

mCode to OMOP Standard Library

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```
library(mOMOP)
library(chariot)
library(tidyverse)
```

```
conn <- chariot::connectAthena()
```

This vignette takes a look at the Value Sets in the mCode Data Dictionary to create the standard library of Oncology concepts, available for loading in this package.

The mCode valuesets are retrieved from the publicly available data dictionary.

```
value_sets <- get_value_sets()
value_sets
```

```
## # A tibble: 6,603 x 5
##   `Value Set Name`   `Code System` `Logical Definitio~ Code `Code Description`
##   <chr>             <chr>         <chr>             <chr> <chr>
## 1 CancerBodyLocatio~ SNOMED CT     includes codes des~ <NA> <NA>
## 2 CancerDiseaseStat~ SNOMED CT     <NA>              3636~ Imaging (procedur~
## 3 CancerDiseaseStat~ SNOMED CT     <NA>              2524~ Histopathology te~
## 4 CancerDiseaseStat~ SNOMED CT     <NA>              7110~ Assessment of sym~
## 5 CancerDiseaseStat~ SNOMED CT     <NA>              5880~ Physical examinat~
## 6 CancerDiseaseStat~ SNOMED CT     <NA>              2507~ Tumor marker meas~
## 7 CancerDiseaseStat~ SNOMED CT     <NA>              3863~ Laboratory data i~
## 8 CancerDisorderVS   ICD-10-CM     <NA>              C000 Malignant neoplas~
## 9 CancerDisorderVS   ICD-10-CM     <NA>              C001 Malignant neoplas~
## 10 CancerDisorderVS  ICD-10-CM     <NA>              C002 Malignant neoplas~
## # ... with 6,593 more rows
```

To map the valuesets to OMOP concepts, it is split by `Code System`, the equivalent to the `vocabulary_id` field in OMOP's Concept table, for joining. As of 2021-01-02, there are 9 `Code Systems` used in mCode.

```
value_sets_by_vocab <- value_sets %>% rubix::split_by(col = `Code System`)
names(value_sets_by_vocab)
```

```
## [1] "ClinVar"
## [2] "http://cancerstaging.org"
## [3] "http://terminology.hl7.org/CodeSystem/v2-0487"
## [4] "http://unitsofmeasure.org"
## [5] "http://varnomen.hgvs.org"
## [6] "http://www.genenames.org/geneId"
## [7] "ICD-10-CM"
## [8] "LOINC"
## [9] "SNOMED CT"
```

Specimen

In mCode, Specimen representation is based on HL7's Code System while in OMOP, the Specimen domain has its own subset of concepts. Furthermore, mCode's Specimen representation maps to other OMOP Concept Ids. The Specimen representation in this package therefore has 2 parts:

1. A map between mCode and OMOP Concepts outside OMOP's Specimen domain
2. OMOP's Specimen domain concepts

Map from mCode to OMOP

Specimen representation is derived from the valuesets.

```
specimen_library <- value_sets_by_vocab$`http://terminology.hl7.org/CodeSystem/v2-0487` %>%
  rubix::format_colnames()
head(specimen_library)
```

```
## # A tibble: 6 x 5
##   value_set_name code_system logical_definit~ code code_description
##   <chr>          <chr>          <chr>          <chr> <chr>
## 1 GeneticSpecimen~ http://terminology.h~ <NA>          AMN Amniotic fluid
## 2 GeneticSpecimen~ http://terminology.h~ <NA>          BIFL Bile Fluid
## 3 GeneticSpecimen~ http://terminology.h~ <NA>          BLD Whole blood
## 4 GeneticSpecimen~ http://terminology.h~ <NA>          BLDA Blood arterial
## 5 GeneticSpecimen~ http://terminology.h~ <NA>          BLDCO Cord blood
## 6 GeneticSpecimen~ http://terminology.h~ <NA>          BLDV Blood venous
```

The specimen description is joined on the Concept Synonym Names in the Concept Synonym table to retrieve the Concept Id the mCode Specimen maps to.

```
specimen_omop_library1 <- join_on_concept_synonym_name(data = specimen_library,
  column = "code_description", case_insensitive = TRUE, conn = conn) %>%
  select(-concept_synonym_name, -language_concept_id) %>% rename(match_concept_id = concept_id)
head(specimen_omop_library1)
```

```
##           value_set_name code_system
## 1 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 2 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 3 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
```

```
## 4 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 5 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 6 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
##   logical_definition code code_description match_concept_id
## 1          <NA> SKN           Skin           1027716
## 2          <NA> PLC           Placenta        1032268
## 3          <NA> SAL           Saliva          1033195
## 4          <NA> SPT           Sputum          1033465
## 5          <NA> WND           Wound           1033739
## 6          <NA> BON           Bone            1585831
```

The complete Concept representation is taken from the Concept Id.

```
specimen_omop_library2 <- join_on_concept_id(data = specimen_omop_library1,
      column = "match_concept_id", conn = conn)

head(specimen_omop_library2)
```

```
##           value_set_name                                code_system
## 1 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 2 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 3 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 4 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 5 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
## 6 GeneticSpecimenTypeVS http://terminology.hl7.org/CodeSystem/v2-0487
##   logical_definition code code_description match_concept_id concept_id
## 1          <NA> SKN           Skin           1027716      1027716
## 2          <NA> PLC           Placenta        1032268      1032268
## 3          <NA> SAL           Saliva          1033195      1033195
## 4          <NA> SPT           Sputum          1033465      1033465
## 5          <NA> WND           Wound           1033739      1033739
## 6          <NA> BON           Bone            1585831      1585831
##           concept_name    domain_id vocabulary_id concept_class_id
## 1              Skin Observation           LOINC      LOINC System
## 2          Placenta Observation           LOINC      LOINC System
## 3              Saliva Observation           LOINC      LOINC System
## 4              Sputum Observation           LOINC      LOINC System
## 5              Wound Observation           LOINC      LOINC System
## 6 Organ Transplant Description: Bone Observation           PPI      Answer
##   standard_concept                concept_code valid_start_date
## 1          <NA>                LP36760-4      1970-01-01
## 2          <NA>                LP7477-5       1970-01-01
## 3          <NA>                LP7565-7       1970-01-01
## 4          <NA>                LP7600-2       1970-01-01
## 5          <NA>                LP7726-5       1970-01-01
## 6          <NA> OrganTransplantDescription_Bone 2017-05-17
##   valid_end_date invalid_reason
## 1    2099-12-31          <NA>
## 2    2099-12-31          <NA>
## 3    2099-12-31          <NA>
## 4    2099-12-31          <NA>
## 5    2099-12-31          <NA>
## 6    2099-12-31          <NA>
```

To maintain a one-to-one representation of the original grain of information, the OMOP mappings are pivoted on the OMOP Domain to see how each mCode Specimen maps to by domain.

```
specimen_omop_library3 <- specimen_omop_library2 %>% merge_strip(into = "concept",
  domain_id) %>% pivot_wider(id_cols = !concept, names_from = domain_id,
  values_from = concept) %>% select(-match_concept_id) %>%
  distinct()
head(specimen_omop_library3)
```

```
## # A tibble: 6 x 14
##   concept_id value_set_name code_system logical_definit~ code code_description
##   <dbl> <chr> <chr> <chr> <chr> <chr>
## 1 1027716 GeneticSpecim~ http://ter~ <NA> SKN Skin
## 2 1032268 GeneticSpecim~ http://ter~ <NA> PLC Placenta
## 3 1033195 GeneticSpecim~ http://ter~ <NA> SAL Saliva
## 4 1033465 GeneticSpecim~ http://ter~ <NA> SPT Sputum
## 5 1033739 GeneticSpecim~ http://ter~ <NA> WND Wound
## 6 1585831 GeneticSpecim~ http://ter~ <NA> BON Bone
## # ... with 8 more variables: Observation <chr>, Condition <chr>, `Spec Anatomic
## # Site` <chr>, Drug <chr>, `Meas Value` <chr>, Measurement <chr>,
## # Specimen <chr>, `NA` <chr>
```

To gather a complete representation of Specimens, any missing Specimen concepts in OMOP are added to the library.

```
omop_specimen <- chariot::queryAthena("SELECT *
  FROM omop_vocabulary.concept
  WHERE domain_id = 'Specimen' AND concept_class_id = 'Specimen';",
  conn = conn) %>% merge_strip(into = "Specimen") %>% select(-Specimen_id)
```

```
head(omop_specimen)
```

```
## # A tibble: 6 x 1
##   Specimen
##   <chr>
## 1 [V] [S] 759813 Physical object specimen [SNOMED 1021000124109] [Specimen] [Sp~
## 2 [V] [S] 759820 Paper specimen [SNOMED 1031000124107] [Specimen] [Specimen]
## 3 [V] [S] 759825 Writing paper specimen [SNOMED 1041000124102] [Specimen] [Spec~
## 4 [V] [S] 759826 Envelope specimen [SNOMED 1051000124100] [Specimen] [Specimen]
## 5 [V] [S] 759829 Package specimen [SNOMED 1061000124103] [Specimen] [Specimen]
## 6 [V] [S] 759869 Clothing specimen [SNOMED 1071000124105] [Specimen] [Specimen]
```

```
specimen_omop_library <- specimen_omop_library3 %>% full_join(omop_specimen,
  by = "Specimen") %>% distinct()
head(specimen_omop_library)
```

```
## # A tibble: 6 x 14
##   concept_id value_set_name code_system logical_definit~ code code_description
##   <dbl> <chr> <chr> <chr> <chr> <chr>
## 1 1027716 GeneticSpecim~ http://ter~ <NA> SKN Skin
## 2 1032268 GeneticSpecim~ http://ter~ <NA> PLC Placenta
## 3 1033195 GeneticSpecim~ http://ter~ <NA> SAL Saliva
## 4 1033465 GeneticSpecim~ http://ter~ <NA> SPT Sputum
## 5 1033739 GeneticSpecim~ http://ter~ <NA> WND Wound
## 6 1585831 GeneticSpecim~ http://ter~ <NA> BON Bone
## # ... with 8 more variables: Observation <chr>, Condition <chr>, `Spec Anatomic
## # Site` <chr>, Drug <chr>, `Meas Value` <chr>, Measurement <chr>,
## # Specimen <chr>, `NA` <chr>
```

This file is written to the data-raw/ folder for distribution if it does not already exist.

```
file <- file.path(getwd(), "data-raw", "specimen.csv")
if (!file.exists(file)) {
  write_csv(x = specimen_omop_library, file = file)
}
```

Cancer Staging

mCode uses the AJCC TNM Staging system, which correlates with NCI concepts in the OMOP vocabulary.

```
cancer_staging_library <- value_sets_by_vocab$`http://cancerstaging.org` %>%
  rubix::format_colnames()
head(cancer_staging_library)
```

```
## # A tibble: 4 x 5
##   value_set_name    code_system logical_definition    code code_description
##   <chr>            <chr>        <chr>                <chr> <chr>
## 1 TNMDistantMetast~ http://cancer~ includes codes from c~ <NA> <NA>
## 2 TNMPPrimaryTumorC~ http://cancer~ includes codes from c~ <NA> <NA>
## 3 TNMRegionalNodes~ http://cancer~ includes codes from c~ <NA> <NA>
## 4 TNMStageGroupVS   http://cancer~ includes codes from c~ <NA> <NA>
```

As it is represented in the OMOP Vocabulary, NCIIt is not separated by Tumor, Node, and Metastasis like it is in mCode.

```
ncit_omop_library <- chariot::queryAthena("SELECT *
      FROM omop_vocabulary.concept
      WHERE vocabulary_id = 'NCIt' AND concept_class_id = 'AJCC Category'",
  conn = conn)

head(ncit_omop_library)
```

```
## # A tibble: 6 x 10
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr>        <chr>        <chr>        <chr>
## 1 1537692 Vulvar Canc~ Measurem~ NCIIt        AJCC Category
## 2 1537693 Retinoblast~ Measurem~ NCIIt        AJCC Category
## 3 1537694 Vulvar Canc~ Measurem~ NCIIt        AJCC Category
## 4 1537695 Nasopharyng~ Measurem~ NCIIt        AJCC Category
## 5 1537700 Adrenal Cor~ Measurem~ NCIIt        AJCC Category
## 6 1537780 Retinoblast~ Measurem~ NCIIt        AJCC Category
## # ... with 5 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>
```

NCIt AJCC Category concepts are therefore grouped based on pattern matching with the Concept Code.

```
ncit_omop_library2 <- ncit_omop_library
ncit_omop_library2$value_set_name <- ""
ncit_omop_library2 <- ncit_omop_library2 %>% mutate(value_set_name = case_when(grepl("^[cp]{1}M",
  concept_code) ~ "TNMDistantMetastasesCategoryVS", grepl("^[cp]{1}N",
  concept_code) ~ "TNMRegionalNodesCategoryVS", grepl("^[cp]{1}T",
  concept_code) ~ "TNMPPrimaryTumorCategoryVS"))
head(ncit_omop_library2)
```

```
## # A tibble: 6 x 11
##   concept_id concept_name domain_id vocabulary_id concept_class_id
```

```
##      <dbl> <chr>      <chr>      <chr>      <chr>
## 1    1537692 Vulvar Canc~ Measurem~ NCIIt      AJCC Category
## 2    1537693 Retinoblast~ Measurem~ NCIIt      AJCC Category
## 3    1537694 Vulvar Canc~ Measurem~ NCIIt      AJCC Category
## 4    1537695 Nasopharyng~ Measurem~ NCIIt      AJCC Category
## 5    1537700 Adrenal Cor~ Measurem~ NCIIt      AJCC Category
## 6    1537780 Retinoblast~ Measurem~ NCIIt      AJCC Category
## # ... with 6 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>,
## #   value_set_name <chr>
```

A correlate to mCode's TNMStageGroupVS value set is not present in the OMOP Vocabularies, but since it can be derived from the TNM categories, it is skipped.

```
cancer_staging_omop_library <- cancer_staging_library %>% left_join(ncit_omop_library2,
  by = "value_set_name")
head(cancer_staging_omop_library)
```

```
## # A tibble: 6 x 15
##   value_set_name code_system logical_definit~ code code_description concept_id
##   <chr>          <chr>      <chr>          <chr> <chr>          <dbl>
## 1 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537692
## 2 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537780
## 3 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537798
## 4 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537804
## 5 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537805
## 6 TNMDistantMet~ http://can~ includes codes ~ <NA> <NA>          1537808
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## #   vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## #   concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## #   invalid_reason <chr>
```

This data is written to a cancer_staging.csv in the data-raw/ folder for distribution if it does not already exist.

```
file <- file.path(getwd(), "data-raw", "cancer_staging.csv")
if (!file.exists(file)) {
  write_csv(x = cancer_staging_omop_library, file = file)
}
```

Genomics

HGVS, HGNC, and ClinVar are collapsed into a single Genomics category.

HGVS

mCode uses HGVS (<http://varnomen.hgvs.org>), but these codes are not in the OMOP Vocabulary and cannot be derived from the website. Therefore, it is skipped for now.

```
hgvs_library <- value_sets_by_vocab$http://varnomen.hgvs.org` %>%
  rubix::format_colnames()
head(hgvs_library)
```

```
## # A tibble: 1 x 5
##   value_set_name code_system logical_definition code code_description
```

```
##   <chr>           <chr>           <chr>           <chr> <chr>
## 1 HGVSVS         http://varnomen~ All codes in http://va~ <NA> <NA>
```

HGNC

Gene Names in mCode are presumed to be derived from HGNC and the entire HGNC subset of the OMOP Vocabularies are included in the library.

```
hgnc_library <- value_sets_by_vocab$`http://www.genenames.org/geneId` %>%
  rubix::format_colnames()
head(hgnc_library)
```

```
## # A tibble: 1 x 5
##   value_set_name code_system      logical_definition      code code_description
##   <chr>          <chr>          <chr>          <chr> <chr>
## 1 HGNCVS       http://www.gene~ All codes in http://ww~ <NA> <NA>
```

```
hgnc_omop_library1 <- chariot::queryAthena("SELECT *
      FROM omop_vocabulary.concept
      WHERE vocabulary_id = 'HGNC';",
      conn = conn)
```

```
head(hgnc_omop_library1)
```

```
## # A tibble: 6 x 10
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr>          <chr>          <chr>          <chr>
## 1  35944910 CCDC77 (coi~ Measurem~ HGNC          Gene
## 2  35944911 INMT (indol~ Measurem~ HGNC          Gene
## 3  35944912 ZNF117 (zin~ Measurem~ HGNC          Gene
## 4  35944913 CKAP2 (cyto~ Measurem~ HGNC          Gene
## 5  35944914 ITPRIPL2 (I~ Measurem~ HGNC          Gene
## 6  35944916 PEX19 (pero~ Measurem~ HGNC          Gene
## # ... with 5 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>
```

The OMOP HGNC concepts are joined to the mCode data dictionary set.

```
hgnc_omop_library2 <- hgnc_omop_library1 %>% mutate(value_set_name = "HGNCVS")
head(hgnc_omop_library2)
```

```
## # A tibble: 6 x 11
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr>          <chr>          <chr>          <chr>
## 1  35944910 CCDC77 (coi~ Measurem~ HGNC          Gene
## 2  35944911 INMT (indol~ Measurem~ HGNC          Gene
## 3  35944912 ZNF117 (zin~ Measurem~ HGNC          Gene
## 4  35944913 CKAP2 (cyto~ Measurem~ HGNC          Gene
## 5  35944914 ITPRIPL2 (I~ Measurem~ HGNC          Gene
## 6  35944916 PEX19 (pero~ Measurem~ HGNC          Gene
## # ... with 6 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>,
## #   value_set_name <chr>
```

```
hgnc_omop_library <- hgnc_library %>% left_join(hgnc_omop_library2,
  by = "value_set_name")
head(hgnc_omop_library)
```

```
## # A tibble: 6 x 15
##   value_set_name code_system logical_definit~ code code_description concept_id
##   <chr>          <chr>          <chr>          <chr> <chr>          <dbl>
## 1 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944910
## 2 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944911
## 3 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944912
## 4 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944913
## 5 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944914
## 6 HGNCVS        http://www~ All codes in ht~ <NA> <NA>          35944916
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## #   vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## #   concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## #   invalid_reason <chr>
```

ClinVar

Like HGNC, the entire ClinVar subset of the OMOP Vocabularies are included in the library.

```
clinvar_library <- value_sets_by_vocab$ClinVar %>% rubix::format_colnames()
head(clinvar_library)
```

```
## # A tibble: 1 x 5
##   value_set_name code_system logical_definition code code_description
##   <chr>          <chr>          <chr>          <chr> <chr>
## 1 ClinVarVS      ClinVar      Includes codes from ClinVar <NA> <NA>
```

```
clinvar_omop_library1 <- chariot::queryAthena("SELECT *
      FROM omop_vocabulary.concept
      WHERE vocabulary_id = 'ClinVar';",
      conn = conn)
```

```
head(clinvar_omop_library1)
```

```
## # A tibble: 6 x 10
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr>          <chr>          <chr>          <chr>
## 1 35968119 NC_000002.1~ Measurem~ ClinVar      Variant
## 2 35968121 NM_000038.6~ Measurem~ ClinVar      Variant
## 3 35968122 NM_000038.6~ Measurem~ ClinVar      Variant
## 4 35968123 NM_000038.6~ Measurem~ ClinVar      Variant
## 5 35968124 NM_000038.6~ Measurem~ ClinVar      Variant
## 6 35968125 NM_000038.6~ Measurem~ ClinVar      Variant
## # ... with 5 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>
```

```
clinvar_omop_library2 <- clinvar_omop_library1 %>% mutate(value_set_name = "ClinVarVS")
head(clinvar_omop_library2)
```

```
## # A tibble: 6 x 11
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr>          <chr>          <chr>          <chr>
## 1 35968119 NC_000002.1~ Measurem~ ClinVar      Variant
## 2 35968121 NM_000038.6~ Measurem~ ClinVar      Variant
## 3 35968122 NM_000038.6~ Measurem~ ClinVar      Variant
## 4 35968123 NM_000038.6~ Measurem~ ClinVar      Variant
## 5 35968124 NM_000038.6~ Measurem~ ClinVar      Variant
```



```
## 6 35968125 NM_000038.6~ Measurem~ ClinVar Variant
## # ... with 6 more variables: standard_concept <chr>, concept_code <chr>,
## # valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>,
## # value_set_name <chr>

clinvar_omop_library <- clinvar_library %>% left_join(clinvar_omop_library2,
  by = "value_set_name")
head(clinvar_omop_library)

## # A tibble: 6 x 15
## value_set_name code_system logical_definit~ code code_description concept_id
## <chr> <chr> <chr> <chr> <chr> <dbl>
## 1 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968119
## 2 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968121
## 3 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968122
## 4 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968123
## 5 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968124
## 6 ClinVarVS ClinVar Includes codes ~ <NA> <NA> 35968125
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## # vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## # concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## # invalid_reason <chr>
```

Final Genomics Library

The final genomics library file is written if it does not already exist.

```
genomics_omop_library <- bind_rows(hgnc_omop_library, clinvar_omop_library)
file <- file.path(getwd(), "data-raw", "genomics.csv")
if (!file.exists(file)) {
  write_csv(x = genomics_omop_library, file = file)
}
```

Units of Measure

Like Specimen, the Units of Measure representation is limited and requires incorporation of the OMOP UCUM vocabulary valueset.

```
uom_library <- value_sets_by_vocab$`http://unitsofmeasure.org` %>%
  rubix::format_colnames()
head(uom_library)

## # A tibble: 6 x 5
## value_set_name code_system logical_definit~ code code_description
## <chr> <chr> <chr> <chr> <chr>
## 1 UnitsOfLengthVS http://unitsofmeasur~ <NA> pm Picometer
## 2 UnitsOfLengthVS http://unitsofmeasur~ <NA> nm Nanometer
## 3 UnitsOfLengthVS http://unitsofmeasur~ <NA> mm Millimeter
## 4 UnitsOfLengthVS http://unitsofmeasur~ <NA> cm Centimeter
## 5 UnitsOfLengthVS http://unitsofmeasur~ <NA> m Meter
## 6 UnitsOfLengthVS http://unitsofmeasur~ <NA> ft-us Foot

ucum_omop_library1 <- chariot::queryAthena("SELECT *
  FROM omop_vocabulary.concept")
```

```

WHERE vocabulary_id = 'UCUM';",
conn = conn)

head(ucum_omop_library1)

## # A tibble: 6 x 10
##   concept_id concept_name domain_id vocabulary_id concept_class_id
##   <dbl> <chr> <chr> <chr> <chr>
## 1      8478 avidity ind~ Unit UCUM Unit
## 2      8479 centipoise Unit UCUM Unit
## 3      8480 Ehrlich unit Unit UCUM Unit
## 4      8481 EV Unit UCUM Unit
## 5      8482 pH Unit UCUM Unit
## 6      8483 counts per ~ Unit UCUM Unit
## # ... with 5 more variables: standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>
uom_omop_library <- uom_library %>% bind_rows(ucum_omop_library1)
head(uom_omop_library)

## # A tibble: 6 x 15
##   value_set_name code_system logical_definit~ code code_description concept_id
##   <chr> <chr> <chr> <chr> <chr> <dbl>
## 1 UnitsOfLength~ http://uni~ <NA> pm Picometer NA
## 2 UnitsOfLength~ http://uni~ <NA> nm Nanometer NA
## 3 UnitsOfLength~ http://uni~ <NA> mm Millimeter NA
## 4 UnitsOfLength~ http://uni~ <NA> cm Centimeter NA
## 5 UnitsOfLength~ http://uni~ <NA> m Meter NA
## 6 UnitsOfLength~ http://uni~ <NA> ft-us Foot NA
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## #   vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## #   concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## #   invalid_reason <chr>

Write the data to a file if it does not already exist.

file <- file.path(getwd(), "data-raw", "unitsofmeasurement.csv")
if (!file.exists(file)) {
  write_csv(x = uom_omop_library, file = file)
}

```

SNOMED

The SNOMED subset of the library contains:

- Explicitly stated concepts
- Concepts that are descendants of a stated code
- Concepts that are ancestors of a stated code

To derive all concepts, the code in the `logical_definition` field is extracted based on the listed scenario above.

```

snomed_library <- value_sets_by_vocab$`SNOMED CT` %>% rubix::format_colnames() %>%
  distinct() %>% mutate_all(as.character) %>% extract(col = logical_definition,

```

```

into = "descendants_of", regex = "includes codes descending from ([0-9]{1,})[^0-9]{1}.*$",
remove = FALSE) %>% extract(col = logical_definition, into = "ancestors_of",
regex = "excludes codes descending from ([0-9]{1,})[^0-9]{1}.*$",
remove = FALSE) %>% mutate(all_codes = coalesce(code, ancestors_of,
descendants_of)) %>% mutate_all(trimws)
head(snomed_library)

```

```

## # A tibble: 6 x 8
##   value_set_name code_system logical_definit~ ancestors_of descendants_of code
##   <chr>          <chr>          <chr>          <chr>          <chr>          <chr>
## 1 CancerBodyLoc~ SNOMED CT includes codes ~ <NA>          123037004      <NA>
## 2 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          3636~
## 3 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          2524~
## 4 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          7110~
## 5 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          5880~
## 6 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          2507~
## # ... with 2 more variables: code_description <chr>, all_codes <chr>

```

OMOP Concept table is joined to the mCode's SNOMED library by code.

```

snomed_omop_library1 <- chariot::join_on_concept_code(kind = "LEFT",
  data = snomed_library, column = "all_codes", where_in_concept_field = "vocabulary_id",
  where_in_concept_field_value = "SNOMED")
snomed_omop_library1 <- snomed_library %>% left_join(snomed_omop_library1,
  by = c("value_set_name", "ancestors_of", "descendants_of",
    "all_codes", "code_system", "logical_definition", "code",
    "code_description")) %>% distinct()

```

```
head(snomed_omop_library1)
```

```

## # A tibble: 6 x 18
##   value_set_name code_system logical_definit~ ancestors_of descendants_of code
##   <chr>          <chr>          <chr>          <chr>          <chr>          <chr>
## 1 CancerBodyLoc~ SNOMED CT includes codes ~ <NA>          123037004      <NA>
## 2 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          3636~
## 3 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          2524~
## 4 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          7110~
## 5 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          5880~
## 6 CancerDisease~ SNOMED CT <NA>          <NA>          <NA>          2507~
## # ... with 12 more variables: code_description <chr>, all_codes <chr>,
## #   concept_id <dbl>, concept_name <chr>, domain_id <chr>, vocabulary_id <chr>,
## #   concept_class_id <chr>, standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>

```

The resultset is then split into 3 based on whether the descendants (a) or ancestors (b) need to be derived, or if the concept is explicitly stated by code (c).

The descendants are derived for those mCode concepts that included descendants.

```

snomed_omop_library2a <- snomed_omop_library1 %>% filter(!is.na(descendants_of))
snomed_omop_library2a2 <- join_for_descendants(kind = "LEFT",
  data = snomed_omop_library2a, ancestor_id_column = "concept_id") %>%
  select(all_of(colnames(snomed_library)), starts_with("descendant_")) %>%
  rename_all(~str_remove_all(., pattern = "^descendant_"))

```

```
head(snomed_omop_library2a2)
```

```

##          value_set_name code_system
## 1 CancerBodyLocationVS    SNOMED CT
## 2 CancerBodyLocationVS    SNOMED CT
## 3 CancerBodyLocationVS    SNOMED CT
## 4 CancerBodyLocationVS    SNOMED CT
## 5 CancerBodyLocationVS    SNOMED CT
## 6 CancerBodyLocationVS    SNOMED CT
##
##          logical_definition ancestors_of
## 1 includes codes descending from 123037004 | Body Structure      <NA>
## 2 includes codes descending from 123037004 | Body Structure      <NA>
## 3 includes codes descending from 123037004 | Body Structure      <NA>
## 4 includes codes descending from 123037004 | Body Structure      <NA>
## 5 includes codes descending from 123037004 | Body Structure      <NA>
## 6 includes codes descending from 123037004 | Body Structure      <NA>
##  descendants_of code code_description all_codes concept_id
## 1      123037004 <NA>                <NA> 123037004    4048384
## 2      123037004 <NA>                <NA> 123037004    4002852
## 3      123037004 <NA>                <NA> 123037004    36717763
## 4      123037004 <NA>                <NA> 123037004    4230944
## 5      123037004 <NA>                <NA> 123037004    42605189
## 6      123037004 <NA>                <NA> 123037004    4097829
##
##          concept_name          domain_id
## 1          Body structure Spec Anatomic Site
## 2          Buccal embrasure Spec Anatomic Site
## 3          Skin structure of left lower eyelid Spec Anatomic Site
## 4          Adenofibrosis      Observation
## 5 Intervertebral foramen of eighteenth thoracic vertebra Spec Anatomic Site
## 6          Transitional cell carcinoma      Observation
##
##          vocabulary_id concept_class_id standard_concept concept_code
## 1          SNOMED      Body Structure      S      123037004
## 2          SNOMED      Body Structure      S      110326006
## 3          SNOMED      Body Structure      S      719884003
## 4          SNOMED Morph Abnormality      S      89115006
## 5 SNOMED Veterinary      Body Structure      S 336621000009107
## 6          SNOMED Morph Abnormality      S      27090000
##
##          valid_start_date valid_end_date invalid_reason
## 1          1970-01-01      2099-12-31      <NA>
## 2          1970-01-01      2099-12-31      <NA>
## 3          2017-01-31      2099-12-31      <NA>
## 4          1970-01-01      2099-12-31      <NA>
## 5          2014-01-31      2099-12-31      <NA>
## 6          1970-01-01      2099-12-31      <NA>

```

The ancestors are derived for those mCode concepts that excluded descendants.

```

snomed_omop_library2b <- snomed_omop_library1 %>% filter(!is.na(ancestors_of))
snomed_omop_library2b2 <- join_for_ancestors(kind = "LEFT", data = snomed_omop_library2b,
  descendant_id_column = "concept_id") %>% filter(min_levels_of_separation !=
  0) %>% select(all_of(colnames(snomed_library)), starts_with("ancestor_")) %>%
  rename_all(~str_remove_all(., pattern = "^ancestor_"))

```

```
head(snomed_omop_library2b2)
```

```

##          value_set_name code_system
## 1 HistologyMorphologyBehaviorVS    SNOMED CT

```

```
## 2 HistologyMorphologyBehaviorVS SNOMED CT
## 3 HistologyMorphologyBehaviorVS SNOMED CT
## 4 HistologyMorphologyBehaviorVS SNOMED CT
## 5 HistologyMorphologyBehaviorVS SNOMED CT
## 6 HistologyMorphologyBehaviorVS SNOMED CT
##
## 1 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## 2 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## 3 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## 4 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## 5 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## 6 excludes codes descending from 399983006 | Papillary neoplasm, pancreatobiliary-type, with high gr
## ancestors_of descendants_of code code_description all_codes concept_id
## 1 399983006 <NA> <NA> <NA> 399983006 4019483
## 2 399983006 <NA> <NA> <NA> 399983006 4030314
## 3 399983006 <NA> <NA> <NA> 399983006 4042149
## 4 399983006 <NA> <NA> <NA> 399983006 4043350
## 5 399983006 <NA> <NA> <NA> 399983006 4048384
## 6 399983006 <NA> <NA> <NA> 399983006 4133294
## concept_name domain_id vocabulary_id
## 1 Adenoma AND/OR adenocarcinoma Observation SNOMED
## 2 Neoplasm Observation SNOMED
## 3 Epithelial neoplasm Observation SNOMED
## 4 Morphologically altered structure Observation SNOMED
## 5 Body structure Spec Anatomic Site SNOMED
## 6 In situ neoplasm Observation SNOMED
## concept_class_id standard_concept concept_code valid_start_date
## 1 Morph Abnormality S 115215004 1970-01-01
## 2 Morph Abnormality S 108369006 1970-01-01
## 3 Morph Abnormality S 118285006 1970-01-01
## 4 Morph Abnormality S 118956008 1970-01-01
## 5 Body Structure S 123037004 1970-01-01
## 6 Morph Abnormality S 127569003 1970-01-01
## valid_end_date invalid_reason
## 1 2099-12-31 <NA>
## 2 2099-12-31 <NA>
## 3 2099-12-31 <NA>
## 4 2099-12-31 <NA>
## 5 2099-12-31 <NA>
## 6 2099-12-31 <NA>
```

The concepts that are explicitly stated do not require any additional derivation.

```
snomed_omop_library2c <- snomed_omop_library1 %>% filter(is.na(descendants_of),
  is.na(ancestors_of))
head(snomed_omop_library2c)
```

```
## # A tibble: 6 x 18
## value_set_name code_system logical_definit~ ancestors_of descendants_of code
## <chr> <chr> <chr> <chr> <chr> <chr>
## 1 CancerDisease~ SNOMED CT <NA> <NA> <NA> 3636~
## 2 CancerDisease~ SNOMED CT <NA> <NA> <NA> 2524~
## 3 CancerDisease~ SNOMED CT <NA> <NA> <NA> 7110~
## 4 CancerDisease~ SNOMED CT <NA> <NA> <NA> 5880~
## 5 CancerDisease~ SNOMED CT <NA> <NA> <NA> 2507~
```

```
## 6 CancerDisease~ SNOMED CT      <NA>          <NA>          <NA>          3863~
## # ... with 12 more variables: code_description <chr>, all_codes <chr>,
## #   concept_id <dbl>, concept_name <chr>, domain_id <chr>, vocabulary_id <chr>,
## #   concept_class_id <chr>, standard_concept <chr>, concept_code <chr>,
## #   valid_start_date <date>, valid_end_date <date>, invalid_reason <chr>
```

All 3 subsets are recombined before writing to file.

```
snomed_omop_library <- bind_rows(snomed_omop_library2a2, snomed_omop_library2b2,
  snomed_omop_library2c)
head(snomed_omop_library)
```

```
##           value_set_name code_system
## 1 CancerBodyLocationVS    SNOMED CT
## 2 CancerBodyLocationVS    SNOMED CT
## 3 CancerBodyLocationVS    SNOMED CT
## 4 CancerBodyLocationVS    SNOMED CT
## 5 CancerBodyLocationVS    SNOMED CT
## 6 CancerBodyLocationVS    SNOMED CT
##                                     logical_definition ancestors_of
## 1 includes codes descending from 123037004 | Body Structure      <NA>
## 2 includes codes descending from 123037004 | Body Structure      <NA>
## 3 includes codes descending from 123037004 | Body Structure      <NA>
## 4 includes codes descending from 123037004 | Body Structure      <NA>
## 5 includes codes descending from 123037004 | Body Structure      <NA>
## 6 includes codes descending from 123037004 | Body Structure      <NA>
## descendants_of code code_description all_codes concept_id
## 1      123037004 <NA>                <NA> 123037004   4048384
## 2      123037004 <NA>                <NA> 123037004   4002852
## 3      123037004 <NA>                <NA> 123037004   36717763
## 4      123037004 <NA>                <NA> 123037004   4230944
## 5      123037004 <NA>                <NA> 123037004   42605189
## 6      123037004 <NA>                <NA> 123037004   4097829
##                                     concept_name      domain_id
## 1                                     Body structure Spec Anatomic Site
## 2                                     Buccal embrasure Spec Anatomic Site
## 3                                     Skin structure of left lower eyelid Spec Anatomic Site
## 4                                     Adenofibrosis      Observation
## 5 Intervertebral foramen of eighteenth thoracic vertebra Spec Anatomic Site
## 6                                     Transitional cell carcinoma      Observation
## vocabulary_id concept_class_id standard_concept concept_code
## 1      SNOMED      Body Structure      S      123037004
## 2      SNOMED      Body Structure      S      110326006
## 3      SNOMED      Body Structure      S      719884003
## 4      SNOMED Morph Abnormality      S      89115006
## 5 SNOMED Veterinary      Body Structure      S 336621000009107
## 6      SNOMED Morph Abnormality      S      27090000
## valid_start_date valid_end_date invalid_reason
## 1      1970-01-01      2099-12-31      <NA>
## 2      1970-01-01      2099-12-31      <NA>
## 3      2017-01-31      2099-12-31      <NA>
## 4      1970-01-01      2099-12-31      <NA>
## 5      2014-01-31      2099-12-31      <NA>
## 6      1970-01-01      2099-12-31      <NA>
```

```

file <- file.path(getwd(), "data-raw", "snomed.csv")
if (!file.exists(file)) {
  write_csv(x = snomed_omop_library, file = file)
}

```

LOINC

```

loinc_library <- value_sets_by_vocab$LOINC %>% rubix::format_colnames() %>%
  mutate_all(as.character) %>% mutate_all(trimws) %>% distinct()
head(loinc_library)

```

```

## # A tibble: 6 x 5
##   value_set_name code_system logical_definiti~ code code_description
##   <chr>          <chr>      <chr>          <chr> <chr>
## 1 TumorMarkerTes~ LOINC      <NA>          1695~ 5-Hydroxyindoleacetate [M~
## 2 TumorMarkerTes~ LOINC      <NA>          3120~ 5-Hydroxyindoleacetate [M~
## 3 TumorMarkerTes~ LOINC      <NA>          1692~ 5-Hydroxyindoleacetate [M~
## 4 TumorMarkerTes~ LOINC      <NA>          1693~ 5-Hydroxyindoleacetate [M~
## 5 TumorMarkerTes~ LOINC      <NA>          1694~ 5-Hydroxyindoleacetate [M~
## 6 TumorMarkerTes~ LOINC      <NA>          7282~ 5-Hydroxyindoleacetate [M~

```

```

loinc_omop_library <- chariot::join_on_concept_code(kind = "LEFT",
  data = loinc_library, column = "code", where_in_concept_field = "vocabulary_id",
  where_in_concept_field_value = "LOINC")
loinc_omop_library <- loinc_library %>% left_join(loinc_omop_library,
  by = c("value_set_name", "code_system", "logical_definition",
    "code", "code_description")) %>% distinct()

```

```
head(loinc_omop_library)
```

```

## # A tibble: 6 x 15
##   value_set_name code_system logical_definit~ code code_description concept_id
##   <chr>          <chr>      <chr>          <chr> <chr>          <dbl>
## 1 TumorMarkerTe~ LOINC      <NA>          1695~ 5-Hydroxyindole~    3005148
## 2 TumorMarkerTe~ LOINC      <NA>          3120~ 5-Hydroxyindole~    3023028
## 3 TumorMarkerTe~ LOINC      <NA>          1692~ 5-Hydroxyindole~    3014670
## 4 TumorMarkerTe~ LOINC      <NA>          1693~ 5-Hydroxyindole~    3021106
## 5 TumorMarkerTe~ LOINC      <NA>          1694~ 5-Hydroxyindole~    3021385
## 6 TumorMarkerTe~ LOINC      <NA>          7282~ 5-Hydroxyindole~    43055680
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## #   vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## #   concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## #   invalid_reason <chr>

```

```

file <- file.path(getwd(), "data-raw", "loinc.csv")
if (!file.exists(file)) {
  write_csv(x = loinc_omop_library, file = file)
}

```

ICD-10 CM


```
icd10cm_library <- value_sets_by_vocab$`ICD-10-CM` %>% rubix::format_colnames() %>%
  mutate_all(as.character) %>% mutate_all(trimws) %>% mutate(code = str_replace_all(string = code,
    pattern = "(^[A-Z]{1}[0-9A-Z]{2}) ([0-9A-Z]{1}.*)$", replacement = "\\1.\\2")) %>%
  distinct()
head(icd10cm_library)
```

```
## # A tibble: 6 x 5
##   value_set_name code_system logical_definiti~ code code_description
##   <chr>          <chr>      <chr>          <chr> <chr>
## 1 CancerDisorde~ ICD-10-CM <NA>          C00.0 Malignant neoplasm of ext~
## 2 CancerDisorde~ ICD-10-CM <NA>          C00.1 Malignant neoplasm of ext~
## 3 CancerDisorde~ ICD-10-CM <NA>          C00.2 Malignant neoplasm of ext~
## 4 CancerDisorde~ ICD-10-CM <NA>          C00.3 Malignant neoplasm of upp~
## 5 CancerDisorde~ ICD-10-CM <NA>          C00.4 Malignant neoplasm of low~
## 6 CancerDisorde~ ICD-10-CM <NA>          C00.5 Malignant neoplasm of lip~
```

```
icd10cm_omop_library <- chariot::join_on_concept_code(kind = "LEFT",
  data = icd10cm_library, column = "code", where_in_concept_field = "vocabulary_id",
  where_in_concept_field_value = "ICD10CM")
icd10cm_omop_library <- icd10cm_library %>% left_join(icd10cm_omop_library,
  by = c("value_set_name", "code_system", "logical_definition",
    "code", "code_description")) %>% distinct()
```

```
head(icd10cm_omop_library)
```

```
## # A tibble: 6 x 15
##   value_set_name code_system logical_definit~ code code_description concept_id
##   <chr>          <chr>      <chr>          <chr> <chr>          <dbl>
## 1 CancerDisorde~ ICD-10-CM <NA>          C00.0 Malignant neopl~ 35206047
## 2 CancerDisorde~ ICD-10-CM <NA>          C00.1 Malignant neopl~ 35206048
## 3 CancerDisorde~ ICD-10-CM <NA>          C00.2 Malignant neopl~ 35206049
## 4 CancerDisorde~ ICD-10-CM <NA>          C00.3 Malignant neopl~ 35206050
## 5 CancerDisorde~ ICD-10-CM <NA>          C00.4 Malignant neopl~ 35206051
## 6 CancerDisorde~ ICD-10-CM <NA>          C00.5 Malignant neopl~ 35206052
## # ... with 9 more variables: concept_name <chr>, domain_id <chr>,
## #   vocabulary_id <chr>, concept_class_id <chr>, standard_concept <chr>,
## #   concept_code <chr>, valid_start_date <date>, valid_end_date <date>,
## #   invalid_reason <chr>
```

```
file <- file.path(getwd(), "data-raw", "icd10cm.csv")
if (!file.exists(file)) {
  write_csv(x = icd10cm_omop_library, file = file)
}
```

```
chariot::dcAthena(conn = conn)
```