Appendix E1 - 2024 SEER Program Coding and Staging Manual

Reportable Examples

As referenced in the Reportability instructions of the 2024 SEER Program Coding and Staging Manual

| Rep | Reportable Malignant Examples | | |
|-----|---|---|--|
| # | Diagnosis/Condition | Notes | |
| 1 | Atypical fibroxanthoma (superficial malignant fibrous histiocytoma) | The information in parentheses provides more detail and confirms a reportable malignancy. | |
| 2 | Positive histology from needle biopsy followed by negative resection | This case is reportable based on positive needle biopsy. | |
| 3 | Biopsy-proven squamous cell carcinoma of the nipple with a subsequent areolar resection showing foreign body granulomatous reaction to suture material and no evidence of residual malignancy in the nipple | This case is reportable. The fact that no residual malignancy was found in the later specimen does not disprove the malignancy diagnosed by the biopsy. | |
| 4 | Ulcerated histologically malignant spindle cell neoplasm, consistent with atypical fibroxanthoma; an exhaustive immunohistochemical work-up shows no melanocytic, epithelial or vascular differentiation | Atypical fibroxanthoma is a superficial form of a malignant fibrous histiocytoma. This case is reportable. The pathologist has the final say on behavior for a particular case. In this case, the pathologist states that this tumor is malignant. | |
| 5 | Aggressive adult granulosa cell tumor with one of two lymph nodes positive for malignant metastatic granulosa cell tumor | This case is reportable because malignant granulosa cell tumor is reportable. The lymph node metastases prove malignancy. | |
| 6 | Carcinoid of the appendix found on appendectomy | Carcinoid tumor, NOS is reportable (8240/3). | |
| 7 | Microcarcinoid tumors of the stomach | Microcarcinoid and carcinoid tumors are reportable. The ICD-O-3.2 histology code is 8240/3. Microcarcinoid is a designation for neuroendocrine tumors of the stomach when they are less than 0.5 cm. in size. Neuroendocrine tumors of the stomach are designated carcinoid when they are 0.5 cm or larger. The term microcarcinoid tumor is not equivalent to carcinoid tumorlet. | |
| 8 | Ovarian mucinous borderline tumor with foci of intraepithelial carcinoma | This case is reportable because there are foci of intraepithelial carcinoma (carcinoma in situ). | |
| 9 | Squamous cell carcinoma of the anus, NOS | Squamous cell carcinoma of the anus (C210) is reportable. Note: Squamous cell carcinoma of the perianal skin (C445) is not reportable. | |
| 10 | Mature teratoma of the testis when diagnosed after puberty (malignant) | For testis: Mature teratoma in adults is malignant (9080/3). Note: Do not report when diagnosed in a child (benign). Do not report mature teratoma of the testis when it is not known whether the patient is prepubescent or postpubescent. Pubescence can take place over a number of years; review physical history and do not rely only on age. | |

| # | Diagnosis/Condition | Notes |
|----|---|--|
| 11 | Well-differentiated neuroendocrine tumor (NET) of the | The WHO classification of digestive system tumors uses the term NET G1 (grade 1) as a synonym for |
| | stomach | carcinoid and well-differentiated NET, 8240/3. |
| 12 | Cystic pancreatic endocrine neoplasm (CPEN) | Assign 8150/3 unless specified as a neuroendocrine tumor, Grade 1 (8240/3) or neuroendocrine |
| | | tumor, Grade 2 (8249/3). |
| 13 | Solid pseudopapillary neoplasm of the pancreas | Assign 8452/3. |
| 14 | Liver cases with an LI-RADS category LR-4 or LR-5 | |
| | | Report based on the American College of Radiology Liver Imaging Reporting and Data System (LI-RADS) <u>definitions.</u> |
| | | Use the date of the LR-4 (Probably HCC) or LR-5 (Definitely HCC) scan as the date of diagnosis when it is the earliest confirmation of the malignancy. |
| | | If there is no statement of the LI-RADS score but there is reference that a lesion is in the Organ Procurement and Transplantation Network (OPTN) 5 category, report based on the OPTN class of 5. |
| | | OPTN class 5 indicates that a nodule meets radiologic criteria for hepatocellular carcinoma. |
| 15 | Mammary analogue secretory carcinoma (MASC) | MASC is a tumor that predominantly arises in the parotid gland. If the primary site is submandibular gland, assign C080. Assign 8502/3. Override any edits triggered by the combination of C080 and 8502/3. |
| 16 | Malignant perivascular epithelioid cell tumor (PEComa) | Assign 8714/3 to malignant PEComa. Some PEComas such as angiomyolipoma and lymphangiomyomatosis have specific ICD-O codes and their malignant counterparts may be coded to 8860/3 and 9174/3, respectively. There are no separate ICD-O codes for other specific PEComas, e.g., clear cell sugar tumor of lung, clear cell myomelanocytic tumor of the falciform ligament, and some unusual clear cell tumors occurring in other organs or for PEComa, NOS. These PEComas may therefore be coded to 8005 as clear cell tumors NOS; in other words, clear cell tumors are not clear cell variants of carcinomas, sarcomas, or other specific tumor type. |
| | | Note: PEComa is non-specific as to behavior. Unless the pathologist states that it is malignant, the default code is 8005/1 (non-reportable). |
| 17 | Noninvasive mucinous cystic neoplasm (MCN) of the | For neoplasms of the pancreas, MCN with high grade dysplasia is the preferred term and mucinous |
| | pancreas with high grade dysplasia | cystadenocarcinoma, noninvasive is a related term (8470/2). |
| 18 | Noninvasive low grade (micropapillary) serous carcinoma (MPSC) of the ovary | Assign code 8460/2, applying the ICD-O-3 matrix concept to this noninvasive carcinoma. Noninvasive can be used as a synonym for in situ, ICD-O-3 behavior code /2. See page 66 in ICD-O-3. |
| | | |

| # | Diagnosis/Condition | Notes |
|----|--|--|
| 19 | Prostate cancer cases with an PI-RADS category 4 or 5 | Report based on the American College of Radiology Prostate Imaging Reporting and Data System (PI-RADS) <u>definitions</u> . PI-RADS categories 4 (high-clinically significant cancer is likely to be present) and 5 (very high-clinically significant cancer is highly likely to be present) are reportable, unless there is other information to the contrary. |
| 20 | Early or evolving melanoma, in situ or invasive | As of 1/1/2021, early or evolving melanoma in situ, or any other early or evolving melanoma, is reportable. |
| 21 | | Report LAMN beginning with January 1, 2022 diagnoses. LAMN is assigned a behavior of /2 or /3 making it reportable. LAMNs are slow-growing neoplasms that have the potential for peritoneal spread and can result in patient death. LAMNs demonstrate an interesting biology in that they do not have hematogenous dissemination risk, but risk for appendiceal perforation, which can result in peritoneal dissemination, repeated recurrences after surgery and even death. |
| 22 | Clear cell papillary renal cell carcinoma | Clear cell papillary renal cell carcinoma (8323/3) is reportable. |
| 23 | grade II, or grade III (with exceptions) • High grade squamous intraepithelial lesion (HGSIL or HSIL) (with exception) • Intraepithelial neoplasia grade II/III; II-III • Squamous dysplasia, high grade (for stomach, small intestine, and esophagus only) | Exceptions (not reportable) Squamous intraepithelial neoplasia (SIN) of cervix (C53_) and skin sites coded to C44_ Carcinoma in situ (CIS) arising in cervix (53_) and perianal skin (C445) Cervical intraepithelial neoplasia (CIN III) of cervix (C53_) High grade squamous intraepithelial lesion (HGSIL or HSIL) arising in perianal skin (C445) AIN II and AIN III (8077) arising in perianal skin (C445) High grade prostatic intraepithelial neoplasia (PIN) See also the 2024 SEER manual, Reportability section, for additional reportable terms. |

| # | Diagnosis/Condition | Notes |
|----|---|-------|
| 24 | 8380/2 (C54_) | |
| | Endometrioid intraepithelial neoplasia (EIN) | |
| | Intraepithelial neoplasm of endometrium | |
| | Atypical hyperplasia of endometrium | |
| 25 | Pancreatic intraepithelial neoplasia (PanIN III) 8148/2 | |
| 26 | Differentiated penile intraepithelial neoplasia 8071/2 | |
| 27 | Intracholecystic papillary neoplasm (ICPN) with high- | |
| | grade dysplasia 8503/2 | |

| # | Diagnosis/Condition | Notes |
|-----|---|---|
| Rep | ortable Non-Malignant Examples | |
| # | Diagnosis/Condition | Notes |
| 28 | Hemangioma, NOS (9120/0) and cavernous hemangioma | Report the CNS site in which the hemangioma originates. |
| | (9121/0) | <i>Note:</i> For cavernous sinus hemangioma, report the site as cerebral meninges C700. |
| 29 | Dermoid cyst of the brain | This condition is reportable for cases diagnosed 2004 and later. Assign 9084/0. |
| 30 | Tectal plate lipoma | This is a reportable brain tumor. It is a benign neoplasm (lipoma) of the mid brain (brain stem) as |
| | | noted by the location "tectal plate." |
| 31 | Lhermitte-Duclos disease | The WHO classification for CNS tumors lists this entity as dysplastic gangliocytoma of the cerebellum |
| | | (Lhermitte-Duclos disease) signifying that the terms are used synonymously. Assign C716, 9493/0. |
| | | |
| 32 | Rathke pouch tumor (C751, 9350/1) | Rathke pouch tumor is a reportable neoplasm for cases diagnosed 2004 and later. Rathke cleft cyst |
| | | and Rathke pouch tumor are different conditions. |
| | | Note: Rathke cleft cyst is not reportable. |

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Non-Reportable Examples

As referenced in the Reportability instructions of the 2024 SEER Program Coding and Staging Manual

| # | Diagnosis/Condition | Notes |
|----|---|---|
| | | Lymph node involvement with sclerosing pneumocytoma, formerly sclerosing hemangioma (an |
| | | obsolete term), is non-malignant. According to the WHO Classification of Thoracic Tumors, 5th |
| | Sclerosing pneumocytoma of the lung with multiple | edition, "most sclerosing pneumocytomas behave in a benign fashion. However, although cases with |
| | regional lymph nodes involved with sclerosing | lymph node metastases and distant organ metastases have occasionally been reported, these |
| 1 | pneumocytoma | findings do not appear to adversely affect prognosis ." |
| | High grade squamous intraepithelial lesion (HGSIL or | |
| | HSIL), carcinoma in situ (CIS), and AIN III (8077) arising in | HGSIL or HSIL, CIS, and AIN III arising in perianal skin are not reportable. Refer to the Reportability |
| 2 | perianal skin (C445) | Section of the main manual. |
| | | Squamous cell carcinoma of sites in C44 is not reportable. Squamous cell carcinoma of the anus |
| 3 | Squamous cell carcinoma of the perianal skin (C445) | (C210) is reportable. |
| 4 | Squamous cell carcinoma of the canthus (C441) | Squamous cell carcinoma in sites coded to C44 is not reportable. |
| 5 | | The American College of Radiology defines Category 4 as "Suspicious." The descriptions in categories 4, 4a, 4b, and 4c are not diagnostic of malignancy. They all represent a percentage of likelihood, the highest being 4c which is greater than 50% but less than 95% likelihood of malignancy. The ACR states "This category is reserved for findings that do not have the classic appearance of malignancy but are sufficiently suspicious to justify a recommendation for biopsy." Category 5 is "Highly Suggestive of Malignancy." "Suggestive" is not reportable ambiguous terminology. ACR states that Category 5 has a "very high probability" of malignancy, but again, it is not diagnostic. |
| | | Lung: Do not use the ACR Lung Imaging Reporting and Data System (Lung-RADS™) to determine |
| 6 | Lung cases designated "Lung-RADS 4A," 4B, or 4X | reportability. Look for reportable terminology from the managing physician or other sources. |
| | Liver cases based only on an LI-RADS category of | |
| 7 | LR-3 | Do not report liver cases based only on an LI-RADS category of LR-3. |
| | | DIPNECH is a generalized proliferation of scattered single cells, small nodules (neuroendocrine |
| | Diffuse idiopathic pulmonary neuroendocrine cell | bodies) or linear proliferation of pulmonary neuroendocrine cells (PNCs) according to the WHO |
| 8 | hyperplasia (DIPNECH) | classification of lung tumors. |
| | Basal cell carcinoma (BCC) with neuroendocrine | |
| | differentiation of the skin | BCC in sites coded to C44 is not reportable to SEER. |
| 10 | Lentiginous melanocytic lesion | Not reportable. |
| | Intraductal papillary mucinous neoplasms with low or | |
| 11 | moderate grade dysplasia (also called IPMN adenomas) | Not reportable. |

| # Diagnosis/Condition | Notes |
|--|---|
| Noninvasive mucinous cystic neoplasm (MCN) of the | |
| 12 pancreas with low or intermediate grade dysplasia | Not reportable. |
| | Subdural hygroma is not a neoplasm; it is a collection of cerebrospinal fluid in the subdural space. It |
| 13 Subdural hygroma | may be related to a head injury. |
| 14 Brain lesions associated with multiple sclerosis | These brain lesions are not neoplastic; they are part of the disease process of multiple sclerosis. |
| | Pubescence can take place over a number of years; review history and physical information and do |
| Mature teratoma of the testis when diagnosed before | not rely only on age. Do not report mature teratoma when it is not known whether the patient is pre- |
| 15 puberty (benign, 9084/0). | or post-pubescent. |
| 16 Mature teratoma of the ovary (9080/0) | Not reportable. |
| | The primary site for venous (hem)angioma arising in the brain is blood vessel (C490). The |
| | combination of 9122/0 and C490 is not reportable. This is a venous abnormality. Previously called |
| 17 Venous angiomas (9122/0) | venous angiomas, these are currently referred to as developmental venous anomalies (DVA). |
| | Previously called multilocular cystic renal cell carcinoma, this diagnosis became non-reportable |
| Multilocular cystic renal neoplasm of low malignant | beginning with the new designation in 2016. Refer to the Solid Tumor Tumor Coding Rules, Kidney |
| 18 potential | Equivalent Terms and Definitions, for histology/morphology information. |
| | Lymphangioma is a malformation of the lymphatic system. Even though it has an ICD-O code, do not |
| 19 Lymphangioma of the brain or CNS | report it. |
| | Carcinoid heart disease is not reportable but this diagnosis indicates that the patient likely has a |
| 20 Carcinoid heart disease based on clinical information | carcinoid tumor which may be reportable. Obtain further information. |
| 21 Carcinoid tumorlet of the lung | Not reportable. |
| | According to WHO, this resembles a typical leiomyoma but it is found in the lungs of women with a |
| | history of typical uterine leiomyomas. A recent article states that because of the hormone-sensitive |
| Pulmonary benign metastasizing leiomyoma (BML) | characteristics of BML, treatments are based on hormonal manipulation along with either surgical or |
| 22 (8898/1) | medical oophorectomy. Tamoxifen treatment is in keeping with the BML diagnosis. |
| | Colloid cysts are endodermal congenital malformations and do not have an ICD-O-3 code. See the |
| 23 Colloid cyst at the foramen of Monro | glossary for registrars at: Colloid cyst |
| | Mammary fibromatosis is not reportable. The WHO classification for breast tumors assigns |
| | mammary fibromatosis a behavior code of /1. According to WHO, mammary fibromatosis is a locally |
| 24 Mammary fibromatosis | infiltrative lesion without metastatic potential. |
| | Amyloidoma (tumoral amyloidosis, amyloid tumor) is a tumor-like deposit of amyloid. It is not |
| 25 Thalamic amyloidoma | neoplastic. Amyloid is a protein derived substance deposited in various clinical settings. |
| | Pseudotumor cerebri is not a neoplasm. The pressure inside the skull is increased and the brain is |
| 26 Pseudotumor cerebri | affected in a way that appears to be a tumor, but it is not a tumor. |

| # | Diagnosis/Condition | Notes |
|----|--|--|
| 27 | | According to our expert pathologist consultant, there has been a lot of debate in the literature about the diagnostic criteria, terminology, and natural history of PAM. The main issue is whether PAM with atypia should be regarded as melanoma in situ. In most studies it appears that PAM with no atypia or mild atypia does not progress to melanoma, and only a small percentage of those with severe atypia do so. PAM, even with atypia, is not melanoma in situ, and should not be reported. For further information, see this article for a review of a large number of patients: Shields, Jerry A, Shields, Carol L, et al. Primary Acquired Melanosis of the Conjunctiva: Experience with 311 Eyes. Trans. Am Ophthalmol Soc 105:61-72, Dec 2007. |
| 28 | Neurofibromatosis type 1 (NF1) and Neurofibromatosis type 2 (NF2) | Genetic disease that produces non-malignant tumors in skin, brain, CNS, and other sites. The brain and CNS tumors spawned by NF1 or NF2 are reportable, the genetic disease is not. |
| 29 | Ovarian mucinous borderline tumor with microinvasion | For an ovarian mucinous borderline tumor, the term "microinvasion" is not an indication of malignancy. Low malignant potential/borderline ovarian tumors are defined by the pathology of the primary tumor and are not affected by microinvasion or invasion in implants. Though a case may be staged, this does not mean it is reportable. |
| 30 | Rathke cleft cyst | Rathke cleft cyst, also called pars intermedia cyst of the parotid gland, is not reportable; whereas, Rathke pouch tumor is reportable. |
| 31 | Colon atypical hyperplasia | Not reportable. |
| | High grade dysplasia in sites other than stomach, small intestine, and esophageal primary sites Ecchordosis physaliphora | Not reportable. Ecchordosis physaliphora, a lesion within the prepontine cistern, is not reportable. |
| | Low to intermediate grade neuroendocrine neoplasm or middle ear adenomatoid tumor (MEANT) dysplasia of lung | Not reportable. |
| 35 | | Not reportable. |
| 36 | | PIN III is not reportable. |