## **TargetSearchData**

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Example GC-MS data for TargetSearch Package

### Description

A TargetSearch example GC-MS data. This package contains raw NetCDF files from a E.coli salt stress experiment, extracted peak list of each NetCDF file and three tab-delimted text files: a sample description, a reference library and a retention index marker definition. The data is a subset of the original data from 200-400 seconds and 85-320 m/z.

#### Usage

data(TargetSearchData)

#### Format

The data contains the following objects:

sampleDescription a tsSample object. The sample description.

refLibrary a tsLib object. The reference library.

rimLimits a tsRim object. The RI markers definition.

**RImatrix** a matrix object. The retention time of the RI markers.

corRI a matrix object. The sample RI.

peakData a tsMSdata object. The intensities and RIs of all the masses that were searched for.

**metabProfile** a tsProfile object. The metabolite profile.

#### **Details**

All files are located in gc-ms-data subdirectory.

#### See Also

 ${\tt ImportLibrary, ImportSamples, ImportFameSettings,}$ 

2 TargetSearchData

#### **Examples**

```
require(TargetSearch)
## The directory with the NetCDF GC-MS files
cdfpath <- file.path(.find.package("TargetSearchData"), "gc-ms-data")</pre>
cdfpath
list.files(cdfpath)
samp.file <- file.path(cdfpath, "samples.txt")</pre>
rim.file <- file.path(cdfpath, "rimLimits.txt")</pre>
lib.file <- file.path(cdfpath, "library.txt")</pre>
# import files from package
sampleDescription <- ImportSamples(samp.file, CDFpath = cdfpath, RIpath = ".")</pre>
               <- ImportLibrary(lib.file)</pre>
refLibrary
                 <- ImportFameSettings(rim.file, mass = 87)
rimLimits
# perform RI correction
RImatrix
                  <- RIcorrect(sampleDescription, rimLimits, massRange = c(85,320),</pre>
                   IntThreshold = 25, pp.method = "ppc", Window = 15)
# update median RI
refLibrary
                  <- medianRILib(sampleDescription, refLibrary)</pre>
# get the sample RI
corRI
                  <- sampleRI(sampleDescription, refLibrary, r_thres = 0.95)
\sharp obtain the peak Intensities of all the masses in the library
peakData <- peakFind(sampleDescription, refLibrary, corRI)</pre>
# make a profile of the metabolite data
metabProfile
              <- Profile(sampleDescription, refLibrary, peakData, r_thres = 0.95)</pre>
# show the metabolite profile
profileInfo(metabProfile)
# show the matrix intensities
Intensity(metabProfile)
```

# **Index**

```
*Topic datasets
    TargetSearchData, 1
.required(TargetSearchData), 1

corRI(TargetSearchData), 1

ImportFameSettings, I
ImportLibrary, I
ImportSamples, I

metabProfile(TargetSearchData), 1

peakData(TargetSearchData), 1

refLibrary(TargetSearchData), 1

RImatrix(TargetSearchData), 1

rimLimits(TargetSearchData), 1

sampleDescription
    (TargetSearchData), 1

TargetSearchData, 1
```