Using reticulate to access CPTAC data in R

reticulate is an R package that allows you to create Python objects and convert them into R objects for use in your R environment.

Thus, even though cptac is a Python package, with the help of reticulate you can load and work with the datasets in R.

Environment setup

First, you need to load the reticulate package, and then tell it which Python environment you want to use to access the cptac package.

```
# Install reticulate if necessary
if (!require(reticulate)) install.packages("reticulate")

# Load the package
library(reticulate)

# Specify to use the environment where the cptac package is installed. Replace
# "dev" with the name of your environment. If you use an environment manager
# besides conda, you'll want this command: use_virtualenv("myenv")
use_condaenv("dev", required = TRUE)
```

Load cptac and access data

Now we will load cptac and get the dataset we want.

```
# Import the package. We pass convert = FALSE so that objects won't be
# converted from Python to R until we explicitly ask for it. This is necessary
# to properly prepare a multiindex dataframe for conversion to R (see below).
cptac <- import("cptac", convert = FALSE)

# Load the dataset
en <- cptac$Endometrial()</pre>
```

Load a dataframe and convert it to an R object

This dataframe just has a regular column index (not a multiindex), so we can directly convert it into an R object after loading it.

When reticulate converts a pandas.DataFrame into R, it converts it to the R data.frame type.

```
# Load the table
prot_py <- en$get_proteomics()

# Convert into R
prot <- py_to_r(prot_py)
print(prot[1:10, 1:8])</pre>
```

```
##
               A1BG
                       A2M A2ML1
                                   A4GALT
                                             AAAS
                                                     AACS
                                                            AADAT
                                                                    AAED1
## C3L-00006 -1.180 -0.8630 -0.802
                                   0.2220 0.2560
                                                   0.6650 1.2800 -0.3390
## C3L-00008 -0.685 -1.0700 -0.684
                                                                   0.1390
                                   0.9840
                                          0.1350
                                                   0.3340 1.3000
## C3L-00032 -0.528 -1.3200 0.435
                                      NaN -0.2400
                                                   1.0400 -0.0213 -0.0479
## C3L-00090 -1.670 -1.1900 -0.443
                                   0.2430 -0.0993
                                                   0.7570
                                                           0.7400 -0.9290
## C3L-00098 -0.374 -0.0206 -0.537
                                   0.3110 0.3750
                                                   0.0131 -1.1000
## C3L-00136 -1.080 -0.7080 -0.126 -0.4260 -0.1140 -0.1110 0.8950
## C3L-00137 -1.320 -0.7080 -0.808 -0.0709 0.1380
                                                   0.6560 -0.2800 -0.1280
## C3L-00139 -0.467 0.3700 -0.339
                                      NaN
                                           0.4340
                                                   0.0358 -0.1750
                                                                   0.1810
## C3L-00143 -1.120 -1.3100 0.912 0.4180 -0.0768
                                                   0.8460 -0.1210
## C3L-00145 -0.716 -0.8850
                            2.820 -0.3430 0.1470
                                                   0.4450 -0.0565 -0.8380
```

Load a multiindex dataframe

If it's a dataframe with a column multiindex, you need to first flatten the index before converting it to an R object.

See tutorial 4 for more information on column multiindexes.

```
# Load the table
phos_py_multiindex <- en$get_phosphoproteomics()

# Load cptac.utils so we can access a helper function to convert the multiindex
# to a single level index.
utils <- import("cptac.utils", convert = FALSE)

# Convert the multiindex to a single level index. This works because we passed
# convert = FALSE when we imported the cptac package, so the dataframe is still
# a Python object at this point (it wasn't automatically converted into an R
# object on loading) and can be passed to this Python function.
phos_py_single_index <- utils$reduce_multiindex(phos_py_multiindex, flatten = TRUE)

# Convert to R
phos <- py_to_r(phos_py_single_index)

print(phos[1:10, 1:6])</pre>
```

```
##
              AAAS S495 AAAS S541 AAAS Y485 AACS S618 AAED1 S12 AAGAB S310
## C3L-00006
                                                  -0.881
                                                            -1.8100
                                                                            NaN
                    NaN
                               NaN
                                          NaN
## C3L-00008
                    NaN
                               NaN
                                          NaN
                                                     NaN
                                                             0.0840
                                                                            NaN
## C3L-00032
                 -0.202
                               NaN
                                                     NaN
                                                            -1.8800
                                                                            NaN
                                          NaN
## C3L-00090
                 -0.002
                               NaN
                                       -0.407
                                                     NaN
                                                                NaN
                                                                            NaN
## C3L-00098
                  0.556
                           -0.0461
                                                                            NaN
                                          NaN
                                                     NaN
                                                             0.9410
## C3L-00136
                    NaN
                               NaN
                                          NaN
                                                     NaN
                                                             0.0796
                                                                            NaN
## C3L-00137
                  0.300
                               NaN
                                          NaN
                                                  -0.371
                                                            -1.1400
                                                                            NaN
## C3L-00139
                  0.490
                               NaN
                                          NaN
                                                     NaN
                                                                {\tt NaN}
                                                                            NaN
## C3L-00143
                 -0.231
                            0.0108
                                          NaN
                                                             0.2420
                                                                            NaN
                                                     NaN
## C3L-00145
                  0.268
                               NaN
                                          NaN
                                                     NaN
                                                            -0.1120
                                                                            NaN
```