Working with Concept Sets in Capr

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library	y(Capr)	

In version 2 of Capr, we have introduced a whole new way of building concept sets. Some highlights of this new system are, we no longer need a database connection, only need OMOP concept ids, and we provide functionality to improve how we map to the vocabulary (i.e. find descendants or exclude). The new interface makes it much easier to build and read information about a concept set. In this vignette we will demonstrate how the new AI works!

0.1 Defining a concept set

The new function cs is a binder function that collects integer OMOP ids into a set. The idea is for the function to look like c() which in R is use to formulate vectors. Say we found several ingredients that are all ace inhibitors. We can combine those OMOP concept ids into a set as follows:

```
ace1 <- cs(1335471, #benazepril
            1340128, #captopril
            1341927, #enalapril
            1308216, #lisinopril
            1363749, #fosinopril
            name = "aceInhibitors")
#> -- <Capr Concept Set> aceInhibitors -
#> # A tibble: 5 x 9
     conceptId conceptCode conceptName domainId vocabularyId standardConcept includeDescendants
#>
          <int> <chr>
                              <chr>>
                                           <chr>
                                                      <chr>
                                                                    <chr>
                                                                                      <lql>
        1335471 ""
                              11 11
#> 1
                                                                                      FALSE
                              11 11
                                           11 11
                                                      11 11
                                                                    11 11
       1340128 ""
                                                                                      FALSE
                              11 11
       1341927 ""
#> 3
                                                                                      FALSE
                                                      11 11
#> 4
       1308216 ""
                                                                    11 11
                                                                                      FALSE
                              11 11
                                            11 11
       1363749 ""
                                                                                      FALSE
#> # i 2 more variables: isExcluded <lql>, includeMapped <lql>
```

0.2 Including Descendants

If you were only add the OMOP id for the ingredient, our query would only look at whether that specific ID is among the drug concepts for a set of patients. With OMOP, we often don't want just the ingredient ID but also its descendants. Descendants are child concepts that map to the adult concepts. For example a brand name or specific dosage of lisinopril would have a different OMOP concept Id but it would map to the ingredient term via the vocabulary hierarchy. This helps us include all variations of linisopril that could be seen in the data. In Capr we can easily add this logic using the descendants command

```
ace2 <- cs(descendants(1335471, 1340128, 1341927, 1363749, 1308216),
            name = "aceInhibitors")
ace2
#> -- <Capr Concept Set> aceInhibitors -----
#> # A tibble: 5 x 9
     concept Id\ concept Code\ concept Name\ domain Id\ vocabulary Id\ standard Concept\ include Descendants
#>
                             <chr>
                                                                   <chr>
          <int> <chr>
                                           <chr>
                                                    <chr>
                                                                                     <lgl>
#> 1
                             11 11
                                           11 11
                                                     11 11
                                                                   11 11
      1335471 ""
                                                                                     TRUE
      1340128 ""
                             11 11
                                           11 11
                                                                   11 11
                                                                                     TRUE
#> 2
                              11 11
                                           11 11
                                                     11 11
                                                                   11 11
#> 3
      1341927 ""
                                                                                     TRUE
#> 4 1363749 ""
                                                                                     TRUE
#> 5 1308216 ""
                              11 11
                                           11 11
                                                     11 11
                                                                                     TRUE
#> # i 2 more variables: isExcluded <lgl>, includeMapped <lgl>
```

0.3 Toggling other logic

A similar strategy can be used to exclude concepts from a set or include mapped. We can also combine this within the cs function, as shown below

```
ace3 <- cs(descendants(1335471, 1340128, 1341927, 1363749), exclude(1308216),
            name = "aceInhibitors")
#> -- <Capr Concept Set> aceInhibitors -----
#> # A tibble: 5 x 9
     conceptId conceptCode conceptName domainId vocabularyId standardConcept includeDescendants
#>
          <int> <chr>
                              \langle chr \rangle
                                             <chr>
                                                       <chr>
                                                                      <chr>
                                                                                         <lql>
                               11 11
                                                                      11 11
#> 1
       1335471 ""
                                                                                         TRUE
                               11 11
                                             11 11
                                                                      11 11
      1340128 ""
                                                       11 11
                                                                                         TRUE
#> 3 1341927 ""
                               _{II} _{II}
                                             11 11
                                                        11 11
                                                                       11 11
                                                                                         TRUE
                               11 11
                                                        11 11
                                                                       11 11
#> 4 1363749 ""
                                                                                         TRUE
                               \eta \eta
                                             _{II} _{II}
#> 5 1308216 ""
                                                                                         FALSE
#> # i 2 more variables: isExcluded <lql>, includeMapped <lql>
```

0.4 Fill in missing concept set details

By default, Capr puts in minimal information for the concept set, the concept id. However it is often helpful to fill out the remaining information of the concept using the OMOP vocabularies. To fill out the remaining information, a user needs to connect to a OMOP CDM to access the vocabulary information for the concept in the concept set. First we show a regular concept set, and the json it renders.

```
diclofenac <- cs(descendants(1124300), name = "diclofenac")
cat(as.json(diclofenac))</pre>
```

Now we add in the CDM connection to get the remaining information:

```
con <- DatabaseConnector::connect(Eunomia::getEunomiaConnectionDetails())
diclofenac <- getConceptSetDetails(diclofenac, con, vocabularyDatabaseSchema = "main")
cat(as.json(diclofenac))</pre>
```