), Gerota's fascia, perinephric fat, or partial/total ureter.
- 70 Any nephrectomy (simple, subtotal, complete, partial, simple, total, radical) in continuity with the resection of other organ(s) (colon, bladder)

The other organs, such as colon or bladder, may be partially or totally removed.
- 80 Nephrectomy, NOS; Ureterectomy, NOS

Specimen sent to pathology from surgical events 20–80.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

BLADDER

C67.0-C67.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
 - 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
 - 15 Intravesical therapy
 - 16 Bacillus Calmette-Guerin (BCG) or other immunotherapy

No specimen sent to pathology from surgical events 10–16.

- 20 Local tumor excision, NOS
 - 26 Polypectomy
 - 27 Excisional biopsy
 - Combination of 20 or 26–27 WITH
 - 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
 - 25 Laser excision
- 30 Partial cystectomy
- 50 Simple/total/complete cystectomy
- 60 Complete cystectomy with reconstruction
 - 61 Radical cystectomy PLUS ileal conduit
 - 62 Radical cystectomy PLUS continent reservoir or pouch, NOS
 - 63 Radical cystectomy PLUS abdominal pouch (cutaneous)
 - 64 Radical cystectomy PLUS in situ pouch (orthotopic)

When the procedure is described as a pelvic exenteration for males, but the prostate is not removed, the surgery should be coded as a cystectomy (code 60-64)

- 70 Pelvic exenteration, NOS
 - 71 Radical cystectomy including anterior exenteration

For females, includes removal of bladder, uterus, ovaries, entire vaginal wall, and entire urethra. For males, includes removal of the prostate. When a procedure is described as a pelvic exenteration for males, but the prostate is not removed, the surgery should be coded as a cystectomy (code 60-64).
 - 72 Posterior exenteration

For females, also includes removal of vagina, rectum and anus. For males, also includes

prostate, rectum and anus

73 Total exenteration
Includes all tissue and organs removed for an anterior and posterior exenteration

74 Extended exenteration
Includes pelvic blood vessels or bony pelvis

80 Cystectomy, NOS

Specimen sent to pathology from surgical events 20–80.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

BRAIN

Meninges C70.0-C70.9, Brain C71.0-C71.9, Spinal Cord, Cranial Nerves

and Other Parts of Central Nervous System C72.0-C72.9
(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Do not code laminectomies for spinal cord primaries.

Codes

00 None; no surgery of primary site; autopsy ONLY

10 Tumor destruction, NOS
No specimen sent to pathology from surgical event 10.

Do not record stereotactic radiosurgery (SRS), Gamma knife, Cyber knife, or Linac radiosurgery as surgical tumor destruction. All of these modalities are recorded in the radiation treatment fields.

20 Local excision of tumor, lesion or mass; excisional biopsy

21 Subtotal resection of tumor, lesion or mass in brain

22 Resection of tumor of spinal cord or nerve

30 Radical, total, gross resection of tumor, lesion or mass in brain

40 Partial resection of lobe of brain, when the surgery can not be coded as 20-30.

55 Gross total resection of lobe of brain (lobectomy)

Codes 30 - 55 are not applicable for spinal cord or spinal nerve primary sites.

Specimen sent to pathology from surgical events 20–55.

90 Surgery, NOS

99 Unknown if surgery performed; death certificate ONLY

**THYROID
C73.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 13 Local tumor destruction, NOS
No specimen sent to pathology from surgical event 13.
- 25 Removal of less than a lobe, NOS
 - 26 Local surgical excision
 - 27 Removal of a partial lobe ONLY
- 20 Lobectomy and/or isthmectomy
 - 21 Lobectomy ONLY
 - 22 Isthmectomy ONLY
 - 23 Lobectomy WITH isthmus
- 30 Removal of a lobe and partial removal of the contralateral lobe
- 40 Subtotal or near total thyroidectomy
- 50 Total thyroidectomy
- 80 Thyroidectomy, NOS

Specimen sent to pathology from surgical events 25–80.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

LYMPH NODES C77.0-C77.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Codes

- 00 None; no surgery of primary site; autopsy ONLY
- 19 Local tumor destruction or excision, NOS
Unknown whether a specimen was sent to pathology for surgical events coded to 19 (principally for cases diagnosed prior to January 1, 2003).
- 15 Local tumor destruction, NOS
No specimen sent to pathology from surgical event 15.
- 25 Local tumor excision, NOS
Less than a full chain, includes an excisional biopsy of a single lymph node.
- 30 Lymph node dissection, NOS
 - 31 One chain
 - 32 Two or more chains
- 40 Lymph node dissection, NOS PLUS splenectomy
 - 41 One chain
 - 42 Two or more chains
- 50 Lymph node dissection, NOS and partial/total removal of adjacent organ(s)
 - 51 One chain
 - 52 Two or more chains
- 60 Lymph node dissection, NOS and partial/total removal of adjacent organ(s) PLUS splenectomy.
(Includes staging laparotomy for lymphoma)
 - 61 One chain
 - 62 Two or more chains

Specimen sent to pathology from surgical events 25–62.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

ALL OTHER SITES

C14.2–C14.8, C17.0–C17.9, C23.9, C24.0–C24.9, C26.0–C26.9, C30.0–C 30.1, C31.0–C31.9, C33.9, C37.9, C38.0–C38.8, C39.0–C39.9, C48.0–C48.8, C51.0–C51.9, C52.9,

C57.0–C57.9, C58.9, C60.0–C60.9, C63.0–C63.9, C68.0–C68.9, C69.0–C69.9, C74.0–C74.9, C75.0–C75.9

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE**Codes**

- 00 None; no surgery of primary site; autopsy ONLY
- 10 Local tumor destruction, NOS
- 11 Photodynamic therapy (PDT)
 - 12 Electrocautery; fulguration (includes use of hot forceps for tumor destruction)
 - 13 Cryosurgery
 - 14 Laser
- No specimen sent to pathology from surgical events 10–14.**
- 20 Local tumor excision, NOS
- 26 Polypectomy
 - 27 Excisional biopsy
- Any combination of 20 or 26–27 WITH
- 21 Photodynamic therapy (PDT)
 - 22 Electrocautery
 - 23 Cryosurgery
 - 24 Laser ablation
- 25 Laser excision
- 30 Simple/partial surgical removal of primary site
- 40 Total surgical removal of primary site; enucleation
- 41 Total enucleation (for eye surgery only)
- 50 Surgery stated to be “debulking”
- 60 Radical surgery
- Partial or total removal of the primary site WITH a resection in continuity (partial or total removal) with other organs.**

Specimen sent to pathology from surgical events 20–60.

- 90 Surgery, NOS
- 99 Unknown if surgery performed; death certificate ONLY

**UNKNOWN AND ILL-DEFINED PRIMARY SITES
C76.0-C76.8, C80.9**

(Except for M-9727, 9733, 9741-9742, 9764-9809, 9832, 9840-9931, 9945-9946, 9950-9967, and 9975-9992)

SURGERY OF PRIMARY SITE

Code

98 All unknown and ill-defined disease sites, WITH or WITHOUT surgical treatment.

Surgical procedures for unknown and ill-defined primaries are to be recorded using the data item *Surgical Procedure/Other Site*.

**APPENDIX L:
ABBREVIATIONS AND SYMBOLS**

ABBREVIATIONS

The PCR requires all cases to include text information to support specific coded fields. Complete and descriptive text is vital to the quality control efforts of the PCR. Often it is necessary to use abbreviations to provide adequate descriptions within the limited size of the text fields. However, a reader may interpret many standard medical abbreviations differently. The PCR will rely on the attached abbreviation list to indicate how PCR staff will interpret the abbreviation when its use is unclear. It is a combination of the North American Association of Central Cancer Registries (NAACCR)'s *Standards for Cancer Registries, Volume II: Data Standards and Data Dictionary, Tenth Edition; Version 11 -- Appendix G: Recommended Abbreviations for Abstractors* and the abbreviation list provided in previous PCR Manuals.

The Abbreviations Listings consist of two main lists word/terms and their recommended abbreviations, as well as a special table delineating context-sensitive abbreviations and one for symbols. The first main listing is ordered by word/term to enable the look-up of a recommended abbreviation for a particular word or term, and the second main listing is ordered by abbreviation to enable the look-up of the word or term for a particular abbreviation. The context-sensitive abbreviations list consists of a subset of the abbreviations from the main lists where a different context for the same abbreviation conveys a different meaning (for example, CA may mean calcium or carcinoma/ML may mean milliliter or middle lobe). For these context-sensitive abbreviations, the meaning of the abbreviation should be readily apparent from the context in which it is used.

The listings are not exhaustive, but many of the most commonly used terms were included. Abbreviations for chemotherapy drugs and/or regimens are not included. For short names and acronyms of antineoplastic drugs, consult the SEER Program *Self Instructional Manual for Tumor Registrars: Book 8-Antineoplastic Drugs, Third Edition* or SEER RX at <http://seer.cancer.gov/tools/seerrx/>.

Please note that although abbreviations are presented in uppercase, either upper- or lowercase may be utilized when entering abbreviations within abstraction software. When abstracting into text fields, the use of abbreviations should be limited to those that appear on these lists whenever practical. Abbreviations and symbols should be used carefully.

The abbreviations list does not include an abbreviation for the word **cancer**. While the abbreviation "CA" is often used in the medical record to mean either the term **cancer** or **carcinoma**, it should be used in text reported to the PCR to indicate the histologic term of carcinoma. This distinction is very important when verifying histologic coding for cancer, NOS (8000/3) and carcinoma, NOS (8010/3).

This appendix contains two tables for abbreviations, one in term order and one in abbreviation order.

ABBREVIATIONS**ORDERED BY WORD/TERM**

WORD/TERM (S)	ABBREVIATION/SYMBOL
Abdomen (abdominal)	ABD
Abdominal hysterectomy	ABD HYST
Abdominal perineal (Abdominoperineal)	AP
Abdominoperineal resection	APR
Abnormal	ABN
Abnormal liver function test	ALFT
Above	^
Above knee (amputation)	AK(A)
Absent/Absence	ABS
Abstract/Abstracted	ABST
Achilles tendon reflex	ATR
Acid phosphatase	ACID PHOS
Acquired Immune Deficiency Syndrome	AIDS
Acral lentiginous melanoma	ALM
Activities of daily living	ADL
Acute erythroleukemia	AEL
Acute granulocytic leukemia	AGL
Acute leukemia	AL
Acute lymphocytic leukemia	ALL
Acute megakaryoblastic leukemia	AMEGL
Acute myeloblastic leukemia	AMBL
Acute myelogenous leukemia	AML
Acute myelomonocytic leukemia	AMML
Acute myocardial infarction	AMI
Acute promyelocytic leukemia	APL
Acute renal failure	ARF
Acute Respiratory Distress (Disease) Syndrome	ARDS
Acute tubular necrosis	ATN
Acute undifferentiated leukemia	AUL
Adenocarcinoma	ADENOCA, ACA
Adenosine triphosphate	ATP
Adjacent	ADJ
Admission/Admit	ADM
Adrenal cortex	AC
Adrenal cortical hormone	ACH
Adrenocorticotrophic hormone	ACTH

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Adult T-cell leukemia	ATL
Adult T-cell leukemia/lymphoma	ATLL
Adult-onset Diabetes Mellitus	AODM
Affirmative	AFF
Against medical advice	AMA
AIDS-related condition (complex)	ARC
AIDS-related disease	ARD
Air contrast barium enema	ACBE
Albumin	ALB
Alcohol	ETOH
Alkaline phosphatase	ALK PHOS
Alpha chain disease	ACD
Alpha-fetoprotein	AFP
Also known as	AKA
Alternate	ALT
Ambulatory	AMB
Amount	AMT
Amputation	AMP
Amyotrophic lateral sclerosis	ALS
Anal intraepithelial neoplasia, grade III	AIN III
Anaplastic	ANAP
And	&
Angioblastic immunoblastic lymphadenopathy	AIL
Angiography/Angiogram	ANGIO
Anterior	ANT
Anteroposterior	AP
Antidiuretic hormone	ADH
Antigen	AG
Aortic stenosis	A-STEN
Apparently	APPL'Y
Appendix	APP
Approximately	APPROX
Arrhythmia	ARRHY
Arterial blood gases	ABG
Arteriosclerosis/Arteriosclerotic	AS
Arteriosclerotic cardiovascular disease	ASCVD
Arteriosclerotic heart disease	ASHD
Arteriosclerotic Peripheral Vascular Disease	ASPVD

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Arteriovenous	AV
Arteriovenous malformation	AVM
Artery (ial)	ART
As soon as possible	ASAP
Ascending	ASC
Ascending colon	A-COLON
Aspiration	ASP
Aspiration biopsy cytology	ABC
Aspirin, Acetylsalicylic acid	ASA
At	@
Atrial fibrillation	A FIB
Atrial flutter	A FLUTTER
Atrial premature complexes	APC
Atrial stenosis/insufficiency/incompetence	AI
Auscultation & percussion	A&P
Autoimmune hemolytic anemia	AIHA
Autologous bone marrow	ABM
Autologous bone marrow transplantation	ABMT
Autonomic nervous system	ANS
Autopsy	AUT
Average	AVG
Axilla(ry)	AX
Bacillus Calmette-Guerin	BCG
Barium	BA
Barium enema	BE
Barium swallow	BAS
Bartholin's, Urethral & Skene's	BUS
Basal cell carcinoma	BCC
Before noon	AM
Below knee (amputation)	BK(A)
Benign prostatic hypertrophy/hyperplasia	BPH
Bilateral	BIL
Bilateral hilar lymphadenopathy	BHL
Bilateral lower lobes	BLL
Bilateral pelvic lymph node dissection	BPLND
Bilateral salpingo-oophorectomy	BSO
Bile duct	BD
Biological response modifier	BRM

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Biopsy	BX
Bipolar affective disorder	BAD
Black female	B/F
Black male	B/M
Bladder outlet obstruction	BOO
Bladder tumor	BT
Blood pressure	BP
Blood urea nitrogen	BUN
Blood volume	BV
Bone Marrow	BM
Bone marrow aspirate	BMA
Bone marrow biopsy	BMBX
Bone Marrow Transplant	BMT
Bowel Movement	BM
Bowel sounds	BS
Breast self examination	BSE
Breath sounds	BRS
Bright red blood	BRB
Bright red blood per rectum	BRBPR
Bronchial lymph node	BLN
Bronchoalveolar washing	BAW
Bronchogenic carcinoma	BGCA
Burkitt lymphoma	BL
Calcium	CA
Capsule (s)	CAP(S)
Carcinoembryonic antigen	CEA
Carcinoma	CA
Carcinoma <i>in situ</i>	CIS
Carcinoma unknown primary	CUP
Cardioesophageal junction	CEJ
Cardiovascular disease	CVD
CAT/CT scan/Computerized axial tomography	CT
Ceased to breath	CTB
Centigram	CGM
Centigray	CGY
Centimeter	CM
Central nervous system	CNS
Cerebrospinal fluid	CSF

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Cerebrovascular accident	CVA
Cervical intraepithelial neoplasia	CIN
Cervical intraepithelial neoplasia, grade III	CIN III
Cervical spine	C-SPINE
Cervical vertebrae	C1-C7
Cervix	CX
Change	CHG
Chemotherapy	CHEMO
Chest X-ray	CXR
Chief complaint	C/C
Cholecystectomy	CHOLE
Chronic	CHR
Chronic granulocytic leukemia	CGL
Chronic leukemia	CL
Chronic lymphocytic leukemia	CLL
Chronic lymphosarcoma leukemia	CLSL
Chronic myelodysplastic syndrome	CMS
Chronic myeloid (myelocytic) leukemia	CML
Chronic myelomonocytic leukemia	CMML
Chronic obstructive lung disease	COLD
Chronic obstructive pulmonary disease	COPD
Chronic renal failure	CRF
Chronic ulcerative colitis	CUC
Cigarettes	CIG
Clear	CLR
Clinical tumor, nodes, metastases	CTNM
Cobalt 60	CO60
Collaborative stage	CS
Colon, Ascending	A-COLON
Colon, Descending	D-COLON
Colon, Sigmoid	SIG-COLON
Colon, Transverse	TRANS-COLON
Colony-stimulating factor	C-SF
Common bile duct	CBD
Complaint (-ning) of	C/O
Complete blood count	CBC
Complete continuous remission	CCR
Computerized axial tomography scan	CT, CAT

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Congenital heart disease	CHD
Congestive heart failure	CHF
Consistent with	C/W
Continue/continuous	CONT
Contralateral	CONTRA
Coronary artery bypass graft	CABG
Coronary artery disease	CAD
Coronary care unit	CCU
Cubic centimeter	CC
Curie	CU
Cutaneous	CUT
Cutaneous T-cell lymphoma	CTCL
Cystic fibrosis	CF
Cystoscopy	CYSTO
Cytology	CYTO
Date of birth	DOB
Date of death	DOD
Dead on arrival	DOA
Debridement	DEB
Decrease(d)	DECR
Deep tendon reflex	DTR
Deep vein thrombosis	DVT
Deoxyribonucleic acid	DNA
Dermatofibrosarcoma protuberans	DFSP
Dermatology	DERM
Descending	DESC
Descending colon	D-COLON
Diabetes mellitus	DM
Diagnosis	DX
Diagnostic laparoscopy	DL
Diameter	DIAM
Died of other causes	DOC
Died with disease	DWD
Diethylstilbestrol	DES
Differentiated/differential	DIFF
Digital rectal examination	DRE
Dilatation and curettage	D&C
Direct extension	DE

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Discharge	DISCH
Discontinue(d)	DC
Disease	DZ
Disease free interval	DFI
Disseminated	DISSEM
Disseminated intravascular coagulopathy	DIC
Distant metastases	DM
Doctor	DR
Ductal carcinoma <i>in situ</i>	DCIS
Dyspnea on exertion	DOE
Ears, nose, and throat	ENT
Electrocardiogram	ECG/EKG
Electroencephalogram	EEG
Electromyogram	EMG
Emergency room	ER
Endoscopic retrograde cholangiopancreatography	ERCP
Enlarged	ENLGD
Equal(s)	=
Esophagogastro-duodenoscopy	EGD
Esophagus	ESO
Estrogen receptor assay	ERA
Evaluation	EVAL
Every	Q
Every day	QD
Examination	EXAM
Examination under anesthesia	EUA
Excision/excised	EXC(D)
Expired	EXP
Exploratory	EXPL
Exploratory laparotomy	EXPL LAP
Extend/extension	EXT
Extended care facility	ECF
External	EX
Extremity	EXTR
Eyes, ears, nose and throat	EENT
Family history	FHX
Family medical history	FMH
Fever of unknown origin	FUO

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Fine needle aspiration	FNA
Fine needle aspiration biopsy	FNAB
Fingerbreadth	FB
Flexible sigmoidoscopy	FLEX SIG
Floor of mouth	FOM
Fluid	FL
Fluoroscopy	FLURO
Follow-up	FU
For example	E.G
Fracture	FX
French-American-British	FAB
Frequent/Frequency	FREQ
Frozen section	FS
Full thickness skin graft	FTSG
Gallbladder	GB
Gastroesophageal	GE
Gastroesophageal reflux disease	GERD
Gastrointestinal	GI
General/Generalized	GEN
Genitourinary	GU
Grade	GR
Gram	GM
Greater/Greater than	>
Gynecology	GYN
Head, eyes, ears, nose, throat	HEENT
Hematocrit	HCT
Hematology	HEMO
Hemoglobin	HGB
Hepatitis A (virus)	HAV
Hepatitis B (virus)	HBV
Hepatitis C (virus)	HCV
Hepatitis D (virus)	HDV
Hepatocellular carcinoma	HCC
Hepatosplenomegaly	HSM
History	HX
History and physical	H&P
History of	H/O
History of present illness	HPI

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Hodgkin disease	HD
Hormone	HORM
Hospital	HOSP
Hour/Hours	HR(S)
Human chorionic gonadotropin	HCG
Human Immunodeficiency Virus	HIV
Human Papilloma Virus	HPV
Human T-Lymphotropic Virus, (Type III)	HTLV
Hypertension	HTN
Hypertensive cardiovascular disease	HCVD
Hypertensive vascular disease	HVD
Hysterectomy	HYST
Idiopathic hypertrophic subaortic stenosis	IHSS
Idiopathic thrombocytopenia	ITP
Immunoglobulin	IG
Immunohistochemical	IHC
Impression	IMP
Inch	IN
Incision & drainage	I&D
Includes/Including	INCL
Increase(d)	INCR
Inferior	INF
Inferior vena cava	IVC
Infiltrating	INFILT
Inflammatory bowel disease	IBD
Inpatient	IP
Insulin-dependent diabetes mellitus	IDDM
Intensive care unit	ICU
Intercostal margin	ICM
Intercostal space	ICS
Intermittent positive pressure breathing	IPPB
Internal	INT
Internal mammary artery	IMA
Interstitial lung disease	ILD
Intra abdominal	IAB
Intramuscular	IM
Intrathecal	IT
Intravenous	IV

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Intravenous cholangiogram	IVCA
Intravenous pyelogram	IVP
Invade(s)/invading/invasion	INV
Involve(s)/involvement/involving	INVL
Iodine	I
Ipsilateral	IPSI
Irregular	IRREG
Joule	J
Jugular venous distention	JVD
Junction	JCT, JX
Juvenile rheumatic arthritis	JRA
Kaposi sarcoma	KS
Kidneys, ureters, bladder	KUB
Kilogram	KG
Kilovolt	KV
Laboratory	LAB
Lactic dehydrogenase	LDH
Laparotomy	LAP
Large	LRG
Large bowel resection	LBR
Large cleaved cell	LCC
Last menstrual period	LMP
Lateral	LAT
Left	LT
Left breast biopsy	LBBX
Left bundle branch block	LBBB
Left costal margin	LCM
Left eye	OS
Left lower extremity	LLE
Left lower lobe	LLL
Left lower quadrant	LLQ
Left salpingo-oophorectomy	LSO
Left upper extremity	LUE
Left upper lobe	LUL
Left upper outer quadrant	LUOQ
Left upper quadrant	LUQ
Left ureteral orifice	LUO
Less/Less than	<

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Licensed practical nurse	LPN
Linear accelerator	LINAC
Liver, kidney, spleen	LKS
Liver, kidney, spleen, bladder	LKSB
Liver/spleen scan	LS SCAN
Lobular carcinoma in situ	LCIS
Lobular in situ	LIS
Lobular neoplasia, grade 2	LN2
Long Term Care Facility	LTCF
Lower extremity	LE
Lower inner quadrant	LIQ
Lower outer quadrant	LOQ
Lower right quadrant	LRQ
Lumbar puncture	LP
Lumbar spine	L-SPINE
Lumbar vertebra	L1-L5
Lumbosacral	LS
Lupus erythematosus	LUP ERYTH
Lymph node biopsy	LNBX
Lymph node dissection	LND
Lymph node resection	LNR
Lymph node(s)	LN(S)
Lymphadenopathy-associated virus	LAV
Lymphangiography/lymphangiogram	LAG
Macrophage colony-stimulating factor	M-CSF
Magnetic resonance cholangiopancreatography	MRCP
Magnetic resonance imaging	MRI
Main stem bronchus	MSB
Malignant	MALIG
Malignant carcinoid syndrome	MCS
Malignant fibrous histiocytoma	MFH
Mandible/mandibular	MAND
Mastectomy	MAST, MX
Maximum	MAX
Medical center	MC
Medical history	MHX
Medication	MED
Melanoma associated antigen	MAA

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Metastatic/Metastasis	METS
Methicillin Resistant Staphylococcus Aureus	MRSA
Microgram	MCG
Microscopic	MICRO
Midclavicular line	MCL
Middle	MID
Middle lobe	ML
Millicurie (hours)	MC(H)
Milligram (hours)	MG(H)
Milliliter	ML
Millimeter	MM
Million electron volts	MEV
Minimum	MIN
Minus	-
Minute	MIN
Mitral valve prolapse	MVP
Mixed combined immunodeficiency	MCID
Mixed connective tissue disease	MCTD
Moderate (ly)	MOD
Moderately differentiated	MD, MOD DIFF
Modified radical mastectomy	MRM
Monoclonal antibody	MC-AB, MCAB, MAB, MOAB
More/More than	>
Multifocal arterial tachycardia	MAT
Multifocal premature ventricular contraction	MPVC
Multiple	MULT
Multiple myeloma	MM
Multiple sclerosis	MS
Myasthenia gravis	MG
Myelodysplasia/myelodysplastic syndrome	MDS
Myeloproliferative disease	MPD
Myocardial infarction	MI
Natural killer	NK
Nausea and vomiting	N&V
Neck vein distention	NVD
Needle biopsy	NBX
Needle liver biopsy	NLBX
Negative	NEG, -

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Neoplasm	NEOPL
Neoplasm embryonic antigen	NEA
Nephrectomy	NX
Nerves, Cranial 1-12	N-I - N-XII
Neurology	NEURO
No acute/active disease	NAD
No evidence of disease	NED
No evidence of recurrence	NER
No significant findings	NSF
Nodular & diffuse lymphoma	NDL
Non small cell carcinoma	NSCCA
Non-Hodgkin malignant lymphoma	NHML
Non-Hodgkins lymphoma	NHL
Non-small cell lung cancer	NSCLC
Normal	NL
Not applicable	NA
Not elsewhere classified/classifiable	NEC
Not otherwise specified	NOS
Not recorded	NR
Number	#
Nursing home	NH
Obstetrics	OB
Obstructed (-ing, -ion)	OBST
Occult primary malignancy	OPM
Oncology	ONC
Operating room	OR
Operation	OP
Operative report	OP RPT
Organic brain syndrome	OBS
Orthopedics	ORTHO
Otology	OTO
Ounce	OZ
Outpatient	OP
Outpatient surgery	OPS
Packs per day	PPD
Palpated (-able)	PALP
Papanicolaou smear	PAP
Papillary	PAP

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Past/personal (medical) history	PMH
Pathologic tumor, nodes, metastases	PTNM
Pathology	PATH
Patient	PT
Pediatrics	PEDS
Pelvic inflammatory disease	PID
Peptic ulcer disease	PUD
Percussion and auscultation	P&A
Percutaneous	PERC
Percutaneous transhepatic cholecystogram	PTC
Peripheral vascular disease	PVD
Phosphorus 32	P32
Physical examination	PE
Physiotherapy/Physical therapy	PT
Plasma cell leukemia	PCL
Platelets	PLT
Plus	+
Polycythemia vera	PCV
Poorly differentiated	PD, POOR DIFF
Positive	POS, +
Positron emission tomography	PET
Possible	POSS
Posterior	POST
Posteroanterior	PA
Postoperative (-ly)	POST OP
Pound(s)	LB(S), #
Premature atrial contraction	PAC
Preoperative (-ly)	PRE OP
Prescription	RX
Present illness	PI
Previous	PREV
Primary medical physician	PMP
Primitive neuroectodermal tumor	PNET
Prior to admission	PTA
Probable (-ly)	PROB
Proctoscopy	PROCTO
Progesterone receptor assay	PRA
Prolymphocytic leukemia	PLL

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Prostatic intraepithelial neoplasia	PIN
Prostatic intraepithelial neoplasia, grade III	PIN III
Prostatic specific antigen	PSA
Pulmonary	PULM
Pulmonary artery	PULM ART
Quadrant	QUAD
Radiation absorbed dose	RAD
Radiation therapy	RT
Radical neck dissection	RND
Radioactive iodine	RAI
Radioimmunoassay	RIA
Received	REC'D
Red blood cells (count)	RBC
Regarding	RE
Regional medical center	RMC
Regular	REG
Regular sinus rhythm	RSR
Resection (ed)	RESEC
Respiratory	RESPIR, RESP
Review of outside films	ROF
Review of outside slides	ROS
Rheumatic heart disease	RHD
Rheumatoid arthritis	RA
Right	RT
Right breast biopsy	RBBX
Right bundle branch block	RBBB
Right costal margin	RCM
Right eye	OD
Right inner quadrant	RIQ
Right lower extremity	RLE
Right lower lobe	RLL
Right lower quadrant	RLQ
Right middle lobe	RML
Right outer quadrant	ROQ
Right salpingo-oophorectomy	RSO
Right upper extremity	RUE
Right upper lobe	RUL
Right upper quadrant	RUQ

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Right ureteral orifice	RUO
Rule out	R/O
Sacral spine	S-SPINE
Sacral vertebra	S1-S5
Salpingo-oophorectomy	SO
Sarcoma	SARC
Satisfactory	SATIS
Sequential multiple analysis	SMA
Serum glutamic oxaloacetic transaminase	SGOT
Serum glutamic pyruvic transaminase	SGPT
Severe combined immunodeficiency syndrome	SCID
Short(ness) of breath	SOB
Sick sinus syndrome	SSS
Sigmoid colon	SIG COLON
Skilled nursing facility	SNF
Small	SM
Small bowel	SB
Small bowel obstruction	SBO
Small bowel resection	SBR
Small cell lung carcinoma	SCLC
Specimen	SPEC
Spine, Cervical	C-SPINE
Spine, Lumbar	L-SPINE
Spine, Sacral	S-SPINE
Spine, Thoracic	T-SPINE
Split thickness skin graft	STSG
Squamous	SQ
Squamous cell carcinoma	SCC
Status post	S/P
Subcutaneous	SUBCU
Summary stage	SS
Superior vena cava	SVC
Surgery/Surgical	SURG
Suspicious/suspected	SUSP
Symptoms	SX
Syndrome of inappropriate	ADH SIADH
Systemic lupus erythematosus	SLE
T-cell acute lymphoblastic leukemia	T-ALL

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
T-cell chronic lymphatic leukemia	T-CLL
Thoracic spine	T-SPINE
Thromboticthrombocytopenia purpura	TTP
Times	X
Total abdominal hysterectomy	TAH
Total abdominal hysterectomy- bilateral salpingo-oophorectomy	TAH-BSO
Total axial (lymph) node irradiation	TANI
Total parenteral nutrition	TPN
Total vaginal hysterectomy	TVH
Transbronchial biopsy	TBBX
Transient ischemic attack	TIA
Transitional cell carcinoma	TCC
Transrectal ultrasound	TRUS
Transrectal ultrasound of prostate	TRUSP
Transurethral resection	TUR
Transurethral resection bladder	TURB
Transurethral resection bladder tumor	TURBT
Transurethral resection prostate	TURP
Transverse colon	TRANS-COLON
Transverse rectus abdominous myocutaneous	TRAM
Treatment	TX
True vocal cord	TVC
Tumor size	TS
Tumor, node, metastasis	TNM
Twice a day (daily)	BID
Ultrasound	US
Undetermined	UNDET
Undetermined origin	UDO
Undifferentiated	UNDIFF
Unilateral salpingo-oophorectomy	USO
Unknown	UNK
Upper extremity	UE
Upper gastrointestinal (series)	UGI
Upper inner quadrant	UIQ
Upper outer quadrant	UOQ
Upper respiratory infection	URI
Upper right quadrant	URQ
Urinary tract infection	UTI

ABBREVIATIONS

WORD/TERM (S)	ABBREVIATION/SYMBOL
Vagina/Vaginal	VAG
Vaginal hysterectomy	VAG HYST
Vaginal intraepithelial neoplasia	VAIN
Vaginal intraepithelial neoplasia (grade III)	VAIN III
Vascular	VASC
Versus	VS
Vulvar intraepithelial neoplasia	VIN
Vulvar intraepithelial neoplasia (grade III)	VIN III
Well differentiated	WD, WELL DIFF
White blood cells (count)	WBC
White female	W/F
White male	W/M
Will follow (in) office	WF-O
Wilms (tumor), aniridia, genitourinary (abnormalities), and (mental)	WAGR
With	W/
Within normal limits	WNL
Without	W/O
Work-up	W/U
Xray	XR
Year	YR
Yolk Sac Tumor	YST

ABBREVIATIONS**ORDER BY ABBREVIATION**

ABBREVIATION	WORD/TERM(S)
A FIB	Atrial fibrillation
A FLUTTER	Atrial flutter
A&P	Auscultation & percussion
ABC	Aspiration biopsy cytology
ABD	Abdomen (abdominal)
ABD HYST	Abdominal hysterectomy
ABG	Arterial blood gases
ABM	Autologous bone marrow
ABMT	Autologous bone marrow transplantation
ABN	Abnormal
ABS	Absent/Absence
ABST	Abstract/Abstracted
AC	Adrenal cortex
ACA	Adenocarcinoma
ACBE	Air contrast barium enema
ACD	Alpha chain disease
ACH	Adrenal cortical hormone
ACID PHOS	Acid phosphatase
A-COLON	Ascending colon
ACTH	Adrenocorticotrophic hormone
ADENOCA, ACA	Adenocarcinoma
ADH	Antidiuretic hormone
ADH SIADH	Syndrome of inappropriate
ADJ	Adjacent
ADL	Activities of daily living
ADM	Admission/Admit
AEL	Acute erythroleukemia
AFF	Affirmative
AFP	Alpha-fetoprotein
AG	Antigen
AGL	Acute granulocytic leukemia
AI	Atrial stenosis/insufficiency/incompetence
AIDS	Acquired Immune Deficiency Syndrome
AIHA	Autoimmune hemolytic anemia
AIL	Angioblastic immunoblastic lymphadenopathy
AIN III	Anal intraepithelial neoplasia, grade III

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
AK(A)	Above knee (amputation)
AKA	Also known as
AL	Acute leukemia
ALB	Albumin
ALFT	Abnormal liver function test
ALK PHOS	Alkaline phosphatase
ALL	Acute lymphocytic leukemia
ALM	Acral lentiginous melanoma
ALS	Amyotrophic lateral sclerosis
ALT	Alternate
AM	Before noon
AMA	Against medical advice
AMB	Ambulatory
AMBL	Acute myeloblastic leukemia
AMEGL	Acute megakaryoblastic leukemia
AMI	Acute myocardial infarction
AML	Acute myelogenous leukemia
AMML	Acute myelomonocytic leukemia
AMP	Amputation
AMT	Amount
ANAP	Anaplastic
ANGIO	Angiography/Angiogram
ANS	Autonomic nervous system
ANT	Anterior
AODM	Adult-onset Diabetes Mellitus
AP	Abdominal perineal (Abdominoperineal)
AP	Anteroposterior
APC	Atrial premature complexes
APL	Acute promyelocytic leukemia
APP	Appendix
APPL'Y	Apparently
APPROX	Approximately
APR	Abdominoperineal resection
ARC	AIDS-related condition (complex)
ARD	AIDS-related disease
ARDS	Acute Respiratory Distress (Disease) Syndrome
ARF	Acute renal failure
ARRHY	Arrhythmia

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
ART	Artery (ial)
AS	Arteriosclerosis/Arteriosclerotic
ASA	Aspirin, Acetylsalicylic acid
ASAP	As soon as possible
ASC	Ascending
ASCVD	Arteriosclerotic cardiovascular disease
ASHD	Arteriosclerotic heart disease
ASP	Aspiration
ASPVD	Arteriosclerotic Peripheral Vascular Disease
A-STEN	Aortic stenosis
ATL	Adult T-cell leukemia
ATLL	Adult T-cell leukemia/lymphoma
ATN	Acute tubular necrosis
ATP	Adenosine triphosphate
ATR	Achilles tendon reflex
AUL	Acute undifferentiated leukemia
AUT	Autopsy
AV	Arteriovenous
AVG	Average
AVM	Arteriovenous malformation
AX	Axilla(ry)
B/F	Black female
B/M	Black male
BA	Barium
BAD	Bipolar affective disorder
BAS	Barium swallow
BAW	Bronchoalveolar washing
BCC	Basal cell carcinoma
BCG	Bacillus Calmette-Guerin
BD	Bile duct
BE	Barium enema
BGCA	Bronchogenic carcinoma
BHL	Bilateral hilar lymphadenopathy
BID	Twice a day (daily)
BIL	Bilateral
BK(A)	Below knee (amputation)
BKA	Below knee amputation
BL	Burkitt lymphoma

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
BLL	Bilateral lower lobes
BLN	Bronchial lymph node
BM	Bone Marrow
BM	Bowel Movement
BMA	Bone marrow aspirate
BMBX	Bone marrow biopsy
BMT	Bone Marrow Transplant
BOO	Bladder outlet obstruction
BP	Blood pressure
BPH	Benign prostatic hypertrophy/hyperplasia
BPLND	Bilateral pelvic lymph node dissection
BRB	Bright red blood
BRBPR	Bright red blood per rectum
BRM	Biological response modifier
BRS	Breath sounds
BS	Bowel sounds
BSE	Breast self examination
BSO	Bilateral salpingo-oophorectomy
BT	Bladder tumor
BUN	Blood urea nitrogen
BUS	Bartholin's, Urethral & Skene's
BV	Blood volume
BX	Biopsy
C/C	Chief complaint
C/O	Complaint (-ning) of
C/W	Consistent with
C1-C7	Cervical vertebrae
CA	Calcium
CA	Carcinoma
CABG	Coronary artery bypass graft
CAD	Coronary artery disease
CAP(S)	Capsule (s)
CBC	Complete blood count
CBD	Common bile duct
CC	Cubic centimeter
CCR	Complete continuous remission
CCU	Coronary care unit
CEA	Carcinoembryonic antigen

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
CEJ	Cardioesophageal junction
CF	Cystic fibrosis
CGL	Chronic granulocytic leukemia
CGM	Centigram
CGY	Centigray
CHD	Congenital heart disease
CHEMO	Chemotherapy
CHF	Congestive heart failure
CHG	Change
CHOLE	Cholecystectomy
CHR	Chronic
CIG	Cigarettes
CIN	Cervical intraepithelial neoplasia
CIN III	Cervical intraepithelial neoplasia, grade III
CIS	Carcinoma <i>in situ</i>
CL	Chronic leukemia
CLL	Chronic lymphocytic leukemia
CLR	Clear
CLSL	Chronic lymphosarcoma leukemia
CM	Centimeter
CML	Chronic myeloid (myelocytic) leukemia
CMML	Chronic myelomonocytic leukemia
CMS	Chronic myelodysplastic syndrome
CNS	Central nervous system
CO60	Cobalt 60
COLD	Chronic obstructive lung disease
CONT	Continue/continuous
CONTRA	Contralateral
COPD	Chronic obstructive pulmonary disease
CRF	Chronic renal failure
CS	Collaborative stage
CSF	Cerebrospinal fluid
C-SF	Colony-stimulating factor
C-SPINE	Cervical spine
CT	CAT/CT scan/Computerized axial tomography
CT, CAT	Computerized axial tomography scan
CTB	Ceased to breath
CTCL	Cutaneous T-cell lymphoma

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
CTNM	Clinical tumor, nodes, metastases
CU	Curie
CUC	Chronic ulcerative colitis
CUP	Carcinoma unknown primary
CUT	Cutaneous
CVA	Cerebrovascular accident
CVD	Cardiovascular disease
CX	Cervix
CXR	Chest X-ray
CYSTO	Cystoscopy
CYTO	Cytology
D&C	Dilatation and curettage
DC	Discontinue(d)
DCIS	Ductal carcinoma <i>in situ</i>
D-COLON	Descending colon
DE	Direct extension
DEB	Debridement
DECR	Decrease(d)
DERM	Dermatology
DES	Diethylstilbestrol
DESC	Descending
DFI	Disease free interval
DFSP	Dermatofibrosarcoma protuberans
DIAM	Diameter
DIC	Disseminated intravascular coagulopathy
DIFF	Differentiated/differential
DISCH	Discharge
DISSEM	Disseminated
DL	Diagnostic laparoscopy
DM	Diabetes mellitus
DM	Distant metastases
DNA	Deoxyribonucleic acid
DOA	Dead on arrival
DOB	Date of birth
DOC	Died of other causes
DOD	Date of death
DOE	Dyspnea on exertion
DR	Doctor

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
DRE	Digital rectal examination
DTR	Deep tendon reflex
DVT	Deep vein thrombosis
DWD	Died with disease
DX	Diagnosis
DZ	Disease
E.G	For example
ECF	Extended care facility
ECG/EKG	Electrocardiogram
EEG	Electroencephalogram
EENT	Eyes, ears, nose and throat
EGD	Esophagogastro-duodenoscopy
EMG	Electromyogram
ENLGD	Enlarged
ENT	Ears, nose, and throat
ER	Emergency room
ERA	Estrogen receptor assay
ERCP	Endoscopic retrograde cholangiopancreatography
ESO	Esophagus
ETOH	Alcohol
EUA	Examination under anesthesia
EVAL	Evaluation
EX	External
EXAM	Examination
EXC(D)	Excision/excised
EXP	Expired
EXPL	Exploratory
EXPL LAP	Exploratory laparotomy
EXT	Extend/extension
EXTR	Extremity
FAB	French-American-British
FB	Fingerbreadth
FHX	Family history
FL	Fluid
FLEX SIG	Flexible sigmoidoscopy
FLURO	Fluoroscopy
FMH	Family medical history
FNA	Fine needle aspiration

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
FNAB	Fine needle aspiration biopsy
FOM	Floor of mouth
FREQ	Frequent/Frequency
FS	Frozen section
FTSG	Full thickness skin graft
FU	Follow-up
FUO	Fever of unknown origin
FX	Fracture
GB	Gallbladder
GE	Gastroesophageal
GEN	General/Generalized
GERD	Gastroesophageal reflux disease
GI	Gastrointestinal
GM	Gram
GR	Grade
GU	Genitourinary
GYN	Gynecology
H&P	History and physical
H/O	History of
HAV	Hepatitis A (virus)
HBV	Hepatitis B (virus)
HCC	Hepatocellular carcinoma
HCG	Human chorionic gonadotropin
HCT	Hematocrit
HCV	Hepatitis C (virus)
HCVD	Hypertensive cardiovascular disease
HD	Hodgkin disease
HDV	Hepatitis D (virus)
HEENT	Head, eyes, ears, nose, throat
HEMO	Hematology
HGB	Hemoglobin
HIV	Human Immunodeficiency Virus
HORM	Hormone
HOSP	Hospital
HPI	History of present illness
HPV	Human Papilloma Virus
HR(S)	Hour/Hours
HSM	Hepatosplenomegaly

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
HTLV	Human T-Lymphotropic Virus, (Type III)
HTN	Hypertension
HVD	Hypertensive vascular disease
HX	History
HYST	Hysterectomy
I	Iodine
I&D	Incision & drainage
IAB	Intra abdominal
IBD	Inflammatory bowel disease
ICM	Intercostal margin
ICS	Intercostal space
ICU	Intensive care unit
IDDM	Insulin-dependent diabetes mellitus
IG	Immunoglobulin
IHC	Immunohistochemical
IHSS	Idiopathic hypertrophic subaortic stenosis
ILD	Interstitial lung disease
IM	Intramuscular
IMA	Internal mammary artery
IMP	Impression
IN	Inch
INCL	Includes/Including
INCR	Increase(d)
INF	Inferior
INFILT	Infiltrating
INT	Internal
INV	Invade(s)/invading/invasion
INVL	Involve(s)/involvement/involving
IP	Inpatient
IPPB	Intermittent positive pressure breathing
IPSI	Ipsilateral
IRREG	Irregular
IT	Intrathecal
ITP	Idiopathic thrombocytopenia
IV	Intravenous
IVC	Inferior vena cava
IVCA	Intravenous cholangiogram
IVP	Intravenous pyelogram

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
J	Joule
JCT	Junction
JRA	Juvenile rheumatic arthritis
JVD	Jugular venous distention
JX	Junction
KG	Kilogram
KS	Kaposi sarcoma
KUB	Kidneys, ureters, bladder
KV	Kilovolt
L1-L5	Lumbar vertebra
LAB	Laboratory
LAG	Lymphangiography/lymphangiogram
LAP	Laparotomy
LAT	Lateral
LAV	Lymphadenopathy-associated virus
LB(S)	Pound(s)
LBBB	Left bundle branch block
LBBX	Left breast biopsy
LBR	Large bowel resection
LCC	Large cleaved cell
LCIS	Lobular carcinoma in situ
LCM	Left costal margin
LDH	Lactic dehydrogenase
LE	Lower extremity
LINAC	Linear accelerator
LIQ	Lower inner quadrant
LIS	Lobular in situ
LKS	Liver, kidney, spleen
LKSB	Liver, kidney, spleen, bladder
LLE	Left lower extremity
LLL	Left lower lobe
LLQ	Left lower quadrant
LMP	Last menstrual period
LN(S)	Lymph node(s)
LN2	Lobular neoplasia, grade 2
LNBX	Lymph node biopsy
LND	Lymph node dissection
LNR	Lymph node resection

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
LOQ	Lower outer quadrant
LP	Lumbar puncture
LPN	Licensed practical nurse
LRG	Large
LRQ	Lower right quadrant
LS	Lumbosacral
LS SCAN	Liver/spleen scan
LSO	Left salpingo-oophorectomy
L-SPINE	Lumbar spine
LT	Left
LTCF	Long Term Care Facility
LUE	Left upper extremity
LUL	Left upper lobe
LUO	Left ureteral orifice
LUOQ	Left upper outer quadrant
LUP ERYTH	Lupus erythematosus
LUQ	Left upper quadrant
MAA	Melanoma associated antigen
MAB	Monoclonal antibody
MALIG	Malignant
MAND	Mandible/mandibular
MAST	Mastectomy
MAT	Multifocal arterial tachycardia
MAX	Maximum
MC	Medical center
MC(H)	Millicurie (hours)
MC-AB, MCAB	Monoclonal antibody
MCG	Microgram
MCID	Mixed combined immunodeficiency
MCL	Midclavicular line
MCS	Malignant carcinoid syndrome
M-CSF	Macrophage colony-stimulating factor
MCTD	Mixed connective tissue disease
MD	Moderately differentiated
MDS	Myelodysplasia/myelodysplastic syndrome
MED	Medication
MED	Medicine
METS	Metastatic/Metastasis

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
MEV	Million electron volts
MFH	Malignant fibrous histiocytoma
MG	Myasthenia gravis
MG(H)	Milligram (hours)
MHX	Medical history
MI	Myocardial infarction
MICRO	Microscopic
MID	Middle
MIN	Minimum
MIN	Minute
ML	Middle lobe
ML	Milliliter
MM	Millimeter
MM	Multiple myeloma
MOAB	Monoclonal antibody
MOD	Moderate (ly)
MOD DIFF	Moderately differentiated
MPD	Myeloproliferative disease
MPVC	Multifocal premature ventricular contraction
MRCP	Magnetic resonance cholangiopancreatography
MRI	Magnetic resonance imaging
MRM	Modified radical mastectomy
MRSA	Methicillin Resistant Staphylococcus Aureus
MS	Multiple sclerosis
MSB	Main stem bronchus
MULT	Multiple
MVP	Mitral valve prolapse
MX	Mastectomy
N&V	Nausea and vomiting
NA	Not applicable
NAD	No acute/active disease
NBX	Needle biopsy
NDL	Nodular & diffuse lymphoma
NEA	Neoplasm embryonic antigen
NEC	Not elsewhere classified/classifiable
NED	No evidence of disease
NEG	Negative
NEOPL	Neoplasm

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
NER	No evidence of recurrence
NEURO	Neurology
NH	Nursing home
NHL	Non-Hodgkins lymphoma
NHML	Non-Hodgkin malignant lymphoma
N-I - N-XII	Nerves, Cranial 1-12
NK	Natural killer
NL	Normal
NLBX	Needle liver biopsy
NOS	Not otherwise specified
NR	Not recorded
NSCCA	Non small cell carcinoma
NSCLC	Non-small cell lung cancer
NSF	No significant findings
NVD	Neck vein distention
NX	Nephrectomy
OB	Obstetrics
OBS	Organic brain syndrome
OBST	Obstructed (-ing, -ion)
OD	Right eye
ONC	Oncology
OP	Operation
OP	Outpatient
OP RPT	Operative report
OPM	Occult primary malignancy
OPS	Outpatient surgery
OR	Operating room
ORTHO	Orthopedics
OS	Left eye
OTO	Otology
OZ	Ounce
P&A	Percussion and auscultation
P32	Phosphorus 32
PA	Posteroanterior
PAC	Premature atrial contraction
PALP	Palpated (-able)
PAP	Papanicolaou smear
PAP	Papillary

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
PATH	Pathology
PCL	Plasma cell leukemia
PCV	Polycythemia vera
PD	Poorly differentiated
PE	Physical examination
PEDS	Pediatrics
PERC	Percutaneous
PET	Positron emission tomography
PI	Present illness
PID	Pelvic inflammatory disease
PIN	Prostatic intraepithelial neoplasia
PIN III	Prostatic intraepithelial neoplasia, grade III
PLL	Prolymphocytic leukemia
PLT	Platelets
PMH	Past/personal (medical) history
PMP	Primary medical physician
PNET	Primitive neuroectodermal tumor
POOR DIFF	Poorly differentiated
POS	Positive
POSS	Possible
POST	Posterior
POST OP	Postoperative (-ly)
PPD	Packs per day
PRA	Progesterone receptor assay
PRE OP	Preoperative (-ly)
PREV	Previous
PROB	Probable (-ly)
PROCTO	Proctoscopy
PSA	Prostatic specific antigen
PT	Patient
PT	Physiotherapy/Physical therapy
PTA	Prior to admission
PTC	Percutaneous transhepatic cholecystogram
PTNM	Pathologic tumor, nodes, metastases
PUD	Peptic ulcer disease
PULM	Pulmonary
PULM ART	Pulmonary artery
PVD	Peripheral vascular disease

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
Q	Every
QD	Every day
QUAD	Quadrant
R/O	Rule out
RA	Rheumatoid arthritis
RAD	Radiation absorbed dose
RAI	Radioactive iodine
RBBB	Right bundle branch block
RBBX	Right breast biopsy
RBC	Red blood cells (count)
RCM	Right costal margin
RE	Regarding
REC'D	Received
REG	Regular
RESEC	Resection (ed)
RESP	Respiratory
RESPIR	Respiratory
RHD	Rheumatic heart disease
RIA	Radioimmunoassay
RIQ	Right inner quadrant
RLE	Right lower extremity
RLL	Right lower lobe
RLQ	Right lower quadrant
RMC	Regional medical center
RML	Right middle lobe
RND	Radical neck dissection
ROF	Review of outside films
ROQ	Right outer quadrant
ROS	Review of outside slides
RSO	Right salpingo-oophorectomy
RSR	Regular sinus rhythm
RT	Radiation therapy
RT	Right
RUE	Right upper extremity
RUL	Right upper lobe
RUO	Right ureteral orifice
RUQ	Right upper quadrant
RX	Prescription

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
S/P	Status post
S1-S5	Sacral vertebra
SARC	Sarcoma
SATIS	Satisfactory
SB	Small bowel
SBO	Small bowel obstruction
SBR	Small bowel resection
SCC	Squamous cell carcinoma
SCID	Severe combined immunodeficiency syndrome
SCLC	Small cell lung carcinoma
SGOT	Serum glutamic oxaloacetic transaminase
SGPT	Serum glutamic pyruvic transaminase
SIG COLON	Sigmoid colon
SLE	Systemic lupus erythematosus
SM	Small
SMA	Sequential multiple analysis
SNF	Skilled nursing facility
SO	Salpingo-oophorectomy
SOB	Short(ness) of breath
SPEC	Specimen
SQ	Squamous
SS	Summary stage
S-SPINE	Sacral spine
SSS	Sick sinus syndrome
STSG	Split thickness skin graft
SUBCU	Subcutaneous
SURG	Surgery/Surgical
SUSP	Suspicious/suspected
SVC	Superior vena cava
SX	Symptoms
TAH	Total abdominal hysterectomy
TAH-BSO	Total abdominal hysterectomy- bilateral salpingo-oophorectomy
T-ALL	T-cell acute lymphoblastic leukemia
TANI	Total axial (lymph) node irradiation
TBBX	Transbronchial biopsy
TCC	Transitional cell carcinoma
T-CLL	T-cell chronic lymphatic leukemia
TIA	Transient ischemic attack

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
TNM	Tumor, node, metastasis
TPN	Total parenteral nutrition
TRAM	Transverse rectus abdominous myocutaneous
TRANS-COLON	Transverse colon
TRUS	Transrectal ultrasound
TRUSP	Transrectal ultrasound of prostate
TS	Tumor size
T-SPINE	Thoracic spine
TTP	Thromboticthrombocytopenia purpura
TUR	Transurethral resection
TURB	Transurethral resection bladder
TURBT	Transurethral resection bladder tumor
TURP	Transurethral resection prostate
TVC	True vocal cord
TVH	Total vaginal hysterectomy
TX	Treatment
UDO	Undetermined origin
UE	Upper extremity
UGI	Upper gastrointestinal (series)
UIQ	Upper inner quadrant
UNDET	Undetermined
UNDIFF	Undifferentiated
UNK	Unknown
UOQ	Upper outer quadrant
URI	Upper respiratory infection
URQ	Upper right quadrant
US	Ultrasound
USO	Unilateral salpingo-oophorectomy
UTI	Urinary tract infection
VAG	Vagina/Vaginal
VAG HYST	Vaginal hysterectomy
VAIN	Vaginal intraepithelial neoplasia
VAIN III	Vaginal intraepithelial neoplasia (grade III)
VASC	Vascular
VIN	Vulvar intraepithelial neoplasia
VIN III	Vulvar intraepithelial neoplasia (grade III)
VS	Versus
W/	With

ABBREVIATIONS

ABBREVIATION	WORD/TERM(S)
W/F	White female W/F
W/M	White male
W/O	Without
W/U	Work-up
WAGR	Wilms (tumor), aniridia, genitourinary (abnormalities), and (mental)
WBC	White blood cells (count)
WD	Well differentiated
WELL DIFF	Well differentiated
WF-O	Will follow (in) office
WNL	Within normal limits
XR	Xray
YR	Year
YST	Yolk Sac Tumor

ABBREVIATIONS

CONTEXT-SENSITIVE ABBREVIATIONS

When using these abbreviations make sure the meaning of the abbreviation is readily apparent in the context in which it is used.

ABBREVIATION	WORD/TERM(S)
AP	Anteroposterior
	Abdominal perineal
BM	Bone marrow
	Bowel movement
CA	Calcium
	Carcinoma
DM	Diabetes mellitus
	Distant metastases
MIN	Minimum
	Minute
ML	Milliliter
	Middle lobe
MM	Millimeter
	Multiple myeloma
OP	Operation
	Outpatient
PAP	Papillary
	Papanicolaou smear
PT	Patient
	Physiotherapy/Physical therapy
RT	Right
	Radiation therapy

ABBREVIATIONS

SYMBOLS

SYMBOL	WORD/TERM (S)
-	Minus
#	Number
&	And
@	At
^	Above
+	Plus, Positive
<	Less/Less than
=	Equal(s)
>	Greater/Greater than, More/more than
X	Times

Page intentionally left blank

**APPENDIX M:
WEB PLUS DOWNLOAD INSTRUCTIONS**

**Procedure: Web Plus file download**

Purpose: To receive files sent from the Pennsylvania Cancer Registry (PCR), via a secure website.

This procedure is used when facilities have been sent a document from the PCR (for example, yearly reconciliation listing). Web Plus replaces the need to send confidential data in an email file attachment or on paper through post-mail.

General Information:

1. Security: Web Plus is an Internet-based application developed by the Centers for Disease Control and Prevention, National Program of Cancer Registries. Web Plus has been designed as a highly secure application that can be used to safely transmit data between reporting facilities and the PCR over the public internet.

Security is achieved by a combination of software features and network infrastructure. Web Plus is hosted on a secure Web server; the communication between the client and the server is encrypted with 128-bit encryption Secure Socket Layer technology.

Security features of the application include:

- Web Plus keeps an extensive log of user logins, data accesses, and updates for auditing purposes.
- User accounts can be locked out if invalid login attempts exceed a threshold value, configurable by the PCR Central Administrator.
- Initial passwords are randomly generated by the system and the user will be forced to change it after their first successful login.
- Current user activities are visible to the PCR Central Administrator through the Current User Activities page.
- User passwords are stored in the database using a one-way hash encryption method.
- The Web Plus configuration file will store the connection string to the SQL Server database in encrypted format.
- The application times out after a specified time period.
- Web Plus uses form-based authentication where users are required to enter their unique user ID and strong password to be authenticated by the application.

2. Screen Resolution: The resolution for Web Plus should be 1024 X 768. If the resolution set on your PC is different you may still be able to use Web Plus, but Web Plus has been designed to be viewed best at 1024 X 768 or higher. You will receive a message on the Web Plus log-in screen if your resolution is not set correctly. Contact your Information Technology (IT) department if you are unsure how to change your screen resolution.

3. Web Plus icon on desk top: It is recommended that your IT department create an icon on your desktop from the Web Plus link for easy access to the application.
4. Email notification: When you have files to retrieve in WebPlus, you will receive an email from your PCR Field Representative telling you about the file and reminding you of these instructions on how to get it.
5. Password protected or encrypted files: Files uploaded to your facility from the PCR via Web Plus will not be password protected or encrypted. The security features of Web Plus replaces the need to password protect or encrypt files.
6. Password changes: You will be prompted to change your password the first time you log into Web Plus and then every 60 days after that. If you forget your password, contact your PCR Field Representative at (800) 272-1850 or (717) 783-2548.
7. User ID or website location: If you forget your User ID or cannot locate the Web Plus website, contact your PCR Field Representative at (800) 272-1850 or (717) 783-2548.

Procedure:

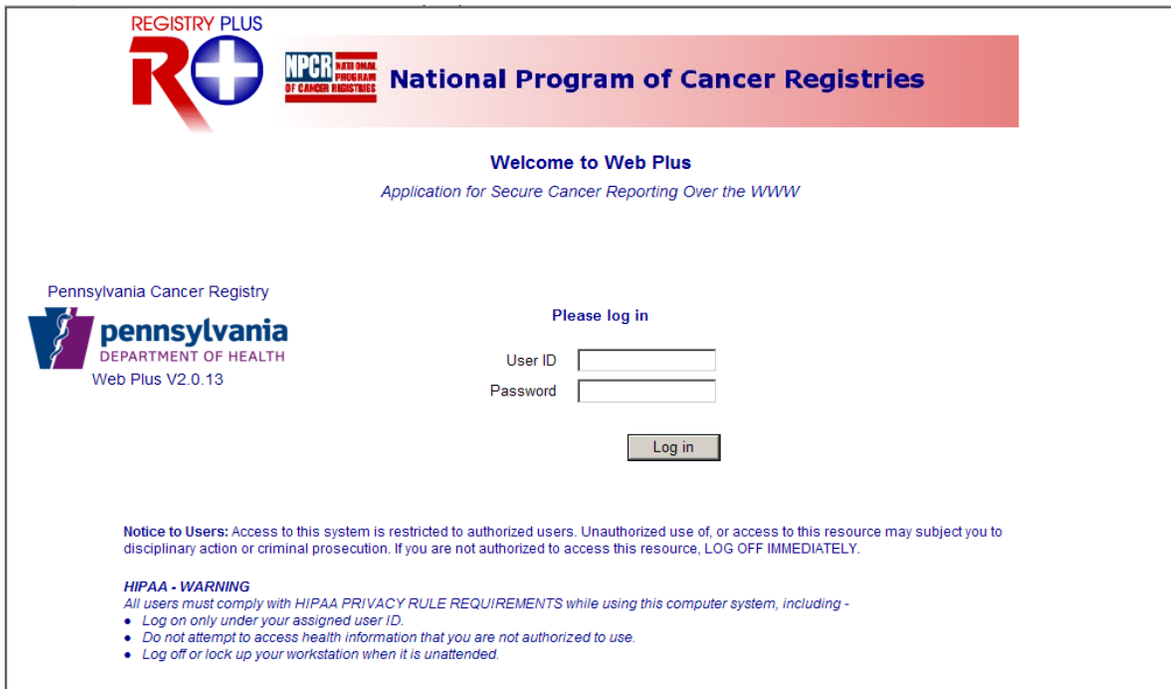
Downloading files

1. Open Web Plus using the link in the email previously sent by the PCR or type the following into your web address line

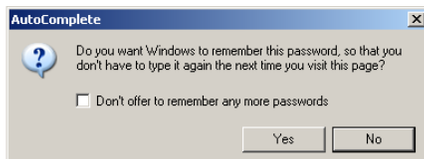
<https://pcr.health.pa.gov/webplus>

2. Type your User ID and password into the appropriate boxes. Note: Your User ID and password were previously sent via email by the PCR.

Click Log in.

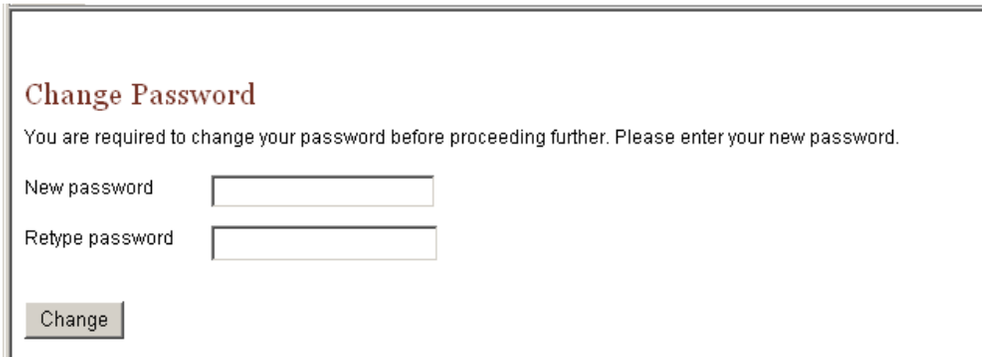


If the following message appears, click No.



*The first time you log in, the screen below will appear forcing you to change your password.

Enter a new password using the following criteria: “Password must be between 8 to 20 characters, contain at least one digit and one alphabetic character, and must not contain any special characters”.

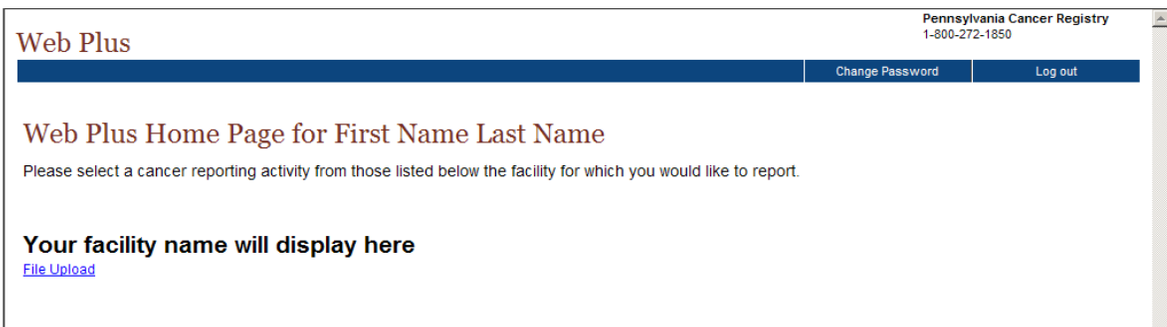


The screenshot shows a web form titled "Change Password" in a red font. Below the title, a message states: "You are required to change your password before proceeding further. Please enter your new password." There are two input fields: "New password" and "Retype password". A "Change" button is located at the bottom left of the form area.

Click on Change.

If the password does not meet the criteria specified above or if the new password does not match the retype password line, you will receive a message ‘Password not changed’. The Change Password screen will remain until the password meets the criteria and the two password lines match.

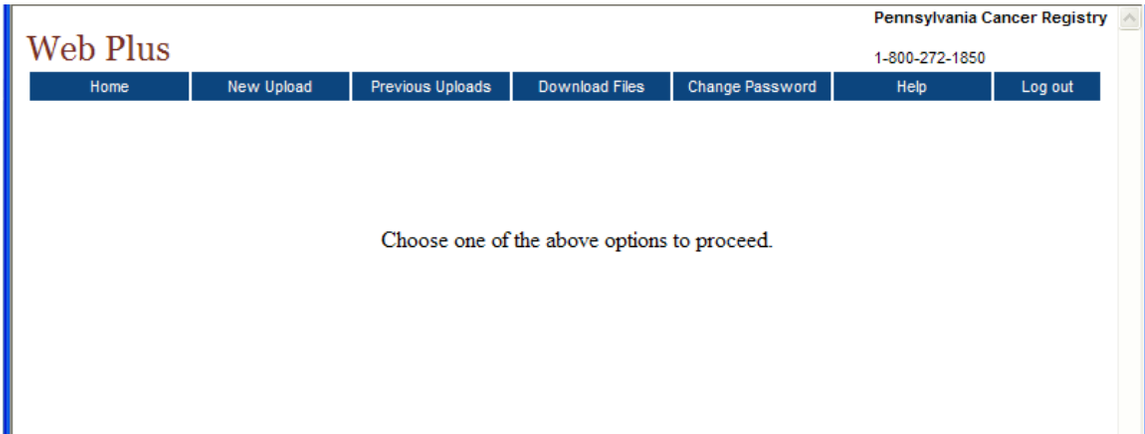
3. The Web Plus home page for your facility opens.



The screenshot shows the "Web Plus" home page. At the top right, it says "Pennsylvania Cancer Registry 1-800-272-1850". Below this is a navigation bar with "Change Password" and "Log out" buttons. The main heading is "Web Plus Home Page for First Name Last Name". Below the heading, it says "Please select a cancer reporting activity from those listed below the facility for which you would like to report." There is a section titled "Your facility name will display here" with a blue link labeled "File Upload".

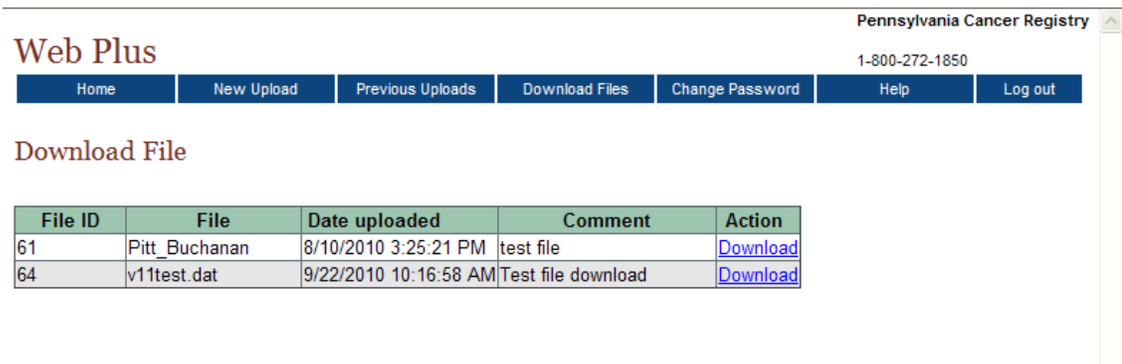
4. Click on File Upload under your Facility name.

The following screen will display:



5. Click on the Download Files blue box.

The following screen will display:



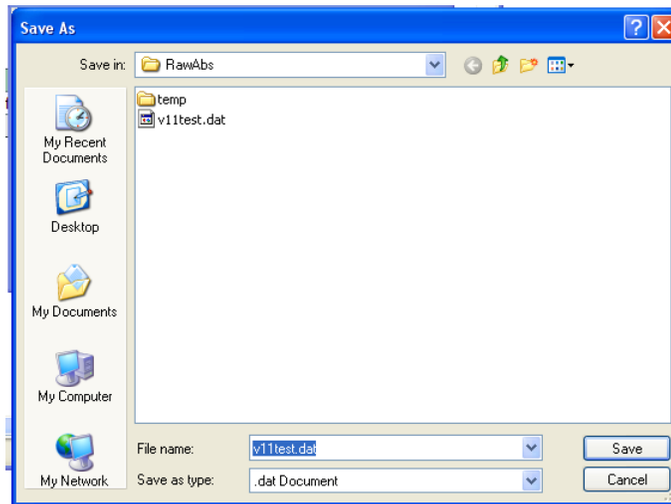
6. Click on Download under Action on the line with the file you want to retrieve.

The following screen will display:



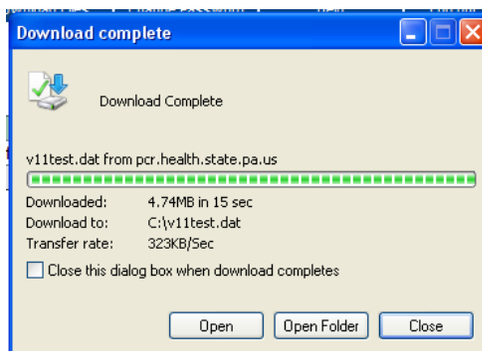
7. Click Save.

The following screen will display:



8. Rename the file (optional).
9. Select the location in which to save the file.
10. Click Save.

The following screen will display when the download is complete.



11. Click Close. Screen will return to display as in #5.
12. Repeat steps #6 to #11 for each additional file.
13. Click on Log Out blue box to close Web Plus when finished.

APPENDIX N:
PCR Manual Page Revisions

Following the initial release of *PCR Manual, 2010 Edition* in September 2010, the manual has undergone some of modifications and revisions. All revisions have been made to the online edition of the manual.

This edition contains all the necessary documentation to support changes in case reporting to accommodate the standard data item changes implemented in 2011 and after. This appendix contains changes introduced since the initial release of the *PCR Manual, 2010 Edition*.

Page #	Section	Change
January 2016 Whole Manual		
	Web links	All web links were updated as needed to current web addresses for standard documents.
January 2016 Part Three		
110	Tumor Size Summary	New Field for 2016; Replaces CS Field
113	Regional Nodes Examined	New Field for 2016; Replaces CS Field
115	Regional Nodes Positive	New Field for 2016; Replaces CS Field
117	Lymph-Vascular Invasion	New Field for 2016; Replaces CS Field
119	Mets at DX-Bone	New Field for 2016; Replaces CS Field
121	Mets at DX-Brain	New Field for 2016; Replaces CS Field
123	Mets at DX-Distant LN	New Field for 2016; Replaces CS Field
125	Mets at DX-Liver	New Field for 2016; Replaces CS Field
127	Mets at DX-Lung	New Field for 2016; Replaces CS Field
129	Mets at DX-Other	New Field for 2016; Replaces CS Field
135	AJCC-TNM Clinical and Pathologic Staging	Direct coding of AJCC-TNM required for 2016
January 2016 Appendix E		
	Effective January 1, 2016	Section was added to list the new terms, synonyms and related terms that need to be added to ICD-O-3 effective January 1, 2016.
January 2016 Appendix F		
<p>Added Items:</p> <p>Tumor Size Summary, Mets at DX-Bone, Brain, Distant LN, Liver, Lung, Other TNM Path T, TNM Path N, TNM Path M, TNM Path Stage Group, TNM Path Descriptor TNM Clin T, TNM Clin N, TNM Clin M, TNM Clin Stage Group, TNM Clin Descriptor TNM Edition Number</p>		
January 2015 Whole Manual		
	Web links	All web links were updated as needed to current web addresses for standard documents.
January 2015 Part One		
5	Carcinoid of the Appendix	Updated to reflect effective January 1, 2015 all carcinoids of the appendix are reportable.
January 2015 Part Three		
51	Sex	Following Codes were added:

Page #	Section	Change
		5- Transsexual, natal male 6- Transsexual, natal female
113	SEER Summary Stage	Revised to reflect the requirement of directly coded SEER Summary Stage for cases diagnosed in 2015 and after.
125	RX Date-Most Definitive Surgery	This is a new data item added for 2015
126	RX Date-Most Definitive Surgery	This is a new data item added for 2015
January 2015 Part Four		
207	Completeness	Deleted reference to Cases Accessioned by the PCR. These lists are no longer generated effective Febuary 2015.
January 2015 Appendix E		
	Effective January 1, 2015	Section was added to list the new terms, synonyms and related terms that need to be added to ICD-O-3 effective January 1, 2015.
January 2014 Whole Manual		
	Web links	All web links were updated as needed to current web addresses for standard documents.
January 2014 Part One		
19	Contacting the PCR	Section was revised to reference PCR Web Site.
January 2014 Part Three		
	Grade Path Value	Data item is no longer required. Pages were deleted.
	Grade Path System	Data item is no longer required. Pages were deleted.
98-104	Grade	Section completely replaced by new grade coding guidelines
110-111	Collaborative Stage	Added V02.05 required for cases diagnosed on or after January 1, 2014 and after.
January 2014 Appendix D		
ICD-9-CM List		<ul style="list-style-type: none"> • <u>Code 173</u> - 173 codes (malignant neoplasm of skin) were listed individually to eliminate non-reportable basal and squamous cell carcinomas of the skin • <u>Code 238.6</u> - deleted from list. Plasmacytoma and solitary myeloma are now coded to 203.8. • <u>Code 288.4</u> – deleted from list. Langerhans cell histiocytosis is now coded to 277.89 • <u>Code 277.89</u> – added to list. This code includes other specified disorders of metabolism (reportable terms include: Hand-Schuller Christian disease, histiocytosis (acute) (chronic), histiocytosis X (chronic) • <u>Supplemental list was deleted</u>
ICD-10-CM List		An ICD-10-CM casefinding list was added. It is effective on October 1, 2014.
January 2014 Appendix E		

Effective January 1, 2014	Section was added to list the new terms, synonyms and related terms that need to be added to ICD-O-3 effective January 1, 2014.	
January 2014 Appendix F		
Deleted:		
<ul style="list-style-type: none"> • Grade Path Value • Grade Path System 		
January 2014 Appendix J		
Breast	SSF 10 and 12 were deleted from list of PCR required SSFs	
January 2014 Appendix P		
A Summary of Changes in Data Collection Standards was added as Appendix P		
January 2013 Part One		
2	PCR Reference Date	The following note was deleted: <i>Note:</i> To assure complete case ascertainment, reference date is not used to determine what cases are reportable to the PCR. See <i>Part One, Date of Diagnosis Reportability</i> .
8	Date of Diagnosis Reportability	Completely revised to reflect cases diagnosed prior to 1985 and cases with an unknown date of diagnosis are not reportable.
14	Non-Reportable Situations	#10 was added: <u>Diagnosed Prior to 1985</u> - Patients with a date of diagnosis prior to January 1, 1985 are not reportable.
14	Non-Reportable Situations	#11 was added: <u>Unknown Date of Diagnosis</u> - Cases with an unknown month, day and year of diagnosis are no longer required to be reported.
16	When in Doubt	Question #2 added: Is the Date of Diagnosis known and is it on or after January 1, 1985?
48	Birthplace-State	Changed from Birth Place to Birthplace-State
49	Birthplace-Country	New data item
January 2013 Part Three		
55	Race	#6 was added- <i>Specific Race and Non-Specific Race</i> - Code only the specific race when both a specific race code and a non-specific race code apply.
205	FIN Coding System	Deleted. No longer a required System Code
	Race Coding System	Deleted. No longer a required System Code
206	First Course Calc Method	Deleted. No longer a required System Code
	COC Coding System Current	Deleted. No longer a required System Code
January 2013 Appendix D		
The following codes were deleted on SEER 2013 Casefinding List. The list was revised to enhance the accuracy of identifying reportable conditions.		
<ul style="list-style-type: none"> • 236.0 • 273.2 • 288.4 		

Page #	Section	Change
January 2013 Appendix F		
Added:		
<ol style="list-style-type: none"> 1. Birthplace-State 2. Birthplace-Country 		
Deleted:		
<ol style="list-style-type: none"> 1. FIN Coding System 2. Race Coding System 3. First Course Calc Method 4. COC Coding System Current 		
January 2013 Appendix I		
Whole appendix		Added state/Country codes for BIRTHPLACE-STATE and BIRTHPLACE-COUNTRY data items.
January 2012 Part Three		
125-129	Rx Summ - Scope Reg LN Surg	Added additional definitions/instructions to ensure the correct coding of regional lymph node surgery.
141	RX Summ- Surg/Rad Seq	Definition for Code 6 is now “ <i>Intraoperative radiation therapy with other therapy administered before and/or after surgery</i> ”. “And” was added.
		Added code 7- <i>Surgery both before and after radiation.</i>
165	RX Summ-Systemic Sur Seq	Definition for Code 6 is now “ <i>Intraoperative systemic therapy with other therapy administered before and/or after surgery</i> ”. “And” was added.
		Added code 7- <i>Surgery both before and after systemic therapy.</i>
167	RX Summ-Other	The rules for coding other treatment for hematopoietic diseases in #1 have been revised to reflect the changes released in the <i>2012 Hematopoietic and Lymphoid Neoplasm Case Reportability and Coding Manual</i> .
August 2011 Appendix J (PCR Required SSFs)		
1	Breast	SSF 16 was deleted as a required SSF.
January 2011 Part One		
1	PCR Manual, 2010 Edition	The following was added <i>Note:</i> “As revisions are made to the <i>PCR Manual, January 2010 Edition</i> the changes will be documented in <i>Appendix O</i> . In addition, the revised pages will have the date the revision is effective in the footer.”
7	Ambiguous Terminology	The following was added to top of page

Page #	Section	Change
		<p>Note: Do not substitute synonyms such as “supposed” for presumed or “equal” for comparable. Do not substitute “likely” for “most likely.”</p> <p>The following was deleted under #3-a-2 “discrepancies”</p> <p>Note: If the word or an equivalent term does not appear on either the reportable or not reportable list or is not a form of a word on the reportable list, the term is not diagnostic of cancer. Do not report the case. Forms of the word include: “Favored” rather than Favor(s); “appeared to be” rather than appears. Do not substitute synonyms such as “supposed” for presumed or “equal” for comparable.</p>
8	Ambiguous Terminology	<p>This note was deleted under #3 “discrepancies”</p> <p>Note: If the word or an equivalent term does not appear on either the reportable or not reportable list or is not a form of a word on the reportable list, the term is not diagnostic of cancer. Do not report the case. Forms of the word include: “Favored” rather than Favor(s); “appeared to be” rather than appears. Do not substitute synonyms such as “supposed” for presumed or “equal” for comparable.</p>
65 and 66	Class of Case, Codes and Definitions	<p>Added “part of first course treatment was done elsewhere” to codes 13 and 21</p> <p>Added “or a decision not to treat was done at the reporting facility” to code 22</p> <p>Added “treatment plan only” to code 30 examples.</p> <p>Added “or hospital provided care that facilitated treatment elsewhere (for example stent placement)” to code 31</p>
<p>January 2011 Part Three</p>		
65 and 66	Class of Case, Codes and Definitions	<p>Added “(active disease)” to code 32</p> <p>Added “(disease not active)” to code 33</p> <p>Added #2: <u>Ownership of Physician Practice</u>- If the hospital has purchased a physician practice, it will be necessary to determine whether the practice is now legally considered part of the hospital (their activity is coded as the hospital’s) or not. If the practice is not legally part of the hospital, it will be necessary to determine whether the physicians involved are staff physicians or not, as with any other physician.</p> <p>The following was added to #6.</p> <p>Note: Most paired sites cannot develop midline tumors.</p>

Page #	Section	Change
		<p>Skin of the trunk can have a midline tumor, but breast cannot. “Midline of the right breast” is coded 1, right; midline in this usage indicates the primary site is C50.8 (overlapping sites).</p> <p>Added #7:</p> <p><u>Brachytherapy with 125 seeds</u>- Assign code 53 for brachytherapy with 125 seeds. Seeds are always low dose therapy because they are left in place and the radioactivity decays over time.</p> <p>Clarified at least two courses of radiation must be given to assign code 4.</p>
67	Recording Class of Case	New data item added for 2011.
90	Recording Laterality	Clarified at least two courses of systemic therapy must be given to assign code 4.
136	RAD-Regional Modality	<p>Added #4:</p> <p>4. <u>PUVA</u>- Assign code 1 in the rare event Psoralen (P) and long-wave ultraviolet radiation (UVA) it is used as treatment for extremely thin melanomas or cutaneous T-cell lymphomas (e.g. mycosis fungoides).</p>
140	RX Summ- Surg/Rad Seq	<p>Expanded #2 to include:</p> <p>a. Patient refused treatment</p> <p>b. Physician decides not to treat for any reason, such as the presence of Comorbidities.</p>
142	Reason for No Radiation	<p>Added #4:</p> <p>5. <u>PUVA</u>- Assign code 1 in the rare event Psoralen (P) and long-wave ultraviolet radiation (UVA) it is used as treatment for extremely thin melanomas or cutaneous T-cell lymphomas (e.g. mycosis fungoides).</p>
164	RX Summ-Systemic Sur Seq	<p>Expanded #2 to include:</p> <p>c. Patient refused treatment</p> <p>d. Physician decides not to treat for any reason, such as the presence of Comorbidities.</p>
167	RX Summ-Other	<p>Added #4:</p> <p>6. <u>PUVA</u>- Assign code 1 in the rare event Psoralen (P) and long-wave ultraviolet radiation (UVA) it is used as treatment for extremely thin melanomas or cutaneous T-cell lymphomas (e.g. mycosis fungoides).</p>
174	RX Summ-Treatment Status	SSF 10 was added as a required SSF.

**January 2011
Appendix D (Reportable ICD-9-CM Codes)**

Page #	Section	Change
<p>The following codes were deleted to SEER 2011 Casefinding List. The list was revised in enhance the accuracy of identifying reportable conditions.</p> <ul style="list-style-type: none"> • 227.9 • 238.81-238.89 • 237.2-237.4 • 288.3 • 237.70-237.79 		
<p>January 2011 Appendix J (PCR Required SSFs)</p>		
1	BileDuctsDistal	Reworded code 66 to state: Excision of an intrahepatic bile duct PLUS partial hepatectomy
2	Breast	Reworded code 75 to state: Extrahepatic bile duct and hepatectomy WITH transplant
5	Testis	Added the following below code 40: A total (simple) mastectomy removes all breast tissue, the nipple, and areolar complex. An axillary dissection is not done, but sentinel lymph nodes may be removed.
<p>January 2011 Appendix K (Surgery to Primary Site Codes)</p>		
19	Liver and Intrahepatic Bile Ducts- Code 66 and 75	Added “and no specimen sent to pathology or unknown if sent” to code 19
		Added “with specimen sent to pathology” to code 21
28	Breast- code 40	Added “and no specimen sent to pathology or unknown if sent” to code 19
29	Breast- code 76	Added “with specimen sent to pathology” to code 21
36	Prostate- code 19	Added “and no specimen sent to pathology or unknown if sent” to code 19
36	Prostate-code 21	Added “with specimen sent to pathology” to code 21
<p>January 2011 Appendix M (TipSheets)</p>		
<p>Appendix M was added for <i>TipSheets</i>. <i>TipSheets</i> provide tips and clarifications. <i>TipSheets</i> on Breast and Genitourinary have been included. More <i>TipSheets</i> will be provided as they are developed.</p>		

APPENDIX O:
Summary of Changes in Data Collection Standards

Data collection standards have changed numerous times over the years. The following table summarizes these changes including the dates the changes became effective.

Data Collection Standard	Effective Date
2016	
Directly Coded AJCC-TNM Clinical and Pathologic Stage required	Date of Diagnosis 1/1/2016 and after
Collaborative Stage – SSF’s, Lymph-vascular Invasion, Regional LN Positive and Regional Lymph Nodes Examined required only	Date of Diagnosis 1/1/2016 and after
ICD-O-3 Changes - New terms, synonyms, and related terms	Date of Diagnosis 1/1/2016 and after
2015	
ICD-O-3 Changes - New terms, synonyms, and related terms	Date of Diagnosis 1/1/2015 and after
Directly Coded SEER Summary Stage required once again	Date of Diagnosis 1/1/2015 and after
2014	
ICD-O-3 Changes - New terms, synonyms, and related terms	Date of Diagnosis 1/1/2014 and after
Grade – New instructions for coding grade	Date of Diagnosis 1/1/2014 and after
2012	
Scope Regional Lymph Node Surgery Instructions revised	Date of Diagnosis 1/1/2012 and after
Hematopoietic and Lymphoid Database and Manual revised	Date of Diagnosis 1/1/2012 and after; current web-based version should be used
2010	
Collaborative Stage Revised (Version 2)	Date of Diagnosis 1/1/2010 and after
Hematopoietic and Lymphoid Database and Manual Implemented	Date of Diagnosis 1/1/2010-12/31/2011
New Reportable Hematopoietic/Lymphoid Conditions and terms added	Date of Diagnosis 1/1/2010 and after
2007	
SEER Multiple Primary and Histology Coding Manual Implemented	Date of Diagnosis 1/1/2007 and after
2005	
SEER RX Implemented	Date of Diagnosis 1/1/2005; current web version should be used
2004	
Collaborative Stage Implemented (Version 1)	Date of Diagnosis 1/1/2004 and after
Benign Brain and Central Nervous System tumors became reportable	Date of Diagnosis 1/1/2004 and after

2001	
ICD-O-3 Implemented	Date of Diagnosis 1/1/2001 and after

ALPHABETICAL INDEX

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