State of Connecticut
Department of Public Health

Reportable List
Connecticut Tumor Registry

All cases with behavior code of “2” or “3” in the International Classification of Diseases of Oncology:
- Reportable skins NOS of the genital sites only (C52.9, C51.0-C51.9, C60.0, C60.9 and C63.2) vagina, clitoris, vulva, prepuce, penis, and scrotum.
- If a “0” or “1” behavior code item in ICD-O is verified by a pathologist as in-situ or malignant, these cases are reportable.
- Behavior code changes from borderline/1 to malignant/3 are reportable (for cases diagnosed 1/1/01 or later).
- Bronchial adenoma carcinoid type or cylindroid type are malignant conditions (M-8240/3 and M-8200/3 respectively).
- Report only malignant cystosarcoma phyllodes (M-9020/3).
- Argentaffin tumors also have a malignant code (M-8241/3).
- Any “Benign” tumor which states “malignant changes” or “malignant degeneration”.
- Any “Pre-1935” diagnosis of a malignancy only if new primary tumor is diagnosed in 1935 or later.
- Any previously diagnosed malignancy mentioned in past history.
- Benign tumors of brain reported in past history.
- Non-resident cases diagnosed after January 1979.
- All autopsy diagnoses.
- All cases considered to be malignant clinically.

The following are exclusions:
- M8000 – M8110 Neoplasms, malignant, NOS of the skin (C44.0 – C44.9)(has applied since 6/1/84).
- M8010 – M8046 Epithelial carcinomas of the skin (C44.0 – C44.9)(has applied since 6/1/84).
- M8050 – M8084 Papillary and squamous cell carcinoma of the skin (C44.0 – C44.9)(has applied since 6/1/84).
- M8090 – M8110 Basal cell carcinoma of any site except genital sites (previously removed from reportable list).
- M8010 – M8077 Carcinoma in-situ (any/2) and CINIII of the cervix (C53.0 – C53.9)(cases diagnosed after 1/1/96).
- M8442 – M8473 Borderline tumors of ovary (cystadenomas) revert to/1(cases diagnosed after 12/31/2000).

Reportable “Benign” tumors listed below:
- All neoplasms of the brain and central nervous system (C70.0 through C72.9).
- Carcinoid and argentaffin tumors of the appendix.
- Papillomas of urinary bladder.
- Argentaffin tumors of the entire digestive tract.
- Polycythemia vera.
- Papillomas of urinary organs ureter, urethra, kidney and renal pelvis.

The following neoplasms have been removed from the benign REPORTABLE list effective January 1, 1994 diagnoses:
- Mixed tumors, salivary gland type.
- Papillary adenomas or mixed papillary and follicular adenomas of the thyroid.
- “Benign” kaposi’s sarcoma.
- Bronchial adenomas.
- Cystosarcoma phyllodes of the breast.
- Granulosa Cell Tumor, NOS (8620/1).
FOR CASES DIAGNOSED 2001 AND LATER

NEWLY REPORTABLE HEMATOPOIETIC DISEASE (NRHD):
Any of the myeloprofilerative or myelodysplastic diseases that changed from /1 borderline to /3 malignant in ICD-O-3

MYELODYSPLASIA
Other names: preleukemia, preleukemic disorders, smoldering leukemia, oligoblastic leukemia, subacute myeloid leukemia, de novo myelodysplastic syndrome, dysmyelopoietic syndrome, homopoietic dysplasia
ICD-O-3 C42.1
ICD-O-3 M9989/3

MYELODYSPLASTIC SYNDROME (NRHD)
ICD-O-3 C42.1
M9980/3 Refractory anemia
M9981/3 Refractory anemia without sideroblasts
M9982/3 Refractory anemia with ringed sideroblasts
M9983/3 Refractory anemia with excess blasts
M9984/3 Refractory anemia with excess blasts In transformation (obs)
M9985/3 Refractory cytopenia with multilineage dysplasia
M9986/3 Myelodysplastic syndrome with 5q-(5q deletion) syndrome
M9987/3 Therapy related myelodysplastic syndrome

CHRONIC MYELOPROLIFERATIVE DISORDER
Other name: chronic myeloproliferative disease
ICD-O-3 C42.1
ICD-O-3 M9960/3

MYELOSCLEROSIS WITH MYELOID METAPLASIA
Other names: myelofibrosis with meyloid metaplasia, primary myelofibrosis (PMF), myelofibrosis as a result of myeloproliferative disease, megakaryocytic meyloidosclerosis, agnogenic myeloid metaplasia (AMM)
ICD-O-3 C42.1
ICD-O-3 M9961/3

ESSENTIAL THROMBOCYTHEMIA
Other names: idiopathic thrombocythemia, essential hemorrhagic thrombocythemia, idiopathic thrombocythemia, Primary thrombocythemia
ICD-O-3 C42.1
ICD-O-3 M9962/3

BEHAVIOR CODE CHANGES BORDERLINE/1 TO MALIGNANT /3
8931/3 Endolymphatic stromal myosis (C54.1) now a synonym for endometrial stromal sarcoma, low grade
9538/3 Papillary meningioma malignant version of clear/chordoid meningioma

REPORTABLE BEHAVIOR FROM CODE CHANGE MALIGNANT/3 TO BORDERLINE/1
Pilocytic/juvenile/piloid astrocytoma/spongioblastoma: listed as M-9421/1 ICD-O-3 will remain reportable as M9421/3 and sequence as malignant in North America (ICD-O-3 Errata and Clarifications 5/22/2001)

BEHAVIOR CODE CHANGES MALIGNANT/3 TO BORDERLINE/1 WHICH ARE NON-REPORTABLE
Borderline tumors of ovary C56.9
M-8442/1 serous cystadenoma borderline
M-8451/1 papillary cystadenoma borderline
M-8462/1 serous papillary cystadenoma borderline
M-8472/1 mucinous cystadenoma borderline
M-8473/1 papillary mucinous cystadenoma borderline

NEWLY REPORTABLE HISTOLOGY TERMS FOR CASES DIAGNOSED 2010 AND LATER

Primary cutaneous follicle centre lymphoma 9597/3
T-cell/histiocyte rich large B-cell lymphoma 9688/3
Intravascular large B-cell lymphoma 9712/3
Systemic EBV positive T-cell lymphoproliferative disease of childhood 9724/3
Hydroa vacciniforme-like lymphoma 9725/3
Primary cutaneous gamma-delta T-cell lymphoma 9726/3
Plasmablastic lymphoma 9735/3
ALK positive large B-cell lymphoma 9737/3
Large B-cell lymphoma arising in HHV8-associated multicentric Castleman disease 9738/3
Fibroblastic reticular cell tumor 9759/3
Mixed phenotype acute leukemia with t(9;22)(q34;q11.2); BCR-ABL1 9806/3
Mixed phenotype acute leukemia with t(v;11q23); MLL rearranged 9807/3
Mixed phenotype acute leukemia, B/myeloid, NOS 9808/3
Mixed phenotype acute leukemia, T/myeloid, NOS 9809/3
B lymphoblastic leukemia/lymphoma, NOS 9811/3
B lymphoblastic leukemia/lymphoma with t(9;22)(q34;q11.2); BCR-ABL1 9812/3
B lymphoblastic leukemia/lymphoma with t(v;11q23); MLL rearranged 9813/3
B lymphoblastic leukemia/lymphoma with t(12;21)(p13;q22); TEL-AML1 (ETV6-RUNX1) 9814/3
B lymphoblastic leukemia/lymphoma with hyperdiploidy 9815/3
B lymphoblastic leukemia/lymphoma with hypodiploidy (hypodiploid ALL) 9816/3
B lymphoblastic leukemia/lymphoma with t(5;14)(q31;q32); IL3-IGH 9817/3
B lymphoblastic leukemia/lymphoma with t(1;19)(q23;p13.3); E2A PBX1 (TCF3 PBX1) 9818/3
T lymphoblastic leukemia/lymphoma 9837/3
Acute myeloid leukemia with t(6;9)(p23;q34) DEK-NUP214 9865/3
Acute myeloid leukemia with inv(3)(q21q26.2) or t(3;3)(q21;q26.2); RPN1EVI1 9869/3
Myeloid leukemia associated with Down Syndrome 9898/3
Acute myeloid leukemia (megakaryoblastic) with t(1;22)(p13;q13); RBM15-MKL1 9911/3
Myeloid and lymphoid neoplasms with PDGFRB rearrangement 9965/3
Myeloid and lymphoid neoplasms with PDGFRB arrangement 9966/3
Myeloid and lymphoid neoplasms with FGFR1 abnormalities 9967/3
Polymorphic PTLD 9971/3
Refractory neutropenia 9991/3
Refractory thrombocytopenia 9992/3

NEWLY REPORTABLE CONDITIONS CHANGES FROM BENIGN/1 TO MALIGNANT/3 2010 AND FORWARD

Langerhans cell histiocytosis, NOS 9751/3
Myeloproliferative neoplasm, unclassifiable/ Myelodysplastic/Myeloproliferative neoplasm, unclassifiable 9975/3
T-cell large granular lymphocytic leukemia/ Chronic lymphoproliferative disorder of NK-cells 9831/3

Please refer to ICD-O-3 Appendix items 1-6 (pg. 219-240) for New Morphology Terms & Synonyms, New Codes and Changes.