# North American Association of Central Cancer Registries

# Standards for Cancer Registries Volume II

# **Data Standards and Data Dictionary**

Ninth Edition Record Layout Version 10.2

> Edited By Lori A. Havener Dianne Hultstrom

> > March 2004

**Sponsoring Organizations** 

American Cancer Society American College of Surgeons American Joint Committee on Cancer Canadian Association of Provincial Cancer Agencies Centers for Disease Control and Prevention Health Canada National Cancer Institute National Cancer Registrars Association Statistics Canada

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Comments and suggestions on this and other NAACCR standards documents are welcome. Please send your comments to the Editor or any member of the NAACCR Board of Directors.

#### The other volumes in the series, Standards for Cancer Registries, are:

#### \* Volume I, Data Exchange Standards and Record Description

Intended for programmers and selected users of central cancer registry data, this Volume provides the record layouts and specifications for a number of standard NAACCR record formats, including: the standard record layouts for data exchange among central cancer registries; an update/correction record layout; and an analysis record layout that provides standard recodes for grouping selected variables such as race and primary site, as well as algorithms for converting data from one version of the International Classification of Diseases for Oncology to another. The Volume is released annually as an electronic document and posted on the NAACCR Web Site.

Volume III, Standards for Completeness, Quality, Analysis, and Management of Data Intended for central registries, this provides detailed standards for many aspects of the operation of a population-based cancer registry.

Copies of all standards documents can be viewed or downloaded from NAACCR's Web Site at: http://www.naaccr.org. For additional paper copies, write to the NAACCR Executive Office at: 2121 W. White Oaks Drive, Suite C, Springfield, IL, 62704-6495.

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# PREFACE TO THE NINTH EDITION

Standardization of cancer registry data is a core component of cancer registration and surveillance. It provides the foundation for developing comparable data among registries that can then be combined for the compilation of national or regional rates. Standardization also allows data from different registries to be used for comparison of variations in cancer rates among different populations and across geographic boundaries.

NAACCR continues its strong commitment to all its members in North America to maintain standardization of cancer registry data, as evidenced in the publication of the Ninth Edition of NAACCR Standards for Cancer Registry Volume II: *Data Standards and Data Dictionary*. This Volume represents a new level of collaboration and commitment among our members to collect timely and accurate uniform data, in response to the needs of changing medical practice and delivery. As in the past, there will be challenges in the implementation of the new and revised standards in this Volume. I hope that these revisions will assist our members in achieving the NAACCR mission, namely, providing current, high-quality, and useful data for the cancer surveillance community and cancer control researchers with the ultimate goal of reducing cancer morbidity and mortality in North America. **Please note that black vertical lines in the outside margins highlight revisions from the previous version**.

This Volume is the result of the collaboration and cooperation of our sponsoring members, many of which set the standards for the differing needs of their organizations. We are especially grateful to the National Cancer Institute's Surveillance, Epidemiology and End Results Program, the Centers for Disease Control and Prevention's National Program for Cancer Registries, the American College of Surgeons' Commission on Cancer, the American Joint Commission on Cancer, and the National Cancer Registrars Association for their collaborative spirit and willingness to compromise in the interest of uniformity and achieving a common goal. On behalf of the NAACCR Board of Directors, I express our gratitude to these organizations for their support of the work that this Volume represents.

This new Edition also represents the voluntary contributions of NAACCR Committees, Subcommittees, and Work Groups. I would like to thank the many individuals for their commitment to this project. Special appreciation goes to the members of the Uniform Data Standards Committee and the Volume II Work Group for their diligence, valuable insights, consensus building, and this final document.

The NAACCR Board of Directors would like to extend a special thanks to Dianne Hultstrom, Chair of the Volume II Work Group. Under her leadership, this group provided detailed documentation for the over-ride flags and reviewed other data items for consistency with manuals published by standard setters and put countless hours of work into completing this revision. The Board of Directors also would like to recognize the leadership of Andrew Stewart, Chair of the Uniform Data Standards Committee. We greatly appreciate their efforts in bringing standard setters together, resolving differences, building consensus, and coordinating all aspects of this project.

Dennis Deapen, DrPH President

# CHAPTER I

# PROBLEM STATEMENT, GOALS, AND SCOPE OF THIS DOCUMENT

# THE PROBLEM

In the late 1980s, increased efforts to pool data collected by different cancer registries for different purposes drew attention to problems encountered as a result of insufficient data standardization. It became clear to the cancer registry community that the lack of standardization had a substantial cost and limited more widespread use of valuable data. Three examples follow:

#### Electronic Submission of Hospital Registry Data to State or Other Central Registries

Central registries recognized that data quality and collection efficiency could be improved with electronic data reporting by means of a diskette, modem, or the Internet. Many registries have established systems for receiving electronic data from multiple sources. Often, these data were collected using different software, different data variables, different codes, and different coding rules. Central registries experienced the frustration of mapping submission files into their own data systems. Software providers were frustrated at the need to prepare submissions for multiple state registries that different from each other and followed different models of electronic data collection.

#### North American Association of Central Cancer Registries Data Evaluation and Publications Committee Activities

The North American Association of Central Cancer Registries (NAACCR) requested statistical analysis files from its member registries in the standard NAACCR Data Exchange Record Layout<sup>1</sup> to prepare descriptive epidemiological data about the participating areas. However, data sets submitted by the participants differed; the original codes, data formats, edits, and coding rules varied; and a significant amount of work was required to produce comparable summary statistics.

#### National Cancer Data Base

The National Cancer Data Base (NCDB) is a joint project of the American College of Surgeons' (ACoS) Commission on Cancer (COC) and the American Cancer Society (ACS) that pools data submitted by participating hospitals to address questions of clinical interest. Discrepancies in codes, format, and data sets, however, required effort and interpretation before the data could successfully be pooled.

Data items used by different registries or software systems varied in their definition and codes, even when they had the same name and were intended to represent the same information. Other problems encountered in pooling data included the lack of standardization regarding the use of blanks in fields and the inconsistent use of blanks, dashes, and defined codes for "unknown" data. More substantial discrepancies were less easy to detect and correct. Hospitals were faced with conflicting standards when they were both reporting to a central registry and maintaining a database consistent with COC standards, and the requirements were not the same.

#### THE SOLUTION

Many of NAACCR's sponsoring organizations, including the National Cancer Institute (NCI), the Centers for Disease Control and Prevention (CDC), and COC recognized that increasing standardization is an essential step in decreasing the costs associated with data collection; making more efficient use of increasingly limited human

resources needed for data collection, management, and analysis; and obtaining more useful data that can be compared across registries and geographic areas.

Preparation of a statement of consensus on data standards for cancer registries was proposed by the NCDB and the NAACCR Data Exchange Committee, and prepared by a subcommittee of NAACCR's Uniform Data Standards Committee. At the same time, CDC entered into an agreement with NAACCR—one of the projects to be accomplished under that agreement was the preparation of broader standards for population-based cancer registries. The two efforts were complementary, producing separate but related documents that together specified NAACCR standards. The continued support from CDC has enabled continued development and maintenance of standards. The results of these efforts are the following standards documents published to date:

# NAACCR Standards Volume I:

Gordon B, editor. Standards for Cancer Registries, Volume I, Data Exchange Standards and Record Description. Version 3.0. Sacramento (CA): American Association of Central Cancer Registries; February 1994.

Gordon B and Seiffert J, editors. Standards for Cancer Registries, Volume I, Data Exchange Standards and Record Description. Version 5.1. Sacramento (CA): North American Association of Central Cancer Registries; 1997.

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North American Association of Central Cancer Registries. Standards for Cancer Registries, Volume I, Data Exchange Standards and Record Description. Version 7. Sacramento (CA): North American Association of Central Cancer Registries; January 1, 1999.

Abe T and Seiffert J, editors. North American Association of Center Cancer Registries, Standards for Cancer Registries, Volume I, Data Exchange Standards and Record Description. Version 9. Springfield, IL: North American Association of Center Cancer Registries, September 7, 2000. (Electronic version only; available at http://www.naaccr.org.)

Havener L, Abe T, Bushhouse S, Gordon B, Hill K, Hurlbut A, Seiffert J, editors. Standards for Cancer Registries Volume I: Data Exchange and Data Descriptions, Version 10.1. Springfield, IL: North American Association of Central Cancer Registries, July 2003.

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Seiffert J, editor. Standards for Cancer Registries Volume II: Data Standards and Data Dictionary, Changed Data Dictionary Entries Only. Sacramento (CA): North American Association of Central Cancer Registries; April 13, 1998.

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North American Association of Central Cancer Registries. Standards for Completeness, Quality, Analysis, and Management of Data, Volume III. Springfield (IL): North American Association of Central Cancer Registries; September 2000.

#### NAACCR Standards Volume IV:

Seiffert J, Capron S, and Tebbel J, editors. Standards for Cancer Registries Volume IV: Standard Data Edits. Sacramento (CA): North American Association of Central Cancer Registries; April 4, 1996.

#### **GOAL OF THIS DOCUMENT**

The goal of this document, which describes and publishes continuing, modified, and new data items and codes as well as the specification for transmission of data in record layout Version 10.2, is to define the NAACCR data standards for cancer registration for use by central registries, hospital-based registries, and other groups in North America as of January 1, 2005. Although the new and modified codes and the layout are available for use on that date, some registries may continue to use compatible earlier versions of the NAACCR record layout.

Objectives of the standardization effort, and of this document, are to:

- Provide a comprehensive reference to ensure uniform data collection
- Reduce the need for redundant coding and data recording between agencies

- Facilitate the collection of comparable data among groups
- Provide a resource document to help registries that are establishing or revising their databases
- Encourage the adoption of these standards by all parties.

This document will be used by new and existing facility-based and central cancer registries to ensure that the definitions and codes used within their programs are standard and consistent with those used by regional and national databases. Other potential users include registry software providers and those using registry data, especially if they are combining data from multiple sources or exchanging data. National standard-setting groups, such as ACoS, CDC, NAACCR, and NCI also will benefit.

The present document uses the same structure and philosophy as NAACCR's data exchange standards. Where a standard exists for an item or type of data, the standard is incorporated by reference. Where a variety of standards are in use, alternate coding schemes are provided, but the different items are kept separate or another data field is used to indicate which coding standard was used.

The NAACCR data exchange layout incorporates several record types that are combinations of standard components, such as demographic information, patient confidential information, and text. Thus, the different purposes and constraints of data exchange can be accommodated without the requirement for separate formats (see Volume I for specifics).

## SCOPE OF THIS DOCUMENT: WHAT STANDARDS ARE INCLUDED?

A variety of standards for cancer registries can be specified. Some standards apply to the data themselves, and other standards record activities in the registration process, such as death clearance procedures, follow-up methods, or quality control. Yet another standard might address the completeness of coverage of a population-based central registry, and still another the qualifications and adequacy of staffing.

The present document is limited to standards regarding data rather than procedures. More specifically, it focuses on a subset of possible data standards that NAACCR considers important to establish. These include:

#### \* Reportability

Reportability specifies the rules for which tumors are to be included in the registry (see Chapter III).

#### ✤ Data Items or Elements To Be Included

Data items or elements consist of required or recommended data items that a registry should collect and include in its database. Chapter IX contains standards for data set items.

Example: "Sex" is a standard data element on the list in Chapter IX.

#### Standardized Item Numbers and Item Names

For ease and consistency of reference, all items are assigned both item numbers and names (e.g., the item "Sex" is assigned the item number 220). The item number is intended to be permanent and will not change in future NAACCR standards publications. Assignment of permanent numbers was necessary because standard-setting organizations have changed item names over time or have applied similar names to items with different definitions. Item numbers allow the required precision of reference. When data items have been deleted the item numbers are retired; item numbers will never be reused for a different data item. Data

item numbers were not assigned consecutively to allow insertion of related items in the future. Ranges of available data item numbers have been assigned to different uses, as follows:

Range	Use
00001 - 04999	Data items in new case layouts, record types I, C, or A.
05000 - 06999	Data items in Analysis/Research record only.
07000 - 08999	Pathology Laboratory record.
09000 - 09099	Data items in Update/Correction record only.
09100 - 09499	Future use.
09500 - 09999	Data items for Local use.
10000 - 10499	System variables for Local use.
20000 - 20999	Data items for International use. (These data items are not within the purview of
	NAACCR, and NAACCR will not use the data item numbers in this range.)
99000 - 999999	Data items for Patient Care Evaluation studies. These may be assigned by ACoS or others. A large range is allotted because many new items may be assigned each year for individual studies.

Refer to NAACCR Standards Volume I for additional information on record layouts.

The NAACCR data item names are assigned to meet the needs of NAACCR and its data standards publications. Where possible, the NAACCR item name is the same as that used by the standard setter. However, the following constraints are placed on the names:

#### • Length

Data item names are limited to 25 characters because that is the maximum length for item names in the EDITS software system (see Chapter IV). Standardized abbreviations, punctuation, and spacing are used when necessary (i.e., the word "first" always is entered "1st," "treatment" is "RX," and so on). Other limitations will be imposed as needed. Thus, item names can be identical in this data standards volume and the NAACCR Metafile.

#### • Consistency

Consistency was attempted in formatting names and in using special characters. The character "--" is used to distinguish among item names built on the same stem name.

*Example: "Sequence Number--Hospital" and "Sequence Number--Central" are the names of two differently defined sequence numbers.* 

#### • Interrelated Items, Fields, and Subfields

To make the relationship among items more apparent, a constant term was consistently added to the stem of the name.

Example: All of the names of treatment fields related to radiation therapy begin with "Rad," so that in a list of item names they will appear together: Rad--No of Treatment Vol Rad--Elapsed RX Days

## Record Layout/Data Exchange

Record layout/data exchange identifies the position of the data item in a standard flat file data exchange record. These positions are indicated in Chapter VIII. Also, see Volume I<sup>1</sup> in this series for information on the data exchange and other NAACCR standard layouts.

Example: "Sex" is in character position 118 in the NAACCR Data Exchange Record Layout Version 10.2.

#### Codes

Codes identify allowable values, their meanings, and data entry formats for data items. Chapters X and XI specify the standard codes for each data item.

*Example for the item "Sex": Codes* 

- 1 Male
- 2 Female
- *3 Other (Hermaphrodite)*
- 4 Transsexual
- 9 Not stated

When it is necessary to collect more specific information than that represented by the standard codes, every effort should be made to ensure that the more specific codes would accurately collapse into the categories represented by the standard codes. This approach permits diversity without compromising inter-registry comparability or meta-analyses.

#### Coding Rules

Coding rules are the rules and interpretations for deciding the correct code for a given tumor. Coding rules are defined in the documentation of other standard-setting organizations. For each data item, Chapters IX and XI list a "Source of Standard," and the documentation of this source should be consulted for coding rule standards.

*Hypothetical Example: A coding rule might state what code to assign for sex when the medical record states the patient is female and the death certificate states male.* 

# **CHAPTER II**

# HISTORICAL BACKGROUND AND STATUS OF U.S. STANDARDS

# STANDARD-SETTING ORGANIZATIONS AND OTHER STANDARDS DOCUMENTS

Several organizations have played a major role in the development of cancer registry standards. They are listed in alphabetical order.

#### **American Cancer Society**

ACS historically has supported the development of standardized cancer classification systems, publishing the first code manual for the morphology of neoplasms in 1951. ACS has long supported the standard-setting programs of ACoS, including the Fundamental Tumor Registry Operations Education Program, the Registry Operations and Data Standards, and the American Joint Committee on Cancer (AJCC).

#### **American College of Surgeons**

Since the 1950s, ACoS has taken a leading role in establishing standards for hospital-based cancer programs and the cancer registries that are a part of such programs. Through its Approvals Program, COC implements its requirements for case management, registry operation and case inclusion, and data set specifications as published in:

- Cancer Program Standards (Standards of the Commission on Cancer, Volume 1),<sup>28</sup> which presents standards for the full range of cancer program activities, including the registry.
- Facility Oncology Registry Data Standards (FORDS) (Standards of the Commission on Cancer, Volume II),<sup>2</sup> which specifies standards for cases to be included in the registry, data items to be collected, and the codes and coding rules for those items.

Beginning with 2003 cases, COC requires approved cancer programs to use the codes published in FORDS.

Through NCDB, COC provides data quality feedback to facilities, software providers, and the general cancer registry community. Hospitals in the Approvals Program are required to submit non-confidential registry data to NCDB, and COC monitors the quality of data submissions in accordance with existing published standards for approved programs.

The NCDB Call for Data announcements, instructions, and technical specifications are available to download at no charge at http://www.facs.org. COC maintains an interactive Inquiry and Response Database to field questions about all cancer-related requirements at the same online site.

#### American Joint Committee on Cancer

AJCC formulates and publishes systems of classification of tumors by their anatomic site and histology through use of the Tumor, Node, Metastasis (TNM) staging system. The TNM staging system is the U.S. standard used by the medical profession to select the most effective treatments and determine prognosis to facilitate the management of cancer care. AJCC is dedicated to the ideal that all cancer cases should be staged, and it publishes the *Cancer Staging Manual*,<sup>5</sup> now in its Sixth Edition as well as the *Collaborative Staging Manual* and *Coding Instructions*.<sup>11</sup>

## National Cancer Registrars Association

An organization of cancer data professionals founded as the National Tumor Registrars Association in 1974, the National Cancer Registrars Association (NCRA) has been instrumental in the training and certification of cancer registrars. NCRA has produced a variety of educational materials, including guidelines for a college curriculum in cancer registry management, a planning manual for registry staffing, training materials for staging of cancer, and a publication on using cancer data to promote the services of the cancer registry. A college-level cancer registry methods textbook also was published (*Cancer Registry Management: Principles & Practice*, 1997).<sup>41</sup>

Since 1983, NCRA has promoted the certification of cancer registrars through a semi-annual examination. More than 4,000 Certified Tumor Registrars (CTRs) successfully have completed the exam, which evaluates technical knowledge of methods of cancer data collection, management, and quality control, as well as *International Classification of Diseases-Oncology* (ICD-O) topography and morphology coding and AJCC and Surveillance, Epidemiology and End Results (SEER) Program staging systems. To maintain their credentials, CTRs are required to complete 20 hours of continuing education every 2 years, which can be obtained by participating in conferences and teleconferences that NCRA has precertified, and by obtaining a passing score on quizzes in NCRA's *Journal of Registry Management*.

Membership in NCRA is open to anyone interested in cancer data collection. For further information, contact NCRA at the address on page xiii or on the Web at: http://www.ncra-usa.org.

#### National Coordinating Council for Cancer Surveillance

Founded in 1995, the National Coordinating Council for Cancer Surveillance (NCCCS) meets biannually to coordinate surveillance activities within the United States through communication and collaboration among major national cancer organizations, ensuring that the needs of cancer patients and the communities in which they live are fully served; that scarce resources are maximally used; and that the burden of cancer in the United States is adequately measured and ultimately reduced. NCCCS includes representatives from ACoS, ACS, AJCC, CDC and its National Center for Health Statistics, NCI, NCRA, NAACCR, and the Armed Forces Institute of Pathology. Current priorities for NCCCS include developing a roadmap between staging systems and establishing a national framework for cancer surveillance.

# **National Program of Cancer Registries**

CDC has worked to improve registry data nationwide since 1992, when Congress authorized the establishment of the National Program of Cancer Registries (NPCR) through the Cancer Registries Amendment Act (Public Law 102-515).<sup>36</sup> CDC provides funds to 45 states, 3 territories, and the District of Columbia to assist in planning or enhancing cancer registries, developing model legislation and regulations for programs to increase the viability of registry operations, setting standards for data, providing training for registry personnel, and helping establish computerized reporting and data processing systems.

CDC has contributed substantially to the development of data standards through its financial support of NAACCR, as well as by funding and developing EDITS, a software system that facilitates the coordination of data standards (see Chapter IV). In administering NPCR, CDC requires participating central registries to collect data items that conform to NAACCR's standards. NPCR staff also continue to maintain Registry Plus<sup>TM</sup>, a suite of publicly accessible free software programs made available by CDC to facilitate the implementation of NPCR.

To maximize the benefits of state-based cancer registries, CDC is implementing the NPCR-Cancer Surveillance System (CSS) for receiving, assessing, enhancing, aggregating, and disseminating data from NPCR-funded registries. This system of cancer statistics will provide valuable feedback to improve the quality and usefulness of registry data and monitor the impact of cancer prevention and control programs. For additional information on NPCR, visit the CDC/NPCR Web Site at: http://www.cdc.gov/cancer/npcr/index.htm.

#### North American Association of Central Cancer Registries

The American Association of Central Cancer Registries (AACCR) was established in 1987, and with the addition in 1995 of Canadian registries as members, the name was changed to the North American Association of Central Cancer Registries. Members are population-based cancer registries in the United States and Canada, national cancer and vital statistics organizations in both countries, and other organizations and individuals interested in cancer registration and surveillance. NAACCR is a professional organization that develops and promotes uniform data standards for cancer registration; provides education and training; certifies population-based registries for high-quality data; evaluates, aggregates, and publishes data from central cancer registries; and promotes the use of cancer surveillance data and systems for cancer in North America. NAACCR welcomes membership from cancer registries and other organizations or individuals that are concerned with the collection, analysis, and publication of data on cancer incidence.

# Surveillance Epidemiology and End Results Program

NCI's SEER Program has collected standardized data to measure progress in cancer prevention and control for more than 25 years. Established by a Federal mandate—the National Cancer Act of 1971—the SEER Program is an organizational descendent of the NCI-sponsored End Results Group (1956-72) and the Third National Cancer Survey (1969-71).

Seven population-based registries have provided data continuously since the SEER Program began in 1973: the States of Connecticut, Iowa, New Mexico, Utah, and Hawaii; and the Metropolitan Areas of Detroit and San Francisco-Oakland. In 1974-75, the regions of Seattle-Puget Sound and Metropolitan Atlanta were added. These areas, plus the rural Georgia region added in 1978, cover about 9.5 percent of the U.S. population. In 1992, the SEER Program added two additional regions in California—Los Angeles and San Jose-Monterey—bringing coverage of the U.S. population to 14 percent. These regions were selected for their epidemiologically significant population subgroups and, in fact, oversample minority populations in the United States. In 2001, four states were added—Kentucky, Louisiana, New Jersey, and the remainder of California—resulting in coverage of about 26 percent of the U.S. population.

The purpose of the SEER Program, as stated in the National Cancer Act legislation, is to collect, analyze, and disseminate data useful in the prevention, diagnosis, and treatment of cancer. The goals of the Program are to:

- Monitor annual cancer incidence trends to identify patterns of cancer occurring in population subgroups
- Provide continuing information on changes over time in the extent of disease (EOD) at diagnosis, trends in therapy, and associated changes in patient survival
- Promote studies to identify factors that can be studied and applied to achieve cancer prevention and control.

These goals illustrate that the aim of the SEER Program is providing cancer surveillance over time. As a result, changes in standards are carefully considered for their impact both on future data and compatibility with previous data.

Participating registries are required to submit data twice per year in a standard format using standardized definitions and codes (currently the *SEER Program Code Manual*, Third Edition, 1998,<sup>3</sup> and *SEER Extent of Disease-1998: Codes and Coding Instructions*, Third Edition).<sup>6</sup> However, the individual SEER registries have

not used standardized data collection methods or data management methods locally, and they differ in the extent to which they impose data requirements on the reporting facilities in their areas.

Standardized edits, developed by SEER and shared with participating registries, are applied to data submissions, and the results are returned to the participating registries.

SEER Program publications relating to data standards include:

- A series of eight self-instructional manuals for cancer registrars<sup>39</sup> covering abstracting, coding, terminology, anatomy, treatment, statistics, and other aspects of cancer registry operations. Book 8 in the series is a comprehensive list of drugs used in treating cancer and is the standard reference for drug-treatment coding rules.
- SEER Extent of Disease-1998: Codes and Coding Instructions, Third Edition.<sup>6</sup> This document includes site-specific codes and coding guidelines to describe spread of tumor in anatomic terms. EOD is a 10-digit code that includes 3 digits for size of tumor, 2 digits for tumor extension, 1 digit for lymph node involvement, 2 digits for the number of regional lymph nodes examined, and 2 digits for the number of positive regional lymph nodes. SEER always has collected EOD information and collapses this information into different staging schemes.
- The SEER Program Code Manual, Third Edition.<sup>3</sup> This manual includes comprehensive codes and coding guidelines for the data elements required by SEER.
- Comparative Staging Guide for Cancer.<sup>4</sup> This guide illustrates the relationships among EOD codes, the summary staging system, and the Third Edition of the TNM Staging System. A revision updating the comparative stagings to the Fifth Edition of the TNM Staging System is in development.
- Summary Staging Guide for the Cancer Surveillance, Epidemiology and End Results Reporting Program.<sup>9</sup> Originally published in April 1977, and most recently reprinted in July 1986, this is the standard for localized-regional-distant staging for tumors diagnosed between 1977 and 2000.
- SEER Summary Staging Manual 2000.<sup>10</sup> Published in 2001, is the standard for summary stage for cases diagnosed January 1, 2001, and after.

There is no charge for single copies of SEER Program publications. To place an order or to obtain further information, contact SEER at the address on page xiii, or go to the SEER Program Web Site at: http://seer.cancer.gov/Publications.

# World Health Organization

The World Health Organization (WHO), an agency of the United Nations, is responsible for publishing and maintaining the international standard for diagnosis coding systems. Selected publications include:

- International Classification of Diseases (ICD-9, the Ninth Revision), as modified by the Health Care Financing Administration<sup>13</sup>
- International Statistical Classification of Diseases and Related Health Problems (ICD-10, the 10th Revision)<sup>12</sup>
- International Classification of Diseases for Oncology.<sup>14, 15</sup>

These publications are world-standard diagnosis coding systems.

ICD-9 was adapted for use in the United States as the Clinical Modification of ICD-9<sup>13</sup> (ICD-9-CM), and is the current standard for coding medical record diagnoses in health information management departments in U.S. health care facilities. ICD-10 was implemented for coding causes of death on death certificates in the United States effective January 1, 1999.

The Second Edition of ICD-O became the standard for coding cancer diagnoses in the United States in 1992. An extensive revision of the morphology codes, especially the Lymphoma and Leukemia Section, was field-tested for the 1999 and 2000 diagnosis years, and the Third Edition of ICD-O<sup>15</sup> has been implemented for 2001 diagnoses.

WHO publications are sold through the following two agencies in the United States:

Q Corporation 49 Sheridan Avenue Albany, NY 12210 (518) 436-9686

College of American Pathologists 325 Waukegan Road Northfield, IL 60076 (800) 323-4040 http://www.cap.org/index.cfm

In the United States, the contact for further information on ICD-O is the Expert on Nomenclature and Coding at SEER (see address on page xiii).

# HISTORICAL BACKGROUND OF STANDARDS COORDINATION

Because the various standard-setting organizations use their data for different purposes, some data elements had different meanings, depending on the organization using the data. A long history of cooperation has been evident among organizations interested in cancer data to resolve the discrepancies between organizations in their interpretation of data elements.

The earliest standard setters were COC and SEER. The End Results Group, predecessor of SEER, published coding rules and guidelines as early as the 1950s; COC published its first data collection manual, the *Supplement on the Tumor Registry*, in conjunction with its *Cancer Program Manual 1981*. At that time, hospital-based cancer registries often used COC's recommended codes and coding rules, and SEER central registries used those of the SEER Program. The two systems were not always in agreement. As a result, COC and SEER began working together in the early 1980s to make the codes and definitions in their manuals consistent.

COC and SEER attempted to define one common set of data item definitions, field lengths, and codes for use by both SEER registries and hospital-based registries. By 1988, the collaboration resulted in the publication of both COC's *Data Acquisition Manual* and the *SEER Program Code Manual*, with data items and codes in substantial agreement. Having more congruent data sets allowed for easier data sharing and data comparisons, especially with the advent of personal computers that were sufficiently powerful to analyze large amounts of cancer data.

This achievement helped set precedents for cooperation in data management, and maintaining congruence whenever possible has continued to be a top priority for these two groups.

During the same period, the California Cancer Registry was developing a statewide automated system that allows facilities to report electronically to the state registry system. One region in California was a SEER registry at that time, and a large number of hospitals maintained COC-approved programs. To facilitate implementation of standards within its program, the California Cancer Registry requested that SEER and COC establish a formal committee to pursue data standardization and requested membership on this committee.

The function of that committee was transferred to NAACCR's Uniform Data Standards Committee (UDSC) when it was established in 1987. Membership was expanded to include all of the major standard-setting organizations and representation from registry software vendors and central registries. This Committee has made enormous progress toward standardization. A major success occurred when all of the participating groups agreed to implement the Second Edition of ICD-O simultaneously for tumors diagnosed in 1992 and later. In 1993, NAACCR convened a multidisciplinary conference to address the issue of collecting data on preinvasive cervical neoplasia, resulting in specific recommendations for member registries to cease collection of cervical carcinoma *in situ*. UDSC provides a national forum to discuss data issues and reach consensus on data standards. Given the extensive effort required to maintain uniform standards, in 2000, a subsidiary of UDSC, the Volume II Work Group, was formed to focus on the annual updates, revisions, and additions to compendiums of national standards.

CDC added another strong voice for standardization. CDC requires that the registries in 45 states, the District of Columbia, and U.S. Territories funded by NPCR use standard data items and codes. CDC is a sponsoring member of NAACCR, and has participated in committee activities of NAACCR. Through its contractor, CDC provides quality control activities for participants in NPCR and has facilitated the setting of standards and encouraged their adoption. The EDITS project described in Chapter IV is an example of the innovative approach CDC has supported.

At the time of this revision to Volume II, the major organizations agree in principle that their data standards will be consistent wherever possible. There are, however, areas where agreement has not been reached. These are discussed in detail in Chapter V. It also must be realized that standardization is not always desirable or feasible. For example, although the NAACCR standard for entry of dates is MM/DD/CCYY, SEER collects only month and year of birth date and date of death. SEER does not want to receive date of birth or death because of potential compromises to patient confidentiality, although individual SEER registries may collect this information.

Despite the progress made toward standardization and the near-universal agreement that standardization is desirable, much remains to be done. Implementation of existing standards is not uniform, and implementation of changes in standards is not always synchronized. SEER and COC will continue to publish separate coding manuals on different update schedules. Coding rules and rule interpretations sometimes are determined informally and documented marginally. Standardized data edits must be updated, maintained, and used by all registries.

In Canada, cancer registries at the provincial and territorial level joined together with Statistics Canada, a national agency, to form the Canadian Council of Cancer Registries. This process started in 1986 and led to the development of common national standards for the Canadian Cancer Registry, which were implemented with a reference date of January 1, 1992. A Data Quality Committee, which reports to the Council, is responsible for making recommendations to set national standards, and will review and monitor data quality and resolve any inconsistencies in procedures, coding, or other activities affecting data comparability.

NAACCR hopes that documenting existing standards, recommending standards where they do not yet exist, and publishing the results in a concise and authoritative form will enable registries and software providers to move forward in achieving comparable data that can be more widely used.

## Schedule of Revisions to NAACCR Standards Documents

The NAACCR Board of Directors established a Standards Implementation Task Force to review the current timeline for changes to data standards and to recommend guidelines for a new timeline that will meet the needs of the standard-setting organizations, central cancer registries, vendors, and reporting facilities. The Standards Implementation Task Force developed guidelines for **major** changes to be implemented on a 3-year cycle. All standard setters will adhere to the same 3-year cycle, implementation of the process began January 2003, with the next implementation date for major changes occurring on January 1, 2006 (i.e., then 2009, 2012, 2015, etc.). These changes will require the publication of a new Version of the NAACCR Volume II Data Dictionary and Data Standards (e.g., from Version 10.x to Version 11.0). **Minor** changes would be implemented on an annual cycle. These changes will be published in an update of the current Version of the NAACCR Volume II Data Dictionary and Data Standards (e.g., Version 10.1 [*Exception:* An updated Version will not be published the year a new Version is published, minor changes will be included in the new Version]). The intent is to allow the ability to fix errors and clarify codes or add new codes should they be necessary during the interval between the scheduled major revisions and updates. See the *Standards Implementation Guidelines*<sup>42</sup> for definitions of major and minor changes and additional information.

# **CHAPTER III**

# STANDARDS FOR TUMOR INCLUSION AND REPORTABILITY

Due to recent efforts by standard-setting organizations, facility-based registries and population-based central registries now follow nearly identical standards for determining tumors that are reportable and are to be included in the registry; however, some differences remain. COC stipulates the tumors that must be included in approved facility registries, while most population-based registries, at a minimum, follow the standards set by SEER or NPCR. The *Cancer Program Standards*,<sup>28</sup> the COC *FORDS* Manual,<sup>2</sup> SEER code manuals,<sup>3, 6</sup> and the NPCR Program Announcement<sup>40</sup> should be consulted for more details.

Standards for tumor reportability are defined by the following criteria:

## **Reference Date**

The reference date is the effective date cancer registration starts in a specified at-risk population or in a specific facility. It is not the date the registry is organized or the date work begins. Tumors diagnosed on or after the reference date must be included. The reference date typically begins on January 1 of a calendar year, but sometimes it is another date.

## Residency

For a population-based registry, it is essential to include all tumors occurring in the at-risk population, and rules must be in place for determining the members of that population. The goal is to use the same rules for the patient's demographic data at the time of diagnosis as those used by the Census Bureau in enumerating the population. For example, a population-based registry must have rules for determining residency of part-year residents, institutionalized persons, homeless persons, military personnel, and students. See the *SEER Program Code Manual*<sup>3</sup> for specific instructions.

NAACCR recommends that population-based registries include in their database tumor reports of non-residents from facilities in their catchment areas to:

- Share tumor information that otherwise may go unreported with the resident's population-based registry.
- Facilitate death clearance and other record linkages.
- Allow preparation of complete and accurate reports to individual facilities.

Hospital-based registries are less concerned with residency of the patient than the reason for admission, and hospital registries might not collect data for certain categories of patients that the central registry must include, such as patients admitted to a hospice unit or transient patients who receive interim care to avoid interrupting a course of therapy. Also, COC does not require complete abstracting of tumors that are "nonanalytic" for the facility. Therefore, for the central registry, clear rules that are well documented, widely distributed, and accepted are essential to prevent missed case reports (source records).

# **Reportable List**

COC, NPCR, and SEER have achieved greater consensus on reportable tumors in the past few years (see Table 1). For all tumors diagnosed from January 1, 1992, through December 31, 2000, all three standard setters required the inclusion of all neoplasms in the International Classification of Diseases for Oncology, Second

Edition<sup>15</sup> (ICD-O-2) with a behavior code of 2 or 3 (*in situ* or malignant), with the exception of squamous cell and basal cell carcinoma of the skin and carcinoma *in situ* of the cervix uteri since 1996. (See the CARCINOMA *IN SITU* OF THE CERVIX, CIN, AND THE BETHESDA SYSTEM Section later in this Chapter).

For all tumors diagnosed on or after January 1, 2001, all three organizations require the inclusion of all neoplasms in the International Classification of Diseases for Oncology, Third Edition<sup>14</sup> (ICD-O-3) with a behavior code of 2 or 3 (*in situ* or malignant), with the exception of squamous cell and basal cell carcinoma of the skin, prostatic intraepithelial neoplasia (PIN) III, carcinoma *in situ* (CIS) of the cervix, and cervical intraepithelial neoplasia (CIN) III. Code M9421 (juvenile astrocytoma, pilocytic astrocytoma, or piloid astrocytoma), with a behavior code of 1 (borderline) in ICD-O-3, is reportable as M9421/3. Prior to 2003, COC considered basal and squamous skin cancers that were AJCC stage group II or higher at diagnosis as reportable.

In addition, all three organizations require the inclusion of all non-malignant primary intracranial and central nervous system (CNS) tumors diagnosed on or after January 1, 2004. Specifically, non-malignant primary intracranial and CNS tumors of any morphology in ICD-O- $3^{14}$  having a behavior code of 0 or 1 (benign/ borderline) occurring in the following sites: brain, meninges, spinal cord, cranial nerves and other parts of the CNS, pituitary gland, pineal gland, and craniopharyngeal duct (see Table 2).

# In Situ/Invasive

It is important to distinguish between the morphologic condition of *in situ* as it is represented in ICD-O-2 or ICD-O-3 behavior codes and Tis as it is defined for the purpose of prognostic staging in the *AJCC Cancer Staging Manual*. Some morphologic and disease descriptive terms that are invasive in ICD-O-2/ICD-O-3 or localized in the SEER *Summary Staging Guide*/SEER *Summary Staging Manual 2000* are Tis in the *AJCC Cancer Staging Manual*. Some examples are:

- Paget's disease of the nipple (8540/3) (an "invasive" code in ICD-O-2 and ICD-O-3) with no underlying tumor is classified as Tis in AJCC Sixth Edition.
- For colon/rectum, "invasion of the lamina propria" (intramucosal) with no extension through the muscularis mucosae into the submucosa is classified as Tis according to AJCC Sixth Edition but localized in SEER Summary Stage 2000.

Whether a tumor diagnosis is *in situ* or invasive is important because it affects how the tumor will be reported in published statistics. Some tumors staged by central cancer registries using SEER Summary Stage or SEER EOD codes as "localized" can be classified as Tis or Stage 0 when coded according to AJCC or when EOD codes are converted to AJCC. Some tumors classified as invasive in the behavior code can be classified as Tis or Stage 0 when coded according to AJCC Sixth Edition. These differences should be considered when data are being compared. For more information on differences in staging classifications and current activities toward improving the situation, see Chapter V.

# **Multiple Primary Rules**

The method used for counting tumors affects the comparability of cancer rates among registries. It is important that identical rules have been used for counting multiple tumors in the patient—whether in the same organ, on opposite sides of paired organs, in different sites or subsites—and whether they were diagnosed at the same or different times. SEER rules are the *de facto* standard in the United States for both central and hospital-based registries. See the *SEER Program Code Manual*<sup>3</sup> for details.

SEER rules are not identical to the international standard recommended by the International Agency for Research on Cancer (IARC) and the International Association of Cancer Registries (IACR).<sup>37</sup> The IARC rules have the effect of defining fewer cases than do the SEER rules.

The following addition to SEER multiple primary rules was reviewed by UDSC and adopted on April 26, 1994, effective with tumors diagnosed in 1995 and later.

**EXCEPTION:** If there is an *in situ* followed by an invasive cancer at the same site more than 2 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 (*SEER Program Code Manual*, Third Edition, page 11).

This important rule affects how the tumor will be counted in published statistics. With the exception of bladder, *in situ* tumors are not usually included in published incidence rates. Without the reporting of these invasive cancers, for example, rates of invasive breast cancer would be underreported. COC, with an emphasis on clinical data, has not adopted this exception to the general rule.

# CARCINOMA IN SITU OF THE CERVIX, CIN, AND THE BETHESDA SYSTEM

The term "pre-invasive cervical neoplasia" refers to carcinoma *in situ* of the cervix and conditions viewed as equivalent to it or on a continuum with it. Diagnostic terminology for pre-invasive cervical neoplasia has changed significantly over time, from the four-tiered system of dysplasia and carcinoma *in situ*, to the three-tiered system of CIN, to the two-tiered Bethesda System, with high- and low-grade squamous intraepithelial lesions (SIL). In the past, cancer registries generally considered carcinoma *in situ* of the cervix reportable, but they differed in which of these other terms they considered synonymous with carcinoma *in situ* and hence reportable. Consequently, data were not comparable over time or across registries.

NAACCR convened a multidisciplinary working group in April 1993 to review the problem and make recommendations for its membership. The recommendation was that "population-based registries discontinue routine collection of data on pre-invasive cervical neoplasia unless there is strong local need and interest and sufficient resources are available to collect all [high-grade squamous intraepithelial lesions] and its equivalent terms."<sup>33</sup>NAACCR and NPCR adopted this recommendation at that time. SEER and COC adopted it effective for cases diagnosed January 1, 1996 forward.

#### **Ambiguous Terminology**

In most circumstances, the diagnosis of cancer, as recorded in the patient's medical record, clearly is synonymous with reportable cancer. However, in those situations where the physician is not certain of the diagnosis, the associated terminology in the medical record reflects that uncertainty and is ambiguous. COC and SEER are in agreement in regard to the list of terms considered as diagnostic of cancer and a list of terms not considered as cancer. These terms are shown in Table 1.

	COC	SEER	NPCR
Reportable	<ol> <li>Behavior code of 2 or 3 in ICD-O-3.</li> <li>Non-malignant (behavior codes 0 and 1)</li> </ol>	1. Behavior code of 2 or 3 in ICD-O-3.	1. Behavior code of 2 or 3 in ICD-O-3 (includes VIN III, VAIN III, AIN III).
Diagnoses	primary intracranial and central nervous system tumors, including juvenile astrocytoma (M9421/3)* for primary sites	2. Non-malignant (behavior codes 0 and 1) primary intracranial and central nervous system tumors,	<ol> <li>Non-malignant (behavior codes 0 and 1) primary intracranial and</li> </ol>
On or after 1/1/2005	as defined in Table 2.	including juvenile astrocytoma (M9421/3)* for primary sites as defined in Table 2.	central nervous system tumors, including juvenile astrocytoma (M9421/3)* for primary sites as defined in Table 2.
	1. Skin cancers (C44) with histology 8000-8110 (after 1/1/2003); prior to that date, only AJCC stage groups 0 and I tumors in this group were not reportable).	1. Skin cancers (C44) with histologies (8000-8005, 8010-8046, 8050-8084, 8090-8110) other than those listed above.	1. Skin cancers (C44) with histologies (8000-8005, 8010- 8046, 8050-8084, 8090-8110) other than those listed above.
Exceptions	2. CIS of the cervix and CIN III (after 1/1/96).	2. CIS of the cervix and CIN III (after 1/1/96).	2. CIS of the cervix and CIN III.
(not reportable)	3. PIN III (after 1/1/96).	3. PIN III (after 1/1/2001).	3. PIN III (after 1/1/2001).
	<ol> <li>VIN III (after 1/1/96).</li> <li>VAIN III (after 1/1/96).</li> </ol>		
	<ul><li>6. AIN (after 1/1/96).</li><li>Follows SEER rules with the following</li></ul>	Follows SEER rules.	Follows SEER rules.
Multiple Primary Rules	exception: when there is an <i>in situ</i> followed by an invasive cancer at the same site more than 2 months apart, do not report the invasive cancer as a second primary if stated by the physician to be a recurrence.	Follows SEEK fules.	Follows SEEK fules.
Ambiguous Terminology Considered as Diagnostic of Cancer	apparent(ly) appears comparable with compatible with consistent with favors malignant appearing most likely presumed probable suspect(ed) suspicious (for) typical of	apparent(ly) appears comparable with compatible with consistent with favors malignant appearing most likely presumed probable suspect(ed) suspicious (for) typical of	Not addressed.
	Exception: if the cytology is reported as "suspicious" and neither a positive biopsy nor a physician's clinical impression supports the cytology findings, do not consider as diagnosis of cancer.	Exception: if the cytology is reported as "suspicious" and neither a positive biopsy nor a physician's clinical impression supports the cytology findings, do not consider as diagnosis of cancer.	
Ambiguous Terminology NOT Considered as Diagnostic of Cancer	cannot be ruled out equivocal possible potentially malignant questionable suggests worrisome	cannot be ruled out equivocal possible potentially malignant questionable suggests worrisome	Not addressed.

Table 1. NAACCR Layout Version 10.2: Comparison of Reportable Cancers: COC, SEER, and NPCR.
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\* Juvenile astrocytomas should be reported as 9421/3.

Table 2. Primary Site Codes for Non-Malignant Primary Intracranial and Central Nervous System Tumors<br/>(non-malignant primary intracranial and central nervous system tumors with a behavior code of 0 or 1<br/>[benign/borderline] regardless of histologic type for these topography codes).

	Topography		
Codes	Description		
	Meninges		
C70.0	Cerebral Meninges		
C70.1	Spinal meninges		
C70.9	Meninges, NOS		
	Brain		
C71.0	Cerebrum		
C71.1	Frontal lobe		
C71.2	Temporal lobe		
C71.3	Parietal lobe		
C71.4	Occipital lobe		
C71.5	Ventricle, NOS		
C71.6	Cerebellum, NOS		
C71.7	Brain stem		
C71.8	Overlapping lesion of brain		
C71.9	Brain, NOS		
	Spinal Cord, Cranial Nerves, and Other Parts of the Central Nervous System		
C72.0	Spinal cord		
C72.1	Cauda equina		
C72.2	Olfactory nerve		
C72.3	Optic nerve		
C72.4	Acoustic nerve		
C72.5	Cranial nerve, NOS		
C72.8	Overlapping lesion of brain and central nervous system		
C72.9	Nervous system, NOS		
	Other Endocrine Glands and Related Structures		
C75.1	Pituitary gland		
C75.2	Craniopharyngeal duct		
C75.3	Pineal gland		

## CHAPTER IV

#### **RECOMMENDED DATA EDITS AND SOFTWARE COORDINATION OF STANDARDS**

#### Definitions

"Data edits" refer to computer software algorithms that check the content of data fields against an encoded set of acceptable codes and subsequently provide feedback on the quality of the data. Data edits verify that only acceptable values are used for codes and, more importantly, enforce relationships between the codes in related data items. Data edits can apply pass/fail criteria to data, so that a particular code or set of entries is determined to be either correct or incorrect. Incorrect data will have to be corrected to pass subsequent edits. Other types of edits indicate possible (or probable) errors that require human review for resolution. Many of these possible errors are tied to over-ride flags that indicate that the data in a record (or records) have been reviewed and, while unlikely, are correct.

Generally, there are three types of edits:

- Single-field edits or item edits are those that look at only one data field at a time. For example, an edit of the item "Sex" would verify that only valid values are used in the field.
- Interfield edits or multifield edits are those that compare the codes of a data item with those in other related data items. For example, a common interfield edit compares the code for "Sex" with the code for "Primary Site," and identifies female prostate cancer as an error.
- Interrecord edits or multirecord edits compare data on more than one record, commonly for those situations where a patient has multiple tumors. They compare the code of a data item in one record for a particular tumor with the same data item in another record or tumor. For example, an interrecord edit compares sequence numbers in multiple tumors to ensure that they have been assigned in chronological order for the patient's cancers.

#### Challenges

There are at least six challenges to the standardization of data edits across central and hospital-based cancer registries. These include:

- Registry systems that encode an edit from standard specifications may be written in different computer languages, with possible differences in translation detail.
- Each implementation of an agreed-upon standard specification may be programmed differently, despite intent to encode a standard meaning.
- Complete edits are not always performed at the time of data entry.
- Documentation of the edit algorithms often is difficult for both data analysts and data collectors to obtain and use.
- Consolidated data collected via different data entry tools may encourage "apples" and "oranges" to be equated, without the users' knowledge.

When standards change, synchronized implementation is difficult, due to the release schedules of software providers and their limited ability to respond to changes at a given time.

Comparable results can only be reasonably expected when identical edits are applied to cancer registry data.

#### The EDITS Software

The EDITS Software Project began with an informal discussion about promoting and supporting data processing standards after a 1990 meeting of the NAACCR Data Evaluation and Publication Committee. A small group of registry operators, software producers, and data consumers identified a missing element of standard setting: an executable version of a standard that could be applied directly to data in a variety of processing scenarios without reinterpretation by programmers. Producers of cancer registry software who intended to adhere to a published standard had to write their own computer code to implement the edit-checking algorithms. The solution would need to be flexible in many dimensions to accommodate the many technical, operational, scientific, economic, and agency considerations that determine the cancer registry milieu.

Although EDITS handles single-field and interfield type edits routinely and interactively, the software's ability to process interrecord edits is limited. CDC has developed EDITS to accommodate interrecord edits. This edit typically is applied as a freestanding batch program and run at the time of data submission.

The EDITS software consists of three main components: EditWriter, the EDITS Application Program Interface (API), and the Generic EDITS Driver Program (GenEDITS).

#### ✤ EditWriter

The EditWriter is a versatile and complete development environment for defining, testing, documenting, and distributing data standards. It also provides a means of maintaining the definition of a standard as it matures and changes over time. Data checking can be as complete and as complicated as the applications require.

The output of EditWriter is the EDITS Metafile, a compiled database that contains all of the logic, tables, and constant values needed to check fields of data for validity. Single-field and interfield checks are included in the NAACCR Metafile. Although EditWriter is an MS-DOS program, the metafiles that it produces can be copied and used on other operating systems, such as UNIX. The metafiles also can be used on hardware platforms other than the PC.

#### EDITS Application Program Interface

The EDITS API can be incorporated into programs of many descriptions, including programs for interactive data entry, after-the-fact verification of data, recoding, reformatting, and vertical or horizontal subsetting. Any language product for Windows should be able to use the EDITS API. Additionally, applications written in C and compiled with modern compilers for MS-DOS, UNIX, and VAX/VMS operating systems can include the EDITS engine. The EDITS API is distributed as a Windows Dynamic Link Library and as C source code.

#### ✤ Generic EDITS Driver Program

GenEDITS is a configurable application for editing any data file with any EDITS Metafile. GenEDITS is the fastest way to apply standard edits to data and obtain a report of data errors. Because GenEDITS already incorporates the EDITS API, no programming is required.

#### The EDITS Language

Using the EDITS language—a simplified programming language designed to validate data—specifies the algorithms that check data. The language includes a collection of powerful and specialized built-in functions that often reduce the complete validation of a data item to a single program statement. When complicated data relationships exist within a record, the EDITS language can express an arbitrarily complex validation schema, including multiple fields, multiple table look-ups, nested control statements, local and global variables, and external functions.

For additional information about EDITS or to download the EDITS software, see CDC's Division of Cancer Prevention and Control Web Site at: http://www.cdc.gov/cancer/edits/editintr.htm.

#### The EDITS Metafile

EDITS Metafiles contain everything needed to edit a data file, except the data. Metafiles provide portability of edits; in that the same edits can be applied to different data formats for different purposes. EDITS Metafiles are created and modified using EditWriter. The key components of a metafile include: agencies, data dictionary, record layouts, edits, edit sets, error messages, and user look-up tables.

#### SEER\*Edits

For many years, the SEER Program has maintained a library of standardized edits written in IBM COBOL,<sup>34</sup> which it applied to data submissions from the participating SEER registries. Over the years as experience and expertise increased, SEER has fine-tuned and expanded the edits and has made these edits available to SEER and other registries. In addition, the logic of the SEER edits has been used as the foundation for the EDITS project where SEER is the source of standard for the item or items.

As more and more computer processing moved away from the mainframe environment, the SEER Program decided to reprogram their edits in C++. This change has allowed the SEER edit engine to be ported to and compiled on a variety of hardware platforms. The edit engine includes the entire field, interfield, and interrecord edits in the COBOL edits plus new and revised edits needed because of the introduction of ICD-O-3. The SEER\*Edits package replaces the COBOL edits and the COBOL edits are no longer being maintained. SEER\*Edits can be used as a stand-alone package for the SEER areas to use before submission of data to SEER, or the edits can be incorporated individually by SEER registries for use in their data entry programs or routine editing of data. Data are input into the stand-alone version of SEER\*Edits in NAACCR format. The SEER\*Edits package also includes report-generating functions including manipulation of errors to facilitate data correction, a follow-up report, and a surveillance report. Any change made to the SEER\*Edits package also is made to the SEER metafile for the EDITS project and vice versa to keep them synchronized.

### NAACCR Standard Edits and the NAACCR Metafile

NAACCR has made increased standardization of data edits a priority, facilitated by the EDITS software, which provides a mechanism for standardized, transportable, and updateable edits to be provided through a "public library." The goals are to help limit standards proliferation when there is no compelling need to be different, and to provide comprehensive public documentation in a current and readily accessible form in those instances where standards must differ.

The NAACCR Metafile is a comprehensive database of cancer registry standards and consists of a collection of tables that contain all the information needed to test data fields for validity and acceptability. The NAACCR Metafile specifically includes the following: look-up tables, translation tables, choice lists, data dictionary of standard fields, local field name table, error messages, executable single- and multifield validation logic, text

descriptions of edits, sets of fields defining standard records, standard-setter list, description of local data storage, data-entry help, standards documentation text, EDITS system help, and EDITS language reference.

NAACCR first made standard edits available in 1996. These edits corresponded to its 1995 record layout and data dictionary, as Volume IV in its Standards series.<sup>29</sup> Since that time, NAACCR has posted standard edits on the Internet that correspond to the annual record layouts and data dictionaries. For example, "Revised Version 6 Metafile--NAACCR6D" refers to the current standard edits in the NAACCR Version 6 record layout. The "D" notation indicates the fourth revision to the Version 6 record layout standard edits. The hardcopy Volume IV has been discontinued in favor of Internet publication. The EDITS Software with general instructions and various current and previous metafiles containing the most recent and historical public standards for cancer registry data are available on the NAACCR Web Site at: http://www.naaccr.org/ Standards/Edits.html.

## CHAPTER V

### **UNRESOLVED ISSUES**

Despite the progress made toward data standardization, some issues remain unresolved. These issues are described in detail below. UDSC will continue to seek consensus on unresolved issues. Before new standards can be agreed upon, all interested parties must be provided sufficient time to study the proposals. Once UDSC approves new standards, there must be adequate time for implementation.

All members are encouraged to present suggestions or comments on proposed changes to the standards to UDSC. The NAACCR Web Site, http://www.naaccr.org, provides the name of the Committee Chair and forms for proposing additions or revisions.

#### **Record Layouts:**

Ten versions of the NAACCR layout have been released. All registries should begin using Version 10.2 in January 2005:

- Version 10.2 (dated March 2004)
- Version 10.1 (dated March 2003)
- Version 10 (dated March 2002)
- Version 9.1 (dated March 2001)
- Version 9 (dated May 2000)
- Version 8 (dated April 1999)
- Version 7 (dated April 13, 1998)
- Version 6 (dated January 23, 1998, and as slightly revised, dated March 20, 1998)
- Version 5.1 (dated March 12, 1997)
- Version 5 (dated April 10, 1996)
- ✤ Version 4 (dated 1994).

Please refer to Table 3 on the following page for more detail.

All versions of the NAACCR layout are compatible, but information is likely to be lost during a conversion. CDC and NAACCR are preparing standardized conversion programs between the versions.

Standards for Tumor Inclusions, Reportability, and Multiple Primary Rules are in Chapter III.

NAACCR	Release Date	Effective Date*	Reference Manuals Accommodated	NAACCR Metafile Version
Version 4	02/14/1994	01/01/1994	COC/ACOS Data Acquisition Manual, 1994 SEER Program Code Manual, 1992 WHO ICD-O-2, 1990 SEER Summary Staging Guide, 1977 AJCC Staging Manual, Fourth Edition, 1992 SEER Extent of Disease Manual, 1992	Metafile Version 4
Version 5	04/10/1996	01/01/1996	COC/ <i>ROADS</i> , 1996 SEER Program Code Manual, 1992 WHO ICD-O-2, 1990 SEER Summary Staging Guide, 1977 AJCC Staging Manual, Fourth Edition, 1992 SEER Extent of Disease Manual, 1992	Metafile Version 5
Version 5.1	03/12/1997	01/01/1997	Same as Version 5	Metafile Version 5
Version 6	01/23/1998 Rev 3/20/1998	01/01/1998	COC/ <i>ROADS</i> , 1996, Rev. 1998 SEER Program Code Manual, 1998 WHO ICD-O-2, 1990 SEER Summary Staging Guide, 1977 AJCC Staging Manual, Fifth Edition, 1997 SEER Extent of Disease Manual, 1998	Metafile Version 6
Version 7	04/13/1998	01/01/1999	Same as Version 6	Metafile Version 7
Version 8	03/30/1999	01/01/2000	Same as Versions 6 and 7	Metafile Version 8
Version 9	05/15/2000	01/01/2001	COC/ <i>ROADS</i> , 1996, Rev. 1998 SEER Program Code Manual, 1998 WHO ICD-O-3, 2000 SEER Summary Staging Manual, 2000 AJCC Staging Manual, Fifth Edition, 1997 SEER Extent of Disease Manual, 1998	Metafile Version 9
Version 9.1	03/21/2001	01/01/2002	Same as Version 9	Metafile Version 9
Version 10	03/20/2002	01/01/2003	COC FORDS SEER Program Code Manual, 2003 WHO ICD-O-3, 2000 SEER Summary Staging Manual, 2000 AJCC Staging Manual, Sixth Edition, 2002 Collaborative Staging Manual and Coding Instructions, Version 1.0 (implementation 01/01/2004)	Metafile Version 10
Version 10.1	03/2003	01/01/2004	Same as Version 10	Metafile Version 10 (most recent)
Version 10.2	03/2004	01/01/2005	Same as Version 10	Metafile Version 10 (most recent)

#### Table 3. Record Layout Table With References.

Bolded text indicates changes from previous version.

\* Either the date of diagnosis or year first seen for this cancer may have been used by some standardsetters. Refer to the Data Dictionary or to the standard-setter reference manuals for clarification of date requirements.

#### County--Current (item 1840)

County--Current was an item in the COC data set prior to 2003. Codes used may have varied among facilities for reasons described in the discussion of County at DX, item 90. Users of pooled data should ascertain what codes were used for this item.

#### County at DX (item 90)

NAACCR has adopted the Federal Information Processing Standards (FIPS) codes for county as the standard in this volume (see Appendix A for codes). However, standards for codes used vary somewhat by standard setter as follows:

- The SEER Program requires the use of FIPS codes for counties in the United States, plus the special code 999. Because SEER collects only cases of residents of the reporting areas, no codes are needed for SEER registries other than the codes for the counties in their areas.
- COC requires the use of FIPS county codes as their standard, plus the special codes 998 and 999. However, the *FORDS Manual* also provides for use of geocodes for countries of residence outside the United States and Canada to be used in this field.
- NPCR requires the use of FIPS codes for counties in the United States, plus the special code 999, starting with cancers diagnosed on or after January 1, 2002. Prior to 2002, NPCR recommended the use of FIPS county codes.
- ✤ NAACCR recommends the use of FIPS codes.

For cancers diagnosed prior to 2002, the use of FIPS codes was not universally adopted. For this reason, users of data should determine which codes were used for coding County at DX in a particular file, because no field indicating "County at DX Coding System" is included in the NAACCR layout.

#### Hispanic Ethnicity (items 190-210)

There is agreement on the standard data item "Spanish/Hispanic Origin" and its codes. However, there has been substantial variation among registries in how the Spanish or Hispanic origin is determined. Procedures for determining ethnicity include:

- Recording ethnicity from information in the medical record.
- Recording ethnicity based on all information available, including the surname, birthplace, or stated ethnicity.
- Recording ethnicity based on a manual or computer matching of surname against a list of Spanish surnames that, in most cases, is based on the 1980 Census. Some registries also perform an additional manual or computer match on the maiden name.
- Recording the ethnicity based on the application of a computer algorithm to surnames to determine ethnicity.

Population-based registries must attempt to categorize their cases using a method that best approximates the method used by the Census Bureau to determine ethnicity of the population denominators, but a standard method has not been determined. NAACCR's UDSC has discussed the issue extensively, and a subcommittee convened a workshop in Atlanta, GA, in January 1996. A report was prepared and is available on the

NAACCR Web Site (http://www.naaccr.org) under the heading "Epidemiologic Reports." In 1999, a research group was formed from representatives of NAACCR to address issues of definition and to produce comparable data for Hispanic ethnicities across the United States. The group operates under the auspice of the Data Evaluation and Publications Committee. Results from the survey of registry practices are available on the NAACCR Web Site.

Based on these discussions, NAACCR has added fields for Computed Ethnicity and Computed Ethnicity Source, and has clarified how the code for Spanish/Hispanic Origin is to be determined. Registries continue to use different methods of coding ethnicity, but users of the data should be able to determine how coding was done in a particular file if the standard codes are used. See the descriptions and notes for items 190-210 for details.

#### Name--Last (item 2230)

The COC *FORDS Manual* allows embedded spaces, hyphens, apostrophes, and punctuation in the last name field. NAACCR standards allow no embedded spaces and punctuation, except hyphens. Neither COC nor NAACCR standards allow the last name field to be blank.

#### Name--Maiden (item 2390)

The COC *ROADS Manual* allowed embedded spaces, hyphens, apostrophes, and punctuation in the maiden name field. NAACCR standards allow no embedded spaces and punctuation, except hyphens. Both COC and NAACCR standards allow the maiden name field to be blank. This data item is not collected in the COC *FORDS Manual*.

#### Occupation and Industry (items 270-330)

Most population-based registries have found the collection of usual occupation and industry data to be difficult and of limited utility. Traditionally, no consensus on data items and codes for occupation and industry had been achieved. In 1992, the Cancer Registries Amendment Act required collection of occupation or industry data to the extent available in the medical record by central registries funded by NPCR.<sup>36</sup> In response to this mandate, CDC sponsored a meeting of experts in occupational health and cancer epidemiology in 1995. Recommendations from the meeting resulted in the adoption of data items and codes by the NAACCR UDSC in August 1995.<sup>25</sup> These agreed-upon standards were included in Versions 6 and later of NAACCR's data standards.

Data on usual occupation and industry are unavailable in an unknown, but significant, proportion of medical records. Additionally, even when available, the quality of the data in the medical record is generally untested and often limited to less useful information such as "retired." Concurrently, this information generally is available in text format on death certificates and, in some states, on the associated state mortality data tapes. Some state mortality data tapes in addition to the text data also contain the associated occupation and industry codes. Software for the automated coding of the text data is available from the Division of Safety Research, National Institute for Occupational Safety and Health, CDC. Regardless, much work remains to be done to improve the availability and capture of this potentially important information.

NAACCR will continue to discuss the quality and completeness of occupation and industry data and will reconsider the inclusion of occupation and industry in its recommended data sets.

### RX Summ--Rad to CNS (item 1370)

SEER and COC had different requirements for this item. SEER no longer collects it for cases diagnosed 1998 and later; however, they retain the codes for older years' cases, and also convert the data into an appropriate code in the RX Summ--Radiation field.

#### Sequence Number (items 380 and 560)

As discussed in Chapter III, SEER, NPCR, and COC have different standards for determining tumors that are reportable and are to be included in the registry. In addition to collecting these required tumors, some registries collect and assign sequence numbers to other tumors such as cervix carcinoma *in situ* or PIN III.

Two sequence number data items, one assigned by the reporting facility, Sequence Number--Hospital [560], and one assigned by the central registry, Sequence Number--Central [380], are now in use. The time period of both Sequence Number data items is a person's lifetime, although with earlier definitions of Sequence Number--Central [380], central registries historically assigned the numbers from the reference date of the registry.

Numerous operational issues, such as storage of multiple facility-specific sequence numbers, appropriate linkage rules, and feedback of data to hospitals, have arisen because of policy differences from state to state. When attempting to use the Sequence Number--Central to identify individuals who have had only one cancer in their lifetime, it is important to realize the definitions used to make that determination vary and that cases may have been handled inconsistently in different systems.

#### Stage, TNM, and EOD (items 760-830, 880-1070)

Currently, five major staging schemes are widely used in cancer registries throughout the United States. The schemes differ in complexity, structure, purpose, definitions, and rules. The five schemes are:

#### ✤ The American Joint Committee on Cancer's TNM System

In its Sixth Edition, the *Cancer Staging Manual* includes a clinically oriented, site-specific staging system that consists of a separate category for the tumor, nodes, and metastases. The TNM categories then are grouped by stage, from 0 to IV. COC standards for approved cancer programs require that the medical record contain the AJCC stage assigned by the managing physician.

#### **\*** SEER Extent of Disease

This site-specific 10-digit coding scheme<sup>6</sup> is required for SEER registries and is used by some other state and central registries as well. EOD was designed to allow collapse of the codes into the stage groupings of several different staging systems, including AJCC stage group.

#### ✤ SEER Summary Stage

This site-specific single-digit coding scheme is required for NPCR registries, and is used by some SEER registries as well. In addition, COC requires the coding of SEER Summary Stage when a corresponding AJCC TNM site code scheme is not available. There are two related data items: SEER Summary Stage 1977 [760] and SEER Summary Stage 2000 [759]. Cancers diagnosed on or after January 1, 2001, should be assigned a summary stage according to the *SEER Summary Staging Manual, 2000*,<sup>10</sup> and the code should be reported in the SEER Summary Stage 2000 [759] data item. Cancers diagnosed before January 1, 2001, should be assigned a summary stage according to *Summary Stage Guide, Cancer Surveillance Epidemiology and End Results Reporting, SEER Program, April 1977*,<sup>9</sup> and the code should be reported in the SEER Summary Stage 1977 [760] data item (see NAACCR Guidelines for Implementation of SEER Summary Stage 2000).

#### ✤ SEER Historic Stage

When SEER stage data are published, the stage categories used are derived from categories used by an earlier program, the End Results Group. The categories are not identical to those in the SEER Summary Stage. However, the Historic Stage variable has been defined consistently over time to facilitate trend analyses.

### Collaborative Stage

The initial focus was to develop a translation between the TNM staging system of the AJCC and the SEER Summary Staging System. The translation would eliminate duplicate data collection by registrars reporting to clinical and epidemiologic registries, address the concerns of clinicians for more clinically relevant data as well as the public health sector's concerns about data reproducibility over time, and provide a higher degree of compatibility between the systems that would expand data-sharing opportunities.

The Collaborative Stage (CS) Data Set is a combination of data items (most of which have traditionally been collected) that include tumor size, extension, lymph node status, metastatic status, evaluation fields that describe the hierarchy of the data collected, and site-specific factors. This unified data set for cancer reporting has an algorithm that derives three different staging systems and resolves staging rule differences. The three systems are AJCC TNM, SEER Summary Stage (SS) (1977 and 2000), and SEER EOD. AJCC TNM staging provides forward flexibility and clinical utility. SEER EOD provides longitudinal stability for epidemiological studies, and SEER Summary Stage provides a population surveillance staging system.

Collaboration among the participating organizations has resulted in resolution of the timing rule and standardized staging rules for one staging information collection model. The timing rule going into effect on January 1, 2004, will be: "use all information through the first course of surgery or 4 months, whichever is longer." This timing rule change will allow the CS Data Set to capture "best stage" combining clinical and pathologic data. SEER currently uses the "4-month rule," and this collaboration brings both SEER and AJCC to one standard. Other rule modifications have been made and are printed in the "site-specific" chapters.

The CS model will improve the quality of data being collected. Uniform rules and standardized training will make it easier for cancer registry personnel to complete staging tasks.

These schemes were designed for different purposes at different times, and are not easily compared. There have been several editions of the *TNM Manual*, and implementation has not been synchronized. SEER has published the *Comparative Staging Guide for Cancer*<sup>4</sup> as an attempt to present comprehensive, site-specific comparisons of the schemes to aid in data collection and interpretation. This guide covers the major cancer sites of colon and rectum, lung and bronchus, breast, female genital, prostate gland, and urinary bladder. According to the guide:

- Changes over time in methods of cancer screening, diagnosis, staging, and treatment have affected the distribution of stage of disease.
- Changes over time in the classification schemes themselves can complicate data analysis and obscure the meaning of time trends. Various other staging schemes also are in use. Several oncology subspecialties have developed staging systems applying to a limited number of cancer sites.

For these reasons, comparing cancer registry data by stage over time or across registries, or using pooled data collected by different registries applying different staging schema, is problematic<sup>4</sup> (page I.3).

The lack of comparability among these systems causes major problems for those collecting the data and for users of the data. For example, hospital cancer registrars often are required to code stage information using more than one scheme to meet requirements of different standard-setting organizations. This increases the training needed for staff and the time needed to code each case. Users of the data may be unaware that the

same term may be defined differently in the schemes, and that data cannot be compared easily. For example, the category of *in situ* carcinoma of the colon includes different cases in TNM and SEER historic stage.

For a discussion of staging issues that affect rules for case inclusion and reportability, see Chapter III, especially the paragraphs "*In Situ*/Invasive" and "Multiple Primary Rules."

#### Surgery, Radiation, Chemotherapy, and Hormonal Treatment for Years 1996-97, 1998-2002, and 2003 Forward (items 1200-1296, 1310-1460, 1510, 1540-1590, 1640-1645, 3200-3280) and Corresponding Fields for Hospital-Specific Treatment and Subsequent Treatment

#### 1996-1997

For the diagnosis years 1996 and 1997, the COC *ROADS*, in preparation for the major revision of the coding of treatment implemented in 1998, separated the concept of non-cancer-directed surgery and reconstructive surgery from the field for cancer-directed surgery, while keeping the same basic codes. Additionally, the data item RX Summ--Reconstruct 1st [1330] was redefined to include reconstruction at any time in the patient's course rather than just in the first course of therapy. Three new fields (Reason for No Radiation [1430], Reason for No Chemo [1440], and Reason for No Hormone [1450]) were added, and codes 7 and 8 (Patient or Patient's Guardian Refused Chemotherapy; and Chemotherapy Recommended, Unknown if Administered) were removed from the corresponding code list. These new fields, codes, and related dates were required of COC-approved programs beginning with 1996 cases. NAACCR added all necessary fields to the Data Exchange Record Layout for 1996.

SEER continued to collect codes 7 and 8 (Patient or Patient's Guardian Refused [treatment modality] and Recommended Unknown if Given, respectively within the specific fields for radiation, chemotherapy, and hormonal therapy [1360, 1390, 1400]), instead of adding separate fields for "Reason for No [treatment modality]." Thus, there were major differences in the coding of treatment among standard-setting organizations for 1996 and 1997 cases. NAACCR revised the meaning of some codes and added a new code to RX Coding System--Current [1460] that indicates how treatment is coded in the record.

#### \* 1998-2002

Effective with cases diagnosed between January 1, 1998, and December 31, 2002, the completed treatment code revisions were implemented by COC, and the NAACCR layout was modified as needed. New fields were added: RX Summ--Scope Reg LN Sur [1292], RX Summ--Surg Oth Reg/Dis [1294], and RX Summ--Reg LN Examined [1296]. Three data items were renamed: RX Summ--CA Dir Surg [item 1290] became RX Summ--Surg Prim Site; Residual Primary Tumor [1320] became RX Summ--Surgical Margins; and Reconstructive Surgery [1330] became RX Summ--Reconstruct 1st. Another data item, RX Summ--Surgical Approch [1310], was redefined. Analogous changes were made to the corresponding fields of RX Hosp and Subsq RX. COC-approved cancer programs were required to implement all of these changes effective with 1998 cases.

SEER adopted some, but not all, of these fields effective with cancers diagnosed January 1, 1998 through December 31, 2002. SEER implemented the new codes for RX Summ--Surg Prim Site [1290]. They added the new items RX Summ--Scope Reg LN Sur [1292], RX Summ--Surg Oth Reg/Dis [1294], and RX Summ--Reg LN Examined [1296], to their required data set. They elected, on a trial basis for 1998, to collect RX Summ--Reconstruct 1st [1330] for breast cancers only. SEER continued to collect codes 7 and 8 (Patient or Patient's Guardian Refused [treatment modality] and Recommended Unknown if Given, respectively) within the specific fields for radiation, chemotherapy, and hormonal therapy [1360, 1390, 1400], instead of adding separate fields for "Reason for No [treatment modality]." COC had dropped codes 7 and 8 from each modality for which they had added a "Reason No…" field (see above). UDSC

has allowed users to either assign codes 7 and 8 or to use the fields "Reason No..." for radiation, chemotherapy, and hormonal therapy.

Historically, NPCR has required the collection of "date and type of first course of definitive treatment when available in the medical record."<sup>30</sup> For the 1996-1997 diagnosis years, NPCR-funded registries were required to collect and process available treatment information using either the (1995 or 1996) SEER Program treatment data set or the (1995 or 1996) COC treatment data set.

For 1998-2000, NPCR had a similar recommendation. NPCR-funded registries were required to adopt either the SEER 1998 or the full COC 1998 treatment data set. The NAACCR data item "RX Coding System--Current" [1460] also was encouraged to indicate how treatment was coded for a specific record.

#### Forward

Beginning in 2003, the COC *FORDS*<sup>2</sup> has redefined some treatment fields and added several more. The new and redefined data fields along with dates of treatment are required. New treatment fields include: Rad--Boost RX Modality [3200]; Rad--Boost Dose cGy [3210]; RX Hosp--Palliative Proc [3280]; RX Hosp--Scope Reg 98-02 [747]; RX Hosp--Surg Oth 98-02 [748]; RX Hosp--Surg Site 98-02 [746]; RX Summ--Palliative Proc [3270]; RX Summ--Scope Reg 98-02 [1647]; RX Summ--Surg Oth 98-02 [1648]; RX Summ--Surg Site 98-02 [1646]; and RX Summ--Transplnt/Endocr [3250]. The following fields were revised for 2003 data collection: Rad--Regional RX Modality [1570]; Rad--Treatment Volume [1540]; Reason for No Radiation [1430]; Reason for No Surgery [1340]; RX Hosp--BRM [720]; RX Summ--BRM [1410]; RX Hosp--Chemo [700]; RX Summ--Chemo [1390]; RX Hosp--DX/Stg Proc [740]; RX Summ--DX/Stg Proc [1350]; RX Hosp--Hormone [710]; RX Summ--Hormone [1400]; RX Hosp--Other [730]; RX Summ--Other [1420]; RX Hosp--Scope Reg LN Sur [672]; RX Summ--Scope Reg LN Sur [1292]; RX Hosp--Surg Oth Reg/Dis [674]; RX Summ--Surg Oth Reg/Dis [1294]; RX Hosp--Surg Prim Site [670]; RX Summ--Surg Prim Site [1290]; and RX Summ--Surgical Margins [1320].

In 2003, field width was expanded to 2 characters and codes 82, 85, 86, 87, and 88 were added to the code list for RX Hosp--Chemo [700], RX Summ--Chemo [1390], RX Hosp--BRM [720], RX Summ--BRM [1410], RX Hosp--Hormone [710]; RX Summ--Hormone [1400]; RX Hosp--Other [730]; and RX Summ--Other [1420] to record the reason if the respective treatment was not provided. The last two codes correspond to the codes 7 and 8 in the former "Reason No…" items for those treatments. Also in 2003, Reason for No Chemo [440] and Reason for No Hormone [1450] fields were discontinued.

SEER will use the same codes as the COC *FORDS* but may not collect all of the fields. For example, SEER areas will not collect Rad--Treatment Volume. See the list of data items (Chapter IX) that SEER areas collect and that SEER requires the SEER registries to transmit to NCI. SEER areas will use the field Rad--Regional RX Modality [1570] from COC hospitals to complete RX Summ--Radiation [1360].

COC Rules for conversion between the various available treatment coding schemes have been developed. It should be emphasized, however, that treatment data collected using pre-1998 treatment coding cannot be completely converted to the 1998 codes without review.

#### Time Period for First Course of Treatment (items 1260, 1270, 1500)

The NAACCR record layout provides two data items that indicate the date of the start of the first course of treatment: Date of 1st CRS RX--COC [1270] as defined by COC, and Date of Initial RX--SEER [1260] as defined by SEER. The primary difference between these two definitions is that COC defines the date the physician decides not to treat the patient as the date of initial treatment, while SEER considers such a decision

to be no treatment and the date is recorded as zeros. The NAACCR record layout contains a data item, First Course Calc Method [1500], to record which definition was used.

The SEER and COC definitions of treatment to be included as "first course" have become increasingly congruent, differing now primarily in their "fall-back" recommendations that apply when no treatment plan is recorded, no standard facility practice applies, no protocol applies, no physician is able to provide assistance, and no record of treatment failure or recurrence of disease is available. In that extreme instance, COC recommends a 4-month cutoff for the beginning of first-course treatment, and SEER applies a 1-year cutoff. Users of historical treatment data should be aware that the definitions of "first course" have changed over time and have been disjointed in the past. The applicable coding manuals and standard-setting organizations should be consulted for specifics.

Users of treatment data also should be aware that registries differ in the amount of treatment data collected in terms of the types of treatment included, non-hospital treatment locations surveyed, items covered (see the previous section), and the use of all codes provided for each item. Thus, treatment data are likely to be inconsistent among registries and to have varying levels of completeness, especially for treatment given in physicians' offices or other non-hospital settings.

#### Tumor Size Rules (item 780)

Both SEER and COC measure the size of the primary tumor (and, for malignant melanomas, the depth of invasion) in millimeters, but SEER defines variations that are not defined by COC:

- COC sets Tumor Size for all Kaposi's sarcoma, Hodgkin's lymphoma and non-Hodgkin's lymphoma cases to unknown (999); SEER uses the field for these cases to indicate HIV/AIDS status.
- SEER defines the code 001 for solid tumors as "microscopic focus or foci only," and 002 as "< 2 mm." COC applies the code 001 for "microscopic focus," but also uses the code to indicate 1 mm.</li>
- *Note:* Through 2001, COC used the same scale of measurement for the depth of invasion of malignant melanomas (whole millimeters) as it did for other tumors; SEER has always used a measurement scale 100 times finer, allowing measurements to the tenth and hundredth of a millimeter. Beginning with cases diagnosed January 1, 2002, COC uses the same measurement scale as SEER.

#### Type of Reporting Source (item 500)

This item is used to identify the source documents used to abstract a cancer case. The existing codes do not distinguish between inpatient and outpatient or clinic records. Many central registries want to keep track in more detail of the types of facilities submitting cases to the registry, especially to monitor shifts in the types of facilities delivering cancer care. UDSC has reviewed suggested enhancements to this item that would provide greater coding detail (e.g., identifying freestanding clinics).

Some central registries have adapted this item to meet changing needs. The California Cancer Registry uses the additional data item Source of Case finding to indicate the type of service or facility where a case was first identified. The NAACCR UDSC may recommend additional data items or codes in the future.

#### Vital Status (item 1760)

Both SEER and COC use code 1 in this 1-digit field to indicate that the patient is alive. However, these programs use codes 4 and 0, respectively, to indicate that the patient is dead. Both programs have long-standing historical reasons to retain their coding. No agreement has been reached on this data item.

#### **Canadian Data**

The NAACCR data standards adopted thus far do not adequately deal with data from places outside the United States. Changes have been made to accommodate postal codes, standard abbreviations for provinces/territories, and other fields in Canadian data. Future versions of this document will review and increasingly incorporate standards established for Canadian cancer registries.

## CHAPTER VI

## PATHOLOGY LABORATORY ELECTRONIC REPORTING RECOMMENDATIONS

Chapter VI, Pathology Laboratory Electronic Reporting Recommendations, is included in NAACCR Standards Volume II, Version 10 (Seventh Edition). This chapter recommended standards and implementation guidelines for electronic transmission of reports from pathology laboratories to central cancer registries. The Pathology Laboratory Electronic Reporting Recommendations currently are being revised and will be moved to the NAACCR Standards for Cancer Registries Volume I: Data Exchange Standards and Record Description, scheduled for distribution in September 2004.

# CHAPTER VII

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## **CHAPTER VIII**

## **RECORD LAYOUT TABLE (COLUMN # ORDER)**

The following table presents Version 10.2 of the NAACCR record layout. The table has column number, length, item number, item name, section, and note fields. Differences from Version 10.1 are marked "Revised" or "New" in the "Note" column of the table. Changes also are summarized in Appendix F. Please note that "Retired" items are not reflected in this table.

Column #	Length	Item #	Item Name	Section	Note
1-1	1	10	Record Type	Record ID	
2-9	8	20	Patient ID Number	Record ID	
10-10	1	30	Registry Type	Record ID	
11-11	1	35	FIN Coding System	Record ID	
12-18	7	37	Reserved 00	Record ID	
19-19	1	50	NAACCR Record Version	Record ID	
20-29	10	40	Registry ID	Record ID	
30-31	2	60	Tumor Record Number	Record ID	
32-51	20	370	Reserved 01	Record ID	
52-71	20	70	Addr at DXCity	Demographic	
72-73	2	80	Addr at DXState	Demographic	
74-82	9	100	Addr at DXPostal Code	Demographic	
83-85	3	90	County at DX	Demographic	
86-91	6	110	Census Tract 1970/80/90	Demographic	
92-92	1	120	Census Cod Sys 1970/80/90	Demographic	
93-98	6	130	Census Tract 2000	Demographic	
99-99	1	362	Census Tract Block Group	Demographic	
100-100	1	364	Census Tr Cert 1970/80/90	Demographic	
101-101	1	365	Census Tr Certainty 2000	Demographic	
102-102	1	150	Marital Status at DX	Demographic	
103-104	2	160	Race 1	Demographic	
105-106	2	161	Race 2	Demographic	
107-108	2	162	Race 3	Demographic	
109-110	2	163	Race 4	Demographic	
111-112	2	164	Race 5	Demographic	
113-113	1	170	Race Coding SysCurrent	Demographic	
114-114	1	180	Race Coding SysOriginal	Demographic	
115-115	1	190	Spanish/Hispanic Origin	Demographic	
116-116	1	200	Computed Ethnicity	Demographic	
117-117	1	210	Computed Ethnicity Source	Demographic	
118-118	1	220	Sex	Demographic	
119-121	3	230	Age at Diagnosis	Demographic	
122-129	8	240	Birth Date	Demographic	
130-132	3	250	Birthplace	Demographic	
133-134	2	260	Religion	Demographic	
135-137	3	270	Occupation CodeCensus	Demographic	
138-140	3	280	Industry CodeCensus	Demographic	
141-141	1	290	Occupation Source	Demographic	
142-142	1	300	Industry Source	Demographic	
143-182	40	310	TextUsual Occupation	Demographic	
183-222	40	320	TextUsual Industry	Demographic	
223-223	1	330	Occup/Ind Coding System	Demographic	

Column #	Length	Item #	Item Name	Section	Note
224-224	1	340	Tobacco History	Demographic	
225-225	1	350	Alcohol History	Demographic	
226-226	1	360	Family History of Cancer	Demographic	
227-228	2	3300	RuralUrban Continuum 1993	Demographic	
229-230	2	3310	RuralUrban Continuum 2000	Demographic	
231-231	1	191	NHIA Derived Hisp Origin	Demographic	New
232-280	50	530	Reserved 02	Demographic	
281-282	2	380	Sequence NumberCentral	Cancer Identification	
283-290	8	390	Date of Diagnosis	Cancer Identification	
291-294	4	400	Primary Site	Cancer Identification	
295-295	1	410	Laterality	Cancer Identification	
296-299	4	420	Histology (92-00) ICD-O-2	Cancer Identification	Subfield
296-300	5	419	MorphType&Behav ICD-O-2	Cancer Identification	Group
300-300	1	430	Behavior (92-00) ICD-O-2	Cancer Identification	Subfield
301-304	4	522	Histologic Type ICD-O-3	Cancer Identification	Subfield
301-305	5	521	MorphType&Behav ICD-O-3	Cancer Identification	Group
305-305	1	523	Behavior Code ICD-O-3	Cancer Identification	Subfield
306-306	1	440	Grade	Cancer Identification	
307-307	1	450	Site Coding SysCurrent	Cancer Identification	
308-308	1	460	Site Coding SysOriginal	Cancer Identification	
309-309	1	470	Morph Coding SysCurrent	Cancer Identification	
310-310	1	480	Morph Coding SysOriginl	Cancer Identification	
311-311	1	490	Diagnostic Confirmation	Cancer Identification	
312-312	1	500	Type of Reporting Source	Cancer Identification	
313-320	8	510	Screening Date	Cancer Identification	
321-321	1	520	Screening Result	Cancer Identification	
322-371	50	680	Reserved 03	Cancer Identification	
372-381	10	538	Reporting Hospital FAN	Hospital-Specific	
382-391	10	540	Reporting Hospital	Hospital-Specific	
392-401	10	3100	Archive FIN	Hospital-Specific	
402-410	9	550	Accession NumberHosp	Hospital-Specific	
411-412	2	560	Sequence NumberHospital	Hospital-Specific	
413-415	3	570	Abstracted By	Hospital-Specific	
416-423	8	580	Date of 1st Contact	Hospital-Specific	
424-431	8	590	Date of Inpatient Adm	Hospital-Specific	
432-439	8	600	Date of Inpatient Disch	Hospital-Specific	
440-440	1	610	Class of Case	Hospital-Specific	
441-444	4	620	Year First Seen This CA	Hospital-Specific	
445-446	2	630	Primary Payer at DX	Hospital-Specific	
447-447	1	640	Inpatient/Outpt Status	Hospital-Specific	
448-448	1	650	Presentation at CA Conf	Hospital-Specific	
449-456	8	660	Date of CA Conference	Hospital-Specific	

Column #	Length	Item #	Item Name	Section	Note
457-458	2	670	RX HospSurg Prim Site	Hospital-Specific	
459-459	1	672	RX HospScope Reg LN Sur	Hospital-Specific	
460-460	1	674	RX HospSurg Oth Reg/Dis	Hospital-Specific	
461-462	2	676	RX HospReg LN Removed	Hospital-Specific	
463-463	1	690	RX HospRadiation	Hospital-Specific	
464-465	2	700	RX HospChemo	Hospital-Specific	
466-467	2	710	RX HospHormone	Hospital-Specific	
468-469	2	720	RX HospBRM	Hospital-Specific	
470-470	1	730	RX HospOther	Hospital-Specific	
471-472	2	740	RX HospDX/Stg Proc	Hospital-Specific	
473-473	1	3280	RX HospPalliative Proc	Hospital-Specific	
474-474	1	742	RX HospScreen/BX Proc1	Hospital-Specific	
475-475	1	743	RX HospScreen/BX Proc2	Hospital-Specific	
476-476	1	744	RX HospScreen/BX Proc3	Hospital-Specific	
477-477	1	745	RX HospScreen/BX Proc4	Hospital-Specific	
478-479	2	746	RX HospSurg Site 98-02	Treatment-1st Course	
480-480	1	747	RX HospScope Reg 98-02	Treatment-1st Course	
481-481	1	748	RX HospSurg Oth 98-02	Treatment-1st Course	
482-527	46	750	Reserved 04	Hospital-Specific	
528-528	1	759	SEER Summary Stage 2000	Stage/Prognostic Factors	
529-529	1	760	SEER Summary Stage 1977	Stage/Prognostic Factors	
530-530	1	770	Loc/Reg/Distant Stage	Stage/Prognostic Factors	
531-533	3	780	EODTumor Size	Stage/Prognostic Factors	Subfield
531-542	12	779	Extent of Disease 10-Dig	Stage/Prognostic Factors	Group
534-535	2	790	EODExtension	Stage/Prognostic Factors	Subfield
536-537	2	800	EODExtension Prost Path	Stage/Prognostic Factors	Subfield
538-538	1	810	EODLymph Node Involv	Stage/Prognostic Factors	Subfield
539-540	2	820	Regional Nodes Positive	Stage/Prognostic Factors	Subfield
541-542	2	830	Regional Nodes Examined	Stage/Prognostic Factors	Subfield
543-555	13	840	EODOld 13 Digit	Stage/Prognostic Factors	
556-557	2	850	EODOld 2 Digit	Stage/Prognostic Factors	
558-561	4	860	EODOld 4 Digit	Stage/Prognostic Factors	
562-562	1	870	Coding System for EOD	Stage/Prognostic Factors	
563-564	2	880	TNM Path T	Stage/Prognostic Factors	
565-566	2	890	TNM Path N	Stage/Prognostic Factors	
567-568	2	900	TNM Path M	Stage/Prognostic Factors	
569-570	2	910	TNM Path Stage Group	Stage/Prognostic Factors	
571-571	1	920	TNM Path Descriptor	Stage/Prognostic Factors	
572-572	1	930	TNM Path Staged By	Stage/Prognostic Factors	
573-574	2	940	TNM Clin T	Stage/Prognostic Factors	
575-576	2	950	TNM Clin N	Stage/Prognostic Factors	
577-578	2	960	TNM Clin M	Stage/Prognostic Factors	

Column #	Length	Item #	Item Name	Section	Note
579-580	2	970	TNM Clin Stage Group	Stage/Prognostic Factors	
581-581	1	980	TNM Clin Descriptor	Stage/Prognostic Factors	
582-582	1	990	TNM Clin Staged By	Stage/Prognostic Factors	
583-584	2	1000	TNM Other T	Stage/Prognostic Factors	
585-586	2	1010	TNM Other N	Stage/Prognostic Factors	
587-588	2	1020	TNM Other M	Stage/Prognostic Factors	
589-590	2	1030	TNM Other Stage Group	Stage/Prognostic Factors	
591-591	1	1040	TNM Other Staged By	Stage/Prognostic Factors	
592-592	1	1050	TNM Other Descriptor	Stage/Prognostic Factors	
593-594	2	1060	TNM Edition Number	Stage/Prognostic Factors	
595-609	15	1070	Other Staging System	Stage/Prognostic Factors	
610-617	8	1080	Date of 1st Positive BX	Stage/Prognostic Factors	
618-618	1	1090	Site of Distant Met 1	Stage/Prognostic Factors	
619-619	1	1100	Site of Distant Met 2	Stage/Prognostic Factors	
620-620	1	1110	Site of Distant Met 3	Stage/Prognostic Factors	
621-622	2	1120	Pediatric Stage	Stage/Prognostic Factors	
623-624	2	1130	Pediatric Staging System	Stage/Prognostic Factors	
625-625	1	1140	Pediatric Staged By	Stage/Prognostic Factors	
626-626	1	1150	Tumor Marker 1	Stage/Prognostic Factors	
627-627	1	1160	Tumor Marker 2	Stage/Prognostic Factors	
628-628	1	1170	Tumor Marker 3	Stage/Prognostic Factors	
629-631	3	2800	CS Tumor Size	Stage/Prognostic Factors	
632-633	2	2810	CS Extension	Stage/Prognostic Factors	
634-634	1	2820	CS Tumor Size/Ext Eval	Stage/Prognostic Factors	
635-636	2	2830	CS Lymph Nodes	Stage/Prognostic Factors	
637-637	1	2840	CS Reg Node Eval	Stage/Prognostic Factors	
638-639	2	2850	CS Mets at DX	Stage/Prognostic Factors	
640-640	1	2860	CS Mets Eval	Stage/Prognostic Factors	
641-643	3	2880	CS Site-Specific Factor 1	Stage/Prognostic Factors	
644-646	3	2890	CS Site-Specific Factor 2	Stage/Prognostic Factors	
647-649	3	2900	CS Site-Specific Factor 3	Stage/Prognostic Factors	
650-652	3	2910	CS Site-Specific Factor 4	Stage/Prognostic Factors	
653-655	3	2920	CS Site-Specific Factor 5	Stage/Prognostic Factors	
656-658	3	2930	CS Site-Specific Factor 6	Stage/Prognostic Factors	
659-660	2	2940	Derived AJCC T	Stage/Prognostic Factors	
661-661	1	2950	Derived AJCC T Descriptor	Stage/Prognostic Factors	
662-663	2	2960	Derived AJCC N	Stage/Prognostic Factors	
664-664	1	2970	Derived AJCC N Descriptor	Stage/Prognostic Factors	
665-666	2	2980	Derived AJCC M	Stage/Prognostic Factors	
667-667	1	2990	Derived AJCC M Descriptor	Stage/Prognostic Factors	
668-669	2	3000	Derived AJCC Stage Group	Stage/Prognostic Factors	
670-670	1	3010	Derived SS1977	Stage/Prognostic Factors	

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671-671	1	3020	Derived SS2000	Stage/Prognostic Factors	
672-672	1	3030	Derived AJCCFlag	Stage/Prognostic Factors	
673-673	1	3040	Derived SS1977Flag	Stage/Prognostic Factors	
674-674	1	3050	Derived SS2000Flag	Stage/Prognostic Factors	
675-679	5	3110	Comorbid/Complication 1	Stage/Prognostic Factors	
680-684	5	3120	Comorbid/Complication 2	Stage/Prognostic Factors	
685-689	5	3130	Comorbid/Complication 3	Stage/Prognostic Factors	
690-694	5	3140	Comorbid/Complication 4	Stage/Prognostic Factors	
695-699	5	3150	Comorbid/Complication 5	Stage/Prognostic Factors	
700-704	5	3160	Comorbid/Complication 6	Stage/Prognostic Factors	
705-710	6	2935	CS Version 1st	Stage/Prognostic Factors	
711-716	6	2936	CS Version Latest	Stage/Prognostic Factors	
717-754	38	1180	Reserved 05	Stage/Prognostic Factors	
755-762	8	1200	RX DateSurgery	Treatment-1st Course	
763-770	8	3170	RX DateMost Defin Surg	Treatment-1st Course	
771-778	8	3180	RX DateSurgical Disch	Treatment-1st Course	
779-786	8	1210	RX DateRadiation	Treatment-1st Course	
787-794	8	3220	RX DateRadiation Ended	Treatment-1st Course	
795-802	8	3230	RX DateSystemic	Treatment-1st Course	
803-810	8	1220	RX DateChemo	Treatment-1st Course	
811-818	8	1230	RX DateHormone	Treatment-1st Course	
819-826	8	1240	RX DateBRM	Treatment-1st Course	
827-834	8	1250	RX DateOther	Treatment-1st Course	
835-842	8	1260	Date of Initial RXSEER	Treatment-1st Course	
843-850	8	1270	Date of 1st Crs RXCOC	Treatment-1st Course	
851-858	8	1280	RX DateDX/Stg Proc	Treatment-1st Course	
859-860	2	1290	RX SummSurg Prim Site	Treatment-1st Course	
861-861	1	1292	RX SummScope Reg LN Sur	Treatment-1st Course	
862-862	1	1294	RX SummSurg Oth Reg/Dis	Treatment-1st Course	
863-864	2	1296	RX SummReg LN Examined	Treatment-1st Course	
865-865	1	1310	RX SummSurgical Approch	Treatment-1st Course	
866-866	1	1320	RX SummSurgical Margins	Treatment-1st Course	
867-867	1	1330	RX SummReconstruct 1st	Treatment-1st Course	
868-868	1	1340	Reason for No Surgery	Treatment-1st Course	
869-870	2	1350	RX SummDX/Stg Proc	Treatment-1st Course	
871-871	1	3270	RX SummPalliative Proc	Treatment-1st Course	
872-872	1	1355	Reserved 22	Treatment-1st Course	
873-873	1	1360	RX SummRadiation	Treatment-1st Course	
874-874	1	1370	RX SummRad to CNS	Treatment-1st Course	
875-875	1	1380	RX SummSurg/Rad Seq	Treatment-1st Course	
876-877	2	3250	RX SummTransplnt/Endocr	Treatment-1st Course	
878-879	2	1390	RX SummChemo	Treatment-1st Course	

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880-881	2	1400	RX SummHormone	Treatment-1st Course	
882-883	2	1410	RX SummBRM	Treatment-1st Course	
884-884	1	1420	RX SummOther	Treatment-1st Course	
885-885	1	1430	Reason for No Radiation	Treatment-1st Course	
886-886	1	1440	Reason for No Chemo	Treatment-1st Course	
887-887	1	1450	Reason for No Hormone	Treatment-1st Course	
888-889	2	1460	RX Coding SystemCurrent	Treatment-1st Course	
890-890	1	1470	Protocol Eligibility Stat	Treatment-1st Course	
891-892	2	1480	Protocol Participation	Treatment-1st Course	
893-893	1	1490	Referral to Support Serv	Treatment-1st Course	
894-894	1	1500	First Course Calc Method	Treatment-1st Course	
895-899	5	1510	RadRegional Dose: CGY	Treatment-1st Course	
900-901	2	1520	RadNo of Treatment Vol	Treatment-1st Course	
902-904	3	1530	RadElapsed RX Days	Treatment-1st Course	
905-906	2	1540	RadTreatment Volume	Treatment-1st Course	
907-907	1	1550	RadLocation of RX	Treatment-1st Course	
908-908	1	1560	RadIntent of Treatment	Treatment-1st Course	
909-910	2	1570	RadRegional RX Modality	Treatment-1st Course	
911-912	2	3200	RadBoost RX Modality	Treatment-1st Course	
913-917	5	3210	RadBoost Dose cGy	Treatment-1st Course	
918-918	1	1580	RadRX Completion Status	Treatment-1st Course	
919-919	1	1590	RadLocal Control Status	Treatment-1st Course	
920-931	12	1635	Reserved 23	Treatment-1st Course	
932-933	2	1640	RX SummSurgery Type	Treatment-1st Course	
934-934	1	1642	RX SummScreen/BX Proc1	Treatment-1st Course	
935-935	1	1643	RX SummScreen/BX Proc2	Treatment-1st Course	
936-936	1	1644	RX SummScreen/BX Proc3	Treatment-1st Course	
937-937	1	1645	RX SummScreen/BX Proc4	Treatment-1st Course	
938-938	1	3190	Readm Same Hosp 30 Days	Treatment-1st Course	
939-940	2	1646	RX SummSurg Site 98-02	Treatment-1st Course	
941-941	1	1647	RX SummScope Reg 98-02	Treatment-1st Course	
942-942	1	1648	RX SummSurg Oth 98-02	Treatment-1st Course	
943-987	45	1190	Reserved 06	Treatment-1st Course	
988-995	8	1660	Subsq RX 2nd Course Date	Treatment-Subsequent & Other	
996-1002	7	1670	Subsq RX 2nd Course Codes	Treatment-Subsequent & Other	Group
996-997	2	1671	Subsq RX 2nd Course Surg	Treatment-Subsequent & Other	Subfield
998-998	1	1672	Subsq RX 2nd Course Rad	Treatment-Subsequent & Other	Subfield
999-999	1	1673	Subsq RX 2nd Course Chemo	Treatment-Subsequent & Other	Subfield
1000-1000	1	1674	Subsq RX 2nd Course Horm	Treatment-Subsequent & Other	Subfield
1001-1001	1	1675	Subsq RX 2nd Course BRM	Treatment-Subsequent & Other	Subfield
1001-1001	1	1676	Subsq RX 2nd Course DRM Subsq RX 2nd Course Oth	Treatment-Subsequent & Other	Subfield
1002-1002	8	1680	Subsq RX 2rd Course Oth Subsq RX 3rd Course Date	Treatment-Subsequent & Other	Subliciu

Column #	Length	Item #	Item Name	Section	Note
1011-1012	2	1691	Subsq RX 3rd Course Surg	Treatment-Subsequent & Other	Subfield
1011-1017	7	1690	Subsq RX 3rd Course Codes	Treatment-Subsequent & Other	Group
1013-1013	1	1692	Subsq RX 3rd Course Rad	Treatment-Subsequent & Other	Subfield
1014-1014	1	1693	Subsq RX 3rd Course Chemo	Treatment-Subsequent & Other	Subfield
1015-1015	1	1694	Subsq RX 3rd Course Horm	Treatment-Subsequent & Other	Subfield
1016-1016	1	1695	Subsq RX 3rd Course BRM	Treatment-Subsequent & Other	Subfield
1017-1017	1	1696	Subsq RX 3rd Course Oth	Treatment-Subsequent & Other	Subfield
1018-1025	8	1700	Subsq RX 4th Course Date	Treatment-Subsequent & Other	
1026-1027	2	1711	Subsq RX 4th Course Surg	Treatment-Subsequent & Other	Subfield
1026-1032	7	1710	Subsq RX 4th Course Codes	Treatment-Subsequent & Other	Group
1028-1028	1	1712	Subsq RX 4th Course Rad	Treatment-Subsequent & Other	Subfield
1029-1029	1	1713	Subsq RX 4th Course Chemo	Treatment-Subsequent & Other	Subfield
1030-1030	1	1714	Subsq RX 4th Course Horm	Treatment-Subsequent & Other	Subfield
1031-1031	1	1715	Subsq RX 4th Course BRM	Treatment-Subsequent & Other	Subfield
1032-1032	1	1716	Subsq RX 4th Course Oth	Treatment-Subsequent & Other	Subfield
1033-1040	8	1720	Subsq RX 5th Course Date	Treatment-Subsequent & Other	
1041-1042	2	1731	Subsq RX 5th Course Surg	Treatment-Subsequent & Other	Subfield
1041-1047	7	1730	Subsq RX 5th Course Codes	Treatment-Subsequent & Other	Group
1043-1043	1	1732	Subsq RX 5th Course Rad	Treatment-Subsequent & Other	Subfield
1044-1044	1	1733	Subsq RX 5th Course Chemo	Treatment-Subsequent & Other	Subfield
1045-1045	1	1734	Subsq RX 5th Course Horm	Treatment-Subsequent & Other	Subfield
1046-1046	1	1735	Subsq RX 5th Course BRM	Treatment-Subsequent & Other	Subfield
1047-1047	1	1736	Subsq RX 5th Course Oth	Treatment-Subsequent & Other	Subfield
1048-1048	1	1677	Subsq RX 2ndScope LN SU	Treatment-Subsequent & Other	
1049-1049	1	1678	Subsq RX 2ndSurg Oth	Treatment-Subsequent & Other	
1050-1051	2	1679	Subsq RX 2ndReg LN Rem	Treatment-Subsequent & Other	
1052-1052	1	1697	Subsq RX 3rdScope LN SU	Treatment-Subsequent & Other	
1053-1053	1	1698	Subsq RX 3rdSurg Oth	Treatment-Subsequent & Other	
1054-1055	2	1699	Subsq RX 3rdReg LN Rem	Treatment-Subsequent & Other	
1056-1056	1	1717	Subsq RX 4thScope LN SU	Treatment-Subsequent & Other	
1057-1057	1	1718	Subsq RX 4thSurg Oth	Treatment-Subsequent & Other	
1058-1059	2	1719	Subsq RX 4thReg LN Rem	Treatment-Subsequent & Other	
1060-1060	1	1737	Subsq RX 5thScope LN SU	Treatment-Subsequent & Other	
1061-1061	1	1738	Subsq RX 5thSurg Oth	Treatment-Subsequent & Other	
1062-1063	2	1739	Subsq RX 5thReg LN Rem	Treatment-Subsequent & Other	
1064-1064	1	1741	Subsq RXReconstruct Del	Treatment-Subsequent & Other	
1065-1114	50	1300	Reserved 07	Treatment-Subsequent & Other	
1115-1115	1	1981	Over-ride SS/NodesPos	Edit Overrides/Conversion History/System Admin	
1116-1116	1	1982	Over-ride SS/TNM-N	Edit Overrides/Conversion History/System Admin	
1117-1117	1	1983	Over-ride SS/TNM-M	Edit Overrides/Conversion History/System Admin	

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1118-1118	1	1984	Over-ride SS/DisMet1	Edit Overrides/Conversion History/System Admin	
1119-1119	1	1985	Over-ride Acsn/Class/Seq	Edit Overrides/Conversion History/System Admin	
1120-1120	1	1986	Over-ride HospSeq/DxConf	Edit Overrides/Conversion History/System Admin	
1121-1121	1	1987	Over-ride COC-Site/Type	Edit Overrides/Conversion History/System Admin	
1122-1122	1	1988	Over-ride HospSeq/Site	Edit Overrides/Conversion History/System Admin	
1123-1123	1	1989	Over-ride Site/TNM-StgGrp	Edit Overrides/Conversion History/System Admin	
1124-1124	1	1990	Over-ride Age/Site/Morph	Edit Overrides/Conversion History/System Admin	
1125-1125	1	2000	Over-ride SeqNo/DxConf	Edit Overrides/Conversion History/System Admin	
1126-1126	1	2010	Over-ride Site/Lat/SeqNo	Edit Overrides/Conversion History/System Admin	
1127-1127	1	2020	Over-ride Surg/DxConf	Edit Overrides/Conversion History/System Admin	
1128-1128	1	2030	Over-ride Site/Type	Edit Overrides/Conversion History/System Admin	
1129-1129	1	2040	Over-ride Histology	Edit Overrides/Conversion History/System Admin	
1130-1130	1	2050	Over-ride Report Source	Edit Overrides/Conversion History/System Admin	
1131-1131	1	2060	Over-ride Ill-define Site	Edit Overrides/Conversion History/System Admin	
1132-1132	1	2070	Over-ride Leuk, Lymphoma	Edit Overrides/Conversion History/System Admin	
1133-1133	1	2071	Over-ride Site/Behavior	Edit Overrides/Conversion History/System Admin	
1134-1134	1	2072	Over-ride Site/EOD/DX Dt	Edit Overrides/Conversion History/System Admin	
1135-1135	1	2073	Over-ride Site/Lat/EOD	Edit Overrides/Conversion History/System Admin	
1136-1136	1	2074	Over-ride Site/Lat/Morph	Edit Overrides/Conversion History/System Admin	
1137-1140	4	1960	Site (73-91) ICD-O-1	Edit Overrides/Conversion History/System Admin	
1141-1144	4	1971	Histology (73-91) ICD-O-1	Edit Overrides/Conversion History/System Admin	Subfield
1141-1146	6	1970	Morph (73-91) ICD-O-1	Edit Overrides/Conversion History/System Admin	Group
1145-1145	1	1972	Behavior (73-91) ICD-O-1	Edit Overrides/Conversion History/System Admin	Subfield
1146-1146	1	1973	Grade (73-91) ICD-O-1	Edit Overrides/Conversion History/System Admin	Subfield
1147-1147	1	1980	ICD-O-2 Conversion Flag	Edit Overrides/Conversion History/System Admin	
1148-1163	16	2082	Reserved 24	Edit Overrides/Conversion History/System Admin	

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1164-1173	10	2081	CRC CHECKSUM	Edit Overrides/Conversion History/System Admin	
1174-1181	8	2090	Date Case Completed	Edit Overrides/Conversion History/System Admin	
1182-1189	8	2100	Date Case Last Changed	Edit Overrides/Conversion History/System Admin	
1190-1197	8	2110	Date Case Report Exported	Edit Overrides/Conversion History/System Admin	
1198-1198	1	2120	SEER Coding SysCurrent	Edit Overrides/Conversion History/System Admin	
1199-1199	1	2130	SEER Coding SysOriginal	Edit Overrides/Conversion History/System Admin	
1200-1201	2	2140	COC Coding SysCurrent	Edit Overrides/Conversion History/System Admin	
1202-1203	2	2150	COC Coding SysOriginal	Edit Overrides/Conversion History/System Admin	
1204-1213	10	2170	Vendor Name	Edit Overrides/Conversion History/System Admin	
1214-1214	1	2180	SEER Type of Follow-Up	Edit Overrides/Conversion History/System Admin	
1215-1216	2	2190	SEER Record Number	Edit Overrides/Conversion History/System Admin	
1217-1218	2	2200	Diagnostic Proc 73-87	Edit Overrides/Conversion History/System Admin	
1219-1226	8	2111	Date Case Report Received	Edit Overrides/Conversion History/System Admin	
1227-1234	8	2112	Date Case Report Loaded	Edit Overrides/Conversion History/System Admin	
1235-1242	8	2113	Date Tumor Record Availbl	Edit Overrides/Conversion History/System Admin	
1243-1243	1	2116	ICD-O-3 Conversion Flag	Edit Overrides/Conversion History/System Admin	
1244-1293	50	1650	Reserved 08	Edit Overrides/Conversion History/System Admin	
1294-1301	8	1750	Date of Last Contact	Follow-up/Recurrence/Death	
1302-1302	1	1760	Vital Status	Follow-up/Recurrence/Death	
1303-1303	1	1770	Cancer Status	Follow-up/Recurrence/Death	
1304-1304	1	1780	Quality of Survival	Follow-up/Recurrence/Death	
1305-1305	1	1790	Follow-Up Source	Follow-up/Recurrence/Death	
1306-1306	1	1800	Next Follow-Up Source	Follow-up/Recurrence/Death	
1307-1326	20	1810	Addr CurrentCity	Follow-up/Recurrence/Death	
1327-1328	2	1820	Addr CurrentState	Follow-up/Recurrence/Death	
1329-1337	9	1830	Addr CurrentPostal Code	Follow-up/Recurrence/Death	
1338-1340	3	1840	CountyCurrent	Follow-up/Recurrence/Death	
1341-1341	1	1850	Unusual Follow-Up Method	Follow-up/Recurrence/Death	
1342-1349	8	1860	Recurrence Date1st	Follow-up/Recurrence/Death	
1350-1350	1	1871	Recurrence Distant Site 1	Follow-up/Recurrence/Death	
1351-1351	1	1872	Recurrence Distant Site 2	Follow-up/Recurrence/Death	

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1352-1352	1	1873	Recurrence Distant Site 3	Follow-up/Recurrence/Death	
1353-1354	2	1880	Recurrence Type1st	Follow-up/Recurrence/Death	
1355-1356	2	1890	Recurrence Type1stOth	Follow-up/Recurrence/Death	
1357-1376	20	1842	Follow-Up ContactCity	Follow-up/Recurrence/Death	
1377-1378	2	1844	Follow-Up ContactState	Follow-up/Recurrence/Death	
1379-1387	9	1846	Follow-Up ContactPostal	Follow-up/Recurrence/Death	
1388-1391	4	1910	Cause of Death	Follow-up/Recurrence/Death	
1392-1392	1	1920	ICD Revision Number	Follow-up/Recurrence/Death	
1393-1393	1	1930	Autopsy	Follow-up/Recurrence/Death	
1394-1396	3	1940	Place of Death	Follow-up/Recurrence/Death	
1397-1446	50	1740	Reserved 09	Follow-up/Recurrence/Death	
1447-1946	500	2220	State/Requestor Items	Special Use	
1947-1971	25	2230	NameLast	Patient-Confidential	
1972-1985	14	2240	NameFirst	Patient-Confidential	
1986-1999	14	2250	NameMiddle	Patient-Confidential	
2000-2002	3	2260	NamePrefix	Patient-Confidential	
2003-2005	3	2270	NameSuffix	Patient-Confidential	
2006-2020	15	2280	NameAlias	Patient-Confidential	
2021-2035	15	2390	NameMaiden	Patient-Confidential	
2036-2085	50	2290	NameSpouse/Parent	Patient-Confidential	
2086-2096	11	2300	Medical Record Number	Patient-Confidential	
2097-2098	2	2310	Military Record No Suffix	Patient-Confidential	
2099-2107	9	2320	Social Security Number	Patient-Confidential	
2108-2147	40	2330	Addr at DXNo & Street	Patient-Confidential	
2148-2187	40	2335	Addr at DXSupplementl	Patient-Confidential	
2188-2227	40	2350	Addr CurrentNo & Street	Patient-Confidential	
2228-2267	40	2355	Addr CurrentSupplementl	Patient-Confidential	
2268-2277	10	2360	Telephone	Patient-Confidential	
2278-2283	6	2380	DC State File Number	Patient-Confidential	
2284-2313	30	2394	Follow-Up ContactName	Patient-Confidential	
2314-2353	40	2392	Follow-Up ContactNo&St	Patient-Confidential	
2354-2393	40	2393	Follow-Up ContactSuppl	Patient-Confidential	
2394-2403	10	2352	Latitude	Patient-Confidential	
2404-2414	11	2354	Longitude	Patient-Confidential	
2415-2464	50	1835	Reserved 10	Patient-Confidential	
2465-2474	10	2430	Last Follow-Up Hospital	Hospital-Confidential	
2475-2484	10	2440	Following Registry	Hospital-Confidential	
2485-2494	10	2410	Institution Referred From	Hospital-Confidential	
2495-2504	10	2420	Institution Referred To	Hospital-Confidential	
2505-2554	50	1900	Reserved 11	Hospital-Confidential	
2555-2562	8	2460	PhysicianManaging	Other-Confidential	
2563-2570	8	2470	PhysicianFollow-Up	Other-Confidential	

Column #	Length	Item #	Item Name	Section	Note
2571-2578	8	2480	PhysicianPrimary Surg	Other-Confidential	
2579-2586	8	2490	Physician 3	Other-Confidential	
2587-2594	8	2500	Physician 4	Other-Confidential	
2595-2644	50	1950	Reserved 12	Other-Confidential	
2645-2844	200	2520	TextDX ProcPE	Text-Diagnosis	
2845-3094	250	2530	TextDX ProcX-ray/Scan	Text-Diagnosis	
3095-3344	250	2540	TextDX ProcScopes	Text-Diagnosis	
3345-3594	250	2550	TextDX ProcLab Tests	Text-Diagnosis	
3595-3844	250	2560	TextDX ProcOp	Text-Diagnosis	
3845-4094	250	2570	TextDX ProcPath	Text-Diagnosis	
4095-4134	40	2580	TextPrimary Site Title	Text-Diagnosis	
4135-4174	40	2590	TextHistology Title	Text-Diagnosis	
4175-4474	300	2600	TextStaging	Text-Diagnosis	
4475-4624	150	2610	RX TextSurgery	Text-Treatment	
4625-4774	150	2620	RX TextRadiation (Beam)	Text-Treatment	
4775-4924	150	2630	RX TextRadiation Other	Text-Treatment	
4925-5124	200	2640	RX TextChemo	Text-Treatment	
5125-5324	200	2650	RX TextHormone	Text-Treatment	
5325-5424	100	2660	RX TextBRM	Text-Treatment	
5425-5524	100	2670	RX TextOther	Text-Treatment	
5525-5874	350	2680	TextRemarks	Text-Miscellaneous	
5875-5924	50	2690	Place of Diagnosis	Text-Miscellaneous	
5925-6694	770	2700	Reserved 19	Text-Miscellaneous	

## **CHAPTER IX**

#### **REQUIRED STATUS TABLE (ITEM # ORDER)**

The following table presents Version 10.2 of the NAACCR required status summarizing the requirements and recommendations for collection of each item by standard-setting groups. Differences from Version 10.1 are marked "Revised," "New," or "Retired" in the "Note" column of the table. Changes are also summarized in Appendix F.

The following abbreviations and symbols are used in the table:

NAACCR Exc	NAACCR committees are reviewing and will make recommendations in Version 11.
NAACCR Inc	NAACCR committees are reviewing and will make recommendations in Version 11.
NAACCR Full	NAACCR committees are reviewing and will make recommendations in Version 11.
NPCR	Refers to requirements and recommendations of the NPCR regarding data items that should be collected or computed by NPCR state registries. Note: Personal identifying data items that are collected are not transmitted to CDC.
COC	Refers to requirements of COC. Facilities should refer to the COC <i>FORDS Manual</i> for further clarification of required fields.
SEER	Refers to requirements of NCI's SEER Program. Facilities and central registries should refer to the <i>SEER Program Code Manual</i> for further clarification of required fields.

**Codes for Recommendations:** R = Required. RH = Historically collected and currently transmitted. RC = Collected by SEER from COC-approved hospitals. RS = Required, site specific. S = Supplementary/recommended. D = Derived. • = Not in data set but available. \* = When available. # = Central registries may code available data using either the SEER or COC data item and associated rules.

^ = These text requirements may be met with one or several text block fields.

Item :	Item Name	<u>NPCR</u>	COC		SEER		Source of	
			Collect	Transmit	Collect	Transmit	Standard	Note
10	Record Type	•	•	R	•	R	NAACCR	
20	Patient ID Number	R	•	•	R	R	Reporting Registry	
30	Registry Type	•	•	•	•	•	NAACCR	
35	FIN Coding System	S	•	•	•	•	NAACCR	
37	Reserved 00	•	•	•	•	•		
40	Registry ID	S	•	•	R	R	NAACCR	
50	NAACCR Record Version	R	•	R	•	•	NAACCR	
60	Tumor Record Number	S	•	•	•	•	NAACCR	
70	Addr at DXCity	R	R	R	R	•	COC	
80	Addr at DXState	R	R	R	R	•	COC	
90	County at DX	R	R	R	R	R	FIPS/SEER	
100	Addr at DXPostal Code	R	R	R	R	•	COC	
110	Census Tract 1970/80/90	RH	•	•	RH	RH	SEER	
120	Census Cod Sys 1970/80/90	RH	•	•	RH	RH	SEER	
130	Census Tract 2000	R	•	•	R	R	NAACCR	
140	Census Tract Cod SysAlt	•	•	•	•	•		Retired
150	Marital Status at DX	S	•	•	R	R	SEER	
160	Race 1	R	R	R	R	R	SEER/COC	
161	Race 2	R	R	R	R	R	SEER/COC	
162	Race 3	R	R	R	R	R	SEER/COC	
163	Race 4	R	R	R	R	R	SEER/COC	
164	Race 5	R	R	R	R	R	SEER/COC	
170	Race Coding SysCurrent	•	R	R	•	•	NAACCR	
180	Race Coding SysOriginal	•	R	R	•	•	NAACCR	
190	Spanish/Hispanic Origin	R	R	R	R	R	SEER/COC	
191	NHIA Derived Hisp Origin	R	•	•	R	R	NAACCR	New
200	Computed Ethnicity	S	•	•	R	R	SEER	
210	Computed Ethnicity Source	S	•	•	R	R	SEER	
220	Sex	R	R	R	R	R	SEER/COC	
230	Age at Diagnosis	R	R	R	R	R	SEER/COC	
240	Birth Date	R	R	R	R	R	SEER/COC	
250	Birthplace	R*	R	R	R	R	SEER/COC	
260	Religion	•	•	•	•	•	Varies	

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Item :	Item Name	<u>NPCR</u>	COC		SEER		Source of	<b>.</b> .
			Collect	Transmit	Collect	Transmit	Standard	Note
270	Occupation CodeCensus	S	•	•	•	•	Census/NPCR	
280	Industry CodeCensus	S	•	•	•	•	Census/NPCR	
290	Occupation Source	S	•	•	•	•	NPCR	
300	Industry Source	S	•	•	•	•	NPCR	
310	TextUsual Occupation	R*	•	•	•	•	NPCR	
320	TextUsual Industry	R*	•	•	•	•	NPCR	
330	Occup/Ind Coding System	S	•	•	•	•	NPCR	
340	Tobacco History	•	•	•	•	•	Varies	
350	Alcohol History	•	•	•	•	•	Varies	
360	Family History of Cancer	•	•	•	•	•	Varies	
362	Census Tract Block Group	•	•	•	•	•	Census	
364	Census Tr Cert 1970/80/90	RH	•	•	RH	RH	SEER	
365	Census Tr Certainty 2000	R	•	•	R	R	NAACCR	
370	Reserved 01	•	•	•	•	•		
380	Sequence NumberCentral	R	•	•	R	R	SEER	
390	Date of Diagnosis	R	R	R	R	R	SEER/COC	
400	Primary Site	R	R	R	R	R	SEER/COC	
410	Laterality	R	R	R	R	R	SEER/COC	
419	MorphType&Behav ICD-O-2							
420	Histology (92-00) ICD-O-2	RH	•	RH	RH	RH	SEER/COC	
430	Behavior (92-00) ICD-O-2	RH	•	RH	RH	RH	SEER/COC	
440	Grade	R	R	R	R	R	SEER/COC	
450	Site Coding SysCurrent	S	R	R	•	•	NAACCR	
460	Site Coding SysOriginal	•	R	R	•	•	NAACCR	
470	Morph Coding SysCurrent	S	R	R	•	•	NAACCR	
480	Morph Coding SysOriginl	•	R	R	•	•	NAACCR	
490	Diagnostic Confirmation	R	R	R	R	R	SEER/COC	
500	Type of Reporting Source	R	•	•	R	R	SEER	
510	Screening Date	•	•	•	•	•	COC	
520	Screening Result	•	•	•	•	•	COC	
521	MorphType&Behav ICD-O-3							
522	Histologic Type ICD-O-3	R	R	R	R	R	SEER/COC	
523	Behavior Code ICD-O-3	R	R	R	R	R	SEER/COC	
530	Reserved 02	•	•	•	•	•		

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 $^{\wedge}$  = These text requirements may be met with one or several text block fields.

Item :	Item Name	NDCD	COC		SEER		Source of	
		<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
538	Reporting Hospital FAN	•	•	•	•	•	COC	
540	Reporting Hospital	S	R	R	R	•	COC	
550	Accession NumberHosp	S	R	R	R	•	COC	
560	Sequence NumberHospital	S	R	R	R	•	COC	
570	Abstracted By	•	R	R	R	•	COC	
580	Date of 1st Contact	R	R	R	•	•	COC	
590	Date of Inpatient Adm	•	٠	•	•	•	NAACCR	
600	Date of Inpatient Disch	•	•	•	•	•	NAACCR	
610	Class of Case	S	R	R	RC	•	COC	
620	Year First Seen This CA	•	٠	•	•	•	COC	
630	Primary Payer at DX	•	R	R	•	•	COC	
640	Inpatient/Outpt Status	•	•	•	•	•	COC	
650	Presentation at CA Conf	•	•	•	•	•	COC	
660	Date of CA Conference	•	•	•	•	•	COC	
670	RX HospSurg Prim Site	•	R	R	R	•	COC	
672	RX HospScope Reg LN Sur	•	R	R	R	•	COC	
674	RX HospSurg Oth Reg/Dis	•	R	R	R	•	COC	
676	RX HospReg LN Removed	•	•	RH	•	•	COC	
680	Reserved 03	•	•	•	•	•		
690	RX HospRadiation	•	•	RH	RH	•	SEER/COC	
700	RX HospChemo	•	R	R	R	•	COC	
710	RX HospHormone	•	R	R	R	•	COC	
720	RX HospBRM	•	R	R	R	•	COC	
730	RX HospOther	•	R	R	R	•	COC	
740	RX HospDX/Stg Proc	•	R	R	•	•	COC	
742	RX HospScreen/BX Proc1	•	•	•	•	•	COC	
743	RX HospScreen/BX Proc2	•	•	•	•	•	COC	
744	RX HospScreen/BX Proc3	•	•	•	•	•	COC	
745	RX HospScreen/BX Proc4	•	•	•	•	•	COC	
746	RX HospSurg Site 98-02	•	•	RH	RH	•	COC	
747	RX HospScope Reg 98-02	•	•	RH	RH	•	COC	
748	RX HospSurg Oth 98-02	•	•	RH	RH	•	COC	
750	Reserved 04	•	•	•	•	•		
759	SEER Summary Stage 2000	RH	R	R	•	•	SEER	

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\* = When available. # = Central registries may code available data using either the SEER or COC data item and associated rules.

^ = These text requirements may be met with one or several text block fields.

<b>T</b> .	T. N	NIDCID	<u>C</u>	<u>OC</u>	S	EER	Source of	
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
760	SEER Summary Stage 1977	RH	RH	RH	•	•	SEER	
770	Loc/Reg/Distant Stage	•	•	•	•	•	Varies	
779	Extent of Disease 10-Dig							
780	EODTumor Size	•	•	RH	RH	RH	SEER/COC	
790	EODExtension	•	•	•	RH	RH	SEER	
800	EODExtension Prost Path	•	•	•	RH	RH	SEER	
810	EODLymph Node Involv	•	•	•	RH	RH	SEER	
820	Regional Nodes Positive	S	R	R	R	R	SEER/COC	
830	Regional Nodes Examined	S	R	R	R	R	SEER/COC	
840	EODOld 13 Digit	•	•	•	RH	RH	SEER	
850	EODOld 2 Digit	•	•	•	RH	RH	SEER	
860	EODOld 4 Digit	•	•	•	RH	RH	SEER	
870	Coding System for EOD	•	•	•	R	R	SEER	
880	TNM Path T	•	R	R	•	•	AJCC	
890	TNM Path N	•	R	R	•	•	AJCC	
900	TNM Path M	•	R	R	•	•	AJCC	
910	TNM Path Stage Group	•	R	R	•	•	AJCC	
920	TNM Path Descriptor	•	R	R	•	•	COC	
930	TNM Path Staged By	•	R	R	•	•	COC	
940	TNM Clin T	•	R	R	•	•	AJCC	
950	TNM Clin N	•	R	R	•	•	AJCC	
960	TNM Clin M	•	R	R	•	•	AJCC	
970	TNM Clin Stage Group	•	R	R	•	•	AJCC	
980	TNM Clin Descriptor	•	R	R	•	•	COC	
990	TNM Clin Staged By	•	R	R	•	•	COC	
1000	TNM Other T	•	•	•	•	•	AJCC	
1010	TNM Other N	•	•	•	•	•	AJCC	
1020	TNM Other M	•	٠	•	•	•	AJCC	
1030	TNM Other Stage Group	•	•	•	•	•	AJCC	
1040	TNM Other Staged By	•	•	•	•	•	COC	
1050	TNM Other Descriptor	•	•	•	•	•	COC	
1060	TNM Edition Number	•	R	R	•	•	COC	
1070	Other Staging System	•	•	•	•	•	COC	
1080	Date of 1st Positive BX	•	•	•	•	•	COC	

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Itaan	Itam Nama	NDCD	<u>C</u>	<u>OC</u>	S	EER	Source of	Nata
Item :	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
1090	Site of Distant Met 1	•	•	RH	•	•	COC	
1100	Site of Distant Met 2	•	•	RH	•	•	COC	
1110	Site of Distant Met 3	•	•	RH	•	•	COC	
1120	Pediatric Stage	•	•	•	•	•	COC	
1130	Pediatric Staging System	•	•	•	•	•	COC	
1140	Pediatric Staged By	•	•	•	•	•	COC	
1150	Tumor Marker 1	•	RH	RH	RH	RH	SEER	
1160	Tumor Marker 2	•	RH	RH	RH	RH	SEER	
1170	Tumor Marker 3	•	RH	RH	RH	RH	SEER	
1180	Reserved 05	•	•	•	•	•		
1190	Reserved 06	•	•	•	•	•		
1200	RX DateSurgery	S	R	R	•	•	COC	
1210	RX DateRadiation	S	R	R	•	•	COC	
1220	RX DateChemo	•	•	•	•	•	NAACCR	
1230	RX DateHormone	•	•	•	•	•	NAACCR	
1240	RX DateBRM	•	•	•	•	•	NAACCR	
1250	RX DateOther	S	R	R	•	•	COC	
1260	Date of Initial RXSEER	#	•	•	R	R	SEER	
1270	Date of 1st Crs RXCOC	#	R	R	•	•	COC	
1280	RX DateDX/Stg Proc	•	R	R	•	•	COC	
1290	RX SummSurg Prim Site	R	R	R	R	R	SEER/COC	
1292	RX SummScope Reg LN Sur	R	R	R	R	R	SEER/COC	
1294	RX SummSurg Oth Reg/Dis	R	R	R	R	R	SEER/COC	
1296	RX SummReg LN Examined	RH	•	RH	RH	RH	SEER/COC	
1300	Reserved 07	•	•	•	•	•		
1310	RX SummSurgical Approch	•	•	RH	•	•	COC	
1320	RX SummSurgical Margins	•	R	R	•	•	COC	
1330	RX SummReconstruct 1st	•	•	•	RH	RH	SEER	
1340	Reason for No Surgery	S	R	R	R	R	SEER/COC	
1350	RX SummDX/Stg Proc	•	R	R	•	•	COC	
1355	Reserved 22							
1360	RX SummRadiation	•	•	RH	R	R	SEER	
1370	RX SummRad to CNS	•	•	•	RH	RH	SEER/COC	
1380	RX SummSurg/Rad Seq	S	R	R	R	R	SEER/COC	

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T.c	I.c. N.c	NDCD	<u>C</u>	<u>OC</u>	S	EER	Source of	Nut
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
1390	RX SummChemo	S	R	R	R	R	SEER/COC	
1400	RX SummHormone	S	R	R	R	R	SEER/COC	
1410	RX SummBRM	S	R	R	R	R	SEER/COC	
1420	RX SummOther	S	R	R	R	R	SEER/COC	
1430	Reason for No Radiation	S	R	R	•	•	COC	
1440	Reason for No Chemo	•	•	•	•	•	COC	
1450	Reason for No Hormone	•	•	•	•	•	COC	
1460	RX Coding SystemCurrent	R	R	R	•	RH	NAACCR	
1470	Protocol Eligibility Stat	•	•	•	•	•	COC	
1480	Protocol Participation	•	•	•	•	•	COC	
1490	Referral to Support Serv	•	•	•	•	•	COC	
1500	First Course Calc Method	•	•	•	•	•	NAACCR	
1510	RadRegional Dose: CGY	•	R	R	•	•	COC	
1520	RadNo of Treatment Vol	•	R	R	•	•	COC	
1530	RadElapsed RX Days	•	•	•	•	•	COC	
1540	RadTreatment Volume	•	R	R	•	•	COC	
1550	RadLocation of RX	•	R	R	•	•	COC	
1560	RadIntent of Treatment	•	•	•	•	•	COC	
1570	RadRegional RX Modality	S	R	R	RC	•	COC	
1580	RadRX Completion Status	•	•	•	•	•	COC	
1590	RadLocal Control Status	•	•	•	•	•	COC	
1600	Chemotherapy Field 1	•	•	•	•	•		Retired
1610	Chemotherapy Field 2	•	•	•	•	•		Retired
1620	Chemotherapy Field 3	•	•	•	•	•		Retired
1630	Chemotherapy Field 4	•	•	•	•	•		Retired
1635	Reserved 23							
1640	RX SummSurgery Type	•	•	•	RH	RH	SEER	
1642	RX SummScreen/BX Proc1	•	•	•	•	•	COC	
1643	RX SummScreen/BX Proc2	•	•	•	•	•	COC	
1644	RX SummScreen/BX Proc3	•	•	•	٠	•	COC	
1645	RX SummScreen/BX Proc4	•	•	•	•	•	COC	
1646	RX SummSurg Site 98-02	RH	•	RH	RH	RH	SEER/COC	
1647	RX SummScope Reg 98-02	RH	•	RH	RH	RH	SEER/COC	
1648	RX SummSurg Oth 98-02	RH	•	RH	RH	RH	SEER/COC	

\* = When available. # = Central registries may code available data using either the SEER or COC data item and associated rules.

T	<b>1</b> 4 <b>N</b>	NDCD	<u>C</u>	<u>OC</u>	<u>S</u>	EER	Source of	Nut
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
1650	Reserved 08	•	•	•	•	•		
1660	Subsq RX 2nd Course Date	•	•	•	•	•	COC	
1670	Subsq RX 2nd Course Codes							
1671	Subsq RX 2nd Course Surg	•	•	•	•	•	COC	
1672	Subsq RX 2nd Course Rad	•	•	•	•	•	COC	
1673	Subsq RX 2nd Course Chemo	•	•	•	•	•	COC	
1674	Subsq RX 2nd Course Horm	•	•	•	•	•	COC	
1675	Subsq RX 2nd Course BRM	•	•	•	•	•	COC	
1676	Subsq RX 2nd Course Oth	•	•	•	•	•	COC	
1677	Subsq RX 2ndScope LN SU	•	•	•	•	•	COC	
1678	Subsq RX 2ndSurg Oth	•	•	•	•	•	COC	
1679	Subsq RX 2ndReg LN Rem	•	•	•	•	•	COC	
1680	Subsq RX 3rd Course Date	•	•	•	•	•	COC	
1690	Subsq RX 3rd Course Codes							
1691	Subsq RX 3rd Course Surg	•	•	•	•	•	COC	
1692	Subsq RX 3rd Course Rad	•	•	•	•	•	COC	
1693	Subsq RX 3rd Course Chemo	•	•	•	•	•	COC	
1694	Subsq RX 3rd Course Horm	•	•	•	•	•	COC	
1695	Subsq RX 3rd Course BRM	•	•	•	•	•	COC	
1696	Subsq RX 3rd Course Oth	•	•	•	•	•	COC	
1697	Subsq RX 3rdScope LN SU	•	•	•	•	•	COC	
1698	Subsq RX 3rdSurg Oth	•	•	•	•	•	COC	
1699	Subsq RX 3rdReg LN Rem	•	•	•	•	•	COC	
1700	Subsq RX 4th Course Date	•	•	•	•	•	COC	
1710	Subsq RX 4th Course Codes							
1711	Subsq RX 4th Course Surg	•	•	•	•	•	COC	
1712	Subsq RX 4th Course Rad	•	•	•	•	•	COC	
1713	Subsq RX 4th Course Chemo	•	•	•	•	•	COC	
1714	Subsq RX 4th Course Horm	•	•	•	•	•	COC	
1715	Subsq RX 4th Course BRM	•	•	•	•	•	COC	
1716	Subsq RX 4th Course Oth	•	•	•	•	•	COC	
1717	Subsq RX 4thScope LN SU	•	•	•	•	•	COC	
1718	Subsq RX 4thSurg Oth	•	•	•	•	•	COC	
1719	Subsq RX 4thReg LN Rem	•	•	•	•	•	COC	1

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<b>T</b> /	T. N	NDCD	C	<u>OC</u>	S	EER	Source of	
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
1720	Subsq RX 5th Course Date	•	•	•	•	•	NAACCR	
1730	Subsq RX 5th Course Codes							
1731	Subsq RX 5th Course Surg	•	•	•	•	•	NAACCR	
1732	Subsq RX 5th Course Rad	•	•	•	•	•	NAACCR	
1733	Subsq RX 5th Course Chemo	•	•	•	•	•	NAACCR	
1734	Subsq RX 5th Course Horm	•	•	•	•	•	NAACCR	
1735	Subsq RX 5th Course BRM	•	•	•	•	•	NAACCR	
1736	Subsq RX 5th Course Oth	•	•	•	•	•	NAACCR	
1737	Subsq RX 5thScope LN SU	•	•	•	•	•	NAACCR	
1738	Subsq RX 5thSurg Oth	•	•	•	•	•	NAACCR	
1739	Subsq RX 5thReg LN Rem	•	•	•	•	•	NAACCR	
1740	Reserved 09	•	•	•	•	•		
1741	Subsq RXReconstruct Del	•	•	•	•	•	COC	
1750	Date of Last Contact	R	R	R	R	R	SEER/COC	
1760	Vital Status	R	R	R	R	R	SEER/COC	
1770	Cancer Status	•	R	R	•	•	COC	
1780	Quality of Survival	•	•	•	•	•	COC	
1790	Follow-Up Source	•	R	R	•	•	COC	
1800	Next Follow-Up Source	•	R	•	•	•	COC	
1810	Addr CurrentCity	•	R	•	R	•	COC	
1820	Addr CurrentState	•	R	•	R	•	COC	
1830	Addr CurrentPostal Code	•	R	•	R	•	COC	
1835	Reserved 10	•	•	•	•	•		
1840	CountyCurrent	•	•	•	•	•	NAACCR	
1842	Follow-Up ContactCity	•	٠	•	R	•	SEER	
1844	Follow-Up ContactState	•	•	•	R	•	SEER	
1846	Follow-Up ContactPostal	•	•	•	R	•	SEER	
1850	Unusual Follow-Up Method	•	•	•	•	•	COC	
1860	Recurrence Date1st	S	R	R	RC	•	COC	
1870	Recurrence Distant Sites							Retired
1871	Recurrence Distant Site 1	•	•	•	•	•	NAACCR	
1872	Recurrence Distant Site 2	•	•	•	•	•	NAACCR	
1873	Recurrence Distant Site 3	•	•	•	•	•	NAACCR	
1880	Recurrence Type1st	S	R	R	RC	•	COC	

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I4	Harry Name	NDCD	<u>C</u>	<u>OC</u>	S	EER	Source of	Nata
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
1890	Recurrence Type1stOth	•	•	•	•	•	COC	
1900	Reserved 11	•	•	•	•	•		
1910	Cause of Death	R	•	•	R	R	SEER	
1920	ICD Revision Number	R	•	•	R	R	SEER	
1930	Autopsy	•	•	•	•	•	NAACCR	
1940	Place of Death	S	•	•	•	•	NPCR	
1950	Reserved 12	•	•	•	•	•		
1960	Site (73-91) ICD-O-1	•	•	RH	RH	RH	SEER	
1970	Morph (73-91) ICD-O-1							
1971	Histology (73-91) ICD-O-1	•	•	RH	RH	RH	SEER	
1972	Behavior (73-91) ICD-O-1	•	•	RH	RH	RH	SEER	
1973	Grade (73-91) ICD-O-1	•	•	RH	RH	RH	SEER	
1980	ICD-O-2 Conversion Flag	•	R	R	RH	RH	SEER	
1981	Over-ride SS/NodesPos	•	•	•	•	•	NAACCR	
1982	Over-ride SS/TNM-N	•	•	•	•	•	NAACCR	
1983	Over-ride SS/TNM-M	•	•	•	•	•	NAACCR	
1984	Over-ride SS/DisMet1	•	•	•	•	•	NAACCR	
1985	Over-ride Acsn/Class/Seq	•	R	R	•	•	COC	
1986	Over-ride HospSeq/DxConf	•	R	R	•	•	COC	
1987	Over-ride COC-Site/Type	•	R	R	•	•	COC	
1988	Over-ride HospSeq/Site	•	R	R	•	•	COC	
1989	Over-ride Site/TNM-StgGrp	•	R	R	•	•	COC	
1990	Over-ride Age/Site/Morph	R	R	R	R	R	SEER	
2000	Over-ride SeqNo/DxConf	R	•	•	R	R	SEER	
2010	Over-ride Site/Lat/SeqNo	S	•	•	R	R	SEER	
2020	Over-ride Surg/DxConf	R	R	R	R	R	SEER	
2030	Over-ride Site/Type	R	R	R	R	R	SEER	
2040	Over-ride Histology	R	R	R	R	R	SEER	
2050	Over-ride Report Source	R	•	•	R	R	SEER	
2060	Over-ride Ill-define Site	R	•	•	R	R	SEER	
2070	Over-ride Leuk, Lymphoma	R	R	R	R	R	SEER	
2071	Over-ride Site/Behavior	R	R	R	R	R	SEER	
2072	Over-ride Site/EOD/DX Dt	S	•	•	R	R	SEER	
2073	Over-ride Site/Lat/EOD	S	•	•	R	R	SEER	

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<b>.</b>	<b>.</b>	NDCD	C	<u>OC</u>	S	EER	Source of	
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
2074	Over-ride Site/Lat/Morph	R	R	R	R	R	SEER	
2080	Reserved 13							Retired
2081	CRC CHECKSUM	•	•	•	•	•	NAACCR	
2082	Reserved 24							
2090	Date Case Completed	•	•	•	•	•	NAACCR	
2100	Date Case Last Changed	•	•	•	•	•	NAACCR	
2110	Date Case Report Exported	S	•	R	•	•	NPCR	
2111	Date Case Report Received	R	•	•	•	•	NPCR	
2112	Date Case Report Loaded	S	•	•	•	•	NPCR	
2113	Date Tumor Record Availbl	S	•	•	•	•	NPCR	
2114	Future Use Timeliness 1	•	•	•	•	•		Retired
2115	Future Use Timeliness 2	•	•	•	•	•		Retired
2116	ICD-O-3 Conversion Flag	R	R	R	R	R	SEER/COC	
2120	SEER Coding SysCurrent	S	•	•	•	•	NAACCR	
2130	SEER Coding SysOriginal	S	•	•	•	•	NAACCR	
2140	COC Coding SysCurrent	S	R	R	•	•	COC	
2150	COC Coding SysOriginal	S	R	R	•	•	COC	
2160	Subsq Report for Primary	•	•	•	•	•		Retired
2161	Reserved 20	•	٠	•	•	•		Retired
2170	Vendor Name	•	•	R	•	•	NAACCR	
2180	SEER Type of Follow-Up	•	•	•	R	R	SEER	
2190	SEER Record Number	•	•	•	•	R	SEER	
2200	Diagnostic Proc 73-87	•	•	•	RH	RH	SEER	
2210	Reserved 14	•	•	•	•	•		Retired
2220	State/Requestor Items	•	•	•	•	•	Varies	
2230	NameLast	R	R	•	R	•	NAACCR	
2240	NameFirst	R	R	•	R	•	NAACCR	
2250	NameMiddle	R	R	•	R	•	COC	
2260	NamePrefix	•	•	•	•	•	SEER	
2270	NameSuffix	•	•	•	R	•	SEER	
2280	NameAlias	S	٠	•	R	•	SEER	
2290	NameSpouse/Parent	•	٠	•	•	•	NAACCR	
2300	Medical Record Number	S	R	•	R	•	COC	
2310	Military Record No Suffix	•	R	•	•	•	COC	

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Items	Iterra Norma	NDCD	<u>C</u>	<u>OC</u>	S	EER	Source of	Nata
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
2320	Social Security Number	R	R	•	R	•	COC	
2330	Addr at DXNo & Street	S	R	•	R	•	COC	
2335	Addr at DXSupplementl	S	R	•	•	•	COC	
2350	Addr CurrentNo & Street	•	R	•	R	•	COC	
2352	Latitude	•	•	•	•	•	NAACCR	
2354	Longitude	•	•	•	•	•	NAACCR	
2355	Addr CurrentSupplementl	•	R	•	•	•	COC	
2360	Telephone	•	R	•	R	•	COC	
2370	DC State	•	•	•	•	•		Retired
2371	Reserved 21							Retired
2380	DC State File Number	S	•	•	•	•	State	
2390	NameMaiden	S	•	•	R	•	SEER	
2392	Follow-Up ContactNo&St	•	•	•	R	•	SEER	
2393	Follow-Up ContactSuppl	•	•	•	•	•	SEER	
2394	Follow-Up ContactName	•	•	•	R	•	SEER	
2400	Reserved 16	•	•	•	•	•		Retired
2410	Institution Referred From	•	R	•	•	•	COC	
2420	Institution Referred To	•	R	•	•	•	COC	
2430	Last Follow-Up Hospital	•	•	•	•	•	NAACCR	
2440	Following Registry	•	R	•	R	•	COC	
2450	Reserved 17	•	•	•	•	•		Retired
2460	PhysicianManaging	•	•	•	•	•	NAACCR	
2470	PhysicianFollow-Up	•	R	•	R	•	COC	
2480	PhysicianPrimary Surg	•	R	•	•	•	COC	
2490	Physician 3	•	R	•	•	•	COC	
2500	Physician 4	•	R	•	•	•	COC	
2520	TextDX ProcPE	R^	•	•	R	•	NPCR	
2530	TextDX ProcX-ray/Scan	R^	•	•	R	•	NPCR	
2540	TextDX ProcScopes	R^	•	•	R	•	NPCR	
2550	TextDX ProcLab Tests	R^	•	•	R	•	NPCR	
2560	TextDX ProcOp	R^	•	•	R	•	NPCR	
2570	TextDX ProcPath	R^	•	•	R	•	NPCR	
2580	TextPrimary Site Title	S	•	•	R	•	NPCR	
2590	TextHistology Title	S	•	•	R	•	NPCR	

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T.	L. N	NIDCD	<u>C</u>	<u>OC</u>	S	EER	Source of	
Item	Item Name	<u>NPCR</u>	Collect	Transmit	Collect	Transmit	Standard	Note
2600	TextStaging	R^	•	•	R	•	NPCR	
2610	RX TextSurgery	R^	•	•	R	•	NPCR	
2620	RX TextRadiation (Beam)	S	•	•	R	•	NPCR	
2630	RX TextRadiation Other	S	•	•	R	•	NPCR	
2640	RX TextChemo	S	•	•	R	•	NPCR	
2650	RX TextHormone	S	•	•	R	•	NPCR	
2660	RX TextBRM	S	•	•	R	•	NPCR	
2670	RX TextOther	S	•	•	R	•	NPCR	
2680	TextRemarks	S	•	•	R	•	NPCR	
2690	Place of Diagnosis	S	•	•	•	•	NPCR	
2700	Reserved 19	•	•	•	•	•		
2800	CS Tumor Size	S	R	R	R	R	AJCC	
2810	CS Extension	R	R	R	R	R	AJCC	
2820	CS Tumor Size/Ext Eval	S	R	R	•	•	AJCC	
2830	CS Lymph Nodes	R	R	R	R	R	AJCC	
2840	CS Reg Node Eval	S	R	R	•	•	AJCC	
2850	CS Mets at DX	R	R	R	R	R	AJCC	
2860	CS Mets Eval	S	R	R	•	•	AJCC	
2880	CS Site-Specific Factor 1	RS	R	R	R	R	AJCC	
2890	CS Site-Specific Factor 2	S	R	R	R	R	AJCC	
2900	CS Site-Specific Factor 3	RS	R	R	R	R	AJCC	
2910	CS Site-Specific Factor 4	S	R	R	R	R	AJCC	
2920	CS Site-Specific Factor 5	S	R	R	R	R	AJCC	
2930	CS Site-Specific Factor 6	S	R	R	R	R	AJCC	
2935	CS Version 1st	R	R	R	R	R	AJCC	
2936	CS Version Latest	R	R	R	R	R	AJCC	
2940	Derived AJCC T	D	R	R	D	D	AJCC	
2950	Derived AJCC T Descriptor	D	R	R	•	•	AJCC	
2960	Derived AJCC N	D	R	R	D	D	AJCC	
2970	Derived AJCC N Descriptor	D	R	R	•	•	AJCC	
2980	Derived AJCC M	D	R	R	D	D	AJCC	
2990	Derived AJCC M Descriptor	D	R	R	•	•	AJCC	
3000	Derived AJCC Stage Group	D	R	R	D	D	AJCC	
3010	Derived SS1977	D	R	R	D	D	AJCC	

\* = When available. # = Central registries may code available data using either the SEER or COC data item and associated rules.

Item :	Item Name	NPCR	<u>C</u>	<u>OC</u>	<u>S</u>	<u>EER</u>	Source of	Note
item :	item ivanie	MICK	Collect	Transmit	Collect	Transmit	Standard	Note
3020	Derived SS2000	D	R	R	D	D	AJCC	
3030	Derived AJCCFlag	D	R	R	D	D	AJCC	
3040	Derived SS1977Flag	D	R	R	D	D	AJCC	
3050	Derived SS2000Flag	D	R	R	D	D	AJCC	
3100	Archive FIN	•	R	R	•	•	COC	
3110	Comorbid/Complication 1	•	R	R	•	•	COC	
3120	Comorbid/Complication 2	•	R	R	•	•	COC	
3130	Comorbid/Complication 3	•	R	R	•	•	COC	
3140	Comorbid/Complication 4	•	R	R	•	•	COC	
3150	Comorbid/Complication 5	•	R	R	•	•	COC	
3160	Comorbid/Complication 6	•	R	R	•	•	COC	
3170	RX DateMost Defin Surg	S	R	R	•	•	COC	
3180	RX DateSurgical Disch	•	R	R	•	•	COC	
3190	Readm Same Hosp 30 Days	•	R	R	•	•	COC	
3200	RadBoost RX Modality	•	R	R	RC	•	COC	
3210	RadBoost Dose cGy	•	R	R	•	•	COC	
3220	RX DateRadiation Ended	•	R	R	•	•	COC	
3230	RX DateSystemic	S	R	R	•	•	COC	
3250	RX SummTransplnt/Endocr	S	R	R	R	R	COC	
3260	Pain Assessment	•	•	•	•	•		Retired
3270	RX SummPalliative Proc	•	R	R	•	•	COC	
3280	RX HospPalliative Proc	•	R	R	•	•	COC	
3300	RuralUrban Continuum 1993	D	•	•	•	•	NAACCR	
3310	RuralUrban Continuum 2000	D	•	•	•	•	NAACCR	

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# **CHAPTER X**

## DATA DESCRIPTOR TABLE (ITEM # ORDER)

The following table presents Version 10.2 of the NAACCR data descriptor table summarizing the item number, item name, format, allowable values, and length of each item. The data type for all data items is "character." Differences from Version 10.1 are marked "Revised," "New," or "Retired" in the "Note" column of the table. Some changes also are summarized in Appendix F. A program that generates a file of records in the NAACCR data exchange format should handle instances where information is unavailable for any given field.

General Rules:

When ALL of the records in the file to be generated contain no information on a specific data item, then the corresponding columns in the exchange record should be left as blanks.

*Example:* You are submitting data in NAACCR 10.2 format, but your registry does not collect data on AJCC stage. The columns in the file you generate that are supposed to contain the information on AJCC stage should all contain blanks.

 When some of the records contain information for a given field, and other records will not contain information for that field, then the code that indicates "unknown," "not available," or "not applicable" (as appropriate) must be written in the corresponding columns in the exchange record.

*Example:* You are submitting data in NAACCR 10.2 format, and you collect information on surgery date. However, in some cases the date is not there because your program stores it as a date-time variable and either no surgery was given, it is unknown whether surgery was given, or it was an autopsy or death certificate-only (DCO) case. Those columns in the file you generate must contain no blanks; instead, the columns should contain "99999999" when it is unknown whether or not surgery was given or when the case was DCO, and "00000000" when no surgery was given or autopsy-only.

*Exception:* You are submitting in the NAACCR 10.2 format, and cases diagnosed in the years 1997-2001 are included. The Morph--Type&Behavior ICD-O-2 fields should contain the original ICD-O-2 codes for cases diagnosed in or before 2000, but the fields should be blank for cases diagnosed in 2001 (unless you have back-translated the ICD-O-3 morphology codes).

All "blanks" must be transmitted as the appropriate number of "spaces" (ASCII 20h), never as nulls or as numeric fields with no value assigned. Nulls may shift the record contents out of column alignment, and numeric fields with no value assigned to them erroneously transmit zeroes as code content.

Date fields are recorded in the month, day, year format (MMDDCCYY), with 99 for unknown day or month and 9999 for unknown year. For example:

- 00000000 No date
- 99999999 Unknown date
- 01992003 Example of date when the month and year are known but the day is unknown
- 99992003 Example of date when the year is known but the month and day are unknown.

Item #	Item Name	Format	Allowable Values	Length	Note
10	Record Type		I, C, A, U, R, M, L	1	
20	Patient ID Number	Right justified, zero filled		8	
30	Registry Type		1-3	1	
35	FIN Coding System		1-3,9	1	
37	Reserved 00			7	
40	Registry ID	Right justified, zero filled	10-digit number. Reference to EDITS table REGID.DBF in Appendix B	10	
50	NAACCR Record Version		Blank, 1, 4-9, A	1	
60	Tumor Record Number	Right justified, zero filled	01-99	2	
70	Addr at DXCity	Mixed case letters, special characters only as allowed by USPS, embedded spaces allowed, left justified, blank filled	City Name or UNKNOWN	20	Revised
80	Addr at DXState	Upper case	Refer to EDITS table STATE.DBF in Appendix B	2	
90	County at DX	Right justified, zero filled	See Appendix A for county codes for each state. For non-U.S. residents, COC uses Appendix B (BPLACE.DBF). Also 998, 999	3	
100	Addr at DXPostal Code	Numbers or upper case letters. No special characters or embedded spaces allowed. Left justified, blank filled	5-digit or 9-digit U.S. ZIP codes; 6-character Canadian postal codes; valid postal codes from other countries, 888888888, 999999999, 88888+4 blanks, 99999+4blanks	9	
110	Census Tract 1970/80/90	Right justified, zero filled	Census Tract Codes 000100-949999, BNA Codes 950100-998999, 000000, 999999, blank	6	
120	Census Cod Sys 1970/80/90		0-3, blank	1	
130	Census Tract 2000	Right justified, zero filled	Census Tract Codes 000101-999998, 000000, 999999, blank	6	
140	Census Tract Cod SysAlt				Retired
150	Marital Status at DX		1-5, 9	1	
160	Race 1	Right justified, zero filled	01-14, 20-22, 25-28, 30-32, 96-99	2	
161	Race 2	Right justified, zero filled	01-14, 20-22, 25-28, 30-32, 88, 96-99, blank	2	
162	Race 3	Right justified, zero filled	01-14, 20-22, 25-28, 30-32, 88, 96-99, blank	2	
163	Race 4	Right justified, zero filled	01-14, 20-22, 25-28, 30-32, 88, 96-99, blank	2	
164	Race 5	Right justified, zero filled	01-14, 20-22, 25-28, 30-32, 88, 96-99, blank	2	
170	Race Coding SysCurrent		1-6, 9	1	
180	Race Coding SysOriginal		1-6, 9	1	
190	Spanish/Hispanic Origin		0-8, 9	1	Revised
191	NHIA Derived Hisp Origin		0-8, blank	1	New

Item #	Item Name	Format	Allowable Values	Length	Note
200	Computed Ethnicity		0-7, blank	1	
210	Computed Ethnicity Source		0-9, blank	1	
220	Sex		1-4, 9	1	
230	Age at Diagnosis	Right justified, zero filled	000-120, 999	3	
240	Birth Date	MMDDCCYY	Valid date or 99999999	8	
250	Birthplace	Right justified, zero filled	Reference to EDITS table BPLACE.DBF in Appendix B	3	Revised
260	Religion	No standard	Any	2	
270	Occupation CodeCensus		Reference Industry and Occupation Coding for Death Certificates	3	
280	Industry CodeCensus		Reference Industry and Occupation Coding for Death Certificates	3	
290	Occupation Source		0-3, 7-9, blank	1	
300	Industry Source		0-3, 7-9, blank	1	
310	TextUsual Occupation	Free text	Neither carriage return nor line feed characters allowed	40	
320	TextUsual Industry	Free text	Neither carriage return nor line feed characters allowed	40	
330	Occup/Ind Coding System		1-4, 7, 9, blank	1	
340	Tobacco History	No standard	Any	1	
350	Alcohol History	No standard	Any	1	
360	Family History of Cancer	No standard	Any	1	
362	Census Tract Block Group	No standard	Refer to Census Bureau	1	
364	Census Tr Cert 1970/80/90		1-5, 9, blank	1	
365	Census Tr Certainty 2000		1-5, 9, blank	1	
370	Reserved 01			20	
380	Sequence NumberCentral	Right justified, zero filled	00-35, 60-87, 88, 98, 99	2	
390	Date of Diagnosis	MMDDCCYY	Valid date or 99999999	8	
400	Primary Site	C followed by 3 digits, no special characters, no embedded blanks	Reference ICD-O-3 for valid entries	4	
410	Laterality		0-4, 9	1	
419	MorphType&Behav ICD- O-2		Reference to ICD-0-2	5	
420	Histology (92-00) ICD-O-2		Reference to ICD-0-2	4	
430	Behavior (92-00) ICD-O-2		Reference to ICD-0-2	1	Revised
440	Grade		1-9	1	
450	Site Coding SysCurrent		1-6, 9	1	
460	Site Coding SysOriginal		1-6, 9	1	
470	Morph Coding SysCurrent		1-7, 9	1	
480	Morph Coding SysOriginl		1-7, 9	1	
490	Diagnostic Confirmation		1, 2, 4-9	1	
500	Type of Reporting Source		1, 3-7	1	
510	Screening Date	MMDDCCYY	Valid date, 00000000, 99999999	8	
520	Screening Result		0-4, 8, 9	1	

Item #	Item Name	Format	Allowable Values	Length	Note
521	MorphType&Behav ICD- O-3		Reference to ICD-O-3	5	
522	Histologic Type ICD-O-3		Reference to ICD-O-3	4	
523	Behavior Code ICD-O-3		Reference to ICD-O-3	1	Revised
530	Reserved 02			50	
538	Reporting Hospital FAN			10	
540	Reporting Hospital	Right justified, zero filled	10-digit number	10	
550	Accession NumberHosp		9-digit number	9	
560	Sequence NumberHospital	Right justified, zero filled	00-35, 60-87, 88, 99	2	
570	Abstracted By	No special characters	Letters and numbers	3	
580	Date of 1st Contact	MMDDCCYY	Valid dates or 99999999	8	
590	Date of Inpatient Adm	MMDDCCYY	Valid dates 00000000, 99999999	8	
600	Date of Inpatient Disch	MMDDCCYY	Valid dates, 00000000, 99999999	8	
610	Class of Case		0-9	1	
620	Year First Seen This CA	ССҮҮ	1944 to current year	4	
630	Primary Payer at DX	Right justified, zero filled	01, 02, 10, 20, 31, 35, 36, 50-56, 99	2	
640	Inpatient/Outpt Status		1-3, 8, 9	1	
650	Presentation at CA Conf		0-9	1	
660	Date of CA Conference	MMDDCCYY	Valid dates, 00000000, 99999999	8	
670	RX HospSurg Prim Site	Right justified, zero filled	00, 10-80, 90, 98, 99 (site specific)	2	
672	RX HospScope Reg LN Sur		0-7,9	1	
674	RX HospSurg Oth Reg/Dis		0-5,9	1	
676	RX HospReg LN Removed		00-90, 95-99	2	
680	Reserved 03			50	
690	RX HospRadiation		0-5,9	1	
700	RX HospChemo	Right justified, zero filled	00-03, 82, 85-88, 99	2	
710	RX HospHormone	Right justified, zero filled	00, 01, 82, 85-88, 99	2	
720	RX HospBRM	Right justified, zero filled	00, 01, 82, 85-88, 99	2	
730	RX HospOther		0-3, 6-9	1	
740	RX HospDX/Stg Proc	Right justified, zero filled	00-07, 09	2	
742	RX HospScreen/BX Proc1		Site-specific: 0 (all cases); 1-3, 5, 9 (breast); 1-4, 9 (prostate)	1	
743	RX HospScreen/BX Proc2		Site-specific: 0 (all cases); 1-7, 9 (breast); 1-3, 9 (prostate)	1	
744	RX HospScreen/BX Proc3		Site-specific: 0 (all cases); 1, 9 (breast); 1-5, 9 (prostate)	1	
745	RX HospScreen/BX Proc4		Site-specific: 0 (all cases); 1-4, 9 (breast); 1-7, 9 (prostate)	1	
746	RX HospSurg Site 98-02	Right justified, zero filled	00, 10-80, 90, 98, 99 (site specific)	2	

Item #	Item Name	Format	Allowable Values	Length	Note
747	RX HospScope Reg 98-02		0-7, 9	1	
748	RX HospSurg Oth 98-02		0-5, 9	1	
750	Reserved 04			46	
759	SEER Summary Stage 2000		0-5, 7, 8, 9	1	
760	SEER Summary Stage 1977		0-5, 7, 8, 9	1	
770	Loc/Reg/Distant Stage		0-3, 9, blank	1	
779	Extent of Disease 10-Dig			12	
780	EODTumor Size	Right justified, zero filled	See respective source references	3	
790	EODExtension	Right justified, zero filled	Reference SEER Extent of Disease manual	2	
800	EODExtension Prost Path	Right justified, zero filled	Reference SEER Extent of Disease manual	2	
810	EODLymph Node Involv		Reference SEER Extent of Disease manual	1	
820	Regional Nodes Positive	Right justified, zero filled	See respective source references	2	
830	Regional Nodes Examined	Right justified, zero filled	See respective source references	2	
840	EODOld 13 Digit	Numeric and special characters		13	
850	EODOld 2 Digit	Numeric plus special characters "&" and "dash" ("-")		2	
860	EODOld 4 Digit			4	
870	Coding System for EOD		0-4	1	
880	TNM Path T	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
890	TNM Path N	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
900	TNM Path M	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
910	TNM Path Stage Group	Upper case, alphanumeric. Convert AJCC Roman numerals to Arabic numerals. Left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, 99, blank	2	
920	TNM Path Descriptor		0-6, 9	1	
930	TNM Path Staged By		0-9	1	
940	TNM Clin T	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
950	TNM Clin N	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
960	TNM Clin M	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, blank	2	
970	TNM Clin Stage Group	Upper case, alphanumeric. Convert AJCC Roman numerals to Arabic numerals. Left justified, blank filled	See AJCC Cancer Staging Manual and FORDS Manual; also 88, 99, blank	2	
980	TNM Clin Descriptor		0-6, 9	1	
990	TNM Clin Staged By		0-9	1	

Item #	Item Name	Format	Allowable Values	Length	Note
1000	TNM Other T	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual; also 88, blank	2	
1010	TNM Other N	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual; also 88, blank	2	
1020	TNM Other M	Upper case, alphanumeric, left justified, blank filled	See AJCC Cancer Staging Manual; also 88, blank	2	
1030	TNM Other Stage Group	Upper case, alphanumeric. Convert AJCC Roman numerals to Arabic numerals. Left justified, blank filled	See AJCC Cancer Staging Manual; also 88, 99, blank	2	
1040	TNM Other Staged By		0-9	1	
1050	TNM Other Descriptor		0-6, 9	1	
1060	TNM Edition Number	Right justified, zero filled	00-06, 88, 99	2	
1070	Other Staging System	Free text	Neither carriage return nor line feed characters allowed	15	
1080	Date of 1st Positive BX	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1090	Site of Distant Met 1		0-9	1	
1100	Site of Distant Met 2		0-9	1	
1110	Site of Distant Met 3		0-9	1	
1120	Pediatric Stage		Reference to EDITS table PEDSTAGE.DBF.CODE in Appendix B	2	
1130	Pediatric Staging System		00-15, 88, 97, 99	2	
1140	Pediatric Staged By		0-9	1	
1150	Tumor Marker 1		0-6, 8, 9	1	
1160	Tumor Marker 2		0-6, 8, 9	1	
1170	Tumor Marker 3		0-6, 8, 9	1	
1180	Reserved 05			38	
1190	Reserved 06			45	
1200	RX DateSurgery	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1210	RX DateRadiation	MMDDCCYY	Valid dates, 00000000, 888888888, 999999999	8	
1220	RX DateChemo	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1230	RX DateHormone	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1240	RX DateBRM	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1250	RX DateOther	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1260	Date of Initial RXSEER	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1270	Date of 1st Crs RXCOC	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1280	RX DateDX/Stg Proc	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1290	RX SummSurg Prim Site	Right justified, zero filled	00, 10-80, 90, 98, 99 (site specific)	2	
1292	RX SummScope Reg LN Sur		0-7, 9	1	

Item #	Item Name	Format	Allowable Values	Length	Note
1294	RX SummSurg Oth Reg/Dis		0-5,9	1	
1296	RX SummReg LN Examined	Right justified, zero filled	00-90, 95-99	2	
1300	Reserved 07			50	
1310	RX SummSurgical Approch		0-9 (site-specific)	1	
1320	RX SummSurgical Margins		0-3, 7-9	1	
1330	RX SummReconstruct 1st		0-9 (site-specific)	1	
1340	Reason for No Surgery		0-2, 5-9	1	
1350	RX SummDX/Stg Proc	Right justified, zero filled	00-07, 09	2	
1355	Reserved 22			1	
1360	RX SummRadiation		0-5, 7-9	1	
1370	RX SummRad to CNS		0, 1, 7-9	1	
1380	RX SummSurg/Rad Seq		0, 2-6, 9	1	
1390	RX SummChemo	Right justified, zero filled	00-03, 82, 85-88, 99	2	Revised
1400	RX SummHormone	Right justified, zero filled	00, 01, 82, 85-88, 99	2	Revised
1410	RX SummBRM	Right justified, zero filled	00, 01, 82, 85-88, 99	2	
1420	RX SummOther		0-3, 6-9	1	
1430	Reason for No Radiation		0-2, 5-9	1	
1440	Reason for No Chemo		0-2, 6-9	1	
1450	Reason for No Hormone		0-2, 6-9	1	
1460	RX Coding SystemCurrent	Right justified, zero filled	00-06, 99	2	
1470	Protocol Eligibility Stat		0-4, 6-9	1	
1480	Protocol Participation	Right justified, zero filled	00-99	2	
1490	Referral to Support Serv		0, 1, 9	1	
1500	First Course Calc Method		1, 2, 9	1	
1510	RadRegional Dose: CGY	Right justified, zero filled	00000-99999	5	
1520	RadNo of Treatment Vol	Right justified, zero filled	00-99	2	
1530	RadElapsed RX Days	Right justified, zero filled	000-999	3	
1540	RadTreatment Volume	Right justified, zero filled	00-41, 50, 60, 98, 99	2	
1550	RadLocation of RX		0-4, 8, 9	1	
1560	RadIntent of Treatment		0-2, 4-6, 8, 9	1	
1570	RadRegional RX Modality	Right justified, zero filled	00, 20-32, 40-43, 50-55, 60- 62, 80, 85, 98, 99	2	
1580	RadRX Completion Status		0-9	1	
1590	RadLocal Control Status		0-4, 8, 9	1	
1600	Chemotherapy Field 1				Retired
1610	Chemotherapy Field 2				Retired
1620	Chemotherapy Field 3				Retired
1630	Chemotherapy Field 4				Retired
1635	Reserved 23			12	
1640	RX SummSurgery Type	Right justified, zero filled	00-99 (site-specific)	2	
1642	RX SummScreen/BX Proc1		Site-specific: 0 (all cases); 1-3, 5, 9 (breast); 1-4, 9 (prostate)	1	

Item #	Item Name	Format	Allowable Values	Length	Note
1643	RX SummScreen/BX Proc2		Site-specific: 0 (all cases); 1-7, 9 (breast); 1-3, 9 (prostate)	1	
1644	RX SummScreen/BX Proc3		Site-specific: 0 (all cases); 1, 9 (breast); 1-5, 9 (prostate)	1	
1645	RX SummScreen/BX Proc4		Site-specific: 0 (all cases); 1-4, 9 (breast); 1-7, 9 (prostate)	1	
1646	RX SummSurg Site 98-02	Right justified, zero filled	00, 10-80, 90, 98, 99 (site specific)	2	
1647	RX SummScope Reg 98-02		0-7, 9	1	
1648	RX SummSurg Oth 98-02		0-5, 9	1	
1650	Reserved 08			50	
1660	Subsq RX 2nd Course Date	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1670	Subsq RX 2nd Course Codes			7	
1671	Subsq RX 2nd Course Surg	Right justified, zero filled	00, 10-90, 99	2	
1672	Subsq RX 2nd Course Rad		0-5, 9	1	
1673	Subsq RX 2nd Course Chemo		0-3, 9	1	
1674	Subsq RX 2nd Course Horm		0-3, 9	1	
1675	Subsq RX 2nd Course BRM		0-9	1	
1676	Subsq RX 2nd Course Oth		0-3, 6-9	1	
1677	Subsq RX 2ndScope LN SU		0-9	1	
1678	Subsq RX 2ndSurg Oth		0-9	1	
1679	Subsq RX 2ndReg LN Rem	Right justified, zero filled	00-90, 95-99	2	
1680	Subsq RX 3rd Course Date	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1690	Subsq RX 3rd Course Codes			7	
1691	Subsq RX 3rd Course Surg	Right justified, zero filled	00, 10-90, 99	2	
1692	Subsq RX 3rd Course Rad		0-5,9	1	
1693	Subsq RX 3rd Course Chemo		0-3, 9	1	
1694	Subsq RX 3rd Course Horm		0-3, 9	1	
1695	Subsq RX 3rd Course BRM		0-9	1	
1696	Subsq RX 3rd Course Oth		0-3, 6-9	1	
1697	Subsq RX 3rdScope LN SU		0-9	1	
1698	Subsq RX 3rdSurg Oth		0-9	1	
1699	Subsq RX 3rdReg LN Rem	Right justified, zero filled	00-90, 95-99	2	
1700	Subsq RX 4th Course Date	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1710	Subsq RX 4th Course Codes			7	
1711	Subsq RX 4th Course Surg	Right justified, zero filled	00, 10-90, 99	2	
1712	Subsq RX 4th Course Rad	-	0-5, 9	1	
1713	Subsq RX 4th Course Chemo		0-3, 9	1	
1714	Subsq RX 4th Course Horm		0-3, 9	1	
1715	Subsq RX 4th Course BRM		0-9	1	

Item #	Item Name	Format	Allowable Values	Length	Note
1716	Subsq RX 4th Course Oth		0-3, 6-9	1	
1717	Subsq RX 4thScope LN SU		0-9	1	
1718	Subsq RX 4thSurg Oth		0-9	1	
1719	Subsq RX 4thReg LN Rem	Right justified, zero filled	00-90, 95-99	2	
1720	Subsq RX 5th Course Date	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1730	Subsq RX 5th Course Codes			7	
1731	Subsq RX 5th Course Surg	Right justified, zero filled	00, 10-90, 99	2	
1732	Subsq RX 5th Course Rad		0-5,9	1	
1733	Subsq RX 5th Course Chemo		0-3, 9	1	
1734	Subsq RX 5th Course Horm		0-3, 9	1	
1735	Subsq RX 5th Course BRM		0-9	1	
1736	Subsq RX 5th Course Oth		0-3, 6-9	1	
1737	Subsq RX 5thScope LN SU		0-9	1	
1738	Subsq RX 5thSurg Oth		0-9	1	
1739	Subsq RX 5thReg LN Rem	Right justified, zero filled	00-90, 95-99	2	
1740	Reserved 09			50	
1741	Subsq RXReconstruct Del		Site-specific	1	
1741	Date of Last Contact	MMDDCCYY	Valid dates or 99999999	8	
1760	Vital Status		0, 1, 4	1	
1700	Cancer Status		1, 2, 9	1	
1770				1	
1780	Quality of Survival		0-4, 8, 9 0-5, 7-9	1	
	Follow-Up Source		· · ·		
1800 1810	Next Follow-Up Source Addr CurrentCity	Mixed case letters, no special characters, embedded spaces allowed, left justified, blank filled	0-5, 8, 9	1 20	
1820	Addr CurrentState	Upper case	See EDITS table STATE.DBF in Appendix B	2	
1830	Addr CurrentPostal Code	Numbers or upper case letters. No special characters or embedded spaces allowed. Left justified, blank filled.	5-digit or 9-digit U.S. ZIP codes; 6- character Canadian postal codes; valid postal codes from other countries, 888888888, 999999999	9	
1835	Reserved 10			50	
1840	CountyCurrent	Right justified, zero filled	See Appendix A for standard FIPS county codes. See EDITS table BPLACE.DBF in Appendix B for geocodes used by CoC for non-U.S. residents. Also 998, 999	3	
1842	Follow-Up ContactCity	Mixed case letters, no special characters, embedded spaces allowed, left justified, blank filled		20	
1844	Follow-Up ContactState	Upper case	See EDITS table STATE.DBF in Appendix B	2	

Item #	Item Name	Format	Allowable Values	Length	Note
1846	Follow-Up ContactPostal	Numbers or upper case letters. No special characters or embedded spaces allowed. Left justified, blank filled	5-digit or 9-digit U.S. ZIP codes; 6- character Canadian postal codes; valid postal codes from other countries, 888888888, 999999999	9	
1850	Unusual Follow-Up Method		0-9	1	
1860	Recurrence Date1st	MMDDCCYY	Valid dates, 00000000, 999999999	8	
1870	Recurrence Distant Sites				Retired
1871	Recurrence Distant Site 1		0-9	1	
1872	Recurrence Distant Site 2		0-9	1	
1873	Recurrence Distant Site 3		0-9	1	
1880	Recurrence Type1st	Right justified, zero filled	00, 04, 06, 10, 13-17, 20-22, 25-27, 30, 36, 40, 46, 51-60, 62, 70, 88, 99	2	
1890	Recurrence Type1stOth		00, 01, 06, 10, 11, 15-17, 20-22, 25-27, 30, 36, 40, 46, 70, 88, 99	2	
1900	Reserved 11			50	
1910	Cause of Death	4 digits (for ICD-7, 8, 9) or upper case letter followed by 3 digits (for ICD-10)	Valid ICD-7, ICD-8, ICD-9, and ICD-10 codes; also 0000, 7777, 7797	4	
1920	ICD Revision Number		0, 1, 7, 8, 9	1	
1930	Autopsy		0-2, 9	1	
1940	Place of Death	Right justified, zero filled	Reference SEER Manual	3	
1950	Reserved 12			50	
1960	Site (73-91) ICD-O-1	Four digits, first digit equals 1	Reference ICD-O-1 for valid entries	4	
1970	Morph (73-91) ICD-O-1		Reference ICD-O-1 for valid entries	6	
1971	Histology (73-91) ICD-O-1		Reference ICD-O-1 for valid entries	4	
1972	Behavior (73-91) ICD-O-1		Reference ICD-O-1 for valid entries	1	
1973	Grade (73-91) ICD-O-1		Reference ICD-O-1 for valid entries	1	
1980	ICD-O-2 Conversion Flag		0-6	1	
1981	Over-ride SS/NodesPos		1 or blank	1	
1982	Over-ride SS/TNM-N		1 or blank	1	
1983	Over-ride SS/TNM-M		1 or blank	1	
1984	Over-ride SS/DisMet1		1 or blank	1	
1985	Over-ride Acsn/Class/Seq		1 or blank	1	
1986	Over-ride HospSeq/DxConf		1 or blank	1	
1987	Over-ride COC-Site/Type		1 or blank	1	
1988	Over-ride HospSeq/Site		1 or blank	1	
1989	Over-ride Site/TNM-StgGrp		1 or blank	1	
1990	Over-ride Age/Site/Morph		1 or blank	1	
2000	Over-ride SeqNo/DxConf		1 or blank	1	
2010	Over-ride Site/Lat/SeqNo		1 or blank	1	
2020	Over-ride Surg/DxConf		1 or blank	1	

Item #	Item Name	Format	Allowable Values	Length	Note
2030	Over-ride Site/Type		1 or blank	1	
2040	Over-ride Histology		1-3 or blank	1	
2050	Over-ride Report Source		1 or blank	1	
2060	Over-ride Ill-define Site		1 or blank	1	
2070	Over-ride Leuk, Lymphoma		1 or blank	1	
2071	Over-ride Site/Behavior		1 or blank	1	
2072	Over-ride Site/EOD/DX Dt		1 or blank	1	
2073	Over-ride Site/Lat/EOD		1 or blank	1	
2074	Over-ride Site/Lat/Morph		1 or blank	1	
2080	Reserved 13				Retired
2081	CRC CHECKSUM		Calculated or blank	10	
2082	Reserved 24			16	
2090	Date Case Completed	MMDDCCYY		8	
2100	Date Case Last Changed	MMDDCCYY		8	
2110	Date Case Report Exported	MMDDCCYY		8	
2111	Date Case Report Received	MMDDCCYY		8	
2112	Date Case Report Loaded	MMDDCCYY		8	
2112	Date Tumor Record Availbl	MMDDCCYY		8	
2113	Future Use Timeliness 1			0	Retired
2115	Future Use Timeliness 2				Retired
2115	ICD-O-3 Conversion Flag		Blank, 0, 1, 3	1	reentea
2120	SEER Coding SysCurrent		0-6	1	
2120	SEER Coding Sys-Original		0-6	1	
2130	COC Coding Sys-Current	Right justified, zero filled	00-08, 99	2	
2140	COC Coding Sys-Original	Right justified, zero filled	00-08, 99	2	
2160	Subsq Report for Primary		00 00, 77	2	Retired
2160	Reserved 20				Retired
2170	Vendor Name	Embedded spaces allowed		10	reenieu
2180	SEER Type of Follow-Up		1-4	1	
2190	SEER Record Number	Right justified, zero filled	01-99	2	
2200	Diagnostic Proc 73-87			2	
2210	Reserved 14			_	Retired
2220	State/Requestor Items			500	itettieu
2230	NameLast	Mixed case, no embedded spaces, left justified, blank filled. Embedded hyphen allowed, but no other special characters		25	
2240	NameFirst	Mixed case, no embedded spaces, no special characters, left justified, blank filled		14	
2250	NameMiddle	Mixed case, no embedded spaces, no special characters, left justified, blank filled		14	
2260	NamePrefix	Mixed case, no special characters		3	
2270	NameSuffix	Mixed case, no special characters		3	
2280	NameAlias	Left justified, blank filled		15	
2290	NameSpouse/Parent	No standard		50	
2300	Medical Record Number	Leading spaces, right justified		11	
2310	Military Record No Suffix	Right justified, zero filled	01-20, 30-69, 98, 99	2	
2320	Social Security Number	9 digits, no dashes	Any 9-digit number except 000000000	9	

Item #	Item Name	Format	Allowable Values	Length	Note
2330	Addr at DXNo & Street	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2335	Addr at DXSupplementl	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2350	Addr CurrentNo & Street	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2352	Latitude	Right justified	See Data Dictionary	10	
2354	Longitude	Right justified	See Data Dictionary	11	
2355	Addr CurrentSupplementl	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2360	Telephone	10-digit number	Any 10-digit number	10	
2370	DC State				Retired
2371	Reserved 21				Retired
2380	DC State File Number		Any characters or blank	6	
2390	NameMaiden	Mixed case, no embedded spaces, left justified, blank filled, embedded hyphen allowed, no other special characters		15	
2392	Follow-Up ContactNo&St	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2393	Follow-Up ContactSuppl	Mixed case, embedded spaces, punctuation limited to periods, slashes, hyphens, and pound signs. Left justified, space filled		40	
2394	Follow-Up ContactName	Mixed case, embedded spaces, no special characters, left justified, blank fill		30	
2400	Reserved 16				Retired
2410	Institution Referred From	Right justified and zero filled	10-digit number	10	
2420	Institution Referred To	Right justified and zero filled	10-digit number	10	
2430	Last Follow-Up Hospital	Right justified and zero filled	10-digit number	10	
2440	Following Registry	Right justified and zero filled	10-digit number	10	
2450	Reserved 17				Retired
2460	PhysicianManaging	Left justified		8	
2470	PhysicianFollow-Up	Left justified		8	
2480	PhysicianPrimary Surg	Left justified		8	
2490	Physician 3	Left justified		8	
2500	Physician 4	Left justified		8	
2520	TextDX ProcPE	Free text	Neither carriage return nor line feed characters allowed	200	
2530	TextDX ProcX-ray/Scan	Free text	Neither carriage return nor line feed characters allowed	250	
2540	TextDX ProcScopes	Free text	Neither carriage return nor line feed characters allowed	250	
2550	TextDX ProcLab Tests	Free text	Neither carriage return nor line feed characters allowed	250	
2560	TextDX ProcOp	Free text	Neither carriage return nor line feed characters allowed	250	
2570	TextDX ProcPath	Free text	Neither carriage return nor line feed characters allowed	250	

Item #	Item Name	Format	Allowable Values	Length	Note
2580	TextPrimary Site Title	Free text	Neither carriage return nor line feed characters allowed	40	
2590	TextHistology Title	Free text	Neither carriage return nor line feed characters allowed	40	
2600	TextStaging	Free text	Neither carriage return nor line feed characters allowed	300	
2610	RX TextSurgery	Free text	Neither carriage return nor line feed characters allowed	150	
2620	RX TextRadiation (Beam)	Free text	Neither carriage return nor line feed characters allowed	150	
2630	RX TextRadiation Other	Free text	Neither carriage return nor line feed characters allowed	150	
2640	RX TextChemo	Free text	Neither carriage return nor line feed characters allowed	200	
2650	RX TextHormone	Free text	Neither carriage return nor line feed characters allowed	200	
2660	RX TextBRM	Free text	Neither carriage return nor line feed characters allowed	100	
2670	RX TextOther	Free text	Neither carriage return nor line feed characters allowed	100	
2680	TextRemarks	Free text	Neither carriage return nor line feed characters allowed	350	
2690	Place of Diagnosis	Free text	Neither carriage return nor line feed characters allowed	50	
2700	Reserved 19			770	
2800	CS Tumor Size	Right justified, zero filled	000-999 (site specific)	3	
2810	CS Extension	Right justified, zero filled	00-99 (site specific)	2	
2820	CS Tumor Size/Ext Eval		0-9 (site specific) and N	1	Revised
2830	CS Lymph Nodes	Right justified, zero filled	00-99 (site specific)	2	
2840	CS Reg Node Eval		0-9(site specific) and N	1	Revised
2850	CS Mets at DX	Right justified, zero filled	00-99 (site specific)	2	
2860	CS Mets Eval		0-9 (site specific) and N	1	Revised
2880	CS Site-Specific Factor 1	Right justified, zero filled	000-999 (site specific)	3	
2890	CS Site-Specific Factor 2	Right justified, zero filled	000-999 (site specific)	3	
2900	CS Site-Specific Factor 3	Right justified, zero filled	000-999 (site specific)	3	
2910	CS Site-Specific Factor 4	Right justified, zero filled	000-999 (site specific)	3	
2920	CS Site-Specific Factor 5	Right justified, zero filled	000-999 (site specific)	3	
2930	CS Site-Specific Factor 6	Right justified, zero filled	000-999 (site specific)	3	
2935	CS Version 1st	6-digit number	Any 6-digit number	6	
2936	CS Version Latest	6-digit number	Any 6-digit number	6	
2940	Derived AJCC T		Derived from Collaborative Stage fields	2	
2950	Derived AJCC T Descriptor		c, p, a, y, N, and blank (derived from Collaborative Stage fields)	1	Revised
2960	Derived AJCC N		Derived from Collaborative Stage fields	2	
2970	Derived AJCC N Descriptor		c, p, a, y, N, and blank (derived from Collaborative Stage fields)	1	Revised

Item #	Item Name	Format	Allowable Values	Length	Note
2980	Derived AJCC M		Derived from Collaborative Stage fields	2	
2990	Derived AJCC M Descriptor		c, p, a, y, N, and blank (derived from Collaborative Stage fields)	1	Revised
3000	Derived AJCC Stage Group		Derived from Collaborative Stage fields	2	
3010	Derived SS1977		0-5, 7, 8, 9 (derived from Collaborative Stage fields)	1	
3020	Derived SS2000		0-5, 7, 8, 9 (derived from Collaborative Stage fields)	1	
3030	Derived AJCCFlag		1, 2, blank	1	
3040	Derived SS1977Flag		1, 2, blank	1	
3050	Derived SS2000Flag		1, 2, blank	1	
3100	Archive FIN	Right justified, zero filled	10-digit number	10	
3110	Comorbid/Complication 1	Left justified, zero filled	00000, 00100-13980, 24000-99990, E8700- E8799, E9300-E9499, V0720-V0739, V1000- V1590, V2220-V2310, V2540, V4400-V4589, V5041-V5049, blank	5	Revised
3120	Comorbid/Complication 2	Left justified, zero filled	00100-13980, 24000-99990, E8700-E8799, E9300- E9499, V0720-V0739, V1000-V1590, V2220- V2310, V2540, V4400- V4589, V5041-V5049, blank	5	Revised
3130	Comorbid/Complication 3	Left justified, zero filled	00100-13980, 24000-99990, E8700-E8799, E9300- E9499, V0720-V0739, V1000-V1590, V2220- V2310, V2540, V4400- V4589, V5041-V5049, blank	5	Revised
3140	Comorbid/Complication 4	Left justified, zero filled	00100-13980, 24000-99990, E8700-E8799, E9300- E9499, V0720-V0739, V1000-V1590, V2220- V2310, V2540, V4400- V4589, V5041-V5049, blank	5	Revised
3150	Comorbid/Complication 5	Left justified, zero filled	00100-13980, 24000-99990, E8700-E8799, E9300- E9499, V0720-V0739, V1000-V1590, V2220- V2310, V2540, V4400- V4589, V5041-V5049, blank	5	Revised
3160	Comorbid/Complication 6	Left justified, zero filled	00100-13980, 24000-99990, E8700-E8799, E9300- E9499, V0720-V0739, V1000-V1590, V2220- V2310, V2540, V4400- V4589, V5041-V5049, blank	5	Revised

Item #	Item Name	Format	Allowable Values	Length	Note
3170	RX DateMost Defin Surg	MMDDCCYY	Valid dates, 00000000, 999999999	8	
3180	RX DateSurgical Disch	MMDDCCYY	Valid dates, 00000000, 999999999	8	
3190	Readm Same Hosp 30 Days		0-3, 9	1	
3200	RadBoost RX Modality	Right justified, zero filled	00, 20-32, 40-43, 50-55, 60- 62, 98, 99	2	
3210	RadBoost Dose cGy	Right justified, zero filled	00000-99999	5	
3220	RX DateRadiation Ended	MMDDCCYY	Valid dates, 00000000, 888888888, 999999999	8	
3230	RX DateSystemic	MMDDCCYY	Valid dates, 00000000, 888888888, 999999999	8	
3250	RX Summ Transplnt/Endocr	Right justified, zero filled	00, 10-12, 20, 30, 40, 82, 85-88, 99	2	
3260	Pain Assessment				Retired
3270	RX SummPalliative Proc		0-7, 9	1	
3280	RX HospPalliative Proc		0-7, 9	1	
3300	RuralUrban Continuum 1993	Right justified, zero filled	00-09, 98, 99, (calculated); blank	2	
3310	RuralUrban Continuum 2000	Right justified, zero filled	00-09, 98, 99, (calculated); blank	2	

# CHAPTER XI

# DATA DICTIONARY

In this chapter, data items are presented in alphabetical order by item names. For each item, a general description, specific codes and meanings are given. For many items, the document provides a brief rationale for collecting the data item or for using the codes listed. The at-a-glance header for each data item has alternate name(s), item number, length, source of standard, and column numbers (for a discussion of NAACCR's standard naming conventions, see Chapter I).

Differences from Version 10.1 are marked "Revised" or "New" following the item name and item number. Black vertical lines in the outside margins highlight changes. Some changes are summarized in Appendix F.

Alternate names by which the same item is called under NAACCR's naming convention are listed in Appendix D.

The Source of Standard implies the reference for detailed coding instructions for many of the data items. A list of references can be found in Chapter VII. Chapter V, Table 3 provides a list of reference manuals for Version 10.2 (and prior versions).

Date fields are recorded in the month, day, year format (MMDDCCYY), with 99 for unknown day or month and 9999 for unknown year. For example:

- 00000000 No date
- 99999999 Unknown date
- 01992003 Example of date when the month and year are known but the day is unknown
- 99992003 Example of date when the year is known but the month and day are unknown.

#### ABSTRACTED BY

Alternate Name	Item #	Length	Source of Standard	Column #
	570	3	COC	413-415

#### Description

An alphanumeric code assigned by the reporting facility that identifies the individual abstracting the case.

#### **ACCESSION NUMBER--HOSP**

Alternate Name	Item #	Length	Source of Standard	Column #
Accession Number (COC)	550	9	COC	402-410

#### Description

Unique number assigned by the hospital registry to identify the patient. The first 4 digits identify the year (in the format CCYY) the patient was first seen at that institution for the diagnosis or treatment of cancer. The first 4 digits must be greater than or equal to 1944.

The last five numbers are the numeric order in which the registry entered the case into the database. Within a registry, all primaries for an individual must have the same accession number.

#### Rationale

Hospitals use this number to identify cases. If the central registry preserves this number, they can refer to it when communicating with the hospital. It also provides a way to link computerized follow-up reports from hospitals into the central database.

#### ADDR AT DX--CITY

Alternate Name	Item #	Length	Source of Standard	Column #
City or Town (pre-96 COC)	70	20	COC	52-71
City/Town at Diagnosis (COC)				

#### Description

Name of the city in which the patient resides at the time the reportable tumor was diagnosed. If the patient resides in a rural area, record the name of the city used in their mailing address. If the patient has multiple primaries, the city of residence may be different for each primary.

#### Codes

UNKNOWN (in addition to valid City)

#### ADDR AT DX--NO & STREET

Alternate Name	Item #	Length	Source of Standard	Column #
Patient Address (Number and Street) at	2330	40	COC	2108-2147
Diagnosis (COC)				
Number and Street (pre-96 COC)				

### Description

The number and street address or the rural mailing address of the patient's residence at the time the reportable tumor was diagnosed. If the patient has multiple tumors, address at diagnosis may be different for each tumor. Additional address information such as facility, nursing home, or name of apartment complex should be entered in Addr at DX--Supplementl [2335]. Do not update this item if patient moves after diagnosis.

U.S. addresses should conform to the U.S. Postal Service (USPS) Postal Addressing Standards. These standards are referenced in USPS Publication 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/Pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current Canadian Postal Address standards may be found at the following Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

#### Rationale

Addresses that are formatted to conform to USPS Postal Addressing Standards can be more properly geocoded by geographic information systems (GIS) software and vendors to the correct census tract, which is required by NPCR and SEER registries. The USPS Standards also address a number of issues that are problematic in producing precise addresses, including the use of punctuation, abbreviations, and proper placement of address elements, such as street direction, apartment and suite numbers, and unusual addressing situations. Spanish-language addresses also are covered by the USPS Standard.

#### **Coding Instructions (summary of USPS guidelines)**

The address should be fully spelled out with standardized use of abbreviations and punctuation per USPS postal addressing standards (USPS *Postal Addressing Standards*, Pub. 28, November 2000). Upper case recommended. Mixed case allowed.

Abbreviations should be limited to those recognized by USPS standard abbreviations, these include but are not limited to (A complete list of recognized street abbreviations is provided in Appendix C of USPS Pub. 28):

APT	apartment	Ν	north
BLDG	building	NE	northeast
FL	floor	NW	northwest
STE	suite	S	south
UNIT	unit	SE	southeast
RM	room	SW	southwest
DEPT	department	E	east

Punctuation marks should be avoided, except when punctuation is necessary to convey the meaning. Punctuation normally is limited to periods when the period carries meaning (e.g., 39.2 RD), slashes for fractional addresses (e.g., 101 1/2 MAIN ST), and hyphens when the hyphen carries meaning (e.g., 289-01 MONTGOMERY AVE). Use of the pound sign (#) to designate address units should be avoided whenever possible. The preferred notation is as follows: 102 MAIN ST APT 101. If a pound sign is used, there must be a space between the pound sign and the secondary number (e.g., 425 FLOWER BLVD # 72).

#### Codes (in addition to valid street address)

UNKNOWN Patient's address is unknown

ADDR AT DXPOSTAL CODE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Postal Code at Diagnosis (COC)	100	9	COC	74-82
Zip Code (pre-COC)				

#### Description

Postal code for the address of the patient's residence at the time the reportable tumor is diagnosed. If the patient has multiple tumors, the postal code may be different for each tumor.

For U.S. residents, use either the 5-digit or the extended 9-digit ZIP code. Blanks follow the 5-digit code. If the 4-digit extension is not collected, then the corresponding characters of an unknown value may be blank.

For Canadian residents, use the 6-character alphanumeric postal code. Blanks follow the 6-character code.

When available, enter the postal code for other countries.

#### Codes (in addition to US and Canadian postal codes)

888888888 Resident of country other than the United States, U.S. possessions or territories, or Canada and the postal code is unknown.

999999999 Resident of the United States (including its possessions, etc.) or Canada and the postal code is unknown

ADDR AT DXSTATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
State (pre-96 COC)	80	2	COC	72-73
State at Diagnosis (COC)				

#### Description

USPS abbreviation for the state, territory, commonwealth, U.S. possession, or Canadian province/territory in which the patient resides at the time the reportable tumor is diagnosed. If the patient has multiple primaries, the state of residence may be different for each tumor.

#### Codes (in addition to USPS abbreviations)

- XX Resident of country other than the United States (including its territories, commonwealths, or possessions) or Canada, and country is known
- YY Resident of country other than the United States (including its territories, commonwealths, or possessions) or Canada, and country is unknown
- ZZ Resident of the United States, NOS (including its territories, commonwealths, or possessions); Canada, NOS; residence unknown

#### ADDR AT DX--SUPPLEMENTL

Alternate Name	Item #	Length	Source of Standard	Column #
Patient Address (Number and Street) at	2335	40	COC	2148-2187
DiagnosisSupplemental (COC)				

#### Description

This data item provides the ability to store additional address information such as the name of a place or facility, a nursing home, or the name of an apartment complex. If the patient has multiple tumors, address at diagnosis may be different for each tumor. Additional address information such as number and street should be entered in Addr at DX--No & Street [2330].

U.S. addresses should conform to the USPS Postal Addressing Standards. These standards are referenced in USPS Pub. 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/Pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current *Canada Postal Guide* may be found at the following Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

#### Rationale

Sometimes the registry receives the name of a facility instead of a proper street address containing the street number, name, direction, and other elements necessary to locate an address on a street file for the purpose of geocoding. By having a second street address field to hold address information, the registry can look up and store the street address and not lose the facility name due to a shortage of space. The presence of a second street address field to hold address information also aids in follow-up.

Addresses that are formatted to conform to USPS Postal Addressing Standards can be more properly geocoded by GIS software and vendors to the correct census tract, which is required by NPCR and SEER registries. The USPS Standards also address a number of issues that are problematic in producing precise addresses, including the use of punctuation, abbreviations, and proper placement of address elements, such as street direction, apartment and suite numbers, and unusual addressing situations. Spanish-language addresses also are covered by the USPS Standard.

#### **Coding Instructions (summary of USPS guidelines)**

The address should be fully spelled out with standardized use of abbreviations and punctuation per USPS postal addressing standards (USPS *Postal Addressing Standards*, Pub. 28, November 2000). Upper case recommended. Mixed case allowed.

Abbreviations should be limited to those recognized by the Postal Service standard abbreviations, these include but are not limited to (a complete list of recognized abbreviations is provided in Appendix C of USPS Pub. 28.):

APT	apartment	Ν	north
BLDG	building	NE	northeast
FL	floor	NW	northwest
STE	suite	S	south
UNIT	unit	SE	southeast
RM	room	SW	southwest
DEPT	department	E	east

#### ADDR CURRENT--CITY

Alternate Name	Item #	Length	Source of Standard	Column #
City/TownCurrent (COC)	1810	20	COC	1307-1326

#### Description

Name of city of the patient's current usual residence. If the patient has multiple tumors, the current city of residence should be the same for all tumors.

#### Rationale

"Current address" can be used to measure the regional "cancer burden" (cost, medical care needs), especially in major retirement regions. Sometimes central registries carry out follow-up by contacting the patients by a letter or telephone calls to ascertain their vital status. The most current reported address and telephone number are needed. This information is also useful for conducting interview studies.

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

# ADDR CURRENT--NO & STREETAlternate NameItem #LengthSource of StandardPatient Address (Number and Street)--<br/>Current (COC)235040COC

#### Description

The number and street address or the rural mailing address of the patient's current usual residence. This can be used to generate a follow-up inquiry, and must correspond to other fields in the current address. If the patient has multiple tumors, the current address should be the same. Additional address information such as facility, nursing home, or name of apartment complex should be entered in item Addr Current-Supplementl [2335].

U.S. addresses should conform to the USPS Postal Addressing Standards. These standards are referenced in USPS Pub. 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/Pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current Canadian Postal Address standards may be found at the following Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

#### Rationale

"Current address" can be used to measure the regional "cancer burden" (cost, medical care needs), especially in major retirement regions. Sometimes central registries carry out follow-up by contacting the patients via letter or telephone calls to ascertain their vital status. The most current reported address and telephone number are needed. This information also is useful for conducting interview studies.

Addresses that are formatted to conform to USPS Postal Addressing Standards can be more properly geocoded by GIS software and vendors to the correct census tract. The USPS Standards also address a number of issues that are problematic in producing precise addresses, including the use of punctuation, abbreviations, and proper placement of address elements, such as street direction, apartment and suite numbers, and unusual addressing situations. Spanish-language addresses also are covered by the USPS Standard.

#### **Coding Instructions (summary of USPS guidelines)**

The address should be fully spelled out with standardized use of abbreviations and punctuation per USPS postal addressing standards (USPS *Postal Addressing Standards*, Pub. 28, November 2000). Upper case recommended. Mixed case allowed.

Abbreviations should be limited to those recognized by USPS standard abbreviations, these include but are not limited to (a complete list of recognized street abbreviations is provided in Appendix C of USPS Pub. 28.):

APT	apartment	Ν	north
BLDG	building	NE	northeast
FL	floor	NW	northwest
STE	suite	S	south
UNIT	unit	SE	southeast
RM	room	SW	southwest
DEPT	department	Е	east

Column #

2188-2227

Punctuation marks should be avoided, except when punctuation is necessary to convey the meaning. Punctuation normally is limited to periods when the period carries meaning (e.g., 39.2 RD), slashes for fractional addresses (e.g., 101 1/2 MAIN ST), and hyphens when the hyphen carries meaning (e.g., 289-01 MONTGOMERY AVE). Use of the pound sign (#) to designate address units should be avoided whenever possible. The preferred notation is as follows: 102 MAIN ST APT 101. If a pound sign is used, there must be a space between the pound sign and the secondary number (e.g., 425 FLOWER BLVD # 72).

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

ADDR CURRENTPOSTAL CODE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Postal CodeCurrent (COC)	1830	9	COC	1329-1337

#### Description

Postal code for the address of the patient's current usual residence. If the patient has multiple tumors, the postal codes should be the same. For U.S. residents, use either the 5-digit or the extended 9-digit ZIP code. Blanks follow the 5-digit code. For Canadian residents, use the 6-character alphanumeric postal code. Blanks follow the 6-character code. When available, enter postal code for other countries.

#### Rationale

"Current address" can be used to measure the regional "cancer burden" (cost, medical care needs), especially in major retirement regions. Sometimes central registries carry out follow-up by contacting the patients by a letter or telephone calls to ascertain their vital status. The most current reported address and telephone number are needed. This information also is useful for conducting interview studies.

#### Codes (in addition to U.S., Canadian, and Foreign postal codes)

- 888888888 Resident of country other than the United States (including its possessions, etc.) or Canada, and postal code unknown
- 999999999 Resident of the United States (including its possessions, etc.) or Canada, and postal code unknown

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

ADDR CURRENTSTATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
StateCurrent (COC)	1820	2	COC	1327-1328

#### Description

USPS abbreviation for the state (including U.S. territories, commonwealths, or possessions) or Canadian province/territory of the patient's current usual residence. If the patient has multiple tumors, the current state of residence should be the same for all tumors.

#### Rationale

"Current address" can be used to measure the regional "cancer burden" (cost, medical care needs), especially in major retirement regions. Sometimes central registries carry out follow-up by contacting the patients via letter or telephone calls to ascertain vital status. The most current reported address and telephone number are needed. This information also is useful for conducting interview studies.

#### Codes (in addition to the U.S. and Canadian postal service abbreviations)

- Resident of country other than the United States (including its territories, commonwealths, or XX possessions) or Canada, and country is known
- YY Resident of country other than the United States (including its territories, commonwealths, or possessions) or Canada, and country is unknown
- ZZ Resident of the United States, NOS (including its territories, commonwealths, or possessions); Canada, NOS; residence unknown

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

#### ADDR CURRENT--SUPPLEMENTL

Alternate Name	Item #	Length	Source of Standard	Column #
Patient Address (Number and Street)	2355	40	COC	2228-2267
CurrentSupplemental (COC)				

#### Description

This data item provides the ability to store additional address information such as the name of a place or facility, a nursing home, or the name of an apartment complex. This can be used to generate a follow-up inquiry, and must correspond to other fields in the current address. If the patient has multiple tumors, the current address should be the same.

U.S. addresses should conform to the USPS Postal Addressing Standards. These standards are referenced in USPS Pub. 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current *Canada Postal Guide* may be found at the follow Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

#### Rationale

Sometimes the registry receives the name of a facility instead of a proper street address containing the street number, name, direction, and other elements necessary to locate an address on a street file for the purpose of geocoding. By having a second street address field to hold address information, the registry can look up and store the street address and not lose the facility name due to a shortage of space. The presence of a second street address field to hold address information also aids in follow-up.

"Current address" can be used to measure the regional "cancer burden" (cost, medical care needs), especially in major retirement regions. Sometimes central registries carry out follow-up by contacting the patients via letter or telephone calls to ascertain their vital status. The most current reported address and telephone number are needed. This information is also useful for conducting interview studies.

#### **Coding Instructions (summary of USPS guidelines)**

The address should be fully spelled out with standardized use of abbreviations and punctuation per USPS postal addressing standards (USPS *Postal Addressing Standards*, Pub. 28, November 2000). Upper case recommended. Mixed case allowed.

Abbreviations should be limited to those recognized by the Postal Service standard abbreviations, these include but are not limited to (a complete list of recognized abbreviations is provided in Appendix C of USPS Pub. 28.):

APT	apartment	Ν	north
BLDG	building	NE	northeast
FL	floor	NW	northwest
STE	suite	S	south
UNIT	unit	SE	southeast
RM	room	$\mathbf{SW}$	southwest
DEPT	department	E	east

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

#### AGE AT DIAGNOSIS

Alternate Name	Item #	Length	Source of Standard	Column #
	230	3	SEER/COC	119-121

#### Description

Age of the patient at diagnosis in complete years. Different tumors for the same patient may have different values.

#### Codes

000	Less than 1 year old
001	1 year old, but less than 2 years
002	2 years old
	(show actual age in completed years)
101	101 years old
120	120 years old
999	Unknown age

#### ALCOHOL HISTORY

Alternate Name	Item #	Length	Source of Standard	Column #
	350	1	Varies	225-225

#### Description

NAACCR has not adopted standards for this item.

#### **ARCHIVE FIN**

Alternate Name	Item #	Length	Source of Standard	Column #
	3100	10	COC	392-401

#### Description

This field identifies the facility that originally accessioned the tumor.

#### Rationale

Each facility's facility identification number (FIN) is unique. It is essential for hospital registries to have the ability to distinguish cases originally accessioned by each registry of a merged unit. This enables the central registry to manage the receipt of historical data and to appropriately attribute these data.

Efforts are underway at the federal level to establish uniform national provider ID numbers. COC and NAACCCR committees will consider the adoption of any federal standards when they become available.

#### **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs in the range of 6020009-6953290 that were assigned by COC before January 1, 2001, the coded FIN will consist of three leading zeroes followed by the full 7-digit number.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001, enter FIN codes of this type as two zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10-digit codes.

AUTOPSY				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1930	1	NAACCR	1393-1393

#### Description

Code indicating whether or not an autopsy was performed.

#### Codes

- 0 Not applicable; patient alive
- 1 Autopsy performed
- 2 No autopsy performed
- 9 Patient expired, unknown if autopsy performed

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### BEHAVIOR (73-91) ICD-O-1

Alternate Name	Item #	Length	Source of Standard	Column #
	1972	1	SEER	1145-1145

#### Description

Area for retaining behavior portion (1 digit) of the ICD-O-1 or field trial morphology codes entered before a conversion to ICD-O-2. See grouped data item Morph (73-91) ICD-O-1 [1970] in Appendix E. The item name includes years 73-91. However, some states may have used the codes for cases before 1973. It is a subfield of the morphology code.

#### Codes

For tumors diagnosed before 1992, contains the ICD-O-1 or field trial 1-digit behavior code as originally coded, if available. Blank for tumors coded directly into ICD-O-2 (I.e., 1992 and later tumors).

#### BEHAVIOR (92-00) ICD-O-2

Alternate Name	Item #	Length	Source of Standard	Column #
	430	1	SEER/COC	300-300

#### Description

Code for the behavior of the tumor being reported using ICD-O-2. NAACCR adopted ICD-O-2 as the standard coding system for tumors diagnosed from January 1, 1992, through December 31, 2000. In addition, NAACCR recommended that cases diagnosed prior to 1992 be converted to ICD-O-2. See Behavior (73-91) ICD-O-1 [1972], for ICD-O-1 and field trial codes.

#### Codes

Valid codes are 0-3. See ICD-0-2,<sup>15</sup> page 22, for behavior codes and definitions.

#### **Clarification of Required Status**

This data item is required by all standard-setting organizations for tumors diagnosed from January 1, 1992, through December 31, 2000, and recommended for tumors diagnosed before 1992.

When the histologic type is coded according to the ICD-O-2, the histology code must be reported in Histology (92-00) ICD-O-2 [420], with behavior coded in Behavior (92-00) ICD-O-2 [430].

For information on required status for related data items for histologic type and behavior when coded according to ICD-O-3, see Histologic Type ICD-O-3 [522] and Behavior Code ICD-O-3 [523].

#### **BEHAVIOR CODE ICD-O-3**

Alternate Name	Item #	Length	Source of Standard	Column #
Behavior Code (COC)	523	1	SEER/COC	305-305

#### Description

Code for the behavior of the tumor being reported using ICD-O-3. NAACCR adopted ICD-O-3 as the standard coding system for tumors diagnosed beginning January 1, 2001, and later recommended that prior cases be converted from ICD-O-2. See Behavior (92-00) ICD-O-2 [430], for ICD-O-2 codes.

Juvenile astrocytoma is coded as borderline in ICD-O-3, North American registries report as 9421/3.

#### Codes

Valid codes are 0-3. See ICD-O-3,<sup>14</sup> page 66, for behavior codes and definitions.

#### **Clarification of Required Status**

Behavior is required by all standard-setting organizations for tumors diagnosed on or after January 1, 2001, and recommended (by conversion from ICD-O-2 codes) for tumors diagnosed before 2001.

When the histologic type is coded according to the ICD-O-3, the histology code must be reported in Histologic Type ICD-O-3 [522], with behavior coded in Behavior Code ICD-O-3 [523].

For information on required status for related data items for histologic type and behavior when coded according to ICD-O-2, see Histology (92-00) ICD-O-2 [420] and Behavior (92-00) ICD-O-2 [430].

#### **BIRTH DATE**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Birth (SEER/COC)	240	8	SEER/COC	122-129

#### Description

Date of birth of the patient. See page 83 for date format. A zero must precede single-digit months and days. If age at diagnosis and year of diagnosis are known, but year of birth is unknown, then year of birth should be calculated and so coded. Month and day would be coded as unknown (99). Estimate date of birth when information is not available. It is better to estimate than to code as an unknown value.

#### BIRTHPLACE

Alternate Name	Item #	Length	Source of Standard	Column #
Place of Birth (SEER/COC)	250	3	SEER/COC	130-132

#### Description

Code for place of birth of the patient. If a patient has multiple tumors, all records should contain the same code.

#### Rationale

Place of Birth is helpful for patient matching and can be used when reviewing race and ethnicity. In addition, adding birthplace data to race and ethnicity allows for a more specific definition of the population being reported. Careful descriptions of ancestry, birthplace, and immigration history of populations studied are needed to make the basis for classification into ethnic groups clear. Birthplace has been associated with variation in genetic, socioeconomic, cultural, and nutritional characteristics that affect patterns of disease. A better understanding of the differences within racial and ethnic categories also can help states develop effective, culturally sensitive public health prevention programs to decrease the prevalence of high-risk behaviors and increase the use of preventive services.

#### Code

See Appendix B (also Appendix B of the *SEER Program Code Manual*) for numeric and alphabetic lists of places and codes.

#### **CANCER STATUS**

Alternate Name	Item #	Length	Source of Standard	Column #
	1770	1	COC	1303-1303

#### Description

Records the cancer status for this primary as of the date entered in Date of Last Contact [1750]. If the patient has multiple primaries, the values may be different for each primary.

#### Rationale

Hospitals use this field to compute survival analysis (disease-free intervals). By maintaining this data item, central registries can assist hospital registries by sharing this information with other hospital registries that serve the same patients, if the state's privacy laws so permit.

#### Codes

- 1 No evidence of this cancer
- 2 Evidence of this cancer
- 9 Unknown, indeterminate whether this cancer is present, not stated in patient record

CAUSE OF DEATH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Underlying Cause of Death (SEER)	1910	4	SEER	1388-1391
Underlying Cause of Death (ICD Code)				
(pre-96 COC)				

Official cause of death as coded from the death certificate in valid ICD-7, ICD-8, ICD-9, and ICD-10 codes.

### Rationale

Cause of death is used for calculation of adjusted survival rates by the life table method. The adjustment corrects for deaths other than from the diagnosed cancer.

## Special codes in addition to ICD-7, ICD-8, ICD-9, and ICD-10 (refer to *SEER Program Code Manual* for additional instructions)

- 0000 Patient alive at last contact
- 7777 State death certificate not available
- 7797 State death certificate available but underlying cause of death is not coded

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### CENSUS COD SYS 1970/80/90

Alternate Name	Item #	Length	Source of Standard	Column #
Census Coding System (COC)	120	1	SEER	92-92
Coding System for Census Tract (pre-96				
SEER/COC)				

#### Description

Identified the set of Census Bureau census tract definitions (boundaries) that were used to code the census tract in Census Tract 1970/80/90 [110] for a specific record.

#### Rationale

Allows for changes in census tracts over time. The census tract definition used to code the case must be recorded so that data are correctly grouped and analyzed. If the coding system were not recorded, the census codes would have to be converted or recorded every time the census tracts were changed.

#### Codes

- 0 Not tracted
- 1 1970 Census Tract Definitions
- 2 1980 Census Tract Definitions
- 3 1990 Census Tract Definitions
- Blank Census Tract 1970/80/90 not coded

#### **Clarification of NPCR Required Status**

<u>Census-1990 data items:</u> Census Tract 1970/80/90 [110] Census Tr Cert 1970/80/90 [364] Census Tract Cod Sys--1970/80/90 [120] <u>Census-2000 data items:</u> Census Tract 2000 [130] Census Tr Certainty 2000 [365]

Information on census tract, census tract certainty, and census tract coding system is required. For tumors diagnosed in or after 2003, Census Tract 2000 [130] and Census Tr Certainty 2000 [365] ("Census-2000 data items") are required. For tumors diagnosed in or before 2002, the requirement can be met by collecting either the Census-1990 data items [110, 364, 120] or the Census-2000 data items, although the Census-2000 data items [130 and 365] are recommended for tumors diagnosed in 1998 through 2002.

#### **CENSUS TR CERT 1970/80/90**

Alternate Name	Item #	Length	Source of Standard	Column #
Census Tract Certainty	364	1	SEER	100-100

#### Description

Code indicating basis of assignment of census tract or block numbering area (BNA) for an individual record. Helpful in identifying cases tracted from incomplete information or P.O. Box. Most of the time, this information is provided by a geocoding vendor service. Alternatively, a central registry staff manually assigns the code. This item is not coded by the hospital. Codes are hierarchical, with lower numbers having priority.

#### Codes

- 1 Census tract/BNA based on complete and valid street address of residence
- 2 Census tract/BNA based on residence ZIP + 4
- 3 Census tract/BNA based on residence ZIP + 2
- 4 Census tract/BNA based on residence ZIP code only
- 5 Census tract/BNA based on ZIP code of P.O. Box
- 9 Unable to assign census tract or BNA based on available information
- Blank Not applicable (e.g., census tracting not attempted); Census Tract Certainty information for 1970/80/90 not coded

#### **Clarification of NPCR Required Status**

<u>Census-1990 data items:</u> Census Tract 1970/80/90 [110] Census Tr Cert 1970/80/90 [364] Census Tract Cod Sys--1970/80/90 [120] <u>Census-2000 data items:</u> Census Tract 2000 [130] Census Tr Certainty 2000 [365]

Information on census tract, census tract certainty, and census tract coding system is required. For tumors diagnosed in or after 2003, Census Tract 2000 [130] and Census Tr Certainty 2000 [365] ("Census-2000 data items") are required. For tumors diagnosed in or before 2002, the requirement can be met by collecting either the Census-1990 data items [110, 364, 120] or the Census-2000 data items, although the Census-2000 data items [130 and 365] are recommended for tumors diagnosed in 1998 through 2002.

#### **CENSUS TR CERTAINTY 2000**

Alternate Name	Item #	Length	Source of Standard	Column #
	365	1	NAACCR	101-101

#### Description

Code indicating basis of assignment of census tract for an individual record. Helpful in identifying cases tracted from incomplete information or P.O. Box. Most of the time, this information is provided by a geocoding vendor service. Alternatively, a central registry staff manually assigns the code. This item is not coded by the hospital. Codes are hierarchical, with lower numbers having priority.

#### Codes

1	Census tract based on complete and valid street address of residence
2	Census tract based on residence ZIP + 4
3	Census tract based on residence $ZIP + 2$
4	Census tract based on residence ZIP code only
5	Census tract based on ZIP code of P.O. Box
9	Unable to assign census tract or bloc numbering based on available information
Blank	Not applicable (e.g., census tracting not attempted); Census Tract Certainty information for 2000
	not coded
Clarifia	ation of NPCR Required Status
Clarine	ation of NFCK Required Status
Conque	1000 data itams: Cansus 2000 data itams:

Census-1990 data items: Census Tract 1970/80/90 [110] Census Tr Cert 1970/80/90 [364] Census Tract Cod Sys--1970/80/90 [120] Census-2000 data items: Census Tract 2000 [130] Census Tr Certainty 2000 [365]

Information on census tract, census tract certainty, and census tract coding system is required. For tumors diagnosed in or after 2003, Census Tract 2000 [130] and Census Tr Certainty 2000 [365] ("Census-2000 data items") are required. For tumors diagnosed in or before 2002, the requirement can be met by collecting either the Census-1990 data items [110, 364, 120] or the Census-2000 data items, although the Census-2000 data items [130 and 365] are recommended for tumors diagnosed in 1998 through 2002.

#### **CENSUS TRACT 1970/80/90**

Alternate Name	Item #	Length	Source of Standard	Column #
Census Tract/Block Numbering Area	110	6	SEER	86-91
(BNA) (SEER)				
Census Tract				

#### Description

Code for the census tract or BNA of the patient's residence at the time of diagnosis. SEER used this field for tumors reported before 1998. If the patient has more than one tumor, the codes may be different for each tumor.

Codes are those used by the U.S. Census Bureau. Census Bureau codes for BNA also are entered in this field.

Both census tracts and BNAs have a 4-digit basic number and also may have a 2-digit suffix. Census tract numbers range from 0001.00 to 9499.99. BNA numbers range from 9501.00 to 9989.99. See the Census Bureau's "Area Classifications" 35 for further details.

#### Rationale

Allows central registries to calculate incidence rates for geographical areas having population estimates. The Census Bureau provides population data for census tracts. Those rates can be used for general surveillance or special geographical and socioeconomic analysis.

#### Codes

Census Tract Codes BNA Codes	000100-949999 950100-998999
000000	Area not census-tracted
999999	Area census-tracted, but census tract is not available
Blank	Census Tract 1970/80/90 not coded

#### **Clarification of NPCR Required Status**

Information on census tract, census tract certainty, and census tract coding system is required. Tumors diagnosed in 2003 or later, must be coded to the 2000 census definitions and recorded in Census Tract 2000 [130] and Census Tr Certainty 2000 [365]. Tumors diagnosed in 2002, or before must be coded to the 2000 census tract definitions OR to 1990 definitions OR to both the 2000 and 1990 census definitions. Census tract, census tract certainty and census tract coding system should be recorded in the year appropriate data item fields. For tumors diagnosed between January 1, 1998 and December 31, 2002 (inclusive) use of the 2000 census tract definitions is recommended.

#### CENSUS TRACT 2000

Alternate Name	Item #	Length	Source of Standard	Column #
Census TractAlternate (pre-2003)	130	6	NAACCR	93-98

#### Description

This field is provided for coding census tract of patient's residence at time of diagnosis. See Census Tract 1970/80/90 [110]. Codes are those used by the U.S. Census Bureau for the Year 2000 Census. Census tract codes have a 4-digit basic number and also may have a 2-digit suffix. Census tract numbers range from 0001.01 to 9999.98. See the Census Bureau's "Area Classifications" at the following Web Site: http://www.census.gov/prod/cen2000/doc/sf1.pdf for further details.

#### Rationale

Census tract codes allow central registries to calculate incidence rates for geographical areas having population estimates. This field allows a central registry to add Year 2000 Census tracts to tumors diagnosed in previous years, without losing the codes in data item 110.

The Census Bureau provides population data for census tracts. Those rates can be used for general surveillance or special geographical and socioeconomic analysis.

Because census tracts for particular cases can change between censuses, the central registry may wish to assign an alternate census tract code to its cases. For example, a registry may code its 1985 cases using both the 1980 and 1990 census tract boundaries. The central registry can use this information for different comparisons.

#### Codes

Census Tract Codes

000000Area not census tracted999999Area census-tracted, but census tract is not availableBlankCensus Tract 2000 not coded		
	999999	Area census-tracted, but census tract is not available

000101-999998

#### **Clarification of NPCR Required Status**

Information on census tract, census tract certainty, and census tract coding system is required. Tumors diagnosed in 2003 or later, must be coded to the 2000 census definitions and recorded in Census Tract 2000 [130] and Census Tr Certainty 2000 [365]. Tumors diagnosed in 2002, or before must be coded to the 2000 census tract definitions OR to 1990 definitions OR to both the 2000 and 1990 census tract definitions. Census tract, census tract certainty and census tract coding system should be recorded in the year appropriate data item fields. For tumors diagnosed between January 1, 1998 and December 31, 2002 (inclusive) use of the 2000 cases tract definitions is recommended.

#### CENSUS TRACT BLOCK GROUP

Alternate Name	Item #	Length	Source of Standard	Column #
	362	1	Census	99-99

#### Description

NAACCR has not adopted standards for this item.

#### CENSUS TRACT COD SYS--ALT

				neenica
Alternate Name	Item #	Length	Source of Standard	Column #
	140			

#### Description

This data item was retired for Version 10 because Census Tract 2000 [130] is expected to contain only Census 2000 codes.

## CHEMOTHERAPY FIELD 1 Retired Alternate Name Item # Length Source of Standard Column # 1600</

#### Description

This field has been listed as in development since 1996. The NAACCR UDSC approved to retire this data item in Version 10.1.

#### CHEMOTHERAPY FIELD 2

CHEMOTHERAFT FIELD 2				Kettreu
Alternate Name	Item #	Length	Source of Standard	Column #
	1610			

#### Description

This field has been listed as in development since 1996. The NAACCR UDSC approved to retire this data item in Version 10.1.

<b>CHEMOTHERAPY FIELD 3</b>				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	1620			

#### Description

This field has been listed as in development since 1996. The NAACCR UDSC approved to retire this data item in Version 10.1.

<b>CHEMOTHERAPY FIELD 4</b>				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	1630			

#### Description

This field has been listed as in development since 1996. The NAACCR UDSC approved to retire this data item in Version 10.1.

Retired

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#### CLASS OF CASE

Alternate Name	Item #	Length	Source of Standard	Column #
	610	1	COC	440-440

#### Description

For a hospital registry, divides cases into two groups: analytic cases are those included in reports on patient treatment and outcomes; nonanalytic cases are those not included in such reports. Class of Case codes 0-2 identify cases that are analytic (i.e., cases that were first diagnosed and/or received all or part of their first course of treatment or had treatment planned at the reporting hospital). Class of Case codes 3-5, 7, 8, and 9 identify cases that are considered nonanalytic (i.e., were first diagnosed and received all of their first course of treatment at a facility other than the reporting institution, or were diagnosed at autopsy or by death certificate only). Class of Case 6 identifies cases that were first diagnosed and received their entire first course of treatment in the same staff physician's office. These cases were considered analytic for diagnosis dates January 1, 1998, through December 31, 1999. For diagnosis dates on or after January 1, 2000, these cases are considered nonanalytic.

Class of Case can be used in conjunction with Type of Reporting Source [500]. Type of Reporting Source is designed to document the source of documents used to abstract the cancer being reported.

#### Codes

- 0 Diagnosis at the reporting facility and all of the first course of treatment was performed elsewhere or the decision not to treat was made at another facility.
- 1 Diagnosis at the reporting facility, and all or part of the first course of treatment was performed at the reporting facility.
- 2 Diagnosis elsewhere, and all or part of the first course of treatment was performed at the reporting facility.
- 3 Diagnosis and all of the first course of treatment was performed elsewhere. Presents at your facility with recurrence or persistent disease.
- 4 Diagnosis and/or first course of treatment were performed at the reporting facility prior to the reference date of the registry.
- 5 Diagnosed at autopsy
- 6 Diagnosis and all of the first course of treatment were completed by the same staff physician in an office setting. "Staff physician" is any medical staff with admitting privileges at the reporting facility.
- 7 Pathology report only. Patient does not enter the reporting facility at any time for diagnosis or treatment. This category excludes tumors diagnosed at autopsy.
- 8 Diagnosis was established by death certificate only. Used by central registries only.
- 9 Unknown. Sufficient detail for determining Class of Case is not stated in patient record. Used by central registries only.

#### COC CODING SYS--CURRENT

Alternate Name	Item #	Length	Source of Standard	Column #
Commission on Cancer Coding System	2140	2	COC	1200-1201
Current (COC)				

#### Description

Code the ACoS COC coding system currently used in the record. COC codes may be converted from an earlier version.

#### Codes

- 00 No COC coding system used
- Pre-1988 (Cancer Program Manual Supplement) 01
- 02 1988 Data Acquisition Manual
- 1989 Data Acquisition Manual Revisions 03
- 1990 Data Acquisition Manual Revisions 04
- 1994 Data Acquisition Manual (Interim/Revised) 05
- ROADS (effective with cases diagnosed 1996-1997) 06
- 07 ROADS and 1998 Supplement (effective with cases diagnosed 1998-2002)
- 08 FORDS 2003/2004 (effective with cases diagnosed 2003 and forward)
- 99 Unknown coding system

#### COC CODING SYS--ORIGINAL

COC CODING SYSORIGINAL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2150	2	COC	1202-1203

#### Description

Code for the ACoS COC coding system originally used to code the record.

#### Codes

- 00 No COC coding system used
- Pre-1988 (Cancer Program Manual Supplement) 01
- 1988 Data Acquisition Manual 02
- 03 1989 Data Acquisition Manual Revisions
- 04 1990 Data Acquisition Manual Revisions
- 05 1994 Data Acquisition Manual (Interim/Revised)
- ROADS (effective with cases diagnosed 1996-1997) 06
- ROADS and 1998 Supplement (effective with cases diagnosed 1998-2002) 07
- 08 FORDS 2003/2004 (effective with cases diagnosed 2003 and forward)
- 99 Unknown coding system

#### CODING SYSTEM FOR EOD

Alternate Name	Item #	Length	Source of Standard	Column #
Coding System for Extent of Disease	870	1	SEER	562-562
(SEER)				

#### Description

Indicates the type of SEER EOD code applied to the tumor. Should be used whenever EOD coding is applied.

#### Rationale

Used in data editing and analysis.

#### Codes

- 0 2-Digit Nonspecific Extent of Disease (1973-82)
- 1 2-Digit Site-Specific Extent of Disease (1973-82)
- 2 13-Digit (expanded) Site-Specific Extent of Disease (1973-1982)
- 3 4-Digit Extent of Disease (1983-87)
- 4 10-Digit Extent of Disease, 1988 (1988+)

<b>COMORBID/COMPLICATION 1</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #1	3110	5	COC	675-679

#### Description

Records the patient's pre-existing medical conditions and/or complications, during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to FORDS for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

00000 No secondary diagnoses documented

<b>COMORBID/COMPLICATION 2</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #2	3120	5	COC	680-684

Records the patient's pre-existing medical conditions and/or complications, during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to FORDS for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V00740-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

*Note:* For comorbid conditions (ICD-9-CM codes 00100-13980 and 24000-99990), there is an assumed decimal point between the third and fourth characters. For complications (ICD-9-CM codes E8700-E8799 and E9300-E9499), there is an assumed decimal point between the fourth and fifth characters. For conditions influencing health status and contact with health services (ICD-9-CM codes V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049), there is an assumed decimal point between the fourth and fifth characters.

<b>COMORBID/COMPLICATION 3</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #3	3130	5	COC	685-689

#### Description

Records the patient's pre-existing medical conditions and/or complications, during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to FORDS for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

<b>COMORBID/COMPLICATION 4</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #4	3140	5	COC	690-694

Records the patient's pre-existing medical conditions and/or complications during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to *FORDS* for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

*Note:* For comorbid conditions (ICD-9-CM codes 00100-13980 and 24000-99990), there is an assumed decimal point between the third and fourth characters. For complications (ICD-9-CM codes E8700-E8799 and E9300-E9499), there is an assumed decimal point between the fourth and fifth characters. For conditions influencing health status and contact with health services (ICD-9-CM codes V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049), there is an assumed decimal point between the fourth and fifth characters.

<b>COMORBID/COMPLICATION 5</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #5	3150	5	COC	695-699

#### Description

Records the patient's pre-existing medical conditions and/or complications during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to FORDS for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

<b>COMORBID/COMPLICATION 6</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Comorbidities and Complications #6	3160	5	COC	700-704

Records the patient's pre-existing medical conditions and/or complications during the patient's hospital stay for the treatment of this cancer. Both are considered secondary diagnoses.

#### Rationale

Comorbidities may affect treatment decisions and influence patient outcomes. Information on comorbidities is used to adjust risk outcome statistics when evaluating patient survival and other outcomes. Complications may be related to the quality-of-care.

#### Codes (refer to *FORDS* for additional instructions)

ICD-9-CM Codes 00100-13980, 24000-99990, E8700-E8799, E9300-E9499, V0720-V0739, V1000-V1590, V2220-V2310, V2540, V4400-V4589, and V5041-V5049

COMPUTED ETHNICITY				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	200	1	SEER	116-116

Code identifying those cases for which ethnicity was determined by matching Name--Last [2230] and Name--Maiden [2390] to a computer list of Spanish/Hispanic names or by a software algorithm. This field was adopted for use for tumors diagnosed 1994 forward.

See also Computed Ethnicity Source [210].

#### Rationale

One method of identifying persons of Hispanic origin is to apply a standard computer list or algorithm to items 2230 and 2390, the patient's surname and/or maiden name. This has advantages across large populations of being reproducible and facilitating comparisons between areas using identical methods. It may sometimes be possible to identify population denominators in which the same method was used to identify Hispanics. Generally, only central registries will have this capability.

This field provides coding to indicate both that such a computerized name-based method was applied and the results of the method. Coding is independent of that in Spanish/Hispanic Origin [190]. The computer-derived ethnicity may be different from the ethnicity reported by registries in Spanish/Hispanic Origin [190] as code 7 (Spanish Surname Only), because that field may include manual review. This field shows the results of computer-derived ethnicity only.

#### Codes

- 0 No match was run (for 1994 and later tumors)
- Non-Hispanic last name and non-Hispanic maiden name 1
- 2 Non-Hispanic last name, did not check maiden name or patient was male
- 3 Non-Hispanic last name, missing maiden name
- 4 Hispanic last name, non-Hispanic maiden name
- 5 Hispanic last name, did not check maiden name or patient was male
- Hispanic last name, missing maiden name 6
- 7 Hispanic Maiden name (females only) (regardless of last name)

Blank 1993 and earlier tumors, no match was run

Note: For SEER, blanks are required for all cases diagnosed before 1994 and blanks are not allowed for any case diagnosed 1994 and after. Other registries may have computed this item for earlier years.

Note: NAACCR recognizes that available definitions and abstracting instructions for the data items Name--Last and Name--Maiden may be inadequate for describing names used in some cultures, including Hispanic cultures. Explicit instructions have not been provided for entering compound names, with or without hyphens or "De." Order of names, use of maternal and paternal names, and use of hyphens can vary across cultures. It is likely, too, that abstracting and coding practice for these items varies across registries. Limitations inherent in these definitions should be kept in mind in any use of the data.

COMPUTED ETHNICITY SOURCE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	210	1	SEER	117-117

Code identifying the method used to determine ethnicity as recorded in Computed Ethnicity [200].

#### Codes

- 0 No match was run, for 1994 and later tumors
- 1 Census Bureau list of Spanish surnames, NOS
- 2 1980 Census Bureau list of Spanish surnames
- 3 1990 Census Bureau list of Spanish surnames
- 4 GUESS Program
- 5 Combination list including South Florida names
- 6 Combination of Census and other locally generated list
- 7 Combination of Census and GUESS, with or without other lists
- 8 Other type of match
- 9 Unknown type of match
- Blank 1993 and earlier tumors, no match was run

*Note:* For SEER, blanks are required for all cases diagnosed before 1994 and blanks are not allowed for any case diagnosed 1994 and after. Other registries may have computed this item for earlier years.

#### COUNTY AT DX

Alternate Name	Item #	Length	Source of Standard	Column #
County (pre-96 SEER/COC)	90	3	FIPS/SEER	83-85
County at Diagnosis (COC)				

#### Description

Code for the county of the patient's residence at the time the tumor was diagnosed. For U.S. residents, standard codes are those of the FIPS publication "Counties and Equivalent Entities of the United States, Its Possessions, and Associated Areas." If the patient has multiple tumors, the county codes may be different for each tumor.

Detailed standards have not been set for Canadian provinces/territories. Use code 998 for Canadian residents.

*Note:* The standard of using FIPS codes for this item has not been adopted by all states. Some states use their own codes for this data item. See Chapter V, Unresolved Issues, for further information.

*Note:* See Appendix A for standard FIPS county codes. See EDITS Table BPLACE.DBF in Appendix B for geocodes used by COC.

*Note:* SEER does not use code 998. COC uses country geocodes for nonresidents of the United States (see Appendix B) and 998 for residents of other states.

#### Codes (in addition to FIPS and Geocodes)

- 998 Known town, city, state, or country of residence but county code not known AND a resident outside of the state of reporting institution (must meet all criteria)
- 999 County unknown

COUNTYCURRENT				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1840	3	NAACCR	1338-1340

#### Description

Code for county of patient's current residence. See Chapter V, Unresolved Issues, for further discussion.

*Note:* This item was used by COC only. COC recommended use of FIPS codes (see Appendix A). The *ROADS Manual* also provided for use of geocodes for countries of residence outside the United States and Canada to be used in the county fields.

#### Rationale

This item may be used in administrative reports to define a referral area.

#### Codes (in addition to FIPS and geocodes)

- 998 Known town, city, state, or country of residence but county code not known AND a resident outside of the state of reporting institution (must meet all criteria)
- 999 County unknown

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### **CRC CHECKSUM**

Alternate Name	Item #	Length	Source of Standard	Column #
	2081	10	NAACCR	1164-1173

#### Description

Cyclic Redundancy Code (CRC) CHECKSUM for the NAACCR record in which it resides. A unique value is calculated for each unique record in a NAACCR file. The value is calculated by applying a CRC algorithm to all data fields of the NAACCR record (excluding the CRC CHECKSUM field). Following a transmission, the CRC CHECKSUM can be recalculated and compared with the transmitted CHECKSUM. Identical values indicate an error-free transmission; differing values indicate an error in transmission.

The algorithm recommended by NAACCR is on the NAACCR Web Site at: http://www.naaccr.org. Users must provide recipients of the data with the algorithm used to create the data transmission file. Otherwise, the item should be left blank.

#### Rationale

The CHECKSUM can be used to determine if a record-level error occurred during transmission and can also be used to correct any such errors. Record-level CRC CHECKSUMs also allow portions of a NAACCR file to be salvaged in the event of a transmission error.

#### **CS EXTENSION**

Alternate Name	Item #	Length	Source of Standard	Column #
	2810	2	AJCC	632-633

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. It is based on and replaces EOD--Extension (790) and EOD--Extension Prost Path (800). This modification for CS is collapsible into AJCC T code according to the Sixth Edition of *AJCC Cancer Staging Manual*. "CS Extension" identifies the primary tumor growth within the organ of origin or its extension into neighboring organs. For certain sites such as ovary, discontinuous metastasis is coded in the CS Extension field.

Site-specific codes provide extensive detail describing disease extent. "CS Extension" is used to derive the Derived AJCC T [2940], Derived AJCC Stage Group [3000], Derived SS1977 [3010], and Derived SS2000 [3020] codes.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

#### CS LYMPH NODES

Alternate Name	Item #	Length	Source of Standard	Column #
	2830	2	AJCC	635-636

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. It is based on and replaces EOD--Lymph Node Involv [810]. This modification for CS is collapsible into AJCC N code according to the Sixth Edition of *AJCC Cancer Staging Manual*. "CS Lymph Nodes" is site-specific and identifies the regional lymph nodes involved with cancer at the time of diagnosis.

Site-specific codes provide extensive detail describing disease extent. "CS Lymph Nodes" is used to derive the Derived AJCC N [2960], Derived AJCC Stage Group [3000], Derived SS1977 [3010], and Derived SS2000 [3020] codes.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

#### CS METS AT DX

Alternate Name	Item #	Length	Source of Standard	Column #
	2850	2	AJCC	638-639

#### Description

This belongs to the set of Collaborative Staging (CS) data items and is part of the detailed site-specific codes for anatomic EOD effective with 2004 diagnosis. It replaces data items 1090, 1100, and 1110 (Site of Distant Met 1-3). This modification for CS is collapsible into AJCC M code according to the Sixth Edition of *AJCC Cancer Staging Manual*. "CS Metastasis at Diagnosis" identifies the site(s) of metastatic involvement at time of diagnosis.

Site-specific codes provide extensive detail describing disease extent. "CS Mets at DX" is used to derive the Derived AJCC M [2980], Derived AJCC Stage Group [3000], Derived SS1977 [3010], and Derived SS2000 [3020] codes.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

### CS METS EVAL

Alternate Name	Item #	Length	Source of Standard	Column #
	2860	1	AJCC	640-640

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. "CS Mets Eval" records how the code for item. "CS Mets at DX" [2850] was determined based on the diagnostic methods employed.

This data item is used in CS to identify whether the M (of AJCC TNM) was clinically or pathologically diagnosed and by what methods, "CS Mets Eval" is used to derive the Derived AJCC M Descriptor [2990].

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

#### **CS REG NODE EVAL**

Alternate Name	Item #	Length	Source of Standard	Column #
	2840	1	AJCC	637-637

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. "CS Reg Nodes Eval" records how the code for the item "CS Lymph Nodes" [2830] was determined based on the diagnostic methods employed.

This data item is used in CS to identify whether the N (of AJCC TNM) was clinically or pathologically diagnosed and by what method "CS Reg Nodes Eval" is used to derive the Derived AJCC N Descriptor [2970].

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2880	3	AJCC	641-643

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2890	3	AJCC	644-646

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR records for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2900	3	AJCC	647-649

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR records for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2910	3	AJCC	650-652

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR records for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2920	3	AJCC	653-655

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR records for these outputs.

#### Codes

Alternate Name	Item #	Length	Source of Standard	Column #
	2930	3	AJCC	656-658

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. The "CS Site-Specific Factor" items (1-6) are used to code additional site-specific information needed to derive TNM or AJCC stage, or to code prognostic factors that have an effect on stage or survival. Some site-specific information that was formerly recorded in "EOD--Tumor Size" [780] (Breslow's Thickness for melanoma; HIV/AIDS status for lymphoma) is now also in "CS Site-Specific Factor" items. Many site-specific schemas do not use any of the Site-Specific Factors; other schemas use from 1 to all 6 of the factors.

#### Rationale

CS Site Specific Factors provide an optional area for coding information needed for stage. CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR records for these outputs.

#### Codes

#### CS TUMOR SIZE

Alternate Name	Item #	Length	Source of Standard	Column #
	2800	3	AJCC	629-631

#### Description

This item belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. It is based on and replaces EOD--Tumor Size [780]. For most sites, CS Tumor Size is used to record the largest dimension, or the diameter of the primary tumor in millimeters (for example: 1 mm = 001, 1 cm = 010). See the CS schemes for site-specific variants. For many sites, the CS algorithm uses this data item to derive the Derived AJCC T [2940] according to the Sixth Edition of *AJCC Cancer Staging Manual*.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

See the *Collaborative Staging Manual and Coding Instructions, Version 1.0*, for site-specific codes and coding rules.

#### CS TUMOR SIZE/EXT EVAL

Alternate Name	Item #	Length	Source of Standard	Column #
	2820	1	AJCC	634-634

#### Description

This belongs to the set of Collaborative Staging (CS) data items effective with 2004 diagnosis. "CS Tumor Size/Ext Eval" records how the codes for "CS Tumor Size" [2800] and "CS Extension" [2810] were determined based on the diagnostic methods employed. This data item is used in CS to identify whether the T (of AJCC TNM) was clinically or pathologically diagnosed and by what method, "CS Tumor Size/Ext Eval" is used to derive the Derived AJCC T Descriptor [2950].

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

#### Codes

#### **CS VERSION 1ST**

Alternate Name	Item #	Length	Source of Standard	Column #
	2935	6	AJCC	705-710

#### Description

This item indicates the number of the version used to initially code CS fields. The CS version number is returned as part of the output of the CS algorithm. As long as the CS algorithm is run and the output values stored at the time of initial abstracting, the returned values from the program should be automatically stored as CS Version 1st. This item may be blank if the CS algorithm has not been run or if this field has not been implemented. When it is implemented, this data item should be entered at the time the CS fields are first coded and the algorithm first applied. If the calculation algorithm is not called at the time of the initial abstracting, the CS Version 1st could also be entered manually by the abstractor.

It is not expected that this field would be updated every time a coded value is changed. However, the field should be available for future updating if, for example, the CS fields for certain records were to be systematically recoded for a special study using a later version, the CS Version 1st could be appropriately updated with the new version. The meaning and interpretation of CS Version 1st will be dependent on vendor implementation and local practices. This field should be interpreted with caution in a dataset where the actual coding procedures are unknown.

#### Codes

CS Version 1st is a 6-digit code. The first two digits represent the major version number; the second two digits represent minor version changes; and, the last two digits represent even less significant changes, such as corrections of typographical errors that do not affect coding or derivation of results.

#### **CS VERSION LATEST**

Alternate Name	Item #	Length	Source of Standard	Column #
	2936	6	AJCC	711-716

#### Description

This item indicates the number of the version of the CS used most recently to derive the CS output fields. This data item is recorded the first time the CS output fields are derived and should be updated each time the CS Derived items are re-computed. The CS version number is returned as part of the output of the CS algorithm. The returned value from the program should be automatically stored as CS Version Latest. This item should not be updated manually.

#### Codes

CS Version Latest is a 6-digit code. The first two digits represent the major version number; the second two digits represent minor version changes; and, the last two digits represent even less significant changes, such as corrections of typographical errors that do not affect coding or derivation results.

This item should not be blank if the CS Derived items contain stored values. This item should be blank if the CS Derived items are empty or the CS algorithm has not been applied.

DATE CASE COMPLETED				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2090	8	NAACCR	1174-1181

The date that: (1) the abstractor decided that the tumor report was complete, and (2) the case passed all edits that were applied. Definitions may vary among registries and software providers. This is a local use field. See page 83 for date format. Standard edits check that no dates are later than the current date.

# DATE CASE LAST CHANGEDRevisedAlternate NameItem #LengthSource of StandardColumn #21008NAACCR1182-1189

#### Description

Date the case was last changed or updated. See page 83 for date format. Standard edits check that no dates are later than the current date. Definitions may vary among areas.

DATE CASE REPORT EXPORTED				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date Case Transmitted (pre-98 NAACCR)	2110	8	NPCR	1190-1197

#### Description

Date the reporting facility exports the electronic abstract to a file for transmission to the central registry via diskette or other electronic medium. See page 83 for date format. Standard edits check that no dates are later than the current date.

Definitions may vary among registries and software providers. This item is not yet well defined for use when a central registry is creating a transmission record for a consolidated tumor record from multiple source records.

DATE CASE REPORT LOADED				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2112	8	NPCR	1227-1234

#### Description

Date the tumor report is loaded into a central registry computerized processing file for initiation of quality control activities (e.g., visual editing, application of computerized edits, etc.). See page 83 for date format.

This item is not yet well defined for use when a central registry is creating a transmission record for a consolidated tumor record from multiple source records.

DATE CASE REPORT RECEIVED				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2111	8	NPCR	1219-1226

Date the electronic or paper abstract (or source record) is received by the central cancer registry for the respective tumor. If multiple reports are received from two or more sources and if a single date is needed, use the date the first abstract (or source record) was received from any source. See page 83 for date format.

#### Rationale

This item is used to assess and monitor the timeliness of reporting. Timeliness of abstracting (and reporting) is a concern for all standard-setting organizations and consequently, timeliness standards have been established. This item can be used with the Date of 1st Contact [580] or the Path--Date of Specimen Collection [7320] to measure timeliness of reporting by individual reporting facilities to central cancer registries. This data item also can be used with the Date Tumor Record Availbl [2113] to measure timeliness of processing within the central cancer registry.

DATE OF 1ST CONTACT				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date of Adm/1st Contact	580	8	COC	416-423

Date of first patient contact, as inpatient or outpatient, with the reporting facility for the diagnosis and/or treatment of the tumor. The date may represent the date of an outpatient visit for a biopsy, x-ray, scan, or laboratory test. See page 83 for date format.

When Pathology Specimen Only (Class of Case 7, Type of Reporting Source 3) tumors are collected, the Path--Date of Specimen Collection [7320] from the pathology report should be used for the Date of 1st Contact. If a pathology-specimen-only case is followed by a patient contact with the facility for the diagnosis and/or treatment of the respective tumor, the Date of 1st Contact is not changed. The date of the initial pathology laboratory specimen collection remains the Date of 1st Contact.

When Death Certificate Only (Class of Case 8, Type of Reporting Source 7) tumors are collected, the date of death should be used as the Date of 1st Contact. When Autopsy Only (Class of Case 5, Type of Reporting Source 6) tumors are collected, the date of death should be used as the Date of 1st Contact.

# Rationale

Timeliness of abstracting (and reporting) is a concern for all standard-setting organizations. Date of 1st Contact is one of several data items that can be used to measure timeliness of reporting by individual facilities to central cancer registries. For tumors that are not diagnosed at the reporting facility (Class of Case 2, 3, or 4), the Date of 1st Contact [580] can be used in conjunction with the Date Case Report Received [2111] to measure timeliness of reporting by individual facilities. To accurately measure the timeliness of data collection and submission of abstracts that are first diagnosed at autopsy (Class of Case 5, Type of Reporting Source 6) the date of death should be used as the Date of 1st Contact since the diagnosis was not determined until the autopsy was performed. Death Certificate Only cases (Class of Case 8, Type of Reporting Source 7) are created only by the central registry. For these cases, Date of 1st Contact should be filled with the date of death, and timeliness for DCO cases should be measured by different criteria.

# DATE OF 1ST CRS RX--COC

Alternate Name	Item #	Length	Source of Standard	Column #
Date of First Course Treatment (COC)	1270	8	COC	843-850
Date Started (pre-96 COC)				

# Description

Date of initiation of the first cancer-directed therapy for the cancer being reported, using the COC definition of first course. The date of first treatment includes the date a decision was made not to treat the patient. See *FORDS* for details. See Chapter V, Unresolved Issues for further discussion of the difference between SEER and COC items. See page 83 for date format.

# Codes (in addition to valid dates)

00000000Diagnosed at autopsy.99999999When it is unknown whether any treatment was administered to the patient, the date is<br/>unknown or the case was identified by death certificate-only.

# **Clarification of NPCR Required Status**

Central registries funded by NPCR are required to collect either Date of Initial RX--SEER [1260] or Date of 1st Crs RX--COC [1270].

# DATE OF 1ST POSITIVE BX

Alternate Name	Item #	Length	Source of Standard	Column #
Date of First Positive Biopsy (COC)	1080	8	COC	610-617

# Description

Date of first positive tissue biopsy/positive histology. See page 83 for date format.

# Codes (in addition to valid dates)

00000000 Positive biopsy never obtained

Note: This data item is no longer supported by COC (as of January 1, 2003).

# DATE OF CA CONFERENCE

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Cancer Conference (COC)	660	8	COC	449-456

#### Description

Date on which the case was first presented at cancer conference at the reporting facility. See page 83 for date format.

#### Rationale

Collection of this item and Presentation at CA Conf [650] allows preparation of reports on the contents of cancer conferences: sites presented, types of presentation for administrative use, quality control, and survey preparation.

#### Special Codes (in addition to valid dates)

00000000Case was never presented at cancer conference999999999Unknown

Note: This data item is no longer supported by COC (as January 1, 2003).

#### **DATE OF DIAGNOSIS**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Initial Diagnosis (COC)	390	8	SEER/COC	283-290

#### Description

Date of initial diagnosis by a recognized medical practitioner for the tumor being reported whether clinically or microscopically confirmed. See page 83 for date format.

For more discussion on determining date of diagnosis, consult the SEER Program Manual or COC FORDS Manual.

#### DATE OF INITIAL RX--SEER

Alternate Name	Item #	Length	Source of Standard	Column #
Date Therapy Initiated (SEER)	1260	8	SEER	835-842
Date Started (SEER)				

#### Description

Date of initiation of the first cancer-directed therapy for the tumor being reported, using the SEER definition of first course. See also Date of 1st Crs RX--COC [1270]. See Chapter V, Unresolved Issues, for further discussion of the difference between SEER and COC items. See page 83 for date format.

#### Codes (in addition to valid dates)

0000000	No cancer-directed therapy
99999999	Unknown if any cancer-directed therapy was administered

#### **Clarification of NPCR Required Status**

Central registries funded by NPCR are required to collect either Date of Initial RX--SEER [1260] or Date of 1st Crs RX--COC [1270].

DATE OF INPATIENT ADM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date of Inpatient Admission (COC)	590	8	NAACCR	424-431

Date of the inpatient admission to the reporting facility for the most definitive surgery. In the absence of surgery, use date of inpatient admission for any other cancer-directed therapy. In the absence of cancer-directed therapy, use date of inpatient admission for diagnostic evaluation. See page 83 for date format.

#### Codes (in addition to valid dates)

00000000Patient was never an inpatient at the reporting facility999999999Unknown

Note: This data item is no longer supported by COC (as of January 1, 2003).

DATE OF INPATIENT DISCH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date of Inpatient Discharge (COC)	600	8	NAACCR	432-439

#### Description

Date of the inpatient discharge from the reporting facility after the most definitive surgery. In the absence of surgery, use date of inpatient discharge for other cancer-directed therapy. In the absence of cancer-directed therapy, use date of inpatient discharge for diagnostic evaluation. This discharge date corresponds to the admission date described by Date of Inpatient Adm [590]. See page 83 for date format.

*Note:* This item is not the same as the old NAACCR item, Date of Discharge, which has been deleted from the NAACCR layout.

#### Special Codes (in addition to a valid date

00000000Patient was never an inpatient at the reporting hospital999999999Unknown

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### DATE OF LAST CONTACT

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Last Contact or Death (COC)	1750	8	SEER/COC	1294-1301
Date of Last Follow-Up or of Death				
(SEER)				

#### Description

Date of last contact with the patient, or date of death. If the patient has multiple tumors, Date of Last Contact should be the same for all tumors. See page 83 for date format.

#### Rationale

Used for Date of Last Contact from active or passive follow-up. Used to record date of death.

DATE TUMOR RECORD AVAILBL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2113	8	NPCR	1235-1242

Date the demographic and tumor identification information on a single primary/reportable neoplasm, compiled from one or more source records, from one or more facilities, is available in the central cancer registry database to be counted as an incident tumor. Cancer identification information includes, at a minimum, site, histology, laterality, behavior, and date of diagnosis. See page 83 for date format.

#### Rationale

This item is used to assess and monitor the timeliness of reporting. Timeliness of abstracting (and reporting) is a concern for all standard-setting organizations and consequently, timeliness standards have been established. This data item can be used with the Date Case Report Received [2111] to measure timeliness of processing within the central cancer registry. This item also can be used with the Date of 1st Contact [580] or the Path-Date of Specimen Collection [7320] to measure overall timeliness.

DC STATE				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2370			

#### Description

See Place of Death [1940].

#### DC STATE FILE NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
	2380	6	State	2278-2283

#### Description

Death certificate identification number as assigned by the vital statistics office in the place recorded in Place of Death [1940].

DERIVED AJCC M				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2980	2	AJCC	665-666

This is the AJCC "M" component that is derived from CS coded fields, using the CS algorithm, effective with 2004 diagnosis.

# Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form and adds several additional fields. When CS data items are coded, a computer algorithm provided by the Task Force allows generation of AJCC Sixth Edition TNM stage, Summary Stage 1977, and Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

The Storage Code column is the value to be stored in the NAACCR record. The 2-character numeric Storage Codes are designed for analysis purposes. The Display String is the corresponding label that should be displayed on the screen or in a report. The meaning of these strings is explained for each site in the AJCC Manual.

M Storage Code	<b>Display String</b>	Comments
99	MX	MX
00	M0	M0
10	M1	M1
11	Mla	M1a
12	M1b	M1b
13	M1c	M1c
19	M1NOS	M1 NOS
88	NA	Not applicable

DERIVED AJCC M DESCRIPTOR				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2990	1	AJCC	667-667

This is the AJCC "M Descriptor" component that is derived from coded fields, using the CS algorithm, effective with 2004 diagnosis.

# Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

This derived output records a "c," "p," "a," "y," or "N" for "clinical," "pathological," "autopsy only," or "y prefix," respectively. For those tumors in which staging classification is performed during or following initial multimodality therapy, the category is identified by a "y prefix" to be derived from the computerized algorithm.

- c Clinical stage
- p Pathologic stage
- a Autopsy stage
- y Pathologic examination of metastatic tissue performed after presurgical systemic treatment or radiation, and extension based on pathologic evidence
- N Not applicable
- Blank Not derived

DERIVED AJCC N				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2960	2	AJCC	662-663

This is the AJCC "N" component that is derived from coded fields, using the CS algorithm, effective with 2004 diagnosis.

# Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

The Storage Code column is the value to be stored in the NAACCR record. The 2-character numeric Storage Codes are designed for analysis purposes. The Display String is the corresponding label that should be displayed on the screen or in a report. The meaning of these strings is explained for each site in the AJCC Manual.

N Storage Codes	<b>Display String</b>	Comments
00	NO	N0
09	N0NOS	N0 NOS
01	N0(i-)	N0(i-)
02	N0(i+)	N0(i+)
03	N0(mol-)	N0(mol-)
04	N0(mol+)	N0(mol+)
10	N1	N1
19	N1NOS	N1 NOS
11	Nla	N1a
12	N1b	N1b
13	N1c	N1c
18	N1mi	N1mi
20	N2	N2
29	N2NOS	N2 NOS
21	N2a	N2a
22	N2b	N2b
23	N2c	N2c
30	N3	N3
39	N3NOS	N3 NOS
31	N3a	N3a
32	N3b	N3b
33	N3c	N3c
88	NA	Not applicable
99	NX	NX

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DERIVED AJCC N DESCRIPTOR				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2970	1	AJCC	664-664

This is the AJCC "N Descriptor" component that is derived from coded fields using the CS algorithm, effective with 2004 diagnosis.

# Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACOS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

This derived output records a "c," "p," "a," "y," or "N" for "clinical," "pathological," "autopsy only," or "y prefix", respectively. For those tumors in which AJCC TNM staging classification is performed during or following initial multimodality therapy, the category is identified by a "y prefix" to be derived from the computerized algorithm.

- c Clinical stage
- p Pathologic stage
- a Autopsy stage
- y Lymph nodes removed for examination after presurgical systemic treatment or radiation, and lymph node evaluation based on pathologic evidence
- N Not applicable
- Blank Not derived

DERIVED AJCC STAGE GROUP				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	3000	2	AJCC	668-669

This is the AJCC "Stage Group" component that is derived from the CS detailed site-specific codes, using the CS from the CS algorithm effective with 2004 diagnosis.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

Codes		
AJCC Storage Code	<b>Display String</b>	Comments
00	0	Stage 0
01	0a	Stage 0a
02	0is	Stage 0is
10	Ι	Stage I
11	INOS	Stage I NOS
12	IA	Stage IA
13	IA1	Stage IA1
14	IA2	Stage IA2
15	IB	Stage IB
16	IB1	Stage IB1
17	IB2	Stage IB2
18	IC	Stage IC
19	IS	Stage IS
23	ISA	Stage ISA (lymphoma only)
24	ISB	Stage ISB (lymphoma only)
20	IEA	Stage IEA (lymphoma only)
21	IEB	Stage IEB (lymphoma only)
22	IE	Stage IE (lymphoma only)
30	II	Stage II
31	IINOS	Stage II NOS
32	IIA	Stage IIA
33	IIB	Stage IIB
34	IIC	Stage IIC
35	IIEA	Stage IIEA (lymphoma only)
36	IIEB	Stage IIEB (lymphoma only)
37	IIE	Stage IIE (lymphoma only)
38	IISA	Stage IISA (lymphoma only)
39	IISB	Stage IISB (lymphoma only)
40	IIS	Stage IIS (lymphoma only)

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41	IIESA	Stage IIESA (lymphoma only)
42	IIESB	Stage IIESB (lymphoma only)
43	IIES	Stage IIES (lymphoma only)
50	III	Stage III
51	IIINOS	Stage III NOS
52	IIIA	Stage IIIA
53	IIIB	Stage IIIB
54	IIIC	Stage IIIC
55	IIIEA	Stage IIIEA (lymphoma only)
56	IIIEB	Stage IIIEB (lymphoma only)
57	IIIE	Stage IIIE (lymphoma only)
58	IIISA	Stage IIISA (lymphoma only)
59	IIISB	Stage IIISB (lymphoma only)
60	IIIS	Stage IIIS (lymphoma only)
61	IIIESA	Stage IIIESA (lymphoma only)
62	IIIESB	Stage IIIESB (lymphoma only)
63	IIIES	Stage IIIES (lymphoma only)
70	IV	Stage IV
71	IVNOS	Stage IV NOS
72	IVA	Stage IVA
73	IVB	Stage IVB
74	IVC	Stage IVC
88	NA	Not applicable
90	OCCULT	Stage Occult
99	UNK	Stage Unknown

DERIVED AJCC T				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2940	2	AJCC	659-660

This is the AJCC "T" component that is derived from CS coded fields, using the CS algorithm, effective with 2004 diagnosis.

# Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

The Storage Code column is the value to be stored in the NAACCR record. The 2-character numeric Storage Codes are designed for analysis purposes. The Display String is the corresponding label that should be displayed on the screen or in a report. The meaning of these strings is explained for each site in the AJCC Manual.

T Storage Code	<b>Display String</b>	Comments
99	TX	TX
00	TO	T0
01	Та	Та
05	Tis	Tis
06	Tispu	Tispu (urethra only)
07	Tispd	Tispd (urethra only)
10	T1	T1
11	T1mic	T1mic
19	T1NOS	T1 NOS
12	T1a	T1a
13	Tlal	Tlal
14	T1a2	T1a2
15	T1b	T1b
16	T1b1	T1b1
17	T1b2	T1b2
18	T1c	T1c
20	T2	T2
29	T2NOS	T2NOS
21	T2a	T2a
22	T2b	T2b
23	T2c	T2c
30	Т3	T3
39	T3NOS	T3 NOS
31	T3a	T3a

32	T3b	T3b
33	T3c	T3c
40	T4	T4
49	T4NOS	T4 NOS
41	T4a	T4a
42	T4b	T4b
43	T4c	T4c
44	T4d	T4d
88	NA	Not applicable

DERIVED AJCC T DESCRIPTOR				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2950	1	AJCC	661-661

This is the AJCC "T Descriptor" component that is derived from CS coded fields, using the CS algorithm, effective with 2004 diagnosis.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

This derived output records a "c," "p," "a," "y," or "N" for "clinical," "pathological," "autopsy only," or "y prefix", respectively. For those cases in which staging classification is performed during or following initial multimodality therapy, the category is identified by a "y prefix" to be derived from the computerized algorithm.

#### Codes

- c Clinical stage
- p Pathologic stage
- a Autopsy stage
- y Surgical resection performed after presurgical systemic treatment or radiation; tumor size/extension based on pathologic evidence
- N Not applicable
- Blank Not derived

DERIVED AJCCFLAG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	3030	1	AJCC	672-672

Flag to indicate whether the derived AJCC stage was coded directly or was derived from CS or EOD codes.

#### Codes

- 1 AJCC Sixth Edition derived from *Collaborative Staging Manual and Coding Instructions, Version 1.0*
- 2 AJCC Sixth Edition derived from EOD (prior to 2004)

Blank Not derived

#### **DERIVED SS1977**

Alternate Name	Item #	Length	Source of Standard	Column #
	3010	1	AJCC	670-670

#### Description

This item is the derived "SEER Summary Stage 1977" from the CS algorithm (or EOD codes) effective with 2004 diagnosis.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

Storage Code	<b>Display String</b>	Comments
	ERROR	Processing error (no storage code needed)
	NONE	None (internal use only, no storage code needed)
0	IS	In situ
1	L	Localized
2	RE	Regional, direct extension
3	RN	Regional, lymph nodes only
4	RE+RN	Regional, extension and nodes
5	RNOS	Regional, NOS
7	D	Distant
8	NA	Not applicable
9	U	Unknown/Unstaged

DERIVED SS1977FLAG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	3040	1	AJCC	673-673

Flag to indicate whether the derived SEER Summary Stage 1977 was coded directly or was derived from CS or EOD codes.

# Codes

1 SS1977 derived from Collaborative Staging Manual and Coding Instructions, Version 1.0

2 SS1977 derived from EOD (prior to 2004)

Blank Not derived

#### **DERIVED SS2000**

Alternate Name	Item #	Length	Source of Standard	Column #
	3020	1	AJCC	671-671

#### Description

This item is the derived "SEER Summary Stage 2000" from the CS algorithm (or EOD codes) effective with 2004 diagnosis.

#### Rationale

CS is a group of data items set up by a joint task force including representatives from SEER, ACoS, CDC, NAACCR, NCRA, and AJCC, designed to provide a single uniform set of codes and rules for coding EOD/ stage information to meet the needs of all of the participating standard setters. CS incorporates all of the fields from SEER 10-digit EOD, in a modified form, and adds several additional fields. When CS data items are coded, a computer algorithm allows generation of AJCC Sixth Edition TNM stage, SEER Summary Stage 1977, and SEER Summary Stage 2000. Separate CS fields are provided in the NAACCR record for these outputs.

Storage Code Display String Comments				
	ERROR	Processing error (no storage code needed)		
	NONE	None (internal use only, no storage code needed)		
0	IS	In situ		
1	L	Localized		
2	RE	Regional, direct extension		
3	RN	Regional, lymph nodes only		
4	RE+RN	Regional, extension and nodes		
5	RNOS	Regional, NOS		
7	D	Distant		
8	NA	Not applicable		
9	U	Unknown/Unstaged		

DERIVED SS2000FLAG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	3050	1	AJCC	674-674

Flag to indicate whether the derived SEER Summary Stage 2000 was coded directly or was derived from CS or EOD codes.

#### Codes

- 1 SS2000 derived from Collaborative Staging Manual and Coding Instructions, Version 1.0
- 2 SS2000 derived from EOD (prior to 2004)

Blank Not derived

#### **DIAGNOSTIC CONFIRMATION**

Alternate Name	Item #	Length	Source of Standard	Column #
	490	1	SEER/COC	311-311

# Description

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

#### Rationale

Diagnostic confirmation is useful to calculate rates based on microscopically confirmed cancers. Full incidence calculations must also include tumors that are only confirmed clinically. The percentage of tumors that are clinically diagnosed only is an indication of whether case finding is including sources outside of pathology reports.

#### Codes

- 1 Positive histology
- 2 Positive cytology, no positive histology
- 4 Positive microscopic confirmation, method not specified
- 5 Positive laboratory test/marker study
- 6 Direct visualization without microscopic confirmation
- 7 Radiography and other imaging techniques without microscopic confirmation
- 8 Clinical diagnosis only (other than 5, 6, or 7)
- 9 Unknown whether or not microscopically confirmed

#### DIAGNOSTIC PROC 73-87

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic Procedures (1973-87 SEER)	2200	2	SEER	1217-1218

#### Description

Data item required by SEER for tumors of certain sites for the years 1973-87. This item is no longer collected. See Appendix D of the *SEER Program Code Manual* for details.

# **EOD--EXTENSION**

Alternate Name	Item #	Length	Source of Standard	Column #
Extension (pre-96 SEER/COC)	790	2	SEER	534-535
Extension (SEER EOD) (96 COC)				

# Description

Part of the 10-digit EOD [779]. Detailed site-specific codes for anatomic EOD used by SEER for tumors diagnosed from 1988 forward.

Codes were revised effective January 1, 1998, to reflect changes in the *AJCC Cancer Staging Manual*, Fifth Edition.

#### Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

# Codes

See SEER Extent of Disease, 1988: Codes and Coding Instructions, Third Edition for site-specific codes and coding rules for all EOD fields.

# EOD--EXTENSION PROST PATH

Alternate Name	Item #	Length	Source of Standard	Column #
	800	2	SEER	536-537

#### Description

Part of the 10-digit EOD [779]. Detailed site-specific codes for anatomic EOD used by SEER for tumors diagnosed from 1988 forward.

Codes were revised effective January 1, 1998, to reflect changes in the *AJCC Cancer Staging Manual*, Fifth Edition.

#### Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

EOD--Extension Prost Path is an additional field for prostate cancer only to reflect information from radical prostatectomy, effective with 1995 diagnoses. The field is left blank for all other primaries.

#### Codes

See SEER Extent of Disease, 1988: Codes and Coding Instructions, Third Edition, for site-specific codes and coding rules for all EOD fields.

# **EOD--LYMPH NODE INVOLV**

Alternate Name	Item #	Length	Source of Standard	Column #
Lymph Nodes (pre-96 SEER/COC)	810	1	SEER	538-538
Lymph Nodes (SEER EOD) (96 COC)				

# Description

Part of the 10-digit EOD [779]. Detailed site-specific codes for anatomic EOD used by SEER for tumors diagnosed from 1988 forward.

Codes were revised effective January 1, 1998, to reflect changes in the *AJCC Cancer Staging Manual*, Fifth Edition.

# Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

# Codes

See SEER Extent of Disease, 1988: Codes and Coding Instructions, Third Edition for site-specific codes and coding rules for all EOD fields.

# EOD--OLD 2 DIGIT

Alternate Name	Item #	Length	Source of Standard	Column #
2-Digit Nonspecific and 2-Digit Site- Specific Extent of Disease (1973-1982 SEER)	850	2	SEER	556-557

# Description

Site-specific codes for EOD used by SEER for tumors diagnosed from January 1, 1973, to December 31, 1982, for cancer sites that did not have a 13-digit scheme see EOD--Old 13 Digit [840].

# Codes

See Extent of Disease: Codes and Coding Instructions (SEER 1977) for codes.

# **EOD--OLD 4 DIGIT**

Alternate Name	Item #	Length	Source of Standard	Column #
4-Digit Extent of Disease (1983-1987	860	4	SEER	558-561
SEER)				

# Description

Codes for site-specific EOD used by SEER for tumors diagnosed from January 1, 1983, to December 31, 1987, for all cancer sites.

# Codes

See SEER Extent of Disease: New 4-Digit Schemes: Codes and Coding Instructions for codes.

# EOD--OLD 13 DIGIT

Alternate Name	Item #	Length	Source of Standard	Column #
13-Digit (Expanded) Site-Specific Extent	840	13	SEER	543-555
of Disease (SEER)				
SEER EEOD (SEER)				

# Description

Detailed site-specific codes for EOD used by SEER for selected sites of cancer for tumors diagnosed 1973-1982, except death-certificate-only cases.

# Codes

See Extent of Disease: Codes and Coding Instructions (SEER 1977) for codes.

#### **EOD--TUMOR SIZE**

Alternate Name	Item #	Length	Source of Standard	Column #
Size of Primary Tumor (SEER)	780	3	SEER/COC	531-533
Size of Tumor (COC)				

#### Description

Part of the 10-digit EOD [779]. Detailed site-specific codes for anatomic EOD used by SEER for tumors diagnosed from 1988 forward.

This field is included in the COC dataset, separate from EOD.

Codes were revised effective January 1, 1998, to reflect changes in the *AJCC Cancer Staging Manual*, Fifth Edition.

#### Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

#### Codes

See *SEER Extent of Disease, 1988: Codes and Coding Instructions*, Third Edition, for site-specific codes and coding rules for all EOD fields. The COC codes for Tumor Size are in the *FORDS* Manual.

Note: See Chapter V, Unresolved Issues, for a discussion of coding differences between COC and SEER.

# **EXTENT OF DISEASE 10-DIG**

Alternate Name	Item #	Length	Source of Standard	Column #
	779	12		531-542

# Description

The name for a group of subfields that contain detailed site-specific codes for the anatomic EOD. SEER uses the subfields for tumors diagnosed from 1988 forward.

Group names appear only in the data dictionary and in Appendix E.

# Subfields

EOD--Tumor Size [780] EOD--Extension [790] EOD--Extension Prost Path [800] EOD--Lymph Node Involv [810] Regional Nodes Positive [820] Regional Nodes Examined [830]

# FAMILY HISTORY OF CANCER

Alternate Name	Item #	Length	Source of Standard	Column #
	360	1	Varies	226-226

#### Description

NAACCR has not adopted standards for this item.

#### FIN CODING SYSTEM

Alternate Name	Item #	Length	Source of Standard	Column #
	35	1	NAACCR	11-11

#### Description

The FIN coding system is a generated code that identifies the coding system used for individual facilities (hospital, clinics, or other providers) submitting data to a registry. This field identifies the coding system used for facilities in the following seven fields of the NAACCR layout:

Registry ID [40] (when Registry Type [30] = 3) Reporting Hospital [540] Institution Referred From [2410] Institution Referred To [2420] Last Follow-Up Hospital [2430] Following Registry [2440] Archive FIN [3100]

Within a single NAACCR record, all of these fields must be coded using the same FIN coding system.

#### Codes

- 1 COC 7-digit codes (assigned by COC until the end of 2000)
- 2 COC FIN 10-digit codes (assigned 2001+)
- 3 NPI 8-digit codes
- 9 Unknown

Note: Code 4, 15-digit codes, has been deleted.

#### FIRST COURSE CALC METHOD

Alternate Name	Item #	Length	Source of Standard	Column #
	1500	1	NAACCR	894-894

#### Description

Codes indicating the time interval for defining the first course of therapy.

- 1 COC definitions
- 2 SEER definitions
- 9 Other, unknown

FOLLOWING REGISTRY				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2440	10	COC	2475-2484

Records registry responsible for following the patient.

#### Rationale

Each FIN is unique. The number is essential to NCDB for monitoring data submissions, ensuring the accuracy of data, and identifying areas for special studies. Efforts are underway at the federal level to establish uniform national provider ID numbers. COC and NAACCR committees will consider adoption of any federal standards when they become available.

#### **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001: Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10 digit codes.

#### Codes (in addition to COC assigned codes)

000000000Case not reported by a facility00999999999Case reported, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN coding system [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

FOLLOW-UP CONTACTCITY				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1842	20	SEER	1357-1376

#### Description

Name of the city of the follow-up contact's current usual residence. If the patient has multiple tumors, the follow-up contact city of residence should be the same for all tumors.

#### Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

FOLLOW-UP CONTACTNAME				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2394	30	SEER	2284-2313

First and last name, in natural order, of a person, other than the patient or a physician, who can be contacted to obtain follow-up information for the patient. See Follow-Up Contact--City [1842] for further explanation.

#### Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

#### FOLLOW-UP CONTACT--NO&ST

rollow-or contact-noast				IXC viscu
Alternate Name	Item #	Length	Source of Standard	Column #
	2392	40	SEER	2314-2353

#### Description

The number and street address or the rural mailing address of the follow-up contact's current usual residence. This can be used to generate a follow-up inquiry, and must correspond to the other fields in the follow-up contact address. If the patient has multiple tumors, Follow-Up Contact--No&St should be the same. See Follow-Up Contact--City [1842] for rationale and further description.

U.S. addresses should conform to the USPS Postal Addressing Standards. These standards are referenced in USPS Pub. 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/Pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current Canadian Postal Address standards may be found at the following Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

#### Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current address in the NAACCR record layout.

Revised

FOLLOW-UP CONTACTPOSTAL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1846	9	SEER	1379-1387

Postal code for the address of the follow-up contact's current usual residence. If the patient has multiple tumors, the follow-up contact postal codes should be the same for all tumors. For U.S. residents, use either the 5-digit or the extended 9-digit ZIP code. Blanks follow the 5-digit code. For Canadian residents, use the 6-character, alphanumeric postal code. Blanks follow the 6-character code. When available, enter postal code for other countries.

# Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

# Codes (in addition to U.S., Canadian, and foreign postal codes)

- 888888888 Resident of country other than the United States (including its possessions, etc.) or Canada, and postal code unknown
- 999999999 Resident of the United States (including its possessions, etc.) or Canada, and postal code unknown

FOLLOW-UP CONTACTSTATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1844	2	SEER	1377-1378

#### Description

USPS abbreviation for the state (including U.S. territories, commonwealths, or possessions), or Canada Post abbreviation for the Canadian province/territory of the follow-up contact's current usual residence. If the patient has multiple tumors, the follow-up contact state should be the same for all tumors.

#### Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

# Codes (in addition to USPS and Canadian Postal Service abbreviations)

- XX Resident of country other than the United States (including its territories, commonwealths, or possessions) or Canada, and country is known
- YY Resident of country other than the United States (including its territories, commonwealths, or possessions) or Canada, and country is unknown
- ZZ Resident of the United States, NOS (including its territories, commonwealths, or possessions); Canada, NOS; residence unknown

FOLLOW-UP CONTACTSUPPL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2393	40	SEER	2354-2393

This data item provides the ability to store additional address information such as the name of a place or facility, a nursing home, or the name of an apartment complex. It can be used to generate a follow-up inquiry, and must correspond to the other fields in the follow-up contact address. If the patient has multiple tumors, Follow-Up Contact--Suppl should be the same. See Follow-Up Contact--City [1842] for rationale and further description.

U.S. addresses should conform to the USPS Postal Addressing Standards. These standards are referenced in USPS Pub. 28, November 2000, *Postal Addressing Standards*. The current USPS Pub. 28 may be found and downloaded from the following Web Site: http://pe.usps.gov/cpim/ftp/pubs/Pub28/pub28.pdf.

Canadian addresses should conform to the *Canada Postal Guide*. The current *Canada Postal Guide* may be found at the following Web Site: http://www.canadapost.ca/tools/pg/manual/b03-e.asp#top.

# Rationale

Sometimes registries carry out follow-up by contacting the patient and other contacts by a letter or phone call to ascertain their vital status. When a patient's current address is unknown or the patient is for some reason not to be contacted (e.g., patient is a minor child), the most current name, address and phone number of another contact, such as a relative or neighbor are needed. This information may also be useful for conducting epidemiological or research studies.

#### **Coding Instructions (summary of USPS guidelines)**

The address should be fully spelled out with standardized use of abbreviations and punctuation per USPS postal addressing standards (USPS *Postal Addressing Standards*, Pub. 28, November 2000). Upper case recommended. Mixed case allowed.

Abbreviations should be limited to those recognized by the Postal Service standard abbreviations, these include but are not limited to (a complete list of recognized abbreviations is provided in Appendix C of USPS Pub. 28):

APT	apartment	Ν	north
BLDG	building	NE	northeast
FL	floor	NW	northwest
STE	suite	S	south
UNIT	unit	SE	southeast
RM	room	SW	southwest
DEPT	department	E	east

#### **FOLLOW-UP SOURCE**

Alternate Name	Item #	Length	Source of Standard	Column #
Follow-Up Method (pre-96 COC)	1790	1	COC	1305-1305

# Description

Records the source from which the latest follow-up information was obtained.

#### Rationale

For registries performing follow-up, this field helps evaluate the success rates of various methods of followup. It also can be used to report to institutions the source of follow-up information that is sent to them. When there is a conflict in follow-up information, knowing the source can help resolve the inconsistency.

#### Codes

- 0 Reported hospitalization
- 1 Readmission
- 2 Physician
- 3 Patient
- 4 Department of Motor Vehicles
- 5 Medicare/Medicaid file
- 7 Death certificate
- 8 Other
- 9 Unknown, not stated in patient record

#### 

#### Description

Reserved for future use for storing date of a central registry processing milestone. No standards have been adopted for this item. The NAACCR UDSC approved to retire this data item in Version 10.1.

FUTURE USE TIMELINESS 2				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2115			

#### Description

Reserved for future use for storing date of a central registry processing milestone. No standards have been adopted for this item. The NAACCR UDSC approved to retire this data item in Version 10.1.

Retired

Column #

# GRADE

Alternate Name	Item #	Length	Source of Standard	Column #
Grade, Differentiation, or Cell Indicator	440	1	SEER/COC	306-306
(SEER)				
Grade/Differentiation (COC)				

# Description

Code for the grade or degree of differentiation of the reportable tumor. For lymphomas and leukemias, field also is used to indicate T-, B-, Null-, or NK-cell origin.

Note: Code 8 was adopted for use with lymphoma cases diagnosed in 1995 and later.

# Codes

See the grade tables on page 67 of ICD-O-3.<sup>14</sup> See also the COC *FORDS* Manual and The *SEER Program Code Manual*, Third Edition, for site-specific coding rules and conversions.

- 1 Grade I
- 2 Grade II
- 3 Grade III
- 4 Grade IV
- 5 T-cell
- 6 B-cell
- 7 Null cell
- 8 NK (natural killer) cell
- 9 Grade/differentiation unknown, not stated, or not applicable

#### **GRADE (73-91) ICD-O-1**

Alternate Name	Item #	Length	Source of Standard	Column #
	1973	1	SEER	1146-1146

#### Description

Area for retaining the grade portion (1 digit) of the ICD-O-1 or field trial grade code entered before a conversion to ICD-O-2. See grouped data item Morph (73-91) ICD-O-1 [1970] in Appendix E. The item name includes years 1973-91. However, some states may have used the codes for cases before 1973.

#### Codes

For cases diagnosed before 1992, contains the ICD-O-1 or field trial 1-digit grade code as originally coded, if available.

# HISTOLOGIC TYPE ICD-O-3

Alternate Name	Item #	Length	Source of Standard	Column #
	522	4	SEER/COC	301-304

# Description

Codes for the histologic type of the tumor being reported using ICD-O-3. NAACCR adopted ICD-O-3 as the standard coding system for tumors diagnosed in 2001 and later, and recommended that prior tumors be converted from ICD-O-2.

Note: See Histology (92-00) ICD-O-2 [420] for ICD-O-2 codes.

# Codes

See ICD-O-3<sup>14</sup>, Morphology Section.

# **Clarification of Required Status**

This data item is required by all standard-setting organizations for tumors diagnosed on or after January 1, 2001, and recommended (by conversion from ICD-O-2 codes when conversion algorithms and tables are available) for tumors diagnosed before 2001.

When the histologic type is coded according to ICD-O-3, the histology code must be reported in Histologic Type ICD-O-3 [522], with behavior coded in Behavior Code ICD-O-3 [523].

For information on required status for related data items for histologic type and behavior when coded according to ICD-O-2, see Histology (92-00) ICD-O-2 [420] and Behavior (92-00) ICD-O-2 [430].

#### HISTOLOGY (73-91) ICD-O-1

Alternate Name	Item #	Length	Source of Standard	Column #
	1971	4	SEER	1141-1144

#### Description

Area for retaining the histology portion (4 digits) of the ICD-O-1 or field trial morphology codes entered before a conversion to ICD-O-2. See grouped data item Morph (73-91) ICD-O-1 [1970], in Appendix E. The item name includes years 1973-91. However, some states may have used the codes for cases before 1973.

# Codes

For cases diagnosed before 1992, contains the ICD-O-1 or field trial 4-digit histology code as originally coded, if available. Blank for tumors coded directly into ICD-O-2 (i.e., 1992 and later cases).

#### HISTOLOGY (92-00) ICD-O-2

Alternate Name	Item #	Length	Source of Standard	Column #
Histology (COC)	420	4	SEER/COC	296-299

#### Description

Codes for the histologic type of the tumor being reported using ICD-O-2. NAACCR adopted ICD-O-2 as the standard coding system for tumors diagnosed in 1992 and later and recommended that prior cases be converted to ICD-O-2.

Note: See Histology (73-91) ICD-O-1 [1971] for ICD-O-1 and field trial codes.

#### Codes

See ICD- O -2,<sup>15</sup> Morphology Section.

#### **Clarification of Required Status**

This data item is required by all standard-setting organizations for tumors diagnosed from January 1, 1992 through December 31, 2000, and recommended for tumors diagnosed before 1992.

When the histologic type is coded according to ICD-O-2, the histology code must be reported in Histology (92-00) ICD-O-2 [420], with behavior coded in Behavior (92-00) ICD-O-2 [430].

For information on required status for related data items for histologic type and behavior when coded according to ICD-O-3, see Histologic Type ICD-O-3 [522] and Behavior Code ICD-O-3 [523].

#### ICD REVISION NUMBER

ICD REVISION NUMBER				
Alternate Name	Item #	Length	Source of Standard	Column #
ICD Code Revision Used for Cause of	1920	1	SEER	1392-1392
Death (SEER)				

#### Description

Indicator for the coding scheme used to code the cause of death.

- 0 Patient alive at last follow-up
- 1 **ICD-10**
- ICD-7 7
- 8 ICDA-8
- 9 ICD-9

# **ICD-O-2 CONVERSION FLAG**

Alternate Name	Item #	Length	Source of Standard	Column #
Review Flag for 1973-91 Cases (SEER)	1980	1	SEER	1147-1147

# Description

Code specifying how the conversion of site and morphology codes from ICD-O-1 and the field trial editions to ICD-O-2 was accomplished. The item names include years 1973-91. However, some states may have used the codes for tumors before 1973. The code also covers morphology conversions from ICD-O-3 to ICD-O-2.

# Codes

- 0 Primary site and morphology originally coded in ICD-O-2
- 1 Primary site and morphology converted without review
- 2 Primary site converted with review; morphology machine-converted without review
- 3 Primary site machine-converted without review, morphology converted with review
- 4 Primary site and morphology converted with review
- 5 Morphology converted from ICD-O-3 without review
- 6 Morphology converted from ICD-O-3 with review
- Blank Not converted

# **ICD-O-3 CONVERSION FLAG**

Alternate Name	Item #	Length	Source of Standard	Column #
	2116	1	SEER/COC	1243-1243

#### Description

Code specifying how the conversion of site and morphology codes from ICD-O-2 to ICD-O-3 was accomplished.

- 0 Morphology (Morph--Type&Behav ICD-O-3 [521] originally coded in ICD-O-3
- 1 Morphology (Morph--Type&Behav ICD-O-3 [521] converted from (Morph--Type&Behav ICD-O-2 [419] without review
- 3 Morphology (Morph--Type&Behav ICD-O-3 [521] converted from (Morph--Type&Behav ICD-O-2 [419] with review
- Blank Not converted

#### INDUSTRY CODE--CENSUS

Alternate Name	Item #	Length	Source of Standard	Column #
	280	3	Census/NPCR	138-140

#### Description

Code for the patient's usual industry, using U.S. Census Bureau codes (2000 Census<sup>26</sup> is preferable) according to coding procedures recommended for death certificates.<sup>25</sup> This data item applies only to patients who are age 14 years or older at the time of diagnosis.

*Note:* Occupation/industry coding should NOT be performed by reporting facilities. This is a central cancer registry data item. Specially trained and qualified personnel should perform coding.

*Note:* 2000 Census codes for occupation and industry are recommended for tumors diagnosed on or after January 1, 2003.<sup>26</sup> The 1990 Census codes are recommended for tumors diagnosed before January 1, 2003.<sup>24</sup> For more information, see the U.S. Census Bureau Web Site at: http://www.census.gov/hhes/www/ioindex/ overview.html.

#### Rationale

Use of the Census Bureau classification system improves consistency of data collected from multiple sources. The Census Bureau industrial classification system is used for coding industry information from death certificates and from the U.S. Census of Population. The system includes specific coding rules.<sup>22-27</sup>

#### Codes

For the 1990 Census codes see Instructional Manual Part 19: *Industry and Occupation Coding for Death Certificates*, 1999<sup>23</sup> and related materials in the reference list, Chapter VII. A similar instruction manual for the 2000 Census codes has not been developed. Software for automated coding of occupation and industry is available from the Division of Safety Research, National Institute for Occupational Safety and Health, CDC. Contact Suzanne Marsh at (304) 285-6009 or at smm2@cdc.gov.

# **INDUSTRY SOURCE**

Alternate Name	Item #	Length	Source of Standard	Column #
	300	1	NPCR	142-142

#### Description

Code that best describes the source of industry information provided on this patient. This is a central cancer registry data item (i.e., codes should be applied by a central or regional registry rather than collected from reporting facilities).

# Rationale

Industry information may come from a variety of sources. The most valid and reliable source of industry information for patients has not yet been determined.

#### Codes

- 0 Unknown industry/no industry available
- 1 Reporting facility records
- 2 Death certificate
- 3 Interview
- 7 Other source
- 8 Not applicable, patient less than 14 years of age at diagnosis
- 9 Unknown source
- Blank Not collected

#### **INPATIENT/OUTPT STATUS**

Alternate Name	Item #	Length	Source of Standard	Column #
Inpatient/Outpatient Status (COC)	640	1	COC	447-447

#### Description

Access point from which the patient first entered the hospital system for either the initial diagnosis or treatment.

#### Codes

- 1 Inpatient only
- 2 Outpatient only
- 3 In- and outpatient\*
- 8 Other, including physician's office
- 9 Unknown

\**Note:* This applies to patients who entered the institution as outpatients and were admitted as inpatients on the same day as well as on different dates.

Note: This data item is no longer supported by COC (as of January 1, 2003).

INSTITUTION REFERRED FROM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Facility Referred From	2410	10	COC	2485-2494

Identifies the facility that referred the patient to the reporting hospital.

#### Rationale

Each facility's FIN is unique. This number is used to document and monitor referral patterns. Efforts are underway to establish uniform national provider ID numbers. COC and NAACCR committees will consider adoption of any federal standards, when they become available.

#### **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001: Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10-digit codes.

#### Codes (in addition to COC assigned codes)

000000000Case not referred from a facility00999999999Case referred from a facility, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN Coding System [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

INSTITUTION REFERRED TO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Facility Referred To	2420	10	COC	2495-2504

#### Description

Identifies the facility to which the patient was referred for further care after discharge from the reporting facility.

#### Rationale

Each facility's FIN is unique. This number is used to document and monitor referral patterns. Efforts are underway to establish uniform national provider ID numbers. COC and NAACCR committees will consider adoption of any federal standards, when they become available.

#### **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001: Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10-digit codes.

# Codes (in addition to COC assigned codes)

000000000	Case not referred to a facility
0099999999	Case referred to a facility, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN Coding System [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

# LAST FOLLOW-UP HOSPITAL

Alternate Name	Item #	Length	Source of Standard	Column #
	2430	10	NAACCR	2465-2474

# Description

Records facility where the patient was last followed.

#### Rationale

Each facility's FIN is unique. The number is essential to NCDB for monitoring data submissions, ensuring the accuracy of data, and identifying areas for special studies.

Efforts are underway at the federal level to establish uniform national provider ID numbers. COC and NAACCR committees will consider adoption of any federal standards, when they become available.

#### **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001: Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10-digit codes.

# Codes (in addition to COC assigned codes)

000000000	Case not reported by a facility
0099999999	Case reported, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN Coding System [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

# LATERALITY

Alternate Name	Item #	Length	Source of Standard	Column #
Laterality at Diagnosis (SEER)	410	1	SEER/COC	295-295

# Description

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

- 0 Not a paired site
- 1 Right: origin of primary
- 2 Left: origin of primary
- 3 Only one side involved, right or left origin unspecified
- 4 Bilateral involvement, lateral origin unknown; stated to be single primary; including both ovaries involved simultaneously, single histology; bilateral retinoblastomas; bilateral Wilms' tumors
- 9 Paired site, but no information concerning laterality, midline tumor

#### LATITUDE

Alternate Name	Item #	Length	Source of Standard	Column #
	2352	10	NAACCR	2394-2403

# Description

Cancer Registry spatial data for a tumor record represents the point location of the individual's residence on the Earth's surface. The point location is expressed as a coordinate pair of latitude and longitude values determined by any one of several methods: for example, geocoding, address matching, global positioning satellite (GPS) readings, and interpolation from paper or electronic maps. Most of the time this information is provided by a geocoding vendor service. Alternatively, a central registry staff manually assigns the code. This item is not coded by the hospital.

# Rationale

Decimal degree coordinate data can be thought of as the universal "currency" of exchange for spatial data to be used (projected or not projected) in GIS. Data in this format can be used by any GIS software and projected for the appropriate area of interest, and would be consistent with formats of data obtained from other sources. Users may not necessarily need to project their data unless they need to preserve properties of area, shape, distance, or direction. Different projections provide one or more of these properties. Some projections are used simply for presentation purposes because they make the map "look" better. Displaying a large area such as a state or province/territory using an unprojected rectangular latitude/longitude decimal degree grid may make the area appear distorted, especially in far northern latitudes.

#### Allowable values and format

Projection and Units -- Spatial data will be exchanged in "unprojected" latitude and longitude coordinates. The data units will be in decimal degrees (and not in degrees, minutes, seconds).

Correct:	Latitude: 41.890949 Longitude: -123.128943
Not this:	Latitude: 41 deg 53' 27" Longitude: -71 deg 7' 44"

The latitude field is a 10-byte numeric field, right justified. This coordinate may be carried out to 6 decimal places with an explicit decimal point. It has the following format: x12.345678, where "x" is reserved for a negative sign if the coordinate represents a location south of the equator.

#### Codes

Latitude and longitude data shall always be stored and exchanged as numeric values. Latitude north of the equator is positive. Longitude west of 0 degrees (the Prime Meridian) and east of 180 degrees (approximately the International Date Line) is negative—this applies to the entire North American continent with the exception of the tip of the Aleutian Islands in Alaska.

*Note:* The datum of the decimal degree data shall be North American Datum of 1983 (NAD 83). Data in NAD 27 shall be converted to NAD 83 prior to data exchange.

## LOC/REG/DISTANT STAGE

Alternate Name	Item #	Length	Source of Standard	Column #
	770	1	Varies	530-530

# Description

For use if no other staging is available. Use may not be consistent among registries.

Note: This is not the same as SEER historic stage. See the Comparative Staging Guide for Cancer.

- 0 In situ
- 1 Local
- 2 Regional
- 3 Distant
- 9 Unstaged

# LONGITUDE

Alternate Name	Item #	Length	Source of Standard	Column #
	2354	11	NAACCR	2404-2414

## Description

Cancer Registry spatial data for a tumor record represents the point location of the individual's residence on the Earth's surface. The point location is expressed as a coordinate pair of latitude and longitude values determined by any one of several methods: for example, geocoding, address matching, GPS readings, and interpolation from paper or electronic maps. Most of the time this information is provided by a geocoding vendor service. Alternatively, a central registry staff manually assigns the code. This item is not coded by the hospital.

## Rationale

Decimal degree coordinate data can be thought of as the universal "currency" of exchange for spatial data to be used (projected or not projected) in GIS. Data in this format can be used by any GIS software and projected for the appropriate area of interest, and would be consistent with formats of data obtained from other sources. Users may not necessarily need to project their data unless they need to preserve properties of area, shape, distance, or direction. Different projections provide one or more of these properties. Some projections are used simply for presentation purposes because they make the map "look" better. Displaying a large area such as a state or province/territory using an unprojected rectangular latitude/longitude decimal degree grid may make the area appear distorted, especially in far northern latitudes.

## Allowable values and format

Projection and Units -- Spatial data are exchanged in "unprojected" latitude and longitude coordinates. The data units are in decimal degrees (and not in degrees, minutes, seconds).

Correct:	Latitude: 41.890949 Longitude: -123.128943
Not this:	Latitude: 41 deg 53' 27" Longitude: -71 deg 7' 44"

The longitude field is an 11-byte numeric field, right justified. This coordinate may be carried out to 6 decimal places with an explicit decimal point. It has the following format: x123.456789, where "x" is reserved for a negative sign if the coordinate represents a location west of 0 degrees (Prime Meridian) and east of 180 degrees.

# Codes

Latitude and longitude data are stored and exchanged as numeric values. Latitude north of the equator is positive. Longitude west of 0 degrees (the Prime Meridian) and east of 180 degrees (approximately the International Date Line) is negative—this applies to the entire North American continent with the exception of the tip of the Aleutian Islands in Alaska.

*Note:* The datum of the decimal degree data is NAD 83. Data in NAD 27 are converted to NAD 83 prior to data exchange.

# MARITAL STATUS AT DX

Alternate Name	Item #	Length	Source of Standard	Column #
Marital Status at Diagnosis (SEER/COC)	150	1	SEER	102-102
Marital Status at Initial Diagnosis (pre-96				
COC)				

## Description

Code for the patient's marital status at the time of diagnosis for the reportable tumor. If the patient has multiple tumors, marital status may be different for each tumor.

## Rationale

Incidence and survival with certain cancers vary by marital status. The item also helps in patient identification.

## Codes

- 1 Single (never married)
- 2 Married (including common law)
- 3 Separated
- 4 Divorced
- 5 Widowed
- 9 Unknown

MEDICAL RECORD NUMBER				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2300	11	COC	2086-2096

## Description

Records medical record number used by the facility to identify the patient. The COC *FORDS* Manual instructs registrars to record numbers assigned by the facility's Health Information Management (HIM) Department only, not department-specific numbers.

## Rationale

This number identifies the patient in a facility. It can be used by a central registry to point back to the patient record, and it helps identify multiple reports on the same patient.

## Codes (in addition to the medical record number)

UNK Medical record number unknown

- RT Radiation therapy department patient without HIM number
- SU 1-day surgery clinic patient without HIM number

*Note:* Other standard abbreviations may be used to indicate departments within the facility for patients without HIM numbers assigned.

## MILITARY RECORD NO SUFFIX

Alternate Name	Item #	Length	Source of Standard	Column #
Military Medical Record Number Suffix	2310	2	COC	2097-2098
(COC)				

## Description

Patient identifier used by military hospitals to record relationship of the patient to the sponsor.

Codes	
01-19	Child
20	Sponsor
30-39	Spouse
40-44	Mother
45-49	Father
50-54	Mother-in-law
55-59	Father-in-law
60-69	Other eligible dependents
98	Civilian emergency (Air Force/Navy)
99	Not classified elsewhere/stillborn
Blank	Not applicable, medical record number not from a military hospital

## MORPH (73-91) ICD-O-1

Alternate Name	Item #	Length	Source of Standard	Column #
	1970	6		1141-1146

## Description

The name for a group of subfields describing the type and behavior of the tumor being reported using ICD-O-1 codes.

Group names appear only in the data dictionary and Appendix E.

## Subfields

Histology (73-91) ICD-O-1 [1971] Behavior (73-91) ICD-O-1 [1972] Grade (73-91) ICD-O-1 [1973]

## MORPH CODING SYS--CURRENT

Alternate Name	Item #	Length	Source of Standard	Column #
	470	1	NAACCR	309-309

## Description

Code that best describes how morphology is currently coded. If converted, this field shows the system it is converted to.

## Codes

- 1 ICD-O, First Edition
- 2 ICD-O, 1986 Field Trial
- 3 ICD-O, 1988 Field Trial
- 4 ICD-O, Second Edition
- 5 ICD-O, Second Edition, plus REAL lymphoma codes effective 1/1/95
- 6 ICD-O, Second Edition, plus FAB codes effective 1/1/98
- 7 ICD-O, Third Edition
- 9 Other

## **MORPH CODING SYS--ORIGINL**

Alternate Name	Item #	Length	Source of Standard	Column #
	480	1	NAACCR	310-310

#### Description

Code that best describes how morphology was originally coded. If later converted, this field shows the original codes used.

- 1 ICD-O, First Edition
- 2 ICD-O, 1986 Field Trial
- 3 ICD-O, 1988 Field Trial
- 4 ICD-O, Second Edition
- 5 ICD-O, Second Edition, plus REAL lymphoma codes effective 1/1/95
- 6 ICD-O, Second Edition, plus FAB codes effective 1/1/98
- 7 ICD-O-3, Third Edition
- 9 Other

## MORPH--TYPE&BEHAV ICD-O-2

Alternate Name	Item #	Length	Source of Standard	Column #
	419	5		296-300

## Description

The name for a group of subfields describing the type and behavior of the tumor being reported using ICD-O-2 codes.

Group names appear only in the data dictionary and Appendix E.

## Subfields

Histology (92-00) ICD-O-2 [420] Behavior (92-00) ICD-O-2 [430]

## MORPH--TYPE&BEHAV ICD-O-3

Alternate Name	Item #	Length	Source of Standard	Column #
	521	5		301-305

## Description

The name for a group of subfields describing the type and behavior of the tumor being reported using ICD-O-3 codes.

Group names appear only in the data dictionary and Appendix E.

## Subfields

Histologic Type ICD-O-3 [522] Behavior Code ICD-O-3 [523]

#### NAACCR RECORD VERSION

NAACCR RECORD VERSION				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	50	1	NAACCR	19-19

#### Description

This item applies only to record types I, C, A and M. Code the NAACCR record version used to create the record.

*Note:* The correction record (U) has its own record version data item.

#### Codes

- 1 1992-1994 Version (Version 2 and Version 3)
- 4 1995 Version (Version 4.0)
- 5 1996 and 1997 Version (Version 5.0 or Version 5.1)
- 1998 Version (Version 6) 6
- 1999 Version (Version 7) 7
- 8 2000 Version (Version 8)
- 9 2001 and 2002 Version (Version 9 and 9.1)
- А 2003, 2004, and 2005 Version (Version 10, 10.1, and 10.2)
- Blank September 1989 Version

Note: Code 4 was assigned to the 1995 Version to synchronize the document version and the layout version numbers. Layout document Versions 2 and 3 are coded as 1.

NAMEALIAS				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Alias (COC)	2280	15	SEER	2006-2020

#### Description

Records an alternate name or "AKA" (also known as) used by the patient, if known. Note that maiden name is entered in item 2390.

#### NAME--FIRST

Alternate Name	Item #	Length	Source of Standard	Column #
First Name (COC)	2240	14	NAACCR	1972-1985

## Description

First name of the patient.

Note: The COC FORDS Manual allows this field to be blank. If facilities with COC-approved cancer programs submit blanks to the central registry, it is suggested that the central registry devise procedures for completing the last and first name with text, such as UNKNOWN, after verifying with the hospital that the field was left intentionally blank.

## NAME--LAST

Alternate Name	Item #	Length	Source of Standard	Column #
Last Name (COC)	2230	25	NAACCR	1947-1971

## Description

Last name of the patient.

*Note:* From *FORDS* Edits: Last Name is required. The last name of the patient must be entered left justified with trailing blanks. Mixed case is allowed. Spaces, hyphens, and apostrophes are allowed. The field may not be completely blank. If the last name is unknown, enter "Unknown."

NAMEMAIDEN				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Maiden Name (COC)	2390	15	SEER	2021-2035

## Description

Maiden name of female patients who are or have been married.

#### Rationale

This is used to link reports on a woman who changed her name between reports. It also is critical when using Spanish surname algorithms to categorize ethnicity.

Note: See Chapter V, Unresolved Issues, for discussion of hyphenated maiden name.

#### NAME--MIDDLE

Alternate Name	Item #	Length	Source of Standard	Column #
Middle Name (COC)	2250	14	COC	1986-1999
Middle Initial (pre-96 COC)				

#### Description

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

NAMEPREFIX				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Name Prefix (COC)	2260	3	SEER	2000-2002

## Description

Abbreviated title that precedes name in a letter (e.g., "Rev.," "Ms.").

#### NAME--SPOUSE/PARENT

NAMESPOUSE/PARENT				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2290	50	NAACCR	2036-2085

#### Description

NAACCR has not adopted standards for this item. Use varies by area.

#### NAME SUFFIX

NAMESUFFIX				Kevised
Alternate Name	Item #	Length	Source of Standard	Column #
Name Suffix (COC)	2270	3	SEER	2003-2005

#### Description

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

#### **NEXT FOLLOW-UP SOURCE**

Alternate Name	Item #	Length	Source of Standard	Column #
Next Follow-Up Method (pre-96 COC)	1800	1	COC	1306-1306

#### Description

Identifies the method planned for the next follow-up.

#### Codes

- Chart requisition 0
- Physician letter 1
- 2 Contact letter
- 3 Phone call
- Other hospital contact 4
- 5 Other, NOS
- 8 Foreign residents (not followed)
- 9 Not followed, other cases for which follow-up is not required

Derrad

NHIA DERIVED HISP ORIGIN				New
Alternate Name	Item #	Length	Source of Standard	Column #
	191	1	NAACCR	231-231

The NAACCR Hispanic Identification Algorithm (NHIA) uses a combination of NAACCR variables to directly or indirectly classify cases as Hispanic for analytic purposes. It is possible to separate Hispanic ancestral subgroups (e.g., Mexican) when indirect assignment results from birthplace information but not from surname match. The algorithm uses the following NAACCR standard variables: Spanish/Hispanic Origin [190], Name--Last [2230], Name--Maiden [2390], Birthplace [250], Race 1 [160], and Sex [220].

Code 7 (Spanish surname only) of the Spanish/Hispanic Origin [190] data item became effective with 1994 diagnosis. It is recommended that NHIA should be run on 1995 and forward diagnoses. However, a central registry may run it on their data for prior years.

## Rationale

Sometimes, despite best efforts to obtain complete information directly from the medical record, information is not available and is reported to the cancer registry as a missing data item. With regard to Hispanic ethnicity, some cancer registries have found it necessary to rely on indirect methods to populate this data element. The registries often have significant numbers or proportions of Hispanic populations in their jurisdiction.

- 0 Non-Hispanic
- 1 Mexican, by birthplace or other specific identifier
- 2 Puerto Rican, by birthplace or other specific identifier
- 3 Cuban, by birthplace or other specific identifier
- 4 South or Central American (except Brazil), by birthplace or other specific identifier
- 5 Other specified Spanish/Hispanic origin (includes European; excludes Dominican Republic), by birthplace or other specific identifier
- 6 Spanish, NOS; Hispanic, NOS; Latino, NOS
- 7 NHIA surname match only
- 8 Dominican Republic
- Blank Algorithm has not been run

# **OCCUP/IND CODING SYSTEM**

Alternate Name	Item #	Length	Source of Standard	Column #
	330	1	NPCR	223-223

# Description

Code that identifies coding system used for occupation and industry. This is a central cancer registry data item (i.e., codes should be applied by a central or regional registry rather than collected from reporting facilities).

# Codes

- 1 1970 Census
- 2 1980 Census
- 3 1990 Census
- 4 2000 Census
- 7 Other coding system
- 9 Unknown coding system

# Blank Not collected

*Note:* 2000 Census codes for occupation and industry are recommended for tumors diagnosed on or after January 1, 2003.<sup>26</sup> The 1990 Census codes are recommended for tumors diagnosed before January 1, 2003.<sup>24</sup> For more information, see the U.S. Bureau of the Census Web Site at: http://www.census.gov/hhes/www/ioindex /overview.html.

## **OCCUPATION CODE--CENSUS**

Alternate Name	Item #	Length	Source of Standard	Column #
	270	3	Census/NPCR	135-137

## Description

Code for the patient's usual occupation, using U.S. Census Bureau codes (2000 Census<sup>26</sup> is preferable) according to coding procedures recommended for death certificates.<sup>25</sup> This data item applies only to patients who are age 14 years or older at the time of diagnosis.

*Note:* Occupation/industry coding should NOT be performed by reporting facilities. This is a central registry data item. Specially trained and qualified personnel should perform coding.

*Note:* 2000 Census codes for occupation and industry are recommended for cancers diagnosed on or after January 1, 2003.<sup>26</sup> The 1990 Census codes are recommended for cancers diagnosed before January 1, 2003.<sup>24</sup> For more information, see the U.S. Bureau of the Census Web Site at: http://www.census.gov/hhes/www/ioindex/overview.html.

## Rationale

Use of the Census Bureau classification system improves consistency of data collected from multiple sources. The Census Bureau occupation classification system is used for coding occupation information from death certificates and from the U.S. Census of Population. The system includes specific coding rules.<sup>22-27</sup>

#### Codes

For the 1990 Census codes, see Instructional Manual Part 19: *Industry and Occupation Coding for Death Certificates*, 1999,<sup>23</sup> and related materials in the reference list, Chapter VII. A similar instruction manual for the 2000 Census codes has not been developed. Software for automated coding of occupation and industry is available from the Division of Safety Research, National Institute for Occupational Safety and Health, CDC. Contact Suzanne Marsh at (304) 285-6009 or at smm2@cdc.gov.

## **OCCUPATION SOURCE**

Alternate Name	Item #	Length	Source of Standard	Column #
	290	1	NPCR	141-141

## Description

Code that best describes the source of occupation information provided on this patient. This is a central cancer registry data item (i.e., codes should be applied by a central or regional registry rather than collected from reporting facilities).

## Rationale

Occupation information may come from a variety of sources. The most valid and reliable source of occupation information for patients has not yet been determined.

## Codes

- 0 Unknown occupation/no occupation available
- 1 Reporting facility records
- 2 Death certificate
- 3 Interview
- 7 Other source
- 8 Not applicable, patient less than 14 years of age at diagnosis
- 9 Unknown source
- Blank Not collected

# **OTHER STAGING SYSTEM**

Alternate Name	Item #	Length	Source of Standard	Column #
	1070	15	COC	595-609

## Description

Field for collecting additional staging classifications (e.g., Dukes, AUA). Text field. User defined.

Note: This data item is no longer supported by COC (as of January 1, 2003).

OVER-RIDE ACSN/CLASS/SEQ				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Accession/Class of	1985	1	COC	1119-1119
Case/Sequence				

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edit in the NAACCR Metafile of the EDITS software: Accession Number, Class of Case, Seq Number (COC).

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# Over-ride Flag as Used in the EDITS Software Package

The edit, Accession Number, Class of Case, Seq Number (COC), checks the following:

- 1. If the case is the only case or the first of multiple cases diagnosed at the facility (Sequence Number-Hospital = 00, 01, 60, or 61, and Class of Case = 0, 1, or 6), then the first 4 characters of the Accession Number must equal the year of the Date of 1st Contact.
- 2. If the case is first diagnosed at autopsy (Class of Case = 5) and the case is the only case or the first of multiple cases for a patient (Sequence Number--Hospital = 00, 01, 60, or 61), then the first 4 characters of the Accession Number must equal the year of the Date of Last Contact AND must equal the year of the Date of 1st Contact.
- 3. If the case is first diagnosed at autopsy (Class of Case = 5) and the case is not the first case for a patient (Sequence Number--Hospital) greater than 01 or greater than 61), then the year of the Date of 1st Contact must equal the year of Date of Last Contact.

There are some exceptions to the above rules. Over-ride Acsn/Class/Seq may be used to override the edit when the circumstances fit the following situation or one similar to it:

1. The case may be the only or the first of multiple malignant cases for a patient (Sequence Number-Hospital = 00 or 01), but there is an earlier benign case (with an earlier year of the Date of 1st Contact) to which the Accession Number applies.

# **Instructions for Coding**

- 1. If edit generates an error or warning message, verify that the Accession Number, Sequence Number, and Class of Case are correct.
- 2. Leave blank if the program does not generate an error message for the edit Accession Number, Class of Case, Seq Number (COC).
- 3. Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- 4. Code 1 if review of accession number, sequence number and class of case verifies that they have been coded correctly and there is an unusual combination of these data items.

## Codes

1 Reviewed Blank Not reviewed or reviewed and corrected

## **OVER-RIDE AGE/SITE/MORPH**

<b>OVER-RIDE AGE/SITE/MORPH</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Age/Site/Histology Interfield Review	1990	1	SEER	1124-1124
(Interfield Edit 15) (SEER #3)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Age, Primary Site, Morphology (SEER IF15)

Age, Primary Site, Morph ICDO3--Adult (SEER)

Age, Primary Site, Morph ICDO3--Pediatric (NPCR)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## **Over-ride Flag as Used in the EDITS Software Package**

Some cancers occur almost exclusively in certain age groups.

Edits of the type Age, Primary Site, Morphology require review if a site/morphology combination occurs in an age group for which it is extremely rare. The edit Age, Primary Site, Morph ICDO3--Adult (SEER) edits cases with an Age at Diagnosis of 15 and older. The edit Age, Primary Site, Morph ICDO3--Pediatric (NPCR) edits cases with an Age at Diagnosis of less than 15. The ICD-O-2 version of the edit is Age, Primary Site, Morphology (SEER IF15) and contains logic for all ages.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Age, Primary Site, 1. Morphology, the edit Age, Primary Site, Morph ICDO3--Adult, or the edit Age, Primary Site, Morph ICDO3--Pediatric.
- 2. Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 as indicated if review of items in the error or warning message confirms that all are correct. 3.

- 1 Reviewed: An unusual occurrence of a particular age/site/histology combination for a given age group has been reviewed.
- Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE COC-SITE/TYPE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1987	1	COC	1121-1121

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Primary Site, Morphology-Type Check (COC)

Primary Site, Morphology-Type ICDO3 (COC)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## **Over-ride Flag as Used in the EDITS Software Package**

Multiple versions of edits of the type Primary Site, Morphology-Type check for "usual" combinations of site and ICD-O-2 or ICD-O-3 histology. The SEER version of the edit is more restrictive than the COC edit, and thus, uses a different over-ride flag. The COC version of the edit will accept Over-ride COC Site/Type or Over-ride Site/Type (the SEER edit) as equivalent.

1. The Site/Histology validation list (available on the SEER Web Site) contains those histologies commonly found in the specified primary site. Histologies that occur only rarely or never are not included. These edits require review of all combinations not listed.

Review of these cases requires investigating whether the combination is biologically plausible or whether cancer registry coding conventions would allow different codes for the diagnosis. Review of these rare combinations often results in a change to either the site or histology.

## **Instructions for Coding**

- 1. Leave blank if the program does not generate an error message for edits of the type Primary Site, Morphology-Type Check.
- 2. Leave blank and correct any errors for the case if an item is discovered to be incorrect.

Code 1 if review of all items in the error or warning message confirms they are correct and coded in conformance with coding rules.

- 1 Reviewed
- Blank Not reviewed or reviewed and corrected

## **OVER-RIDE HISTOLOGY**

<b>OVER-RIDE HISTOLOGY</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Histology/Behavior Interfield Review	2040	1	SEER	1129-1129
(Field Item Edit Morph) (SEER #2)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Diagnostic Confirmation, Behavior Code (COC) Diagnostic Confirmation, Behavior Code (SEER IF31) Diagnostic Confirmation, Behavior ICDO3 (COC) Diagnostic Confirmation, Behavior ICDO3 (SEER IF31) Morph (1973-91) ICD-O-1 (SEER MORPH) Morphology--Type/Behavior (COC) Morphology--Type/Behavior (SEER MORPH) Morphology--Type/Behavior ICDO3 (COC) Morphology--Type/Behavior ICDO3 (SEER MORPH)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## **Over-ride Flags as Used in the EDITS Software Package**

Edits of the type Diagnostic Confirmation, Behavior Code differ in the use of ICD-O-2 or ICD-O-3 and check that, for *in situ* cases (Behavior = 2), Diagnostic Confirmation specifies microscopic confirmation (1, 2, or 4).

The distinction between *in situ* and invasive is very important to a registry, because prognosis is so different. Because the determination that a neoplasm has not invaded surrounding tissues (i.e., in situ, is made microscopically), cases coded *in situ* in behavior should have a microscopic confirmation code. However, very rarely, a physician will designate a case noninvasive or *in situ* without microscopic evidence.

- 1. If an edit of the type, Diagnostic Confirmation, Behavior Code, gives an error message or warning, check that Behavior and Diagnostic Confirmation have been coded correctly. Check carefully for any cytologic or histologic evidence that may have been missed in coding.
- The following histologies are not accepted as *in situ*: 8000-8005, 8020, 8021, 8331, 8320, 8800-9055, 2. 9062, 9082, 9083, 9110-9493, 9501-9989.

Edits of the type, Morphology--Type/Behavior, perform the following check:

- Codes listed in ICD-O-2 or ICD-O-3 with behavior codes of only 0 or 1 are considered valid, since 1. the behavior matrix of ICD-O-2 and ICD-O-3 allows for the elevation of the behavior of such histologies when the tumor is *in situ* or malignant. This edit forces review of these rare cases to verify that they are indeed in situ or malignant.
- 2. If a Morphology-Type/Behavior edit produces an error or warning message and the case is one in which the 4-digit morphology code is one that appears in ICD-O-2 or ICD-O-3 only with behavior

codes of 0 or 1, verify the coding of morphology and that the behavior should be coded malignant or *in situ*. The registrar may need to consult a pathologist or medical advisor in problem cases.

## Exceptions:

If year of Date of Diagnosis > 2000, then a behavior code of 1 is valid for the following ICD-O-2 histologies and no over-ride flag is needed: 8931, 9393, 9538, 9950, 9960-9962, 9980-9984, and 9989. Similarly, the following ICD-O-3 histologies are valid with a behavior code of 1: 8442, 8451, 8462, 8472, and 8473.

If year of Date of Diagnosis > 2003, the following ICD-O-3 benign histologies will pass without review: 8146, 8271, 8861, 8897, 9121, 9122, 9131, 9161, 9350, 9351, 9352, 9360, 9361, 9383, 9384, 9394, 9412, 9413, 9444, 9492, 9493, 9506, 9531, 9532, 9533, 9534, 9537, 9541, 9550, 9562, and 9570.

3. Grade 5-8 with histologies not in the rage of 9590-9948 is impossible.

## **Instructions for Coding**

- 1. Leave blank if the program does not generate an error message for the edits of the types, Diagnostic Confirmation, Behav Code or Morphology--Type/Behavior.
- 2. Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- 3. Code 1, 2, or 3 as indicated if review of all items (Diagnostic Confirmation, Behavior, and Morphology) in the error or warning message confirms that all are correct.

- 1 Reviewed: The behavior code of the histology is designated as "benign" or "uncertain" in ICD-O-2 or ICD-O-3, and the pathologist states the primary to be "*in situ*" or "malignant" (flag for a "Morphology Type & Behavior" edit)
- 2 Reviewed: The behavior code is "*in situ*," but the case is not microscopically confirmed (flag for a "Diagnostic Confirmation, Behavior Code" edit)
- 3 Reviewed: Conditions 1 and 2 above both apply
- Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE HOSPSEQ/DXCONF</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Hospital Sequence/Diagnostic	1986	1	COC	1120-1120
Confirmation				

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edit in the NAACCR Metafile of the EDITS software: Diagnostic Confirm, Seq Num--Hosp (COC)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# Over-ride Flag as Used in the EDITS Software Package

The edit, Diagnostic Confirm, Seq Num--Hosp (CoC), does the following:

- 1. If any case is one of multiple primaries and is not microscopically confirmed or lacks a positive laboratory test/marker study (i.e., Diagnostic Confirmation > 5 and Sequence Number--Hospital > 00 [more than one primary]), review is required.
- 2. If Primary Site specifies an ill-defined or unknown primary (C760-C768, C809), no further checking is done.
- 3. If Sequence Number--Hospital is in the range of 60-88, this edit is skipped.

It is important to verify that the non-microscopically confirmed case is indeed a separate primary from any others that may have been reported. This edit forces review of multiple primary cancers when one of the primaries is coded to a site other than ill-defined or unknown and is not microscopically confirmed or confirmed by a positive laboratory test/marker study.

- 1. If the suspect case is confirmed accurate as coded and if the number of primaries is correct, set the Over-ride HospSeq/DxConf to 1. Do not set the over-ride flag on the patient's other primary cancers.
- 2. If it turns out that the non-microscopically confirmed cancer is considered a manifestation of one of the patient's other cancers, delete the non-microscopically confirmed case. Check the sequence numbers of remaining cases, correcting them if necessary. Also check for other data items on the remaining cases that may need to be changed as a result of the corrections, such as stage and treatment.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Diagnostic Confirm, Seq Num--Hosp (COC).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of all items in the error or warning message confirms that all are correct.

# Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE HOSPSEQ/SITE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Hospital Sequence/Site	1988	1	COC	1122-1122

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Seq Num--Hosp, Primary Site, Morph (COC)

Seq Num--Hosp, Primary Site, Morph ICDO3 (COC)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

Edits of the type Seq Num--Hosp, Primary Site, Morph differ in use of ICD-O-2 or ICD-O-3 morphology. They force review of multiple primary cancers when one of the primaries is coded to a site/morphology combination that could indicate a metastic site rather than a primary site.

- 1. If Sequence Number--Hospital indicates the person has had more than one primary, then any case with one of the following site/histology combinations requires review:
  - C760-C768 (ill-defined sites) or C809 (unknown primary) and ICD-O-2 or ICD-O-3 histology < 9590. Look for evidence that the unknown or ill-defined primary is a secondary site from one of the patient's other cancers. For example, a clinical discharge diagnosis of "abdominal carcinomatosis" may be attributable to the patient's primary ovarian cystadenocarcinoma already in the registry, and should not be entered as a second primary.
  - C770-C779 (lymph nodes) and ICD-O-2 histology not in range 9590-9717 or ICD-O-3 histology not in the range 9590-9729; or C420-C424 and ICD-O-2 histology not in range 9590-9941 or ICD-O-3 histology not in the range 9590-9989. That combination is most likely a metastatic lesion. Check whether the lesion could be a manifestation of one of the patient's other cancers.
  - Any site and ICD-O-2 histology in the range 9720-9723, 9740-9741 or ICD-O-3 histology in the range 9740-9758. Verify that these diagnoses are coded correctly and are indeed separate primaries form the others.
- 2. If it turns out that the suspect tumor is a manifestation of one of the patient's other cancers, delete the metastatic or secondary case, re-sequence remaining cases, and correct the coding on the original case as necessary.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for an edit of the type Seq Num-Hosp, Primary Site, Morph.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of all items in the error or warning message confirms that hospital sequence number and site are both correct.

## Codes

1 Reviewed Blank Not reviewed or reviewed and corrected

## **OVER-RIDE ILL-DEFINE SITE**

OVER-RIDE ILL-DEFINE SITE		Keviscu		
Alternate Name	Item #	Length	Source of Standard	Column #
Sequence Number/Ill-defined Site Interfield Review (Interfield Edit 22) (SEER #8)	2060	1	SEER	1131-1131

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Seq Num--Central, Primary Site, Morph (NAACCR IF22)

Seq Num--Central, Primary Site, Morph (SEER IF22)

Seq Num--Central, Prim Site, Morph ICDO3 (NAACCR IF22)

Seq Num--Central, Prim Site, Morph ICDO3 (SEER IF22)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

If Sequence Number-Central indicates the person has had more than one primary, then any case with one of the following Primary Site/Histologic Type ICD-O-3 combinations requires review:

- C760-C768 (ill-defined sites) or C809 (unknown primary) and Histologic Type ICD-O-3 < 9590.
- C77\_ and Histologic Type ICD-O-3 not in range 9590-9729.
- C420-C424 and Histologic Type ICD-O-3 not in range 9590-9989.
- Any site code and Histologic Type ICD-O-3 in range 9740-9758.

## **Instructions for Coding**

• Code 1 can be used if a second or subsequent primary reporting with an ill-defined primary site has been reviewed and is indeed an independent primary.

## Codes

1 Reviewed: A second or subsequent primary reported with an ill-defined primary site (C76.0-C76.8, C80.9) has been reviewed and is an independent primary.

Blank Not reviewed or reviewed and corrected

Dovisod

<b>OVER-RIDE LEUK, LYMPHOMA</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Leukemia or Lymphoma/Diagnostic	2070	1	SEER	1132-1132
Confirmation Interfield Review (Interfield				
Edit 48) (SEER #9)				

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Diagnostic Confirmation, Histol Typ (COC)

Diagnostic Confirmation, Histologic Typ (SEER IF48)

Diagnostic Confirmation, Histol Typ ICDO3 (COC)

Diagnostic Confirmation, Histology ICDO3 (SEER IF48)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

#### Over-ride Flag as Used in the EDITS Software Package

Edits of the type Diagnostic Confirmation, Histol Typ differ in use of ICD-O-2 or ICD-O-3 and check the following:

- 1. Because lymphoma and leukemia are almost exclusively microscopic diagnoses, this edit forces review of any cases of lymphoma that have diagnostic confirmation of direct visualization or clinical, and any leukemia with a diagnostic confirmation of direct visualization.
- 2. If histology = 9590-9717 for ICD-O-2 or 9590-9729 for ICD-O-3 (lymphoma) then Diagnostic Confirmation cannot be 6 (direct visualization) or 8 (clinical).
- 3. If histology = 9720-9941 for ICD-O-2 or 9731-9948 for ICD-O-3 (leukemia and other) then Diagnostic Confirmation cannot be 6 (direct visualization).

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edits of the type Diagnostic Confirmation, Histol Typ.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- If the edit produces an error or warning message, verify that the Histologic Type and Diagnostic Confirmation are correctly coded. Remember that positive hematologic findings and bone marrow specimens are included as histologic confirmation (code 1 in Diagnostic Confirmation) for leukemia. Code 1 indicates that a review has taken place and histologic type and diagnostic confirmation are correctly coded.

- 1 Reviewed
- Blank Not reviewed or reviewed and corrected

## **OVER-RIDE REPORT SOURCE**

<b>OVER-RIDE REPORT SOURCE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Type of Reporting Source/Sequence	2050	1	SEER	1130-1130
Number Interfield Review (Interfield Edit				
04) (Seer #7)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Type of Report Srce(DC), Seq Num--Cent (NAACCR IF04)

Type of Report Srce(DC), Seq Num--Central (SEER IF04)

Type of Rep Srce(DC), Seq Num--Cent, ICDO3 (NAACCR)

Type of Rep Srce(DC), Seq Num--Cent, ICDO3 (SEER IF04)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Date Edits and Software Coordination of Standards.

## **Over-ride Flag as Used in the EDITS Software Package**

The edit Report Source checks that if the case is a death-certificate-only case and the histology is not a lymphoma, leukemia, immunoproliferative or myeloproliferative disease (morphology is less than 9590), then the tumor sequence number must specify one primary only (sequence "00").

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the report source edit.
- Code 1 if review of type of reporting source, histologic type and tumor sequence number verified that • a second or subsequent primary with a reporting source of death-certificate-only has been reviewed and is indeed an independent primary.

## Codes

Reviewed 1 Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE SEQNO/DXCONF</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Sequence Number/Diagnostic	2000	1	SEER	1125-1125
Confirmation Interfield Review (Interfield				
Edit 23) (SEER #4)				

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Diagnostic Confirm, Seq Num--Central (NAACCR IF23)

Diagnostic Confirm, Seq Num--Central (SEER IF23)

## Rationale

Some edits check for code combinations that are impossible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

- The edit checks if the case is one of multiple primaries and is not microscopically confirmed or has only positive laboratory test/marker studies (i.e., diagnostic confirmation > 5) and tumor sequence number > 00 (more than one primary).
- The edit is skipped if the sequence number is in the range of 60-99.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the Diagnostic Confirmation and Sequence Number Central edit.
- Code 1 if the cases have been reviewed and it is verified that there are multiple primaries of special sites in which at least one diagnosis has not been microscopically confirmed.

- 1 Reviewed
- Blank Not reviewed or reviewed and corrected

# **OVER-RIDE SITE/BEHAVIOR**

<b>OVER-RIDE SITE/BEHAVIOR</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Flag for Site/Behavior (IF39)	2071	1	SEER	1133-1133
(SEER #11)				

# Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Primary Site, Behavior Code (COC)

Primary Site, Behavior Code (SEER IF39)

Primary Site, Behavior Code ICDO3 (COC)

Primary Site, Behavior Code ICDO3 (SEER IF39)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## **Over-ride Flag as Used in the EDITS Software Package**

Edits of the type, Primary Site, Behavior Code, require review of the following primary sites with a behavior of *in situ* (ICD-O-2 or ICD-O-3 behavior = 2):

C269	Gastrointestinal tract, NOS
C399	Ill-defined sites within respiratory system
C559	Uterus, NOS
C579	Female genital tract, NOS
C639	Male genital organs, NOS
C689	Urinary system, NOS
C729	Nervous system, NOS
C759	Endocrine gland, NOS
C760-C768	Ill-defined sites
C809	Unknown primary site

Since the designation of *in situ* is very specific and almost always requires microscopic confirmation, ordinarily specific information should also be available regarding the primary site. Conversely, if inadequate information is available to determine a specific primary site, it is unlikely that information about a cancer being in situ is reliable.

1. If an *in situ* diagnosis is stated, try to obtain a more specific primary site. A primary site within an organ system can sometimes be identified based on the diagnostic procedure or treatment given or on the histologic type. If no more specific site can be determined, it is usually preferable to code a behavior code of 3. In the exceedingly rare situation in which it is certain that the behavior is *in situ* and no more specific site code is applicable, set Over-ride Site/Behavior to 1.

# **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Primary Site, Behavior Code (COC) and/or the edit Primary Site, Behavior Code ICD-O-3 (COC).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of site and behavior verifies that the patient has an *in situ* cancer of a nonspecific site and no further information about the primary site is available.

## Codes

- 1 Reviewed
- Blank Not reviewed or reviewed and corrected

*Note:* The IF 39 edit does not allow *in situ* cases of nonspecific sites, such as gastrointestinal tract, NOS; uterus, NOS; female genital tract, NOS; male genital organs, NOS; and others. The over-ride indicates that the conflict has been reviewed.

# **OVER-RIDE SITE/EOD/DX DT**

				IC VISCU
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Flag for Site/EOD/Diagnosis	2072	1	SEER	1134-1134
Date (IF40) (SEER #13)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Primary Site, EOD (SEER IF40) Primary Site, EOD, ICDO3 (SEER IF40)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

Edits of this type Site/EOD/DX DT do not allow "localized" disease with nonspecific sites, such as mouth, NOS; colon, NOS (except histology 8220); bone, NOS; female genital system, NOS; male genital organs, NOS; and others.

# **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Site/EOD/DX DT.
- Code 1 if the case has been reviewed and it has been verified that the patient had "localized" disease with a nonspecific site and no further information about the primary site is available.

# Codes

1 Reviewed Blank Not reviewed or reviewed and corrected Revised

## **OVER-RIDE SITE/LAT/EOD**

<b>OVER-RIDE SITE/LAT/EOD</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Flag for Site/Laterality/EOD	2073	1	SEER	1135-1135
(IF41) (SEER #12)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Primary Site, Laterality, EOD (SEER IF41) Primary Site, Laterality, EOD, ICDO3 (SEER IF41)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

Edits of this type Site/Lat/EOD apply to paired organs and do not allow EOD to be specified as *in situ*, localized, or regional by direct extension if laterality is coded as "bilateral, site unknown," or "laterality unknown."

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Site/Laterality/EOD.
- Code 1 if the case has been reviewed and it has been verified that the patient had laterality coded • nonspecifically and EOD coded specifically.

# Codes

Reviewed 1

Blank Not reviewed or reviewed and corrected

## **OVER-RIDE SITE/LAT/MORPH**

Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Flag for	2074	1	SEER	1136-1136
Site/Laterality/Morphology (IF42) (SEER				
#13)				

## Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

Laterality, Primary Site, Morphology (NAACCR IF42)

Laterality, Primary Site, Morphology (SEER IF42)

Laterality, Primary Site, Morph ICDO3 (NAACCR IF42)

Laterality, Primary Site, Morph ICDO3 (SEER IF42)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# Over-ride Flag as Used in the EDITS Software Package

Edits of the type Laterality, Primary Site, Morph differ in whether they produce a warning or an error message and in use of ICD-O-2 (including the Field Trial Editions) or ICD-O-3 morphology and do the following:

- 1. If the Primary Site is a paired organ and Behavior is *in situ* (2), then laterality must be 1, 2, or 3.
- 2. If diagnosis year less than 1988 and histology > = 9590, no further editing is performed.
- 3. If diagnosis year greater than 1987 and histology = 9140, 9700, 9701, 9590-9980, no further editing is performed.

The intent of this edit is to force review of *in situ* cases for which laterality is coded 4 (bilateral) or 9 (unknown laterality) as to origin.

1. In rare instances when the tumor is truly midline (9) or the rare combination is otherwise confirmed correct, enter a code 1 for Override Site/Lat/Morph.

# **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Laterality, Primary site, Morphology (SEER IF 42) and/or the edit Laterality, Primary site, Morph ICDO3 (SEER IF42).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if review of site, laterality and morphology verifies that the case had behavior code of "*in situ*" and laterality is not stated as "right: origin of primary;" "left: origin of primary;" or "only one side involved, right or left origin not specified".

# Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

Dovisod

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# Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following Interrecord Edit from the SEER Program:

Verify Same Primary Not Reported Twice for a Person (SEER IR09)

Presently, documentation on interrecord edits is not included in the EDITS software.

# Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

Edits of the type Site/Lat/SeqNo applies to paired organs and does not allow two cases with the same primary site group, laterality and three digit histology code. This edit verifies that the same primary is not reported twice for a person.

# **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Site/Laterality/EOD.
- Code 1 if the case has been reviewed and it has been verified that the patient had multiple primaries of the same histology (3 digit) in the same primary site group.

# Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE SITE/TNM-STGGRP</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1989	1	COC	1123-1123

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edit in the NAACCR Metafile of the EDITS software: Primary Site, AJCC Stage Group - Edition 6 (COC)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the override flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

The edit, Primary Site, AJCC Stage Group - Edition 6 (COC), checks that the pathologic and clinical AJCC stage group codes are valid for the site and histology group according to the *AJCC Cancer Staging Manual* Sixth Edition, using the codes described for the items Clinical Stage Group [970] and Pathologic Stage Group [910]. Combinations of site and histology not represented in any AJCC schema must be coded 88. Unknown codes must be coded 99. Blanks are not permitted.

Since pediatric cancers whose sites and histologies have an AJCC scheme may be coded according to a pediatric scheme instead, Override Site/TNM-Stage Group is used to indicate pediatric cases not coded according to the AJCC Manual. Pediatric Stage groups should not be recorded in the Clinical Stage Group or Pathologic Stage Group items. When neither clinical nor pathologic AJCC staging is used for pediatric cases, code all AJCC items 88. When any components of either is used to stage a pediatric case, follow the instructions for coding AJCC items and leave Override Site/TNM-Stage Group blank.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit, Primary Site, AJCC Stage Group Edition 6 (COC).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if the case is confirmed to be a pediatric case that was coded using a pediatric coding system.

## Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

<b>OVER-RIDE SITE/TYPE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Site/Type Interfield Review (Interfield Edit	2030	1	SEER	1128-1128
25) (SEER #1)				

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Primary Site, Morphology-Type Check (SEER IF25) Primary Site, Morphology-Type Check ICDO3 (SEER IF25)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

Multiple versions of edits of the type Primary site, Morphology-Type check for "usual" combinations of site and ICD-O-2 or ICD-O-3 histology. The SEER version of the edit is more restrictive than the COC edit, and thus uses a different over-ride flag. The COC version of the edit will accept Over-ride COC-Site/Type or Over-ride Site/Type as equivalent.

- 1. The Site/Histology validation list (available on the SEER web site) contains those histologies commonly found in the specified primary site. Histologies that occur only rarely or never are not included. These edits require review of all combinations not listed.
- 2. Since basal and squamous cell carcinomas of non-genital skin sites are not reportable to SEER, these site/histology combinations do not appear on the SEER validation list. For the COC version of the edit, if Primary Site is in the rage C440-C449 (skin), and Histologic Type ICD-O-3 is in the range 8000-8005 (neoplasms, malignant, NOS), 8010-8046 (epithelial carcinomas), 8050-8084 (papillary and squamous cell carcinomas), or 8090-8110 (basal cell carcinomas), no further editing is done. No over-ride is necessary for these cases in the COC version of the edit.

Review of these cases requires investigating whether a) the combination is biologically implausible, or b) there are cancer registry coding conventions that would dictate different codes for the diagnosis. Review of these rare combinations often results in changes to the primary site and/or morphology, rather than a decision that the combination is correct.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit Primary Site, Morphology-Type Check (SEER IF25) and/or the edit Primary Site, Morphology-Type ICDO3 (SEERIF25).
- Leave blank and correct any errors for the case if an item is discovered to be incorrect.
- Code 1 if the case has been reviewed and both the site and histology are correct.

# Codes

1 Reviewed Blank Not reviewed or reviewed and corrected

# **OVER-RIDE SS/DISMET1**

OVER-KIDE 55/DISMETT				Keviseu
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Summary Stage/Distant	1984	1	NAACCR	1118-1118
Metastasis 1				

#### Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Summary Stage, Site Dist Met 1 (NAACCR) Summary Stage 2000, Site Dist Met 1 (NAACCR)

#### Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

The edit SS/DisMet1 checks Summary Stage against the Distant Metastatic Site 1 and generates an error if there is an incompatibility between the two data items.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit SS/Distant Metastasis 1.
- Code 1 if the case has been reviewed and it has been verified that SEER Summary Stage and Distant Site Metastasis 1 have been coded correctly.

## Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

Dovisod

<b>OVER-RIDE SS/NODESPOS</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Summary Stage/Nodes Positive	1981	1	NAACCR	1115-1115

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Summary Stage, Regional Nodes Pos (NAACCR)

Summary Stage 2000, Regional Nodes Pos (NAACCR)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## Over-ride Flag as Used in the EDITS Software Package

The edit SS/NodesPos checks Summary Stage against the Regional Nodes Positive and generates an error if there is an incompatibility between the two data items.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit SS/Nodes Positive.
- Code 1 if the case has been reviewed and it has been verified that the case has both SEER Summary Stage and Nodes Positive coded correctly.

- 1 Reviewed
- Blank Not reviewed or reviewed and corrected

OVER-RIDE SS/TNM-M				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Summary Stage/TNM-M	1983	1	NAACCR	1117-1117

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Summary Stage, TNM-M (NAACCR) Summary Stage 2000, TNM-M (NAACCR)

## Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

The edit SS/TNM-M checks the Summary Stage against the TNM-M and generates an error if the Summary Stage is "distant" and the TNM-M is "0". It also checks if the Summary Stage is not "distant" and the TNM-M is greater than or equal to "1".

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit SS and TNM-M.
- Code 1 if the case has been reviewed and it has been verified that both SEER Summary Stage and TNM-M have been coded correctly.

# Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

OVER-RIDE SS/TNM-N				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Over-ride Summary Stage/TNM-N	1982	1	NAACCR	1116-1116

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software: Summary Stage, TNM-N (NAACCR) Summary Stage 2000, TNM-N (NAACCR)

# Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

# **Over-ride Flag as Used in the EDITS Software Package**

The edit SS/TNM-N checks Summary Stage against the TNM-N and generates an error if the Summary Stage indicates regional nodal involvement and the TNM-N does not. It also generates an error if the Summary Stage is "*in situ*" or "localized" and the TNM-N is greater than or equal to "1".

# **Instructions for Coding**

- Leave blank if the program does not generate an error message for the edit SS and TNM-N.
- Code 1 if the case has been reviewed and it has been verified that both SEER Summary Stage and TNM-N have been coded correctly.

# Codes

1 Reviewed

Blank Not reviewed or reviewed and corrected

#### **OVER-RIDE SURG/DXCONE**

OVER-RIDE SURG/DXCONF				
Alternate Name	Item #	Length	Source of Standard	Column #
Surgery/Diagnostic Confirmation Interfield	2020	1	SEER	1127-1127
Review (Interfield Edit 46) (SEER #6)				

#### Description

Some computer edits identify errors. Others indicate possible errors that require manual review for resolution. To eliminate the need to review the same cases repeatedly, over-ride flags have been developed to indicate that data in a record (or records) have been reviewed and, while unusual, are correct.

This over-ride is used with the following edits in the NAACCR Metafile of the EDITS software:

RX Summ--Surg Prim Site, Diag Conf (NAACCR IF76)

RX Summ--Surg Prim Site, Diag Conf (SEER IF76)

RX Summ--Surgery Type, Diag Conf (SEER IF46)

#### Rationale

Some edits check for code combinations that are possible, but quite rare. If the code combination generates an error message and review of the case indicates that the codes are correct for the case, then the over-ride flag is used to skip the edit in the future. See Chapter IV, Recommended Data Edits and Software Coordination of Standards.

## **Over-ride Flag as Used in the EDITS Software Package**

Edits of the type RX Summ--Surg Prim Site, Diag Conf check that cases with a primary site surgical procedure coded 20-90 are histologically confirmed.

If the patient had a surgical procedure, most likely there was a microscopic examination of the cancer. Verify the surgery and diagnostic confirmation codes, and correct any errors. Sometimes there are valid reasons why no microscopic confirmation is achieved with the surgery; for example, the tissue removed may be inadequate for evaluation.

## **Instructions for Coding**

- Leave blank if the program does not generate an error message for edits of the type, RX Summ--Surg Prim Site, Diag Conf.
- Leave blank and correct any errors for the case if an item is discovered to be incorrect. •
- Code 1 if review of the Surgery of Primary Site and Diagnostic Confirmation confirms that they are • correct. The patient had (cancer-directed) surgery, but the tissue removed was not sufficient for microscopic confirmation.

#### Codes

Reviewed 1

Blank Not reviewed or reviewed and corrected

PAIN ASSESSMENT				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	3260			

## Description

This data item was published in FORDS but later withdrawn by COC and never implemented. The NAACCR UDSC approved the COC proposal to retire this data item in September 2002.

### PATIENT ID NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
	20	8	<b>Reporting Registry</b>	2-9

### Description

Unique number assigned to an individual patient by the central registry. The central registry will assign this same number to all of the patient's subsequent tumors (records).

Patient ID Number will only differ when multiple central registries accession the same patient. Each central registry will assign their unique Patient ID Number.

NAACCR recommends that the registry should not reissue or reuse this number when a patient's record is deleted from the files.

In the transmit file (data exchange) this number will be the Patient ID Number assigned by the sending registry as defined in Registry ID [40].

#### Rationale

Provides the central registry with a unique identification number that will link all records (multiple tumors) for the same patient. The unique number also allows the central registry to identify the patient when there are multiple reports from different hospitals.

#### PEDIATRIC STAGE

Alternate Name	Item #	Length	Source of Standard	Column #
	1120	2	COC	621-622

#### Description

Code for stage of pediatric tumor in an AJCC stage scheme, a pediatric intergroup study scheme, or a pediatric cooperative group scheme.

#### Rationale

Staging of pediatric tumors requires very different schemes from those used to stage adult tumors.

### Codes

See the ROADS Manual for allowable codes for this field.

### PEDIATRIC STAGED BY

Alternate Name	Item #	Length	Source of Standard	Column #
Staged By (Pediatric Stage) (COC)	1140	1	COC	625-625

# Description

Code for person who documented the pediatric staging system and stage.

### Codes

- 0 Not staged
- 1 Managing physician
- 2 Pathologist
- 3 Other physician
- 4 Any combination of 1, 2, or 3
- 5 Registrar
- 6 Any combination of 5 with 1, 2, or 3
- 7 Other
- 8 Staged, individual not specified
- 9 Unknown if staged

# PEDIATRIC STAGING SYSTEM

Alternate Name	Item #	Length	Source of Standard	Column #
Type of Staging System (Pediatric) (COC)	1130	2	COC	623-624

# Description

Staging system used to assign the Pediatric Stage.

### Rationale

Staging of pediatric tumors requires very different schemes from those used to stage adult tumors.

### Codes

- 00 None
- 01 AJCC
- 02 Ann Arbor
- 03 Children's Cancer Group (CCG)
- 04 Evans
- 05 General Summary
- 06 Intergroup Ewings
- 07 Intergroup Hepatoblastoma
- 08 Intergroup Rhabdomyosarcoma
- 09 International System
- 10 Murphy
- 11 NCI (pediatric oncology)
- 12 National Wilms's Tumor Study
- 13 Pediatric Oncology Group (POG)
- 14 Reese-Ellsworth
- 15 SEER Extent of Disease
- 88 Not applicable (not pediatric case)
- 97 Other
- 99 Unknown

Note: This data item is no longer supported by COC (as of January 1, 2003).

PHYSICIAN 3				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Physician #3 (COC)	2490	8	COC	2579-2586
Other Physician (pre-96 COC)				

### Description

Code for another physician involved in the care of the patient. Registry may use physicians' medical license numbers or may create individual numbering systems. See *FORDS* Manual for suggested use of this item and detailed instructions.

### Codes in addition to medical license numbers or facility-generated codes

 Patient did not have radiation therapy or a radiation therapy consult.
 Physician who performed a radiation therapy procedure was not a radiation oncologist (diagnostic radiologist or surgeon).
 Primary radiation oncologist is unknown or an identification number is not assigned.

PHYSICIAN 4				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Physician #4 (COC)	2500	8	COC	2587-2594
Other Physician (pre-96 COC)				

### Description

Code for another physician involved in the care of the patient. Registry may use physicians' medical license numbers or may create individual numbering systems. See *FORDS* Manual for suggested use of this item and detailed instructions.

#### Codes in addition to medical license numbers or facility-generated codes

00000000	Patient did not receive systemic therapy or a medical oncology consult.
00000000	
88888888	Physician who gave systemic therapy was not a medical oncologist (radiation oncologist,
	general practitioner, or surgeon).
99999999	Primary medical oncologist is unknown or an identification number is not assigned.

### PHYSICIAN--FOLLOW-UP

Alternate Name	Item #	Length	Source of Standard	Column #
Following Physician (COC)	2470	8	COC	2563-2570
Follow-Up Physician (pre-96 COC)				

#### Description

Code for the physician currently responsible for the patient's medical care. Registry may use physicians' medical license numbers or may create individual numbering systems.

### Codes in addition to medical license numbers or facility-generated codes

99999999 Follow-up physician unknown or ID number not assigned

PHYSICIANMANAGING				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Managing Physician (COC)	2460	8	NAACCR	2555-2562
Attending Physician (pre-96 COC)				

### Description

Code for the physician who is responsible for the overall management of the patient during diagnosis and/or treatment for this cancer. Registry may use physicians' medical license numbers or may create individual numbering systems.

### Codes in addition to medical license numbers or facility-generated codes

99999999 Managing physician unknown or ID number not assigned

### PHYSICIAN--PRIMARY SURG

Alternate Name	Item #	Length	Source of Standard	Column #
Primary Surgeon (COC)	2480	8	COC	2571-2578

### Description

Code for physician who performed the most definitive surgical procedure. Registry may use physician's medical license numbers or may create individual numbering systems.

#### Codes in addition to medical license numbers or facility-generated codes

00000000	Patient had no surgery and no surgical consultation.
88888888	Physician who performed a surgical procedure was not a surgeon (i.e., radiation oncologist,
	diagnostic radiologist, or general practitioner)
00000000	Primary Surgeon unknown or ID number not assigned

99999999	Primary Surgeon unknown or ID number not assigned
----------	---

PLACE OF DEATH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1940	3	NPCR	1394-1396

#### Description

State or country where the patient died and where certificate of death is filed.

#### Rationale

This field also helps carry out death clearance. When a hospital reports a place of death, the information can help in death certificate matching. It can also signal an out-of-state death for which the death certificate is to be requested.

### Codes in addition to geocodes

- 997 Not applicable, patient alive
- 999 Place of death unknown

Note: See Appendix B of the SEER Program Code Manual or the COC ROADS Manual, Appendix C.

PLACE OF DIAGNOSIS				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2690	50	NPCR	5875-5924

## Description

Text area for manual documentation of the facility, physician office, city, state, or county where the diagnosis was made.

## Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

### Suggestions for text:

- The complete name of the hospital or the physician office where diagnosis occurred. The initials of a hospital are not adequate.
- For out-of-state residents and facilities, include the city and the state where the medical facility is located.

### Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item Number
Reporting Hospital	540
RX Hosp-DX/Stg Proc	740
RX Hosp-Surg Prim Site	670
Type of Reporting Source	500

Class of Case	620
Institution Referred From	2410
Institution Referred To	2420

## PRESENTATION AT CA CONF

Alternate Name	Item #	Length	Source of Standard	Column #
Presentation at Cancer Conference (COC)	650	1	COC	448-448

#### Description

Documents presentation of the case at a cancer conference at the reporting facility and the type or format of the presentation.

### Rationale

Collection of this item and Date of CA Conference [660] allows preparation of reports on the number of cancer conferences, sites presented and types of presentation for administrative use, quality control, and survey preparation.

### Codes

- 0 Not presented
- 1 Prospective presentation (diagnostic)
- 2 Prospective presentation (treatment)
- 3 Prospective presentation (follow-up care)
- 4 Prospective presentation (combinations of 1, 2, or 3)
- 5 Prospective, NOS
- 6 Retrospective presentation
- 7 Follow-up presentation
- 8 Presentation, NOS
- 9 Unknown

## PRIMARY PAYER AT DX

Alternate Name	Item #	Length	Source of Standard	Column #
Primary Payer at Diagnosis (COC)	630	2	COC	445-446

## Description

Primary payer/insurance carrier at the time of initial diagnosis and/or treatment.

### Rationale

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

### Codes

- 01 Not insured
- 02 Not insured, self-pay
- 10 Insurance, NOS
- 20 Managed care, HMO, PPO
- 31 Medicaid
- 35 Medicaid administered through a Managed Care plan
- 36 Medicaid with Medicare supplement
- 50 Medicare
- 51 Medicare with supplement
- 52 Medicare with Medicaid supplement
- 53 TRICARE
- 54 Military
- 55 Veterans Affairs
- 56 Indian/Public Health Service
- 99 Insurance status unknown

### PRIMARY SITE

Alternate Name	Item #	Length	Source of Standard	Column #
	400	4	SEER/COC	291-294

#### Description

Code for the primary site of the tumor being reported using either ICD-O-2 or ICD-O-3. NAACCR adopted ICD-O-2 as the standard coding system for tumors diagnosed beginning January 1, 1992. In addition, NAACCR recommended that tumors diagnosed prior to 1992 be converted to ICD-O-2. The topography (primary site) codes have not changed between ICD-O-2 and ICD-O-3.

### Codes

See ICD-O-2,<sup>14</sup> or ICD-O-3,<sup>13</sup> Topography Section, for the codes for primary site.

Note: See Site (73-91) ICD-O-1 [1960] for ICD-O-1 cases.

### PROTOCOL ELIGIBILITY STAT

Alternate Name	Item #	Length	Source of Standard	Column #
Protocol Eligibility Status (COC)	1470	1	COC	890-890

### Description

Code for eligibility status of patient to be entered into a protocol.

#### Codes

- 0 Protocol not available
- 1 On protocol
- 2 Patient ineligible (age, stage, etc.)
- 3 Patient ineligible (comorbidity, pre-existing condition)
- 4 Patient entered but withdrawn from study
- 6 Patient eligible, but not entered, reason not specified
- 7 Patient eligible, patient or patient's guardian refused
- 8 Protocol not recommended
- 9 Unknown if on protocol

Note: This data item is not longer supported by COC (as of January 1, 2003).

### PROTOCOL PARTICIPATION

Alternate Name	Item #	Length	Source of Standard	Column #
	1480	2	COC	891-892

### Description

Code indicating agency or group that established the protocol in which the patient is participating.

### Codes

00 Not on/not applicable

- National Protocols:
- 01 NSABP
- 02 GOG
- 03 RTOG
- 04 SWOG
- 05 ECOG
- 06 POG
- 07 CCG
- 08 CALGB
- 09 NCI
- 10 ACS
- 11 National protocol, NOS
- 12 ACOS-OG
- 13-50 National trials
- 51-98 Locally defined trials
- 99 Unknown

## **QUALITY OF SURVIVAL**

Alternate Name	Item #	Length	Source of Standard	Column #
	1780	1	COC	1304-1304

### Description

Records patient's ability to carry on the activities of daily living at the date of last contact.

#### Codes

- 0 Normal activity
- 1 Symptomatic and ambulatory
- 2 Ambulatory more than 50 percent of the time, occasionally needs assistance
- 3 Ambulatory less than 50 percent of the time, nursing care needed
- 4 Bedridden, may require hospitalization
- 8 Not applicable, dead
- 9 Unknown or unspecified

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### RACE 1

Alternate Name	Item #	Length	Source of Standard	Column #
Race	160	2	SEER/COC	103-104

#### Description

Code the patient's race. Race is coded separately from Spanish/Hispanic Origin [190]. All tumors for the same patient should have the same race code. If the patient is multiracial, code all races using RACE 2 through RACE 5 [161-164].

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

### Rationale

Because race has a significant association with cancer rates and outcomes, a comparison between areas with different racial distributions may require an analysis of race to interpret the findings. The race codes listed correspond closely to race categories used by the U.S. Census Bureau to allow calculation of race-specific incidence rates. The full coding system should be used to allow accurate national comparison and collaboration, even if the state population does not include many of the race categories.

- 01 White
- 02 Black
- 03 American Indian, Aleutian, or Eskimo (includes all indigenous populations of the Western hemisphere)
- 04 Chinese
- 05 Japanese
- 06 Filipino
- 07 Hawaiian
- 08 Korean
- 09 Asian Indian, Pakistani
- 10 Vietnamese

- 11 Laotian
- 12 Hmong
- 13 Kampuchean
- 14 Thai
- 20 Micronesian, NOS
- 21 Chamorran
- 22 Guamanian, NOS
- 25 Polynesian, NOS
- 26 Tahitian
- 27 Samoan
- 28 Tongan
- 30 Melanesian, NOS
- 31 Fiji Islander
- 32 New Guinean
- 96 Other Asian, including Asian, NOS and Oriental, NOS
- 97 Pacific Islander, NOS
- 98 Other
- 99 Unknown

*Note:* Codes 20-97 were adopted for use effective with 1991 diagnoses. Code 14 was adopted for use effective with 1994 diagnoses.

# RACE 2

Alternate Name	Item #	Length	Source of Standard	Column #
	161	2	SEER/COC	105-106

### Description

Code the patient's race. Race is coded separately from Spanish/Hispanic Origin [190]. All tumors for the same patient should have the same race code. If the patient is multiracial, code all races using RACE 2 through RACE 5 [161-164].

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

### Rationale

Because race has a significant association with cancer rates and outcomes, a comparison between areas with different racial distributions may require an analysis of race to interpret the findings. The race codes listed correspond closely to race categories used by the U.S. Census Bureau to allow calculation of race-specific incidence rates. The full coding system should be used to allow accurate national comparison and collaboration, even if the state population does not include many of the race categories.

- 01 White
- 02 Black
- 03 American Indian, Aleutian, or Eskimo (includes all indigenous populations of the Western hemisphere)
- 04 Chinese
- 05 Japanese
- 06 Filipino

- 07 Hawaiian
- 08 Korean
- 09 Asian Indian, Pakistani
- 10 Vietnamese
- 11 Laotian
- 12 Hmong
- 13 Kampuchean
- 14 Thai
- 20 Micronesian, NOS
- 21 Chamorran
- 22 Guamanian, NOS
- 25 Polynesian, NOS
- 26 Tahitian
- 27 Samoan
- 28 Tongan
- 30 Melanesian, NOS
- 31 Fiji Islander
- 32 New Guinean
- 88 No further race documented
- 96 Other Asian, including Asian, NOS and Oriental, NOS
- 97 Pacific Islander, NOS
- 98 Other
- 99 Unknown

*Note:* Codes 20-97 were adopted for use effective with 1991 diagnoses. Code 14 was adopted for use effective with 1994 diagnoses. Code 88 was adopted for use effective with 2000 diagnoses.

*Note:* If diagnosed prior to 2000 and any race code (Race 2, 3, 4, or 5) is blank, all subsequent race codes must be blank. If diagnosed after 1999 and any race code (for Race 2, 3, 4, and 5) is 88 (no further race documented), then all subsequent race codes also must be 88. If any race equals 99, then all race codes (Race 1, 2, 3, 4, and 5) must be 99.

RACE 3				
Alternate Name	Item #	Length	Source of Standard	Column #
	162	2	SEER/COC	107-108

### Description

Code the patient's race. Race is coded separately from Spanish/Hispanic Origin [190]. All tumors for the same patient should have the same race code. If the patient is multiracial, code all races using RACE 2 through RACE 5 [161-164].

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

### Rationale

Because race has a significant association with cancer rates and outcomes, a comparison between areas with different racial distributions may require an analysis of race to interpret the findings. The race codes listed correspond closely to race categories used by the U.S. Census Bureau to allow calculation of race-specific

incidence rates. The full coding system should be used to allow accurate national comparison and collaboration, even if the state population does not include many of the race categories.

Codes

- 01 White
- 02 Black
- 03 American Indian, Aleutian, or Eskimo (includes all indigenous populations of the Western hemisphere)
- 04 Chinese
- 05 Japanese
- 06 Filipino
- 07 Hawaiian
- 08 Korean
- 09 Asian Indian, Pakistani
- 10 Vietnamese
- 11 Laotian
- 12 Hmong
- 13 Kampuchean
- 14 Thai
- 20 Micronesian, NOS
- 21 Chamorran
- 22 Guamanian, NOS
- 25 Polynesian, NOS
- 26 Tahitian
- 27 Samoan
- 28 Tongan
- 30 Melanesian, NOS
- 31 Fiji Islander
- 32 New Guinean
- 88 No further race documented
- 96 Other Asian, including Asian, NOS and Oriental, NOS
- 97 Pacific Islander, NOS
- 98 Other
- 99 Unknown

*Note:* Codes 20-97 were adopted for use effective with 1991 diagnoses. Code 14 was adopted for use effective with 1994 diagnoses. Code 88 was adopted for use effective with 2000 diagnoses.

*Note:* If diagnosed prior to 2000 and any race code (Race 2, 3, 4, or 5) is blank, all subsequent race codes must be blank. If diagnosed after 1999 and any race code (for Race 2, 3, 4, and 5) is 88 (no further race documented), then all subsequent race codes also must be 88. If any race equals 99, then all race codes (Race 1, 2, 3, 4, and 5) must be 99.

# RACE 4

Alternate Name	Item #	Length	Source of Standard	Column #
	163	2	SEER/COC	109-110

## Description

Code the patient's race. Race is coded separately from Spanish/Hispanic Origin [190]. All tumors for the same patient should have the same race code. If the patient is multiracial, code all races using RACE 2 through RACE 5 [161-164].

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

# Rationale

Because race has a significant association with cancer rates and outcomes, a comparison between areas with different racial distributions may require an analysis of race to interpret the findings. The race codes listed correspond closely to race categories used by the U.S. Census Bureau to allow calculation of race-specific incidence rates. The full coding system should be used to allow accurate national comparison and collaboration, even if the state population does not include many of the race categories.

- 01 White
- 02 Black
- 03 American Indian, Aleutian, or Eskimo (includes all indigenous populations of the Western hemisphere)
- 04 Chinese
- 05 Japanese
- 06 Filipino
- 07 Hawaiian
- 08 Korean
- 09 Asian Indian, Pakistani
- 10 Vietnamese
- 11 Laotian
- 12 Hmong
- 13 Kampuchean
- 14 Thai
- 20 Micronesian, NOS
- 21 Chamorran
- 22 Guamanian, NOS
- 25 Polynesian, NOS
- 26 Tahitian
- 27 Samoan
- 28 Tongan
- 30 Melanesian, NOS
- 31 Fiji Islander
- 32 New Guinean
- 88 No further race documented
- 96 Other Asian, including Asian, NOS and Oriental, NOS
- 97 Pacific Islander, NOS
- 98 Other
- 99 Unknown

*Note:* Codes 20-97 were adopted for use effective with 1991 diagnoses. Code 14 was adopted for use effective with 1994 diagnoses. Code 88 was adopted for use effective with 2000 diagnoses.

Note If diagnosed prior to 2000 and any race code (Race 2, 3, 4, or 5) is blank, all subsequent race codes must be blank. If diagnosed after 1999 and any race code (for Race 2, 3, 4, and 5) is 88 (no further race documented), then all subsequent race codes also must be 88. If any race equals 99, then all race codes (Race 1, 2, 3, 4, and 5) must be 99.

# RACE 5

Alternate Name	Item #	Length	Source of Standard	Column #
	164	2	SEER/COC	111-112

### Description

Code the patient's race. Race is coded separately from Spanish/Hispanic Origin [190]. All tumors for the same patient should have the same race code. If the patient is multiracial, code all races using RACE 2 through RACE 5 [161-164].

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

## Rationale

Because race has a significant association with cancer rates and outcomes, a comparison between areas with different racial distributions may require an analysis of race to interpret the findings. The race codes listed correspond closely to race categories used by the U.S. Census Bureau to allow calculation of race-specific incidence rates. The full coding system should be used to allow accurate national comparison and collaboration, even if the state population does not include many of the race categories.

- 01 White
- 02 Black
- 03 American Indian, Aleutian, or Eskimo (includes all indigenous populations of the Western hemisphere)
- 04 Chinese
- 05 Japanese
- 06 Filipino
- 07 Hawaiian
- 08 Korean
- 09 Asian Indian, Pakistani
- 10 Vietnamese
- 11 Laotian
- 12 Hmong
- 13 Kampuchean
- 14 Thai
- 20 Micronesian, NOS
- 21 Chamorran
- 22 Guamanian, NOS
- 25 Polynesian, NOS
- 26 Tahitian
- 27 Samoan

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

*Note:* Codes 20-97 were adopted for use effective with 1991 diagnoses. Code 14 was adopted for use effective with 1994 diagnoses. Code 88 was adopted for use effective with 2000 diagnoses.

*Note:* If diagnosed prior to 2000 and any race code (Race 2, 3, 4, or 5) is blank, all subsequent race codes must be blank. If diagnosed after 1999 and any race code (for Race 2, 3, 4, and 5) is 88 (no further race documented), then all subsequent race codes also must be 88. If any race equals 99, then all race codes (Race 1, 2, 3, 4, and 5) must be 99.

## **RACE CODING SYS--CURRENT**

Alternate Name	Item #	Length	Source of Standard	Column #
	170	1	NAACCR	113-113

### Description

Code describes how race currently is coded. If the data have been converted, this field shows the system to which it has been converted.

### Rationale

Race 1 - 5 codes [160 - 164] have changed over time. To be able to accurately group and analyze the data, it is necessary to record the system used to record the race codes.

- 1 4-value coding: 1 = White, 2 = Black, 3 = Other, 9 = Unknown
- 2 SEER < 1988 (1-digit)
- 3 1988-1990 SEER & COC (2-digit)
- 4 1991-1993 SEER & COC (added codes 20-97, additional Asian and Pacific Islander codes)
- 5 1994-1999 SEER & COC (added code 14, Thai)
- 6 2000+ SEER & COC (added code 88 for Race 2, 3, 4, and 5)
- 9 Other

## RACE CODING SYS--ORIGINAL

Alternate Name	Item #	Length	Source of Standard	Column #
	180	1	NAACCR	114-114

### Description

Code that best describes how Race [160] originally was coded. If data have been converted, this field identifies the coding system originally used to code the case.

#### Rationale

Race 1 - 5 codes [160 - 164] have changed over time. Identifying both original and current coding systems used to code race promotes accurate data grouping and analysis.

#### Codes

- 1 4-value coding: 1 = White, 2 = Black, 3 = Other, 9 = Unknown
- 2 SEER < 1988 (1-digit)
- 3 1988-1990 SEER & COC (2-digit)
- 4 1991-1993 SEER & COC (added codes 20-97, additional Asian and Pacific Islander codes)
- 5 1994-1999 SEER & COC (added code 14, Thai)
- 6 2000+ SEER & COC (added code 88 for Race 2, 3, 4, and 5)
- 9 Other

#### **RAD--BOOST DOSE CGY**

Alternate Name	Item #	Length	Source of Standard	Column #
Boost Radiation Dose: cGy	3210	5	COC	913-917

#### Description

Records the additional dose delivered to that part of the treatment volume encompassed by the boost fields or devices. The unit of measure is centiGray (cGy).

#### Rationale

To evaluate patterns of radiation oncology care, it is necessary to describe the prescribed boost radiation dose. As in chemotherapy, outcomes are strongly related to the dose delivered.

### Codes (in addition to value dose)

- (Fill blanks) Record the actual boost dose delivered
  00000 Boost radiation therapy was not administered
  88888 Not applicable, brachytherapy or radioisotopes administered to the patient
- 99999 Boost radiation therapy administered, boost dose unknown

## **RAD--BOOST RX MODALITY**

Alternate Name	Item #	Length	Source of Standard	Column #
Boost Radiation Treatment Modality	3200	2	COC	911-912

## Description

Records the radiation treatment--boost modality used to deliver the most clinically significant dose to the primary volume of interest during the first course of treatment. This is accomplished with external beam fields of reduced size (relative to the regional treatment fields), implants, stereotactic radiosurgery, conformal therapy, or intensity-modulated radiation therapy. External beam boosts may consist of two or more successive phases with progressively smaller fields, and they are generally coded as a single entity. This field is used with Rad--Regional RX Modality [1570].

## Rationale

Radiation treatment frequently is delivered in two or more phases that can be summarized as regional and boost treatments. For outcomes analysis, the modalities used for each of these phases can be very important.

- 00 No boost treatment
- 20 External beam, NOS
- 21 Orthovoltage
- 22 Cobalt-60, Cesium-137
- 23 Photons (2-5 MV)
- 24 Photons (6-10 MV)
- 25 Photons (11-19 MV)
- 26 Photons (> 19 MV)
- 27 Photons (mixed energies)
- 28 Electrons
- 29 Photons and electrons mixed
- 30 Neutrons, with or without photons/electrons
- 31 IMRT
- 32 Conformal or 3-D therapy
- 40 Protons
- 41 Stereotactic radiosurgery, NOS
- 42 Linac radiosurgery
- 43 Gamma Knife
- 50 Brachytherapy, NOS
- 51 Brachytherapy, Intracavitary, LDR
- 52 Brachytherapy, Intracavitary, HDR
- 53 Brachytherapy, Interstitial, LDR
- 54 Brachytherapy, Interstitial, HDR
- 55 Radium
- 60 Radio-isotopes, NOS
- 61 Strontium 89
- 62 Strontium 90
- 98 Other, NOS
- 99 Unknown

# RAD--ELAPSED RX DAYS

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Elapsed Treatment Time (Days)	1530	3	COC	902-904
(COC)				

## Description

Actual number of radiation treatment days during first course of treatment, including weekend days and intervals of rest. See also RX Summ--Radiation [1360].

## **Special codes**

- 000 No radiation therapy administered
- 999 Radiation therapy administered, but number of treatment days is unknown; unknown if radiation therapy given

Note: This data item is no longer supported by COC (as of January 1, 2003).

# **RAD--INTENT OF TREATMENT**

Alternate Name	Item #	Length	Source of Standard	Column #
Intent of Treatment (Radiation) (COC)	1560	1	COC	908-908

## Description

Code for intent of radiation treatment during first course of therapy. See also RX Summ--Radiation [1360].

### Codes

- 0 No radiation treatment
- 1 Curative (primary)
- 2 Curative (adjuvant)
- 4 Palliative (pain control)
- 5 Palliative (other, cosmetic)
- 6 Prophylactic (no symptoms, preventive)
- 8 Other, NOS
- 9 Intent unknown; unknown if radiation therapy given

# **RAD--LOCAL CONTROL STATUS**

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Therapy Local Control Status	1590	1	COC	919-919
(Irradiated Volume) (COC)				

# Description

Code for results of radiation therapy during first course of therapy in terms of disease control within the irradiated volume. See also RX Summ--Radiation [1360].

## Codes

- 0 No radiation treatment
- 1 Tumor control status not evaluable
- 2 Tumor/symptoms controlled
- 3 Tumor/symptoms have returned
- 4 Tumor/symptoms never adequately controlled
- 8 Other, NOS
- 9 Status unknown; unknown if radiation therapy given

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **RAD--LOCATION OF RX**

Alternate Name	Item #	Length	Source of Standard	Column #
Location of Radiation Treatment (COC)	1550	1	COC	907-907

### Description

Code for location where radiation treatment was administered during first course of therapy. See also RX Summ--Radiation [1360].

- 0 No radiation treatment
- 1 All radiation treatment at this facility
- 2 Regional treatment at this facility, boost elsewhere
- 3 Boost radiation at this facility, regional elsewhere
- 4 All radiation treatment elsewhere
- 8 Other, NOS
- 9 Unknown

# RAD--NO OF TREATMENT VOL

Alternate Name	Item #	Length	Source of Standard	Column #
Number of Treatments to this Volume	1520	2	COC	900-901
(COC)				

# Description

Records the total number of treatment sessions (fractions) administered during the first course of therapy. See also RX Summ--Radiation [1360].

Codes

00None01-98Number of treatments.

99 Unknown

# **RAD--REGIONAL DOSE: CGY**

Alternate Name	Item #	Length	Source of Standard	Column #
Regional Dose: cGy (COC)	1510	5	COC	895-899

# Description

The dominant or most clinically significant total dose of regional radiation therapy delivered to the patient during the first course of treatment. The unit of measure is centiGray (cGy). See also Rad--Regional RX Modality [1570].

# Codes (in addition to actual doses)

(Fill spaces) Record the actual regional dose delivered
00000 Radiation therapy was not administered
88888 Not applicable, brachytherapy or radioisotopes administered to the patient
99999 Regional radiation therapy was administered, but the dose is unknown

## **RAD--REGIONAL RX MODALITY**

Alternate Name	Item #	Length	Source of Standard	Column #
Regional Treatment Modality (COC)	1570	2	COC	909-910

## Description

Records the dominant modality of radiation therapy used to deliver the clinically most significant regional dose to the primary volume of interest during the first course of treatment.

### Rationale

Radiation treatment frequently is delivered in two or more phases that can be summarized as regional and boost treatments. To evaluate patterns of radiation oncology care, it is necessary to know which radiation resources were employed in the delivery of therapy. For outcomes analysis, the modalities used for each of these phases can be very important.

## Codes

- 00 No radiation treatment
- 20 External beam, NOS
- 21 Orthovoltage
- 22 Cobalt-60, Cesium-137
- 23 Photons (2-5 MV)
- 24 Photons (6-10 MV)
- 25 Photons (11-19 MV)
- 26 Photons (> 19 MV)
- 27 Photons (mixed energies)
- 28 Electrons
- 29 Photons and electrons mixed
- 30 Neutrons, with or without photons/electrons
- 31 IMRT
- 32 Conformal or 3-D therapy
- 40 Protons
- 41 Stereotactic radiosurgery, NOS
- 42 Linac radiosurgery
- 43 Gamma Knife
- 50 Brachytherapy, NOS
- 51 Brachytherapy, Intracavitary, Low Dose Rate (LDR)
- 52 Brachytherapy, Intracavitary, High Dose Rate (HDR)
- 53 Brachytherapy, Interstitial, Low Dose Rate (LDR)
- 54 Brachytherapy, Interstitial, High Dose Rate (HDR)
- 55 Radium
- 60 Radio-isotopes, NOS
- 61 Strontium 89
- 62 Strontium 90
- 80\* Combination modality, specified
- 85\* Combination modality, NOS
- 98 Other, NOS
- 99 Unknown

*Note:* For tumors diagnosed prior to January 1, 2003, the codes reported in this data item describe any radiation administered to the patient as part or all of the first course of therapy. Codes 80 and 85 describe

specific converted descriptions of radiation therapy coded according to *Volume II ROADS*, and *DAM* rules and should only be used to record regional radiation for tumors diagnosed prior to January 1, 2003.

# **RAD--RX COMPLETION STATUS**

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Treatment Completion Status	1580	1	COC	918-918
(COC)				

# Description

Code indicating whether or not the patient's radiation therapy was completed as outlined in the initial treatment plan. See also RX Summ--Radiation [1360].

# Codes

- 0 No radiation treatment
- 1 Treatment completed
- 2 Radiation not complete, patient health
- 3 Radiation not complete, patient expired
- 4 Radiation not complete, patient choice
- 5 Radiation not complete, family choice
- 6 Radiation not complete, complications
- 7 Radiation not complete, cytopenia
- 8 Radiation not complete, other reason
- 9 Radiation not complete, reason unknown; unknown if radiation therapy given

## **RAD--TREATMENT VOLUME**

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Treatment Volume (COC)	1540	2	COC	905-906

## Description

Identifies the volume or anatomic target of the most clinically significant regional radiation therapy delivered to the patient during the first course of therapy. See also Rad--Regional RX Modality [1570].

- 00 No radiation therapy, not applicable
- 01 Eye/orbit
- 02 Pituitary
- 03 Brain (NOS)
- 04 Brain (limited)
- 05 Head and neck (NOS)
- 06 Head and neck (limited)
- 07 Glottis
- 08 Sinuses
- 09 Parotid
- 10 Chest/lung (NOS)
- 11 Lung (limited)
- 12 Esophagus
- 13 Stomach
- 14 Liver
- 15 Pancreas
- 16 Kidney
- 17 Abdomen (NOS)
- 18 Breast
- 19 Breast/lymph nodes
- 20 Chest wall
- 21 Chest wall/lymph nodes
- 22 Mantle, mini-mantle
- 23 Lower extended field
- 24 Spine
- 25 Skull
- 26 Ribs
- 27 Hip
- 28 Pelvic bones
- 29 Pelvis (NOS)
- 30 Skin
- 31 Soft tissue
- 32 Hemibody
- 33 Whole body
- 34 Bladder and pelvis
- 35 Prostate and pelvis
- 36 Uterus and Cervix
- 37 Shoulder
- 38 Extremities bone, NOS
- 39 Inverted Y

- 40 Spinal cord
- 41 Prostate
- 50 Thyroid
- 60 Lymph node region, NOS
- 98 Other
- 99 Unknown

### **READM SAME HOSP 30 DAYS**

Alternate Name	Item #	Length	Source of Standard	Column #
Readmission to the Same Hospital Within	3190	1	COC	938-938
30 Days of Surgical Discharge				

### Description

Records a readmission to the same hospital within 30 days of discharge following hospitalization for surgical resection of the primary site when readmission is related to the treatment of this cancer.

#### Rationale

This data item provides information related to the quality-of-care. A patient may have a readmission related to the primary diagnosis on discharge if the length of stay was too short, and then needed to return due to problems or complications. A patient may also need to be readmitted if discharge planning and/or follow-up instructions were ineffective. It is important to distinguish a planned from an unplanned readmission, since a planned readmission is not an indicator of quality-of-care problems.

- 0 No surgical procedure of the primary site was performed. Patient not readmitted to the same hospital within 30 days of discharge.
- 1 Patient was surgically treated and was then readmitted to the same hospital within 30 days of being discharged. This readmission was unplanned.
- 2 Patient was surgically treated and was then readmitted to the same hospital within 30 days of being discharged. This readmission was planned (chemotherapy port insertion, revision of colostomy, etc.).
- 3 Patient was surgically treated and, within 30 days of being discharged, had both a planned and an unplanned readmission to the same hospital.
- 9 It is unknown whether surgery of the primary site was recommended or performed. It is unknown whether the patient was readmitted to the same hospital within 30 days of discharge. Death certificate only.

## **REASON FOR NO CHEMO**

Alternate Name	Item #	Length	Source of Standard	Column #
Reason for No Chemotherapy (COC)	1440	1	COC	886-886

## Description

Code for reason patient did not receive chemotherapy as part of first course of therapy. See also RX Summ--Chemo [1390].

### Codes

- 0 Chemotherapy administered
- 1 Chemotherapy not recommended
- 2 Chemotherapy contraindicated because of other conditions; autopsy-only case
- 6 Reason unknown for no chemotherapy
- 7 Patient or patient's guardian refused chemotherapy
- 8 Chemotherapy recommended, unknown if administered
- 9 Unknown if chemotherapy recommended or performed; death certificate-only case

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **REASON FOR NO HORMONE**

Alternate Name	Item #	Length	Source of Standard	Column #
Reason for No Hormone Therapy (COC)	1450	1	COC	887-887

### Description

Code for reason patient did not receive hormone therapy as part of first course of therapy. See also RX Summ--Hormone [1400].

### Codes

- 0 Hormone therapy administered
- 1 Hormone therapy not recommended
- 2 Hormone therapy contraindicated because of other conditions; autopsy-only case
- 6 Reason unknown for no hormone therapy
- 7 Patient or patient's guardian refused hormone therapy
- 8 Hormone therapy recommended, unknown if administered
- 9 Unknown if hormone therapy recommended or performed; death-certificate-only case

# **REASON FOR NO RADIATION**

Alternate Name	Item #	Length	Source of Standard	Column #
Reason for No Regional Radiation Therapy	1430	1	COC	885-885

# Description

Code the reason the patient did not receive radiation treatment as part of first course of therapy. See also RX--Regional RX Modality [1570].

- 0 Radiation therapy was administered.
- 1 Radiation therapy was not administered because it was not part of the planned first-course treatment.
- 2 Radiation therapy was not administered because it was contraindicated due to patient risk factors (comorbid conditions, advanced age, etc).
- 5 Radiation therapy was not administered because the patient died prior to planned or recommended treatment.
- 6 Radiation therapy was not administered; it was recommended by the patient's physician, but was not administered as part of the first-course therapy. No reason was noted in the patient's record.
- 7 Radiation therapy was not administered; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 8 Radiation therapy was recommended, but it is unknown if it was administered.
- 9 It is unknown if radiation therapy was recommended or administered. Death-certificate-only and autopsy-only cases.

# REASON FOR NO SURGERY

Alternate Name	Item #	Length	Source of Standard	Column #
Reason for No Cancer-Directed Surgery	1340	1	SEER/COC	868-868
(SEER)				
Reason for No CA Dir Surgery (COC)				
Reason for No Surgery to Primary Site				

# Description

Records the reason that no surgery was performed on the primary site.

- 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 therapy. No reason was noted in the patient's record.
- 7 Surgery of the primary site was not performed; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 8 Surgery of the primary site was recommended, but it is unknown if it was performed. Further followup is recommended.
- 9 It is unknown if surgery of the primary site was recommended or performed. Death certificate-only cases and autopsy-only cases.

## **RECORD TYPE**

Alternate Name	Item #	Length	Source of Standard	Column #
	10	1	NAACCR	1-1

## Description

Generated field that identifies which of the seven NAACCR data exchange record types is being used in a file of data exchange records. A file should have records of only one type.

### Codes

- I Incidence-only record type (nonconfidential coded data) Length = 1946
- C Confidential record type (incidence record plus confidential data) Length = 2644
- A Full case Abstract record type (incidence and confidential data plus text summaries; used for reporting to central registries) Length = 6694
- U Correction/Update record type (short format record used to submit corrections to data already submitted)

Length = 850

- R Analysis/**R**esearch record type (incidence record plus appended error flags and recoded values) Length = 2215
- M Record Modified since previous submission to central registry (identical in format to the "A" record type)
  - Length = 6694
- L Pathology Laboratory

# **RECURRENCE DATE--1ST**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of First Recurrence (COC)	1860	8	COC	1342-1349

### Description

The date of the first recurrence of this tumor. See page 83 for date format.

Codes

00000000 Patient became disease-free after treatment, never had a recurrence, or patient was never disease-free. Diagnosed at autopsy.

99999999 Unknown if the patient had a first recurrence or the tumor was identified by DCO.

<b>RECURRENCE DISTANT SITE 1</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1871	1	NAACCR	1350-1350

### Description

Code for the distant site or sites in which the tumor has recurred.

#### Codes

- 0 None or none known
- 1 Peritoneum
- 2 Lung
- 3 Pleura
- 4 Liver
- 5 Bone
- 6 Central nervous system
- 7 Skin
- 8 Lymph nodes (distant)
- 9 Other, generalized, NOS, carcinomatosis

Note: When carcinomatosis is present, all three fields—Recurrence Distant Site 1, 2, and 3—are coded 9.

Note: This data item is no longer supported by COC (as of January 1, 2003).

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RECORDENCE DISTANT SITE 2				Keviseu
Alternate Name	Item #	Length	Source of Standard	Column #
	1872	1	NAACCR	1351-1351

#### Description

Code for the distant site or sites in which the tumor has recurred.

#### Codes

- 0 None or none known
- 1 Peritoneum
- 2 Lung
- 3 Pleura
- 4 Liver
- 5 Bone
- 6 Central nervous system
- 7 Skin
- 8 Lymph nodes (distant)
- 9 Other, generalized, NOS, carcinomatosis

*Note:* When carcinomatosis is present, all three fields—Recurrence Distant Site 1, 2, and 3—are coded 9. If Recurrence Distant Site 1 [1871] is coded to 0, then this field also must be coded to 0.

Note: This data item is no longer supported by COC (as of January 1, 2003).

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<b>RECURRENCE DISTANT SITE 3</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1873	1	NAACCR	1352-1352

### Description

Code for the distant site or sites in which the tumor has recurred.

#### Codes

- 0 None or none known
- Peritoneum 1
- 2 Lung
- 3 Pleura
- 4 Liver
- 5 Bone
- Central nervous system 6
- 7 Skin
- 8 Lymph nodes (distant)
- 9 Other, generalized, NOS, carcinomatosis

Note: When carcinomatosis is present, all three fields—Recurrence Distant Site 1, 2, and 3—are coded 9. If Recurrence Distant Site 1 [1871] is coded to 0, then this field also must be coded to 0.

*Note:* This data item is no longer supported by COC (as of January 1, 2003).

# RECURRENCE DISTANT SITES

RECURRENCE DISTANT SITES				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	1870			

#### Description

The name for a group of subfields that describe a distant site or sites in which a tumor has recurred. The subfields are edited as three separate 1-digit fields and as a single field.

Only the group item has retired. The subfields are not retired.

Group names appear only in the data dictionary and Appendix E.

#### Subfields

Recurrence Distant Site 1 [1871] Recurrence Distant Site 2 [1872] Recurrence Distant Site 3 [1873]

## **RECURRENCE TYPE--1ST**

Alternate Name	Item #	Length	Source of Standard	Column #
Type of First Recurrence (COC)	1880	2	COC	1353-1354

## Description

Code for the type of first recurrence after a period of documented disease free intermission or remission.

- 00 Patient became disease-free after treatment and has not had a recurrence; leukemia's that are in remission.
- 04 *In situ* recurrence of an invasive tumor.
- 06 In situ recurrence of an *in situ* tumor.
- 10 Local recurrence and there is insufficient information available to code to 13-17. Recurrence is confined to the remnant of the organ of origin; to the organ of origin; to the anastomosis; or to scar tissue where the organ previously existed.
- 13 Local recurrence of an invasive tumor.
- 14 Trocar recurrence of an invasive tumor. Includes recurrence in the trocar path or entrance site following prior surgery.
- 15 Both local and trocar recurrence of an invasive tumor (both 13 and 14)
- 16 Local recurrence of an *in situ* tumor.
- 17 Both local and trocar recurrence of an *in situ* tumor.
- 20 Regional recurrence, and there is insufficient information available to code to 21-27.
- 21 Recurrence of an invasive tumor in adjacent tissue or organ(s) only.
- 22 Recurrence of an invasive tumor in regional lymph nodes only.
- 25 Recurrence of an invasive tumor in adjacent tissue or organ(s)) and in regional lymph nodes (both 21 and 22) at the same time.
- 26 Regional recurrence of an *in situ* tumor, NOS.
- 27 Recurrence of an *in situ* tumor in adjacent tissue or organ(s) and in regional lymph nodes at the same time.
- 30 Both regional recurrence of an invasive tumor in adjacent tissue or organ(s) and/or regional lymph nodes (20-25) and local and/or trocar recurrence (10, 13, 14, or 15).
- 36 Both regional recurrence of an *in situ* tumor in adjacent tissue or organ(s) and/or regional lymph nodes (26 or 27) and local and/or trocar recurrence (16 or 17).
- 40 Distant recurrence and there is insufficient information available to code to 46-62.
- 46 Distant recurrence of an *in situ* tumor.
- 51 Distant recurrence of an invasive tumor in the peritoneum only. Peritoneum includes peritoneal surfaces of all structures within the abdominal cavity and/or positive ascitic fluid.
- 52 Distant recurrence of an invasive tumor in the lung only. Lung includes the visceral pleura.
- 53 Distant recurrence of an invasive tumor in the pleura only. Pleura includes the pleural surface of all structures within the thoracic cavity and/or positive pleural fluid.
- 54 Distant recurrence of an invasive tumor in the liver only.
- 55 Distant recurrence of an invasive tumor in bone only. This includes bones other than the primary site.
- 56 Distant recurrence of an invasive tumor in the CNS only. This includes the brain and spinal cord, but not the external eye.
- 57 Distant recurrence of an invasive tumor in the skin only. This includes skin other than the primary site.
- 58 Distant recurrence of an invasive tumor in lymph node only. Refer to the staging scheme for a description of lymph nodes that are distant for a particular site.

- 59 Distant systemic recurrence of an invasive tumor only. This includes leukemia, bone marrow metastasis, carcinomatosis, and generalized disease.
- 60 Distant recurrence of an invasive tumor in a single distant site (51-58) and local, trocar, and/or regional recurrence (10-15, 20-25, or 30).
- 62 Distant recurrence of an invasive tumor in multiple sites (recurrences that can be coded to more than one category 51-59).
- 70 Since diagnosis, patient has never been disease-free. This includes cases with distant metastasis at diagnosis, systemic disease, unknown primary, or minimal disease that is not treated.
- 88 Disease has recurred, but the type of recurrence is unknown.
- 99 It is unknown whether the disease has recurred or if the patient was ever disease-free.

#### **RECURRENCE TYPE--1ST--OTH**

Alternate Name	Item #	Length	Source of Standard	Column #
Other Type of First Recurrence (COC)	1890	2	COC	1355-1356

#### Description

Code for an additional type of first recurrence. If more than one type of first recurrence, code one in Recurrence Type--1st [1880], and one in this field. Otherwise, this field is coded 00.

#### Codes

- 00 None, disease free
  - 01 In situ
  - 06 In situ recurrence following diagnosis of an in situ lesion of the same site
- 10 Local
  - 11 Trochar site
  - 15 Combination of 10 and 11
  - 16 Local recurrence following an *in situ* lesion of the same site
  - 17 Combination of 16 with 10, 11, and/or 15
- 20 Regional, NOS
  - 21 Regional Tissue
  - 22 Regional lymph nodes
  - 25 Combination of 21 and 22
  - 26 Regional recurrence following an *in situ* lesion of the same site
  - 27 Combination of 26 with 21, 22, and/or 25
- 30 Any combination of 10 and/or 11 and 20, 21, or 22
- 36 Any combination of recurrence following an *in situ* lesion of the same site with 10, 11, 20, 21, or 22
- 40 Distant
- 46 Distant recurrence following an *in situ* lesion of the same site
- 70 Never disease free
- 88 Recurred, site unknown
- 99 Unknown if recurred

## **REFERRAL TO SUPPORT SERV**

Alternate Name	Item #	Length	Source of Standard	Column #
Referral to Support Services (COC)	1490	1	COC	893-893

## Description

Code for whether or not patient was referred to any specified support services.

### Codes

0 No

1 Yes

9 Unknown, not specified

Note: This data item is no longer supported by COC (as of January 1, 2003).

REGIONAL NODES EXAMINED				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Number of Regional Lymph Nodes	830	2	SEER/COC	541-542
Examined (SEER)				
Pathologic Review of Regional Lymph				
Nodes (SEER)				
Regional Lymph Nodes Examined				

### Description

Part of the 10-digit EOD [779], detailed site-specific codes for anatomic EOD used by SEER for cases diagnosed from 1988 forward.

This field is included in the COC dataset, separate from EOD.

# Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

### Codes

See *SEER Extent of Disease, 1988: Codes and Coding Instructions*, Third Edition, 1998 for site-specific codes and coding rules for all EOD fields. Codes were revised effective January 1, 1998. COC codes for Regional Nodes Examined are in the *FORDS* Manual.

- 00 No regional lymph nodes examined
- 01-89 regional lymph nodes examined (code the exact number of regional lymph nodes examined)90 or more regional lymph nodes examined
- 95 No regional lymph node(s) removed but aspiration of regional lymph node(s) performed
- 96 Regional lymph node removal documented as a sampling and number of lymph nodes examined unknown/not stated

- 97 Regional lymph node removal documented as dissection and number of lymph nodes examined unknown/not stated
- 98 Regional lymph nodes surgically removed but number of lymph nodes examined unknown/not stated and not documented as sampling or dissection
- 99 Unknown; not stated; death certificate only

Note: See Chapter V, Unresolved Issues, for a discussion of coding differences between COC and SEER.

<b>REGIONAL NODES POSITIVE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Number of Positive Regional Lymph	820	2	SEER/COC	539-540
Nodes (SEER)				
Pathologic Review of Regional Lymph				
Nodes (SEER)				
Regional Lymph Nodes Positive				

### Description

Part of the 10-digit EOD [779], detailed site-specific codes for anatomic EOD used by SEER for cases diagnosed from 1988 forward.

This field is included in the COC dataset, separate from EOD.

### Rationale

Site-specific EOD codes provide extensive detail describing disease extent. The EOD codes can be grouped into different stage categories for analysis (e.g., historical summary stage categories consistent with those used in published SEER data since 1973, or more recently, AJCC stage groupings). The codes are updated as needed, but updates are usually backward compatible with old categories. See *Comparative Staging Guide for Cancer*.

### Codes

See *SEER Extent of Disease, 1988: Codes and Coding Instructions*, Third Edition, 1998 for site-specific codes and coding rules for all EOD fields. Codes were revised effective January 1, 1998 to reflect changes in the *AJCC Cancer Staging Manual*, Fifth Edition. COC codes for Regional Nodes Positive are in the *FORDS* Manual.

- 00 All regional lymph nodes examined negative
- 01-89 1-89 regional lymph nodes positive (code exact number of nodes positive)
- 90 90 or more regional lymph nodes positive
- 95 Positive aspiration of lymph node(s) was performed
- 97 Positive regional lymph nodes are documented, but number not specified
- 98 No regional lymph nodes examined
- 99 Unknown if regional lymph nodes are positive or negative; not applicable

Note: See Chapter V, Unresolved Issues, for a discussion of coding differences between COC and SEER.

# **REGISTRY ID**

Alternate Name	Item #	Length	Source of Standard	Column #
	40	10	NAACCR	20-29

# Description

A unique code assigned to each data source identifying who is sending the record and what population it is based on.

# Rationale

For registry types 2 and 3, each facility's FIN is unique. The number is essential to NCDB for monitoring data submissions, ensuring the accuracy of data, and identifying areas for special studies.

For Registry Type 1, the number notes which central registry generated the record transmission of data.

# **Instructions for Coding**

COC maintains the codes for Registry Types 2 and 3, including those for non-hospital sources of reporting. If the registry type is 1 (central registry), refer to REGID.DBF, Appendix B of this volume. If the registry type is 2 or 3, refer to FIN codes maintained by COC.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001. Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code. These sometimes are called COC FIN 10-digit codes.

For Registry Type 1, NAACCR maintains the codes for REGID.DBF.

# Codes (in addition to COC assigned codes or NAACCR assigned codes)

0000000000 Case not reported by a facility

00999999999 Case reported, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN Coding System [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

### **REGISTRY TYPE**

Alternate Name	Item #	Length	Source of Standard	Column #
	30	1	NAACCR	10-10

### Description

A computer-generated code that best describes the type of registry generating the record; used when cases are pooled from multiple registries (a hospital-based registry reporting to a state should have a "3" in this field).

### Rationale

Allows the data from multiple registries to be pooled.

### Codes

- 1 Central registry (population-based)
- 2 Central registry or hospital consortium (not population-based)
- 3 Single hospital/freestanding center

#### RELIGION

Alternate Name	Item #	Length	Source of Standard	Column #
	260	2	Varies	133-134

### Description

NAACCR has not adopted standards for this item.

## **REPORTING HOSPITAL**

Alternate Name	Item #	Length	Source of Standard	Column #
Institution ID Number (COC)	540	10	COC	382-391
Facility Identification Number (COC)				

### Description

Code for the facility reporting the tumor.

### Rationale

Each facility's FIN is unique. The number is used to identify a reporting facility in the central registry database and is useful in monitoring data submission, ensuring the accuracy of data and identifying areas for special studies. The codes for this item are assigned by COC.

## **Instructions for Coding**

COC maintains the codes, including those for non-hospital sources of reporting.

For facilities with 7-digit FINs, consisting of a constant "6" followed by 6-digit facility-specific codes in the range of 6020009-6953290 that were assigned by COC before January 1, 2001: Enter all FIN codes of this type as 3 zeroes, followed by the constant "6" and the 6-digit facility-specific codes.

For facilities with 8-digit FINs greater than or equal to 10000000 that were assigned by COC after January 1, 2001: Enter FIN codes of this type as 2 zeroes followed by the full 8-digit code.

## Codes (in addition to COC assigned codes)

0000000000 Case not reported by a facility

0099999999 Case reported, but facility number is unknown

*Note:* When this special code is being used, the length in 9s should correspond to the length indicated by the code in FIN Coding System [35]. The 9s must be right justified in the field, and the remaining spaces should be filled with leading zeroes to a total length of 10.

### **REPORTING HOSPITAL FAN**

Alternate Name	Item #	Length	Source of Standard	Column #
	538	10	COC	372-381

### Description

The facility association number (FAN) identifies country/state (3 characters), type of institution (2 characters), and facility "ownership" (5 characters).

#### Rationale

Data can be grouped for reporting from country/state, type of institution (freestanding surgery center, pathology laboratory, hospital), or institution group ID code (Kaiser, Humana, Columbia, etc.).

### Codes

COC maintains the codes. The number is entered without dashes. When used, the number reads similar to a social security number with dashes (000-00-00000), for ease of generating reports.

Note: This data item was added to the dataset in 1998, but was never used.

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### **RESERVED 00**

Alternate Name	Item #	Length	Source of Standard	Column #
	37	7		12-18

#### **RESERVED 01**

Alternate Name	Item #	Length	Source of Standard	Column #
	370	20		32-51

#### **RESERVED 02**

Alternate Name	Item #	Length	Source of Standard	Column #
	530	50		232-280

#### **RESERVED 03**

Alternate Name	Item #	Length	Source of Standard	Column #
	680	50		322-371

#### **RESERVED 04**

Alternate Name	Item #	Length	Source of Standard	Column #
	750	46		482-527

### **RESERVED 05**

Alternate Name	Item #	Length	Source of Standard	Column #
	1180	38		717-754

RESERVED 06				
Alternate Name	Item #	Length	Source of Standard	Column #
	1190	45		943-987

### **RESERVED 07**

Alternate Name	Item #	Length	Source of Standard	Column #
	1300	50		1065-1114

### **RESERVED 08**

Alternate Name	Item #	Length	Source of Standard	Column #
	1650	50		1244-1293

### **RESERVED 09**

Alternate Name	Item #	Length	Source of Standard	Column #
	1740	50		1397-1446

### **RESERVED 10**

Alternate Name	Item #	Length	Source of Standard	Column #
	1835	50		2415-2464

#### **RESERVED 11**

Alternate Name	Item #	Length	Source of Standard	Column #
	1900	50		2505-2554

#### **RESERVED 12**

Alternate Name	Item #	Length	Source of Standard	Column #
	1950	50		2595-2644

RESERVED 13				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2080			

RESERVED 14				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2210			

RESERVED 16				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2400			

RESERVED 17				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2450			

# **RESERVED 19**

Alternate Name	Item #	Length	Source of Standard	Column #
	2700	770		5925-6694

RESERVED 20				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2161			

RESERVED 21				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2371			

#### **RESERVED 22**

Alternate Name	Item #	Length	Source of Standard	Column #
	1355	1		872-872

## **RESERVED 23**

Alternate Name	Item #	Length	Source of Standard	Column #
	1635	12		920-931

### **RESERVED 24**

Alternate Name	Item #	Length	Source of Standard	Column #
	2082	16		1148-1163

### **RURALURBAN CONTINUUM 1993**

Alternate Name	Item #	Length	Source of Standard	Column #
Beale Code	3300	2	NAACCR	227-228

### Description

The "RuralUrban Continuum 1993" code, often referred to as the "Beale Code," is generated programmatically using Addr at DX--State [80] and County at DX [90]. It contains the Rural-Urban Continuum code as provided by the Office of Management and Budget (OMB) in 1993.

The code is a 10-point continuum (00-09) measuring urban-rural status. Abstractors do not enter these codes.

The code has been expanded to 2 digits to accommodate areas that are not included in the Rural Urban Continuum code table, such as Canadian provinces/territories and U.S. territories. These areas will be coded with a value of 98. Records for nonresidents of the state of reporting institution (County at DX = 998) also will be coded 98. If Addr at DX--State is XX, YY, or ZZ, the Rural Urban Continuum 93 code will be coded as 99. If County at DX equals 999, the Rural Urban Continuum 1993 code will be coded as 99.

### Rationale

RuralUrban Continuum 1993 codes are provided for each county by the OMB and consist of a 1-character rural-urban status, which is very useful for incidence and mortality data analysis.

## Codes

Metropolitan Counties (00-03)

- 00 Central counties of metropolitan areas of 1 million population or more
- 01 Fringe counties of metropolitan areas of 1 million population or more
- 02 Counties in metropolitan areas of 250,000-1,000,000 population
- 03 Counties in metropolitan areas of less than 250,000 population

Nonmetropolitan Counties (04-09)

- 04 Urban population of 20,000 or more, adjacent to a metropolitan area
- 05 Urban population of 20,000 or more, not adjacent to a metropolitan area
- 06 Urban population of 2,500-19,999, adjacent to a metropolitan area
- 07 Urban population of 2,500-19,999, not adjacent to a metropolitan area
- 08 Completely rural (no places with a population of 2,500 or more) adjacent to a metropolitan area
- 09 Completely rural (no places with a population of 2,500 or more) not adjacent to a metropolitan area
- 98 Program run, but: (1) area is not included in Rural-Urban Continuum code table, or (2) record is for resident outside of state of reporting institution
- 99 Unknown
- Blank Program not run; record not coded

### **RURALURBAN CONTINUUM 2000**

Alternate Name	Item #	Length	Source of Standard	Column #
Beale Code	3310	2	NAACCR	229-230

### Description

The "RuralUrban Continuum 2000" code, often referred to as the "Beale Code," is generated programmatically using Addr at DX--State [80] and County at DX [90]. It contains the Rural-Urban Continuum code as provided by OMB based on the 2000 Census.

The code is a 10-point continuum (00-09) measuring urban-rural status. Abstractors do not enter these codes.

The code has been expanded to 2 digits to accommodate areas that are not included in Rural Urban Continuum code table, such as Canadian provinces/territories and U.S. territories. These areas will be coded with a value of 98. Records for nonresidents of the state of reporting institution (County at DX = 998) will also be coded 98. If Addr at DX--State is XX, YY, or ZZ, the Rural Urban Continuum 2000 code will be coded as 99. If County at DX equals 999, the Rural Urban Continuum 2000 code will be coded as 99.

RuralUrban Continuum 2000 codes are provided for each county by OMB and consist of a 1-character ruralurban status, which is very useful for incidence data analysis.

### Rationale

RuralUrban Continuum 2000 codes are provided for each county by OMB and consist of a 1-character ruralurban status, which is very useful for incidence data analysis.

### Codes

Metropolitan Counties (00-03)

- 00 Central counties of metropolitan areas of 1 million population or more Counties in metropolitan areas of 250,000-1,000,000 population
- 01 Fringe counties of metropolitan areas of 1 million population or more
- 02 Counties in metropolitan areas of 250,000-1,000,000 population
- 03 Counties in metropolitan areas of less than 250,000 population

#### Nonmetropolitan Counties (04-09)

- 04 Urban population of 20,000 or more, adjacent to a metropolitan area
- 05 Urban population of 20,000 or more, not adjacent to a metropolitan area
- 06 Urban population of 2,500-19,999, adjacent to a metropolitan area
- 07 Urban population of 2,500-19,999, not adjacent to a metropolitan area
- 08 Completely rural (no places with a population of 2,500 or more) adjacent to a metropolitan area
- 09 Completely rural (no places with a population of 2,500 or more) not adjacent to a metropolitan area
- 98 Program run, but: (1) area is not included in Rural-Urban Continuum code table, or (2) record is for resident outside of state of reporting institution
- 99 Unknown
- Blank Program not run; record not coded

## **RX CODING SYSTEM--CURRENT**

Alternate Name	Item #	Length	Source of Standard	Column #
	1460	2	NAACCR	888-889

## Description

Code describing how treatment for this tumor now is coded.

### Codes

- 00 Treatment data not coded/transmitted (i.e., all treatment fields [items 1200-1450 and 1500-1645] blank\*)
- 01 Treatment data coded using 1-digit surgery codes (obsolete)
- 02 Treatment data coded according to 1983-1992 SEER manuals and 1983-1995 COC manuals
- 03 Treatment data coded according to 1996 ROADS Manual
- 04 Treatment data coded according to 1998 *ROADS* Supplement
- 05 Treatment data coded according to 1998 SEER Manual
- 06 Treatment data coded according to *FORDS* Manual
- 99 Other coding, including partial or nonstandard coding

RX DATEBRM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date Immunotherapy Started (COC)	1240	8	NAACCR	819-826

### Description

Date of initiation for immunotherapy that is part of the first course of treatment. See also RX Summ--BRM [1410]. See page 83 for date format.

#### Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first course of therapy and to reconstruct the sequence of first-course treatment modes.

### Codes (in addition to valid dates)

00000000No immunotherapy administered; autopsy-only case99999999Unknown if any immunotherapy administered; date unknown, or death certificate-only<br/>case

Note: Beginning January 1, 2003, the COC will no longer support this data item.

RX DATECHEMO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date Chemotherapy Started (COC)	1220	8	NAACCR	803-810

### Description

Date of initiation of chemotherapy that is part of the first course of treatment. See also RX Summ--Chemo [1390]. See page 83 for date format.

### Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

#### **Codes (in addition to valid dates)**

0000000	No chemotherapy administered; autopsy-only case
99999999	Unknown if any chemotherapy administered; date unknown, or death certificate only-case.

Note: This data item is no longer supported by COC (as of January 1, 2003).

### **RX DATE--DX/STG PROC**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Non Cancer-Directed Surgery	1280	8	COC	851-858
(COC)				
Date of Diagnostic, Staging or Palliative				
Procedures (1996-2002)				
Date of Surgical Diagnostic and Staging				
Procedure (COC)				
RX DateDX/Stg/Pall Proc				

### Description

Records the date on which the surgical diagnostic and/or staging procedure was performed. See Surgical and Diagnostic Staging Procedure [1350]. See page 83 for date format.

#### **Codes (in addition to valid dates)**

00000000No diagnostic or staging procedure performed; autopsy-only case99999999Unknown if any diagnostic or staging procedure performed; date unknown, or death<br/>certificate-only case

*Note:* This is a COC item and for tumors diagnosed from January 1, 1996, through December 31, 2002, may date on which diagnostic, staging, and palliative procedures were performed. Beginning with tumors diagnosed on or after January 1, 2003, palliative procedures are collected in RX Summ--Palliative Proc [3270] and RX Hosp--Palliative Proc [3280].

<b>RX DATEHORMONE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Date Hormone Therapy Started (COC)	1230	8	NAACCR	811-818

## Description

Date of initiation for hormone therapy that is part of the first course of treatment. See also RX Summ-Hormone [1400]. See page 83 for date format.

## Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

## Codes (in addition to valid dates)

00000000	No hormone therapy administered; autopsy-only case
99999999	Unknown if any hormone therapy administered; date unknown, or death certificate-only case

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **RX DATE--MOST DEFIN SURG**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Most Definitive Surgical Resection	3170	8	COC	763-770
of the Primary Site				

### Description

Date of most definitive surgical resection of the primary site performed as part of the first course of treatment. See page 83 for date format.

### Rationale

This item is used to measure lag time between diagnosis and the most definitive surgery of the primary site or survival following the procedure. It also is used in conjunction with Date of Surgical Discharge [3180] to calculate the duration of hospitalization following the most definitive primary site surgical procedure to evaluate treatment efficacy.

### Special Codes (in addition to valid dates)

- 00000000 When no surgical resection of the primary site is performed and for cases diagnosed at autopsy.
- 99999999 When it is unknown if any surgical procedure of the primary site was performed, the date is unknown or the case was identified by death certificate-only.

### **RX DATE--OTHER**

Alternate Name	Item #	Length	Source of Standard	Column #
Date Other Treatment Started (COC)	1250	8	COC	827-834

### Description

Date of initiation for other treatment that is part of the first course of treatment at any facility. See RX Summ-Other [1420]. See page 83 for date format.

### Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

#### Codes (in addition to valid dates)

00000000 No other treatment administered; autopsy-only case99999999 Unknown if any other treatment administered; date unknown, or death certificate-only case

### **RX DATE--RADIATION**

Alternate Name	Item #	Length	Source of Standard	Column #
Date Radiation Started (COC)	1210	8	COC	779-786

#### Description

Records the date on which radiation therapy began at any facility that is part of the first course of treatment. See page 83 for date format.

#### Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

#### Codes (in addition to valid dates)

00000000	No radiation therapy administered; autopsy-only case.
88888888	When radiation therapy is planned as part of the first course of therapy, but had not been
	started at the time of the most recent follow-up. The date should be revised at the next follow-
	up.
99999999	When it is unknown whether any radiation therapy was administered; the date is unknown, or
	the case was identified by death certificate-only.

## **RX DATE--RADIATION ENDED**

Alternate Name	Item #	Length	Source of Standard	Column #
Date Radiation Ended	3220	8	COC	787-794

### Description

The date on which the patient completes or receives the last radiation treatment at any facility. See page 83 for date format.

### Rationale

The length of time over which radiation therapy is administered to a patient is a factor in tumor control and treatment morbidity. It is useful in evaluating the quality-of-care and the success of patient support programs designed to maintain continuity of treatment.

### Codes (in addition to valid dates)

00000000	Radiation therapy was not administered or case diagnosed at autopsy
88888888	Radiation was administered and was ongoing at the time of most recent follow-up. The date
	should be revised at the next follow-up.
99999999	Unknown if radiation therapy was administered, or the date radiation ended is unknown.
	Death certificate-only cases.

### **RX DATE--SURGERY**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Cancer-Directed Surgery (COC)	1200	8	COC	755-762
Date of Surgery				
Date of First Surgical Procedure (COC)				

### Description

Date the first surgery of the type described under Surgery of Primary Site, Scope of Regional Lymph Node Surgery, or Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Nodes was performed. See also RX Summ--Surg Prim Site [1290], RX Summ--Scope Reg LN Sur [1292], and RX Summ--Surg Oth Reg/Dis [1294]. See page 83 for date format.

### Rationale

The dates on which different treatment modalities were started are used to evaluate whether the treatments were part of first-course therapy and to reconstruct the sequence of first-course treatment modes.

### Codes (in addition to valid dates)

00000000 No surgical procedure was performed; autopsy-only case

99999999 When it is unknown if any surgical procedure of the primary site was performed, the date is unknown or the case was identified by death certificate-only

### **RX DATE--SURGICAL DISCH**

Alternate Name	Item #	Length	Source of Standard	Column #
Date of Surgical Discharge	3180	8	COC	771-778

### Description

Records the date the patient was discharged following primary site surgery. The date corresponds to the event recorded in Surgical Procedure of Primary Site [1290], and Date of Most Definitive Surgical Resection [3170]. See page 83 for date format.

### Rationale

Length of stay is an important quality-of-care and financial measure among hospital administrations, those who fund public and private health care, and public health users. This date, in conjunction with the data item "Date of Most Definitive Surgical Resection" [3170], will allow for the calculation of a patient's length of hospitalization associated with primary site surgery.

### Special Codes (in addition to valid dates)

00000000 When no surgical treatment of the primary site was performed. Diagnosed at autopsy. 99999999 When it is unknown whether surgical treatment was performed, the date is unknown or the case was identified by death certificate only.

### **RX DATE--SYSTEMIC**

Alternate Name	Item #	Length	Source of Standard	Column #
Date Systemic Therapy Started	3230	8	COC	795-802

### Description

Date of initiation of systemic therapy that is part of the first course of treatment. Systemic therapy includes the administration of chemotherapy agents, hormone agents, biological response modifiers, bone marrow transplants, stem cell harvests, and surgical and/or radiation endocrine therapy. See page 83 for date format.

### Rationale

Collecting dates for each treatment modality allows the sequencing of multiple treatments and aids in the evaluation of time intervals from diagnosis to treatment and from treatment to recurrence.

### Codes (in addition to valid dates)

- When no systemic therapy was administered, or the case was diagnosed at autopsy.
   When systemic therapy is planned as part of the first course of therapy, but had not been started at the time of the most recent follow-up. The date should be revised at the next follow-up.
   When it is unknown if any systemic therapy was administered, the date is unknown, or the
- When it is unknown if any systemic therapy was administered, the date is unknown, or the case was identified by death certificate-only.

## RX HOSP--BRM

Alternate Name	Item #	Length	Source of Standard	Column #
Immunotherapy at this Facility (COC)	720	2	COC	468-469

## Description

Records whether immunotherapeutic agents (biologic response modifiers) were administered as first-course treatment at this facility or the reason they were not given. Immunotherapy consists of biological or chemical agents that alter the immune system or change the host's response to tumor cells.

## Rationale

Systemic therapy may involve the administration of one or a combination of agents. This data item allows for the evaluation of the administration of immunotherapeutic agents as part of the first course of therapy.

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes (refer to FORDS for additional instructions)

- 00 None, immunotherapy was not part of the first course of therapy; not customary therapy for this cancer.
- 01 Immunotherapy
- 82 Immunotherapy was not recommended/administered because it was contraindicated due to patient risk factors (comorbid conditions, advanced age, etc.).
- 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 first-course therapy. No reason was noted in the patient record.
- 87 Immunotherapy was not administered; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 88 Immunotherapy was recommended, but it is unknown if it was administered.
- 99 It is unknown if immunotherapy was recommended or administered; death certificate-only cases.

*Note:* For tumors diagnosed on or after January 1, 2003, information on bone marrow transplants and stem cell transplants should be coded in the new field RX Summ--Transplnt/Endocr [3250]. Codes 02-06 should not be used for tumors diagnosed on or after January 1, 2003.

### **RX HOSP--CHEMO**

Alternate Name	Item #	Length	Source of Standard	Column #
Chemotherapy at this Facility (COC)	700	2	COC	464-465

### Description

Defines the type of chemotherapy the patient received as a part of the initial treatment for the reportable tumor at the reporting facility or the reason chemotherapy was not given.

### Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

### Codes (refer to FORDS for additional instructions)

- 00 None, chemotherapy was not part of the first course of therapy; not customary therapy for this cancer.
- 01 Chemotherapy, NOS
- 02 Chemotherapy, single agent
- 03 Chemotherapy, multiple agents
- 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 first-course therapy. No reason was stated in the patient record.
- 87 Chemotherapy was not administered; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 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 cases.

# RX HOSP--DX/STG PROC

Alternate Name	Item #	Length	Source of Standard	Column #
Non Cancer-Directed Surgery at this	740	2	COC	471-472
Facility (COC)				
Surgical Diagnostic & Staging Procedure				
at this Facility (1996-2002)				
RX HospDX/Stg/Pall Proc				

# Description

Identifies the surgical procedure(s) performed in an effort to diagnose and/or stage disease at this facility. Used for cases diagnosed in 1996 and later. Earlier data may be converted into this field. See also RX Hosp--Surg Prim Site [670].

# Rationale

If central registries wish to study the procedures given at particular hospitals, the hospital-level fields must be used. The summary fields, conversely, combine information across all hospitals that provide for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

# Codes (refer to FORDS for additional instructions)

- 00 No surgical diagnostic or staging procedure was performed.
- 01 A biopsy (incisional, needle, or aspiration) was done to a site other than the primary site. No exploratory procedure was done.
- 02 A biopsy (incisional, needle, or aspiration) was done of the primary site.
- 03 A surgical exploratory only. The patient was not biopsied or treated.
- 04 A surgical procedure with a bypass was performed, but no biopsy was done.
- 05 An exploratory procedure was performed, and a biopsy of either the primary site or another site was done.
- 06 A bypass procedure was performed, and a biopsy of either the primary site or another site was done.
- 07 A procedure was done, but the type of procedure is unknown.
- 09 No information about whether a diagnostic or staging procedure was performed.

*Note:* This item has been used for tumors diagnosed in 1996 and later. For cases diagnosed before 1996, this item may have been converted, and cases with cancer-directed surgery would have been converted to 09 in this field. For cases diagnosed between 1996 and 2002, this field may have described palliative care. For tumors diagnosed on or after January 1, 2003 palliative care is coded in a new field RX Hosp--Palliative Proc [3280].

### **RX HOSP--HORMONE**

Alternate Name	Item #	Length	Source of Standard	Column #
Hormone Therapy at this Facility (COC)	710	2	COC	466-467

### Description

Records whether systemic hormonal agents were administered as first-course treatment at this facility or the reason they were not given. 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.

### Rationale

Systemic therapy may involve the administration of one or a combination of agents. This data item allows for the evaluation of the administration of hormonal agents as part of the first course of therapy. If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

### Codes (refer to FORDS for additional instructions)

- 00 None, hormone therapy was not part of the first course of therapy.
- 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 first-course therapy. No reason was stated in the patient record.
- 87 Hormone therapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 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 the patient record; death certificate-only cases.

*Note:* Any therapy codes 02-03 should have been converted to the appropriate code in the new field RX Summ--Transplnt/Endocr [3250]. Codes 02-03 should not be used for tumors diagnosed on or after January 1, 2003.

### **RX HOSP--OTHER**

Alternate Name	Item #	Length	Source of Standard	Column #
Other Treatment at this Facility (COC)	730	1	COC	470-470

## Description

Identifies other treatment given at this facility that cannot be defined as surgery, radiation, or systemic therapy according to the defined data items in this manual. 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, transfusions, and aspirin.

## Rationale

Information on other therapy is used to describe and evaluate the quality-of-care and treatment practices. If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

### Codes

- 0 None
- 1 Other
- 2 Other Experimental
- 3 Other-Double Blind.
- 6 Other-Unproven
- 7 Refusal
- 8 Recommended; unknown if administered.
- 9 Unknown

*Note:* Aspirin (also known as acetylsalicylic acid [ASA] or by a brand name) is used as a treatment for essential thrombocythemia. Record ONLY aspirin therapy to thin the blood for symptomatic control of thrombocythemia. 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 325-1,000mg 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.

Phlebotomy may be called blood removal, bloodletting, or venisection. Transfusions may include whole blood, red blood cells, platelets, plateletpheresis, fresh frozen plasma, plasmapheresis, and cryoprecipitate.

## **RX HOSP--PALLIATIVE PROC**

Alternate Name	Item #	Length	Source of Standard	Column #
Palliative Procedure at this Facility	3280	1	COC	473-473

## Description

Identifies any procedure performed at the reporting facility in an effort to palliate or alleviate symptoms. Palliative procedures are performed to relieve symptoms and may include surgery, radiation therapy, systemic therapy (chemotherapy, hormone therapy, or other systemic drugs), and/or pain management therapy.

## Rationale

This data item allows reporting facilities to track the use of procedures that are considered palliative rather than therapeutic, diagnostic, or staging.

## Codes

- 0 No palliative care provided
- 1 Surgery (which may involve a bypass procedure) performed to alleviate symptoms, but no attempt to diagnose, stage, or treat the primary tumor is made
- 2 Radiation therapy given to alleviate symptoms, but no attempt to diagnose, stage, or treat the primary tumor is made
- 3 Chemotherapy, hormone therapy, or other systemic drugs given to alleviate symptoms, but no attempt to diagnose, stage or treat the primary tumor is made
- 4 Pain management therapy with no other palliative care
- 5 Any combination of codes 1, 2, and/or 3 without code 4
- 6 Any combination of codes 1, 2 and/or 3 with code 4
- 7 Palliative care was performed, but no information on the type of procedure is available in the patient record
- 9 Unknown if palliative care was performed; not stated in patient record.

<b>RX HOSPRADIATION</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Radiation at this Facility (COC)	690	1	SEER/COC	463-463

## Description

Defines the type of radiation therapy the patient received as a part of the initial treatment for the reportable tumor at the reporting facility.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

### Codes

- 0 None
- 1 Beam radiation
- 2 Radioactive implants
- 3 Radioisotopes
- 4 Combination of 1 with 2 or 3
- 5 Radiation, NOS—method or source not specified
- 9 Unknown if radiation therapy administered

Note: This data item is no longer supported by COC (as of January 1, 2003).

# RX HOSP--REG LN REMOVED

Alternate Name	Item #	Length	Source of Standard	Column #
Number of Regional Lymph Nodes	676	2	COC	461-462
Examined at this Facility (COC)				
RX HospReg LN Examined				

## Description

Describes number of regional lymph nodes removed as part of the first course of treatment. This item reflects that portion of the first course of treatment given at the reporting facility.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing the extent of treatment given at a particular hospital also helps resolve coding issues.

## Codes

- 00 No regional lymph nodes removed
- 01 One regional lymph node removed
- 02 Two regional lymph nodes removed
- ••
- 90 Ninety or more regional lymph nodes removed
- 95 No regional lymph node(s) removed, but aspiration of regional lymph node(s) was performed
- 96 Regional lymph node removal documented as a sampling and number of lymph nodes unknown/not stated
- 97 Regional lymph node removal documented as a dissection and number of lymph 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
- 99 Unknown; not stated; death certificate-only

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

# RX HOSP--SCOPE REG 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Scope of Regional Lymph Node Surgery at	747	1	COC	480-480
this Facility (COC)				

# Description

Describes the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event at the reporting facility. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Scope of Regional Lymph Node Surgery at the reporting facility for all tumors diagnosed before January 1, 2003.

# Rationale

In evaluating quality-of-care and treatment practices it is important to identify the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event.

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing the extent of treatment given at a particular hospital also helps resolve coding issues.

# Codes

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the SEER Program Code Manual, RX Coding System [1460] code 5, 1998 for site-specific codes.

# RX HOSP--SCOPE REG LN SUR

Alternate Name	Item #	Length	Source of Standard	Column #
Scope of Regional Lymph Node Surgery at	672	1	COC	459-459
this Facility (COC)				

# Description

Describes the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event at the reporting facility.

# Rationale

In evaluating quality-of-care and treatment practices it is important to identify the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event.

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing the extent of treatment given at a particular hospital also helps resolve coding issues.

# Codes (refer to FORDS for additional instructions)

- 0 No regional lymph nodes removed
- 1 Biopsy or aspiration of regional lymph node, NOS
- 2 Sentinel lymph node biopsy
- 3 Number of regional lymph nodes removed unknown, not stated; regional lymph nodes removed, NOS
- 4 1 to 3 regional lymph nodes removed
- 5 4 or more regional lymph nodes removed
- 6 Sentinel node biopsy and code 3, 4, or 5 at same time or timing not stated
- 7 Sentinel node biopsy and code 3, 4, or 5 at different times
- 9 Unknown or not applicable

*Note:* One important use of registry data is the tracking of treatment patterns over time. To compare contemporary treatment to previously published treatment based on former codes, or to data unmodified from pre-1998 definitions, the ability to differentiate surgeries in which four or more regional lymph nodes are removed is desirable. However, it is very important to note that the distinction between codes 4 and 5 is made to permit comparison of current surgical procedures with procedures coded in the past when the removal of fewer than 4 nodes was not reflected in surgery codes. It is not intended to reflect clinical significance when applied to a particular surgical procedure. It is important to avoid inferring, by data presentation or other methods, that one category is preferable to another within the intent of these items.

# RX HOSP--SCREEN/BX PROC1

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	742	1	COC	474-474
2001 COC)				
RX HospDiag/Stage Proc1 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for primary site biopsy procedures.

# Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

# Codes

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **RX HOSP--SCREEN/BX PROC2**

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	743	1	COC	475-475
2001 COC)				
RX HospDiag/Stage Proc2 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for use of guidance procedures for the primary site biopsy.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

# RX HOSP--SCREEN/BX PROC3

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	744	1	COC	476-476
2001 COC)				
RX HospDiag/Stage Proc3 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for palpability of a breast primary or the approach for a prostate primary site biopsy.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **RX HOSP--SCREEN/BX PROC4**

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	745	1	COC	477-477
2001 COC)				
RX HospDiag/Stage Proc4 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for first detection of a breast primary or a non-primary site biopsy for a prostate primary.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

# RX HOSP--SURG OTH 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Surgery of Other Regional Site(s), Distant	748	1	COC	481-481
Site(s), or Distant Lymph Node(s) at this				
Facility (COC)				
Surgical Procedure/Other Site at this				
Facility				

# Description

Records the surgical removal of distant lymph nodes or other tissue(s)/organ(s) beyond the primary site at this facility. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Surgery Other Regional/Distant Sites at the reporting facility for all tumors diagnosed before January 1, 2003.

# Rationale

The removal of non-primary tissue documents the extent of surgical treatment and is useful in evaluating the extent of metastatic involvement. If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

# Codes

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the SEER Program Code Manual, RX Coding System [1460] code 5, 1998 for site-specific codes.

# RX HOSP--SURG OTH REG/DIS

Alternate Name	Item #	Length	Source of Standard	Column #
Surgery of Other Regional Site(s), Distant	674	1	COC	460-460
Site(s), or Distant Lymph Node(s) at this				
Facility (COC)				
Surgical Procedure/Other Site at this				
Facility				

# Description

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

## Rationale

The removal of non-primary tissue documents the extent of surgical treatment and is useful in evaluating the extent of metastatic involvement. If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes (refer to FORDS for additional instructions)

- 0 None
- 1 Non-primary surgical procedure performed
- 2 Non-primary surgical procedure to other regional sites
- 3 Non-primary surgical procedure to distant lymph node(s)
- 4 Non-primary surgical procedure to distant site
- 5 Any combination of codes 2, 3, or 4
- 9 Unknown

# **RX HOSP--SURG PRIM SITE**

Alternate Name	Item #	Length	Source of Standard	Column #
Cancer-Directed Surgery at this Facility	670	2	COC	457-458
(pre-96 COC)				
RX HospCA Dir Surgery (pre-96				
NAACCR)				
Surgical Procedure of Primary Site				

# Description

Describes surgical procedures used to treat the primary site of the reportable tumor. This item records that portion of the first course of treatment given at the reporting facility. See Chapter V, Unresolved Issues, for a discussion of differences in treatment coding among groups and over time.

# Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

# Codes in addition to the site-specific codes (refer to FORDS for additional instructions)

- 00 None. No surgical procedure of primary site. Autopsy only.
- 10-19 Site-specific codes. Tumor destruction; no pathologic specimen produced.
- 20-80 Site-specific codes. Resection. Path specimen produced.
- 90 Surgery, NOS
- 98 Site specific codes; special
- 99 Unknown. Death certificate-only.

## RX HOSP--SURG SITE 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Cancer-Directed Surgery at this Facility	746	2	COC	478-479
(pre-96 COC)				
RX HospCA Dir Surgery (pre-96				
NAACCR)				
Surgical Procedure of Primary Site				

# Description

Describes surgical procedures used to treat the primary site of the reportable tumor. This item records that portion of the first course of treatment given at the reporting facility. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Surgery Primary Site at the reporting facility for all tumors diagnosed before January 1, 2003. See Chapter V, Unresolved Issues, for a discussion of differences in treatment coding among groups and over time.

## Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes (in addition to the site-specific codes)

- 00 No cancer-directed surgery performed.
- 99 Unknown if cancer-directed surgery performed.

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the *SEER Program Code Manual*, RX Coding System [1460] code 5, 1998 for site-specific codes.

# RX SUMM--BRM

Alternate Name	Item #	Length	Source of Standard	Column #
Immunotherapy (SEER/COC)	1410	2	SEER/COC	882-883
Biological Response Modifiers (pre-96				
SEER)				

## Description

Records whether immunotherapeutic (biologic response modifiers) agents were administered as first-course treatment at this facility or the reason they were not given. Immunotherapy consists of biological or chemical agents that alter the immune system or change the host's response to tumor cells.

## Rationale

Systemic therapy may involve the administration of one or a combination of agents. This data item allows for the evaluation of the administration of immunotherapeutic agents as part of the first course of therapy.

## Codes (refer to FORDS and the SEER Program Code Manual for additional instructions)

- 00 None, immunotherapy was not part of the planned first course of therapy.
- 01 Immunotherapy administered as first course therapy.
- 82 Immunotherapy was not recommended/administered because it was contraindicated due to patient risk factors (i.e., comorbid conditions, advanced age).
- 85 Immunotherapy was not administered because the patient died prior to planned or recommended therapy.
- 86 Immunotherapy was not administered. It was recommended by the patient's physician, but was not administered as part of first-course therapy. No reason was stated in the patient record.
- 87 Immunotherapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 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 cases.

## Instructions for Storing/Converting Historical Codes

SEER recommends that the 1-digit historical codes be stored in the second character position preceded by a zero. COC recommends that the historic codes be converted to the current codes, using the algorithm it has developed.

Historically (before 2003), this was a 1-character field with the following codes:

- 0 None
- 1 Biological response modifier
- 2 Bone marrow transplant—autologous
- 3 Bone marrow transplant—allogeneic
- 4 Bone marrow transplant, NOS
- 5 Stem cell transplant
- 6 Combination of 1 and any 2, 3, 4 or 5
- 7 Patient or patient's guardian refused
- 8 Biological response modifier recommended, unknown if administered
- 9 Unknown if immunotherapy given

*Note:* For tumors diagnosed on or after January 1, 2003, information on bone marrow transplants and stem cell transplants should be coded in the new field, RX Summ--Transplnt/Endocr [3250]. The COC standards for hospitals do not allow use of codes 02-06 in tumors diagnosed on or after January 1, 2003.

RX SUMMCHEMO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Chemotherapy (SEER/COC)	1390	2	SEER/COC	878-879

## Description

Codes for chemotherapy given as part of the first course of treatment or the reason chemotherapy was not given. Includes treatment given at all facilities as part of the first course.

### Codes (refer to FORDS for additional instructions)

- 00 None, chemotherapy was not part of the planned first course of therapy.
- 01 Chemotherapy, NOS
- 02 Chemotherapy, single agent.
- 03 Chemotherapy, multiple agents.
- 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 first-course therapy. No reason was stated in the patient record.
- 87 Chemotherapy was not administered; it was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 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 cases.

# RX SUMM--DX/STG PROC

Alternate Name	Item #	Length	Source of Standard	Column #
Non Cancer-Directed Surgery (COC)	1350	2	COC	869-870
Surgical Diagnostic and Staging Procedure				
(1996-2002)				
RX SummDX/Stg/Pall Proc				

# Description

Identifies the surgical procedure(s) performed in an effort to diagnose and/or stage disease. COC recommends this item for tumors diagnosed 1996 and forward. For tumors diagnosed before 1996, this item may have been converted, and tumors with cancer-directed surgery would have been converted to 09 in this field. See also RX Summ--Surg Prim Site [1290] and RX Summ--Reconstruct 1st [1330]. For SEER and pre-1996 COC, see RX Summ--Surgery Type [1640].

# Codes (refer to FORDS for additional instructions)

- 00 No surgical diagnostic or staging procedure was performed.
- 01 A biopsy (incisional, needle, or aspiration) was done to a site other than the primary site. No exploratory procedure was done.
- 02 A biopsy (incisional, needle, or aspiration) was done of the primary site.
- 03 A surgical exploratory only. The patient was not biopsied or treated.
- 04 A surgical procedure with a bypass was performed, but no biopsy was done.
- 05 An exploratory procedure was performed, and a biopsy of either the primary site or another site was done.
- 06 A bypass procedure was performed, and a biopsy of either the primary site or another site was done.
- 07 A procedure was done, but the type of procedure is unknown.
- 09 No information about whether a diagnostic or staging procedure was performed.

*Note:* This item has been used for tumors diagnosed in 1996 and later. For tumors diagnosed before 1996, tumors with cancer-directed surgery would have been converted to 09 in this field. For tumors diagnosed between 1996 and 2002 this field may have described palliative care. For tumors diagnosed on or after January 1, 2003 palliative care is coded in a new field RX Summ--Palliative Proc [3270].

RX SUMMHORMONE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Hormone Therapy (SEER/COC)	1400	2	SEER/COC	880-881
Endocrine (Hormone/Steroid) Therapy				
(pre-96 SEER)				

## Description

Records whether systemic hormonal agents were administered as first-course treatment at any facility, or the reason they were not given. 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.

## Rationale

Systemic therapy may involve the administration of one or a combination of agents. This data item allows for the evaluation of the administration of hormonal agents as part of the first course of therapy.

## Codes (refer to FORDS and the SEER Program Code Manual for additional instructions)

- 00 None, hormone therapy was not part of the planned first course of therapy.
- 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 first-course therapy. No reason was stated in the patient record.
- 87 Hormone therapy was not administered. It was recommended by the patient's physician, but this treatment was refused by the patient, the patient's family member, or the patient's guardian. The refusal was noted in the patient record.
- 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 the patient record. Death certificate-only cases.

\**Note:* For COC, codes 7 and 8 were used for tumors diagnosed before 1996, but should have been converted to 0 in this field and to the appropriate code in the new field Reason for No Hormone [1450]. The COC standards for hospitals do not allow use of codes 7 and 8 in 1996 and later. SEER continues to use codes 7 and 8 for all years. See Chapter V, Unresolved Issues, for further discussion.

*Note:* For tumors diagnosed on or after January 1, 2003, information on endocrine surgery and/or endocrine radiation should be coded in the new field, RX Summ--Transplnt/Endocr [3250]. The COC standards for hospitals do not allow use of codes 02-03 in tumors diagnosed on or after January 1, 2003.

## **RX SUMM--OTHER**

Alternate Name	Item #	Length	Source of Standard	Column #
Other Treatment (COC)	1420	1	SEER/COC	884-884
Other Cancer-Directed Therapy				
(SEER/pre-96 COC)				

## Description

Identifies other treatment given at this facility that cannot be defined as surgery, radiation, or systemic therapy according to the defined data items in this manual. 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, transfusions, and aspirin.

## Rationale

Information on other therapy is used to describe and evaluate the quality-of-care and treatment practices.

## Codes (refer to FORDS for additional coding instructions)

- 0 None
- 1 Other
- 2 Other Experimental
- 3 Other-Double Blind
- 6 Other-Unproven
- 7 Refusal
- 8 Recommended
- 9 Unknown; unknown if administered

## **RX SUMM--PALLIATIVE PROC**

Alternate Name	Item #	Length	Source of Standard	Column #
Palliative Procedure	3270	1	COC	871-871

## Description

Identifies any procedure performed in an effort to palliate or alleviate symptoms. Palliative procedures are performed to relieve symptoms and may include surgery, radiation therapy, systemic therapy (chemotherapy, hormone therapy, or other systemic drugs), and/or pain management therapy.

## Rationale

This data item allows the tracking of the use of procedures that are considered palliative rather than therapeutic, diagnostic, or staging.

## Codes

- 0 No palliative care provided.
- 1 Surgery (which may involve a bypass procedure) to alleviate symptoms, but no attempt to diagnose, stage, or treat the primary tumor is made.
- 2 Radiation therapy to alleviate symptoms, but no attempt to diagnose, stage, or treat the primary tumor is made.
- 3 Chemotherapy, hormone therapy, or other systemic drugs to alleviate symptoms, but no attempt to diagnose, stage, or treat the primary tumor is made.
- 4 Patient received or was referred for pain management therapy with no other palliative care.
- 5 Any combination of codes 1, 2, and/or 3 without code 4.
- 6 Any combination of codes 1, 2 and/or 3 with code 4.
- 7 Palliative care was performed or referred, but no information on the type of procedure is available in the patient record.
- 9 Unknown if palliative care was performed or referred; not stated in patient record.

# RX SUMM--RAD TO CNS

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Therapy to CNS (COC)	1370	1	SEER/COC	874-874
Radiation to the Brain and/or Central				
Nervous System (SEER)				

# Description

For lung and leukemia cases only, codes for radiation given to the brain or central nervous system. Includes treatment given at all facilities as part of the first course. See Chapter V, Unresolved Issues, for more information.

*Note:* SEER does not collect this data item beginning with 1998 cases. They retain the codes for older cases in this field, and they have also recoded radiation coded here as radiation in RX Summ--Radiation [1360]. COC does not collect this data item beginning with 1996 cases.

# Codes

For Lung and Leukemia Cases only:

- 0 No radiation to the brain and/or central nervous system
- 1 Radiation
- 7 Patient or patient's guardian refused
- 8 Radiation recommended, unknown if administered
- 9 Unknown
- For all other cases (primaries other than lung or leukemia):
- 9 Not applicable

## **RX SUMM--RADIATION**

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation (SEER/COC)	1360	1	SEER	873-873
Radiation Therapy (pre-96 COC)				

# Description

Codes for the type of radiation therapy performed as part of the first course of treatment.

*Note:* Radiation to brain and central nervous system for leukemia and lung cases is coded as radiation in this field.

## Codes

- 0 None
- 1 Beam radiation
- 2 Radioactive implants
- 3 Radioisotopes
- 4 Combination of 1 with 2 or 3
- 5 Radiation, NOS—method or source not specified
- 7 Patient or patient's guardian refused\*
- 8 Radiation recommended, unknown if administered\*
- 9 Unknown if radiation administered

\**Note:* For COC, codes 7 and 8 were used for tumors diagnosed before 1996, but should have been converted to 0 in this field and to the appropriate code in the new field Reason for No Radiation [1430]. The COC standards for hospitals do not allow use of codes 7 and 8 in 1996 and later. SEER continues to use codes 7 and 8 for all years. See Chapter V, Unresolved Issues, for further discussion.

RX SUMMRECONSTRUCT 1ST				
Alternate Name	Item #	Length	Source of Standard	Column #
ReconstructionFirst Course (SEER)	1330	1	SEER	867-867
Reconstruction/RestorationFirst Course				
(COC)				

## Description

Codes for surgical procedures done to reconstruct, restore, or improve the shape and appearance or function of body structures that are missing, defective, damaged, or misshapen by cancer or cancer-directed therapies. Reconstructive/restorative procedures are coded here when started during the first course of cancer-directed therapy.

COC introduced site-specific codes for this item in the COC *ROADS Manual* 1998 Supplement. RX Coding System--Current [1460] identifies which coding system applies.

SEER collects reconstructive procedures for breast cancer tumors only.

For reconstructive/restorative procedures performed later, see Subseq RX--Reconstruct Del [1741]. See also RX Summ--Surgery Type [1640].

# RX SUMM--REG LN EXAMINED

Alternate Name	Item #	Length	Source of Standard	Column #
Number of Regional Lymph Nodes	1296	2	SEER/COC	863-864
Examined (SEER/COC)				
Number of Regional Lymph Nodes				
Removed (COC)				

# Description

Codes for the number of regional lymph nodes examined in conjunction with surgery performed as part of the first-course treatment. This includes treatment given at all facilities as part of the first course of treatment. See also RX Summ--Scope Reg LN Sur [1292].

# Codes

- 00 No regional lymph nodes examined
- 01 One regional lymph node examined
- 02 Two regional lymph nodes examined
- ••

90 90 or more regional lymph nodes examined

- 95 No regional lymph node(s) removed, but aspiration of regional lymph node(s) was performed
- 96 Regional lymph node removal documented as sampling, and number of lymph nodes unknown/not stated
- 97 Regional lymph node removal documented as a dissection, and number of lymph 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
- 99 Unknown; not stated; death certificate-only

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

# RX SUMM--SCOPE REG 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Scope of Regional Lymph Node Surgery	1647	1	SEER/COC	941-941
(SEER/COC)				

# Description

Describes the removal, biopsy or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event at all facilities. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Scope of Regional Lymph Node Surgery at all facilities for all tumors diagnosed before January 1, 2003.

# Rationale

In evaluating quality-of-care and treatment practices it is important to identify the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event.

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing the extent of treatment given at a particular hospital also helps resolve coding issues.

# Codes

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the SEER Program Code Manual, RX Coding System [1460] code 5, 1998 for site-specific codes.

# **RX SUMM--SCOPE REG LN SUR**

Alternate Name	Item #	Length	Source of Standard	Column #
Scope of Regional Lymph Node Surgery	1292	1	SEER/COC	861-861
(SEER/COC)				

# Description

Describes the removal, biopsy or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event at all facilities.

# Rationale

In evaluating quality-of-care and treatment practices it is important to identify the removal, biopsy, or aspiration of regional lymph node(s) at the time of surgery of the primary site or during a separate surgical event.

# Codes (refer to FORDS and SEER Program Code Manual for additional instructions)

- 0 None
- 1 Biopsy or aspiration of regional lymph node, NOS
- 2 Sentinel lymph node biopsy
- 3 Number of regional lymph nodes removed unknown, not stated; regional lymph nodes removed, NOS
- 4 1 to 3 regional lymph nodes removed
- 5 4 or more regional lymph nodes removed
- 6 Sentinel node biopsy and code 3, 4, or 5 at same time or timing not noted.
- 7 Sentinel node biopsy and code 3, 4, or 5 at different times
- 9 Unknown or not applicable

*Note:* One important use of registry data is the tracking of treatment patterns over time. To compare contemporary treatment to previously published treatment based on former codes, or to data unmodified from pre-1998 definitions, the ability to differentiate surgeries in which four or more regional lymph nodes are removed is desirable. However, it is very important to note that the distinction between codes 4 and 5 is made to permit comparison of current surgical procedures with procedures coded in the past when the removal of fewer than 4 nodes was not reflected in surgery codes. It is not intended to reflect clinical significance when applied to a particular surgical procedure. It is important to avoid inferring, by data presentation or other methods, that one category is preferable to another within the intent of these items.

# RX SUMM--SCREEN/BX PROC1

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	1642	1	COC	934-934
2001 COC)				
RX SummDiag/Stage Proc1 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for primary site biopsy procedure.

## Codes

For detailed site-specific 1-digit codes for each field for breast and prostate tumors, see the COC *ROADS Manual*, 1998 Supplement.

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

## RX SUMM--SCREEN/BX PROC2

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	1643	1	COC	935-935
2001 COC)				
RX SummDiag/Stage Proc2 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for use of guidance procedures for the primary site biopsy.

## Codes

For detailed site-specific 1-digit codes for each field for breast and prostate cases, see the COC *ROADS Manual*, 1998 Supplement.

For all cases other than breast and prostate:

0 Not applicable

# RX SUMM--SCREEN/BX PROC3

Alternate Name	Item #	Length	Source of Standard	Column #	
Diagnostic and Staging Procedures (pre-	1644	1	COC	936-936	
2001 COC)					
RX SummDiag/Stage Proc3 (pre-2001)					
Screening or Biopsy Procedures (COC)					

# Description

Site-specific field with codes for palpability of a breast primary or the approach for a prostate primary site biopsy.

# Codes

For detailed site-specific 1-digit codes for each field for breast and prostate tumors, see the COC *ROADS Manual*, 1998 Supplement.

For all tumors other than breast and prostate:

0 Not applicable

Note: This data item is no longer supported by COC (as of January 1, 2003).

## **RX SUMM--SCREEN/BX PROC4**

Alternate Name	Item #	Length	Source of Standard	Column #
Diagnostic and Staging Procedures (pre-	1645	1	COC	937-937
2001 COC)				
RX SummDiag/Stage Proc4 (pre-2001)				
Screening or Biopsy Procedures (COC)				

## Description

Site-specific field with codes for first detection of a breast primary or a non-primary site biopsy for a prostate primary.

# Codes

For detailed site-specific 1-digit codes for each field for breast and prostate tumors, see the COC *ROADS Manual*, 1998 Supplement.

For all tumors other than breast and prostate:

0 Not applicable

# RX SUMM--SURG OTH 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Surgery of Other Regional Site(s), Distant	1648	1	SEER/COC	942-942
Site(s) or Distant Lymph Nodes				
(SEER/COC)				
Surgical Procedure/Other Site				

# Description

Records the surgical removal of distant lymph nodes or other tissue(s)/organ(s) beyond the primary site given at all facilities as part of the first course of treatment. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Surgery Regional/Distant Sites at all facilities for all tumors diagnosed before January 1, 2003.

## Rationale

The removal of non-primary tissue documents the extent of surgical treatment and is useful in evaluating the extent of metastatic involvement. If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

## Codes

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the SEER Program Code Manual, RX Coding System [1460] code 5, 1998 for site-specific codes.

## **RX SUMM--SURG OTH REG/DIS**

Alternate Name	Item #	Length	Source of Standard	Column #
Surgery of Other Regional Site(s), Distant	1294	1	SEER/COC	862-862
Site(s) or Distant Lymph Nodes				
(SEER/COC)				
Surgical Procedure/Other Site				

## Description

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

## Rationale

The removal of non-primary tissue documents the extent of surgical treatment and is useful in evaluating the extent of metastatic involvement.

## Codes (refer to FORDS and SEER Program Code Manual for additional instructions)

- 0 None; diagnosed at autopsy
- 1 Non-primary surgical procedure performed
- 2 Non-primary surgical procedure to other regional sites
- 3 Non-primary surgical procedure to distant lymph node(s)
- 4 Non-primary surgical procedure to distant site
- 5 Any combination of codes 2, 3, or 4
- 9 Unknown; death certificate only

# **RX SUMM--SURG PRIM SITE**

Alternate Name	Item #	Length	Source of Standard	Column #
Cancer-Directed Surgery (pre-96 COC)	1290	2	SEER/COC	859-860
Surgery of Primary Site (SEER/COC)				

# Description

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

# Codes (in addition to the site-specific codes; refer to *FORDS* and *SEER Program Code Manual* for additional instructions)

00 None.

- 10-19 Site-specific codes. Tumor destruction
- 20-80 Site-specific codes. Resection.
- 90 Surgery, NOS
- 98 Site specific codes; special
- 99 Unknown

# RX SUMM--SURG/RAD SEQ

Alternate Name	Item #	Length	Source of Standard	Column #
Radiation Sequence with Surgery (pre-96	1380	1	SEER/COC	875-875
SEER/COC)				
Radiation/Surgery Sequence (COC)				

## Description

Codes for the sequencing of radiation and surgery given as part of the first course of treatment. See also RX Summ--Surg Prim Site [1290], RX Summ--Scope LN Surg [1292], RX Summ--Surg Oth Reg/Dis [1294], and RX Summ--Radiation [1360].

## Codes

- 0 No radiation and/or no cancer-directed surgery
- 2 Radiation before surgery
- 3 Radiation after surgery
- 4 Radiation both before and after surgery
- 5 Intraoperative radiation
- 6 Intraoperative radiation with other radiation given before or after surgery
- 9 Sequence unknown, but both surgery and radiation were given

## **RX SUMM--SURGERY TYPE**

Alternate Name	Item #	Length	Source of Standard	Column #
SiteSpecific Surgery (pre-98 SEER)	1640	2	SEER	932-933

## Description

Field for pre-1996 surgery codes for COC and pre-1998 surgery codes for SEER. Surgery codes used 1998 and later can be backward converted into the older codes and the converted value can be stored in this field. See Chapter V, Unresolved Issues, for discussion of COC/SEER differences in coding treatment.

#### **RX SUMM--SURGICAL APPROCH**

Alternate Name	Item #	Length	Source of Standard	Column #
Surgical Approach (COC)	1310	1	COC	865-865

#### Description

Codes for method used to approach the surgical field for the primary site. COC requires coding for tumors diagnosed 1996 and forward. COC introduced site-specific codes for this item in the COC *ROADS Manual* 1998 Supplement. See also item RX Summ--Surg Prim Site [1290].

#### Codes

See the COC ROADS Manual, 1998 Supplement, for site-specific codes.

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

#### **RX SUMM--SURGICAL MARGINS**

Alternate Name	Item #	Length	Source of Standard	Column #
Surgical Margins (COC)	1320	1	COC	866-866
Residual Primary Tumor Following				
Cancer-Directed Surgery (pre-96 COC)				

#### Description

Codes describe the final status of surgical margins after resection of the primary tumor. See also RX Summ--Surg Prim Site [1290].

#### Rationale

This item serves as a quality measure for pathology reports, is used for staging, and may be a prognostic factor in recurrence. This item is not limited to cases that have been staged. It applies to all cases that have a surgical procedure of the primary site.

#### Codes (refer to FORDS for additional instructions)

- 0 No residual tumor
- 1 Residual tumor, NOS
- 2 Microscopic residual tumor
- 3 Macroscopic residual tumor
- 7 Margins not evaluable.
- 8 No primary site surgery.
- 9 Unknown or not applicable.

Note: Codes were site specific (1998-2002), and have been changed to be generic across all disease sites.

# RX SUMM--SURG SITE 98-02

Alternate Name	Item #	Length	Source of Standard	Column #
Cancer-Directed Surgery (pre-96 COC)	1646	2	SEER/COC	939-940
Surgery of Primary Site (SEER/COC)				

# Description

Site-specific codes for the type of surgery to the primary site performed as part of the first course of treatment. This includes treatment given at all facilities as part of the first course of treatment. This field is to be used for *ROADS* codes after the *ROADS* to *FORDS* conversion. It is also to be used to code Surgery Primary Site at all facilities for all tumors diagnosed before January 1, 2003.

# Rationale

If central registries wish to study the treatment given at particular hospitals, the hospital-level treatment fields must be used. The summary treatment fields, conversely, combine information across all hospitals that provide first course of treatment for the tumor. Hospital-specific fields allow studies of detailed referral patterns and treatment by type of hospital. Knowing what part of the treatment was given at a particular hospital also helps resolve coding issues.

# Codes (in addition to the site-specific codes)

- 00 No primary site surgery performed.
- 99 Unknown if primary site surgery performed.

*Note:* See the COC *ROADS Manual*, 1998 Supplement, COC Coding System [2140] code 7, and the *SEER Program Code Manual*, RX Coding System [1460] code 5, 1998 for site-specific codes.

# RX SUMM--TRANSPLNT/ENDOCR

Alternate Name	Item #	Length	Source of Standard	Column #
Hematologic Transplant and Endocrine	3250	2	COC	876-877
Procedures				

## Description

Identifies systemic therapeutic procedures administered as part of the first course of treatment at this facility and all other facilities or the reason they were not used. These include bone marrow transplants, stem cell harvests, and surgical and radiation endocrine therapy.

## Rationale

This data item allows the evaluation of patterns of treatment, which involve the alteration of the immune system or change the patient's response to tumor cells but do not involve the administration of antineoplastic agents.

## Codes (refer to FORDS for additional instructions)

- 00 No transplant procedure or endocrine therapy was administered as part of first course therapy
- 10 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
- 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 first-course therapy. No reason was stated in the 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, the patient's family member, or the patient's guardian. The refusal was noted in
- 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 cases and autopsyonly cases.

RX TEXTBRM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2660	100	NPCR	5325-5424

## Description

Text area for manual documentation of information regarding the treatment of the tumor being reported with biological response modifiers or immunotherapy.

## Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

## Instructions

- Date treatment was started.
- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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

## Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RX-SEER	1260
Date of 1st Crs RX-COC	1270
RX Hosp-BRM	720
RX Date Systemic	3230
RX Summ-TranpInt/Endocr	3250
RX Summ-BRM	1410
RX Date-BRM	1240

# **RX TEXT--CHEMO**

KA IEAICHEMU				Kevisea
Alternate Name	Item #	Length	Source of Standard	Column #
	2640	200	NPCR	4925-5124

## Description

Text area for manual documentation of information regarding chemotherapy treatment of the reported tumor.

## Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

## Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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

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## Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RX-SEER	1260
Date of 1st Crs RX-COC	1270
RX Summ-Chemo	1390
RX Hosp-Chemo	700
RX Date-Systemic	3230
RX Date-Chemo	1220
Reason for No Chemo	1440

RX TEXTHORMONE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2650	200	NPCR	5125-5324

#### Description

Text area for information about hormonal cancer-directed treatment.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## **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

#### Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RX-SEER	1260
Date of 1st Crs RX-COC	1270
RX Summ-Hormone	1400
RX Hosp-Hormone	710
RX Date-Systemic	3230
RX Date-Hormone	1230
Reason For No Hormone	1450

## V TEVT OTHER

KX TEXT-OTHER			Revised	
Alternate Name	Item #	Length	Source of Standard	Column #
	2670	100	NPCR	5425-5524

#### Description

Text area for manual documentation of information regarding the treatment of the tumor being reported with treatment that cannot be defined as surgery, radiation, or systemic therapy. This includes experimental treatments (when the mechanism of action for a drug is unknown), and blinded clinical trials. If the mechanism of action for the experimental drug is known, code to the appropriate treatment field.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

## Instructions

- Prioritize entered information in the order of the fields listed below. •
- Text automatically generated from coded data is not acceptable. •
- NAACCR-approved abbreviations should be utilized (see Appendix G). •
- Do not repeat information from other text fields.

- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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

#### Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RX-SEER	1260
Date of 1st Crs RX-COC	1270
RX Summ-Other	1420
RX Date-Other	1250
RX Hosp-Other	730

<b>RX TEXTRADIATION (BEAM)</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2620	150	NPCR	4625-4774

#### Description

Text area for manual documentation of information regarding treatment of the tumor being reported with beam radiation.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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 5 treatments; unknown if radiation was given

## Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RXSEER	1260
Date of 1st Crs RXCOC	1270
RX Summ-Radiation	1360
RX Summ-Surg/Rad Seq	1380
Reason For No Radiation	1430
RX DateRadiation	1210
Rad Regional RX Modality	1570
RX HospRadiation	690
RX Date Radiation Ended	3220
RX SummRad to CNS	1370
RadNo of Treatment Vol	1520
RadRegional Dose cGy	1510
Rad Elapsed Days	1530
Rad Treatment Volume	1540
Rad Location of RX	1550
Rad Intent of Treatment	1560
Rad Boost RX Modality	3200
Rad Boost Dose cGy	3210
Rad RX Completion Status	1580
Rad Local Control Status	1590

<b>RX TEXTRADIATION OTHER</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2630	150	NPCR	4775-4924

#### Description

Text area for manual documentation of information regarding treatment of the tumor being reported with radiation other than beam radiation. This includes brachytherapy and systemic radiation therapy.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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

#### Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Date of Initial RXSEER	1260
Date of 1st Crs RXCOC	1270
RX SummRadiation	1360
RX SummSurg/Rad Seq	1380

Reason For No Radiation	1430
RX DateRadiation	1210
Rad Regional RX Modality	1570
RX HospRadiation	690
RX Date Radiation Ended	3220
RX Summ-Rad to CNS	1370
RadNo of Treatment Vol	1520
RadRegional Dose cGy	1510
Rad Elapsed Days	1530
Rad Treatment Volume	1540
Rad Location of RX	1550
Rad Intent of Treatment	1560
Rad Boost RX Modality	3200
Rad Boost Dose cGy	3210
Rad RX Completion Status	1580
Rad Local Control Status	1590

RX TEXTSURGERY				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2610	150	NPCR	4475-4624

#### Description

Text area for information describing all surgical procedures performed as part of treatment.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

## 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

## Data Item(s) to be verified/validated using the text entered in this field

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
RX Date Surgery	1200
RX SummSurg Prim Site	1290
RX HospSurg Prim Site	670
RX SummScope Reg LN Sur	1292
RX HospScope Reg LN Sur	672
RX SummSurg Oth Reg/Dis	1294
RX HospSurg Oth Reg/Dis	674
Date of Initial RXSEER	1260
Date of 1st Crs RXCOC	1270
EODExtension	790
Site of Distant Met 1-3	1090-1110
Reason for No Surgery	1340
RX SummSurgical Margins	1320
RX HospPalliative Proc	3280
RX SummPalliative Proc	3270
Place of Diagnosis	2690

## SCREENING DATE

Alternate Name	Item #	Length	Source of Standard	Column #
	510	8	COC	313-320

## Description

Most recent date on which the patient participated in a screening program related to this primary cancer.

## Codes (in addition to appropriate dates)

00000000Patient did not participate in screening program related to this primary cancer999999999Patient participated in screening program related to this primary cancer; date is unknown

## SCREENING RESULT

Alternate Name	Item #	Length	Source of Standard	Column #
	520	1	COC	321-321

#### Description

Code the findings from screening recorded in Screening Date [510].

#### Codes

- 0 Within normal limits
- 1 Abnormal/not suggestive of cancer
- 2 Abnormal/suggestive of cancer
- 3 Equivocal/no follow-up necessary
- 4 Equivocal/evaluation recommended
- 8 Not applicable
- 9 Unknown, result not specified

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### SEER CODING SYS--CURRENT

Alternate Name	Item #	Length	Source of Standard	Column #
	2120	1	NAACCR	1198-1198

#### Description

This shows the SEER coding system best describing the majority of SEER items as they are in the record (after conversion).

#### Codes

- 0 No SEER coding
- 1 1987 SEER Coding Manual
- 2 May 1988 SEER Coding Manual
- 3 January 1989 SEER Coding Manual
- 4 January 1992 SEER Coding Manual
- 5 January 1998 SEER Coding Manual
- 6 January 2003 SEER Coding Manual

# SEER CODING SYS--ORIGINAL

Alternate Name	Item #	Length	Source of Standard	Column #
	2130	1	NAACCR	1199-1199

## Description

This shows the SEER coding system best describing the way the majority of SEER items in the record were originally coded.

## Codes

- 0 No SEER coding
- 1 1987 SEER Coding Manual
- 2 May 1988 SEER Coding Manual
- 3 January 1989 SEER Coding Manual
- 4 January 1992 SEER Coding Manual
- 5 January 1998 SEER Coding Manual
- 6 January 2003 SEER Coding Manual

# SEER RECORD NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
Record Number (SEER)	2190	2	SEER	1215-1216

## Description

A unique sequential number assigned by the SEER participant to each record for the person for each submission. The number may change from submission to submission. See also Tumor Record Number [60].

## Codes

- 01 One or first of more than one record for person
- 02 Second record for person
- .. ..
- nn Last of nn records for person

## SEER SUMMARY STAGE 1977

Alternate Name	Item #	Length	Source of Standard	Column #
General Summary Stage (SEER/COC)	760	1	SEER	529-529

## Description

Code for summary stage at the initial diagnosis or treatment of the reportable tumor. This has traditionally been used by central registries to monitor time trends. For hospital registries, COC requires its use in the absence of a defined AJCC classification. For site-specific definitions of categories, see the SEER Summary Staging Guide.

SEER Summary Stage 1977 is limited to information available within 2 months of the date of diagnosis. NAACCR approved extension of this time period to 4 months for prostate tumors diagnosed beginning January 1, 1995.

## Rationale

Stage information is important when evaluating the effects of cancer control programs. It is crucial for understanding whether changes over time in incidence rates or outcomes are due to earlier detection of the cancers. In addition, cancer treatment cannot be studied without knowing the stage at diagnosis.

To study historical trends in stage, the coding system must be relatively unchanged (stable) over time. AJCC's TNM system is updated periodically to maintain clinical relevance with changes in diagnosis and treatment. The surveillance registries often rely on the Summary Stage, which they consider to be more "stable." Summary Stage has been in widespread use, either as the primary staging scheme or a secondary scheme, in most central and hospital registries since 1977.

## Codes

- 0 In situ
- 1 Localized
- 2 Regional, direct extension only
- 3 Regional, regional lymph nodes only
- 4 Regional, direct extension and regional lymph nodes
- 5 Regional, NOS
- 7 Distant
- 8 Not applicable
- 9 Unstaged

*Note:* Code 8 has been added in Version 10.1 to be used when there is not an applicable code to reflect stage (e.g., benign brain, borderline ovarian).

*Note:* See also the item Derived SS1977 [3010] for the value of SEER Summary Stage 1977 as generated by the Collaborative Staging algorithm.

# **Clarification of NAACCR and NPCR Required Status**

Summary stage is required. The correct data item to use (and corresponding code manual) is determined by the year in which the cancer was diagnosed. Tumors diagnosed on or after January 1, 2004, should be assigned a summary stage based upon the Collaborative Stage data item algorithms and retained in Derived SS2000 [3020]. Tumors diagnosed on or after January 1, 2001 should be assigned a summary stage according to the SEER *Summary Staging Manual 2000*, and the code should be reported in SEER Summary Stage 2000

[759]. Tumors diagnosed before January 1, 2001, should be assigned a summary stage according to *SEER Summary Stage Guide 1977*, and the code should be reported in SEER Summary Stage 1977 [760].

## SEER SUMMARY STAGE 2000

Alternate Name	Item #	Length	Source of Standard	Column #
	759	1	SEER	528-528

## Description

Code for summary stage at the initial diagnosis or treatment of the reportable tumor. For hospital registries, COC requires its use in the absence of a defined AJCC classification. For site-specific definitions of categories, see SEER *Summary Staging Manual 2000*.

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

#### Rationale

Stage information is important when evaluating the effects of cancer control programs. It is crucial in understanding whether changes over time in incidence rates or outcomes are due to earlier detection of the cancers. In addition, cancer treatment cannot be studied without knowing the stage at diagnosis.

#### Codes

- 0 In situ
- 1 Localized
- 2 Regional, direct extension only
- 3 Regional, regional lymph nodes only
- 4 Regional, direct extension and regional lymph nodes
- 5 Regional, NOS
- 7 Distant
- 8 Not applicable
- 9 Unstaged

*Note:* Code 8 has been added in Version 10.1 to be used when there is not an applicable code to reflect stage (e.g., benign brain, borderline ovarian).

*Note:* See also the item Derived SS2000 [3020] for the value of SEER Summary Stage 2000 as generated by the Collaborative Staging algorithm.

# **Clarification of NAACCR and NPCR Required Status**

Summary stage is required. The correct data item to use (and corresponding code manual) is determined by the year in which the cancer was diagnosed. Tumors diagnosed on or after January 1, 2004, should be assigned a summary stage based upon the Collaborative Stage data item algorithms and retained in Derived SS2000 [3020]. Tumors diagnosed on or after January 1, 2001 should be assigned a summary stage according to the SEER *Summary Staging Manual 2000*, and the code should be reported in SEER Summary Stage 2000 [759]. Tumors diagnosed before January 1, 2001, should be assigned a summary stage according to *SEER Summary Stage Guide 1977*, and the code should be reported in SEER Summary Stage 1977 [760].

# **SEER TYPE OF FOLLOW-UP**

Alternate Name	Item #	Length	Source of Standard	Column #
Type of Follow-Up (SEER)	2180	1	SEER	1214-1214

# Description

Codes for the type of follow-up expected for a SEER case.

# Codes

- 1 "Autopsy-Only" or "Death Certificate-Only" case
- 2 Active follow-up case
- 3 In situ cancer of the cervix uteri only
- 4 Case not originally in active follow-up, but in active follow-up now (San Francisco-Oakland only)

SEQUENCE NUMBERCENTRAL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Sequence Number (pre-96 SEER)	380	2	SEER	281-282

## Description

Code indicates the sequence of all reportable neoplasms over the lifetime of the person. This data item differs from Sequence Number--Hospital [560], because the definitions of reportable neoplasms often vary between a hospital and a central registry. Each neoplasm is assigned a different number. Sequence Number 00 indicates that the person has had only one *in situ* or one malignant neoplasm as defined by the Federal reportable list (regardless of central registry reference date). Sequence Number 01 indicates the first of two or more reportable neoplasms, while 02 indicates the second of two or more reportable neoplasms, and so on. Because the time period of Sequence Number is a person's lifetime, reportable neoplasms not included in the central registry (those that occur outside the registry catchment area or before the reference date) also are allotted a sequence number. For example, a registry may contain a single record for a patient with a sequence number of 02 because the first reportable neoplasm preceded the central registry's reference date.

## Reporting Requirements: Federally Required and State/Province Defined

The Federal or SEER/NPCR standard defining which neoplasms are reportable is described in Chapter III, Standards For Tumor Inclusion and Reportability. It is assumed that this shared standard is the "minimum" definition of reportability. Individual central cancer registries may define additional neoplasms as reportable.

Numeric codes in the 00-35 range indicate the sequence of neoplasms of *in situ* or malignant behavior (2 or 3) at the time of diagnosis, which SEER/NPCR standards require to be reported. Codes 60 to 87 indicate the sequence of non-malignant tumors (as defined in Chapter III) and any other neoplasms that the central registry has defined as reportable. Neoplasms required by SEER/NPCR with an *in situ* or malignant behavior at the time of diagnosis are sequenced completely independently of this higher-numbered category. Sequence Number--Hospital does not affect Sequence Number--Central. The two notational systems are independent but central registries should take Sequence Number--Hospital [560] into account when coding Sequence Number Central.

# **Timing Rule**

The sequence number may change over the lifetime of the patient. If an individual previously diagnosed with a single reportable malignant neoplasm is subsequently diagnosed with a second reportable malignant neoplasm, the sequence code for the first neoplasm changes from 00 to 01. A central registry might also discover that an individual with one or more known neoplasms had an earlier reportable neoplasm that had

been unknown to the registry. Typically, a re-evaluation of all related sequence numbers is required whenever an additional neoplasm is identified.

If two or more reportable neoplasms are diagnosed at the same time, the lowest sequence number is to be assigned to the diagnosis with the worst prognosis. If no difference in prognosis is evident, the decision is arbitrary. Timing rules for sequencing neoplasms coded in the 60-87 range are the same as timing rules for sequencing of required *in situ* or malignant neoplasms.

If a registry collects any central registry-defined neoplasms, the codes 60-87 should be used. The codes 60-87 also should be used for non-malignant tumor diagnosed on or after January 1, 2004. Timing rules for sequencing these neoplasms are the same as timing rules for sequencing of required *in situ* or invasive neoplasms.

# Rationale

The purpose of sequencing based on the patient's lifetime is to truly identify the 00s, the people who only had one malignant primary in their lifetimes for survival analysis. If a central registry sequences by just what is reported to them, then it will be unclear whether 00 means the person only had one malignant primary in his lifetime or the person had one malignant primary since the central registry started collecting data. The Federally required reportable list has changed throughout the years, so the registry must use the appropriate reportable list for the year of diagnosis. The central registry reference date will not affect Sequence Number-Central.

# Codes

In Situ/Malignant as Federally Required based on Diagnosis Year:

- 00 One primary in the patient's lifetime
- 01 First of two or more primaries
- 02 Second of two or more primaries
- ••
- 35 Thirty-fifth of thirty-five or more primaries
- 99 Unspecified or unknown sequence number of Federally required *in situ* or malignant tumors. Sequence number 99 can be used if there is a malignant tumor and its sequence number is unknown. (If there is known to be more than one malignant tumor, then the tumors must be sequenced.)

Non-malignant Tumor as Federally Required based on Diagnosis Year or State/Province Defined:

- 60 One non-malignant tumor or central registry-defined neoplasm
- 61 First of two or more non-malignant tumor or central registry-defined neoplasms
- 62 Second of two or more non-malignant tumor or central registry-defined neoplasms
- 88 Unspecified or unknown sequence number for non-malignant tumor or central registry-defined neoplasms. (Sequence number 88 can be used if there is a non-malignant tumor and its sequence number is unknown. If there is known to be more than one non-malignant tumor, then the tumors must be sequenced.)
- 98 Cervix carcinoma *in situ* (CIS)/CIN III, Diagnosis Years 1996-2002.

Neoplasm	SeqNum-Central
In Situ/Malignant as Federally Required based on Diagnosis	(Numeric Series)
Year	
<i>In Situ</i> (behavior code = 2) (Cervix CIS/CIN III, Diagnosis Year	00 - 35
before 1996) (includes VIN III, VAIN III, AIN III)	
Malignant (behavior code = 3)	00 - 35
Juvenile Astrocytoma, Diagnosis Year 2001+ (*)	00 - 35
Invasive following In SituNew primary as defined by COC	00 - 35
Invasive following In SituNew primary as defined by SEER	00 - 35
Unspecified Federally Required Sequence Number or Unknown	99
Non-malignant Tumor as Federally Required based on	
<b>Diagnosis Year or State/Province Registry-Defined</b>	
Examples:	
Non-malignant Tumor/Benign Brain	60 - 87
Borderline Ovarian, Diagnosis Year 2001+	60 - 87
Other Borderline/Benign	60 - 87
Skin SCC/BCC	60 - 87
PIN III	60 - 87
Cervix CIS/CIN III, Diagnosis Year 2003+	60 - 87
Unspecified Non-malignant Tumor or Central Registry-Defined	88
Sequence Number	
Cervix CIS/CIN III, Diagnosis Year 1996-2002	98

The table below shows which sequence number series to use by type of neoplasm

\*Juvenile astrocytomas should be reported as 9421/3.

Note: See the section on Sequence Number--Central in The SEER Program Code Manual.

*Note:* Conversion Guidance: The sequence numbers for neoplasms whose histologies were associated with behavior codes that changed from *in situ*/malignant to benign/borderline or vice versa during the conversion from ICD-O-2 to ICD-O-3 should not be re-sequenced.

SEQUENCE NUMBERHOSPITAL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Sequence Number (COC)	560	2	COC	411-412

## Description

Code indicates the sequence of all malignant and non-malignant neoplasms over the lifetime of the patient. This item differs from the Sequence Number--Central [380] because the definitions of reportable neoplasms often vary between a hospital and a central registry. Each neoplasm is assigned a different number. Sequence Number 00 indicates that the person has only one malignant neoplasm in his lifetime (regardless of hospital registry reference date). Sequence Number 01 indicates the first of two or more malignant neoplasms, while 02 indicates the second of two or more malignant neoplasms, and so on. Because the time period of Sequence Number is a person's lifetime, reportable neoplasms not included in the hospital registry are also allotted a sequence number. For example, a registry may contain a single record for a patient with a sequence number of 02 because the first reportable neoplasm occurred before the hospital registry's reference date. Similarly, Sequence Number 60 indicates the patient has only one non-malignant neoplasm, and Sequence Number 61 represents the first of multiple non-malignant neoplasms.

## Reporting Requirements: COC, State/Province, and The Hospital Cancer Committee

The COC standard defining which neoplasms are reportable is described in Chapter III, Standards For Case Inclusion and Reportability; it is assumed that this standard is the "minimum" definition of reportability. In addition to the COC-required reportable neoplasms, hospital cancer registries have to meet the reporting requirements of the central cancer registry and the hospital cancer committee. These neoplasms often are called "reportable-by-agreement" in COC publications. Any tumor in the patient's past that is reportable or reportable-by-agreement must be taken into account when sequencing subsequently accessioned tumors. Sequence numbers should be reassigned if the facility subsequently learns of an unaccessioned tumor that affects sequencing. Sequence Number--Central [380] does not affect Sequence Number--Hospital. The two notational systems are independent.

## **Timing Rule**

If two or more malignant tumors are diagnosed at the same time, the lowest sequence number will be assigned to the diagnosis with the worst prognosis. Likewise, if two or more non-malignant tumors are diagnosed at the same time, the lowest sequence number is assigned to the diagnosis with the worse prognosis. If no difference in prognosis is evident, the decision is arbitrary.

# Codes

In Situ and Malignant Tumors:

- 00 One malignant primary only in the patient's lifetime
- 01 First of two or more malignant primaries
- 02 Second of two or more malignant primaries
- .. (Actual number of this malignant primary)
- 35 Thirty-fifth of thirty-five malignant primaries
- 99 Unspecified sequence number of a primary malignant tumor or unknown (when a patient has multiple tumors with unspecified/unknown sequence numbers code 99 should only be used once.)

Nonmalignant Tumors

- 60 Only one non-malignant tumor in the patient's lifetime
- 61 First of two or more non-malignant tumors
- 62 Second of two or more non-malignant tumors

88 Unspecified number of non-malignant tumors (when a patient has multiple unspecified neoplasms in this category code 88 should only be used once.)

The table below shows which sequence number series to use by type of neoplasm

Neoplasm	SeqNum-Hospital
In situ and Malignant	(code range)
One <i>in situ</i> (behavior code = 2) or malignant (behavior code =3) primary tumor only in the patient's lifetime	00
First of multiple <i>in situ</i> or malignant primary tumors in the patient's lifetime	01
Actual sequence of two or more <i>in situ</i> or malignant primary tumors	02 - 35
Unspecified malignant sequence number or unknown	99
Non-Malignant	
One benign (behavior code = 0) or borderline (behavior code = 1) primary tumor only in the patient's lifetime	60
First of two or more benign or borderline primary tumors in the patient's lifetime	61
Actual sequence of two or more non-malignant primary tumors	62 - 87
Unspecified non-malignant sequence number or unknown	88

\*Juvenile astrocytomas should be reported as 9421/3

Note: See the section on Sequence Number in COC (FORDS) Manual.

#### SEX

..

Alternate Name	Item #	Length	Source of Standard	Column #
	220	1	SEER/COC	118-118

## Description

Code for the sex of the patient.

#### Codes

- 1 Male
- 2 Female
- 3 Other (hermaphrodite)
- 4 Transsexual
- 9 Not stated/Unknown

## SITE (73-91) ICD-O-1

Alternate Name	Item #	Length	Source of Standard	Column #
Primary Site (1973-91) (SEER)	1960	4	SEER	1137-1140

## Description

Area for retaining the ICD-O-1 primary site code entered before conversion to ICD-0-2. The item name includes years 1973-91. However, some states may have used the codes for cases before 1973.

## Codes

For tumors diagnosed before 1992, contains the ICD-O-1 site code as originally coded, if available. Blank for tumors coded directly into ICD-O-2 (i.e., 1992 and later tumors).

## SITE CODING SYS--CURRENT

Alternate Name	Item #	Length	Source of Standard	Column #
	450	1	NAACCR	307-307

## Description

Code that best describes how the primary site currently is coded. If converted, this field shows the system to which it is converted.

#### Codes

- 1 ICD-8 and MOTNAC
- 2 ICD-9
- 3 ICD-O, First Edition
- 4 ICD-O, Second Edition
- 5 ICD-O, Third Edition
- 6 ICD-10
- 9 Other

## SITE CODING SYS--ORIGINAL

Alternate Name	Item #	Length	Source of Standard	Column #
	460	1	NAACCR	308-308

## Description

Code that best describes how primary site was originally coded. If converted, this field shows the original coding system used.

## Codes

- 1 ICD-8 and MOTNAC
- 2 ICD-9
- 3 ICD-O, First Edition
- 4 ICD-O, Second Edition
- 5 ICD-O, Third Edition
- 6 ICD-10
- 9 Other

## SITE OF DISTANT MET 1

Alternate Name	Item #	Length	Source of Standard	Column #
Site of Distant Metastasis #1 (COC)	1090	1	COC	618-618

#### Description

Codes for a site of distant metastasis at initial diagnosis. There are three individual fields, each with a 1-digit code for a site of metastasis.

#### Codes

- 0 None
- 1 Peritoneum
- 2 Lung
- 3 Pleura
- 4 Liver
- 5 Bone
- 6 Central nervous system
- 7 Skin
- 8 Lymph nodes (distant)
- 9 Other, generalized, carcinomatosis, disseminated, not specified, unknown

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

## SITE OF DISTANT MET 2

Alternate Name	Item #	Length	Source of Standard	Column #
Site of Distant Metastasis #2 (COC)	1100	1	COC	619-619

#### Description

Codes for a site of distant metastasis at initial diagnosis. There are three individual fields, each with a 1-digit code for a site of metastasis.

#### Codes

- 0 None
- 1 Peritoneum
- 2 Lung
- 3 Pleura
- 4 Liver
- 5 Bone
- 6 Central nervous system
- 7 Skin
- 8 Lymph nodes (distant)
- 9 Other, generalized, carcinomatosis, disseminated, not specified, unknown

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

## SITE OF DISTANT MET 3

Alternate Name	Item #	Length	Source of Standard	Column #
Site of Distant Metastasis #3 (COC)	1110	1	COC	620-620

## Description

Codes for a site of distant metastasis at initial diagnosis. There are three individual fields, each with a 1-digit code for a site of metastasis.

## Codes

- 0 None
- 1 Peritoneum
- 2 Lung
- 3 Pleura
- 4 Liver 5
- Bone
- Central nervous system 6
- Skin 7
- 8 Lymph nodes (distant)
- Other, generalized, carcinomatosis, disseminated, not specified, unknown 9

Note: As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

## SOCIAL SECURITY NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
	2320	9	COC	2099-2107

## Description

Records patient's social security number. The number is entered without dashes and without any letter suffix. This is not always identical to the Medicare claim number.

## Codes (in addition to social security number)

9999999999 Unknown

## SPANISH/HISPANIC ORIGIN

SPANISH/HISPANIC ORIGIN				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Spanish OriginAll Sources (96 COC)	190	1	SEER/COC	115-115
Spanish Surname or Origin (SEER)				

## Description

Code identifying persons of Spanish or Hispanic origin. This code is used by hospital and central registries to show the "best guess" as to whether or not the person should be classified as Hispanic for purposes of calculating cancer rates. If the patient has multiple tumors, all records should have the same code.

Reference to Census 2000 definitions for ethnicity and race: http://www.census.gov/prod/cen2000/doc/sf2.pdf

All information resources should be used to determine the correct code, including:

- Stated ethnicity in the medical record
- Stated Hispanic origin on the death certificate •
- Birthplace •
- Information about life history and/or language spoken found during the abstracting process
- Patient's last name [2230] or maiden name [2390] found on a list of Hispanic names. •

Some registries code the information from the medical record, others code ethnicity based on Spanish names, and others use a combination of methods.

Persons of Spanish or Hispanic origin may be of any race, but these categories generally are not used for Native Americans, Filipinos, etc., who may have Spanish names. If a patient has an Hispanic name, but there is reason to believe they are not Hispanic (e.g., the patient is Filipino, or the patient is a woman known to be non-Hispanic who has a Hispanic married name), the code in this field should be 0 (non-Spanish, non-Hispanic). The code in item Computed Ethnicity [200], however, would reflect the Hispanic name.

Assign code 7 if Hispanic ethnicity is based strictly on a computer list or algorithm (unless contrary evidence is available) and also code in Computed Ethnicity [200].

See also Computed Ethnicity [200].

Note: NAACCR recognizes that available definitions and abstracting instructions for Name--Last [2230] and Name--Maiden [2390] may be inadequate for describing names used in some cultures, including Hispanic cultures. Explicit instructions have not been provided for entering compound names, with or without hyphens or "De." Order of names, use of maternal and paternal names, and use of hyphens can vary across cultures. It is likely that abstracting and coding practice for these items varies across registries. Limitations inherent in these definitions should be kept in mind when using the data.

## Rationale

See the rationales for the Race 1-5 [160-164] and Computed Ethnicity [200]. Ethnic origin has a significant association with cancer rates and outcomes. Hispanic populations have different patterns of occurrence of cancer from other populations that may be included in the "white" category of Race [160].

## Codes

- 0 Non-Spanish; non-Hispanic
- Mexican (includes Chicano) 1
- 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 of the categories 1-5.

7 Spanish surname only (Code 7 is ordinarily for central registry use only, hospital registrars may use code 7 if using a list of Hispanic surnames provided by their central registry; otherwise, code 9 "unknown whether Spanish or not" should be used.)

# The only evidence of the person's Hispanic origin is the surname or maiden name and there is no contrary evidence that the patient is not Hispanic.

- 8 Dominican Republic
- 9 Unknown whether Spanish or not

Note: Code 7 was adopted for use effective with 1994 diagnosis and modified December 1994.

#### **STATE/REQUESTOR ITEMS**

Alternate Name	Item #	Length	Source of Standard	Column #
	2220	500	Varies	1447-1946

#### Description

Old fields, Site-Specific Studies, and State-Specific Items were combined into this area and renamed. The area also was expanded. Reserved for use by special studies, or for items defined in individual states or central registries. COC uses this area for Patient Care Evaluation Studies.

SUBSQ REPORT FOR PRIMARY				Retired
Alternate Name	Item #	Length	Source of Standard	Column #
	2160			

SUBSQ RX 2ND COURSE BRM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1675	1	COC	1001-1001

#### Description

Codes for the type of biological response modifier therapy given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Immunotherapy, *1998 ROADS Manual*, p. 243. See also First Course Calc Method [1500].

SUBSQ RX 2ND COURSE CHEMO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1673	1	COC	999-999

## Description

Codes for the type of chemotherapy given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Chemotherapy, *1998 ROADS Manual*, p. 228. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

## SUBSQ RX 2ND COURSE CODES

Alternate Name	Item #	Length	Source of Standard	Column #
	1670	7		996-1002

#### Description

The name for a group of subfields that describe the second course or set of subsequent therapy. As of January 1, 2003, COC no longer supports Subsequent Therapy data items.

Group names appear only in the data dictionary and Appendix E.

#### Subfields

Subsq RX 2nd Course Surg [1671] Subsq RX 2nd Course Rad [1672] Subsq RX 2nd Course Chemo [1673] Subsq RX 2nd Course Horm [1674] Subsq RX 2nd Course BRM [1675] Subsq RX 2nd Course Oth [1676]

SUBSQ RX 2ND COURSE DATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Second Course of Therapy-Date Started	1660	8	COC	988-995
(pre-96 COC)				

## Description

Date of initiation of second-course treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. See page 83 for date format.

#### Codes (in addition to valid dates)

00000000	No subsequent therapy
99999999	Unknown if any subsequent therapy

SUBSQ RX 2ND COURSE HORM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1674	1	COC	1000-1000

## Description

Codes for the type of hormonal therapy given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Hormone Therapy, *1998 ROADS Manual*, p. 238. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 2ND COURSE OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1676	1	COC	1002-1002

## Description

Codes for the type of other treatment given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Other Treatment, *1998 ROADS Manual*, p. 246. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 2ND COURSE RAD				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1672	1	COC	998-998

## Description

Codes for the type of radiation given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Radiation, *1998 ROADS Manual*, p. 199. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 2ND COURSE SURG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1671	2	COC	996-997

# Description

Codes for the type of primary site surgery given as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Surgery of Primary Site, *1998 ROADS Manual*, p. 187. See also First Course Calc Method [1500].

SUBSQ RX 2NDREG LN REM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1679	2	COC	1050-1051

Codes for the number of regional lymph nodes removed as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Number of Regional Lymph Nodes Removed, *1998 ROADS Manual*, p. 193. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 2NDSCOPE LN SU				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1677	1	COC	1048-1048

#### Description

Codes for the type of surgery performed to remove regional lymph nodes as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Scope of Regional Lymph Node Surgery, *1998 ROADS Manual*, p. 192. See also First Course Calc Method [1500].

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 2NDSURG OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1678	1	COC	1049-1049

# Description

Codes for the type of surgery performed on tissue or organs other than the primary site and regional lymph nodes as part of the second course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s), *1998 ROADS Manual*, p. 194. See also First Course Calc Method [1500].

SUBSQ RX 3RD COURSE BRM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1695	1	COC	1016-1016

Codes for the type of biological response modifier therapy given as part of the third course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Immunotherapy, *1998 ROADS Manual*, p. 243.

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 3RD COURSE CHEMO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1693	1	COC	1014-1014

# Description

Codes for the type of chemotherapy given as part of the third course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Chemotherapy, *1998 ROADS Manual*, p. 228.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# SUBSQ RX 3RD COURSE CODES

Alternate Name	Item #	Length	Source of Standard	Column #
	1690	7		1011-1017

# Description

The name for a group of subfields that describe the third course or set of subsequent therapy. As of January 1, 2003, COC no longer supports Subsequent Therapy data items.

Group names appear only in the data dictionary and Appendix E.

# Subfields

Subsq RX 3rd Course Surg [1691] Subsq RX 3rd Course Rad [1692] Subsq RX 3rd Course Chemo [1693] Subsq RX 3rd Course Horm [1694] Subsq RX 3rd Course BRM [1695] Subsq RX 3rd Course Oth [1696]

SUBSQ RX 3RD COURSE DATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1680	8	COC	1003-1010

Date of initiation of third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. See page 83 for date format.

# Codes

00000000	No subsequent therapy
99999999	Unknown if any subsequent therapy

SUBSQ RX 3RD COURSE HORM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1694	1	COC	1015-1015

# Description

Codes for the type of hormonal therapy given as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Hormone Therapy, 1998 ROADS Manual, p. 238.

*Note:* This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 3RD COURSE OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1696	1	COC	1017-1017

# Description

Codes for the type of other treatment given as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Other Treatment, 1998 ROADS Manual, p. 246.

Note: This data item is no longer supported by COC (as of January 1, 2003).

#### SUBSO RX 3RD COURSE RAD

SUBSQ RX 3RD COURSE RAD				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1692	1	COC	1013-1013

# Description

Codes for the type of radiation given as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Radiation, 1998 ROADS Manual, p. 1999.

SUBSQ RX 3RD COURSE SURG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1691	2	COC	1011-1012

Codes for the type of primary site surgery given as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Surgery of Primary Site, 1998 ROADS Manual, p. 187.

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 3RDREG LN REM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1699	2	COC	1054-1055

# Description

Codes for the number of regional lymph nodes removed as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Number of Regional Lymph Nodes Removed, 1998 ROADS Manual, p. 193.

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 3RDSCOPE LN SU				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1697	1	COC	1052-1052

# Description

Codes for the type of surgery performed to remove regional lymph nodes as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Scope of Regional Lymph Node Surgery, 1998 ROADS Manual, p. 192.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# SUBSO BY 3PD SUDC OTH

SUBSQ RX 3RDSURG OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1698	1	COC	1053-1053

# Description

Codes for the type of surgery performed on tissue or organs other than the primary site and regional lymph nodes as part of the third course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s), 1998 ROADS Manual, p. 194.

SUBSQ RX 4TH COURSE BRM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1715	1	COC	1031-1031

Codes for the type of biological response modifier therapy given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Immunotherapy, *1998 ROADS Manual*, p. 243.

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 4TH COURSE CHEMO				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1713	1	COC	1029-1029

#### Description

Codes for the type of chemotherapy given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. The codes are the same as those for Chemotherapy, *1998 ROADS Manual*, p. 228.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# SUBSQ RX 4TH COURSE CODES

Alternate Name	Item #	Length	Source of Standard	Column #
	1710	7		1026-1032

#### Description

The name for a group of subfields that describe the fourth course or set of subsequent therapy. As of January 1, 2003, COC no longer supports Subsequent Therapy data items.

Group names appear only in the data dictionary and Appendix E.

#### Subfields

Subsq RX 4th Course Surg [1711] Subsq RX 4th Course Rad [1712] Subsq RX 4th Course Chemo [1713] Subsq RX 4th Course Horm [1714] Subsq RX 4th Course BRM [1715] Subsq RX 4th Course Oth [1716]

SUBSQ RX 4TH COURSE DATE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1700	8	COC	1018-1025

Date of initiation of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. See page 83 for date format.

# **Codes (in addition to valid dates)**

00000000	No subsequent therapy
99999999	Unknown if any subsequent therapy

SUBSQ RX 4TH COURSE HORM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1714	1	COC	1030-1030

#### Description

Codes for the type of hormonal therapy given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Hormone Therapy, 1998 ROADS Manual, p. 238.

*Note:* This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 4TH COURSE OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1716	1	COC	1032-1032

# Description

Codes for the type of other treatment given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Other Treatment, 1998 ROADS Manual, p. 246.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# SUBSO RX 4TH COURSE RAD

SUBSQ RX 4TH COURSE RAD				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1712	1	COC	1028-1028

# Description

Codes for the type of radiation given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Radiation, 1998 ROADS Manual, p. 199.

SUBSQ RX 4TH COURSE SURG				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1711	2	COC	1026-1027

Codes for the type of primary site surgery given as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Surgery of Primary Site, 1998 ROADS Manual, p. 187.

*Note:* This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 4THREG LN REM				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1719	2	COC	1058-1059

#### Description

Codes for the number of regional lymph nodes removed as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Number of Regional Lymph Nodes Removed, 1998 ROADS Manual, p. 193.

Note: This data item is no longer supported by COC (as of January 1, 2003).

SUBSQ RX 4THSCOPE LN SU				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1717	1	COC	1056-1056

#### Description

Codes for the type of surgery performed to remove regional lymph nodes as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Scope of Regional Lymph Node Surgery, 1998 ROADS Manual, p. 192.

*Note:* This data item is no longer supported by COC (as of January 1, 2003).

#### SUBSO RX 4TH--SURG OTH

SUBSQ RX 4THSURG OTH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	1718	1	COC	1057-1057

# Description

Codes for the type of surgery performed on tissue or organs other than the primary site and regional lymph nodes as part of the fourth course of treatment. Central registries currently collecting this data item should follow the 1998 ROADS Manual coding instructions. The codes are the same as those for Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s), 1998 ROADS Manual, p. 194.

# SUBSQ RX 5TH COURSE BRM

Alternate Name	Item #	Length	Source of Standard	Column #
	1735	1	NAACCR	1046-1046

# Description

Codes for the type of biological response modifier therapy given as part of the fifth course of treatment. The codes are the same as those for Immunotherapy, *1998 ROADS Manual*, p. 243

The COC ROADS Manual does not include fifth course of treatment.

# SUBSQ RX 5TH COURSE CHEMO

Alternate Name	Item #	Length	Source of Standard	Column #
	1733	1	NAACCR	1044-1044

# Description

Codes for the type of chemotherapy given as part of the fifth course of treatment. The codes are the same as those for Chemotherapy, *1998 ROADS Manual*, p. 228.

Note: The COC ROADS Manual does not include fifth course of treatment.

# SUBSQ RX 5TH COURSE CODES

Alternate Name	Item #	Length	Source of Standard	Column #
	1730	7		1041-1047

# Description

The name for a group of subfields that describe the fifth course or set of subsequent therapy.

Group names appear only in the data dictionary and Appendix E.

# Subfields

Subsq RX 5th Course Surg [1731] Subsq RX 5th Course Rad [1732] Subsq RX 5th Course Chemo [1733] Subsq RX 5th Course Horm [1734] Subsq RX 5th Course BRM [1735] Subsq RX 5th Course Oth [1736]

# SUBSQ RX 5TH COURSE DATE

Alternate Name	Item #	Length	Source of Standard	Column #
	1720	8	NAACCR	1033-1040

#### Description

Date of initiation of fifth course of treatment. See page 83 for date format.

The COC ROADS Manual does not include fifth course of treatment.

#### Codes (in addition to valid dates)

00000000	No subsequent therapy
99999999	Unknown if any subsequent therapy

#### SUBSQ RX 5TH COURSE HORM

Alternate Name	Item #	Length	Source of Standard	Column #
	1734	1	NAACCR	1045-1045

#### Description

Codes for the type of hormonal therapy given as part of the fifth course of treatment. The codes are the same as those for Hormone Therapy, *1998 ROADS Manual*, p. 238.

The COC ROADS Manual does not include fifth course of treatment.

#### SUBSQ RX 5TH COURSE OTH

Alternate Name	Item #	Length	Source of Standard	Column #
	1736	1	NAACCR	1047-1047

#### Description

Codes for the type of other treatment given as part of the fifth course of treatment. The codes are the same as those for Other Treatment, *1998 ROADS Manual*, p. 246.

The COC ROADS Manual does not include fifth course of treatment.

#### SUBSQ RX 5TH COURSE RAD

Alternate Name	Item #	Length	Source of Standard	Column #
	1732	1	NAACCR	1043-1043

#### Description

Codes for the type of radiation therapy given as part of the fifth course of treatment. The codes are the same as those for Radiation, *1998 ROADS Manual*, p. 199.

The COC ROADS Manual does not include fifth course of treatment.

### SUBSQ RX 5TH COURSE SURG

Alternate Name	Item #	Length	Source of Standard	Column #
	1731	2	NAACCR	1041-1042

#### Description

Codes for the type of primary site surgery given as part of the fifth course of treatment. The codes are the same as those for Surgery of Primary Site, *1998 ROADS Manual* p. 187.

The COC ROADS Manual does not include fifth course of treatment.

#### SUBSQ RX 5TH--REG LN REM

Alternate Name	Item #	Length	Source of Standard	Column #
	1739	2	NAACCR	1062-1063

#### Description

Codes for the number of regional lymph nodes removed as part of the fifth course of treatment. The codes are the same as those for Number of Regional Lymph Nodes Removed, *1998 ROADS Manual*, p. 193.

The COC ROADS Manual does not include fifth course of treatment.

#### SUBSQ RX 5TH--SCOPE LN SU

Alternate Name	Item #	Length	Source of Standard	Column #
	1737	1	NAACCR	1060-1060

#### Description

Codes for the type of surgery performed to remove regional lymph nodes as part of the fifth course of treatment. The codes are the same as those for Scope of Regional Lymph Node Surgery, *1998 ROADS Manual*, p. 192.

The COC ROADS Manual does not include fifth course of treatment.

#### SUBSQ RX 5TH--SURG OTH

Alternate Name	Item #	Length	Source of Standard	Column #
	1738	1	NAACCR	1061-1061

#### Description

Codes for the type of surgery performed on tissue or organs other than the primary site and regional lymph nodes as part of the fifth course of treatment. The codes are the same as those for Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Node(s), *1998 ROADS Manual*, p. 194.

The COC ROADS Manual does not include fifth course of treatment.

SUBSQ RXRECONSTRUCT DEL				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
Reconstruction/RestorationDelayed	1741	1	COC	1064-1064
(COC)				

Code for surgical procedure done to reconstruct, restore, or improve shape and appearance or function of body structures that are missing, defective, damaged, or misshapen by cancer or cancer-directed therapies. Reconstructive/restorative procedures are coded here when started after the first course of cancer-directed therapy. Central registries currently collecting this data item should follow the *1998 ROADS Manual* coding instructions. For reconstructive/restorative procedures started during the first course of therapy, see RX Summ--Reconstruct 1st [1330]. See also RX Summ--Surgery Type [1640].

# Codes

See the COC ROADS Manual, 1998 Supplement, for site-specific codes.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# TELEPHONE

Alternate Name	Item #	Length	Source of Standard	Column #
	2360	10	COC	2268-2277

# Description

Current telephone number with area code for the patient. Number is entered without dashes.

# Codes (in addition to valid telephone number)

000000000Patient does not have a telephone9999999999Telephone number unavailable or unknown

*Note:* Prior to Version 5, Follow-Up Contact fields may have been used for patient current telephone in the NAACCR record layout.

TEXTDX PROCLAB TESTS				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2550	250	NPCR	3345-3594

Text area for manual documentation of information from laboratory examinations other than cytology or histopathology.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# 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).

#### Data Item(s) to be verified/validated using the text entered in this field:

After manual entry of the text field, ensure that the text entered both agrees with the coded values and clearly justifies the selected codes in the following fields:

Item name	Item number
Primary Site	400
Grade	440
Diagnostic Confirmation	490
Laterality	410
Collaborative Stage variables	2800-2930
Date of Diagnosis	390

TEXTDX PROCOP				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2560	250	NPCR	3595-3844

#### Description

Text area for manual documentation of all surgical procedures that provide information for staging.

#### Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# 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

# Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Date of 1st Positive Bx	1080
Date of Diagnosis	390
RX SummDx/Stg Proc	1350
Diagnostic Confirmation	490
Primary Site	400
RX HospDx/Stg Proc	740
RX SummSurg Prim Site	1290
Collaborative Stage variables	2800-2930
SEER Summary Stage 1977	760
SEER Summary Stage 2000	759

TEXTDX PROCPATH				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2570	250	NPCR	3845-4094

Text area for manual documentation of information from cytology and histopathology reports.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable. •
- NAACCR-approved abbreviations should be utilized (see Appendix G). •
- Do not repeat information from other text fields. •
- Additional comments can be continued in empty text fields, including Remarks. For text • documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing. •
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

Note: For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

#### **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 resection margins
- Number of lymph nodes involved and examined
- Record both positive and negative findings. Record positive test results first.
- Note if pathology report is a slide review or a second opinion from an outside source, i.e., AFIP, • Mavo, etc.
- Record any additional comments from the pathologist, including differential diagnoses considered • and any ruled out or favored

# Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Date of Diagnosis	390
Primary Site	400
Laterality	410
Histologic Type ICD-O-3	522
Histology (92-00) ICD-O-2	420
Grade	440
Collaborative Stage variables	2800-2930
Diagnostic confirmation	490

TEXTDX PROCPE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2520	200	NPCR	2645-2844

Text area for manual documentation from the history and physical examination about the history of the current tumor and the clinical description of the tumor.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below. •
- Text automatically generated from coded data is not acceptable. •
- NAACCR-approved abbreviations should be utilized (see Appendix G). •
- Do not repeat information from other text fields. •
- Additional comments can be continued in empty text fields, including Remarks. For text • documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

Note: For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# **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 •

# Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Date of 1st Contact	580
Date of Diagnosis	390
Age at Diagnosis	230
Race 1 - 5	160-164
Spanish Hispanic Origin	190
Sex	220
Primary Site	400
Laterality	410
Histology (92-00) ICD-O-2	420
Histology ICD-O-3	522
Sequence Number-Central	380
Collaborative Stage variables	2800-2930
SEER Summary Stage 1977	760
SEER Summary Stage 2000	759

TEXTDX PROCSCOPES				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2540	250	NPCR	3095-3344

Text area for manual documentation from endoscopic examinations that provide information for staging and treatment.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below. •
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G). •
- Do not repeat information from other text fields. •
- Additional comments can be continued in empty text fields, including Remarks. For text • documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect. •

Note: For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

#### **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

#### Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Date of Diagnosis	390
Date of 1st Positive Bx	1080
RX Summ-Dx/Stg Proc	1350
Diagnostic Confirmation	490
Primary Site	400
Laterality	410
Histology (92-00) ICD-O-2	420
Histology ICD-O-3	522
Collaborative Stage variables	2800-2930
SEER Summary Stage 1977	760
SEER Summary Stage 2000	759

TEXTDX PROCX-RAY/SCAN				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2530	250	NPCR	2845-3094

Text area for manual documentation from all X-rays, scan, and/or other imaging examinations that provide information about staging.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# Suggestions for text:

- Date(s) of X-ray/Scan(s)
- Age, sex, race/ethnicity (when given)
- 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

# Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Date of Diagnosis	390
Sex	220
Birth Date	240
RxSumm-Dx/Stg Proc	1350
Primary Site	400
Laterality	410
Histology (92-00) ICD-O-2	420
Histology ICD-O-3	522
Collaborative Stage variables	2800-2930
SEER Summary Stage 1977	760
SEER Summary Stage 2000	759

TEXTHISTOLOGY TITLE				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2590	40	NPCR	4135-4174

Text area for manual documentation of information regarding the histologic type, behavior, and grade (differentiation) of the tumor being reported.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.

Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# Suggestions for text:

- Information on histologic type and behavior
- Information on differentiation from scoring systems such as Gleason's Score, Bloom-Richardson Grade, etc.

# Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Histology (92-00) ICD-O-2	420
Behavior (92-00) ICD-O-2	430
Histologic Type ICD-O-3	522
Behavior Code ICD-O-3	523
Grade	440

<b>TEXTPRIMARY SITE TITLE</b>				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2580	40	NPCR	4095-4134

Text area for manual documentation of information regarding the primary site and laterality of the tumor being reported.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

#### Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

#### **Suggestions for text:**

- Include information on the location of the primary site of the tumor
- Include available information on tumor laterality

#### Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
Primary site	400
Laterality	410

TEXTREMARKS				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2680	350	NPCR	5525-5874

Text area for information that is given only in coded form elsewhere or for which the abstract provides no other place. Overflow data can also be placed here. Problematic coding issues can also be discussed in this section.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

# Suggestions for text:

- Smoking history
- Family and 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
- Place of birth
- Justification of over-ride flags

TEXTSTAGING				Revised
Alternate Name	Item #	Length	Source of Standard	Column #
	2600	300	NPCR	4175-4474

Additional text area for staging information not already entered in the Text--DX Proc areas.

# Rationale

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. High-quality text documentation facilitates consolidation of information from multiple reporting sources at the central registry.

The text field must contain a description that has been entered by the abstractor independently from the code(s). If cancer abstraction software generates text automatically from codes, the text cannot be utilized to check coded values. Information documenting the disease process should be entered manually from the medical record and should not be generated electronically from coded values.

# Instructions

- Prioritize entered information in the order of the fields listed below.
- Text automatically generated from coded data is not acceptable.
- NAACCR-approved abbreviations should be utilized (see Appendix G).
- Do not repeat information from other text fields.
- Additional comments can be continued in empty text fields, including Remarks. For text documentation that is continued from one text field to another, use asterisks or other symbols to indicate the connection with preceding text.
- If information is missing from the record, state that it is missing.
- Do not include irrelevant information.
- Do not include information that the registry is not authorized to collect.

*Note:* For abstracting software that allows unlimited text, NAACCR recommends that the software indicate to the abstractor the portion of the text that will be transmitted to the central registry.

#### **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 specialty and comments

#### Data Item(s) to be verified/validated using the text entered in this field

Item name	Item number
RX DateDX/Stg Proc	1280
Collaborative Stage variables	2800-2930
SEER Summary Stage 1977	760
SEER Summary Stage 2000	759
EODTumor Size	780
EODLymph Node Involv	810
Regional Nodes Positive	820
Regional Nodes Examined	830
Behavior Code ICD-O-3	523
Behavior (92,00) ICD, O, 2	430
Behavior Code ICD-O-3	523
Behavior (92-00) ICD-O-2	430
Site of Distant Met 1-3	1090-1110

# **TEXT--USUAL INDUSTRY**

Alternate Name	Item #	Length	Source of Standard	Column #
	320	40	NPCR	183-222

# Description

Text area for information about the patient's usual industry, also known as usual kind of business/industry.

# Rationale

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

The data item "usual industry" is defined identically as on death certificates and conforms to the 1989 revision of the U.S. Standard Certificate of Death.<sup>25</sup> See related materials in reference list, Chapter VII.

# **Abstracting Instructions**

Record the primary type of activity carried on by the business/industry at the location where the patient was employed for the most number of years before diagnosis of this tumor. Be sure to distinguish among "manufacturing," "wholesale," "retail," and "service" components of an industry that performs more than one of these components.

If the primary activity carried on at the location where the patient worked is unknown, it may be sufficient for facility registrars to record the name of the company (with city or town) in which the patient performed his/her usual industry. In these situations, if resources permit, a central or regional registry may be able to use the employer name and city/town to determine the type of activity conducted at that location.

As noted in the Text--Usual Occupation [310] section, in those situations where the usual occupation is not available or is unknown, the patient's current or most recent occupation is recorded, if available. The information for industry should be based upon the information in occupation. Therefore, if current or most recent occupation rather than usual occupation was recorded, record the patient's current or most recent business/industry.

If later documentation in the patient's record provides an industry that is more likely to be the usual industry than what was originally recorded, facility registrars are encouraged to update the abstract with the new information. However, it is not the responsibility of the facility registrars to update abstracts with industry

information provided on death certificates. Comparison with death certificate information should be the function of a central or regional registry.

There should be an entry for Text--Usual Industry if any occupation is recorded. If no information is available regarding the industry in which the reported occupation was carried out, record "unknown." If the patient was not a student or housewife and had never worked, record "never worked" as the usual industry. This data item usually is collected only for patients who are age 14 years or older at the time of diagnosis.

# **TEXT--USUAL OCCUPATION**

Alternate Name	Item #	Length	Source of Standard	Column #
	310	40	NPCR	143-182

# Description

Text area for information about the patient's usual occupation, also known as usual type of job or work.

# Rationale

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

The data item "usual occupation" is defined identically as on death certificates and conforms to the 1989 revision of the U.S. Standard Certificate of Death.<sup>25</sup> See related materials in reference list, Chapter VII.

# **Abstracting Instructions**

Record the patient's usual occupation (i.e., the kind of work performed during most of the patient's working life before diagnosis of this tumor). Do not record "retired." If usual occupation is not available or is unknown, record the patient's current or most recent occupation, or any available occupation.

If later documentation in the patient's record provides an occupation that is more likely to be the usual occupation than what was originally recorded, facility registrars are encouraged to update the abstract with the new information. However, it is not the responsibility of the facility registrars to update abstracts with occupation information provided on death certificates. Comparison with death certificate information should be the function of a central or regional registry.

If the patient was a househusband/housewife and also worked outside the home during most of his/her adult life, record the usual occupation outside the home; if the patient was a househusband/housewife and did not work outside the home for most of his/her adult life, record "househusband" or "housewife." If the patient was not a student or housewife and had never worked, record "never worked" as the usual occupation.

If no information is available, record "unknown."

This data item usually is collected only for patients who are age 14 years or older at the time of diagnosis.

# TNM CLIN DESCRIPTOR

Alternate Name	Item #	Length	Source of Standard	Column #
Clinical Stage (Prefix/Suffix) Descriptor	980	1	COC	581-581
(COC)				

# Description

Identifies the AJCC clinical stage (prefix/suffix) descriptor as recorded by the physician. AJCC stage descriptors identify special cases that need separate data analysis. The descriptors are adjuncts to and do not change the stage group.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout. COC defines a descriptor and "Staged By" item for each of these three areas.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes

- 0 None
- 1 E (Extranodal, lymphomas only)
- 2 S (Spleen, lymphomas only)
- 3 M (Multiple primary tumors in a single site)
- 4 Y (Classification during or after initial multimodality therapy)—pathologic staging only
- 5 E & S (Extranodal and spleen, lymphomas only)
- 6 M & Y (Multiple primary tumors and initial multimodality therapy)
- 9 Unknown, not stated in patient record

# TNM CLIN M

Alternate Name	Item #	Length	Source of Standard	Column #
Clinical M (COC)	960	2	AJCC	577-578

# Description

Detailed site-specific codes for the clinical metastases (M) as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

#### Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

#### Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

This field can be left blank if a physician did not record the stage element.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

#### TNM CLIN N

Alternate Name	Item #	Length	Source of Standard	Column #
Clinical N (COC)	950	2	AJCC	575-576

#### Description

Detailed site-specific codes for the clinical nodes (N) as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

#### Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

#### Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

This field can be left blank if a physician did not record the stage element.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM CLIN STAGE GROUP

Alternate Name	Item #	Length	Source of Standard	Column #
Clinical Stage Group (COC)	970	2	AJCC	579-580

# Description

Detailed site-specific codes for the clinical stage group as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

- 88 Not applicable.
- 99 Unknown, not staged

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM CLIN STAGED BY

Alternate Name	Item #	Length	Source of Standard	Column #
Staged By (Clinical Stage) (COC)	990	1	COC	582-582

# Description

Identifies the person who recorded the clinical AJCC staging elements and the stage group in the patient's medical record.

# Rationale

Data captured in this field can be used to evaluate the accuracy and completeness of physician staging and form the basis for quality management and improvement studies. This item is used to monitor compliance with the COC Staging Standard. The medical record contains the AJCC stage assigned/initialed by the managing physician.

# Codes (refer to FORDS for additional coding instructions)

- 0 Not staged.
- 1 Managing physician
- 2 Pathologist
- 3 Pathologist and managing physician
- 4 Cancer Committee chair, cancer liaison physician, or registry physician advisor.
- 5 Cancer registrar
- 6 Cancer registrar and physician
- 7 Staging assigned at another facility.
- 8 Case is not eligible for staging.
- 9 Unknown; not stated in patient record

# TNM CLIN T

Alternate Name	Item #	Length	Source of Standard	Column #
Clinical T (COC)	940	2	AJCC	573-574

# Description

Detailed site-specific codes for the clinical tumor (T) as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

This field can be left blank if a physician did not record the stage element.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM EDITION NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
	1060	2	COC	593-594

# Description

A code that indicates the edition of the AJCC manual used to stage the case. This applies to the manually coded AJCC fields. It does not apply to the Derived AJCC T, N, M and AJCC Stage Group fields [2940, 2960, 2980, and 3000].

# Rationale

TNM codes have changed over time and conversion is not always simple. Therefore, a case-specific indicator is needed to allow grouping of cases for comparison.

# Codes

- 00 Not staged (cases that have AJCC staging scheme and staging was not done)
- 01 First Edition
- 02 Second Edition Second Edition (published 1983)
- 03 Third Edition (published 1988)
- 04 Fourth Edition (published 1992), recommended for use for cases diagnosed 1993-1997
- 05 Fifth Edition (published 1997), recommended for use for cases diagnosed 1998-2002
- 06 Sixth Edition (published 2002), recommended for use for cases diagnosed 2003+
- 88 Not applicable (cases that do not have an AJCC staging scheme)
- 99 Edition Unknown

# TNM OTHER DESCRIPTOR

Alternate Name	Item #	Length	Source of Standard	Column #
Other Stage (Prefix/Suffix) Descriptor	1050	1	COC	592-592
(COC)				

# Description

AJCC stage descriptors identify special cases that need separate data analysis. The descriptors are adjuncts to and do not change the stage group.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout. COC defines a descriptor and "Staged By" item for each of these three areas.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes

- 0 None
- 1 E (Extranodal, lymphomas only)
- 2 S (Spleen, lymphomas only)
- 3 M (Multiple primary tumors in a single site)
- 4 Y (Classification during or after initial multimodality therapy)
- 5 E & S (Extranodal and spleen, lymphomas only)
- 6 M & Y (Multiple primary tumors and initial multimodality therapy)
- 9 Unknown, not stated in patient record

# TNM OTHER M

Alternate Name	Item #	Length	Source of Standard	Column #
Other M (COC)	1020	2	AJCC	587-588

# Description

Detailed site-specific codes for the other metastases (M) as defined by AJCC.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

*Note:* See the *AJCC Cancer Staging Manual*, Fifth Edition for site-specific categories for the TNM elements and stage groups. See the *ROADS Manual*, 1998 Supplement, for specifications for codes and data entry rules.

# TNM OTHER N

Alternate Name	Item #	Length	Source of Standard	Column #
Other N (COC)	1010	2	AJCC	585-586

# Description

Detailed site-specific codes for the other nodes (N) as defined by AJCC.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

*Note:* See the *AJCC Cancer Staging Manual*, Fifth Edition for site-specific categories for the TNM elements and stage groups. See the *ROADS Manual*, 1998 Supplement, for specifications for codes and data entry rules.

# TNM OTHER STAGE GROUP

Alternate Name	Item #	Length	Source of Standard	Column #
Other Stage Group (COC)	1030	2	AJCC	589-590

# Description

Detailed site-specific codes for the other stage group as defined by AJCC.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

- 88 Not applicable.
- 99 Unknown, not staged

*Note:* See the *AJCC Cancer Staging Manual*, Fifth Edition for site-specific categories for the TNM elements and stage groups. See the *ROADS Manual*, 1998 Supplement, for specifications for codes and data entry rules.

# TNM OTHER STAGED BY

Alternate Name	Item #	Length	Source of Standard	Column #
Staged By (Other Stage) (COC)	1040	1	COC	591-591

# Description

AJCC "Staged By" fields identify the person who documented the AJCC staging elements and stage group. COC requires analytic cases to be staged by the managing physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout. COC defines a descriptor and "Staged By" item for each of these three areas.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes

- 0 Not staged
- 1 Managing physician
- 2 Pathologist
- 3 Other physician
- 4 Any combination of 1, 2, or 3
- 5 Registrar
- 6 Any combination of 5 with 1, 2, or 3
- 7 Other
- 8 Staged, individual not specified
- 9 Unknown if staged

# TNM OTHER T

Alternate Name	Item #	Length	Source of Standard	Column #
Other T (COC)	1000	2	AJCC	583-584

# Description

Detailed site-specific codes for the other tumor (T) as defined by AJCC.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

The "other" TNM components and Stage group allow collection of retreatment staging. They also may be used for historic data such as the surgical TNM and stage group items. The "other" categories may be used for converted data if the staging basis is not identified.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

*Note:* See the *AJCC Cancer Staging Manual*, Fifth Edition for site-specific categories for the TNM elements and stage groups. See the *ROADS Manual*, 1998 Supplement, for specifications for codes and data entry rules.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# TNM PATH DESCRIPTOR

Alternate Name	Item #	Length	Source of Standard	Column #
Pathologic Stage (Prefix/Suffix) Descriptor	920	1	COC	571-571
(COC)				

# Description

Identified the AJCC clinical stage (prefix/suffix) descriptor as recorded by the physician. AJCC stage descriptors identify special cases that need separate data analysis. The descriptors are adjuncts to and do not change the stage group.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout. COC defines a descriptor and "Staged By" item for each of these three areas.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes

- 0 None
- 1 E (Extranodal, lymphomas only)
- 2 S (Spleen, lymphomas only)
- 3 M (Multiple primary tumors in a single site)
- 4 Y (Classification during or after initial multimodality therapy)—pathologic staging only
- 5 E & S (Extranodal and spleen, lymphomas only)
- 6 M & Y (Multiple primary tumors and initial multimodality therapy)
- 9 Unknown, not stated in patient record

# TNM PATH M

Alternate Name	Item #	Length	Source of Standard	Column #
Pathologic M (COC)	900	2	AJCC	567-568

# Description

Detailed site-specific codes for the pathologic metastases (M) as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

This field can be left blank if a physician did not record the stage element.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM PATH N

Alternate Name	Item #	Length	Source of Standard	Column #
Pathologic N (COC)	890	2	AJCC	565-566

# Description

Detailed site-specific codes for the pathologic nodes (N) as defined by AJCC and recorded by physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

# 88 Not applicable

This field can be left blank if a physician did not record the stage element.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM PATH STAGE GROUP

Alternate Name	Item #	Length	Source of Standard	Column #
Pathologic Stage Group (COC)	910	2	AJCC	569-570

# Description

Detailed site-specific codes for the pathologic stage group as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

- 88 Not applicable.
- 99 Unknown, unstaged

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# TNM PATH STAGED BY

Alternate Name	Item #	Length	Source of Standard	Column #
Staged By (Pathologic Stage) (COC)	930	1	COC	572-572

# Description

Identifies the person who recorded the pathologic AJCC staging elements and the stage group in the patient's medical record.

# Rationale

Data captured in this field can be used to evaluate the accuracy and completeness of physician staging and form the basis for quality management and improvement studies. This item is used to monitor compliance with the COC Staging Standard. The medical record contains the AJCC stage assigned/initialed by the managing physician.

# Codes (refer to FORDS for additional coding instructions)

- 0 Not staged.
- 1 Managing physician
- 2 Pathologist
- 3 Pathologist and managing physician.
- 4 Cancer Committee chair, cancer liaison physician, or registry physician advisor.
- 5 Cancer registrar
- 6 Cancer registrar and physician.
- 7 Staging assigned at another facility.
- 8 Case is not eligible for staging.
- 9 Unknown; not stated in patient record.

# TNM PATH T

Alternate Name	Item #	Length	Source of Standard	Column #
Pathologic T (COC)	880	2	AJCC	563-564

# Description

Detailed site-specific codes for the pathologic tumor (T) as defined by AJCC and recorded by the physician.

Pathologic, clinical, and other stage data are given three separate areas in the NAACCR Data Exchange Record Layout.

# Rationale

COC requires that AJCC TNM staging be used in its approved cancer programs. AJCC developed its staging system for evaluating trends in the treatment and control of cancer. This staging is used by physicians to estimate prognosis, to plan treatment, to evaluate new types of therapy, to analyze outcome, to design follow-up strategies, and to assess early detection results.

# Codes (in addition to those published in the AJCC Cancer Staging Manual)

88 Not applicable.

*Note:* See the *AJCC Cancer Staging Manual*, current edition for site-specific categories for the TNM elements and stage groups. See the *FORDS* Manual for specifications for codes and data entry rules.

# **TOBACCO HISTORY**

Alternate Name	Item #	Length	Source of Standard	Column #
	340	1	Varies	224-224

# Description

NAACCR has not adopted standards for this item.

# **TUMOR MARKER 1**

Alternate Name	Item #	Length	Source of Standard	Column #
Tumor Marker One (COC)	1150	1	SEER	626-626

# Description

Records prognostic indicators for specific sites or histologies. COC uses these codes for cases diagnosed 1996 and forward. See the COC *ROADS Manual*, 1998 Supplement, for a list of specific sites and histologies. For tumors diagnosed before January 1, 1996, Tumor Marker 1 is coded only for estrogen receptor status of breast cancers.

For SEER requirements for the specific sites, histologies, and diagnosis years for which this item is coded, see the 1998 SEER Program Code Manual.

# Codes

- 0 None done (SX)
- 1 Positive/elevated
- 2 Negative/normal; within normal limits (S0)
- 3 Borderline; undetermined whether positive/elevated or negative/normal

Three-tiered system:

- 4 Range 1 (S1)
- 5 Range 2 (S2)
- 6 Range 3 (S3)
- 8 Ordered, but results not in chart
- 9 Unknown or no information

For sites for which Tumor Marker 1 is not collected:

9 Not applicable

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

# TUMOR MARKER 2

Alternate Name	Item #	Length	Source of Standard	Column #
Tumor Marker Two (COC)	1160	1	SEER	627-627

# Description

Records prognostic indicators for specific sites or histologies. COC uses these codes for cases diagnosed 1996 and forward. See the COC *ROADS Manual*, 1998 Supplement, for a list of specific sites and histologies. For tumors diagnosed before January 1, 1996, Tumor Marker 2 is coded only for progesterone receptor status of breast cancers.

For SEER requirements for the specific sites, histologies, and diagnosis years for which this item is coded, see the 1998 SEER Program Code Manual.

# Codes

- 0 None done (SX)
- 1 Positive/elevated
- 2 Negative/normal; within normal limits (S0)
- 3 Borderline; undetermined whether positive/elevated or negative/normal

Three-tiered system:

- 4 Range 1 (S1)
- 5 Range 2 (S2)
- 6 Range 3 (S3)
- 8 Ordered, but results not in chart
- 9 Unknown or no information

For sites for which Tumor Marker 2 is not collected:

9 Not applicable

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

# TUMOR MARKER 3

Alternate Name	Item #	Length	Source of Standard	Column #
Tumor Marker Three (COC)	1170	1	SEER	628-628

# Description

Records prognostic indicators for specific sites or histologies. COC uses these codes for tumors diagnosed 1998 and forward. See the COC *ROADS Manual*, 1998 Supplement, for a list of specific sites and histologies.

For SEER requirements for the specific sites, histologies, and diagnosis years for which this item is coded, see the 1998 SEER Program Code Manual.

# Codes

- 0 None done (SX)
- 1 Positive/elevated
- 2 Negative/normal; within normal limits (S0)
- 3 Borderline; undetermined whether positive/elevated or negative/normal

Three-tiered system:

- 4 Range 1 (S1)
- 5 Range 2 (S2)
- 6 Range 3 (S3)
- 8 Ordered, but results not in chart
- 9 Unknown or no information

For sites for which Tumor Marker 3 is not collected:

9 Not applicable

*Note:* As of January 1, 2003, this data item is no longer required or recommended by COC. However, the item was collected in the past and it is recommended that historic data be retained.

# TUMOR RECORD NUMBER

Alternate Name	Item #	Length	Source of Standard	Column #
	60	2	NAACCR	30-31

# Description

A system-generated number assigned to each tumor. The number should never change even if the tumor sequence is changed or a record (tumor) is deleted.

# Rationale

This is a unique number that identifies a specific tumor so data can be linked. "Sequence Number" cannot be used as a link because the number is changed if a report identifies an earlier tumor or if a tumor record is deleted.

# **TYPE OF REPORTING SOURCE**

Alternate Name	Item #	Length	Source of Standard	Column #
	500	1	SEER	312-312

# Description

Code identifying source documents used to abstract the tumor being reported. This may not be the source of original case finding (for example, if a case is identified through a pathology laboratory report review and all source documents used to abstract the case are from the physician's office, code this item 4).

Type of Reporting Source can be used in conjunction with Class of Case [610]. Class of case is designed to differentiate between analytic and non-analytic cases at the hospital level.

See Chapter V, Unresolved Issues, for a discussion of inadequacies in this item.

# Rationale

The code in this field can be used to explain why information may be incomplete on a case. The field also is used to monitor the success of non-hospital case reporting and follow-back mechanisms. All population-based registries should have some death certificate-only cases where no hospital admission was involved, but too high a percentage can imply that follow-back to uncover missed hospital reports was not complete.

# Codes

- 1 Hospital inpatient/outpatient or clinic
- 3 Laboratory only (hospital or private)
- 4 Physician's office/private medical practitioner (LMD)
- 5 Nursing/convalescent home/hospice
- 6 Autopsy only
- 7 Death certificate only

Note: Coding is hierarchical. Within codes 1-5, assign codes in the following priority: 1, 4, 5, 3.

# UNUSUAL FOLLOW-UP METHOD

Alternate Name	Item #	Length	Source of Standard	Column #
	1850	1	COC	1341-1341

# Description

User-defined numeric codes used to flag cases that need unusual follow-up methods.

# Codes

User-defined

Note: This data item is no longer supported by COC (as of January 1, 2003).

# VENDOR NAME

Alternate Name	Item #	Length	Source of Standard	Column #
	2170	10	NAACCR	1204-1213

# Description

System-generated. Name of the computer services vendor who programmed the system submitting the data. Abbreviate as necessary and keep a consistent name throughout all submissions. Include software version number where available. Code is self-assigned by vendor.

# Rationale

This is used to track which vendor and which software version submitted the case. It helps define the source and extent of a problem discovered in data submitted by a software provider.

# VITAL STATUS

Alternate Name	Item #	Length	Source of Standard	Column #
	1760	1	SEER/COC	1302-1302

# Description

Vital status of the patient as of the date entered in Date of Last Contact [1750]. If the patient has multiple tumors, vital status should be the same for all tumors.

# Codes

0 Dead (COC)

- 1 Alive
- 4 Dead (SEER)

# YEAR FIRST SEEN THIS CA

Alternate Name	Item #	Length	Source of Standard	Column #
Accession Year (pre-96 COC)	620	4	COC	441-444
Year First Seen for this Primary (COC)				

# Description

Year patient was first seen at the reporting institution for diagnosis and/or treatment of this primary, since the reference date of the registry. It is not the year that the registrar accessioned the case.

# Rationale

This data item is used by hospital registries to organize their case reporting into individual years. It differs from the first 4 digits of the Accession Number, because this variable is tumor-specific rather than patient-specific, and from the diagnosis year because it relates to the specific facility and not the tumor. Central registries that wish to compare their data with hospital case lists can make use of this field to create equivalent reports.

Note: This data item is no longer supported by COC (as of January 1, 2003).

# APPENDIX A

# FIPS CODES FOR COUNTIES AND EQUIVALENT ENTITIES

[Ed. Note: The information in this table is from FIPS Publication Number 6-4, "Counties and Equivalent Entities of the United States, its Possessions, and Associated Areas," as reissued December 21, 1992, and made available electronically on the National Institute of Standards and Technology Web Site (http://www.itl.nist.gov). We compared two versions of the file against printed lists to reconcile apparent errors and discrepancies.]

	E NAME:	087	Macon	090	Fairbanks North Star	023	Santa Cruz
ALAB		089	Madison		(B)	025	Yavapai
	ABETIC CODE: AL	091	Marengo	100	Haines (B)	027	Yuma
NUMI	ERIC CODE: 01	093	Marion	110	Juneau (B)		
		095	Marshall	122	Kenai Peninsula (B)		was established from
	E COUNTY NAME	097	Mobile	130	Ketchikan Gateway	part of	Yuma (1/1/83).
001	Auatauga	099	Monroe		(B)		
003	Baldwin	101	Montgomery	150	Kodiak Island (B)		
005	Barbour	103	Morgan	164	Lake and Peninsula	STATE	E NAME:
007	Bibb	105	Perry		(B)	ARKA	NSAS
009	Blount	107	Pickens	170	Matanuska-Susitna	ALPH	ABETIC CODE: AR
011	Bullock	109	Pike		(B)	NUME	RIC CODE: 05
013	Butler	111	Randolph	180	Nome (C)		
015	Calhoun	113	Russell	185	North Slope (B)	CODE	COUNTY NAME
017	Chambers	115	St. Clair	188	Northwest Arctic	001	Arkansas
019	Cherokee	117	Shelby		(B)	003	Ashley
021	Chilton	119	Sumter	201	Prince of Wales-	005	Baxter
023	Choctaw	121	Talladega		Outer Ketchikan (C)	007	Benton
025	Clarke	123	Tallapoosa	220	Sitka (B)	009	Boone
027	Clay	125	Tuscaloosa	232	Skagway-Hoonah-	011	Bradley
029	Cleburne	127	Walker		Angoon (C)	013	Calhoun
031	Coffee	129	Washington	240	Southeast Fairbanks	015	Carroll
033	Colbert	131	Wilcox		(C)	017	Chicot
035	Conecuh	133	Winston	261	Valdez-Cordova (C)	019	Clark
037	Coosa			270	Wade Hampton (C)	021	Clay
039	Covington			280	Wrangell-Petersburg	023	Cleburne
041	Crenshaw	STAT	E NAME: ALASKA		(C)	025	Cleveland
043	Cullman		ABETIC CODE:	282	Yakutat (B)	027	Columbia
045	Dale	AK		290	Yukon-Koyukuk (C)	029	Conway
047	Dallas		ERIC CODE: 02	220	r unon rioj unun (c)	031	Craighead
049	DeKalb					033	Crawford
051	Elmore	Note: 7	The following is a	STATE	E NAME:	035	Crittenden
053	Escambia		ete list of all current	ARIZO		037	Cross
055	Etowah	1	county equivalents		ABETIC CODE: AZ	039	Dallas
055	Fayette		(B) identifies a		RIC CODE: 03	041	Desha
059	Franklin		h and (C) identifies a			043	Drew
061	Geneva		area per FIPS	CODE	COUNTY NAME	045	Faulkner
063	Greene		ation Change Notice	001	Apache	047	Franklin
065	Hale		ie 12/21/92).	001	Cochise	049	Fulton
067	Henry	(101330	ie 12/21/92).	005	Coconino	051	Garland
069	Houston	CODE	BOROUGH/	005	Gila	053	Grant
071	Jackson	CODE	CENSUS AREA	009	Graham	055	Greene
071	Jefferson	013	Aleutians East (B)	011	Greenlee	055	Hempstead
075	Lamar	013	Aleutians West (C)	011	LaPaz	057	Hot Spring
073	Lauderdale	010	Anchorage (B)	012	Maricopa	059	Howard
077	Lauderdale	020	Bethel (C)	015	Mohave	061	Independence
079	Lawrence	030	Bristol Bay (B)	013	Navajo	065	Izard
081		060		017	Navajo Pima	065	Jackson
083	Limestone Lowndes	068	Denali (B)	019	Pinal	067	Jackson Jefferson
085	Lownues	070	Dillingham (C)	021	r mäi	009	JEHEISOH

071	Johnson
073	Lafayette
075	Lawrence
077	Lee
079	Lincoln
081	Little River
083	Logan
085	Lonoke
087	Madison
089	Marion
091	Miller
093	Mississippi
095	Monroe
097	Montgomery
099	Nevada
101	Newton
103	Ouachita
105	Perry
107	Phillips
109	Pike
111	Poinsett
113	Polk
115	Pope
117	Prairie
119	Pulaski
121	Randolph
123	St. Francis
125	Saline
127	Scott
129	Searcy
131	Sebastian
133	Sevier
135	Sharp
137	Stone
139	Union
141	Van Buren
143	Washington
145	White
147	Woodruff
149	Yell
-	

### STATE NAME: CALIFORNIA ALPHABETIC CODE: CA **NUMERIC CODE: 06**

#### CODE COUNTY NAME 001 Alameda

001	1 manneau
003	Alpine
005	Amador
007	Butte
009	Calaveras
011	Colusa
013	Contra Costa
015	Del Norte
017	El Dorado
019	Fresno
021	Glenn
023	Humboldt
025	Imperial
027	Inyo
029	Kern
031	Kings

033	Lake
035	Lassen
037	Los Angeles
039	Madera
041	Marin
043	Mariposa
045	Mendocino
047	Merced
049	Modoc
051	Mono
053	Monterey
055	Napa
057	Nevada
059	Orange
061	Placer
063	Plumas
065	Riverside
067	Sacramento
069	San Benito
071	San Bernardino
073	San Diego
075	San Francisco
077	San Joaquin
079	San Luis Obispo
081	San Mateo
083	Santa Barbara
085	Santa Clara
087	Santa Cruz
089	Shasta
091	Sierra
093	Siskiyou
095	Solano
097	Sonoma
099	Stanislaus
101	Sutter
103	Tehama
105	Trinity
107	Tulare
109	Tuolomne
111	Ventura
113	Yolo
115	Yuba

STATE NAME: **COLORADO ALPHABETIC CODE:** со NUMERIC CODE: 08

### CODE COUNTY NAME

Adams 001 003 Alamosa 005 Arapahoe 007 Archuleta 009 Baca 011 Bent 013 Boulder 014 Broomfield 015 Chaffee 017 Cheyenne 019 Clear Creek 021 Conejos Costilla 023

025 Crowley 027 Custer 029 Delta 031 Denver 033 Dolores 035 Douglas 037 Eagle 039 Elbert 041 El Paso 043 Fremont 045 Garfield 047 Gilpin 049 Grand 051 Gunnison 053 Hinsdale 055 Huerfano 057 Jackson 059 Jefferson 061 Kiowa 063 Kit Carson 065 Lake 067 La Plata 069 Larimer 071 Las Animas 073 Lincoln 075 Logan 077 Mesa 079 Mineral 081 Moffat 083 Montezuma 085 Montrose 087 Morgan 089 Otero 091 Ouray 093 Park 095 Phillips 097 Pitkin 099 Prowers 101 Pueblo 103 Rio Blanco 105 Rio Grande 107 Routt 109 Saguache 111 San Juan 113 San Miguel 115 Sedgwick Summit 117 119 Teller 121 Washington 123 Weld 125 Yuma Broomfield County, Colorado, has been created from parts of Adams (001), Boulder (013), Jefferson (059) and Weld (123)

a code of 014 for FIPS 6-4. STATE NAME: CONNECTICUT ALPHABETIC CODE: CT NUMERIC CODE: 09

sequences of counties,

Broomfield County will have

### CODE COUNTY NAME

001 Fairfield 003 Hartford 005 Litchfield 007 Middlesex 009 New Haven 011 New London 013 Tolland 015 Windham

### STATE NAME: DELAWARE ALPHABETIC CODE: DE **NUMERIC CODE: 10**

CODE COUNTY NAME 001 Kent

003 New Castle 005 Sussex

### STATE NAME: DISTRICT OF **COLUMBIA ALPHABETIC CODE: DC** NUMERIC CODE: 11

## CODE SUBDIVISION NAME

001 District of Columbia

Name was reported incorrectly as "Washington" in FIPS PUB 6-3. The District has no first-order subdivisions, and therefore "District of Columbia" also serves as the countyequivalent entity.

### STATE NAME: FLORIDA ALPHABETIC CODE: FL NUMERIC CODE: 12

CODE COUNTY NAME 001 Alachua 003 Baker 005 Bay 007 Bradford 009 Brevard Broward 011 013 Calhoun

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

counties effective November

15, 2001. The boundaries of

the boundaries of Broomfield

Broomfield County reflect

city legally in effect on

November 15, 2001. To

maintain alphanumeric

015	Charlotte	Conver	t Dade County 025 to
017	Citrus	Miami-	Dade County 086.
019	Clay	Edits sł	nould only allow for
021	Collier	code 08	36.
023	Columbia		I
027	DeSoto		
029	Dixie	STATE	E NAME:
02)	Duval	GEOR	
033	Escambia		ABETIC CODE:
		GA	ABETIC CODE:
035	Flagler		DIC CODE 12
037	Franklin	NUME	RIC CODE: 13
039	Gadsden	~~~~	~~~~~
041	Gilchrist		COUNTY NAME
043	Glades	001	Appling
045	Gulf	003	Atkinson
047	Hamilton	005	Bacon
049	Hardee	007	Baker
051	Hendry	009	Baldwin
053	Hernando	011	Banks
055	Highlands	013	Barrow
057	Hillsborough	015	Bartow
059	Holmes	017	Ben Hill
061	Indian River	019	Berrien
063	Jackson	021	Bibb
065	Jefferson	023	Bleckley
067	Lafayette	025	Brantley
069	Lake	027	Brooks
071	Lee	029	Bryan
073	Leon	031	Bulloch
075	Levy	033	Burke
077	Liberty	035	Butts
079	Madison	035	Calhoun
081	Manatee	039	Camden
081	Marion	039	Candler
085	Martin	045	Carroll
085	Miami-Dade	043	Catoosa
080	Monroe	049	Charlton
089	Nassau	051	Chatham
089	Okaloosa	053	Chattahoochee
091	Okeechobee	055	
095		055	Chattooga Cherokee
093	Orange Osceola	059	Clarke
097	Palm Beach	059	
101		061	Clay
	Pasco		Clayton
103	Pinellas	065	Clinch
105	Polk	067	Cobb
107	Putnam	069	Coffee
109	St. Johns	071	Colquitt
111	St. Lucie	073	Columbia
113	Santa Rosa	075	Cook
115	Sarasota	077	Coweta
117	Seminole	079	Crawford
119	Sumter	081	Crisp
121	Suwannee	083	Dade
123	Taylor	085	Dawson
125	Union	087	Decatur
127	Volusia	089	DeKalb
129	Wakulla	091	Dodge
131	Walton	093	Dooly
133	Washington	095	Dougherty
		097	Douglas
		099	Early
		101	Echols
		103	Effingam

105	Elbert	235
107	Emanuel	237
109	Evans	239
111	Fannin	241
113	Fayette	243
115	Floyd	245
117	Forsyth	247
119	Franklin	249
121	Fulton	251
123	Gilmer	253
125	Glascock	255
127 129	Glynn Gordon	257 259
129	Grady	239
133	Greene	263
135	Gwinnett	265
137	Habersham	267
139	Hall	269
141	Hancock	271
143	Haralson	273
145	Harris	275
147	Hart	277
149	Heard	279
151	Henry	281
153	Houston	283
155	Irwin	285
157	Jackson	287
159	Jasper	289
161	Jeff Davis	291
163 165	Jefferson Jenkins	293 295
167	Johnson	293 297
169	Jones	299
171	Lamar	301
173	Lanier	303
175	Laurens	305
177	Lee	307
179	Liberty	309
181	Lincoln	311
183	Long	313
185	Lowndes	315
187	Lumpkin	317
189	McDuffie	319
191	McIntosh	321
193	Macon	M
195 197	Madison Marion	Mus
197	Meriwether	(con
201	Miller	(510
201	Mitchell	(510
207	Monroe	
209	Montgomery	STA
211	Morgan	ALI
213	Murray	NUI
215	Muscogee	
217	Newton	CO
219	Oconee	001
221	Oglethorpe	003
223	Paulding	005
225	Peach	007
227	Pickens	009
229	Pierce	17.1
231 233	Pike Polk	Kala
233	I UIK	own

235	Pulaski
237	Putnam
239	Quitman
241	Rabun
243	Randolph
245	Richmond
247	Rockdale
249	Schley
251	Screven
253	Seminole
255	Spalding
257	Stephens
259	Stewart
261	Sumter
263	Talbot
265	Taliaferro
267	Tattnall
269	Taylor
271	Telfair
273	Terrell
275	Thomas
277	Tift
279	Toombs
281	Towns
283	Treutlen
285	Troup
285	Turner
289	Twiggs
20)	Union
293	Upson
295	Walker
293	Walton
299	Ware
301	
303	Warren
305	Washington
	Wayne
307	Webster
309	Wheeler
311	White
313	Whitfield
315	Wilcox
317	Wilkes
319	Wilkinson
321	Worth
м	
wiuscog	ee was reported
incorrec	tly as "Columbus
	dated government)"
(510) in	FIPS PUB6-3.

### STATE NAME: HAWAII ALPHABETIC CODE: HI NUMERIC CODE: 15

CODE	COUNTY NAME
001	Hawaii
003	Honolulu
005	Kalawao
007	Kauai
009	Maui

Kalawao does not have its own local government; it is

administered by the State of
Hawaii. It may be included
with Maui for statistical
purposes.

### STATE NAME: IDAHO ALPHABETIC CODE: ID **NUMERIC CODE: 16**

# CODE COUNTY NAME

001	Ada
003	Adams
005	Bannock
007	Bear Lake
009	Benewah
011	Bingham
013	Blaine
015	Boise
017	Bonner
019	Bonneville
021	Boundary
023	Butte
025	Camas
027	Canyon
029	Caribou
031	Cassia
033	Clark
035	Clearwater
037	Custer
039	Elmore
041	Franklin
043	Fremont
045	Gem
047	Gooding
049	Idaho
051	Jefferson
053	Jerome
055	Kootenai
057	Latah
059	Lemhi
061	Lewis
063	Lincoln
065	Madison
067	Minidoka
069	Nez Perce
071	Oneida
073	Owyhee
075	Payette
077	Power
079	Shoshone
081	Teton
083	Twin Falls
085	Valley
087	Washington
STATE	NAME ILLING

### STATE NAME: ILLINOIS ALPHABETIC CODE: IL NUMERIC CODE: 17

### CODE COUNTY NAME

001 Adams 003 Alexander

005	Bond
007	Boone
009	Brown
011	Bureau
013	Calhoun
015	Carroll
017	Cass
019	Champaign
021	Christian
023	Clark
025	Clay
027	Clinton
029	Coles
031	Cook
033	Crawford
035	Cumberland
037	DeKalb
039	De Witt
041	Douglas
043	DuPage
045	Edgar
047	Edwards
049	Effingham
051	Fayette
	•
053	Ford
055	Franklin
057	Fulton
059	Gallatin
061	Greene
063	Grundy
065	Hamilton
067	Hancock
069	Hardin
071	Henderson
073	Henry
075	Iroquois
077	Jackson
079	Jasper
081	Jefferson
083	Jersey
085	Jo Daviess
087	Johnson
089	Kane
091	Kankakee
093	Kendall
095	Knox
097	Lake
099	La Salle
101	Lawrence
103	Lee
105	Livingston
107	Logan
	U
109	McDonough
111	McHenry
111 113	
113	McLean
113 115	McLean Macon
113	McLean
113 115	McLean Macon
113 115 117 119	McLean Macon Macoupin Madison
113 115 117 119 121	McLean Macon Macoupin Madison Marion
<ol> <li>113</li> <li>115</li> <li>117</li> <li>119</li> <li>121</li> <li>123</li> </ol>	McLean Macon Macoupin Madison Marion Marshall
<ol> <li>113</li> <li>115</li> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> </ol>	McLean Macon Macoupin Madison Marion Marshall Mason
<ol> <li>113</li> <li>115</li> <li>117</li> <li>119</li> <li>121</li> <li>123</li> </ol>	McLean Macon Macoupin Madison Marion Marshall
<ol> <li>113</li> <li>115</li> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> </ol>	McLean Macon Macoupin Madison Marion Marshall Mason Massac
<ol> <li>113</li> <li>115</li> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> </ol>	McLean Macon Macoupin Madison Marion Marshall Mason

133	Monroe
135	Montgomery
137	Morgan
139	Moultrie
141	Ogle
143	Peoria
145	Perry
147	Piatt
149	Pike
151	Pope
153	Pulaski
155	Putnam
157	Randolph
159	Richland
161	Rock Island
163	St. Clair
165	Saline
167	Sangamon
169	Schuyler
171	Scott
173	Shelby
175	Stark
177	Stephenson
179	Tazewell
181	Union
183	Vermilion
185	Wabash
187	Warren
189	Washington
191	Wayne
193	White
195	Whiteside
197	Will
199	Williamson
201	Winnebago
203	Woodford
	E NAME: INDIANA
	ABETIC CODE: IN
NUME	RIC CODE: 18
CODE	COUNTY NAME
001	Adams
001	Allen
005	Bartholomew
003	Benton
007	Blackford
011	Boone
~ 1 1	

### 011 Boone Brown 013 015 Carroll 017 Cass 019 Clark 021 Clay 023 Clinton 025 Crawford 027 Daviess 029 Dearborn 031 Decatur DeKalb 033 035 Delaware 037 Dubois 039 Elkhart Fayette 041

043 045 047 049 051 053 055 057 059 061 063 065 067 079 071 073 075 077 079 081 083 085 087 099 101 103 105 107 109 111 113 115 117 119 121 123 125 127 131 133 135 137 139 141 143 145 147 149 151 157 159 161 163 155 157 159 161 163 155 157 159 161 163 165 167 169 161 163 165 167 169 107 107 107 107 107 107 107 107	Floyd Fountain Franklin Fulton Gibson Grant Greene Hamilton Hancock Harrison Hendricks Henry Howard Huntington Jackson Jasper Jay Jefferson Jasper Jay Jefferson Jackson Jasper Jay Jefferson Knox Kosciusko Lagrange Lake LaPorte Lawrence Madison Marion Marion Marshall Martin Martin Miami Monroe Montgomery Morgan Newton Noble Ohio Orange Owen Parke Perry Pike Porter Porsey Pulaski Putnam Randolph Ripley Rush Scott Shelby Spencer Starke Steuben Sullivan Switzerland Tippecanoe Tipton Unon debrev
155	Switzerland
157	Tippecanoe
159	Tipton
161	Union
163	Vanderburgh
165	Vermillion
167	Vigo
169	Wabash

171	Warren	099	Jasper	015	Butler	143	Ottawa
173	Warrick	101	Jefferson	017	Chase	145	Pawnee
175	Washington	103	Johnson	019	Chautauqua	147	Phillips
177	Wayne	105	Jones	021	Cherokee	149	Pottawatomie
179	Wells	107	Keokuk	023	Cheyenne	151	Pratt
181	White	109	Kossuth	025	Clark	153	Rawlins
183	Whitley	111	Lee	023	Clay	155	Reno
165	whitey		Linn	027	Cloud	155	Republic
		113					1
		115	Louisa	031	Coffey	159	Rice
	TE NAME:	117	Lucas	033	Comanche	161	Riley
IOV		119	Lyon	035	Cowley	163	Rooks
	PHABETIC CODE: IA	121	Madison	037	Crawford	165	Rush
NUN	MERIC CODE: 19	123	Mahaska	039	Decatur	167	Russell
		125	Marion	041	Dickinson	169	Saline
COI	DE COUNTY NAME	127	Marshall	043	Doniphan	171	Scott
001	Adair	129	Mills	045	Douglas	173	Sedgwick
003	Adams	131	Mitchell	047	Edwards	175	Seward
005	Allamakee	133	Monona	049	Elk	177	Shawnee
007	Appanoose	135	Monroe	051	Ellis	179	Sheridan
009	Audubon	135	Montgomery	053	Ellsworth	181	Sherman
011	Benton	139	Muscatine	055	Finney	181	Smith
		139	O'Brien		5	185	
013	Black Hawk			057	Ford		Stafford
015	Boone	143	Osceola	059	Franklin	187	Stanton
017	Bremer	145	Page	061	Geary	189	Stevens
019	Buchanan	147	Palo Alto	063	Gove	191	Sumner
021	Buena Vista	149	Plymouth	065	Graham	193	Thomas
023	Butler	151	Pocahontas	067	Grant	195	Trego
025	Calhoun	153	Polk	069	Gray	197	Wabaunsee
027	Carroll	155	Pottawattamie	071	Greeley	199	Wallace
029	Cass	157	Poweshiek	073	Greenwood	201	Washington
031	Cedar	159	Ringgold	075	Hamilton	203	Wichita
033	Cerro Gordo	161	Sac	077	Harper	205	Wilson
035	Cherokee	163	Scott	079	Harvey	207	Woodson
037	Chickasaw	165	Shelby	081	Haskell	207	Wyandotte
		167	Sioux	081		209	w yandotte
039	Clarke				Hodgeman		
041	Clay	169	Story	085	Jackson	6 m + m	
043	Clayton	171	Tama	087	Jefferson		E NAME:
045	Clinton	173	Taylor	089	Jewell		TUCKY
047	Crawford	175	Union	091	Johnson		ABETIC CODE:
049	Dallas	177	Van Buren	093	Kearny	KY	
051	Davis	179	Wapello	095	Kingman	NUM	ERIC CODE: 21
053	Decatur	181	Warren	097	Kiowa		
055	Delaware	183	Washington	099	Labette	CODE	COUNTY NAME
057	Des Moines	185	Wayne	101	Lane	001	Adair
059	Dickinson	187	Webster	103	Leavenworth	003	Allen
061	Dubuque	189	Winnebago	105	Lincoln	005	Anderson
063	Emmet	191	Winneshiek	107	Linn	007	Ballard
065	Fayette	193	Woodbury	109	Logan	009	Barren
067	Floyd	195	Worth	111	Lyon	011	Bath
069	Franklin	195	Wright	113	McPherson	013	Bell
009	Fremont	197	wiight	115	Marion	015	Boone
073	Greene			117	Marshall	017	Bourbon
075	Grundy		E NAME: KANSAS	119	Meade	019	Boyd
077	Guthrie		ABETIC CODE: KS	121	Miami	021	Boyle
079	Hamilton	NUM	ERIC CODE: 20	123	Mitchell	023	Bracken
081	Hancock			125	Montgomery	025	Breathitt
083	Hardin	CODE	COUNTY NAME	127	Morris	027	Breckinridge
085	Harrison	001	Allen	129	Morton	029	Bullitt
087	Henry	003	Anderson	131	Nemaha	031	Butler
089	Howard	005	Atchison	133	Neosho	033	Caldwell
091	Humboldt	007	Barber	135	Ness	035	Calloway
093	Ida	009	Barton	135	Norton	037	Campbell
095	Iowa	011	Bourbon	139	Osage	039	Carlisle
097	Jackson	013	Brown	141	Osborne	041	Carroll
091	JUCKSOII	015	DIOWII	141	Osoonie	041	Carton

043	Carter
045	Casey
047	Christian
049	Clark
051	Clay
	Cluy
053	Clinton
055	Crittenden
057	Cumberland
059	Daviess
061	Edmonson
063	Elliott
065	Estill
067	Fayette
069	Fleming
071	Floyd
073	Franklin
075	Fulton
077	Gallatin
079	Garrard
081	Grant
083	Graves
	-
085	Grayson
087	Green
089	Greenup
091	Hancock
093	Hardin
095	Harlan
097	Harrison
099	Hart
101	Henderson
103	Henry
105	Hickman
107	Hopkins
109	Jackson
111	Jefferson
113	Jessamine
115	Johnson
117	Kenton
119	Knott
121	Knox
123	Larue
125	Laurel
127	Lawrence
129	Lee
131	Leslie
133	
	Letcher
135	Lewis
137	Lincoln
139	Livingston
141	Logan
	-
143	Lyon
145	McCracken
147	McCreary
149	McLean
151	Madison
153	Magoffin
155	Marion
157	Marshall
159	M C
	Martin
161	Mason
161 163	Mason Meade
161 163 165	Mason Meade Menifee
161 163	Mason Meade
161 163 165	Mason Meade Menifee

171	Monroe
173	Montgomery
175	Morgan
177	Muhlenberg
179	Nelson
181	Nicholas
183	Ohio
185	Oldham
187	Owen
189	Owsley
	•
191	Pendleton
193	Perry
195	Pike
197	Powell
199	Pulaski
201	Roberston
203	Rockcastle
205	Rowan
207	Russell
209	Scott
211	Shelby
213	Simpson
	Spencer
215	
217	Taylor
219	Todd
221	Trigg
223	Trimble
225	Union
227	Warren
229	Washington
231	Wayne
233	Webster
235	Whitley
237	Wolfe
237	Wolfe Woodford
	Wolfe Woodford
237 239	Woodford
237 239 STATE	Woodford E NAME:
237 239 STATE LOUIS	Woodford E NAME: JANA
237 239 STATE LOUIS ALPHA	Woodford E NAME: JANA ABETIC CODE: LA
237 239 STATE LOUIS ALPHA	Woodford E NAME: JANA
237 239 STATE LOUIS ALPHA NUME	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22
237 239 STATE LOUIS ALPHA NUME CODE	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME
237 239 STATE LOUIS ALPHA NUME	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22
237 239 STATE LOUIS ALPHA NUME CODE	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME
237 239 STATE LOUIS ALPHA NUME CODE 001	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia
237 239 STATE LOUIS ALPHA NUME CODE 001 003 005	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension
237 239 STATE LOUIS ALPHA NUME CODE 001 003 005 007	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption
237 239 STATE LOUIS ALPH/ NUME CODE 001 003 005 007 009	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011	Woodford E NAME: HANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011	Woodford E NAME: HANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017	Woodford E NAME: HANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell
237 239 STATE LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023	Woodford <b>C NAME:</b> <b>JANA</b> <b>ABETIC CODE: LA</b> <b>RIC CODE: 22</b> <b>COUNTY NAME</b> Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025	Woodford E NAME: HANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025	Woodford E NAME: HANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027	Woodford E NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031	Woodford <b>C NAME:</b> <b>JANA</b> <b>ABETIC CODE: LA</b> <b>RIC CODE: 22</b> <b>COUNTY NAME</b> Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne Concordia DeSoto
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	Woodford <b>C NAME:</b> <b>JANA</b> <b>ABETIC CODE: LA</b> <b>RIC CODE: 22</b> <b>COUNTY NAME</b> Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne Concordia DeSoto East Baton Rouge
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035	Woodford <b>C NAME:</b> <b>JANA</b> <b>ABETIC CODE: LA</b> <b>RIC CODE: 22</b> <b>COUNTY NAME</b> Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne Concordia DeSoto East Baton Rouge East Carroll
237 239 <b>STATE</b> LOUIS ALPH/ NUME CODE 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	Woodford C NAME: JANA ABETIC CODE: LA RIC CODE: 22 COUNTY NAME Acadia Allen Ascension Assumption Avoylelles Beauregard Bienville Bossier Caddo Calcasieu Caldwell Cameron Catahoula Claiborne Concordia DeSoto East Baton Rouge

039

041

Evangeline

Franklin

043 Grant 045 Iberia 047 Iberville 049 Jackson 051 Jefferson 053 Jefferson Davis 055 Lafayette 057 Lafourche 059 La Salle 061 Lincoln 063 Livingston 065 Madison 067 Morehouse 069 Natchitoches 071 Orleans 073 Ouachita 075 Plaquemines 077 Pointe Coupee 079 Rapides 081 Red River 083 Richland 085 Sabine 087 St. Bernard 089 St. Charles 091 St. Helena 093 St. James 095 St. John the Baptist 097 St. Landry 099 St. Martin 101 St. Mary 103 St. Tammany 105 Tangipahoa 107 Tensas 109 Terrebonne 111 Union 113 Vermilion 115 Vernon 117 Washington 119 Webster 121 West Baton Rouge 123 West Carroll 125 West Feliciana 127 Winn STATE NAME: MAINE **ALPHABETIC CODE:** ME NUMERIC CODE: 23 CODE COUNTY NAME 001 Androscoggin 003 Aroostook 005 Cumberland 007 Franklin 009 Hancock 011 Kennebec 013 Knox 015 Lincoln 017 Oxford 019 Penobscot 021 Piscataquis

027 Waldo Washington 029 031 York STATE NAME: MARYLAND **ALPHABETIC CODE:** MD NUMERIC CODE: 24 CODE COUNTY NAME 001 Allegany 003 Anne Arundel 005 Baltimore 009 Calvert 011 Caroline 013 Carroll 015 Cecil 017 Charles 019 Dorchester 021 Frederick 023 Garrettt 025 Harford 027 Howard 029 Kent Montgomery 031 033 Prince George's 035 Queen Anne's 037 St. Mary's 039 Somerset 041 Talbot 043 Washington 045 Wicomico 047 Worcester CODE INDEPENDENT CITY 510 Baltimore (City) STATE NAME: MASSACHUSETTS ALPHABETIC CODE: MA **NUMERIC CODE: 25** CODE COUNTY NAME 001 Barnstable 003 Berkshire 005 Bristol 007 Dukes 009 Essex 011 Franklin 013 Hampden 015 Hampshire 017 Middlesex 019 Nantucket 021 Norfolk 023 Plymouth

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

Sagadahoc

Somerset

023

025

		11/
MICH		119
	ABETIC CODE: MI	121
NUMI	ERIC CODE: 26	123
		125
CODE	COUNTY NAME	127
001	Alcona	129
003	Alger	131
005	Allegan	133
007	Alpena	135
009	Antrim	137
011	Arenac	139
013	Baraga	141
015	Barry	143
017	Bay	145
019	Benzie	147
021	Berrien	149
023	Branch	151
025	Calhoun	153
027 029	Cass Charlevoix	155 157
029	Cheboygan	157
033	Chippewa	159
035	Clare	163
035	Clinton	165
039	Crawford	105
035	Delta	
043	Dickinson	STA
045	Eaton	MIN
047	Emmet	ALI
049	Genesee	MN
051	Gladwin	NUI
053	Gogebic	
055	Grand Traverse	CO
057	Gratiot	001
059	Hillsdale	003
061	Houghton	005
063	Huron	007
065	Ingham	009
067	Ionia	011
069	Iosco	013
071	Iron	015
073	Isabella	017
075	Jackson	019
077	Kalamazoo	021
079	Kalkaska	023
081	Kent	025
083	Keweenaw	027
085	Lake	029
087	Lapeer Leelanau	031 033
089 091	Lenawee	035
091	Livingston	033
095	Luce	039
097	Mackinac	041
099	Macomb	041
101	Manistee	045
101	Marquette	043
105	Mason	049
107	Mecosta	051
109	Menominee	053
111	Midland	055
113	Missaukee	057
115	Monroe	059

STATE NAME:

17	Montcalm
19	Montmorency
21	Muskegon
23	Newaygo
25	Oakland
27	Oceana
29	Ogemaw
31	Ontonagon
33	Osceola
35	Oscoda
37	Otsego
39	Ottawa
41	Presque Isle
43	Roscommon
45	Saginaw
47	St. Clair
49	St. Joseph
51	Sanilac
53	Schoolcraft
55	Shiawassee
57	Tuscola
59	Van Buren
61	Washtenaw
63	Wayne
65	Wexford
татр	NAME.

1

#### STATE NAME: MINNESOTA **ALPHABETIC CODE:** MN NUMERIC CODE: 27 CODE COUNTY NAME 001 Aitkin 003 Anoka 005 Becker 007 Beltrami 009 Benton 011 Big Stone 013 Blue Earth 015 Brown 017 Carlton 019 Carver 021 Cass 023 Chippewa 025 Chisago 027 Clay 029 Clearwater 031 Cook 033 Cottonwood 035 Crow Wing 037 Dakota 039 Dodge Douglas 041 043 Faribault 045 Fillmore 047 Freeborn 049 Goodhue 051 Grant 053 Hennepin

061 Itasca 063 Jackson Kanabec 065 067 Kandiyohi 069 Kittson 071 Koochiching 073 Lac qui Parle 075 Lake 077 Lake of the Woods 079 Le Sueur 081 Lincoln 083 Lyon 085 McLeod 087 Mahnomen 089 Marshall 091 Martin 093 Meeker 095 Mille Lacs 097 Morrison 099 Mower 101 Murray 103 Nicollet 105 Nobles 107 Norman 109 Olmsted 111 Otter Tail 113 Pennington 115 Pine 117 Pipestone 119 Polk 121 Pope 123 Ramsey 125 Red Lake 127 Redwood 129 Renville 131 Rice 133 Rock 135 Roseau 137 St. Louis 139 Scott 141 Sherburne 143 Sibley 145 Stearns 147 Steele 149 Stevens 151 Swift 153 Todd 155 Traverse 157 Wabasha 159 Wadena 161 Waseca 163 Washington 165 Watonwan 167 Wilkin 169 Winona 171 Wright 173 Yellow Medicine

STATE NAME: MISSISSIPPI **ALPHABETIC CODE:** MS NUMERIC CODE: 28 CODE COUNTY NAME 001 Adams 003 Alcorn 005 Amite 007 Attala 009 Benton 011 Bolivar 013 Calhoun 015 Carroll 017 Chickasaw 019 Choctaw 021 Claiborne 023 Clarke 025 Clay 027 Coahoma 029 Copiah 031 Covington 033 DeSoto 035 Forrest 037 Franklin 039 George 041 Greene 043 Grenada 045 Hancock 047 Harrison 049 Hinds 051 Holmes 053 Humphreys 055 Issaquena Itawamba 057 059 Jackson 061 Jasper 063 Jefferson 065 Jefferson Davis 067 Jones 069 Kemper 071 Lafayette 073 Lamar 075 Lauderdale 077 Lawrence 079 Leake 081 Lee 083 Leflore 085 Lincoln 087 Lowndes 089 Madison 091 Marion 093 Marshall 095 Monroe 097 Montgomery 099 Neshoba 101 Newton 103 Noxubee 105 Oktibbeha 107 Panola 109 Pearl River 111 Perry 113 Pike

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

Houston

Hubbard

Isanti

115	Pontotoc
117	Prentiss
119	Quitman
121	Rankin
123	Scott
125	Sharkey
127	Simpson
129	Smith
131	Stone
133	Sunflower
135	Tallahatchie
137	Tate
139	Tippah
141	Tishomingo
143	Tunica
145	Union
147	Walthall
149	Warren
151	Washington
153	Wayne
155	Webster
157	Wilkinson
159	Winston
161	Yalobusha
163	Yazoo
MO	ABETIC CODE: RIC CODE: 29
NUME	AIC CODE. 29
CODE	
	COUNTY NAME
<b>CODE</b> 001	COUNTY NAME Adair
001	Adair
001 003 005	Adair Andrew Atchison
001 003 005 007	Adair Andrew Atchison Audrain
001 003 005 007 009	Adair Andrew Atchison Audrain Barry
001 003 005 007 009 011	Adair Andrew Atchison Audrain Barry Barton
001 003 005 007 009 011 013	Adair Andrew Atchison Audrain Barry Barton Bates
001 003 005 007 009 011	Adair Andrew Atchison Audrain Barry Barton Bates Benton
001 003 005 007 009 011 013	Adair Andrew Atchison Audrain Barry Barton Bates
001 003 005 007 009 011 013 015	Adair Andrew Atchison Audrain Barry Barton Bates Benton
001 003 005 007 009 011 013 015 017	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger
001 003 005 007 009 011 013 015 017 019 021	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan
001 003 005 007 009 011 013 015 017 019 021 023	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler
001 003 005 007 009 011 013 015 017 019 021 023 025	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell
001 003 005 007 009 011 013 015 017 019 021 023 025 027	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark Clay
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark Clay Clinton
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark Clay Clinton Cole
001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051 053	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark Clay Clinton Cole Cooper
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001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051 053 055	Adair Andrew Atchison Audrain Barry Barton Bates Benton Bollinger Boone Buchanan Butler Caldwell Callaway Camden Cape Girardeau Carroll Carter Cass Cedar Chariton Christian Clark Clay Clinton Cole Cooper Crawford

Daviess DeKalb Dent Douglas Dunklin Franklin Gasconade Gentry Greene Grundy Harrison Henry Hickory Holt Howard Howell Iron Jackson Jasper Jefferson Johnson Knox Laclede Lafayette Lawrence Lewis Lincoln Linn Livingston McDonald Macon Madison Maries Marion Mercer Miller Mississippi Moniteau Monroe Montgomery Morgan New Madrid Newton Nodaway Oregon Osage Ozark Pemiscot Perry Pettis Phelps Pike Platte Polk Pulaski Putnam Ralls Randolph Ray Reynolds Ripley St. Charles St. Clair Ste. Genevieve

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187	St. Francois	0
189	St. Louis County	0
195	Saline	0
197	Schuyler	0
199	Scotland	0
201	Scott	0
203	Shannon	0
205	Shebly	0
207	Stoddard	0
209	Stone	0
211	Sullivan	0
213 215	Taney	0
215	Texas Vernon	0 0
217	Warren	0
21)	Washington	0
223	Wayne	0
225	Webster	0
227	Worth	0
229	Wright	1
		1
CODE	INDEPENDENT	1
CITY		1
510	St. Louis City	1
		1
STATI	E NAME:	Ν
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	ABETIC CODE:	Y
МТ		Ν
	CRIC CODE: 30	G
		C
	COUNTY NAME	Y
001	Beaverhead	л
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003	Big Horn	e
003 005	Big Horn Blaine	
003 005 007	Big Horn Blaine Broadwater	e
003 005 007 009	Big Horn Blaine Broadwater Carbon	e S
003 005 007 009 011	Big Horn Blaine Broadwater Carbon Carter	e S N
003 005 007 009 011 013	Big Horn Blaine Broadwater Carbon Carter Cascade	e S N A
003 005 007 009 011 013 015	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau	e S N
003 005 007 009 011 013	Big Horn Blaine Broadwater Carbon Carter Cascade	e S N A N
003 005 007 009 011 013 015 017	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer	e S N A
003 005 007 009 011 013 015 017 019	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels	ed S N A N
003 005 007 009 011 013 015 017 019 021	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson	e S N A N C
003 005 007 009 011 013 015 017 019 021 023 025 027	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus	e S N A N C 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead	e S N A N C 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill	C C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite	e S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
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003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin	C S N A N C O 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin Lake	C S N A N C O 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin Lake Lewis and Clark	C C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin Lake Lewis and Clark Liberty	C S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051 053	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin Lake Lewis and Clark Liberty Lincoln	C S N A N C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045 047 049 051 053 055	Big Horn Blaine Broadwater Carbon Carter Cascade Chouteau Custer Daniels Dawson Deer Lodge Fallon Fergus Flathead Gallatin Garfield Glacier Golden Valley Granite Hill Jefferson Judith Basin Lake Lewis and Clark Liberty Lincoln	E A S S N A A N C O 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0

)63 Missoula Musselshell )65 )67 Park )69 Petroleum )71 Phillips )73 Pondera )75 Powder River )77 Powell )79 Prairie )81 Ravalli )83 Richland )85 Roosevelt )87 Rosebud )89 Sanders )91 Sheridan )93 Silver Bow )95 Stillwater )97 Sweet Grass )99 Teton 01 Tooke 03 Treasure 05 Valley 07 Wheatland 09 Wibaux 11 Yellowstone

NIST has been notified by he Bureau of Census that Yellowstone National Park, MT, is legally part of Gallatin County and Park County. This eliminates Yellowstone National Park FIPS Code 113) as a county equivalent.

### STATE NAME: NEBRASKA ALPHABETIC CODE: NE NUMERIC CODE: 31

CODE COUNTY NAME 001 Adams 003 Antelope )05 Arthur 007 Banner )09 Blaine )11 Boone )13 Box Butte )15 Boyd )17 Brown )19 Buffalo )21 Burt Butler )23 )25 Cass )27 Cedar )29 Chase )31 Cherry Cheyenne )33 )35 Clay )37 Colfax 039 Cuming Custer 041

Mineral

061

043	Dakota
045	Dawes
047	Dawson
049	Deuel
051	Dixon
053	Dodge
055	Douglas
057	Dundy
059	Fillmore
061	Franklin
063	Frontier
065	Furnas
067	Gage
069	Garden
	Garfield
071	
073	Gosper
075	Grant
077	Greeley
079	Hall
081	Hamilton
083	Harlan
085	Hayes
087	Hitchcock
089	Holt
091	Hooker
093	Howard
095	Jefferson
097	Johnson
099	Kearney
101	Keith
103	Keya Paha
105	Kimball
107	Knox
109	Lancaster
111	Lincoln
113	Logan
115	Loup
117	McPherson
119	Madison
121	Merrick
123	Morrill
125	Nance
	Nemaha
127	
129	Nuckolls
131	Otoe
133	
	Pawnee
135	Perkins
137	Phelps
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139	Pierce
141	Platte
143	Polk
145	Red Willow
147	Richardson
149	Rock
151	Saline
153	Sarpy
155	Saunders
157	Scotts Bluff
159	Seward
161	Sheridan
163	Sherman
165	Sioux
167	Stanton
169	Thayer

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171	Thomas
173	Thurston
175	Valley
177	Washington
179	Wayne
181	Webster
183	Wheeler
185	York
STATE	NAME: NEVADA
ALPHA	ABETIC CODE: NV
	RIC CODE: 32
CODE	COUNTY NAME
001	Churchill
003	Clark
005	Doulgas
007	Elko
009	Esmeralda
009	Eureka
013	Humboldt
015	Lander
017	Lincoln
	Lyon
021	Mineral
023	Nye
027	Pershing
029	Storey
031	Washoe
033	White Pine
	INDEPENDENT
CITY	
510	Carson City
Carson	City does not include
	lesignation (such as
"city").	
	NAME:
NEW H	IAMPSHIRE
ALPHA	ABETIC CODE:
NH	
NUME	RIC CODE: 33
CODE	· ·
001	Belknap
003	Carroll
005	Cheshire
007	Coos
009	Grafton
011	Hillsborough
013	Merrimack
015	Rockingham
017	Strafford
019	Sullivan

STATE	E NAME:	053	Socorro
NEW J	ERSEY	055	Taos
ALPH	ABETIC CODE: NJ	057	Torrance
NUME	RIC CODE: 34	059	Union
		061	Valencia
CODE	COUNTY NAME		
001	Atlantic	Cibola	was established
003	Bergen	part of	Valencia (6/19/
005	Burlington		
007	Camden		
009	Cape May	STATI	E NAME:
011	Cumberland	NEW Y	YORK
013	Essex	ALPH.	ABETIC COD
015	Gloucester	NUME	RIC CODE: 3
017	Hudson		
019	Hunterdon	CODE	COUNTY NA
021	Mercer	001	Albany
023	Middlesex	003	Allegany
025	Monmouth	005	Bronx
027	Morris	007	Broome
029	Ocean	009	Cattaraugus
031	Passaic	011	Cayuga
033	Salem	013	Chautauqua
035	Somerset	015	Chemung
037	Sussex	017	Chenango
039	Union	019	Clinton
041	Warren	021	Columbia
		023	Cortland
		025	Delaware
	E NAME:	027	Dutchess
	MEXICO	029	Erie
	ABETIC CODE:	031	Essex
NM	DIC CODE AF	033	Franklin
NUME	RIC CODE: 35	035	Fulton
CODE		037	Genesee
	COUNTY NAME	039	Greene
001 003	Bernalillo Catron	041 043	Hamilton Herkimer
005	Chaves	045	Jefferson
005	Cibola	043	Kings
000	Colfax	047	Lewis
007	Curry	049	Livingston
011	DeBaca	053	Madison
013	Dona Ana	055	Monroe
015	Eddy	057	Montgomery
017	Grant	059	Nassau
019	Guadalupe	061	New York
021	Harding	063	Niagara
023	Hidalgo	065	Oneida
025	Lea	067	Onondaga
027	Lincoln	069	Ontario
028	Los Alamos	071	Orange
029	Luna	073	Orleans
031	McKinley	075	Oswego
033	Mora	077	Otsego
035	Otero	079	Putnam
037	Quay	081	Queens
039	Rio Arriba	083	Rensselaer
041	Roosevelt	085	Richmond
043	Sandoval	087	Rockland
045	San Juan	089	St. Lawrence
047	San Migual	001	Sarataga

corro )S rrance ion lencia established from encia (6/19/81). ME: RK TIC CODE: NY CODE: 36 DUNTY NAME bany legany onx oome ttaraugus yuga autauqua emung enango nton lumbia rtland laware tchess e sex nklin lton enesee eene milton rkimer ferson ıgs wis ingston ndison onroe ontgomery ssau w York igara eida ondaga tario

047

049

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San Miguel

Santa Fe

Sierra

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Saratoga

Schenectady

Schoharie

097 099	Schuyler Seneca	085 087	Harnett Haywood	DAKO		ALPH	E NAME: OHIO ABETIC CODE:
101	Steuben	089	Henderson	ALPHABETIC CODE: ND		OH	
103	Suffolk	091	Hertford	NUME	CRIC CODE: 38	NUME	RIC CODE: 39
105	Sullivan	093	Hoke				
107	Tioga	095	Hyde		COUNTY NAME		COUNTY NAME
109	Tompkins	097	Iredell	001	Adams	001	Adams
111	Ulster	099	Jackson	003	Barnes	003	Allen
113	Warren	101	Johnston	005	Benson	005	Ashland
115	Washington	103	Jones	007	Billings	007	Ashtabula
117	Wayne	105	Lee	009	Bottineau	009	Athens
119	Westchester	107	Lenoir	011	Bowman	011	Auglaize
121	Wyoming	109	Lincoln	013	Burke	013	Belmont
123	Yates	111	McDowell	015	Burleigh	015	Brown
		113	Macon	017	Cass	017	Butler
	ENAME NODTH	115	Madison	019	Cavalier	019	Carroll
	E NAME: NORTH	117	Martin	021	Dickey Divide	021	Champaign Clark
CARO		119	Mecklenburg	023	Divide Dunn	023 025	Clark
	ABETIC CODE: NC ERIC CODE: 37	121 123	Mitchell	025 027	Eddy	023	Clinton
NUM	ERIC CODE: 37	125	Montgomery Moore	027	Emmons	027	Columbiana
CODE	COUNTY NAME	123	Nash	029	Foster	029	Coshocton
001	Alamance	127	New Hanover	031	Golden Valley	031	Crawford
001	Alexander	129	Northampton	035	Grand Forks	035	Cuyahoga
005	Alleghany	131	Onslow	035	Grant	035	Darke
005	Anson	135	Orange	039	Griggs	039	Defiance
009	Ashe	133	Pamlico	041	Hettinger	041	Delaware
011	Avery	139	Pasquotank	043	Kidder	043	Erie
013	Beaufort	141	Pender	045	LaMoure	045	Fairfield
015	Bertie	143	Perquimans	047	Logan	047	Fayette
017	Bladen	145	Person	049	McHenry	049	Franklin
019	Brunswick	147	Pitt	051	McIntosh	051	Fulton
021	Buncombe	149	Polk	053	McKenzie	053	Gallia
023	Burke	151	Randolph	055	McLean	055	Geauga
025	Cabarrus	153	Richmond	057	Mercer	057	Greene
027	Caldwell	155	Robeson	059	Morton	059	Guernsey
029	Camden	157	Rockingham	061	Mountrail	061	Hamilton
031	Carteret	159	Rowan	063	Nelson	063	Hancock
033	Caswell	161	Rutherford	065	Oliver	065	Hardin
035	Catawba	163	Sampson	067	Pembina	067	Harrison
037	Chatham	165	Scotland	069	Pierce	069	Henry
039	Cherokee	167	Stanly	071	Ramsey	071	Highland
041	Chowan	169	Stokes	073	Ransom	073	Hocking
043	Clay	171	Surry	075	Renville	075	Holmes
045	Cleveland	173	Swain	077	Richland	077	Huron
047	Columbus	175 177	Transylvania	079	Rolette	079	Jackson Jefferson
049	Craven		Tyrrell	081	Sargent	081	
051 053	Cumberland Currituck	179 181	Union Vance	083 085	Sheridan Sioux	083 085	Knox Lake
055	Dare	181	Wake	083	Slope	083	Lawrence
055	Davidson	185	Warren	087	Stark	087	Licking
059	Davie	185	Washington	0091	Steele	082	Logan
061	Duplin	189	Watauga	093	Stutsman	093	Lorain
063	Dupham	191	Wayne	095	Towner	095	Lucas
065	Edgecombe	193	Wilkes	095	Traill	095	Madison
065	Forsyth	195	Wilson	099	Walsh	099	Mahoning
069	Franklin	197	Yadkin	101	Ward	101	Marion
071	Gaston	199	Yancey	101	Wells	101	Medina
073	Gates		2	105	Williams	105	Meigs
075	Graham					107	Mercer
077	Granville					109	Miami
079	Greene					111	Monroe
081	Guilford					113	Montgomery
083	Halifax					115	Morgan

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

117	Morrow	051
119	Muskingum	053
121	Noble	055
123	Ottawa	057
125	Paulding	059
127	Perry	061
129	Pickaway	063
131	Pike	065
133	Portage	067
135	Preble	069
137	Putnam	071
139	Richland	073
141	Ross	075
143	Sandusky	077
145	Scioto	079
147	Seneca	081
149	Shelby	083
151	Stark	085
153	Summit	087
155 157	Trumbull Tuscarawas	089 091
157	Union	091
161	VanWert	095
163	Vinton	093
165	Warren	099
167	Washington	101
169	Wayne	103
171	Williams	105
173	Wood	107
175	Wyandot	109
		111
		113
STATI	E NAME:	115
OKLA	НОМА	115 117
OKLA ALPH		117 119
OKLA ALPH OK	HOMA ABETIC CODE:	117 119 121
OKLA ALPH OK	НОМА	117 119 121 123
OKLA ALPH OK NUME	HOMA ABETIC CODE: CRIC CODE: 40	117 119 121 123 125
OKLA ALPH OK NUME	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME	117 119 121 123 125 127
OKLA ALPH OK NUME CODE 001	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair	117 119 121 123 125 127 129
OKLA ALPH OK NUME CODE 001 003	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa	117 119 121 123 125 127 129 131
OKLA ALPH OK NUME CODE 001 003 005	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa Atoka	117 119 121 123 125 127 129 131 133
OKLA ALPH OK NUME CODE 001 003 005 007	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa Atoka Beaver	117 119 121 123 125 127 129 131 133 135
OKLA ALPH. OK NUME 001 003 005 007 009	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa Atoka Beaver Beckham	117 119 121 123 125 127 129 131 133 135 137
OKLA ALPH OK NUME CODE 001 003 005 007 009 011	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine	117 119 121 123 125 127 129 131 133 135 137 139
OKLA ALPH OK NUME 001 003 005 007 009 011 013	HOMA ABETIC CODE: CRIC CODE: 40 COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan	117 119 121 123 125 127 129 131 133 135 137 139 141
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo	117 119 121 123 125 127 129 131 133 135 137 139 141 143
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian	117 119 121 123 125 127 129 131 133 135 137 139 141 143 145
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter	117 119 121 123 125 127 129 131 133 135 137 139 141 143
OKLA ALPH: OK NUME 001 003 005 007 009 011 013 015 017 019	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian	117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee	117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw	117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron	117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche	1117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151 153 STA
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton	1117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151 153 STA ALL
OKLA ALPH: OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig	<ul> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> <li>129</li> <li>131</li> <li>133</li> <li>135</li> <li>137</li> <li>139</li> <li>141</li> <li>143</li> <li>145</li> <li>147</li> <li>149</li> <li>151</li> <li>153</li> <li>STA</li> <li>ALLI</li> <li>OR</li> </ul>
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig Creek	1117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151 153 STA ALL
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig Creek Custer	<ul> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> <li>129</li> <li>131</li> <li>133</li> <li>135</li> <li>137</li> <li>139</li> <li>141</li> <li>143</li> <li>145</li> <li>147</li> <li>149</li> <li>151</li> <li>153</li> <li>STA</li> <li>ALH</li> <li>OR</li> <li>NUM</li> </ul>
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig Creek Custer Delaware	1117 119 121 123 125 127 129 131 133 135 137 139 141 143 145 147 149 151 153 STA ALLI OR NUN
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig Creek Custer Delaware Dewey	<ul> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> <li>129</li> <li>131</li> <li>133</li> <li>135</li> <li>137</li> <li>139</li> <li>141</li> <li>143</li> <li>145</li> <li>147</li> <li>149</li> <li>151</li> <li>153</li> <li>STA</li> <li>ALH</li> <li>OR</li> <li>NUN</li> <li>COI</li> <li>001</li> </ul>
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043 045	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Coal Comanche Cotton Craig Creek Custer Delaware Dewey Ellis	<ul> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> <li>129</li> <li>131</li> <li>133</li> <li>135</li> <li>137</li> <li>139</li> <li>141</li> <li>143</li> <li>145</li> <li>147</li> <li>149</li> <li>151</li> <li>153</li> <li>STA</li> <li>ALH</li> <li>OR</li> <li>NUN</li> <li>COI</li> <li>001</li> <li>003</li> </ul>
OKLA ALPH. OK NUME 001 003 005 007 009 011 013 015 017 019 021 023 025 027 029 031 033 035 037 039 041 043	HOMA ABETIC CODE: COUNTY NAME Adair Alfalfa Atoka Beaver Beckham Blaine Bryan Caddo Canadian Carter Cherokee Choctaw Cimarron Cleveland Coal Comanche Cotton Craig Creek Custer Delaware Dewey	<ul> <li>117</li> <li>119</li> <li>121</li> <li>123</li> <li>125</li> <li>127</li> <li>129</li> <li>131</li> <li>133</li> <li>135</li> <li>137</li> <li>139</li> <li>141</li> <li>143</li> <li>145</li> <li>147</li> <li>149</li> <li>151</li> <li>153</li> <li>STA</li> <li>ALH</li> <li>OR</li> <li>NUN</li> <li>COI</li> <li>001</li> </ul>

1	Grady
3	
5	Grant
5 7	Greer
	Harmon
9	Harper
1	Haskell
3	Hughes
5	Jackson
7	Jefferson
9	Johnston
1	Kay
3	Kingfisher
5	Kiowa
7	
/	Latimer
9	Le Flore
1	Lincoln
3	Logan
3 5 7	Love
7	McClain
9	McCurtain
1	McIntosh
3	Major
5	Marshall
7	Mayes
9	
	Murray
1	Muskogee
3	Noble
5	Nowata
7	Okfushee
9	Oklahoma
1	
	Okmulgee
3 5 7	Osage
5	Ottawa
7	Pawnee
9	Payne
1	Pittsburg
3	
	Pontotoc
5 7	Pottawatomie
	Pushmataha
9	Roger Mills
1	Rogers
	Seminole
3 5 7	
5	Sequoyah
	Stephens
9	Texas
1	Tillman
3	Tulsa
5	Wagoneer
7	•
	Washington
9	Washita
1	Woods
3	Woodward
LPHA R	NAME: OREGON ABETIC CODE: RIC CODE: 41
ODE	COUNTY NAME
1	
	Balzer
	Baker
3	Benton
3 5	Benton Clackamas
3 5 7	Benton

009	Columbia
011	Coos
013	Crook
015	Curry
017	Deschutes
019	Douglas
021	Gilliam
023	Grant
025	Harney
027	Hood River
029	Jackson
031	Jefferson
033	Josephine
035	Klamath
037	Lake
039	Lane
041	Lincoln
043	Linn
045	Malheur
047	Marion
049	Morrow
051	Multnomah
053	Polk
055	Sherman
057	Tillamook
059	Umatilla
061	Union
063	Wallowa
065	Wasco
067	Washington
069	Wheeler
071	Yamhill

### STATE NAME: PENNSYLVANIA ALPHABETIC CODE: PA NUMERIC CODE: 42

### CODE COUNTY NAME 001 Adams

001	1 I dddiiio
003	Allegheny
005	Armstrong
007	Beaver
009	Bedford
011	Berks
013	Blair
015	Bradford
017	Bucks
019	Butler
021	Cambria
023	Cameron
025	Carbon
027	Centre
029	Chester
031	Clarion
033	Clearfield
035	Clinton
037	Columbia
039	Crawford
041	Cumberland
043	Dauphin
045	Delaware
047	Elk

049	Erie
051	Fayette
053	Forest
055	Franklin
057	Fulton
059	Greene
061	Huntingdon
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
093	Montour
095	Northampton
097	Northumberland
099	Perry
101	Philadelphia
103	Pike
105	Potter
107	Schuylkill
109	Snyder
111	Somerset
113	Sullivan
115	Susquehanna
117	Tioga
119	Union
121	Venango
123	Warren
125	Washington
127	Wayne
129	Westmoreland
131	Wyoming
122	V1-

133 York

### STATE NAME: RHODE ISLAND ALPHABETIC CODE: RI NUMERIC CODE: 44

### CODE COUNTY NAME

- 001 Bristol
- 003 Kent
- 005 Newport
- 007 Providence
- 009 Washington

STAT	E NAME: SOUTH	011	Brookings	STAT	E NAME:	117	Marshall
CAROLINA		013	Brown	TENN	ESSEE	119	Maury
ALPH	LPHABETIC CODE: SC         015         Brule         ALPHABETIC CODE: TN		121	Meigs			
NUM	ERIC CODE: 45	017	Buffalo	NUM	ERIC CODE: 47	123	Monroe
		019	Butte			125	Montgomery
CODI	E COUNTY NAME	021	Campbell		E COUNTY NAME	127	Moore
001	Abbeviille	023	Charles Mix	001	Anderson	129	Morgan
003	Aiken	025	Clark	003	Bedford	131	Obion
005	Allendale	027	Clay	005	Benton	133	Overton
007	Anderson	029	Codington	007	Bledsoe	135	Perry
009	Bamberg	031	Corson	009	Blount	137	Pickett
011	Barnwell	033	Custer	011	Bradley	139	Polk
013	Beaufort	035	Davison	013	Campbell	141	Putnam
015	Berkeley	037	Day	015	Cannon	143	Rhea
017	Calhoun	039	Deuel	017	Carroll	145	Roane
019	Charleston	041	Dewey	019 021	Carter Cheatham	147	Robertson
021	Cherokee	043 045	Douglas	021		149	Rutherford
023 025	Chester	045 047	Edmunds		Chester	151	Scott
023	Chesterfield Clarendon	047	Fall River Faulk	025 027	Claiborne Clay	153 155	Sequatchie Sevier
027	Colleton	049	Grant	029	Cocke	155	Shelby
029	Darlington	051	Gregory	029	Coffee	157	Smith
033	Dillon	055	Haakon	033	Crockett	159	Stewart
035	Dorchester	055	Hamlin	035	Cumberland	163	Sullivan
037	Edgefield	059	Hand	035	Davidson	165	Sumner
039	Fairfield	061	Hanson	039	Decatur	167	Tipton
041	Florence	063	Harding	041	DeKalb	169	Trousdale
043	Georgetown	065	Hughes	043	Dickson	171	Unicoi
045	Greenville	067	Hutchinson	045	Dyer	173	Union
047	Greenwood	069	Hyde	047	Fayette	175	Van Buren
049	Hampton	071	Jackson	049	Fentress	177	Warren
051	Horry	073	Jerauld	051	Franklin	179	Washington
053	Jasper	075	Jones	053	Gibson	181	Wayne
055	Kershaw	077	Kingsbury	055	Giles	183	Weakley
057	Lancaster	079	Lake	057	Grainger	185	White
059	Laurens	081	Lawrence	059	Greene	187	Williamson
061	Lee	083	Lincoln	061	Grundy	189	Wilson
063	Lexington	085	Lyman	063	Hamblen		
065	McCormick	087	McCook	065	Hamilton		
067	Marion	089	McPherson	067	Hancock	STAT	E NAME: TEXAS
069	Marlboro	091	Marshall	069	Hardeman	ALPH	ABETIC CODE: TX
071	Newberry	093	Meade	071	Hardin	NUMI	ERIC CODE: 48
073	Oconee	095	Mellette	073	Hawkins		
075	Orangeburg	097	Miner	075	Haywood		E COUNTY NAME
077	Pickens	099	Minnehaha	077	Henderson	001	Anderson
079	Richland	101	Moody	079	Henry	003	Andrews
081	Saluda	103	Pennington	081	Hickman	005	Angelina
083	Spartanburg	105	Perkins	083	Houston	007	Aransas
085 087	Sumter Union	107 109	Potter Roberts	085 087	Humphreys Jackson	009 011	Archer Armstrong
087	Williamsburg	109	Sanborn	087	Jefferson	011	Atascosa
089	York	111	Shannon	089	Johnson	013	Austin
091	TOIK	115	Spink	093	Knox	013	Bailey
		115	Stanley	095	Lake	019	Bandera
STAT	'E NAME: SOUTH	117	Sully	095	Lauderdale	01)	Bastrop
DAK		119	Todd	099	Lawrence	021	Baylor
	IABETIC CODE: SD	121	Tripp	101	Lewis	025	Bee
	ERIC CODE: 46	125	Turner	101	Lincoln	023	Bell
		127	Union	105	Loudon	029	Bexar
CODE	E COUNTY NAME	129	Walworth	107	McMinn	031	Blanco
003	Aurora	135	Yankton	109	McNairy	033	Borden
005	Beadle	137	Ziebach	111	Macon	035	Bosque
007	Bennett			113	Madison	037	Bowie
009	Bon Homme			115	Marion	039	Brazoria

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

041	Brazos	169	Garza	297	Live Oak	425	Somervell
043	Brewster	171	Gillespie	299	Llano	427	Starr
045	Briscoe	173	Glasscock	301	Loving	429	Stephens
047	Brooks	175	Goliad	303	Lubbock	431	Sterling
049	Brown	177	Gonzales	305	Lynn	433	Stonewall
051	Burleson	179	Gray	307	McCulloch	435	Sutton
053	Burnet	181	Grayson	309	McLennan	437	Swisher
055	Caldwell	181	5	311	McMullen	439	Tarrant
			Gregg				
057	Calhoun	185	Grimes	313	Madison	441	Taylor
059	Callahan	187	Guadalupe	315	Marion	443	Terrell
061	Cameron	189	Hale	317	Martin	445	Terry
063	Camp	191	Hall	319	Mason	447	Throckmorton
065	Carson	193	Hamilton	321	Matagorda	449	Titus
067	Cass	195	Hansford	323	Maverick	451.	Tom Green
069	Castro	197	Hardeman	325	Medina	453	Travis
071	Chambers	199	Hardin	327	Menard	455	Trinity
073	Cherokee	201	Harris	329	Midland	457	Tyler
075	Childress	203	Harrison	331	Milam	459	Upshur
075	Clay	205	Hartley	333	Mills	461	Upton
079	Cochran	203	-	335	Mitchell	463	Uvalde
			Haskell				
081	Coke	209	Hays	337	Montague	465	Val Verde
083	Coleman	211	Hemphill	339	Montgomery	467	Van Zandt
085	Collin	213	Henderson	341	Moore	469	Victoria
087	Collingsworth	215	Hidalgo	343	Morris	471	Walker
089	Colorado	217	Hill	345	Motley	473	Waller
091	Comal	219	Hockley	347	Nacogdoches	475	Ward
093	Comanche	221	Hood	349	Navarro	477	Washington
095	Concho	223	Hopkins	351	Newton	479	Webb
097	Cooke	225	Houston	353	Nolan	481	Wharton
099	Corvell	227	Howard	355	Nueces	483	Wheeler
	5	229			Ochiltree		
101	Cottle		Hudspeth	357		485	Wichita
103	Crane	231	Hunt	359	Oldham	487	Wilbarger
105	Crockett	233	Hutchinson	361	Orange	489	Willacy
107	Crosby	235	Irion	363	Palo Pinto	491	Williamson
109	Culberson	237	Jack	365	Panola	493	Wilson
111	Dallam	239	Jackson	367	Parker	495	Winkler
113	Dallas	241	Jasper	369	Parmer	497	Wise
115	Dawson	243	Jeff Davis	371	Pecos	499	Wood
117	Deaf Smith	245	Jefferson	373	Polk	501	Yoakum
119	Delta	247	Jim Hogg	375	Potter	503	Young
121	Denton	249	Jim Wells	377	Presidio	505	Zapata
121	DeWitt	251	Johnson	379	Rains	505	Zavala
125					Randall	507	Zavala
	Dickens	253	Jones	381			
127	Dimmit	255	Karnes	383	Reagan		
129	Donley	257	Kaufman	385	Real		E NAME: UTAH
131	Duval	259	Kendall	387	Red River		ABETIC CODE: UT
133	Eastland	261	Kenedy	389	Reeves	NUME	CRIC CODE: 49
135	Ector	263	Kent	391	Refugio		
137	Edwards	265	Kerr	393	Roberts	CODE	COUNTY NAME
139	Ellis	267	Kimble	395	Robertson	001	Beaver
141	El Paso	269	King	397	Rockwall	003	Box Elder
143	Erath	271	Kinney	399	Runnels	005	Cache
145	Falls	273	Kleberg	401	Rusk	007	Carbon
147	Fannin	275	Knox	403	Sabine	009	Daggett
149		277		405		011	Davis
	Fayette	277	Lamar Lamb	403	San Augustine San Jacinto		Duchesne
151	Fisher					013	
153	Floyd	281	Lampasas	409	San Patricio	015	Emery
155	Foard	283	La Salle	411	San Saba	017	Garfield
157	Fort Bend	285	Lavaca	413	Schleicher	019	Grand
159	Franklin	287	Lee	415	Scurry	021	Iron
161	Freestone	289	Leon	417	Shackleford	023	Juab
163	Frio	291	Liberty	419	Shelby	025	Kane
165	Gaines	293	Limestone	421	Sherman	027	Millard
167	Galveston	295	Lipscomb	423	Smith	029	Morgan
			-				5

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

031	Piute
033	Rich
035	Salt Lake
037	San Juan
039	Sanpete
041	Sevier
043	Summit
045	Tooele
047	Uintah
049	Utah
051	Wasatch
053	Washington
055	Wayne
057	Weber

### STATE NAME: VERMONT ALPHABETIC CODE: VT **NUMERIC CODE: 50**

### CODE COUNTY NAME 001 Addison

001	ruuison
003	Bennington
005	Caldedonia
007	Chittenden
009	Essex
011	Franklin
013	Grand Isle
015	Lamoille
017	Orange
019	Orleans
021	Rutland
023	Washington
025	Windham
027	Windsor

### STATE NAME: VIRGINIA ALPHABETIC CODE: VA **NUMERIC CODE: 51**

### CODE COUNTY NAME

CODE	COUNTINAME
001	Accomack
003	Albermarle
005	Alleghany
007	Amelia
009	Amherst
011	Appomattox
013	Arlington
015	Augusta
017	Bath
019	Bedford
021	Bland
023	Botetourt
025	Brunswick
027	Buchanan
029	Buckingham
031	Campbell
033	Caroline
035	Carroll
036	Charles City
037	Charlotte

041	
041	
	Chesterfield
	<u>a</u> 1 1
043	Clarke
0.45	Craig
045	Craig
047	Culpeper
047	Culpeper
049	Cumberland
051	Dickenson
053	Dinwiddie
0.57	г
057	Essex
059	Fairfax
039	Faillax
061	Fauquier
	1
063	Floyd
0.6	
065	Fluvanna
0(7	Franklin
067	гтанкни
069	Frederick
071	Giles
073	Gloucester
075	Goochland
077	Grayson
0//	Grayson
079	Greene
081	Greensville
083	Halifax
	Uanovar
085	Hanover
087	Henrico
007	
089	Henry
091	Highland
002	Isle of Wight
093	Isle of wight
095	James City
097	King And Queen
099	King George
101	
101	King William
103	Lancaster
105	Lee
107	Loudoun
100	T autian
109	Louisa
111	Lunenburg
	•
113	Madison
115	Mathews
115 117	Mecklenburg
117	Mecklenburg
117 119	Mecklenburg Middlesex
117 119	Mecklenburg Middlesex
117 119 121	Mecklenburg Middlesex Montgomery
117 119	Mecklenburg Middlesex
117 119 121 125	Mecklenburg Middlesex Montgomery Nelson
117 119 121	Mecklenburg Middlesex Montgomery
117 119 121 125 127	Mecklenburg Middlesex Montgomery Nelson New Kent
<ol> <li>117</li> <li>119</li> <li>121</li> <li>125</li> <li>127</li> <li>131</li> </ol>	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton
<ol> <li>117</li> <li>119</li> <li>121</li> <li>125</li> <li>127</li> <li>131</li> </ol>	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton
117 119 121 125 127 131 133	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland
117 119 121 125 127 131 133	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland
117 119 121 125 127 131 133 135	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway
117 119 121 125 127 131 133	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland
117 119 121 125 127 131 133 135 137	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange
117 119 121 125 127 131 133 135 137 139	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page
117 119 121 125 127 131 133 135 137 139	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page
117 119 121 125 127 131 133 135 137 139 141	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick
117 119 121 125 127 131 133 135 137 139	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick
117 119 121 125 127 131 133 135 137 139 141 143	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania
117 119 121 125 127 131 133 135 137 139 141	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick
117 119 121 125 127 131 133 135 137 139 141 143 145	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan
117 119 121 125 127 131 133 135 137 139 141 143 145 147	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward
117 119 121 125 127 131 133 135 137 139 141 143 145 147	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George
117 119 121 125 127 131 133 135 137 139 141 143 145 147	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155 157	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155 157	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski
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117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155 157 159 161	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock
117 119 121 125 127 131 133 135 137 139 141 143 145 147 149 153 155 157 159 161	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock Richmond Roanoke
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$\begin{array}{c} 117 \\ 119 \\ 121 \\ 125 \\ 127 \\ 131 \\ 133 \\ 135 \\ 137 \\ 139 \\ 141 \\ 143 \\ 145 \\ 147 \\ 149 \\ 153 \\ 155 \\ 157 \\ 159 \\ 161 \\ 163 \\ 165 \\ 167 \\ \end{array}$	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock Richmond Roanoke Rockbridge Rockingham Russell
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$\begin{array}{c} 117\\ 119\\ 121\\ 125\\ 127\\ 131\\ 133\\ 135\\ 137\\ 139\\ 141\\ 143\\ 145\\ 147\\ 149\\ 153\\ 155\\ 157\\ 159\\ 161\\ 163\\ 165\\ 167\\ 169\\ 171\\ \end{array}$	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock Richmond Roanoke Rockbridge Rockingham Russell Scott Shenandoah
$\begin{array}{c} 117\\ 119\\ 121\\ 125\\ 127\\ 131\\ 133\\ 135\\ 137\\ 139\\ 141\\ 143\\ 145\\ 147\\ 149\\ 153\\ 155\\ 157\\ 159\\ 161\\ 163\\ 165\\ 167\\ 169\\ 171\\ \end{array}$	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock Richmond Roanoke Rockbridge Rockingham Russell Scott
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$\begin{array}{c} 117\\ 119\\ 121\\ 125\\ 127\\ 131\\ 133\\ 135\\ 137\\ 139\\ 141\\ 143\\ 145\\ 147\\ 149\\ 153\\ 155\\ 157\\ 159\\ 161\\ 163\\ 165\\ 167\\ 169\\ 171\\ \end{array}$	Mecklenburg Middlesex Montgomery Nelson New Kent Northampton Northumberland Nottoway Orange Page Patrick Pittsylvania Powhatan Prince Edward Prince George Prince William Pulaski Rappahannock Richmond Roanoke Rockbridge Rockingham Russell Scott Shenandoah

177	Spotsylvania
179	Stafford
181	Surry
183	Sussex
185	Tazewell
187	Warren
191 193	Washington Westmoreland
195	Wise
195	Wythe
199	York
CODE	
INDEP 510	ENDENT CITY Alexandria (city)
515	Bedford (city)
520	Bristol (city)
530	Buena Vista (city)
540	Charlottsville (city)
550	Chesapeake (city)
560	Clifton Forge (city)
570	Colonial Heights
(city)	
580	Covington (city)
590	Danville (city)
595 600	Emporia (city)
610	Fairfax (city) Falls Church (city)
620	Franklin (city)
630	Fredericksburg
(city)	Treatmensourg
640	Galax (city)
650	Hampton (city)
660	Harrisonburg (city)
670	Hopewell (city)
678	Lexington (city)
680	Lynchburg (city)
683	Manassas (city)
685 690	Manassas Park (city) Martinsville (city)
700	Newport News
(city)	Newport News
710	Norfolk (city)
720	Norton (city)
730	Petersburg (city)
735	Poquoson (city)
740	Portsmouth (city)
750	Radford (city)
760	Richmond (city)
770 775	Roanoke (city)
775 790	Salem (city) Staunton (city)
800	Suffolk (city)
810	Virginia Beach
(city)	
820	Waynesboro (city)
830	Williamsburg (city)
840	Winchester (city)
The cod	les for Charles City
	arlotte Counties,
reported	d respectively as 037
	in FIPS PUB 6-3,
have be	en corrected. The

Bureau of Economic Analysis, U.S. Department of Commerce has defined codes in the 900 series to represent county/independent city combination in Virginia.

The FIPS county code of 780 for South Boston, VA, is deleted. South Boston will be incorporated within Halifax County rather than a separate county-equivalent surrounded by Halifax County.

The independent city (county-equivalent) of Clifton Forge has reverted to town status, effective July 1, 2001. Clifton Forge is now an incorporated place within Alleghany County, rather than a separate countyequivalent surrounded by Alleghany County. The FIPS county code of 560 for Clifton Forge is deleted.

STATE NAME: WASHINGTON **ALPHABETIC CODE:** WA **NUMERIC CODE: 53** 

### CODE COUNTY NAME 001

Adams 003 Asotin 005 Benton 007 Chelan 009 Clallam 011 Clark 013 Columbia 015 Cowlitz 017 Douglas 019 Ferry 021 Franklin 023 Garfield 025 Grant 027 Gravs Harbor 029 Island 031 Jefferson 033 King 035 Kitsap 037 Kittitas 039 Klickitat 041 Lewis 043 Lincoln 045 Mason 047 Okanogan 049 Pacific 051 Pend Oreille 053 Pierce

Version 10.2 -- Appendix A: FIPS Codes for Counties and Equivalent Entities

087

055	San Juan
057	Skagit
059	Skamania
061	Snohomish
063	Spokane
065	Stevens
067	Thurston
069	Wahkiakum
071	Walla Walla
073	Whatcom
075	Whitman
077	Yakima

### STATE NAME: WEST VIRGINIA **ALPHABETIC CODE:** WV **NUMERIC CODE: 54**

### CODE COUNTY NAME 001 Barbour 003 Berkelev 005 Boone 007 Braxton 000 Brooke

009	Brooke
011	Cabell
013	Calhoun
015	Clay
017	Doddridge
019	Fayette
021	Gilmer
023	Grant
025	Greenbrier
027	Hampshire
029	Hancock
031	Hardy
033	Harrison
035	Jackson
037	Jefferson
039	Kanawha
041	Lewis
043	Lincoln
045	Logan
047	McDowell
049	Marion
051	Marshall
053	Mason
055	Mercer
057	Mineral
059	Mingo
061	Monongalia
063	Monroe
065	Morgan
067	Nicholas
069	Ohio
071	Pendleton
073	Pleasants
075	Pocahontas
077	Preston
079	Putnam
081	Raleigh
083	Randolph
085	Ritchie

087	Roane
089	Summers
091	Taylor
093	Tucker
095	Tyler
097	Upshur
099	Wayne
101	Webster
103	Wetzel
105	Wirt
107	Wood
109	Wyoming

### STATE NAME: WISCONSIN ALPHABETIC CODE: WI **NUMERIC CODE: 55**

## CODE COUNTY NAME

001 Adams 003 Ashland 005 Barron 007 Bayfield 009 Brown 011 Buffalo 013 Burnett 015 Calumet 017 Chippewa 019 Clark 021 Columbia 023 Crawford 025 Dane 027 Dodge 029 Door 031 Douglas 033 Dunn 035 Eau Claire 037 Florence 039 Fond du Lac 041 Forest 043 Grant 045 Green 047 Green Lake 049 Iowa 051 Iron 053 Jackson 055 Jefferson 057 Juneau 059 Kenosha 061 Kewaunee 063 La Crosse 065 Lafayette 067 Langlade 069 Lincoln 071 Manitowoc 073 Marathon 075 Marinette 077 Marquette 078 Menominee 079 Milwaukee 081 Monroe 083 Oconto 085 Oneida

089	Ozaukee
091	Pepin
093	Pierce
095	Polk
097	Portage
099	Price
101	Racine
103	Richland
105	Rock
107	Rusk
109	St. Croix
111	Sauk
113	Sawyer
115	Shawano
117	Sheboygan
119	Taylor
121	Trempealeau
123	Vernon
125	Vilas
127	Walworth
129	Washburn
131	Washington
133	Waukesha
135	Waupaca
137	Waushara
139	Winnebago

Outagamie

### STATE NAME: WYOMING **ALPHABETIC CODE:** WY **NUMERIC CODE: 56**

Wood

141

## CODE COUNTY NAME

001 Albany 003 Big Horn 005 Campbell 007 Carbon 009 Converse 011 Crook 013 Fremont 015 Goshen 017 Hot Springs 019 Johnson 021 Laramie 023 Lincoln 025 Natrona 027 Niobrara 029 Park 031 Platte 033 Sheridan 035 Sublette 037 Sweetwater 039 Teton 041 Uinta Washakie 043 045 Weston

### APPENDIX A

### **AREA NAME:** AMERICAN SAMOA ALPHABETIC CODE: AS **NUMERIC CODE: 60**

### CODE DISTRICT/ISLAND NAME

010 Eastern (District) 020 Manu'a (District) 030 Rose Island 040 Swains Island 050 Western (District)

"Island" is part of the name of Rose Island and Swains Island. The entities called "counties" in American Samoa are subdivisions of the districts, and therefore are second-order subdivisions of American Samoa.

### AREA NAME: GUAM ALPHABETIC CODE: GU **NUMERIC CODE: 66**

### CODE SUBDIVISION NAME 010 Guam

Guam has no first-order subdivisions, and therefore "Guam" also serves as the county-equivalent entity.

AREA NAME: NORTHERN MARINA **ISLANDS ALPHABETIC CODE:** MP **NUMERIC CODE: 69** 

### CODE

MUNICIPALITY NAME 085 Northern Islands 100 Rota 110 Saipan 120 Tinian

### **AREA NAME: PALAU** ALPHABETIC CODE: PW **NUMERIC CODE: 70**

CODE STATE NAME 002 Aimeliik

004	Airai
010	Angaur
050	Hatoboheit
100	Kayangel
150	Koror
212	Melekeok
214	Ngaraard
218	Ngarchelong
222	Ngardmau
224	Ngatpang
226	Ngchesar
227	Ngernmlengui
228	Ngiwal
350	Peleliu
370	Sonsorol

Palau also is known as Beau. and may be referred to as the Republic of ... " Changes since recognition of Palau in Change Notice No. 9 to FIPS PUB 6-3. The first-order subdivisions of Palau have been revised from municipalities to states; the name of Melekeiok has been revised to Melekeok; the name and code for Ngaremlengui (223) have been revised to Ngeremlengui (227); the name and code for Tobi (380) have been revised to Hatobohei (050); the Palau Islands (unorganized territory) (300) is no longer included because that area is part of Koror and Peleliu.

### **AREA NAME: PUERTO** RICO ALPHABETIC CODE: PR **NUMERIC CODE: 72**

CODE MUNICIPALITY NAME 001 Adjuntas Aguada 003 005 Aguadilla 007 Aguas Buenas 009 Aibonito 011 Anasco 013 Arecibo 015 Arroyo 017 Barceloneta 019 Barranquitas 021 Bayamo'n 023 Cabo Rojo 025 Caguas 027 Camuy 029 Canovanas 031 Carolina 033 Catano

035 Cayey 037 Ceiba 039 Ciales 041 Cidra 043 Coamo 045 Comerio 047 Corozal 049 Culebra 051 Dorado 053 Fajardo 054 Florida 057 Guayama 059 Guavanilla Guaynabo 061 063 Gurabo 065 Hatillo Hormigueros 067 069 Humacao 071 Isabela 073 Jayuya 075 Juana Diaz 077 Juncos 079 Lajas 081 Lares 083 Las Marias 085 Las Piedras 087 Loiza 089 Luquillo 091 Manati 093 Maricao 095 Maunabo 097 Mayaguez 099 Moca 101 Morovis 103 Naguabo 105 Naranjito Orocovis 109 Patillas Penuelas Ponce 113 115 Ouebradillas Rincon 119 Rio Grande 121 Sabana Grande 123 Salinas 125 San German 127 San Juan 129 San Lorenzo San Sebastian 131 133 Santa Isabel 135 Toa Alta 137 Toa Baja Trujillo Alto 139 141 Utuado 143 Vega Alta 145 Vega Baja 147 Vieques 149 Villalba Yabucoa 151 153 Yauco

107

111

117

AREA NAME: U.S. **OUTLYING ISLANDS ALPHABETIC CODE:** UM **NUMERIC CODE: 74** 

# CODE ISLAND NAME

050 Baker Island 100 Howland Island 150 Jarvis Island 200 Johnston Island 250 Kingman Reef 300 Midway Islands 350 Navassa Island 400 Palmyra Atoll 450 Wake Island

An FIPS State numeric code is available for each area: FIPS PUB 5-2 identifies the codes and explains their usage. The State codes can be used in combination with the "county" codes listed here.

### AREA NAME: VIRGIN **ISLANDS OF THE** UNITED STATES ALPHABETIC CODE: VI **NUMERIC CODE: 78**

CODE ISLAND NAME 010 St. Croix 020 St. John 030 St. Thomas

### APPENDIX B

AREA NAME: FEDERATED STATES OF MICRONESIA ALPHABETIC CODE: FM **NUMERIC CODE: 64** 

### CODE STATE NAME

002 Chuuk 005 Kosrae 040 Pohnpeit 060 Yap

The Federated States of Micronesia (FSM) became a freely associated state on 11/3/86. Its first-order subdivisions are called states. Changes since recognition of the FSM in Change Notice No. 9 to FIPS PUB 6-3. Ponape was renamed Pohnpei (11/8/84), and retained code 040; Truk

(050) was renamed Chuuk (10/1/89).

### **AREA NAME:** MARSHALL ISLANDS **ALPHABETIC CODE:** MH NUMERIC CODE: 68

CODE

### MUNICIPALITY NAME

007 Ailinginaie 010 Ailinglaplap 030 Ailuk 040 Arno 050 Aur 060 Bikar 070 Bikini 073 Bokak 080 Ebon 090 Enewetak 100 Erikub 110 Jabat 120 Jaluit 130 Jemo 140 Kili 150 Kwajalein 160 Lae 170 Lib 180 Likiep 190 Majuro 300 Maloelap 310 Mejit Mili 320 330 Namorik 340 Namu 350 Rongelap 360 Rongrik 385 Toke 390 Ujae 400 Ujelang 410 Utrik Wotho 420 430 Wotle

The Marshall Islands became a freely associated state on 11/3/86. Its first-order subdivisions also may be referred to as "islands" and "atolls." Since the recognition of the Marshall Islands in Change Notice No. 9. Jemo has been revised from Jemo Island to a municipality. Toke also may be spelled "Taka."

# **APPENDIX B**

# EDITS TABLES FOR SELECTED DATA ITEMS

## Table Name: BPLACE.DBF (SEER GEOCODES FOR CODING PLACE OF BIRTH)

### CONTINENTAL UNITED STATES AND HAWAII

### 000 United States

- 001 New England and New Jersey
  - 002 Maine
  - 003 New Hampshire
  - 004 Vermont
  - 005 Massachusetts
  - 006 Rhode Island
  - 007 Connecticut
  - 008 New Jersey
- 010 North Mid-Atlantic States
  - 011 New York
  - 014 Pennsylvania
  - 017 Delaware

### 020 South Mid-Atlantic States

- 021 Maryland
- 022 District of Columbia
- 023 Virginia
- 024 West Virginia
- 025 North Carolina
- 026 South Carolina
- 030 Southeastern States
  - 031 Tennessee
  - 033 Georgia
  - 035 Florida
  - 037 Alabama
  - 039 Mississippi
- 040 North Central States
  - 041 Michigan
    - 043 Ohio
    - 045 Indiana
    - 047 Kentucky
- 050 Northern Midwest States
  - 051 Wisconsin
  - 052 Minnesota
  - 053 Iowa
  - 054 North Dakota
  - 055 South Dakota
  - 056 Montana

060 Central Midwest States

- 061 Illinois
- 063 Missouri
- 065 Kansas
- 067 Nebraska

- 070 Southern Midwest States
  - 071 Arkansas
  - 073 Louisiana
  - 075 Oklahoma
  - 077 Texas
- 080 Mountain States
  - 081 Idaho
    - 082 Wyoming
    - 083 Colorado
    - 084 Utah
    - 085 Nevada
    - 086 New Mexico
    - 087 Arizona
- 090 Pacific Coast States
  - 091 Alaska
    - 093 Washington
    - 095 Oregon
    - 097 California
    - 099 Hawaii

### UNITED STATES POSSESSIONS

When SEER geocodes were originally assigned during the 1970s, the United States owned or controlled islands in the Pacific. Since then, many of these islands have either been given their independence or had control turned over to another country. In order to maintain consistent information over time, these islands are still to be coded to the original codes. Earlier designations are listed in parentheses.

- 100 Atlantic/Caribbean Area
  - 101 Puerto Rico
  - 102 U.S. Virgin Islands
  - 109 Other Atlantic/Caribbean Area
- 110 Canal Zone
- 120 Pacific Area
  - 121 American Samoa
  - 122 Kiribati (Canton and Enderbury Islands, Gilbert Islands, Southern Line Islands, Phoenix Islands)
  - 123 Micronesia [Federated States of] (Caroline Islands, Trust Territory of Pacific Islands)
  - 124 Cook Islands (New Zealand)
  - 125 Tuvalu (Ellice Islands)
  - 126 Guam
  - 127 Johnston Atoll
  - 129 Mariana Islands (Trust Territory of Pacific Islands)
  - 131 Marshall Islands (Trust Territory Pacific Islands)
  - 132 Midway Islands
  - 133 Nampo-Shoto, Southern

- 134 Ryukyu Islands (Japan)
- 135 Swan Islands
- 136 Tokelau Islands (New Zealand)
- 137 Wake Island
- 139 Palau (Trust Territory of Pacific Islands)

# NORTH AND SOUTH AMERICA, EXCLUSIVE OF THE UNITED STATES AND ITS POSSESSIONS

- 210 Greenland
- 220 Canada
  - 221 Labrador Maritime provinces New Brunswick Newfoundland and Labrador Nova Scotia Prince Edward Island
  - 222 Quebec
  - 223 Ontario
  - 224 Prairie provinces Alberta Manitoba Saskatchewan
  - 225 Northwest Territories Yukon Territory
  - 226 British Columbia
  - 227 Nunavut (Nunavut became an official Territory of Canada on April 1, 1999.)
- 230 Mexico
- 240 North American Islands
  - 241 Cuba
  - 242 Haiti
  - 243 Dominican Republic
  - 244 Jamaica
  - 245 Other Caribbean Islands Anguilla Antigua and Barbuda Barbados British Virgin Islands Cayman Islands Dominica Grenada Guadeloupe
    - Martinique
    - Montserrat
    - Netherlands Antilles
    - St. Kitts and Nevis
    - St. Lucia
    - St. Vincent and the Grenadines
    - Trinidad and Tobago
    - Turks and Caicos
    - Antilles, NOS
    - British West Indies, NOS
    - Carribean, NOS Leeward islands, NOS
    - West Indies, NOS
    - Windward islands, NOS
  - 246 Bermuda
  - 247 Bahamas
  - 249 St. Pierre and Miquelon

- 250 Central America
  - 251 Guatemala
    - 252 Belize (British Honduras)
  - 253 Honduras
  - 254 El Salvador
  - 255 Nicaragua 256 Costa Rica
  - 256 Costa Rica257 Panama
- 260 North America, NOS
- 265 Latin America, NOS
- 300 South America, NOS
  - 311 Colombia
  - 321 Venezuela
  - 331 Guyana (British Guiana)
  - 332 Suriname (Dutch Guiana)
  - 333 French Guiana
  - 341 Brazil
  - 345 Ecuador 351 Peru
  - 351 Peru355 Bolivia
  - 361 Chile
  - 365 Argentina
  - 371 Paraguay
  - 375 Uruguay
- 380 South American Islands381 Falkland Islands

### EUROPE

Former or alternative names are in parentheses

Europe, NOS (See code 499) \* \* Effective tumors diagnosed 1/1/92.

- 400 United Kingdom, NOS 401 England Channel Islands
  - Isle of Man
  - 402 Wales
  - 403 Scotland
  - 404 Northern Ireland (Ulster)
- 410 Ireland (Eire) Ireland, NOS Republic of Ireland
- 420 Scandinavia
  - Lapland, NOS 421 Iceland
  - 423 Norway
    - Svalbard
    - Jan Mayen
    - 425 Denmark
    - Faroe Islands 427 Sweden
    - 429 Finland

430	Germ	anic countries		463 Ba	altic Republic(s), NOS
	431	Germany			altic States, NOS)
		(East Germany including East Berlin)	470		inland Europe
		(West Germany including West Berlin)			reece
	432	Netherlands			ungary
	433	Belgium			ibania
	434	Luxembourg			braltar
		•		405 01	bianai
	435	Switzerland	400	04 M	1
	436	Austria	490		editerranean islands
	437	Liechtenstein			alta
					/prus
440	Roma	ance-language countries			irope, NOS*
	441	France			entral Europe, NOS
		Corsica		Ea	stern Europe, NOS
		Monaco		Nc	orthern Europe, NOS
	443	Spain		So	outhern Europe, NOS
		Andorra		We	estern Europe, NOS
		Balearic Islands			•
		Canary Islands	* Effe	ctive tumo	ors diagnosed 1/1/92.
	445	Portugal	-,,, ,		
	445	Azores			
			AFR		
		Cape Verde Islands	АГК	CA	
		Madeira Islands	500	1.C	100
	447	Italy	500	Africa, N	
		San Marino			Africa, NOS
		Sardinia		Equatoria	al Africa, NOS
		Sicily			
		Vatican City (Holy See)	510	North Afi	rica, NOS
	449	Romania		511 Mo	orocco
				513 Al	geria
450	Slavi	c countries		515 Tu	inisia
	451	Poland		517 Lil	bya
	452	(former) Czechoslovakia region			yrenaica)
	102	Bohemia			ripoli)
		Czech Republic			ripolitania)
		Moravia			gypt (United Arab Republic)
				519 Eg	gypt (Ollited Alab Republic)
		Slovak Republic	520	G 1	
		Slovakia	520		e countries
	453	(former) Yugoslavia region			Faso (Upper Volta)
		Bosnia-Herzogovina		Chad	
		Croatia		Mali	
		Dalmatia		Mauritan	ia
		Montenegro		Niger	
		Macedonia		Sudan (A	nglo-Egyptian Sudan)
		Serbia		Western (	(Spanish) Sahara
		Slavonia			
		Slovenia	530	West Afr	ica, NOS
	454	Bulgaria			Vest Africa, NOS
	455	Russia			geria
	100	Russian Federation			ther West African Countries
		(former) U.S.S.R.			enin (Dahomey)
		Russia, NOS			ameroon (Kameroon)
					entral African Republic (French
	150	(Russian S.F.S.R.)			1
	456	Ukraine and Moldova			uatorial Africa)
		(Bessarabia)			ote d-Ivoire (Ivory Coast)
		Moldavia			ongo (Congo-Brazzaville, French Congo)
		(Moldavian S.S.R.)			uatorial Guinea (Spanish Guinea) (Bioko [Fernando
		(Ukranian S.S.R.)			oo], Rio Muni)
	457	Belarus			ambia
		(Byelorussian S.S.R.)		Ga	abon
		(White Russia)		Gh	nana
	458	Estonia (Estonian S.S.R.)		Gu	linea
	459	Latvia (Latvian S.S.R.)			uinea Bissau (Portuguese Guinea)
	461	Lithuania			beria
		(Lithuanian S.S.R.)			enegal
		(		50	· O···

Togo

- 540 South Africa, NOS
  - 541 Zaire (Congo-Leopoldville, Belgian Congo, Congo/Kinshasa)
  - 543 Angola (Sao Tome, Principe, Cabinda)
  - 545 Republic of South Africa (Bophuthatswana, Cape Colony, Ciskei, Natal, Free State [Orange Free State], Transkei, Transvaal, Venda) Botswana (Bechuanaland) Lesotho (Basutoland)
    - Namibia (South West Africa)
  - Swaziland
  - 547 Zimbabwe (Rhodesia, Southern Rhodesia)
  - 549 Zambia (Northern Rhodesia)
  - 551 Malawi (Nyasaland)
  - 553 Mozambique
  - 555 Madagascar (Malagasy Republic)
- 570 East Africa
  - 571 Tanzania (Tanganyika, Tanzanyika, Zanzibar)
  - 573 Uganda
  - 575 Kenya
  - 577 Rwanda (Ruanda)
  - 579 Burundi (Urundi)
  - 581 Somalia (Somali Republic, Somaliland)
  - 583 Djibouti (French Territory of the Afars and Issas, French Somaliland)
  - 585 Ethiopia (Abyssinia) Eritrea
- 580 African Coastal Islands (previously included in 540) Comoros
  - Mauritius Mayotte Reunion St. Helena
  - Seychelles
- \* Effective tumors diagnosed 1/1/92.

### ASIA

- 600 Asia, NOS\*
- 610 Near East
  - Mesopotamia, NOS
    - 611 Turkey
      - Anatolia Asia Minor, NOS
- 620 Asian Arab Countries Iraq-Saudi Arabia Neutral Zone
  - 621 Syria
  - 623 Lebanon
  - 625 Jordan (Transjordan, former Arab Palestine)
  - 627 Iraq
  - 629 Arabian Peninsula
    - Bahrain
    - Kuwait
    - Oman and Muscat
    - Persian Gulf States, NOS
  - Qatar Saudi Arabia
  - United Arab Emirates (Trucial States)
    - Yemen (Aden, People's Democratic Republic of

Yemen, Southern Yemen)

- 631 Israel and former Jewish Palestine
   Gaza
   Palestine, NOS
   Palestine (Palestinian National Authority [PNA])
  - West Bank
- 633 Caucasian Republics of the former U.S.S.R. Armenia Azerbaijan (Nagorno-Karabakh) Georgia
- 634 Other Asian Republics of the former U.S.S.R. Kazakhstan (Kazakh S.S.R.) Kyrgystan (Kirghiz S.S.R., Kyrgyz) Tajikistan (Tadzhik S.S.R.) Turkmenistan (Turkmen S.S.R.) Uzbekistan (Uzbek S.S.R.)
- 637 Iran (Persia)
- 638 Afghanistan
- 639 Pakistan (West Pakistan)
- 640 Mid-East Asia, NOS
  - Maldives
    - 641 India, Andaman Islands
    - 643 Nepal, Bhutan, Sikkim
    - 645 Bangladesh (East Pakistan)
    - 647 Sri Lanka (Ceylon)
    - 649 Myanmar (Burma)
- 650 Southeast Asia 651 Thailand (Siam)
- 660 Indochina
  - 661 Laos
    - 663 Cambodia, Kampuchea
    - 665 Vietnam (Tonkin, Annam, Cochin China)
    - 671 Malaysia, Singapore, Brunei
    - 673 Indonesia (Dutch East Indies)
    - 675 Philippines (Philippine Islands)
- 680 East Asia
  - 681 China, NOS
  - 682 China (People's Republic of China)
  - 683 Hong Kong
  - 684 Taiwan (Formosa, Republic of China)
  - 685 Tibet
  - 686 Macao (Macau)
  - 691 Mongolia
  - 693 Japan
  - 695 Korea North Korea South Korea
    - South Kor
- \* Effective tumors diagnosed 1/1/92.

## AUSTRALIA AND OCEANIA

- 711 Australia and Australian New Guinea
- 715 New Zealand
- Niue
- 720 Pacific Islands Oceania, NOS Polynesia, NOS
- 721 Melanesian Islands Solomon Islands Fiji Fotuna New Hebrides Vanuatu Wallis
- 723 Micronesian Islands
- 725 Polynesian Islands
- 750 Antarctica

Except possessions of the United States.

## PLACE OF BIRTH UNKNOWN

- 998 Place of Birth stated not to be in United States, but no other information available
- 999 Place of Birth unknown

References: *CIA World Factbook*, 1995. U.S. Bureau of the Census Place of Birth Technical Documentation, 1997.

### ALPHABETICAL LISTING

\* Effective tumors diagnosed 1/1/92.

#### А 585 Abyssinia 629 Aden 583 Afars and Issas 638 Afghanistan 500 Africa 570 Africa, East 510 Africa, North 540 Africa, South 545 Africa, South West 530 Africa, West 580 African Coastal Islands (previously included in 540) 037 Alabama 091 Alaska 481 Albania 224 Alberta 513 Algeria 250 America, Central 260 America, North (see also North America) 300 America, South 121 American Samoa 611 Anatolia 641 Andaman Islands 443 Andorra 543 Angola 245 Anguilla 665 Annam 750 Antarctica 245 Antigua Antilles, NOS 245 245 Antilles, Netherlands 625 Arab Palestine 629 Arabia, Saudi 629 Arabian Peninsula 365 Argentina 087 Arizona 071 Arkansas 633 Armenia (U.S.S.R.) 611 Armenia (Turkey) 750 Antarctica 245 Aruba 600 Asia, NOS\* 680 Asia, East 640 Asia, Mid-East 610 Asia Minor, NOS 610 Asia, Near-East 650 Asia, Southeast 634 Asian Republics of the former U.S.S.R. 620 Asian Arab countries 100 Atlantic/Caribbean area, U.S. possessions 109 Atlantic/Caribbean area, other U.S. possessions 711 Australia 711 Australian New Guinea 436 Austria 633 Azerbaijan

633	Azerbaizhan S.S.R.
445	Azores
	В
247	Bahamas
629	Bahrain
443	Balearic islands
463	Baltic Republic, NOS
463	Baltic States, NOS
645	Bangladesh
245	Barbados
245	Barbuda
431	Bavaria
545	Basutoland
545	Bechuanaland
457	Belarus
541	Belgian Congo
433	Belgium
252	Belize
539	Benin
246	Bermuda
456	Bessarabia
643	Bhutan
539	Bioko (Fernando Poo)
452	Bohemia
355	Bolivia
545	Bophuthatswana
673	Borneo
453	Bosnia-Herzogovina
545	Botswana
341	Brazil
226	British Columbia
331	British Guiana
252	British Honduras
245	British Virgin Islands
245	British West Indies, NOS
671	Brunei
454	Bulgaria
520	Burkina Faso (Upper Volta)
649	Burma
	(see Myanmar)
579	Burundi
457	Byelorussian S.S.R.
	С
543	Cabinda
245	Caicos Islands
097	California
663	Cambodia
520	Companyon

539

220

110

443

122

545

445

245

245

123

711

633

245

Cameroon

Canal Zone

Canary islands

Canton islands

Cape Verde islands

Caribbean islands, other

Caucasian Republics of the

Caribbean, NOS

Caroline Islands

Cartier Islands

former U.S.S.R.

Cayman Islands

Cape Colony

Canada

Central Africa, NOS
Central African Republic
Central America
Central Europe, NOS
Central Midwest States
Ceylon
Chad
Channel Islands (British)
Chile
China
(not otherwise specified)
China, Cochin
China, People's Republic of
China, Republic of
Christmas Island
Ciskel
Cochin China
Cocos (Keeling) Islands
Colombia
Colorado
Comoros
Columbia, British
Columbia, District of
Congo-Brazzaville
Congo-Leopoldville
Congo, Belgian
Congo, French
Congo Kinshasa
Connecticut
Cook Islands
Corsica
Costa Rica
Cote d'Ivoire (Ivory Coast)
Crete
Croatia
Cuba
Curacao
Cyprus
Cyrenaica
Czechoslovakia
Czech Republic

### D

539	Dahomey
453	Dalmatia
017	Delaware
425	Denmark
022	District of Columbia
583	Djibouti
449	Dobruja
245	Dominica
243	Dominican Republic
673	Dutch East Indies
332	Dutch Guiana
	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

425	Faroe (Faeroe) Islands
381	Falkland Islands
431	Federal Republic of Germany
539	Fernando Poo
721	Fiji
429	Finland
035	Florida
684	Formosa
721	Fotuna
441	France
545	Free State (Orange Free State)
539	French Congo
333	French Guiana
725	French Polynesia
583	French Somaliland
530	French West Africa, NOS
245	French West Indies

# 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	Guadaloupe
126	Guam
251	Guatamala
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
551	Sujulu
	Н
242	Haiti
099	Hawaii
432	Holland
253	Honduras
252	Honduras, British
683	Hong Kong
475	Hungary
	Ι
421	Iceland
081	Idaho
061	Illinois
641	India
045	Indiana
673	Indies, Dutch East
660	Indochina
673	Indonesia
053	Iowa
637	Iran
627	Iraq
620	Iraq-Saudi Arabian Neutral Zone
410	Ireland (Eire)
404	Ireland, Northern
410	Ireland, NOS
410	Ireland, Republic of
401	Isle of Man
631	Israel
583	Issas
447	Italy
539	Ivory Coast
	T

# J

423	Jan Mayen
244	Jamaica
693	Japan
673	Java
401	Jersey
631	Jewish Palestine
127	Johnston Atoll
625	Jordan
453	Jugoslavia
	K
539	Kameroon
663	Kampuchea

559	Kameroon
663	Kampuchea
065	Kansas
634	Kazakh S.S.R.
634	Kazakhstan
047	Kentucky
575	Kenya
634	Kirghiz S.S.R.

122	Kiribati
695	Korea
695	Korea, North
695	Korea, South
629	Kuwait
634	Kyrgystan
634	Kyrgyz
	L
221	Labrador
661	Laos
265	Latin America, NOS
420	Lapland, NOS
459	Latvia
459	Latvian S.S.R. (Latvia)
623	Lebanon
245	Leeward island, NOS
545	Lesotho
539	Liberia
517	Libya
437	Liechtenstein
122	Line Islands, Southern
461	Lithuania
461	Lithuanian S.S.R. (Lithuania)
073	Louisiana
434	Luxembourg

## М

686	Macao
686	Macau
453	Macedonia
555	Madagascar
445	Madeira islands
002	Maine
555	Malagasy Republic
551	Malawi
671	Malay Peninsula
671	Malaysia
640	Maldives
520	Mali
491	Malta
224	Manitoba
129	Mariana Islands
221	Maritime provinces, Canada
131	Marshall Islands
245	Martinique
021	Maryland
005	Massachusetts
520	Mauritania
580	Mauritius
580	Mayotte
490	Mediterranean Islands, Other
721	Melanesian islands
610	Mesopotamia, NOS
230	Mexico
041	Michigan
123	Micronesian islands
640	Mid-East Asia
132	Midway Islands
052	Minnesota
249	Miquelon
039	Mississippi
063	Missouri

(Canada)

456	Moldavia
456	Moldavian S.S.R.
456	Moldova
441	Monaco
691	Mongolia
056	Montana
453	Montenegro
245	Montserrat
452	Moravia
511	Morocco
080	Mountain States
553	Mozambique
629	Muscat
649	Myanmar
	(See Burma)

### Ν

545	Namibia
133	Nampo-shoto, Southern
545	Natal
723	Nauru
610	Near-East Asia
067	Nebraska
643	Nepal
432	Netherlands
245	Netherlands Antilles
332	Netherlands Guiana
085	Nevada
245	Nevis
221	New Brunswick
725	New Caledonia
001	New England
673	New Guinea, except
	Australian and North East
711	New Guinea, Australian
711	New Guinea, North East
003	New Hampshire
721	New Hebrides
008	New Jersey
086	New Mexico
011	New York
715	New Zealand
221	Newfoundland
255	Nicaragua
520	Niger
531	Nigeria
715	Niue
711	Norfolk Island
671	North Borneo (Malaysia)
510	North Africa, NOS
260	North America, NOS (use more
	specific term if possible)
240	North American islands
025	North Carolina
040	North Central States
054	North Dakota
711	North East New Guinea
695	North Korea
010	North Mid-Atlantic States
499	Northern Europe, NOS
404	Northern Ireland
129	Northern Mariana Islands
050	Northern Midwest States
549	Northern Rhodesia
225	Northwest Territories

100	(Canada)	
423	Norway	
998	Not United States, NOS	
221	Nova Scotia	
227	Nunavut	
551	Nyasaland	
	5	
	0	
043	Ohio	
075	Oklahoma	
629	Oman	
223	Ontario	
545	Orange Free State	
095	Oregon	
403	Orkney Islands	
405	Orkney Islands	
	Р	
120	Pacific area, U.S. possessions	
720	Pacific islands	
123	Pacific Islands, Trust Territory of	
	the (code to specific islands if	
	possible)	
090	Pacific Coast States	
639	Pakistan	
645	Pakistan, East	
639	Pakistan, West	
139	Palau (Trust Territory of the	
	Pacific Islands)	
625	Palestine, Arab	
631	Palestine, Jewish	
631	Palestine, NOS	
631	Palestinian National Authority	
001	(PNA)	
257	Panama	
711	Papua New Guinea	
371	Paraguay	
014	Pennsylvania	
629	People's Democratic Republic	
02)	of Yemen	
682	People's Republic of China	
637	Persia	
629	Persian Gulf States, NOS	
351	Peru	
675	Philippine Islands	
675	**	
	Philippines Bitagirm	
725	Pitcairn	
451	Poland	
725	Polynesian islands	
445	Portugal	
539	Portuguese Guinea	
224	Prairie Provinces, Canada	
221	Prince Edward Island	
543	Principe	
101	Puerto Rico	
	Q	
620	Oatar	
629 222	Qatar Quebec	
<i>LLL</i>	Quebec	
	R	
	is a second seco	

684

545

Republic of China

Republic of South Africa

580	Reunion
006	Rhode Island
547	Rhodesia
549	Rhodesia, Northern
547	Rhodesia, Southern
539	Rio Muni
440	Romance-language countries
449	Romania
449	Roumania
577	Ruanda
449	Rumania
455	Russia, NOS
457	Russia, White
455	Russian Federation
	(former U.S.S.R.)
455	Russian S.F.S.R.
577	Rwanda
134	Ryukyu Islands
	S

### S

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	Sevchelles
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
581	Somaliland
583	Somaliland, French
540	South Africa
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

Republics (U.S.S.R.) (see

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
	Τ
634	Tadzhik S.S.R.
684	Taiwan

054	Tauzink D.D.R.
684	Taiwan
634	Tajikistan
571	Tanzania
571	Tanganyika
571	Tanzanyika
031	Tennessee
077	Texas
651	Thailand (Siam)
685	Tibet
245	Tobago
539	Togo
136	Tokelau Islands
725	Tonga
665	Tonkin
625	Trans-Jordan
545	Transkei
545	Transvaal
449	Transylvania
245	Trinidad
517	Tripoli
517	Tripolitania
629	Trucial States
515	Tunisia
611	Turkey
634	Turkmen S.S.R.
634	Turkmenistan
245	Turks Islands
125	Tuvalu
	U
573	Uganda
456	Ukraine
456	Ukranian S.S.R.
404	Ulster
545	Union of South Afri

	Republics (0.5.5.R.) (see
	individual republics)
629	United Arab Emirates
519	United Arab Republic
400	United Kingdom
000	United States
102	U.S. Virgin Islands
999	Unknown
520	Upper Volta
375	Uruguay
579	Urundi
084	Utah
634	Uzbekistan
634	Uzbek S.S.R.

### V

721	Vanuatu
447	Vatican City
545	Venda
321	Venezuela
004	Vermont
665	Vietnam
102	Virgin Islands (U.S.)
245	Virgin Islands (British)
102	Virgin Islands (U.S.)
245	Virgin Islands (British)
023	Virginia

### W

Wake Island
Wales
Wallis
Wallachia
Washington (state)
Washington D.C.
West Africa, NOS
West African countries, other
West Bank
West Germany
West Indies, NOS (see also
individual islands)
West Pakistan
West Virginia
Western Europe, NOS
Western Sahara
Western Samoa
White Russia
Windward islands
Wisconsin
Wyoming
Y
Yemen
Yemen, People's Democratic

- Republic of
- 453 Yugoslavia (former
- Yugoslavia region) 225 Yukon Territory

Z

456	Ukranian S.S.R.	541	Zaire
404	Ulster	549	Zambia
545	Union of South Africa	571	Zanzibar
	Union of Soviet Socialist	547	Zimbabwe

Version 10.2 -- Appendix B: EDITS Tables for Selected Data Items

# Table Name: PEDSTAGE.DBF

1	Stage I
1A	Stage IA
1B	Stage IB
2	Stage II
2A	Stage IIA
2B	Stage IIB
2C	Stage IIC
3	Stage III
3A	Stage IIIA
3B	Stage IIIB
3C	Stage IIIC
3D	Stage IIID
3E	Stage IIIE
4	Stage IV
4A	Stage IVA
4B	Stage IVB
4S	Stage IVS
5	Stage V
A	Stage A

- Stage B В
- С Stage C
- Stage D D
- Stage DS DS
- Not applicable (not pediatric case) 88
- Unstaged, unknown 99

#### **Table Name: REGID.DBF**

000000200 Maine Cancer Incidence Registry 0000000300 New Hampshire State Cancer Registry 0000000400 Vermont Cancer Registry 0000000500 Massachusetts Cancer Registry 0000000580 Southeast Massachusetts Cancer Registry 0000000581 Greater Lowell Cancer Program 0000000600 Rhode Island Cancer Registry 000000700 Connecticut Tumor Registry 0000000800 New Jersey State Cancer Registry 0000001100 New York State Cancer Registry 0000001180 Rochester Regional Tumor Registry 0000001400 Pennsylvania Cancer Registry 0000001480 Pennsylvania-Northeast Regional Cancer Ctr. 0000001480 Northeast Regional Cancer Center 0000001500 National Cancer Institute SEER Program 0000001500 SEER Program, National Cancer Institute 0000001501 SEER San Francisco-Oakland SMSA 0000001502 SEER Connecticut 0000001520 SEER Metropolitan Detroit 0000001521 SEER Hawaii 0000001522 SEER Iowa 0000001523 SEER New Mexico 0000001525 SEER Seattle-Puget Sound 0000001526 SEER Utah 0000001527 SEER Metropolitan Atlanta 0000001529 SEER Alaska Native 0000001531 SEER San Jose-Monterey 0000001533 SEER Arizona Indians 0000001535 SEER Los Angeles 0000001537 SEER Rural Georgia 0000001541 SEER California except LA, SF-Oak, and San Jose/Monterey 0000001542 SEER Kentucky 0000001543 SEER Louisiana 0000001544 SEER New Jersey 0000001551 Cherokee Nation-Oklahoma (NCI funded) 0000001680 National Cancer Data Base 0000001700 Delaware State Cancer Registry 0000001801 Central Brain Tumor Registry of the U.S. 0000001900 U.S. Army Central Registry (ACTUR) 0000001900 Automated Central Tumor Registry (ACTUR) 0000002100 Maryland Cancer Registry 0000002200 District of Columbia Central Cancer Registry 0000002300 Virginia Cancer Registry 0000002400 West Virginia Cancer Registry 0000002500 North Carolina Central Cancer Registry 0000002600 South Carolina Central Cancer Registry 0000002601 Savannah River Region Cancer Registry in SC 0000002601 South Carolina - Savannah River Region in SC 0000003100 Tennessee Cancer Reporting System 0000003300 Georgia Center for Cancer Statistics 0000003300 Georgia Cancer Registry 0000003301 Georgia-Metropolitan Atlanta Cancer Registry 0000003301 Metropolitan Atlanta Cancer Registry 0000003302 Georgia-Rural Georgia Cancer Registry 0000003302 Rural Georgia Cancer Registry 0000003303 Georgia-Savannah River Region Cancer Registry 0000003303 Savannah River Region Cancer Registry in GA 0000003500 Florida Cancer Data System 0000003700 Alabama State Cancer Registry 0000003900 Mississippi State Cancer Registry

0000004100 Michigan Cancer Surveillance System 0000004101 Michigan Cancer Foundation, CA Surveillance Detroit Metropolitan Area 0000004101 Detroit Metropolitan 0000004300 Ohio Bureau of Chronic Disease 0000004301 Cancer Data System, Inc. 0000004301 Ohio-Cancer Data System, Inc. 0000004500 Indiana State Cancer Registry 0000004700 Kentucky Cancer Registry 0000005100 Wisconsin Cancer Reporting System 0000005200 Minnesota Cancer Surveillance System 0000005300 Iowa State Health Registry 0000005300 State Health Registry of Iowa 0000005400 North Dakota Cancer Registry 0000005500 South Dakota Cancer Registry 0000005600 Montana Central Tumor Registry 0000006100 Illinois State Cancer Registry 0000006300 Missouri Cancer Registry 0000006500 Kansas-Cancer Data Service 0000006500 Cancer Data Service 0000006700 Nebraska Cancer Registry 0000007100 Arkansas CART I 0000007300 Louisiana Tumor Registry 0000007301 New Orleans Regional Cancer Registry 0000007301 Louisiana Region I 0000007302 Baton Rouge Regional Tumor Registry 0000007302 Louisiana Region II 0000007303 Southeast Louisiana Regional Cancer Registry 0000007303 Louisiana Region III 0000007304 Acadiana Tumor Registry 0000007304 Louisiana Region IV 0000007305 Southwest Louisiana Regional Tumor Registry 0000007305 Louisiana Region V 0000007306 Central Louisiana Regional Tumor Registry 0000007306 Louisiana Region VI 0000007307 Northwest Louisiana Regional Tumor Registry 0000007307 Louisiana Region VII 0000007308 Northeast Louisiana Regional Tumor Registry 0000007308 Louisiana Region VIII 0000007309 New Orleans/Southeast Louisiana Reg. CA RegLouisiana's regions I and III combined 0000007310 North Louisiana Regional Tumor Registry; Louisiana's regions VI, VII, and VIII 0000007500 Oklahoma State Department of Health 0000007580 Eastern Oklahoma Regional Registry 0000007580 Oklahoma-Eastern Regional Registry 0000007700 Texas Cancer Incidence Reporting System 0000008100 Cancer Data Registry of Idaho 0000008100 Idaho Cancer Data Registry 0000008200 Wyoming Central Tumor Registry 0000008300 Colorado Central Cancer Registry 0000008400 Utah Cancer Registry 0000008500 Nevada Statewide Cancer Registry 0000008600 New Mexico Tumor Registry 0000008601 Arizona Indians; data collected by New Mexico Tumor Reg. 0000008700 Arizona Cancer Registry 0000009100 Alaska State Cancer Registry 0000009101 Alaska Area Native Health Service 0000009300 Washington State Cancer Registry

0000009301 Cancer Surveillance System Fred Hutchinson; Seattle Puget Sound area, 13 counties 0000009301 Washington-Seattle-Puget Sound 0000009302 Eastern Washington State Cancer Registry 0000009302 Washington - Eastern State Cancer Registry 0000009380 Spokane Central Tumor Registry (multihospital) 0000009380 Washington - Spokane Central Tumor Registry (multihospital) 0000009500 Oregon State Cancer Registry 0000009580 Sisters of Providence Cancer Registry 0000009580 Oregon-Sisters of Providence Cancer Reg. 0000009700 California Cancer Registry 0000009701 California Region 1 0000009701 San Jose-Monterey 0000009701 Greater Bay Area Cancer Registry (Region 1) 0000009702 California Region 2 0000009702 Cancer Registry of Central California 0000009703 California Region 3 0000009703 Cancer Surveillance Program, Region 3 0000009704 California Region 4 0000009704 Tri-Counties Regional Cancer Registry 0000009705 California Region 5 0000009705 Cancer Surveillance Program, Region 5 0000009706 California Region 6 0000009706 Cancer Registry of Northern California 0000009707 California Region 7 0000009707 San Diego/Imperial Org. for Cancer Control 0000009708 California Region 8 0000009708 San Francisco-Oakland SMSA 0000009708 Greater Bay Area Cancer Registry (Region 8) 0000009709 California Region 9 0000009709 Cancer Surveillance Program of Los Angeles 0000009709 Los Angeles 0000009710 California Region 10 0000009710 Cancer Surveillance Program of Orange County 0000009711 Greater Bay Area Cancer Registry; California's Regions 1 and 8 combined 0000009711 California Greater Bay Area Cancer Registry 0000009712 California CSPOC and SANDIOCC; California's Regions 7 and 10 combined 0000009900 Hawaii Tumor Registry 0010100000 Puerto Rico Central Cancer Registry 0022000000 Canadian Cancer Registry 0022001000 Newfoundland Cancer Treatment & Research Fnd. 0022001100 Prince Edward Island Cancer Registry 0022001200 Nova Scotia Cancer Registry 0022001300 New Brunswick Provincial Cancer Registry 0022002400 Fichier Des Tumeurs Du Quebec 0022002400 Quebec Cancer Registry 0022003500 Ontario Cancer Registry 0022004600 Manitoba Cancer Registry 0022004700 Saskatchewan Cancer Foundation 0022004800 Alberta Cancer Registry 0022005900 British Columbia Cancer Registry 0022006000 Yukon Bureau of Statistics 0022006100 Northwest Territories Department of Health 0088820020 Veterans Health Administration

#### Table Name: STATE.DBF

4.15	A 11
AB	Alberta
AK	Alaska
AL	Alabama
AR	Arkansas
AS	American Samoa
AZ	Arizona
BC	British Columbia
CA	California
CO	Colorado
СТ	Connecticut
DC	District of Columbia
DE	Delaware
FL	Florida
FM	Federated States of Micronesia
GA	Georgia
GU	Guam
HI	Hawaii
IA	Iowa
ID	Idaho
IL	Illinois
IN	Indiana
KS	Kansas
KY	Kentucky
LA	Louisiana
MA	Massachusetts
MB	Manitoba
MD	Maryland
ME	Maine
MH	Marshall Islands
MI	Michigan
MN	Minnesota
MO	Missouri
MP	Northern Mariana Islands
MS	Mississippi
MT	Montana
NB	New Brunswick
NC	North Carolina
ND	North Dakota
NE	Nebraska
NL	Newfoundland and Labrador
NH	New Hampshire
NJ	New Jersey
NM	New Mexico
NS	Nova Scotia
NT	Northwest Territories
NU	Nunavut
NV	Nevada
NY	New York
OH	Ohio
OK	Oklahoma
ON	Ontario
OR	Oregon
PA	Pennsylvania
PE	Prince Edward Island
PR	Puerto Rico
PW	Palau
QC RI	Quebec Rhode Island
SC	NINGLE ISTALL
	South Carolina
SD SK	

TN	Tennessee
TT	Trust Territories
TX	Texas
UM	US Minor Outlying Islands
UT	Utah
VA	Virginia
VI	Virgin Islands
VT	Vermont
WA	Washington
WI	Wisconsin
WV	West Virginia
WY	Wyoming
XX	Country Known, Not U.S., Not Canada
YT	Yukon Territories
YY	Country Unknown, Not U.S., Not Canada
ZZ	U.S., NOS; Canada, NOS; Country Unknown
AA	APO/FPO for Armed Services America
AE	APO/FPO for Armed Services Europe
AP	APO/FPO for Armed Services Pacific

Version 10.2 -- Appendix B: EDITS Tables for Selected Data Items

# **APPENDIX C**

## ABBREVIATIONS AND ACRONYMS USED

AACCR	American Association of Central Cancer Registries
ACoS	American College of Surgeons
ACS	American Cancer Society
AJCC	American Joint Committee on Cancer
BNA	Block Numbering Area
CDC	Centers for Disease Control and Prevention
CIN	Cervical intraepithelial neoplasia
CIS	Carcinoma <i>in situ</i>
CLIA	Clinical Laboratory Improvement Act
COC	Commission on Cancer (of ACoS)
CPT	Current Procedural Terminology (codes)
CRC	Cyclic redundancy code
CS	
CTR	Collaborative Staging
	Certified Tumor Registrar
DAM	Data Acquisition Manual (of ACoS)
DCO	Death Certificate Only
EOD	Extent of Disease
FIPS	Federal Information Processing Standards
FORDS	Facility Oncology Registry Data Standards (manual of ACoS)
FTRO	Fundamental Tumor Registry Operations Program (of ACoS)
GenEDITS	Generic EDITS Driver Program
GIS	Geographic Information System
HCFA	Health Care Finance Administration
HIM	Health Information Management
IACR	International Association of Cancer Registrars
IARC	International Agency for Research on Cancer
ICD	International Classification of Diseases
ICD-O	International Classification of Diseases for Oncology
ICD-O-1	International Classification of Diseases for Oncology, First Edition
ICD-O-2	International Classification of Diseases for Oncology, Second Edition
ICD-O-3	International Classification of Diseases for Oncology, Third Edition
NAACCR	North American Association of Central Cancer Registries
NCDB	National Cancer Data Base
NCI	National Cancer Institute
NCRA	National Cancer Registrars Association
N.d.	No date (bibliographic term: no ascertainable place of publication)
NHIA	NAACCR Hispanic Identification Algorithm
PIN	Prostatic intraepithelial neoplasia
ROADS	Registry Operations and Data Standards (manual of ACoS)
SEER	Surveillance, Epidemiology, and End Results Program of NCI
SIL	Squamous intraepithelial lesion
SS	Summary Stage
TNM	Tumor, Nodes and Metastasis: staging system of AJCC and UICC
UDSC	Uniform Data Standards Committee of NAACCR
UICC	Union Internationale Contre le Cancer (in English, International Union Against Cancer)
USPS	United States Postal Service
WHO	World Health Organization

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# **APPENDIX D**

### ALTERNATE NAMES

Following the item name are other names by which the same item is called, including the name used by the standard setter for the item. All other names are followed by the source of each name indicated with the following labels:

COC COC pre-96 COC pre-98 NAACCR pre-98 SEER SEER pre-98 Previously used name appearing in the COC *ROADS Manual* and Supplements Previously used name appearing in the COC *ROADS Manual* Previously used name appearing in the COC *ROADS Manual* before 1998 Previously used name appearing in NAACCR standards before 1998 Name in the *SEER Program Code Manual*, Third Edition (1998) Previously used name appearing in SEER Manual before 1998

Item #	Item Name	Alternate Names
70	Addr at DXCity	City or Town (pre-96 COC) City/Town at Diagnosis (COC)
80	Addr at DXState	State (pre-96 COC) State at Diagnosis (COC)
90	County at DX	County (pre-96 SEER/COC) County at Diagnosis (COC)
100	Addr at DXPostal Code	Postal Code at Diagnosis (COC) ZIP Code (pre-COC)
110	Census Tract 1970/80/90	Census Tract/Block Numbering Area (BNA) (SEER) Census Tract
120	Census Cod Sys 1970/80/90	Census Coding System (COC) Coding System for Census Tract (pre-96 SEER/COC)
130	Census Tract 2000	Census TractAlternate
150	Marital Status at DX	Marital Status at Diagnosis (SEER/COC) Marital Status at Initial Diagnosis (pre-96 COC)
160	Race 1	Race
190	Spanish/Hispanic Origin	Spanish OriginAll Sources (96 COC) Spanish Surname or Origin (SEER)
240	Birth Date	Date of Birth (SEER/COC)
250	Birthplace	Place of Birth (SEER/COC)
364	Census Tr Cert 1970/80/90	Census Tract Certainty
380	Sequence NumberCentral	Sequence Number (pre-96 SEER)
390	Date of Diagnosis	Date of Initial Diagnosis (COC)
410	Laterality	Laterality at Diagnosis (SEER)
420	Histology (92-00) ICD-O-2	Histology (COC)
440	Grade	Grade, Differentiation, or Cell Indicator (SEER) Grade/Differentiation (COC)

Item #	Item Name	Alternate Names
523	Behavior Code ICD-O-3	Behavior Code (COC)
540	Reporting Hospital	Institution ID Number (COC) Facility Identification Number (COC)
550	Accession NumberHosp	Accession Number (COC)
560	Sequence NumberHospital	Sequence Number (COC)
580	Date of 1st Contact	Date of Adm/1st Contact
590	Date of Inpatient Adm	Date of Inpatient Admission (COC)
600	Date of Inpatient Disch	Date of Inpatient Discharge (COC)
620	Year First Seen This CA	Accession Year (pre-96 COC) Year First Seen for this Primary (COC)
630	Primary Payer at DX	Primary Payer at Diagnosis (COC)
640	Inpatient/Outpt Status	Inpatient/Outpatient Status (COC)
650	Presentation at CA Conf	Presentation at Cancer Conference (COC)
660	Date of CA Conference	Date of Cancer Conference (COC)
670	RX HospSurg Prim Site	Cancer-Directed Surgery at this Facility (pre-96 COC) RX HospCA Dir Surgery (pre 96 NAACCR) Surgical Procedure of Primary Site
672	RX HospScope Reg LN Sur	Scope of Regional Lymph Node Surgery at this Facility (COC)
674	RX HospSurg Oth Reg/Dis	Surgery of Other Regional Site(s), Distant Site(s), or Distant Lymph Node(s) at this Facility (COC) Surgical Procedure/Other Site at this Facility
676	RX HospReg LN Removed	Number of Regional Lymph Nodes Examined at this Facility (COC) RX HospReg LN Examined
690	RX HospRadiation	Radiation at this Facility (COC)
700	RX HospChemo	Chemotherapy at this Facility (COC)
710	RX HospHormone	Hormone Therapy at this Facility (COC)
720	RX HospBRM	Immunotherapy at this Facility (COC)
730	RX HospOther	Other Treatment at this Facility (COC)
740	RX HospDX/Stg Proc	Non Cancer-Directed Surgery at this Facility (COC) Surgical Diagnostic & Staging Procedure at this Facility (1996-2002) RX HospDX/Stg/Pall Proc
742	RX HospScreen/BX Proc1	Diagnostic and Staging Procedures (pre-2001 COC) RX HospDiag/Stage Proc1 (pre-2001) Screening or Biopsy Procedures (COC)
743	RX HospScreen/BX Proc2	Diagnostic and Staging Procedures (pre-2001 COC) RX HospDiag/Stage Proc2 (pre-2001) Screening or Biopsy Procedures (COC)
744	RX HospScreen/BX Proc3	Diagnostic and Staging Procedures (pre-2001 COC) RX HospDiag/Stage Proc3 (pre-2001) Screening or Biopsy Procedures (COC)

Item #	Item Name	Alternate Names
745	RX HospScreen/BX Proc4	Diagnostic and Staging Procedures (pre-2001 COC) RX HospDiag/Stage Proc4 (pre-2001) Screening or Biopsy Procedures (COC)
746	RX HospSurg Site 98-02	Cancer-Directed Surgery at this Facility (pre-96 COC) RX HospCA Dir Surgery (pre-96 NAACCR) Surgical Procedure of Primary Site
747	RX HospScope Reg 98-02	Scope of Regional Lymph Node Surgery at this Facility (COC)
748	RX HospSurg Oth 98-02	Surgery of Other Regional Site(s), Distant Site(s), or Distant Lymph Node(s) at this Facility (COC) Surgical Procedure/Other Site at this Facility
760	SEER Summary Stage 1977	General Summary Stage (SEER/COC)
780	EODTumor Size	Size of Primary Tumor (SEER) Size of Tumor (COC)
790	EODExtension	Extension (pre-96 SEER/COC) Extension (SEER EOD) (96 COC)
810	EODLymph Node Involv	Lymph Nodes (pre-96 SEER/COC) Lymph Nodes (SEER EOD) (96 COC)
820	Regional Nodes Positive	Number of Positive Regional Lymph Nodes (SEER) Pathologic Review of Regional Lymph Nodes (SEER) Regional Lymph Nodes Positive
830	Regional Nodes Examined	Number of Regional Lymph Nodes Examined (SEER) Pathologic Review of Regional Lymph Nodes (SEER) Regional Lymph Nodes Examined
840	EODOld 13 Digit	13-Digit (Expanded) Site-Specific Extent of Disease (SEER) SEER EEOD (SEER)
850	EODOld 2 Digit	2-Digit Nonspecific and 2-Digit Site-Specific Extent of Disease (1973-1982 SEER)
860	EODOld 4 Digit	4-Digit Extent of Disease (1983-1987 SEER)
870	Coding System for EOD	Coding System for Extent of Disease (SEER)
880	TNM Path T	Pathologic T (COC)
890	TNM Path N	Pathologic N (COC)
900	TNM Path M	Pathologic M (COC)
910	TNM Path Stage Group	Pathologic Stage Group (COC)
920	TNM Path Descriptor	Pathologic Stage (Prefix/Suffix) Descriptor (COC)
930	TNM Path Staged By	Staged By (Pathologic Stage) (COC)
940	TNM Clin T	Clinical T (COC)
950	TNM Clin N	Clinical N (COC)
960	TNM Clin M	Clinical M (COC)
970	TNM Clin Stage Group	Clinical Stage Group (COC)
980	TNM Clin Descriptor	Clinical Stage (Prefix/Suffix) Descriptor (COC)
990	TNM Clin Staged By	Staged By (Clinical Stage) (COC)

Item #	Item Name	Alternate Names
1000	TNM Other T	Other T (COC)
1010	TNM Other N	Other N (COC)
1020	TNM Other M	Other M (COC)
1030	TNM Other Stage Group	Other Stage Group (COC)
1040	TNM Other Staged By	Staged By (Other Stage) (COC)
1050	TNM Other Descriptor	Other Stage (Prefix/Suffix) Descriptor (COC)
1080	Date of 1st Positive BX	Date of First Positive Biopsy (COC)
1090	Site of Distant Met 1	Site of Distant Metastasis #1 (COC)
1100	Site of Distant Met 2	Site of Distant Metastasis #2 (COC)
1110	Site of Distant Met 3	Site of Distant Metastasis #3 (COC)
1130	Pediatric Staging System	Type of Staging System (Pediatric) (COC)
1140	Pediatric Staged By	Staged By (Pediatric Stage) (COC)
1150	Tumor Marker 1	Tumor Marker One (COC)
1160	Tumor Marker 2	Tumor Marker Two (COC)
1170	Tumor Marker 3	Tumor Marker Three (COC)
1200	RX DateSurgery	Date of Cancer-Directed Surgery (COC) Date of Surgery Date of First Surgical Procedure (COC)
1210	RX DateRadiation	Date Radiation Started (COC)
1220	RX DateChemo	Date Chemotherapy Started (COC)
1230	RX DateHormone	Date Hormone Therapy Started (COC)
1240	RX DateBRM	Date Immunotherapy Started (COC)
1250	RX DateOther	Date Other Treatment Started (COC)
1260	Date of Initial RXSEER	Date Therapy Initiated (SEER) Date Started (SEER)
1270	Date of 1st Crs RXCOC	Date of First Course Treatment (COC) Date Started (pre-96 COC)
1280	RX DateDX/Stg Proc	Date of Non Cancer-Directed Surgery (COC) Date of Diagnostic, Staging or Palliative Procedures (1996-2002) Date of Surgical Diagnostic and Staging Procedure (COC) RX DateDX/Stg/Pall Proc
1290	RX SummSurg Prim Site	Cancer-Directed Surgery (pre 96 COC) Surgery of Primary Site (SEER/COC)
1292	RX SummScope Reg LN Sur	Scope of Regional Lymph Node Surgery (SEER/COC)

Item #	Item Name	Alternate Names
1294	RX SummSurg Oth Reg/Dis	Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Nodes (SEER/COC) Surgical Procedure/Other Site
1296	RX SummReg LN Examined	Number of Regional Lymph Nodes Examined (SEER/COC) Number of Regional Lymph Nodes Removed (COC)
1310	RX SummSurgical Approch	Surgical Approach (COC)
1320	RX SummSurgical Margins	Surgical Margins (COC) Residual Primary Tumor Following Cancer-Directed Surgery (pre-96 COC)
1330	RX SummReconstruct 1st	ReconstructionFirst Course (SEER) Reconstruction/RestorationFirst Course (COC)
1340	Reason for No Surgery	Reason for No Cancer-Directed Surgery (SEER) Reason for No CA Dir Surgery (COC) Reason for No Surgery of the Primary Site
1350	RX SummDX/Stg Proc	Non Cancer-Directed Surgery (COC) Surgical Diagnostic and Staging Procedure (1996-2002) RX SummDX/Stg/Pall Proc
1360	RX SummRadiation	Radiation (SEER/COC) Radiation Therapy (pre 96 COC)
1370	RX SummRad to CNS	Radiation Therapy to CNS (COC) Radiation to the Brain and/or Central Nervous System (SEER)
1380	RX SummSurg/Rad Seq	Radiation Sequence with Surgery (pre-96 SEER/COC) Radiation/Surgery Sequence (COC)
1390	RX SummChemo	Chemotherapy (SEER/COC)
1400	RX SummHormone	Hormone Therapy (SEER/COC) Endocrine (Hormone/Steroid) Therapy (pre-96 SEER)
1410	RX SummBRM	Immunotherapy (SEER/COC) Biological Response Modifiers (pre-96 SEER)
1420	RX SummOther	Other Treatment (COC) Other Cancer-Directed Therapy (SEER/pre-96 COC)
1430	Reason for No Radiation	Reason for No Regional Radiation Therapy
1440	Reason for No Chemo	Reason for No Chemotherapy (COC)
1450	Reason for No Hormone	Reason for No Hormone Therapy (COC)
1470	Protocol Eligibility Stat	Protocol Eligibility Status (COC)
1490	Referral to Support Serv	Referral to Support Services (COC)
1510	RadRegional Dose: CGY	Regional Dose: CGY (COC)
1520	RadNo of Treatment Vol	Number of Treatments to this Volume (COC)
1530	RadElapsed RX Days	Radiation Elapsed Treatment Time (Days) (COC)
1540	RadTreatment Volume	Radiation Treatment Volume (COC)
1550	RadLocation of RX	Location of Radiation Treatment (COC)
1560	RadIntent of Treatment	Intent of Treatment (Radiation) (COC)
1570	RadRegional RX Modality	Regional Treatment Modality (COC)

Item #	Item Name	Alternate Names
1580	RadRX Completion Status	Radiation Treatment Completion Status (COC)
1590	RadLocal Control Status	Radiation Therapy Local Control Status (Irradiated Volume) (COC)
1640	RX SummSurgery Type	SiteSpecific Surgery (pre-98 SEER)
1642	RX SummScreen/BX Proc1	Diagnostic and Staging Procedures (pre-2001 COC) RX SummDiag/Stage Proc1 (pre-2001) Screening or Biopsy Procedures (COC)
1643	RX SummScreen/BX Proc2	Diagnostic and Staging Procedures (pre-2001 COC) RX SummDiag/Stage Proc2 (pre-2001) Screening or Biopsy Procedures (COC)
1644	RX SummScreen/BX Proc3	Diagnostic and Staging Procedures (pre-2001 COC) RX SummDiag/Stage Proc3 (pre-2001) Screening or Biopsy Procedures (COC)
1645	RX SummScreen/BX Proc4	Diagnostic and Staging Procedures (pre-2001 COC) RX SummDiag/Stage Proc4 (pre-2001) Screening or Biopsy Procedures (COC)
1646	RX SummSurg Site 98-02	Cancer-Directed Surgery (pre-96 COC) Surgery of Primary Site (SEER/COC)
1647	RX SummScope Reg 98-02	Scope of Regional Lymph Node Surgery (SEER/COC)
1648	RX SummSurg Oth 98-02	Surgery of Other Regional Site(s), Distant Site(s) or Distant Lymph Nodes (SEER/COC) Surgical Procedure/Other Site
1660	Subsq RX 2nd Course Date	Second Course of Therapy-Date Started (pre-96 COC)
1741	Subsq RXReconstruct Del	Reconstruction/RestorationDelayed (COC)
1750	Date of Last Contact	Date of Last Contact or Death (COC) Date of Last Follow-Up or of Death (SEER)
1790	Follow-Up Source	Follow-Up Method (pre-96 COC)
1800	Next Follow-Up Source	Next Follow-Up Method (pre-96 COC)
1810	Addr CurrentCity	City/TownCurrent (COC)
1820	Addr CurrentState	StateCurrent (COC)
1830	Addr CurrentPostal Code	Postal CodeCurrent (COC)
1860	Recurrence Date1st	Date of First Recurrence (COC)
1880	Recurrence Type1st	Type of First Recurrence (COC)
1890	Recurrence Type1stOth	Other Type of First Recurrence (COC)
1910	Cause of Death	Underlying Cause of Death (SEER) Underlying Cause of Death (ICD Code) (pre-96 COC)
1920	ICD Revision Number	ICD Code Revision Used for Cause of Death (SEER)
1960	Site (73-91) ICD-O-1	Primary Site (1973-91) (SEER)
1980	ICD-O-2 Conversion Flag	Review Flag for 1973-91 Cases (SEER)
1981	Over-ride SS/NodesPos	Over-ride Summary Stage/Nodes Positive

Item #	Item Name	Alternate Names
1982	Over-ride SS/TNM-N	Over-ride Summary Stage/TNM-N
1983	Over-ride SS/TNM-M	Over-ride Summary Stage/TNM-M
1984	Over-ride SS/DisMet1	Over-ride Summary Stage/Distant Metastasis 1
1985	Over-ride Acsn/Class/Seq	Over-ride Accession/Class of Case/Sequence
1986	Over-ride HospSeq/DxConf	Over-ride Hospital Sequence/Diagnostic Confirmation
1988	Over-ride HospSeq/Site	Over-ride Hospital Sequence/Site
1990	Over-ride Age/Site/Morph	Age/Site/Histology Interfield Review (Interfield Edit 15) (SEER #3)
2000	Over-ride SeqNo/DxConf	Sequence Number/Diagnostic Confirmation Interfield Review (Interfield Edit 23) (SEER #4)
2010	Over-ride Site/Lat/SeqNo	Site/Histology/Laterality/Sequence Number Interrecord Review (Interrecord Edit 09) (SEER #5)
2020	Over-ride Surg/DxConf	Surgery/Diagnostic Confirmation Interfield Review (Interfield Edit 46) (SEER #6)
2030	Over-ride Site/Type	Site/Type Interfield Review (Interfield Edit 25) (SEER #1)
2040	Over-ride Histology	Histology/Behavior Interfield Review (Field Item Edit Morph) (SEER #2)
2050	Over-ride Report Source	Type of Reporting Source/Sequence Number Interfield Review (Interfield Edit 04) (SEER #7)
2060	Over-ride Ill-define Site	Sequence Number/III-defined Site Interfield Review (Interfield Edit 22) (SEER #8)
2070	Over-ride Leuk, Lymphoma	Leukemia or Lymphoma/Diagnostic Confirmation Interfield Review (Interfield Edit 48) (SEER #9)
2071	Over-ride Site/Behavior	Over-ride Flag for Site/Behavior (IF39) (SEER #11)
2072	Over-ride Site/EOD/DX Dt	Over-ride Flag for Site/EOD/Diagnosis Date (IF40) (SEER #13)
2073	Over-ride Site/Lat/EOD	Over-ride Flag for Site/Laterality/EOD (IF41) (SEER #12)
2074	Over-ride Site/Lat/Morph	Over-ride Flag for Site/Laterality/Morphology (IF42) (SEER #13)
2110	Date Case Report Exported	Date Case Transmitted (pre 98 NAACCR)
2140	COC Coding SysCurrent	Commission on Cancer Coding SystemCurrent (COC)
2180	SEER Type of Follow-Up	Type of Follow-Up (SEER)
2190	SEER Record Number	Record Number (SEER)
2200	Diagnostic Proc 73-87	Diagnostic Procedures (1973-87 SEER)
2230	NameLast	Last Name (COC)
2240	NameFirst	First Name (COC)
2250	NameMiddle	Middle Name (COC) Middle Initial (pre-96 COC)
2260	NamePrefix	Name Prefix (COC)

Item #	Item Name	Alternate Names
2270	NameSuffix	Name Suffix (COC)
2280	NameAlias	Alias (COC)
2310	Military Record No Suffix	Military Medical Record Number Suffix (COC)
2330	Addr at DXNo & Street	Patient Address (Number and Street) at Diagnosis (COC) Number and Street (pre-96 COC)
2335	Addr at DXSupplementl	Patient Address (Number and Street) at DiagnosisSupplemental (COC)
2350	Addr CurrentNo & Street	Patient Address (Number and Street)Current (COC)
2355	Addr CurrentSupplementl	Patient Address (Number and Street) CurrentSupplemental (COC)
2390	NameMaiden	Maiden Name (COC)
2410	Institution Referred From	Facility Referred From
2420	Institution Referred To	Facility Referred To
2460	PhysicianManaging	Managing Physician (COC) Attending Physician (pre-96 COC)
2470	PhysicianFollow-Up	Following Physician (COC) Follow-Up Physician (pre-96 COC)
2480	PhysicianPrimary Surg	Primary Surgeon (COC)
2490	Physician 3	Physician #3 (COC) Other Physician (pre-96 COC)
2500	Physician 4	Physician #4 (COC) Other Physician (pre-96 COC)
2820	CS Tumor Size/Ext Eval	CS Tumor Size/Extension Evaluation
2830	CS Lymph Nodes	CS Lymph Nodes (SEER EOD)
2840	CS Reg Nodes Eval	CS Regional Nodes Evaluation
2850	CS Mets at DX	CS Metastasis at Diagnosis
2860	CS Mets Eval	CS Metastasis Evaluation
2940	Derived AJCC T	Derived T
2950	Derived AJCC T Descriptor	Derived T Descriptor
2960	Derived AJCC N	Derived N
2970	Derived AJCC N Descriptor	Derived N Descriptor
2980	Derived AJCC M	Derived M
2990	Derived AJCC M Descriptor	Derived M Descriptor
3000	Derived AJCC Stage Group	Derived Stage Group
3010	Derived SS1977	Derived General Summary Stage (SEER) 1977

Item #	Item Name	Alternate Names
3020	Derived SS2000	Derived SEER Summary Stage 2000
3030	Derived AJCCFlag	AJCC Conversion Flag
3040	Derived SS1977Flag	SS 1977 Conversion Flag
3050	Derived SS2000Flag	SS 2000 Conversion Flag
3110	Comorbid/Complication 1	Comorbidities and Complications #1
3120	Comorbid/Complication 2	Comorbidities and Complications #2
3130	Comorbid/Complication 3	Comorbidities and Complications #3
3140	Comorbid/Complication 4	Comorbidities and Complications #4
3150	Comorbid/Complication 5	Comorbidities and Complications #5
3160	Comorbid/Complication 6	Comorbidities and Complications #6
3170	RX DateMost Defin Surg	Date of Most Definitive Surgical Resection of the Primary Site
3180	RX DateSurgical Disch	Date of Surgical Discharge
3190	Readm Same Hosp 30 Days	Readmission to the Same Hospital Within 30 Days of Surgical Discharge
3200	RadBoost RX Modality	Boost Radiation Treatment Modality
3210	RadBoost Dose cGy	Boost Radiation Dose cGy
3220	RX DateRadiation Ended	Date Radiation Ended
3230	RX DateSystemic	Date Systemic Therapy Started
3250	RX SummTransplnt/Endocr	Hematologic Transplant and Endocrine Procedures
3270	RX SummPalliative Proc	Palliative Procedure
3280	RX HospPalliative Proc	Palliative Procedure at this Facility
3300	RuralUrban Continuum 1993	Beale Code
3310	RuralUrban Continuum 2000	Beale Code

# **APPENDIX E**

## **GROUPED DATA ITEMS**

Item Name [Item#]	Length	Column #
Extent of Disease 10-Dig [779]	12	531-542
Subfields:	12	551 542
EODTumor Size[780]	3	531-533
EODExtension [790]	2	534-535
EODExtension Prost Path [800]	2	536-537
EODLymph Node Involv [810]	1	538-538
Regional Nodes Positive [820]	2	539-540
Regional Nodes Examined [830]	2	541-542
Morph (73-91) ICD-O-1 [1970]	6	1141-1146
Subfields:		
Histology (73-91) ICD-O-1 [1971]	4	1141-1144
Behavior (73-91) ICD-O-1 [1972]	1	1145-1145
Grade (73-91) ICD-O-1 [1973]	1	1146-1146
MorphType&Behav ICD-O-2 [419] Subfields:	5	296-300
Histology (92-00) ICD-O-2 [420]	4	296-299
Behavior (92-00) ICD-O-2 [430]	1	300-300
MorphType&Behav ICD-O-3 [521]	5	301-305
Subfields:	4	301-304
Histologic Type ICD-O-3 [522] Behavior Type ICD-O-3 [523]	4	301-304 305-305
Benavior Type ICD-O-5 [525]	1	303-303
Subsq RX 2nd Course Codes [1670]	7	996-1002
Subsq RX 2nd Course Surg [1671]	2	996-997
Subsq RX 2nd Course Rad [1672]	1	998-998
Subsq RX 2nd Course Chemo [1673]	1	999-999
Subsq RX 2nd Course Horm [1674]	1	1000-1000
Subsq RX 2nd Course BRM [1675]	1	1001-1001
Subsq RX 2nd Course Oth [1676]	1	1002-1002
Subsq RX 3rd Course Codes [1690]	7	1011-1017
Subsq RX 3rd Course Surg [1691]	2	1011-1012
Subsq RX 3rd Course Rad [1692]	1	1013-1013
Subsq RX 3rd Course Chemo [1693]	1	1014-1014
Subsq RX 3rd Course Horm [1694]	1	1015-1015
Subsq RX 3rd Course BRM [1695]	1	1016-1016
Subsq RX 3rd Course Oth [1696]	1	1017-1017

Item Name [Item#]	Length	Column #
Subsq RX 4th Course Codes [1710]	7	1026-1032
Subsq RX 4th Course Surg [1711]	2	1026-1027
Subsq RX 4th Course Rad [1712]	1	1028-1028
Subsq RX 4th Course Chemo [1713]	1	1029-1029
Subsq RX 4th Course Horm [1714]	1	1030-1030
Subsq RX 4th Course BRM [1715]	1	1031-1031
Subsq RX 4th Course Oth [1716]	1	1032-1032
Subsq RX 5th Course Codes [1730]	7	1041-1047
Subsq RX 5th Course Surg [1731]	2	1041-1042
Subsq RX 5th Course Rad [1732]	1	1043-1043
Subsq RX 5th Course Chemo [1733]	1	1044-1044
Subsq RX 5th Course Horm [1734]	1	1045-1045
Subsq RX 5th Course BRM [1735]	1	1046-1046
Subsq RX 5th Course Oth [1736]	1	1047-1047

# **APPENDIX F**

# TABLES AND DATA DICTIONARY REVISIONS

Item #	Item Name	Record Layout Note	Required Status Note	Data Descriptor Note	Data Dictionary Note
50	NAACCR Record Version				Revised
70	Addr at DXCity			Revised	
80	Addr at DXState				Revised
100	Addr at DXPostal Code				Revised
190	Spanish/Hispanic Origin			Revised	Revised
191	NHIA Derived Hisp Origin	New	New	New	New
200	Computed Ethnicity				Revised
210	Computed Ethnicity Source				Revised
250	Birthplace			Revised	
380	Sequence NumberCentral				Revised
430	Behavior (92-00) ICD-O-2			Revised	
523	Behavior Code ICD-O-3			Revised	
560	Sequence NumberHospital				Revised
580	Date of 1st Contact				Revised
590	Date of Inpatient Adm				Revised
600	Date of Inpatient Disch				Revised
690	RX HospRadiation				Revised
820	Regional Nodes Positive				Revised
830	Regional Nodes Examined				Revised
1220	RX DateChemo				Revised
1230	RX DateHormone				Revised
1240	RX DateBRM				Revised
1330	RX SummReconstruct 1st				Revised
1390	RX SummChemo			Revised	Revised
1400	RX SummHormone			Revised	Revised
1660	Subsq RX 2nd Course Date				Revised
1671	Subsq RX 2nd Course Surg				Revised
1672	Subsq RX 2nd Course Rad				Revised
1673	Subsq RX 2nd Course Chemo				Revised
1674	Subsq RX 2nd Course Horm				Revised
1675	Subsq RX 2nd Course BRM				Revised
1676	Subsq RX 2nd Course Oth				Revised
1677	Subsq RX 2ndScope LN SU				Revised
1678	Subsq RX 2ndSurg Oth				Revised
1679	Subsq RX 2ndReg LN Rem				Revised
1680	Subsq RX 3rd Course Date				Revised
1691	Subsq RX 3rd Course Surg				Revised

Item #	Item Name	Record Layout Note	Required Status Note	Data Descriptor Note	Data Dictionary Note
1692	Subsq RX 3rd Course Rad				Revised
1693	Subsq RX 3rd Course Chemo				Revised
1694	Subsq RX 3rd Course Horm				Revised
1695	Subsq RX 3rd Course BRM				Revised
1696	Subsq RX 3rd Course Oth				Revised
1697	Subsq RX 3rdScope LN SU				Revised
1698	Subsq RX 3rdSurg Oth				Revised
1699	Subsq RX 3rdReg LN Rem				Revised
1700	Subsq RX 4th Course Date				Revised
1711	Subsq RX 4th Course Surg				Revised
1712	Subsq RX 4th Course Rad				Revised
1713	Subsq RX 4th Course Chemo				Revised
1714	Subsq RX 4th Course Horm				Revised
1715	Subsq RX 4th Course BRM				Revised
1716	Subsq RX 4th Course Oth				Revised
1717	Subsq RX 4thScope LN SU				Revised
1718	Subsq RX 4thSurg Oth				Revised
1719	Subsq RX 4thReg LN Rem				Revised
1741	Subsq RXReconstruct Del				Revised
1820	Addr CurrentState				Revised
1830	Addr CurrentPostal Code				Revised
1840	CountyCurrent				Revised
1842	Follow-Up ContactCity				Revised
1844	Follow-Up ContactState				Revised
1846	Follow-Up ContactPostal				Revised
1871	Recurrence Distant Site 1				Revised
1872	Recurrence Distant Site 2				Revised
1873	Recurrence Distant Site 3				Revised
1910	Cause of Death				Revised
1920	ICD Revision Number				Revised
1930	Autopsy				Revised
1940	Place of Death				Revised
1981	Over-ride SS/NodesPos				Revised
1982	Over-ride SS/TNM-N				Revised
1983	Over-ride SS/TNM-M				Revised
1984	Over-ride SS/DisMet1				Revised
1985	Over-ride Acsn/Class/Seq				Revised
1986	Over-ride HospSeq/DxConf				Revised
1987	Over-ride COC-Site/Type				Revised
1988	Over-ride HospSeq/Site				Revised
1989	Over-ride Site/TNM-StgGrp				Revised
1909	Over-ride Age/Site/Morph				Revised

Item #	Item Name	Record Layout Note	Required Status Note	Data Descriptor Note	Data Dictionary Note
2000	Over-ride SeqNo/DxConf				Revised
2010	Over-ride Site/Lat/SeqNo				Revised
2020	Over-ride Surg/DxConf				Revised
2030	Over-ride Site/Type				Revised
2040	Over-ride Histology				Revised
2050	Over-ride Report Source				Revised
2060	Over-ride Ill-define Site				Revised
2070	Over-ride Leuk, Lymphoma				Revised
2071	Over-ride Site/Behavior				Revised
2072	Over-ride Site/EOD/DX Dt				Revised
2073	Over-ride Site/Lat/EOD				Revised
2074	Over-ride Site/Lat/Morph				Revised
2090	Date Case Completed				Revised
2100	Date Case Last Changed				Revised
2110	Date Case Report Exported				Revised
2111	Date Case Report Received				Revised
2112	Date Case Report Loaded				Revised
2113	Date Tumor Record Availbl				Revised
2150	COC Coding SysOriginal				Revised
2260	NamePrefix				Revised
2270	NameSuffix				Revised
2280	NameAlias				Revised
2290	NameSpouse/Parent				Revised
2300	Medical Record Number				Revised
2390	NameMaiden				Revised
2392	Follow-Up ContactNo&St				Revised
2393	Follow-Up ContactSuppl				Revised
2394	Follow-Up ContactName				Revised
2410	Institution Referred From				Revised
2420	Institution Referred To				Revised
2440	Following Registry				Revised
2460	PhysicianManaging				Revised
2490	Physician 3				Revised
2500	Physician 4				Revised
2520	TextDX ProcPE				Revised
2530	TextDX ProcX-ray/Scan				Revised
2540	TextDX ProcScopes				Revised
2550	TextDX ProcLab Tests				Revised
2560	TextDX ProcOp				Revised
2570	TextDX ProcPath				Revised
2580	TextPrimary Site Title				Revised
2590	TextHistology Title				Revised

Item #	Item Name	Record Layout Note	Required Status Note	Data Descriptor Note	Data Dictionary Note
2600	TextStaging				Revised
2610	RX TextSurgery				Revised
2620	RX TextRadiation (Beam)				Revised
2630	RX TextRadiation Other				Revised
2640	RX TextChemo				Revised
2650	RX TextHormone				Revised
2660	RX TextBRM				Revised
2670	RX TextOther				Revised
2680	TextRemarks				Revised
2690	Place of Diagnosis				Revised
2820	CS Tumor Size/Ext Eval			Revised	
2840	CS Reg Node Eval			Revised	
2860	CS Mets Eval			Revised	
2940	Derived AJCC T				Revised
2950	Derived AJCC T Descriptor			Revised	Revised
2960	Derived AJCC N				Revised
2970	Derived AJCC N Descriptor			Revised	Revised
2980	Derived AJCC M				Revised
2990	Derived AJCC M Descriptor			Revised	Revised
3000	Derived AJCC Stage Group				Revised
3030	Derived AJCCFlag				Revised
3040	Derived SS1977Flag				Revised
3050	Derived SS2000Flag				Revised
3110	Comorbid/Complication 1			Revised	Revised
3120	Comorbid/Complication 2			Revised	Revised
3130	Comorbid/Complication 3			Revised	Revised
3140	Comorbid/Complication 4			Revised	Revised
3150	Comorbid/Complication 5			Revised	Revised
3160	Comorbid/Complication 6			Revised	Revised

### APPENDIX G

### **RECOMMENDED ABBREVIATIONS FOR ABSTRACTORS**

The use of abbreviations in cancer abstraction is becoming more commonplace as the demands on abstractors increase. Abbreviations often are used by cancer abstractors to shorten the written narratives entered into text fields to facilitate the electronic storage and transmission of the information. However, abbreviations can generate confusion, because abbreviations may vary among different institutions and even between different specialties within the same institution. To be useful, an abbreviation must be clearly understood by any individual who encounters it. Consequently, the use of abbreviations is a useful abstracting practice only if universally recognized and understood abbreviations are used.

The NAACCR Recommended Abbreviations Listings were developed for utilization by cancer report abstractors and the agencies to which they submit their data. These lists were compiled to reduce some of the confusion that can result from the use of common and not-so-common abbreviations when abstracting reports of cancer from the medical record. Although the lists may shed some light on abbreviations used in the medical record, please note that these lists are intended to be used as a primary reference by the cancer abstractor, to help abstract necessary information into a limited number of text fields for storage and transmission of cancer information.

The NAACCR Recommended Abbreviations Listings consist of two main lists of almost 500 word/terms and their recommended abbreviations/symbols, as well as a special table delineating context-sensitive abbreviations. 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/symbol to enable the look-up of the word or term for a particular abbreviation or symbol. 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 were compiled from abbreviation lists from SEER Book 3, the NAACCR Pathology Committee, the Veterans Administration, Dr. Jay Piccirillo's comorbid conditions training materials, the Florida Cancer Data System, and the California Cancer Registry. Terms included in the lists are limited to those that are commonly utilized when abstracting cancer information. 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. 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. Any questions or suggestions for new/modified abbreviations may be e-mailed to either of the current Chairpersons of the NAACCR Registry Operations Committee: Susan Gershman, susan.gershman@state.ma.us or Susan Bolick-Aldrich, bolicks@dhec.sc.gov.

WORD/TERM(S)	ABBREVIATION/SYMBOL
Abdomen (abdominal)	ABD
Abdominal perineal	AP
Abnormal	ABN
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
Activities of daily living	ADL
Acute granulocytic leukemia	AGL
Acute lymphocytic leukemia	ALL
Acute myelogenous leukemia	AML
Acute myocardial infarction	AMI
Acute Respiratory Distress (Disease) Syndrome	ARDS
Acute tubular necrosis	ATN
Acute renal failure	ARF
Adenocarcinoma	ADENOCA
Adenosine triphosphate	ATP
Adjacent	ADJ
Adult-onset Diabetes Mellitus	AODM
Admission/Admit	ADM
Adrenal cortical hormone	АСН
Adrenal cortex	AC
Adrenocorticotrophic hormone	АСТН
Affirmative	AFF
Against medical advice	АМА
AIDS-related condition (complex)	ARC
AIDS-related disease	ARD
Air contrast barium enema	ACBE
Albumin	ALB
Alcohol	ETOH
Alkaline phosphatase	ALK PHOS
Alpha-fetoprotein	AFP
Also known as	AKA
Ambulatory	AMB
Amount	AMD
Amputation	AMP
Amyotrophic lateral sclerosis	ALS
Anyotrophic lateral scienciss Anal intraepithelial neoplasia, grade III	ALS AIN III

### NAACCR RECOMMENDED ABBREVIATION LIST ORDERED BY WORD/TERM(S)

WORD/TERM(S)	ABBREVIATION/SYMBOL
Anaplastic	ANAP
And	&
Angiography/Angiogram	ANGIO
Anterior	ANT
Anteroposterior	AP
Antidiuretic hormone	ADH
Antigen	AG
Aortic stenosis	A-STEN
Appendix	APP
Apparently	APPL'Y
Approximately	APPROX
Arrhythmia	ARRHY
Arterial blood gases	ABG
Arteriosclerotic cardiovascular disease	ASCVD
Arteriosclerotic heart disease	ASHD
Arteriosclerotic Peripheral Vascular Disease	ASPVD
Arteriosclerosis/Arteriosclerotic	AS
Arteriovenous	AV
Arteriovenous malformation	AVM
Artery (ial)	ART
Ascending colon	A-COLON
Aspiration	ASP
Aspirin, Acetylsalicylic acid	ASA
As soon as possible	ASAP
At	(a)
Atrial fibrillation	A FIB
Atrial flutter	A FLUTTER
Atrial stenosis/insufficiency/incompetence	AI
Atrial premature complexes	APC
Auscultation & percussion	A&P
Autonomic nervous system	ANS
Autopsy	AUT
Autoimmune hemolytic anemia	AIHA
Average	AVG
Axilla(ry)	AX
Bacillus Calmette-Guerin	BCG
Barium	BA
Barium enema	BE
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

WORD/TERM(S)	ABBREVIATION/SYMBOL
Bilateral salpingo-oophorectomy	BSO
Bile duct	BD
Biological response modifier	BRM
Biopsy	BX
Bipolar affective disorder	BAD
Black female	B/F
Black male	B/M
Bladder tumor	BT
Blood pressure	BP
Blood urea nitrogen	BUN
Blood volume	BV
Bone marrow	BM
Bone marrow transplant	BMT
Bowel movement	BM
Brother	BRO
Calcium	СА
Capsule (s)	CAP(S)
Carcinoembryonic antigen	CEA
Carcinoma	СА
Carcinoma in situ	CIS
Cardiovascular disease	CVD
CAT/CT scan/Computerized axial tomography	СТ
Centimeter	СМ
Central nervous system	CNS
Cerebrospinal fluid	CSF
Cerebrovascular accident	CVA
Cervical intraepithelial neoplasia	CIN
Cervical intraepithelial neoplasia, grade III	CIN III
Cervical vertebrae	C1-C7
Cervical spine	C-SPINE
Change	CHG
Chemotherapy	СНЕМО
Chest X-ray	CXR
Chronic	CHR
Chronic granulocytic leukemia	CGL
Chronic lymphocytic leukemia	CLL
Chronic myeloid (myelocytic) leukemia	CML
Chronic obstructive lung disease	COLD
Chronic obstructive pulmonary disease	COPD
Chronic renal failure	CRF
Chronic ulcerative colitis	CUC
Cigarettes	CIG
Clear	CLR
Cobalt 60	CO60

WORD/TERM(S)	ABBREVIATION/SYMBOL
Collaborative stage	CS
Colon, Ascending	A-COLON
Colon, Descending	D-COLON
Colon, Sigmoid	SIG COLON
Colon, Transverse	TRANS-COLON
Colony-stimulating factor	C-SF
Complaint (-ning) of	С/О
Complete blood count	CBC
Congenital heart disease	СНД
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
Cystoscopy	CYSTO
Cytology	СҮТО
Cystic fibrosis	CF
Date of birth	DOB
Date of death	DOD
Dead on arrival	DOA
Decrease(d)	DECR
Deep tendon reflex	DTR
Deep vein thrombosis	DVT
Deoxyribonucleic acid	DNA
Descending colon	D-COLON
Dermatology	DERM
Diabetes mellitus	DM
Diagnosis	DX
Diagnosis	DIAM
Diethylstilbestrol	DES
Differentiated/differential	DES
Digital rectal examination	DRE
Dilatation and curettage	D&C
Discharge	DISCH
Discontinue(d)	DC DZ
Disease	DZ
Disseminated intravascular coagulopathy	DIC
Ductal carcinoma <i>in situ</i>	DCIS
Dyspnea on exertion	DOE
Ears, nose, and throat	ENT

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

WORD/TERM(S)	ABBREVIATION/SYMBOL
Electrocardiogram	ECG/EKG
Electroencephalogram	EEG
Electromyogram	EMG
Emergency room	ER
Endoscopic retrograde cholangiopancreatography	ERCP
End stage renal disease	ESRD
Enlarged	ENLGD
Equal(s)	=
Esophagogastro-duodenoscopy	EGD
Estrogen receptor (assay)	ER, ERA
Evaluation	EVAL
Every	Q
Every day	QD
Examination	EXAM
Excision/excised	EXC(D)
Expired	EXP
Exploratory	EXPL
Exploratory laparotomy	EXPL LAP
Extend/extension	EXT
Fever of unknown origin	FUO
Fine needle aspiration	FNA
Fine needle aspiration biopsy	FNAB
Floor of mouth	FOM
Fluid	FL
Fluoroscopy	FLURO
Follow-up	FU
For example	E.G.
Fracture	FX
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
Greater/Greater than	>
Gynecology	GYN
Hematocrit	НСТ
Hemoglobin	HGB

WORD/TERM(S)	ABBREVIATION/SYMBOL
Hepatitis A (virus)	HAV
Hepatitis B (virus)	HBV
Hepatitis C (virus)	HCV
Hepatitis D (virus)	HDV
Hepatosplenomegaly	HSM
History	НХ
History and physical	Н&Р
History of	Н/О
Hormone	HORM
Hospital	HOSP
Hour/Hours	HR(S)
Human chorionic gonadotropin	HCG
Human Immunodeficiency Virus	HIV
Human Papilloma Virus	HPV
Human T-Lymphotrophic 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
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
Interstitial lung disease	ILD
Intramuscular	IM
Intrathecal	IT
Intravenous	IV
Intravenous cholangiogram	IVCA
Intravenous pyelogram	IVP

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

WORD/TERM(S)	ABBREVIATION/SYMBOL	
Invade(s)/invading/invasion	INV	
Involve(s)/involvement/involving	INVL	
Ipsilateral	IPSI	
Irregular	IRREG	
Jugular venous distention	JVD	
Juvenile rheumatic arthritis	JRA	
Kaposi sarcoma	KS	
Kidneys, ureters, bladder	KUB	
Kilogram	KG	
Kilovolt	KV	
laboratory	LAB	
Lactic dehydrogenase	LDH	
Laparotomy	LAP	
Large	LRG	
Last menstrual period	LMP	
Lateral	LAT	
Left	LT	
Left bundle branch block	LBBB	
Left costal margin	LCM	
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 quadrant	LUQ	
Left upper outer quadrant	LUOQ	
Less/Less than	<	
Licensed practical nurse	LPN	
Linear accelerator	LINAC	
Liver/spleen scan	LS SCAN	
Lower extremity	LE	
Lower inner quadrant	LIQ	
Lower outer quadrant	LOQ	
Lumbar vertebra	L1-L5	
Lumbar spine	L-SPINE	
Lumbosacral	LS	
Lymphadenopathy-associated virus	LAV	
Lymph node(s)	LN(S)	
Lymph node dissection	LND	
Lupus erythematosus	LUP ERYTH	

WORD/TERM(S)	ABBREVIATION/SYMBOL	
Macrophage colony-stimulating factor	M-CSF	
Magnetic resonance imaging	MRI	
Magnetic resonance cholangiopancreatography	MRCP	
Main stem bronchus	MSB	
Malignant	MALIG	
Mandible/mandibular	MAND	
Maximum	MAX	
Medical center	MC	
Medication	MED	
Metastatic/Metastasis	METS	
Methicillin Resistant Staphylococcus Aureus	MRSA	
Microgram	MCG	
Microscopic	MICRO	
Middle lobe	ML	
Millicurie (hours)	MC(H)	
Milligram (hours)	MG(H)	
Milliliter	ML	
Millimeter	ММ	
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	
More/More than	>	
Multifocal arterial tachycardia	MAT	
Multifocal premature ventricular contraction	MPVC	
Multiple	MULT	
Multiple sclerosis	MS	
Multiple myeloma	MM	
Myasthenia gravis	MG	
Myocardial infarction	MI	
Neck vein distention	NVD	
Negative	NEG	
Negative	-	
Neoplasm	NEOPL	
Neurology	NEURO	
No evidence of disease	NED	
No significant findings	NSF	
Non-Hodgkins lymphoma	NHL	

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

WORD/TERM(S)	ABBREVIATION/SYMBOL
Normal	NL
Non small cell carcinoma	NSCCA
Not applicable	NA
Not otherwise specified	NOS
Not recorded	NR
Number	#
Nursing home	NH
Obstetrics	OB
Obstructed (-ing, -ion)	OBST
Operating room	OR
Operative report	OP RPT
Organic brain syndrome	OBS
Orthopedics	ORTHO
Otology	ОТО
Ounce	OZ
Outpatient	OP
Packs per day	PPD
Palpated (-able)	PALP
Papanicolaou smear	PAP
Papillary	РАР
Past/personal (medical) history	РМН
Pathology	РАТН
Patient	РТ
Pediatrics	PEDS
Pelvic inflammatory disease	PID
Peptic ulcer disease	PUD
Percutaneous	PERC
Percutaneous transhepatic cholecystogram	РТС
Peripheral vascular disease	PVD
Prescription	RX
Primary medical physician	РМР
Phosphorus 32	P32
Physical examination	PE
Physiotherapy/Physical therapy	РТ
Platelets	PLT
Plus	+
Poorly differentiated	PD, POOR DIFF
Positive	POS
Positive	+
Positron emission tomography	PET
Possible	POSS
Posterior	POST
Postoperative (-ly)	POST OP

WORD/TERM(S)	ABBREVIATION/SYMBOL	
Pound(s)	LB(S)	
Pound(s)	#	
Premature atrial contraction	PAC	
Preoperative (-ly)	PRE OP	
Previous	PREV	
Prior to admission	РТА	
Probable (-ly)	PROB	
Proctoscopy	PROCTO	
Progesterone receptor (assay)	PR, PRA	
Prostatic intraepithelial neoplasia, grade III	PIN III	
Prostatic specific antigen	PSA	
Pulmonary	PULM	
Quadrant	QUAD	
Dediction should do a	RAD	
Radiation absorbed dose	RT RT	
Radiation therapy		
Radioimmunoassay Received	RIA REC'D	
Red blood cells (count)	RBC	
Regarding	RE	
Regional medical center Regular	RMC REG	
Regular sinus rhythm	RSR	
Regular sinus mythm Resection (ed)	RESEC	
Review of outside films	ROF	
Review of outside slides	ROS	
Review of outside sindes	RA	
Rheumatic heart disease	RHD	
	RT	
Right Right bundle branch block	RBBB	
	RCM	
Right costal margin Right inner quadrant		
Right lower extremity	RIQ RLE	
Right lower lobe	RLL	
Right lower quadrant	RLQ	
Right niddle lobe	RML	
Right outer quadrant		
	ROQ	
Right salpingo-oophorectomy	RSO	
Right upper extremity	RUE	
Right upper lobe Right upper quadrant	RUL	
Right upper quadrant Right optimized and Right	RUQ R/O	
Sacral spine	S-SPINE	

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

WORD/TERM(S)	ABBREVIATION/SYMBOL	
Sacral vertebra	S1-S5	
Salpingo-oophorectomy	SO	
Satisfactory	SATIS	
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	
Small	SM	
Small bowel	SB	
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	
Thoracic spine	T-SPINE	
Thromboticthrombocytopenia purpura	ТТР	
Times	Х	
Total abdominal hysterectomy	ТАН	
Total abdominal hysterectomy- bilateral salpingo-oophorectomy	TAH-BSO	
Total vaginal hysterectomy	ТVН	
Transient ischemic attack	TIA	
Transitional cell carcinoma	ТСС	
Transurethral resection	TUR	
Transurethral resection bladder	TURB	
Transurethral resection prostate	TURP	
Transverse colon	TRANS-COLON	
Treatment	TX	
True vocal cord	TVC	
Tuberculosis	ТВ	
Twice a day (daily)	BID	

WORD/TERM(S)	ABBREVIATION/SYMBOL	
Ultrasound	US	
Undifferentiated	UNDIFF	
Unknown	UNK	
Upper extremity	UE	
Upper gastrointestinal (series)	UGI	
Upper inner quadrant	UIQ	
Upper outer quadrant	UOQ	
Upper respiratory infection	URI	
Urinary tract infection	UTI	
Vagina/Vaginal	VAG	
Vaginal hysterectomy	VAG HYST	
Vaginal intraepithelial neoplasia (grade III)	VAIN III	
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	
With	W/	
Within normal limits	WNL	
Without	W/O	
Wolff-Parkinson-White syndrome	WPW	
Work-up	W/U	
Xray	XR	
Year	YR	

### NAACCR RECOMMENDED ABBREVIATION LIST ORDERED BY ABBREVIATION/SYMBOL

ABBREVIATION/SYMBOL	WORD/TERM(S)
^	above
@	at
&	and
<	less, less than
=	equals
>	greater than, more, more than
-	negative, minus
#	number, pound(s)
+	plus, positive
Х	times
A-COLON	Ascending colon
A FIB	Atrial fibrillation
A FLUTTER	Atrial flutter
A-STEN	Aortic stenosis
A&P	Auscultation & percussion
ABD	Abdomen (abdominal)
ABG	Arterial blood gases
ABN	Abnormal
ABS	Absent/Absence
ABST	Abstract/Abstracted
AC	Adrenal cortex
ACBE	Air contrast barium enema
АСН	Adrenal cortical hormone
ACID PHOS	Acid phosphatase
АСТН	Adrenocorticotrophic hormone
ADENOCA	Adenocarcinoma
ADH	Antidiuretic hormone
ADJ	Adjacent
ADL	Activities of daily living
ADM	Admission/Admit
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
AIN III	Anal intraepithelial neoplasia, grade III
AK(A)	Above knee (amputation)
AKA	Also known as
ALB	Albumin

ABBREVIATION/SYMBOL	WORD/TERM(S)
ALK PHOS	Alkaline phosphatase
ALL	Acute lymphocytic leukemia
ALS	Amyotrophic lateral sclerosis
AM	Before noon
AMA	Against medical advice
AMB	Ambulatory
AMI	Acute myocardial infarction
AML	Acute myelogenous leukemia
AMP	Amputation
AMT	Amount
ANAP	Anaplastic
ANGIO	Angiography/Angiogram
ANS	Autonomic nervous system
ANT	Anterior
AODM	Adult-onset Diabetes Mellitus
AP	Abdominal perineal
AP	Anteroposterior
APC	Atrial premature complexes
APP	Appendix
APPL'Y	Apparently
APPROX	Approximately
ARC	AIDS-related condition (complex)
ARD	AIDS-related disease
ARDS	Acute Respiratory Distress (Disease) Syndrome
ARF	Acute renal failure
ARRHY	Arrhythmia
ART	Artery (ial)
AS	Arteriosclerosis/Arteriosclerotic
ASA	Aspirin, Acetylsalicylic acid
ASAP	As soon as possible
ASCVD	Arteriosclerotic cardiovascular disease
ASHD	Arteriosclerotic heart disease
ASP	Aspiration
ASPVD	Arteriosclerotic Peripheral Vascular Disease
ATN	Acute tubular necrosis
ATP	Adenosine triphosphate
ATR	Achilles tendon reflex
AUT	Autopsy
AV	Arteriovenous
AVG	Average
AVM	Arteriovenous malformation
AX	Axilla(ry)
B/F	Black female
B/M	Black male

ABBREVIATION/SYMBOL	WORD/TERM(S)
BA	Barium
BAD	Bipolar affective disorder
BCC	Basal cell carcinoma
BCG	Bacillus Calmette-Guerin
BD	Bile duct
BE	Barium enema
BID	Twice a day (daily)
BIL	Bilateral
BK(A)	Below knee (amputation)
BM	Bone marrow
BM	Bowel movement
BMT	Bone marrow transplant
BP	Blood pressure
BPH	Benign prostatic hypertrophy/hyperplasia
BRM	Biological response modifier
BRO	Brother
BSO	Bilateral salpingo-oophorectomy
BT	Bladder tumor
BUN	Blood urea nitrogen
BUS	Bartholin's, Urethral & Skene's
BV	Blood volume
BX	Biopsy
С/О	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
CC	Cubic centimeter
CCU	Coronary care unit
CEA	Carcinoembryonic antigen
CF	Cystic fibrosis
CGL	Chronic granulocytic leukemia
CHD	Congenital heart disease
СНЕМО	Chemotherapy
CHF	Congestive heart failure
CHG	Change
CHR	Chronic
CIG	Cigarettes
CIN	Cervical intraepithelial neoplasia
CIN III	Cervical intraepithelial neoplasia Cervical intraepithelial neoplasia, grade III

ABBREVIATION/SYMBOL	WORD/TERM(S)
CIS	Carcinoma in situ
CLL	Chronic lymphocytic leukemia
CLR	Clear
СМ	Centimeter
CML	Chronic myeloid (myelocytic) leukemia
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
СТ	CAT/CT scan/Computerized axial tomography
CUC	Chronic ulcerative colitis
CVA	Cerebrovascular accident
CVD	Cardiovascular disease
CXR	Chest X-ray
CYSTO	Cystoscopy
СҮТО	Cytology
D-COLON	Descending colon
D&C	Dilatation and curettage
DC	Discontinue(d)
DCIS	Ductal carcinoma in situ
DECR	Decrease(d)
DERM	Dermatology
DES	Diethylstilbestrol
DIAM	Diameter
DIC	Disseminated intravascular coagulopathy
DIFF	Differentiated/differential
DISCH	Discharge
DM	Diabetes mellitus
DNA	Deoxyribonucleic acid
DOA	Dead on arrival
DOB	Date of birth
DOD	Date of death
DOE	Dyspnea on exertion
DRE	Digital rectal examination
DTR	Deep tendon reflex
DVT	Deep vein thrombosis
DX	Diagnosis

ABBREVIATION/SYMBOL	WORD/TERM(S)
DZ	Disease
E.G.	For example
ECG/EKG	Electrocardiogram
EEG	Electroencephalogram
EGD	Esophagogastro-duodenoscopy
EMG	Electromyogram
ENLGD	Enlarged
ENT	Ears, nose, and throat
ER	Emergency room
ER, ERA	Estrogen receptor (assay)
ERCP	Endoscopic retrograde cholangiopancreatography
ESRD	End stage renal disease
ЕТОН	Alcohol
EVAL	Evaluation
EXAM	Examination
EXC(D)	Excision/excised
EXP	Expired
EXPL	Exploratory
EXPL LAP	Exploratory laparotomy
EXT	Extend/extension
FL	Fluid
FLURO	Fluoroscopy
FNA	Fine needle aspiration
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
GERD	Gastrointestinal
GR	Grade
GU	Genitourinary
GU GYN	Gynecology
0110	Gynecology
H&P	History and physical
11001	History and physical

ABBREVIATION/SYMBOL	WORD/TERM(S)
HAV	Hepatitis A (virus)
HBV	Hepatitis B (virus)
HCG	Human chorionic gonadotropin
НСТ	Hematocrit
HCV	Hepatitis C (virus)
HCVD	Hypertensive cardiovascular disease
HDV	Hepatitis D (virus)
HGB	Hemoglobin
HIV	Human Immunodeficiency Virus
HORM	Hormone
HOSP	Hospital
HPV	Human Papilloma Virus
HR(S)	Hour/Hours
HSM	Hepatosplenomegaly
HTLV	Human T-Lymphotrophic Virus, (Type III)
HTN	Hypertension
HVD	Hypertensive vascular disease
HX	History
HYST	Hysterectomy
I&D	Incision & drainage
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
IMP	Impression
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

ABBREVIATION/SYMBOL	WORD/TERM(S)
IV	Intravenous
IVC	Inferior vena cava
IVCA	Intravenous cholangiogram
IVP	Intravenous pyelogram
JRA	Juvenile rheumatic arthritis
JVD	Jugular venous distention
J V D	
KG	Kilogram
KS	Kaposi sarcoma
KUB	Kidneys, ureters, bladder
KV	Kilovolt
L-SPINE	Lumbar spine
L1-L5	Lumbar vertebra
LAB	laboratory
LAP	Laparotomy
LAT	Lateral
LAV	Lymphadenopathy-associated virus
LB	Pound
LBBB	Left bundle branch block
LCM	Left costal margin
LDH	Lactic dehydrogenase
LE	Lower extremity
LINAC	Linear accelerator
LIQ	Lower inner quadrant
LLE	Left lower extremity
LLL	Left lower lobe
LLQ	Left lower quadrant
LMP	Last menstrual period
LN(S)	Lymph node(s)
LND	Lymph node dissection
LOQ LPN	Lower outer quadrant Licensed practical nurse
LRG	Large
LS	Lunge
LS SCAN	Liver/spleen scan
LSO	Left salpingo-oophorectomy
LT	Left
LUE	Left upper extremity
LUL	Left upper lobe
LUOQ	Left upper outer quadrant
LUP ERYTH	Lupus erythematosus
LUQ	Left upper quadrant

ABBREVIATION/SYMBOL	WORD/TERM(S)
M-CSF	Macrophage colony-stimulating factor
MALIG	Malignant
MAND	Mandible/mandibular
MAT	Multifocal arterial tachycardia
MAX	Maximum
MC	Medical center
MC(H)	Millicurie (hours)
MCG	Microgram
MCID	Mixed combined immunodeficiency
MCTD	Mixed connective tissue disease
MD	Moderately differentiated
MED	Medication
METS	Metastatic/Metastasis
MEV	Million electron volts
MG	Myasthenia gravis
MG(H)	Milligram (hours)
MI	Myocardial infarction
MICRO	Microscopic
MIN	Minimum
MIN	Minute
ML	Middle lobe
ML	Milliliter
MM	Millimeter
MM	Multiple myeloma
MOD	Moderate (ly)
MOD DIFF	Moderately differentiated
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
NA	Not applicable
NED	No evidence of disease
NEG	Negative
NEOPL	Neoplasm
NEURO	Neurology
NH	Nursing home
NHL	Non-Hodgkins lymphoma
NL	Normal
NOS	Not otherwise specified

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

ABBREVIATION/SYMBOL	WORD/TERM(S)
NR	Not recorded
NSCCA	Non small cell carcinoma
NSF	No significant findings
NVD	Neck vein distention
OB	Obstetrics
OBS	Organic brain syndrome
OBST	Obstructed (-ing, -ion)
OP	Outpatient
OP RPT	Operative report
OR	Operating room
ORTHO	Orthopedics
ОТО	Otology
OZ	Ounce
P32	Phosphorus 32
PAC	Premature atrial contraction
PALP	Palpated (-able)
PAP	Papanicolaou smear
PAP	Papillary
PATH	Pathology
PD	Poorly differentiated
PE	Physical examination
PEDS	Pediatrics
PERC	Percutaneous
PET	Positron emission tomography
PID	Pelvic inflammatory disease
PIN III	Prostatic intraepithelial neoplasia, grade III
PLT	Platelets
PMH	Past/personal (medical) history
PMP	Primary medical physician
POOR DIFF	Poorly differentiated
POS	Positive
POSS	Possible
POST POST OP	Posterior Posterorativo ( lu)
	Postoperative (-ly)
PPD PR, PRA	Packs per day Progesterone receptor (assay)
PRE OP	Properative (-ly)
PREV	Previous
PROB	Probable (-ly)
PROCTO	Proctoscopy
PSA	Prostatic specific antigen
PT	Patient
PT	Physiotherapy/Physical therapy

ABBREVIATION/SYMBOL	WORD/TERM(S)
РТА	Prior to admission
РТС	Percutaneous transhepatic cholecystogram
PUD	Peptic ulcer disease
PULM	Pulmonary
PVD	Peripheral vascular disease
Q	Every
QD	Every day
QUAD	Quadrant
R/O	Rule out
RA	Rheumatoid arthritis
RAD	Radiation absorbed dose
RBBB	Right bundle branch block
RBC	Red blood cells (count)
RCM	Right costal margin
RE	Regarding
REC'D	Received
REG	Regular
RESEC	Resection (ed)
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
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
RUQ	Right upper quadrant
RX	Prescription
S/P	Status post
S1-S5	Sacral vertebra
S-SPINE	Sacral spine
SATIS	Satisfactory
SB	Small bowel

Version 10.2 -- Appendix G: Recommended Abbreviations for Abstractors

ABBREVIATION/SYMBOL	WORD/TERM(S)
SCC	Squamous cell carcinoma
SCID	Severe combined immunodeficiency syndrome
SGOT	Serum glutamic oxaloacetic transaminase
SGPT	Serum glutamic pyruvic transaminase
SIADH	Syndrome of inappropriate ADH
SIG COLON	Sigmoid colon
SLE	Systemic lupus erythematosus
SM	Small
SO	Salpingo-oophorectomy
SOB	Short(ness) of breath
SPEC	Specimen
SQ	Squamous
SS	Summary stage
SSS	Sick sinus syndrome
STSG	Split thickness skin graft
SUBCU	Subcutaneous
SURG	Surgery/Surgical
SUSP	Suspicious/suspected
SVC	Superior vena cava
SX	Symptoms
T-SPINE	Thoracic spine
ТАН	Total abdominal hysterectomy
TAH-BSO	Total abdominal hysterectomy- bilateral
ТВ	Tuberculosis
TCC	Transitional cell carcinoma
TIA	Transient ischemic attack
TRANS-COLON	Transverse colon
TTP	Thromboticthrombocytopenia purpura
TUR	Transurethral resection
TURB	Transurethral resection bladder
TURP	Transurethral resection prostate
TVC	True vocal cord
TVH	Total vaginal hysterectomy
TX	Treatment
UE	Upper extremity
UGI	Upper gastrointestinal (series)
UIQ	Upper inner quadrant
UNDIFF	Undifferentiated
UNK	Unknown
UOQ	Upper outer quadrant
URI	Upper respiratory infection
US	Ultrasound
UTI	Urinary tract infection

ABBREVIATION/SYMBOL	WORD/TERM(S)
VAG	Vagina/Vaginal
VAG HYST	Vaginal hysterectomy
VAIN III	Vaginal intraepithelial neoplasia (grade III)
VIN III	Vulvar intraepithelial neoplasia (grade III)
W/	With
W/F	White female
W/M	White male
W/O	Without
W/U	Work-up
WBC	White blood cells (count)
WD	Well differentiated
WELL DIFF	Well differentiated
WNL	Within normal limits
WPW	Wolff-Parkinson-White syndrome
XR	Xray
YR	Year

NAACCR RECOMMENDED ABBREVIATION LIST	
CONTEXT-SENSITIVE ABBREVIATIONS	

ABBREVIATION/SYMBOL	WORD/TERM(S)
AP	Anteroposterior
AP	Abdominal perineal
BM	Bone marrow
BM	Bowel movement
СА	Calcium
СА	Carcinoma
MIN	Minimum
MIN	Minute
ML	Milliliter
ML	Middle lobe
MM	Millimeter
MM	Multiple myeloma
PAP	Papillary
PAP	Papanicolaou smear
РТ	Patient
РТ	Physiotherapy/Physical therapy
RT	Right
RT	Radiation therapy

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