Package 'msconverteR'

May 24, 2019

- ·
Type Package
Title Convert vendor specific mass spectrometry files to mzML
Version 0.1.0
Description Use the pwiz docker container to convert vendor specific mass spectrometry files to .mzML from within the R console on Linux systems.
License GPL (>= 3)
Imports stevedore, stringr, crayon, clisymbols
Encoding UTF-8
LazyData true
RoxygenNote 6.1.1
Roxygen list(markdown = TRUE)
NeedsCompilation no
Author Tom Wilson [aut, cre]
Maintainer Tom Wilson <tpw2@aber.ac.uk></tpw2@aber.ac.uk>
R topics documented:
convert_files
Index
convert_files

Description

Convert vendor specific mass spectrometry files to the open .mzML format. This functions makes a system call to docker in order to convert files. Conversion arguments should be supplied to the args parameter exactly as they would be for msconvert but omitting the --filter prefix.

2 get_pwiz_container

Usage

```
convert_files(files, args = c("peakPicking true1-", "polarity positive"))
```

Arguments

files the absolute filepath vendor specific files to be converted

args a character vector of msconvert arguments.

Details

```
For example to convert a file with vendor specific centroiding only;

convert_files(rawFiels,args = 'peakPicking true1-')

To only retain positive mode data;

convert_files(rawFiels,args = c('peakPicking true1-','polarity positive'))
```

get_pwiz_container

Get Proteowizard (pwiz) Docker Container

Description

Pull the pwiz docker container (chambm/pwiz-skyline-i-agree-to-the-vendor-licenses:latest) from Docker Hub.

Usage

```
get_pwiz_container()
```

Index

```
convert_files, 1
get_pwiz_container, 2
```