Package 'PtH2O2lipids'

May 16, 2024

Title P. tricornutum HPLC-ESI-MS Lipid Data from van Creveld et al. (2015)

Version 1.31.0 **Date** 2016-07-06

Depends R (>= 3.3), xcms, CAMERA, LOBSTAHS, methods, utils

Suggests gplots, RColorBrewer, cluster, vegan

Description Annotated HPLC-ESI-MS lipid data in positive ionization mode from an experiment in which cultures of the marine diatom Phaeodactylum tricornutum were treated with various concentrations of hydrogen peroxide (H2O2) to induce oxidative stress. The experiment is described in Graff van Creveld, et al., 2015, ``Early perturbation in mitochondria redox homeostasis in response to environmental stress predicts cell fate in diatoms," ISME Journal 9:385-395. PtH2O2lipids consists of two objects: A CAMERA xsAnnotate object (ptH2O2lipids\$xsAnnotate) and LOBSTAHS LOBSet object (ptH2O2lipids\$xsAnnotate\$LOBSet). The LOBSet includes putative compound assignments from the default LOBSTAHS database. Isomer annotation is recorded in three other LOBSet slots.

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URL http://dx.doi.org/10.1038/ismej.2014.136,
 https://github.com/vanmooylipidomics/PtH202lipids,
 http://www.whoi.edu/page.do?pid=133616&tid=282&cid=192529

BugReports https://github.com/vanmooylipidomics/PtH2O2lipids/issues/new

biocViews ReproducibleResearch, CellCulture, MassSpectrometryData, Phaeodactylum_tricornutum_data

NeedsCompilation no

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ptH2O2lipids

```
git_url https://git.bioconductor.org/packages/PtH2O2lipids
git_branch devel
git_last_commit 30faaa8
git_last_commit_date 2024-04-30
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```

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Description

Positive ionization mode HPLC-ESI-MS lipid data from an experiment in which cultures of the marine diatom *Phaeodactylum tricornutum* were treated with various concentrations of hydrogen peroxide (H2O2) to induce oxidative stress. ptH2O2lipids\$LOBSet was generated from ptH2O2lipids\$xsAnnotate using the LOBSTAHS function doLOBscreen.

Usage

```
data(ptH202lipids)
```

Format

A list object containing the lipid data in two forms:

ptH202lipids\$L0BSet A 1.2 MB object of formal class "L0BSet" containing screened peak data to which compound assignments and isomer identifications have been made. The structure of ptH202lipids\$L0BSet is:

```
Formal class 'LOBSet' [package "LOBSTAHS"] with 9 slots
 ..@ peakdata
                          :'data.frame': 2056 obs. of 54 variables
 ..@ iso_C3r
                          :List of 2056
 ..@ iso_C3f
                          :List of 2056
 ..@ iso_C3c
                          :List of 2056
  ..@ LOBscreen_diagnostics:'data.frame': 6 obs. of 4 variables:
 ..@ LOBisoID_diagnostics :'data.frame': 3 obs. of 2 variables:
 ..@ LOBscreen_settings
                         :List of 6
 ....$ database : chr "default"
 ....$ remove.iso : logi TRUE
 ....$ rt.restrict : logi TRUE
```

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```
....$ rt.windows : chr "default"
....$ exclude.oddFA: logi TRUE
....$ match.ppm : num 2.5
...@ polarity : Factor w/ 1 level "positive": 1
...@ sampnames : chr [1:16] "0uM_24h_Orbi_0468" "0uM_24h_Orbi_0473" "0uM_4h_Orbi_0476" "0uM_24h_Orbi_0476" "0uM_24h_Orbi_0478" "0uM_4h_Orbi_0476" "0uM_24h_Orbi_0478" "0uM_4h_Orbi_0478" "0uM_4h_All "0uM_4h_Orbi_0478" "0uM_4h_All "0uM_4h_4
```

ptH202lipids\$xsAnnotate An 80 MB object of formal class "xsAnnotate" containing 18,314 peakgroups in 5,080 pseudospectra. This is the object from which ptH202lipids\$L0BSet was created using doL0Bscreen. It includes annotation of possible isotope peaks from findIsotopes. The xcmsSet from which the xsAnnotate object was created (64.5 MB) can be accessed at ptH202lipids\$xsAnnotate@xcmsSet.

Details

ptH202lipids\$LOBSet includes compound identifications assigned from the default LOBSTAHS positive mode database. ptH202lipids\$LOBSet also includes in the slots iso_C3r, iso_C3f, and iso_C3c the various possible isomers identified for each compound. Note that all other slots in the ptH2O2lipids object can be accessed using the accessor functions described for the "LOBSet-class" object class.

The dataset contains peaks from 16 samples that span three H2O2 treatments (0, 30 and 150 μ M) and three timepoints (+4, +8, and +24 hours) in duplicate. The dataset contains only one replicate sample for the 0 and 150 μ M treatments at +4h.

The mzXML files and Thermo .raw files from which these objects are derived can be accessed at https://github.com/vanmooylipidomics/PtH202lipids/tree/master/mzXML and http://www.whoi.edu/page.do?pid=133616&tid=282&cid=192529, respectively.

Users should note that the LOBSet in this package does not include any PUA (polyunsaturated aldehyde) identifications.

Source

http://www.nature.com/ismej/journal/v9/n2/full/ismej2014136a.html

References

Collins, J.R., B.R. Edwards, H.F. Fredricks, and B.A.S. Van Mooy. 2016. LOBSTAHS: An adduct-based lipidomics strategy for discovery and identification of oxidative stress biomarkers. Analytical Chemistry.

Graff van Creveld, et al., 2015, "Early perturbation in mitochondria redox homeostasis in response to environmental stress predicts cell fate in diatoms", *ISME Journal* 9:385-395

See Also

LOBSet-class, LOBSet, doLOBscreen, getLOBpeaklist, xcmsSet, xsAnnotate

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Examples

```
## generate the object in ptH202lipids$LOBSet using ptH202lipids$xsAnnotate as
## input
library(PtH202lipids)

## yields output identical to ptH202lipids$LOBSet
myPtH202LOBSet = doLOBscreen(ptH202lipids$xsAnnotate, polarity = "positive",
database = NULL, remove.iso = TRUE, rt.restrict = TRUE, rt.windows = NULL,
exclude.oddFA = TRUE, match.ppm = 2.5)

## access xsAnnotate object
ptH202lipids$xsAnnotate

## access xcmsSet
ptH202lipids$xsAnnotate@xcmsSet
```

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