patRoon handbook

Rick Helmus

2022-01-13

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1 Introduction

Nowadays there are various software tools available to process data from non-target analysis (NTA) experiments. Individual tools such as ProteoWizard, XCMS, OpenMS, MetFrag and mass spectrometry vendor tools are often combined to perform a complete data processing workflow. During this workflow, raw data files may undergo pre-treatment (e.g. conversion), chromatographic and mass spectral data are combined to extract so called features (or 'peaks') and finally annotation is performed to elucidate chemical identities. The aim of patRoon is to harmonize the many available tools in order to provide a consistent user interface without the need to know all the details of each individual software tool and remove the need for tedious conversion of data when multiple tools are used. The name is derived from a Dutch word that means pattern and may also be an acronym for hyPhenated mAss specTROmetry nOn-target aNalysis. The workflow of non-target analysis is typically highly dependent on several factors such as the analytical instrumentation used and requirements of the study. For this reason, patRoon does not enforce a certain workflow. Instead, most workflow steps are optional, are highly configurable and algorithms can easily be mixed or even combined. Furthermore, patRoon supplies a straightforward interface to easily inspect, select, visualize and report all data that is generated during the workflow.

The documentation of patRoon consists of three parts:

- 1. A tutorial (accessible at here)
- 2. This handbook
- 3. The reference manual (accessible in R with ?`patRoon-package` or online here)

New users are highly recommended to start with the tutorial: this document provides an interactive introduction in performing a basic NTA processing workflow with patRoon. The handbook provides a more thorough overview of all concepts, functionalities and provides instructions and many examples on working with patRoon. Finally, the reference manual provides all the gritty details for all functionalities, and is meant if you want to know more details or need a quick reminder how a function should be used.

2 Installation

patRoon depends on various other software tools to perform the non-target analysis workflow steps and to implement various other functionality. Most of these dependencies are automatically installed when you install the patRoon R package, however, some may need to be manually installed and/or configured.

NOTE It is highly recommended to perform installation steps in a 'clean' R session to avoid errors when installing or upgrading packages. As such it is recommended to close all open (R Studio) sessions and open a plain R console to perform the installation.

2.1 Automatic installation (Windows only)

An installation script is provided that automatically installs and configures all dependencies and finally installs patRoon itself. At this moment, this script only works with Microsoft Windows. You don't have to install anything else to use it, simply open R and execute these commands:

```
source("https://raw.githubusercontent.com/rickhelmus/patRoon/master/install_patRoon.R")
installPatRoon()
```

A simple text based wizard will start and asks you what to install and how to do it. You can re-run this installer at any time, for instance, if something went wrong or you want to install additional dependencies.

2.2 Docker image

Docker images are provided to easily install a reproducible environment with R, patRoon and nearly all of its dependencies. This section assumes you have a basic understanding of Docker and have it installed. If not, please refer to the many guides available on the Internet. The Docker images of patRoon were originally only used for automated testing, however, since these contain a complete working environment of patRoon they are also suitable for using the software. They come with all external dependencies (except ProteoWizard), R dependencies and MetFrag libraries. Furthermore, the Docker image also contains RStudio server, which makes using patRoon even easier.

Below are some example shell commands on how to run the image.

```
# run an interactive R console session
docker run --rm -it patroonorg/patroonrs

# run a linux shell, from which R can be launched
docker run --rm -it patroonorg/patroonrs bash
```

Note that the first two commands run as the default user rstudio, while the last two as root. The last commands launch RStudio server. You can access it by browsing to localhost:8787 and logging in with user rstudio and the password defined by the PASSWORD variable from the command (yourpasswordhere in the above example). The last command also links a local volume in order to obtain persistence of files in the container's home directory. The Docker image is based on the excellent work from the rocker project. For more information on RStudio related options see their documentation for the RStudio image.

2.3 Manual installation

The manual installation is for users who don't use Windows or Docker, prefer to do a manual installation or simply want to know what happens behind the scenes. The manual installation consists of three phases:

- 1. Installing some prerequisite R packages
- 2. Install and configure (non-R) dependencies
- 3. Install patRoon

2.3.1 R prerequisites

When installing patRoon Windows users have the option to install from a customized (miniCRAN) repository (patRoonDeps). This repository provides a central repository for patRoon and all its R packages. An advantage is that installation will be faster and you will not need to install Rtools. Note that you will need to have the latest R version installed in order to use this repository.

When you decide to use the patRoonDeps repository you can simply *skip* this step. **Otherwise** (i.e. you will use regular repositories instead), the following will install the R dependencies.

```
# NOTE: a fork is installed that fixes some issues with the latest R versions
remotes::install_github("BSchamberger/RDCOMClient")

# only when using the R interface (not recommended by default)
remotes::install_github("c-ruttkies/MetFragR/metfRag")

# needed for feature quality calculations with MetaClean
BiocManager::install(c("BiocStyle", "Rgraphviz"))
remotes::install_github("KelseyChetnik/MetaClean")
```

Note that only the CAMERA installation is mandatory, the rest involves installation of *optional* packages. If you are unsure which you need then you can always install the packages at a later stage.

2.3.2 Other dependencies

Depending on which functionality is used, the following external dependencies may need to be installed:

Software	Remarks
Java JDK	Mandatory for e.g. plotting structures and using MetFrag.
Rtools	Necessary on Window and when $\mathtt{patRoon}$ is not installed from
D + 1177 1	patRoonDeps.
ProteoWizard	Needed for automatic data-pretreatment (e.g. data file
	conversion and centroiding, Bruker users may use
	DataAnalysis integration instead).
OpenMS	Recommended. Used for e.g. finding and grouping features.
MetFrag CL	Recommended. Used for annotation with MetFrag.
MetFrag CompTox DB	Database files necessary for usage of the CompTox database
	with MetFrag. Note that a recent version of MetFrag
	(>=2.4.5) is required. Note that the lists with additions for
	smoking metadata and wastewater metadata are also
	supported.
MetFrag PubChemLite DB	Database file needed to use PubChemLite with MetFrag.
SIRIUS	For obtaining feature data and formula and/or compound
	annotation.
BioTransformer	For prediction of transformation products. See the
	BioTransformer page for installation details. If you have
	trouble compiling the jar file you can download it from here.
SAFD	For finding features with SAFD. Please follow all the
	installation on the SAFD webpage.
OpenBabel	Used in some cases for suspect screening (e.g. to calculate
	molecular masses for suspects with only InChI information).
	Otherwise optional.
pngquant	Used to reduce size of HTML reports, definitely optional.

Most of these packages are optional and only needed if their algorithms are used during the workflow. If you are unsure which are needed, simply skip them for now and install them at a later stage.

After installation of these dependencies you may need to configure their filepaths (OpenMS and ProteoWizard are usually found automatically). To configure the file locations you should set some global package options with the options () R function, for instance:

```
options(patRoon.path.pwiz = "C:/ProteoWizard") # location of ProteoWizard installation

→ folder

options(patRoon.path.SIRIUS = "C:/sirius-win64-3.5.1") # location where SIRIUS was

→ extracted

options(patRoon.path.OpenMS = "/usr/local/bin") # directory with the OpenMS binaries

options(patRoon.path.pngquant = "~/pngquant") # directory containing pngquant binary

options(patRoon.path.MetFragCL = "~/MetFrag2.4.5-CL.jar") # full location to the jar file

options(patRoon.path.MetFragCompTox = "C:/CompTox_17March2019_SelectMetaData.csv") # full

→ location to desired CompTox CSV file

options(patRoon.path.MetFragPubChemLite = "~/PubChemLite_14Jan2020_tier0.csv") # full

→ location to desired PubChemLite CSV file

options(patRoon.path.BioTransformer = "~/biotransformer/biotransformer-3.0.1.jar")

options(patRoon.path.obabel = "C:/Program Files/OpenBabel-3.0.0") # directory with

→ OpenBabel binaries
```

These commands have to be executed everytime you start a new R session (e.g. as part of your script). However, it is probably easier to add them to your ~/.Rprofile file so that they are executed automatically when you start R. If you don't have this file yet you can simply create it yourself (for more information see e.g. this SO answer).

2.3.3 patRoon installation

Finally, it is time to install patRoon itself. As mentioned before, Windows users (who have the latest R version) can install patRoon and all its package dependencies from the patRoonDeps repository:

Otherwise, installation occurs directly from GitHub:

```
remotes::install_github("rickhelmus/patRoon")
```

Optional example data is installed via GitHub:

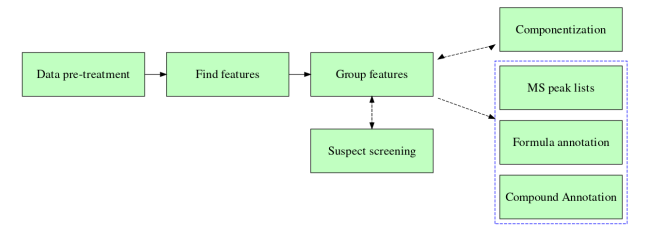
```
# optional example data
remotes::install_github("rickhelmus/patRoonData")
```

Afterwards, you can run the verifyDependencies() function to see if patRoon can find all its dependencies (you may need to restart R beforehand):

```
patRoon::verifyDependencies()
```

3 Workflow concepts

In a non-target workflow both chromatographic and mass spectral data is automatically processed in order to provide a comprehensive chemical characterization of your samples. While the exact workflow is typically dependent on the type of study, it generally involves of the following steps:



Note that patRoon supports flexible composition of workflows. In the scheme above you can recognize optional steps by a *dashed line*. The inclusion of each step is only necessary if a further steps depends on its data. For instance, annotation and componentization do not depend on each other and can therefore be executed in any order or simply be omitted. A brief description of all steps is given below.

During data pre-treatment raw MS data is prepared for further analysis. A common need for this step is to convert the data to an open format so that other tools are able to process it. Other pre-treatment steps may involve re-calibration of m/z data or performing advanced filtering operations.

The next step is to extract **features** from the data. While different terminologies are used, a feature in patRoon refers to a single chromatographic peak in an extracted ion chromatogram for a single m/z value (within a defined tolerance). Hence, a feature contains both chromatographic data (e.g. retention time and peak height) and mass spectral data (e.g. the accurate m/z). Note that with mass spectrometry multiple m/z values may be detected for a single compound as a result of adduct formation, natural isotopes and/or insource fragments. Some algorithms may try to combine these different masses in a single feature. However, in patRoon we generally assume this is not the case (and may optionally be done afterwards during the componentization step described below). Features are sometimes simply referred to as 'peaks.'

Features are found per analysis. Hence, in order to compare a feature across analyses, the next step is to group them. This step is essential as it finds equal features even if their retention time or m/z values slightly differ due to analytical variability. The resulting **feature groups** are crucial input for subsequent workflow steps. Prior to grouping, retention time alignment between analyses may be performed to improve grouping of features, especially when processing multiple analysis batches at once. Outside **patRoon** feature groups may also be defined as profiles, aligned or grouped features or buckets.

Depending on the study type, **suspect screening** is then performed to limit the features that should be considered for further processing. As its name suggests, with suspect screening only those features which are suspected to be present are considered for further processing. These suspects are retrieved from a suspect list which contains the m/z and (optionally) retention times for each suspect. Typical suspect lists may be composed from databases with known pollutants or from predicted transformation products. Note that for a 'full' non-target analysis no suspect screening is performed, hence, this step is simply omitted and all features are to be considered.

The feature group data may then be subjected to **componentization**. A **component** is defined as a collection of multiple feature groups that are somehow related to each other. Typical examples are features that belong to the same chemical compound (i.e. with different m/z values but equal retention time), such as adducts, isotopes and in-source fragments. Other examples are homologous series and features that display a similar intensity trend across samples. If adducts or isotopes were annotated during componentization then this data may be used to prioritize the feature groups.

The last step in the workflow commonly involves **annotation**. During this step MS and MS/MS data are collected in so called **MS peak lists**, which are then used as input for formula and compound annotation.

Formula annotation involves automatic calculation of possible formulae for each feature based on its m/z, isotopic pattern and MS/MS fragments, whereas compound annotation (or identification) involves the assignment of actual chemical structures to each feature. Note that during formula and compound annotation typically multiple candidates are assigned to a single feature. To assist interpretation of this data each candidate is therefore ranked on characteristics such as isotopic fit, number of explained MS/MS fragments and metadata from an online database such as number of scientific references or presence in common suspect lists.

To summarize:

- Data-pretreatment involves preparing raw MS data for further processing (e.g. conversion to an open format)
- Features describe chromatographic and m/z information (or 'peaks') in all analyses.
- A **feature group** consists of equal features across analyses.
- With suspect screening only features that are considered to be on a suspect list are considered further in the workflow.
- Componentization involves consolidating different feature groups that have a relationship to each other in to a single component.
- MS peak lists Summarizes all MS and MS/MS data that will be used for subsequent annotation.
- During formula and compound annotation candidate formulae/structures will be assigned and ranked for each feature.

The next chapters will discuss how to generate this data and process it. Afterwards, several advanced topics are discussed such as combining positive and negative ionization data, screening for transformation products and other advanced functionality.

4 Generating workflow data

Each step in the non-target workflow is performed by a function that performs the heavy lifting of a workflow step behind the scenes and finally return the results. An important goal of patRoon is to support multiple algorithms for each workflow step, hence, when such a function is called you have to specify which algorithm you want to use. The available algorithms and their characteristics will be discussed in the next sections. An overview of all functions involved in generating workflow data is shown in the table below.

Workflow step	Function	Output S4 class
Data pre-treatment	<pre>convertMSFiles(), recalibrarateDAFiles()</pre>	-
Finding features	<pre>findFeatures()</pre>	features
Grouping features	<pre>groupFeatures()</pre>	featureGroups
Suspect screening	screenSuspects()	featureGroupsScreening
Componentization	<pre>generateComponents()</pre>	components
MS peak lists	generateMSPeakLists()	MSPeakLists
Formula annotation	generateFormulas()	formulas
Compound annotation	<pre>generateCompounds()</pre>	compounds

All of these functions store their output in objects derived from so called S4 classes. Knowing the details about the S4 class system of R is generally not important when using patRoon (and well written resources are available if you want to know more). In brief, usage of this class system allows a general data format that is used irrespective of the algorithm that was used to generate the data. For instance, when features have been found by OpenMS or XCMS they both return the same data format.

Another advantage of the S4 class system is the usage of so called *generic functions*. To put simply: a generic function performs a certain task for different types of data objects. A good example is the plotSpectrum()

function which plots an (annotated) spectrum from data of MS peak lists or from formula or compound annotation:

```
# mslists, formulas, compounds contain results for MS peak lists and
# formula/compound annotations, respectively.

plotSpectrum(mslists, ...) # plot raw MS spectrum
plotSpectrum(formulas, ...) # plot annotated spectrum from formula annotation data
plotSpectrum(compounds, ...) # likewise but for compound annotation.
```

The next sections will further detail on how to actually perform the non-target workflow steps to generate data.

4.1 Preparations

4.1.1 Data pre-treatment

Prior to performing the actual non-target data processing workflow some preparations often need to be made. Often data has to be pre-treated, for instance, by converting it to an open format that is usable for subsequent workflow steps or to perform mass re-calibration. Some common functions are listed below.

Task	Function	Algorithms	Supported file formats
Conversion	convertMSFiles()	OpenMS, ProteoWizard, DataAnalysis	All com- mon (algo- rithm depen-
Advanced (e.g. spectral filtering)	convertMSFiles()	ProteoWizard	$egin{array}{l} \operatorname{dent}) \ \operatorname{All} \ \operatorname{com-} \end{array}$
Mass re-calibration	recalibrarateDAFil	es (DataAnalysis	mon Bruker

The convertMSFiles() function supports conversion between many different file formats typically used in non-target analysis. Furthermore, other pre-treatment steps are available (e.g. centroiding, filtering) when the ProteoWizard algorithm is used. For an overview of these functionalities see the MsConvert documentation. Some examples:

NOTE Most algorithms further down the workflow require the *mzML* or *mzXML* file format and additionally require that mass peaks have been centroided. When using the ProteoWizard algorithm (the default), centroiding by vendor algorithms is generally recommended (i.e. by setting centroid="vendor" as shown in the above example).

When Bruker MS data is used it can be automatically re-calibrated to improve its mass accuracy. Often this is preceded by calling the setDAMethod() function to set a DataAnalysis method to all files in order to configure automatic re-calibration. The recalibrarateDAFiles() function performs the actual re-calibration. The getDACalibrationError() function can be used at anytime to request the current calibration error of each analysis. An example of these functions is shown below.

```
# anaInfo is a data.frame with information on analyses (see next section)
setDAMethod(anaInfo, "path/to/DAMethod.m") # configure Bruker files with given method

→ that has automatic calibration setup
recalibrarateDAFiles(anaInfo) # trigger re-calibration for each analysis
getDACalibrationError(anaInfo) # get calibration error for each analysis (NOTE: also

→ shown when previous function is finished)
```

4.1.2 Analysis information

The final bits of preparation is constructing the information for the analyses that need to be processed. In patRoon this is referred to as the *analysis information* and often stored in a variable anaInfo (of course you are free to choose a different name!). The analysis information should be a data.frame with the following columns:

- path: the directory path of the file containing the analysis data
- analysis: the name of the analysis. This should be the file name without file extension.
- **group**: to which *replicate group* the analysis belongs. All analysis which are replicates of each other get the same name.
- blank: which replicate group should be used for blank subtraction.
- conc (optional, advanced) A numeric value describing the concentration or any other value for which the intensity in this sample may correlate, for instance, dilution factor, sampling time etc. This column is only required when you want to obtain quantitative information (e.g. concentrations) using the as.data.table() method function (see ?featureGroups for more information).

The generateAnalysisInfo() function can be used to (semi-)automatically generate a suitable data.frame that contains all the required information for a set of analysis. For, instance:

```
# Take example data from patRoonData package (triplicate solvent blank + triplicate

standard)

generateAnalysisInfo(paths = patRoonData::exampleDataPath(),

groups = c(rep("solvent-pos", 3), rep("standard-pos", 3)),

blanks = "solvent-pos")
```

```
#> path analysis group blank
#> 1 /usr/local/lib/R/site-library/patRoonData/extdata/pos solvent-pos-1 solvent-pos solvent-pos
#> 2 /usr/local/lib/R/site-library/patRoonData/extdata/pos solvent-pos-2 solvent-pos solvent-pos
#> 3 /usr/local/lib/R/site-library/patRoonData/extdata/pos solvent-pos-3 solvent-pos solvent-pos
#> 4 /usr/local/lib/R/site-library/patRoonData/extdata/pos standard-pos-1 standard-pos solvent-pos
#> 5 /usr/local/lib/R/site-library/patRoonData/extdata/pos standard-pos-2 standard-pos solvent-pos
#> 6 /usr/local/lib/R/site-library/patRoonData/extdata/pos standard-pos-3 standard-pos solvent-pos
```

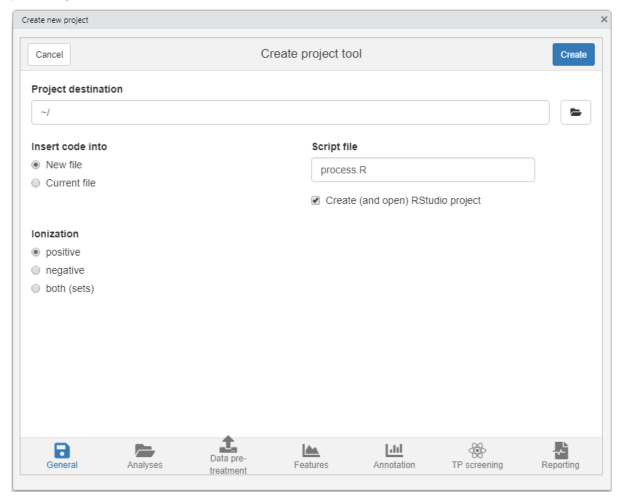
(Note that for the example data the patRoonData::exampleAnalysisInfo() function can also be used.)

Alternatively, the newProject() function discussed in the next section can be used to interactively construct this information.

4.1.3 Automatic project generation with newProject()

The previous sections already highlighted some steps that have to be performed prior to starting a new non-target analysis workflow: data pre-treatment and gathering information on the analysis. Most of the times you will put this and other R code a script file so you can recall what you have done before (i.e. reproducible research).

The newProject() function can be used to setup a new project. When you run this function it will launch a small tool (see screenshot below) where you can select your analyses and configure the various workflow steps which you want to execute (e.g. data pre-treatment, finding features, annotation etc). After setting everything up the function will generate a template script which can easily be edited afterwards. In addition, you have the option to create a new RStudio project, which is advantegeous as it neatly seperates your data processing work from the rest.



NOTE At the moment newProject() only works with RStudio.

4.2 Features

Collecting features from the analyses consists of finding all features, grouping them across analyses (optionally after retention time alignment), and if desired suspect screening:



4.2.1 Finding and grouping features

Several algorithms are available for finding features. These are listed in the table below alongside their usage and general remarks.

Algorithm	Usage	Remarks
OpenMS	findFeatures(algorithm = "openms",)	Uses the Feature-
		FinderMetabo
		algorithm
KCMS	<pre>findFeatures(algorithm = "xcms",)</pre>	Uses
		xcms::xcmsSet()
		function
XCMS (import)	<pre>importFeatures(algorithm = "xcms",)</pre>	Imports an existing
		xcmsSet object
XCMS3	<pre>findFeatures(algorithm = "xcms3",)</pre>	Uses
		xcms::findChromPeaks()
		from the new
		XCMS3 interface
XCMS3 (import)	<pre>importFeatures(algorithm = "xcms3",)</pre>	Imports an existing
		${\tt XCMSnExp}\ { m object}$
enviPick	<pre>findFeatures(algorithm = "envipick",)</pre>	Uses
		enviPick::enviPickwrap
KPIC2	<pre>findFeatures(algorithm = "kpic2",)</pre>	Uses the KPIC2 R
		package
KPIC2 (import)	<pre>importFeatures(algorithm = "kpic2",)</pre>	Imports features
		from KPIC2
SIRIUS	<pre>findFeatures(algorithm = "sirius",)</pre>	Uses SIRIUS to
		find features
SAFD	<pre>findFeatures(algorithm = "safd",)</pre>	Uses the SAFD
		algorithm
		(experimental)
OataAnalysis	<pre>findFeatures(algorithm = "bruker",)</pre>	Uses Find
		Molecular Features
		from DataAnalysis
		(Bruker only)

Most often the performance of these algorithms heavily depend on the data and parameter settings that are used. Since obtaining a good feature dataset is crucial for the rest of the workflow, it is highly recommended to experiment with different settings (this process can also be automated, see the feature optimization section for more details). Some common parameters to look at are listed in the table below. However, there are many more parameters that can be set, please see the reference documentation for these (e.g. ?findFeatures).

Algorithm	Common parameters
OpenMS	noiseThrInt, chromSNR, chromFWHM, mzPPM, minFWHM, maxFWHM (see ?findFeatures)
XCMS /	peakwidth, mzdiff, prefilter, noise (assuming default centWave algorithm, see
XCMS3	<pre>?findPeaks.centWave / ?CentWaveParam)</pre>
enviPick	dmzgap, dmzdens, drtgap, drtsmall, drtdens, drtfill, drttotal, minpeak, minint,
	maxint (see ?enviPickwrap)
KPIC2	kmeans, level, min_snr (see ?findFeatures and ?getPIC / ?getPIC.kmeans)
SIRIUS	The sirius algorithm is currently parameterless
SAFD	<pre>mzRange, maxNumbIter, resolution, minInt (see ?findFeatures)</pre>
DataAnalysis	See $Find \rightarrow Parameters \rightarrow Molecular Features$ in DataAnalysis.

NOTE Support for SAFD is still experimental and some extra work is required to set everything up. Please see the reference documentation for this algorithm (?findFeatures).

NOTE DataAnalysis feature settings have to be configured in DataAnalysis prior to calling findFeatures().

Similarly, for grouping features across analyses several algorithms are supported.

Algorithm	Usage	Remarks
OpenMS	<pre>groupFeatures(algorithm = "openms",)</pre>	Uses the FeatureLinkerUnlabeled algorithm (and MapAlignerPoseClustering for retention alignment)
XCMS	<pre>groupFeatures(algorithm = "xcms",)</pre>	Uses xcms::group() xcms::retcor() functions
XCMS (import)	<pre>importFeatureGroupsXCMS()</pre>	Imports an existing xcmsSet object.
XCMS3	<pre>groupFeatures(algorithm = "xcms3",)</pre>	<pre>Uses xcms::groupChromPeaks() and xcms::adjustRtime() functions</pre>
XCMS3 (import)	<pre>importFeatureGroupsXCMS3()</pre>	Imports an existing XCMSnExp object.
KPIC2	<pre>groupFeatures(algorithm = "kpic2",)</pre>	Uses the KPIC2 package
KPIC2 (import)	<pre>importFeatureGroupsKPIC2()</pre>	Imports a PIC set object
SIRIUS	<pre>groupFeatures(anaInfo, algorithm = "sirius")</pre>	Finds and groups features with SIRIUS
ProfileAnalysis	<pre>importFeatureGroups(algorithm = "brukerpa",)</pre>	Import .csv file exported from Bruker ProfileAnalysis
TASQ	<pre>importFeatureGroups(algorithm = "brukertasq",)</pre>	Imports a Global result table (exported to Excel file and then saved as .csv file)

NOTE: Grouping features with the sirius algorithm will perform both finding and grouping features with SIRIUS. This algorithm cannot work with features from another algorithm.

Just like finding features, each algorithm has their own set of parameters. Often the defaults are a good start but it is recommended to have look at them. See ?groupFeatures for more details.

When using the XCMS algorithms both the 'classical' interface and latest XCMS3 interfaces are supported. Currently, both interfaces are mostly the same regarding functionalities and implementation. However, since future developments of XCMS are primarily focused the latter this interface is recommended.

Some examples of finding and grouping features are shown below.

```
# The anaInfo variable contains analysis information, see the previous section
# Finding features
fListOMS <- findFeatures(anaInfo, "openms") # OpenMS, with default settings
fListOMS2 <- findFeatures(anaInfo, "openms", noiseThrInt = 500, chromSNR = 10) # OpenMS,
\rightarrow adjusted minimum intensity and S/N
fListXCMS <- findFeatures(anaInfo, "xcms", ppm = 10) # XCMS
fListXCMSImp <- importFeatures(anaInfo, "xcms", xset) # import XCMS xcmsSet object
fListXCMS3 <- findFeatures(anaInfo, "xcms3", CentWaveParam(peakwidth = c(5, 15))) # XCMS3
fListEP <- findFeatures(anaInfo, "envipick", minint = 1E3) # enviPick</pre>
fListKPIC2 <- findFeatures(anaInfo, "kpic2", kmeans = TRUE, level = 1E4) # KPIC2
fListSIRIUS <- findFeatures(anaInfo, "sirius") # SIRIUS
# Grouping features
fGroupsOMS <- groupFeatures(fListOMS, "openms") # OpenMS grouping, default settings
fGroupsOMS2 <- groupFeatures(fListOMS2, "openms", rtalign = FALSE) # OpenMS grouping, no
\hookrightarrow RT alignment
fGroupsOMS3 <- groupFeatures(fListXCMS, "openms", maxGroupRT = 6) # group XCMS features
→ with OpenMS, adjusted grouping parameter
# group enviPick features with XCMS3, disable minFraction
fGroupsXCMS <- groupFeatures(fListEP, "xcms3",</pre>
                             xcms::PeakDensityParam(sampleGroups = analInfo$group,
                                                     minFraction = 0))
# group with KPIC2 and set some custom grouping/aligning parameters
fGroupsKPIC2 <- groupFeatures(fListKPIC2, "kpic2", groupArgs = list(tolerance = c(0.002,
\rightarrow 18)),
                              alignArgs = list(move = "loess"))
fGroupsSIRIUS <- groupFeatures(anaInfo, "sirius") # find/group features with SIRIUS
```

4.2.2 Suspect screening

NOTE: you may need to install OpenBabel, for instance, when only InChI data is available for mass calculation.

After features have been grouped a so called suspect screening step may be performed to find features that may correspond to suspects within a given suspect list. The screenSuspects() function is used for this purpose, for instance:

4.2.2.1 Suspect list format The example above has a very simple suspect list with just three compounds. The format of the suspect list is quite flexible, and can contain the following columns:

- name: The name of the suspect. Mandatory and should be unique and file-name compatible (if not, the name will be automatically re-named to make it compatible).
- rt: The retention time in seconds. Optional. If specified any feature groups with a different retention time will not be considered to match suspects.
- mz, SMILES, InChI, formula, neutralMass: at least one of these columns must hold data for each suspect row. The mz column specifies the ionized mass of the suspect. If this is not available then data from any of the other columns is used to determine the suspect mass.
- adduct: The adduct of the suspect. Optional. Set this if you are sure that a suspect should be matched by a particular adduct ion and no data in the mz column is available.
- \bullet fragments_mz and fragments_formula: optional columns that may assist suspect annotation.

In most cases a suspect list is best made as a csv file which can then be imported with e.g. the read.csv() function. This is exactly what happen when you specify a suspect list when using the newProject() function.

Quite often, the ionized masses are not readily available and these have to be calculated. In this case, data in any of the SMILES/InChI/formula/neutralMass columns should be provided. Whenever possible, it is strongly recommended to fill in SMILES column (or InChI), as this will assist annotation. Applying this to the above example:

Since suspect matching now occurs by the neutral mass it is required that the adduct information for the feature groups are set. This is done either by setting the adduct function argument to screenSuspects or by feature group adduct annotations.

Finally, when the adduct is known for a suspect it can be specified in the suspect list:

To summarize:

- If a suspect has data in the mz column it will be directly matched with the m/z value of a feature group.
- Otherwise, if the suspect has data in the adduct column, the m/z value for the suspect is calculated from its neutral mass and the adduct and then matched with the m/z of a feature group.
- Otherwise, suspects and feature groups are matched by their the neutral mass.

The fragments_mz and fragments_formula columns in the suspect list can be used to specify known fragments for a suspect, which can help suspect annotation. The former specifies the ionized m/z of known MS/MS peaks, whereas the second specifies known formulas. Multiple values can be given by separating them with a semicolon:

4.2.2.2 Removing feature groups without hits Note that any feature groups that were not matched to a suspect are *not* removed by default. If you want to remove these, you can use the onlyHits parameter:

The advantage of removing non-hits is that it may significantly reduce the complexity of your dataset. On the other hand, retaining all features allows you to mix a full non-target analysis with a suspect screening workflow. The filter() function (discussed here) can also be used to remove feature groups without a hit at a later stage.

4.2.2.3 Combining screening results The amend function argument to screenSuspects can be used to combine screening results from different suspect lists.

```
fGroupsSusp <- screenSuspects(fGroups, suspects)
fGroupsSusp <- screenSuspects(fGroupsSusp, suspects2, onlyHits = TRUE, amend = TRUE)</pre>
```

In this example the suspect lists defined in suspects and suspects2 are both used for screening. By setting amend=TRUE the original screening results (i.e. from suspects) are preserved. Note that onlyHits should only be set in the final call to screenSuspects to ensure that all feature groups are screened.

4.3 Componentization

In patRoon componentization refers to grouping related feature groups together in components. There are different methodologies to generate components:

- Similarity on chromatographic elution profiles: feature groups with similar chromatographic behaviour which are assuming to be the same chemical compound (e.g. adducts or isotopologues).
- Homologous series: features with increasing m/z and retention time.
- Intensity profiles: features that follow a similar intensity profile in the analyses.
- MS/MS similarity: feature groups with similar MS/MS spectra are clustered.
- Transformation products: Components are formed by grouping feature groups that have a parent/transformation product relationship. This is further discussed in its own chapter.

The following algorithms are currently supported:

Algorithm	Usage	Remarks
CAMERA	<pre>generateComponents(algorithm = "camera",)</pre>	Clusters feature groups with similar chromatographic elution profiles and annotate by known chemical rules (adducts, isotopologues, in-source fragments).
RAMClustR	<pre>generateComponents(algorithm = "ramclustr",)</pre>	As above.
cliqueMS	<pre>generateComponents(algorithm = "cliquems",)</pre>	As above, but using feature components.
OpenMS	<pre>generateComponents(algorithm = "openms",)</pre>	As above. Uses MetaboliteAdductDecharger.
nontarget	<pre>generateComponents(algorithm = "nontarget",)</pre>	Uses the nontarget R package to perform unsupervised homologous series detection.
Intensity clustering	<pre>generateComponents(algorithm = "intclust",)</pre>	Groups features with similar intensity profiles across analyses by hierarchical clustering.
MS/MS clustering	<pre>generateComponents(algorithm = "specclust",)</pre>	Clusters feature groups with similar MS/MS spectra.
Transformation products	<pre>generateComponents(algorithm = "tp",)</pre>	Discussed in its own chapter.

4.3.1 Features with similar chromatographic behaviour

Isotopes, adducts and in-source fragments typically result in detection of multiple mass peaks by the mass spectrometer for a single chemical compound. While some feature finding algorithms already try to collapse (some of) these in to a single feature, this process is often incomplete (if performed at all) and it is not uncommon that multiple features will describe the same compound. To overcome this complexity several algorithms can be used to group features that undergo highly similar chromatographic behavior but have different m/z values. Basic chemical rules are then applied to the resulting components to annotate adducts, in-source fragments and isotopologues, which may be highly useful for general identification purposes.

Note that some algorithms were primarily designed for datasets where features are generally present in the majority of the analyses (as is relatively common in metabolomics). For environmental analyses, however, this is often not the case. For instance, consider the following situation with three feature groups that chromatographically overlap and therefore could be considered a component:

Feature group	m/z	analysis 1	analysis 2	analysis 3
#1	100.08827	Present	Present	Absent
#2	122.07021	Present	Present	Absent
#3	138.04415	Absent	Absent	Present

Based on the mass differences from this example a cluster of [M+H]+, [M+Na]+ and [M+K]+ could be assumed. However, no features of the first two feature groups were detected in the third sample analysis, whereas the third feature group wasn't detected in the first two sample analysis. Based on this it seems unlikely that feature group #3 should be part of the component.

For the algorithms that operate on a 'feature group level' (CAMERA and RAMClustR), the relMinReplicates argument can be used to remove feature groups from a component that are not abundant. For instance, when this value is 0.5 (the default), and all the features of a component were

detected in four different replicate groups in total, then only those feature groups are kept for which its features were detected in at least two different replicate groups (i.e. half of four).

Another approach to reduce unlikely adduct annotations is to use algorithms that operate on a 'feature level' (cliqueMS and OpenMS). These algorithms generate components for each sample analysis individually. The 'feature components' are then merged by a consensus approach where unlikely annotations are removed (the algorithm is described further in the reference manual, ?generateComponents).

Each algorithm supports many different parameters that may significantly influence the (quality of the) output. For instance, care has to be taken to avoid 'over-clustering' of feature groups which do not belong in the same component. This is often easily visible since the chromatographic peaks poorly overlap or are shaped differently. The checkComponents function (discussed here) can be used to quickly verify componentization results. For a complete listing all arguments see the reference manual (e.g. ?generateComponents).

Once the components with adduct and isotopes annotations are generated this data can be used to prioritize and improve the workflow.

Some example usage is shown below.

4.3.2 Homologues series

Homologues series can be automatically detected by interfacing with the nontarget R package. Components are made from feature groups that show increasing m/z and retention time values. Series are first detected within each replicate group. Afterwards, series from all replicates are linked in case (partial) overlap occurs and this overlap consists of the *same* feature groups (see figure below). Linked series are then finally merged if this will not cause any conflicts with other series: such a conflict typically occurs when two series are not only linked to each other.

The series that are linked can be interactively explored with the plotGraph() function (discussed here).

Common function arguments to generateComponents() are listed below.

Argument	Remarks
ionization	Ionization mode: "positive" or "negative". Not needed if adduct annotations are available.

Argument	Remarks
rtRange, mzRange	Retention and m/z increment range. Retention times can be negative to allow series with increasing m/z values and decreasing retention times.
elements	Vector with elements to consider.
rtDev, absMzDev	Maximum retention time and m/z deviation.
•••	Further arguments passed to the homol.search() function.

4.3.3 Intensity and MS/MS similarity

The previous componentization methods utilized chemical properties to relate features. The two componentization algorithms described in this section use a statistical approach based on hierarchical clustering. The first algorithm normalizes all feature intensities and then clusters features with similar intensity profiles across sample analyses together. The second algorithm compares all MS/MS spectra from all feature groups, and then uses hierarchical clustering to generate components from feature groups that have a high MS/MS spectrum similarity.

Some common arguments to generateComponents() are listed below. It is recommended to test various settings (especially for method) to optimize the clustering results.

Argument	Algorithm	Default	Remarks
method	All	"complete"	Clustering method. See ?hclust
metric	intclust	"euclidean"	Metric used to calculate the distance matrix. See ?daisy.
normFunc	intclust	max	Function used to normalize data. Feature intensities within a feature group are divided by the result of when this function is called with their intensity values.
average	intclust	TRUE	Whether intensities of replicates should first be averaged.
MSPeakLists	specclust	; -	The MS peak lists object used for spectral similarity calculations
specSimParams	specclust	getDefSpecSir	mPREnewsner ers used for spectral similarity calculation.
<pre>maxTreeHeight, deepSplit, minModuleSize</pre>	All	1, TRUE, 1	Used for dynamic cluster assignment. See ?cutreeDynamicTree.

The components are generated by automatically assigning clusters using the dynamicTreeCut R package. However, the cluster assignment can be performed manually or with different parameters, as is demonstrated below.

The resulting components are stored in an object from the componentsIntClust or componentsSpecClust S4 class, which are both derived from the componentsClust class (which in turn is derived from the components

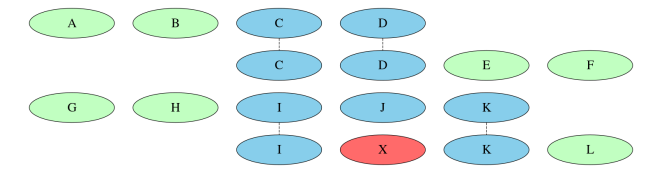


Figure 1: **Linking of homologues series** top: partial overlap and will be linked; bottom: no linkage due to different feature in overlapping series.

class). Several methods are defined that can be used on these objects to re-assign clusters, perform plotting operations and so on. Below are some examples. For plotting see the relevant visualization section. More info can be found in the reference manual (e.g. ?componentsIntClust, ?componentsSpecClust and ?componentsClust).

```
# generate intensity profile components with default settings
componInt <- generateComponents(fGroups, "intclust")

# manually re-assign clusters
componInt <- treeCut(componInt, k = 10)

# automatic re-assignment of clusters (adjusted max tree height)
componInt <- treeCutDynamic(componInt, maxTreeHeight = 0.7)

# MS/MS similarity components
componMSMS <- generateComponents(fGroups, "specclust", MSPeakLists = mslists)</pre>
```

4.4 Incorporating adduct and isotopic data

With mass spectrometry it is common that multiple m/z values are detected for a single compound. These may be different adducts (e.g. [M+H]+, [M+Na]+, [M-H]-), the different isotopes of the molecule or a combination thereof. When multiple m/z values are measured for the same compound, the feature finding algorithm may yield a distinct feature for each, which adds complexity to the data. In the previous section it was discussed how componentization can help to find feature groups that belong to the same adduct and/or isotope clusters. This section explains how this data can be used to simplify the feature dataset. Furthermore, this section also covers adduct annotations for feature groups which may improve and simplify the general workflow.

4.4.1 Selecting features with preferential adducts/isotopes

The selectIons function forms the bridge between feature group and componentization data. This function uses the adduct and isotope annotations to select preferential feature groups. For adduct clusters this means that only the feature group that has a preferential adduct (e.g. [M+H]+) is kept while others (e.g. [M+Na]+) are removed. If none of the adduct annotations are considered preferential, the most intense feature group is kept instead. For isotopic clusters typically only the feature group with the monoisotopic mass (i.e. M0) is kept.

The behavior of selections is configurable with the following parameters:

Argument	Remarks
prefAdduct The preferential adduct. Usually "[M+H]+" or "[M-H]-".	
${\tt onlyMonoIso}$	If TRUE and a feature group is with isotopic annotations then it is only kept if it
	is monoisotopic.
chargeMismatch	How charge mismatches between adduct and isotope annotations are dealt with. Valid options are "isotope", "adduct", "none" or "ignore". See the reference manual for selectIons for more details.

In case componentization did not lead to an adduct annotation for a feature group it will never be removed and simply be annotated with the preferential adduct. Similarly, when no isotope annotations are available and onlyMonoIso=TRUE, the feature group will not be removed.

Although selectIons operates fairly conservative, it is still recommended to verify the componentization results in advance, for instance with the checkComponents function discussed here. Furthermore, the next subsection explains how adduct annotations can be corrected manually if needed.

An example usage is shown below.

"[M+H]+"

#>

```
fGroupsSel <- selectIons(fGroups, componCAM, "[M+H]+")
```

```
#> No isotope annotations available!
```

- #> Removed 21 feature groups detected as unwanted adducts/isotopes
- #> Annotated 13 feature groups with adducts
- #> Remaining 111 feature groups set as default adduct [M+H]+

4.4.2 Setting adduct annotations for feature groups

The adducts() function can be used to obtain a character vector with adduct annotations for each feature group. When no adduct annotations are available it will simply return an empty character vector.

When the selectIons function is used it will automatically add adduct annotations based on the componentization data. In addition, the adducts()<- function can be used to manually add or change adduct annotations.

```
adducts(fGroups) # no adduct annotations

#> character(0)

adducts(fGroupsSel)[1:5] # adduct annotations set by selectIons()

#> M109_R192_20 M111_R330_23 M114_R269_25 M116_R317_29 M120_R268_30
#> "[M+H]+" "[M+H]+" "[M+H]+" "[M+H]+" "[M+K]+"

adducts(fGroupsSel)[3] <- "[M+Na]+" # modify annotation
adducts(fGroupsSel)[1:5] # verify</pre>
```

NOTE Adduct annotations are always available with sets workflows.

#> M109_R192_20 M111_R330_23 M114_R269_25 M116_R317_29 M120_R268_30

"[M+Na]+"

"[M+H]+"

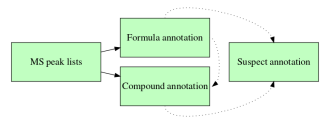
"[M+H]+"

4.4.3 Using adduct annotations in the workflow

When feature groups have adduct annotations available this may simplify and improve the workflow. The adduct and ionization arguments used for suspect screening, formula/compound annotation and some componentization algorithms do not have to be set anymore, since this data can be obtained from the adduct annotations. Furthermore, these algorithms may improve their results, since the algorithms are now able to use adduct information for each feature group individually, instead of assuming that all feature groups have the same adduct.

4.5 Annotation

The annotation consists of collecting MS peak lists and then formula and/or compound annotation:



Note that compound annotation is normally not dependent upon formula annotation. However, formula data can be used to improve ranking of candidates afterwards by the addFormulaScoring() function, which will be discussed later in this section. Furthermore, suspect annotation is not mandatory, and may use data from peak lists, formulae and/or comounds.

4.5.1 MS peak lists

Algorithm	Usage	Remarks
mzR	<pre>generateMSPeakLists(algorithm = "mzr",)</pre>	Uses mzR for spectra retrieval. Recommended default.
v	sisgenerateMSPeakLists(algorithm = "bruker",) sisgenerateMSPeakLists(algorithm = "brukerfmf",)	Loads data after automatically generating MS and MS/MS spectra in DataAnalysis Uses spectra from the find molecular features algorithm.

The recommended default algorithm is mzr: this algorithm is generally faster and is not limited to a vendor data format as it will read the open mzML and mzXML file formats. On the other hand, when DataAnalysis is used with Bruker data the spectra can be automatically background subtracted and there is no need for file conversion. Note that the brukerfmf algorithm only works when findFeatures() was called with the bruker algorithm.

When generateMSPeakists() is called it will

- 1. Find all MS and MS/MS spectra that 'belong' to a feature. For MS spectra this means that all spectra close to the retention time of a feature will be collected. In addition, for MS/MS normally only spectra will be considered that have a precursor mass close to that of the feature (however, this can be disabled for data that was recorded with data independent acquisition (DIA, MS^E, bbCID, ...)).
- 2. Average all MS and MS/MS spectra to produce peak lists for each feature.
- 3. Average all peak lists for features within the same group.

Data from either (2) or (3) is used for subsequent annotation steps. Formula calculation can use either (as a trade-off between possibly more accurate results by outlier removal vs speed), whereas compound annotation will always use data from (3) since annotating single features (as opposed to their groups) would take a very long time.

There are several common function arguments to generateMSPeakLists() that can be used to optimize its behaviour:

Argument	Algorithm(s)	Remarks
maxMSRtWindow	mzr, bruker	Maximum time window +/- the feature retention time (in seconds) to collect spectra for averaging. Higher values may significantly increase processing times.
precursorMzWindow	mzr	Maximum precursor m/z search window to find MS/MS spectra. Set to NULL to disable (i.e. for DIA experiments).
topMost	mzr	Only retain feature data for no more than this amount analyses with highest intensity. For instance, a value of 1 will only keep peak lists for the feature with highest intensity in a feature group.
bgsubtr	bruker	Perform background subtraction (if the spectra type supports this, e.g. MS and bbCID)
minMSIntensity,	bruker,	Minimum MS and MS/MS intensity. Note that
minMSMSIntensity	brukerfmf	DataAnalysis reports many zero intensity peaks so a value of at least 1 is recommended.
MSMSType	bruker	The type of spectra that should be used for MSMS: "BBCID" for bbCID experiments, otherwise "MSMS" (the default).

In addition, several parameters can be set that affect spectral averaging. These parameters are passed as a list to the avgFeatParams (mzr algorithm only) and avgFGroupParams arguments, which affect averaging of feature and feature group data, respectively. Some typical parameters include:

- clusterMzWindow: Maximum m/z window used to cluster mass peaks when averaging. The better the MS resolution, the lower this value should be.
- topMost: Retain no more than this amount of most intense mass peaks. Useful to filter out 'noisy' peaks.
- minIntensityPre / minIntensityPost: Mass peaks below this intensity will be removed before/after averaging.

See ?generateMSPeakLists for all possible parameters.

A suitable list object to set averaging parameters can be obtained with the getDefAvgPListParams() function.

4.5.2 Formulae

Formulae can be automatically calculated for all features using the generateFormulas() function. The following algorithms are currently supported:

Algorithm	Usage	Remarks
GenForm	<pre>generateFormulas(algorithm = "genform",)</pre>	Bundled with patRoon. Reasonable default.
SIRIUS	<pre>generateFormulas(algorithm = "sirius",)</pre>	Requires MS/MS data.
DataAnalysi	s generateFormulas(algorithm = "bruker",)	Requires FMF features (i.e. findFeatures(algorithm = "bruker",)). Uses SmartFormula algorithms.

Calculation with GenForm is often a good default. It is fast and basic rules can be applied to filter out obvious non-existing formulae. A possible drawback of GenForm, however, is that may become slow when many candidates are calculated, for instance, due to a relative high feature m/z (e.g. >600) or loose elemental restrictions. More thorough calculation is performed with SIRIUS: this algorithm often yields fewer and often more plausible results. However, SIRIUS requires MS/MS data (hence features without will not have results) and formula prediction may not work well for compounds that structurally deviate from the training sets used by SIRIUS. Calculation with DataAnalysis is only possible when features are obtained with DataAnalysis as well. An advantage is that analysis files do not have to be converted, however, compared to other algorithms calculation is often relative slow.

There are two methods for formula assignment:

- 1. Formulae are first calculated for each individual feature within a feature group. These results are then pooled, outliers are removed and remaining formulae are assigned to the feature group (i.e. calculateFeatures = TRUE).
- 2. Formulae are directly calculated for each feature group by using group averaged peak lists (see previous section) (i.e. calculateFeatures = FALSE).

The first method is more thorough and the possibility to remove outliers may sometimes result in better formula assignment. However, the second method is much faster and generally recommended for large number of analyses.

By default, formulae are either calculated by only MS/MS data (SIRIUS) or with both MS and MS/MS data (GenForm/Bruker). The latter also allows formula calculation when no MS/MS data is present. Furthermore, with Bruker algorithms, data from both MS and MS/MS formula data can be combined to allow inclusion of candidates that would otherwise be excluded by e.g. poor MS/MS data. However, a disadvantage is that formulae needs to be calculated twice. The MSMode argument (listed below) can be used to customize this behaviour.

An overview of common parameters that are typically set to customize formula calculation is listed below.

Argument	Algorithm(s)	Remarks
relMzDev	genform, sirius	The maximum relative m/z deviation for a formula to be considered (in ppm).
elements	genform, sirius	Which elements to consider. By default "CHNOP". Try to limit possible elements as much as possible.
calculateFeature	s genform, sirius	Whether formulae should be calculated first for all features (see discussion above) (always TRUE with DataAnalysis).

Argument	Algorithm(s)	Remarks
featThreshold	AnnAll	Minimum relative amount $(0-1)$ that a candidate formula for a feature group should be found among all annotated features (e.g. 1 means that a candidate is only considered if it was assigned to all annotated features).
adduct	All	The adduct to consider for calculation (e.g. "[M+H]+", "[M-H]-", more details in the adduct section). Don't set this when adduct annotations are available.
MSMode	genform, bruker	Whether formulae should be generated only from MS data ("ms"), MS/MS data ("msms") or both ("both"). The latter is default, see discussion above.
profile	sirius	Instrument profile, e.g. "qtof", "orbitrap", "fticr".

Some typical examples:

```
formulasGF <- generateFormulas(fGroups, mslists, "genform") # GenForm, default settings
formulasGF2 <- generateFormulas(fGroups, mslists, "genform", calculateFeatures = FALSE) #

\( \to \) direct feature group assignment (faster)
formulasSIR <- generateFormulas(fGroups, mslists, "sirius", elements = "CHNOPSC1Br") #

\( \to \) SIRIUS, common elements for pollutant
formulasSIR2 <- generateFormulas(fGroups, mslists, "sirius", adduct = "[M-H]-") # SIRIUS,
\( \to \) negative ionization
formulasBr <- generateFormulas(fGroups, mslists, "bruker", MSMode = "MSMS") # Only
\( \to \) consider MSMS data (SmartFormula3D)
```

4.5.3 Compounds

An important step in a typical non-target workflow is structural identification for features of interest. Afterall, this information may finally reveal what a feature is. The first step is to find all possible structures in a database that may be assigned to the feature (based on e.g. monoisotopic mass or formula). These candidates are then scored to rank likely candidates, for instance, on correspondence with in-silico or library MS/MS spectra and environmental relevance.

Structure assignment in patRoon is performed automatically for all feature groups with the generateCompounds() function. Currently, this function supports two algorithms:

Algorithm	Usage	Remarks
MetFrag	<pre>generateCompounds(algorithm = "metfrag",)</pre>	Supports many databases (including custom) and scorings for candidate ranking.
SIRIUS with CSI:FingerID	<pre>generateCompounds(algorithm = "sirius",)</pre>	Incorporates prior comprohensive formula calculations.

Compound annotation is often a relative time and resource intensive procedure. For this reason, features are not annotated individually, but instead a feature group as a whole is annotated, which generally saves significant amounts of computational requirements. Nevertheless, it is not uncommon that this is the most time consuming step in the workflow. For this reason, prioritization of features is highly important, even more so to avoid 'abusing' servers when an online database is used for compound retrieval.

Selecting the right database is important for proper candidate assignment. Afterall, if the 'right' chemical compound is not present in the used database, it is impossible to assign the correct structure. Luckily,

however, several large databases such as PubChem and ChemSpider are openly available which contain tens of millions of compounds. On the other hand, these databases may also lead to many unlikely candidates and therefore more specialized (or custom databases) may be preferred. Which database will be used is dictated by the database argument to generateCompounds(), currently the following options exist:

Database	Algorithm(s)	Remarks	
pubchem	"metfrag", "sirius"	PubChem is currently the largest compound database and is used by default.	
chemspider	"metfrag"	ChemSpider is another large database. Requires security token from here (see next section).	
comptox	"metfrag"	The EPA CompTox contains many compounds and scorings relevant to environmental studies. Needs manual download (see next section).	
pubchemlite	"metfrag"	A specialized subset of the PubChem database. Needs manual download (see next section).	
for-ident	"metfrag"	The FOR-IDENT (STOFF-IDENT) database for water related substances.	
kegg	"metfrag", "sirius"	The KEGG database for biological compounds	
hmdb	"metfrag", "sirius"	The HMDB contains many human metabolites.	
bio	"sirius"	Selects all supports biological databases.	
csv, psv, sdf	"metfrag"	Custom database (see next section). CSV example.	

4.5.3.1 Configuring MetFrag databases and scoring Some extra configuration may be necessary when using certain databases with MetFrag. In order to use the ChemSpider database a security token should be requested and set with the chemSpiderToken argument to generateCompounds(). The CompTox and PubChemLite databases need to be manually downloaded from CompTox (or variations with smoking or wastewater metadata) and PubChemLite. The file location of this and other local databases (csv, psv, sdf) needs to be manually configured, see the examples below and/or ?generateCompounds for more information on how to do this.

An example of a custom .csv database can be found here.

With MetFrag compound databases are not only used to retrieve candidate structures but are also used to obtain metadata for further ranking. Each database has its own scorings, a table with currently supported scorings can be obtained with the compoundScorings() function (some columns omitted):

name	metfrag	database	default
score	Score		TRUE
fragScore	FragmenterScore		TRUE
metFusionScore	OfflineMetFusionScore		TRUE
individual MoNAS core	OfflineIndividualMoNAScore		TRUE
numberPatents	PubChemNumberPatents	pubchem	TRUE
numberPatents	Patent_Count	pubchemlite	TRUE
pubMedReferences	PubChemNumberPubMedReferences	pubchem	TRUE
pubMedReferences	ChemSpiderNumberPubMedReferences	chemspider	TRUE
pubMedReferences	NUMBER OF PUBMED ARTICLES	comptox	TRUE
pubMedReferences	PubMed_Count	pubchemlite	TRUE
extReferenceCount	ChemSpiderNumberExternalReferences	chemspider	TRUE
dataSourceCount	ChemSpiderDataSourceCount	chemspider	TRUE
referenceCount	ChemSpiderReferenceCount	chemspider	TRUE
RSCCount	ChemSpiderRSCCount	chemspider	TRUE
formulaScore			FALSE
smartsInclusionScore	SmartsSubstructureInclusionScore		FALSE
smartsExclusionScore	SmartsSubstructureExclusionScore		FALSE
suspectListScore	SuspectListScore		FALSE
retentionTimeScore	RetentionTimeScore		FALSE
CPDATCount	CPDAT_COUNT	comptox	TRUE
TOXCASTActive	TOXCAST PERCENT ACTIVE	comptox	TRUE
dataSources	DATA SOURCES	comptox	TRUE
pubChemDataSources	PUBCHEM DATA SOURCES	comptox	TRUE
EXPOCASTPredExpo	EXPOCAST MEDIAN EXPOSURE PREDICTION MG/KG-BW/DAY	comptox	TRUE
ECOTOX	ECOTOX	comptox	TRUE
NORMANSUSDAT	NORMANSUSDAT	comptox	TRUE
MASSBANKEU	MASSBANKEU	comptox	TRUE
TOX21SL	TOX21SL	comptox	TRUE
TOXCAST	TOXCAST	comptox	TRUE
KEMIMARKET	KEMIMARKET	comptox	TRUE
MZCLOUD	MZCLOUD	comptox	TRUE
pubMedNeuro	PubMedNeuro	comptox	TRUE
CIGARETTES	CIGARETTES	comptox	TRUE
INDOORCT16	INDOORCT16	comptox	TRUE
SRM2585DUST	SRM2585DUST	comptox	TRUE
SLTCHEMDB	SLTCHEMDB	comptox	TRUE
THSMOKE	THSMOKE	comptox	TRUE
ITNANTIBIOTIC	ITNANTIBIOTIC	comptox	TRUE
STOFFIDENT	STOFFIDENT	comptox	TRUE
KEMIMARKET EXPO	KEMIMARKET EXPO	comptox	TRUE
_	_	*	TRUE
KEMIMARKET_HAZ	KEMIMARKET_HAZ	comptox	
REACH2017	REACH2017	comptox	TRUE

$KEMIWW_WDUIndex$	KEMIWW_WDUIndex	comptox	TRUE
$KEMIWW_StpSE$	KEMIWW_StpSE	comptox	TRUE
${\tt KEMIWW_SEHitsOverDL}$	KEMIWW_SEHitsOverDL	comptox	TRUE
ZINC15PHARMA	ZINC15PHARMA	comptox	TRUE
PFASMASTER	PFASMASTER	comptox	TRUE
peakFingerprintScore	AutomatedPeakFingerprintAnnotationScore	•	FALSE
lossFingerprintScore	AutomatedLossFingerprintAnnotationScore		FALSE
agroChemInfo	AgroChemInfo	pubchemlite	FALSE
bioPathway	BioPathway	pubchemlite	FALSE
drugMedicInfo	DrugMedicInfo	pubchemlite	FALSE
foodRelated	FoodRelated	pubchemlite	FALSE
pharmacoInfo	PharmacoInfo	pubchemlite	FALSE
safetyInfo	SafetyInfo	pubchemlite	FALSE
toxicityInfo	ToxicityInfo	pubchemlite	FALSE
knownUse	KnownUse	pubchemlite	FALSE
disorderDisease	DisorderDisease	pubchemlite	FALSE
identification	Identification	pubchemlite	FALSE
annoTypeCount	FPSum	pubchemlite	TRUE
annorypecount	11 Julii	pubenemnte	11000
anno Type Count	AnnoTypeCount	pubchemlite	TRUE

The first two columns contain the generic and original MetFrag naming schemes for each scoring type. While both naming schemes can be used, the generic is often shorter and harmonized with other algorithms (e.g. SIRIUS). The *database* column specifies for which databases a particular scoring is available (empty if not database specific). Most scorings are selected by default (as specified by the *default* column), however, this behaviour can be customized by using the scoreTypes argument:

By default ranking is performed with equal weight (i.e. 1) for all scorings. This can be changed by the scoreWeights argument, which should be a vector containing the weights for all scorings following the order of scoreTypes, for instance:

Sometimes thousands or more structural candidates are found when annotating a feature group. In this situation processing all these candidates will too involving (especially when external databases are used). To avoid this a default cut-off is set: when the number of candidates exceed a certain amount the search will be aborted and no results will be reported for that feature group. The maximum number of candidates can be set with the maxCandidatesToStop argument. The default value is relative conservative, especially for local databases it may be useful to increase this number.

4.5.3.2 Timeout and error handling The use of online databases has the drawback that an error may occur, for instance, as a result of a connection error. Furthermore, MetFrag typically returns an error when too many candidates are found (as set by the maxCandidatesToStop argument). By default

processing is restarted if an error has occurred (configured by the errorRetries argument). Similarly, the timeoutRetries and timeout arguments can be used to avoid being 'stuck' on obtaining results, for instance, due to an unstable internet connection.

If no compounds could be assigned due to an error a warning will be issued. In this case it is best to see what went wrong by manually checking the log files, which by default are stored in the log/metfrag folder.

4.5.3.3 Formula scoring Ranking of candidate structures may further be improved by incorporating formula information by using the addFormulaScoring() function:

```
comps <- addFormulaScoring(coms, formulas, updateScore = TRUE)</pre>
```

Here, corresponding formula and explained fragments will be used to calculate a *formulaScore* for each candidate. Note that SIRIUS candidates are already based on calculated formulae, hence, running this function on SIRIUS results is less sensable unless scoring from another formula calculation algorithm is desired.

4.5.3.4 Further options and parameters There are *many* more options and parameters that affect compound annotation. For a full overview please have a look at the reference manual (e.g. by running ?generateCompounds).

4.5.4 Suspect annotation

The data obtained during the previously described annotation steps can be used to improve a suspect screening workflow. The annotateSuspects() method uses the annotation data to calculate various annotation properties for each suspect, such as their rank in formula/compound candidates, which fragments from the suspect list were matched, and a *rough* indication of the identification level according to Schymanski et al. (2014)

The calculation of identification levels is performed by a set of pre-defined rules. The genIDLevelRulesFile() can be used to inspect the default rules or to create your own rules file, which can subsequently passed to annotateSuspects() with the IDFile argument. See ?annotateSuspects for more details on the file format and options. The default identification levels can be summarized as follows:

Level	Description	Rules
1	Target match	Retention time deviates <12 seconds from suspect list. At least 3 (or all if the suspect list contains less) fragments from the suspect list must match.
2a	Good MS/MS library match	Suspect is top ranked in the compounds results. The individualMoNAScore is at least 0.9 and all other candidates have no MoNA library score.
3a	Fair library match	The individualMoNAScore is at least 0.4.
3b	Known MS/MS match	At least 3 (or all if the suspect list contains less) fragments from the suspect list must match.
3c	Good in-silico MS/MS match	The annotation MS/MS similarity (annSimComp column) is at least 0.7.

Level	Description	Rules
4a	Good formula MS/MS match	Suspect is top ranked formula candidate, annotation MS/MS similarity (annSimForm column) is at least 0.7 and isotopic match (isoScore) of at least 0.5. The latter two scores are at least 0.2 higher than next best ranked candidate.
4b	Good formula isotopic pattern match	Suspect is top ranked formula candidate and isotopic match (isoScore) of at least 0.9 and at least 0.2 higher than next best ranked candidate.
5	Unknown	All else.

In general, the more data provided by the suspect list and to annotateSuspects(), the better identification level estimation works. For instance, when considering the default rules, either the fragments_mz or fragments_formula column is necessary to be able assign a level 3b. Similarly, the suspect list needs retention times (as well as fragment data) to be able to assign level 1. As you can imagine, providing the annotation workflow objects (i.e. MSPeakLists, formulas, compounds) to annotateSuspects() is necessary for calculation of most levels.

The annotateSuspects() function will log decisions for identification level assignments to the log/ sub-directory in the current working directory. This is useful to inspect level assignments and especially useful when you customized any rules.

 ${\it NOTE}$: The current identification level rules are ${\it only}$ optimized for GenForm and MetFrag annotation algorithms.

5 Processing workflow data

The previous chapter mainly discussed how to create workflow data. This chapter will discuss how to use the data.

5.1 Inspecting results

Several generic functions exist that can be used to inspect data that is stored in a particular object (e.g. features, compounds etc):

Generic	Classes	Remarks
length()	All	Returns the length of the object (e.g. number of features, compounds etc)
algorithm()	All	Returns the name of the algorithm used to generate the object.
groupNames()	All	Returns all the unique identitifiers (or names) of the feature groups for which this object contains results.
names()	$\begin{array}{c} \texttt{featureGroups}, \\ \texttt{components} \end{array}$	Returns names of the feature groups (similar to groupNames()) or components
show()	All	Prints general information.

Generic	Classes	Remarks
"[[" / "\$" operators	All	Extract general information, see below.
<pre>as.data.table() / as.data.frame()</pre>	All	Convert data to a data.table or data.frame, see below.
<pre>analysisInfo(), analyses(),</pre>	features,	Returns the analysis information,
replicateGroups()	featureGroups	analyses or replicate groups for which this object contains data.
<pre>groupInfo()</pre>	featureGroups	Returns feature group information (m/z) and retention time values).
<pre>screenInfo()</pre>	featureGroupsScreening	Returns information on hits from suspect screening.
<pre>componentInfo()</pre>	components	Returns information for all components.
annotatedPeakList()	formulas, compounds	Returns a table with annotated mass peaks (see below).

The common R extraction operators "[[", "\$" can be used to obtain data for a particular feature groups, analysis etc:

```
# Feature table (only first columns for readability)
fList[["standard-1"]][, 1:6]
```

#> NULL

```
# Feature group intensities
fGroups$M120_R268_30
```

#> [1] 264836 245372 216560

```
fGroups[[1, "M120_R268_30"]] # only first analysis
```

#> [1] 264836

```
# obtains MS/MS peak list (feature group averaged data)
mslists[["M120_R268_30"]]$MSMS
```

```
#>
      ID
               mz intensity precursor
#> 1: 5 105.0698
                    6183.111
                                FALSE
#> 2: 6 106.0653
                    7643.556
                                FALSE
#> 3: 8 107.0728
                    7760.667
                                FALSE
#> 4: 15 120.0556 168522.667
                                 TRUE
#> 5: 17 121.0587 13894.667
                                FALSE
#> 6: 18 121.0884 10032.889
                                FALSE
#> 7: 19 122.0964 147667.778
                                FALSE
#> 8: 20 123.0803 36631.111
                                FALSE
#> 9: 21 123.0996 15482.444
                                 FALSE
#> 10: 22 124.0805 35580.667
                                 FALSE
```

```
# get all formula candidates for a feature group
formulas[["M120_R268_30"]][, 1:7]
```

```
# get all compound candidates for a feature group
compounds[["M120_R268_30"]][, 1:4]
```

```
#>
       explainedPeaks
                           score neutralMass
                                                                 SMILES
#>
    1:
                    0 2.8673850
                                                     C1=CC2=NNN=C2C=C1
                                    119.0483
#>
    2:
                    0 1.4569407
                                    119.0483 C1=CC=C(C=C1)N=[N+]=[N-]
#>
    3:
                    0 1.1555020
                                    119.0483
                                                   C1=CC2=C(N=C1)N=CN2
                    0 1.0911519
                                                   C1=CC2=C(C=NN2)N=C1
#>
   4:
                                    119.0483
#>
    5:
                    0 1.0910125
                                    119.0483
                                                     C1=CNC2=NC=NC=C21
#>
                    0 0.8722331
#> 26:
                                    119.0483
                                                     C1=CN=C(C=N1)CC#N
#> 27:
                    0 0.8617554
                                    119.0483
                                                     C1=CC2=NC=NN2C=C1
#> 28:
                    0 0.8580143
                                    119.0483
                                                     C1=CC2=CN=NN2C=C1
#> 29:
                    0 0.7249048
                                    119.0483 C1=CC(=[N+]=[N-])C=CC1=N
#> 30:
                    0 0.6442352
                                    119.0483
                                                      C(C#N)C(CC#N)C#N
```

```
# get a table with information of a component
components[["CMP7"]][, 1:6]
```

```
#>
              group
                                   mz isogroup isonr charge
                        ret
#> 1: M143 R206 64 205.787 143.0700
                                            NA
                                                  NA
#> 2: M159_R208_103 208.280 159.0650
                                                  NA
                                                          NΑ
#> 3: M161_R208_104 207.582 161.0806
                                                          NA
                                            NΑ
                                                  NΑ
#> 4: M181 R209 159 208.580 181.0469
                                            NA
                                                  NA
                                                          NA
```

A more sophisticated way to obtain data from a workflow object is to use as.data.table() or as.data.frame(). These functions will convert *all* information within the object to a table (data.table or data.frame) and allow various options to add extra information. An advantage is that this common data format can be used with many other functions within R. The output is in a tidy format.

NOTE If you are not familiar with data.table and want to know more see data.table. Briefly, this is a more efficient and largely compatible alternative to the regular data.frame.

NOTE The as.data.frame() methods defined in patRoon simply convert the results from as.data.table(), hence, both functions are equal in their usage and are defined for the same object classes.

Some typical examples are shown below.

```
# obtain table with all features (only first columns for readability)
as.data.table(fList)[, 1:6]
```

```
#>
              analysis
                                           ID
                                                  ret
                                                                    area intensity
                                                           \mathtt{mz}
#>
     1: solvent-pos-1 f_4025525033642747772 13.176 98.97537 4345232.0
                                                                            391476
                                               7.181 100.11197 797112.1
#>
     2: solvent-pos-1 f 6703224702672402688
                                                                            426956
     3: solvent-pos-1 f_5186620951722599099 192.178 100.11211 9609998.0
#>
                                                                            750532
     4: solvent-pos-1 f_1685696430399603217 19.171 100.11217 5784411.0
#>
                                                                            370376
     5: solvent-pos-1 f 9942721423467665216 4.786 100.11220 551723.6
#>
                                                                            567312
#> 2920: standard-pos-3 f 2072982058656265121 318.892 425.18866 666531.5
                                                                           232636
#> 2921: standard-pos-3 f_3705009681110839484
                                                9.114 427.03242 362024.1
                                                                            114744
#> 2922: standard-pos-3 f_9021412140160071733 318.892 427.18678 200193.5
                                                                            77768
#> 2923: standard-pos-3 f_7380080213749915164 382.682 432.23984 217612.9
                                                                             97648
#> 2924: standard-pos-3 f_17512800645305435413
                                              9.114 433.00457 3086864.0
                                                                            912920
# Returns group info and intensity values for each feature group
as.data.table(fGroups, average = TRUE) # average intensities for replicates
#>
               group
                          ret
                                    mz standard-pos
#>
    1: M109_R192_20 191.8729 109.0759
                                         183482.67
#>
    2: M111_R330_23 330.4093 111.0439
                                          84598.67
    3: M114 R269 25 268.6919 114.0912
                                          85796.00
    4: M116 R317 29 316.7348 116.0527
#>
                                          766888.00
    5: M120 R268 30 268.4092 120.0554
                                          242256.00
#> ---
#> 138: M316_R363_635 363.4894 316.1741
                                          89904.00
#> 139: M318_R349_638 349.1087 318.1450
                                          83320.00
#> 140: M352_R335_664 334.9418 352.2019
                                           74986.67
#> 141: M407_R239_672 239.3580 407.2227
                                        186568.00
#> 142: M425_R319_676 319.4959 425.1885
                                          214990.67
# As above, but with extra suspect screening information
# (select some columns to simplify the output below)
as.data.table(fGroupsSusp, average = TRUE, collapseSuspects = NULL,
             onlyHits = TRUE)[1:3, c("group", "susp_name", "susp_compRank",

    "susp_annSimBoth", "susp_estIDLevel")]

#>
                           susp_name susp_compRank susp_annSimBoth susp_estIDLevel
            group
#> 1: M120 R268 30
                    1H-benzotriazole
                                               1
                                                         0.0000000
                                                                                5
#> 2: M137 R249 53
                       N-Phenyl urea
                                                 1
                                                         0.6443557
#> 3: M146_R309_68 2-Hydroxyquinoline
                                                         0.9896892
                                                                                Зс
# Returns all peak lists for each feature group
as.data.table(mslists)
#>
               group type ID
                                   mz intensity precursor
#>
    1: M120_R268_30 MS 1 100.1120 178952.381
                                                     FALSE
#>
    2: M120 R268 30
                       MS 2 102.1277 202359.667
                                                     FALSE
    3: M120_R268_30
#>
                       MS 3 114.0912 37647.548
                                                     FALSE
    4: M120_R268_30
#>
                       MS 4 115.0752 66685.238
                                                     FALSE
#>
    5: M120_R268_30
                       MS 5 120.0554 113335.857
                                                    TRUE
```

FALSE

#> 235: M192_R355_191 MS 51 299.1274 44083.126

```
#> 236: M192 R355 191
                        MS 52 299.1471
                                         7390.267
                                                      FALSE
#> 237: M192_R355_191 MSMS 14 119.0496 588372.444
                                                      FALSE
#> 238: M192 R355 191 MSMS 18 120.0524 70273.333
                                                      FALSE
#> 239: M192_R355_191 MSMS 31 192.1384
                                        71978.667
                                                       TRUE
# Returns all formula candidates for each feature group with scoring
# information, neutral loss etc
as.data.table(formulas)[, 1:6]
              group neutral_formula ion_formula neutralMass ion_formula_mz
                                                                                error
#> 1: M120_R268_30
                             C6H5N3
                                         C6H6N3
                                                   119.0483
                                                                  120.0556 1.80000000
#> 2: M137 R249 53
                            C7H8N2O
                                        C7H9N2O
                                                   136.0637
                                                                  137.0709 2.90000000
#> 3: M146_R309_68
                                       C9H8NO
                                                   145.0528
                                                                  146.0600 1.66666667
                             C9H7NO
#> 4: M192_R355_191
                           C12H17NO
                                       C12H18NO
                                                   191.1310
                                                                  192.1383 0.03333333
# Returns all compound candidates for each feature group with scoring and other metadata
as.data.table(compounds)[, 1:4]
#>
                group explainedPeaks
                                         score neutralMass
#>
     1: M120_R268_30
                                   0 2.8673850
                                                  119.0483
     2: M120 R268 30
                                   0 1.4569407
                                                  119.0483
```

```
3: M120_R268_30
#>
                                   0 1.1555020
                                                  119.0483
    4: M120 R268 30
                                  0 1.0911519
                                                  119.0483
#>
                                  0 1.0910125
#>
     5:
        M120 R268 30
                                                 119.0483
#>
#> 248: M192 R355 191
                                  0 0.6851763
                                                  191.1310
#> 249: M192 R355 191
                                  0 0.6764365
                                                  191.1310
#> 250: M192_R355_191
                                  0 0.6713679
                                                  191.1310
#> 251: M192 R355 191
                                  0 0.6519220
                                                  191.1310
#> 252: M192_R355_191
                                  0 0.6497224
                                                  191.1310
```

Returns table with all components (including feature group info, annotations etc) as.data.table(components)[, 1:6]

```
analysis size
#>
       name cmp_ret cmp_retsd
                                      neutral_mass
#>
  1:
       CMP1 347.2929 0.0000000
                                              <NA> standard-pos-2
                                                                     2
#> 2: CMP1 347.2929 0.0000000
                                              <NA> standard-pos-2
#> 3: CMP2 349.6343 4.6805095 225.1589/188.20157 standard-pos-3
       CMP2 349.6343 4.6805095 225.1589/188.20157 standard-pos-3
#> 5: CMP2 349.6343 4.6805095 225.1589/188.20157 standard-pos-3
#> ---
#> 86: CMP28 313.3489 0.3104371
                                              <NA> standard-pos-2
                                                                     3
#> 87: CMP28 313.3489 0.3104371
                                              <NA> standard-pos-2
                                                                     3
                                         81.08705 standard-pos-1
                                                                     3
#> 88: CMP29 268.3442 0.3843289
#> 89: CMP29 268.3442 0.3843289
                                         81.08705 standard-pos-1
                                                                     3
#> 90: CMP29 268.3442 0.3843289
                                         81.08705 standard-pos-1
                                                                     3
```

Finally, the annotatedPeakList() function is useful to inspect annotation results for a formula or compound candidate:

```
#>
                 mz intensity precursor ion_formula dbe ion_formula_mz error neutral_loss
                                                                                      CHNO
#>
      2 94.06500 9406.111
                                  FALSE
                                              C6H8N 3.5
                                                              94.06513
                                                                        1.30
   1:
   2: 6 98.97522
                     2212.000
                                  FALSE
                                               <NA>
                                                                           NA
                                                                                      <NA>
                                                    NA
                                                                                      <NA>
#>
  3: 7 105.06971 1662.111
                                  FALSE
                                               <NA> NA
                                                                           NA
                                                                    NA
   4: 14 120.04434
                     7176.222
                                  FALSE
                                             C7H6NO 5.5
                                                              120.04439
                                                                        0.40
                                                                                       H3N
   5: 19 122.07222
                     2246.000
                                  FALSE
                                               <NA> NA
                                                                    NA
                                                                           NA
                                                                                      <NA>
  6: 21 135.08004
                    1565.556
                                  FALSE
                                               <NA> NA
                                                                    NA
                                                                           NA
                                                                                      <NA>
#> 7: 23 137.07039
                     5348.667
                                   TRUE
                                            C7H9N2O 4.5
                                                              137.07094
                                                                        3.35
#> 8: 24 137.09572 2026.889
                                  FALSE
                                               <NA> NA
                                                                     NA
                                                                           NA
                                                                                      <NA>
#> 9: 26 138.09116 12356.667
                                  FALSE
                                               < NA >
                                                    NA
                                                                     NA
                                                                           NA
                                                                                      <NA>
#> 10: 27 139.07503 5020.667
                                  FALSE
                                               <NA>
                                                    NA
                                                                    NA
                                                                           NA
                                                                                      <NA>
```

```
#>
                 mz intensity precursor ion_formula ion_formula_MF neutral_loss score
       ID
#>
    1: 2
           94.06500
                     9406.111
                                   FALSE
                                                C6H8N
                                                        [C6H6N+H]+H+
                                                                              CHNO
                                                                                     405
#>
    2: 6 98.97522
                     2212.000
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NA
   3: 7 105.06971
                     1662.111
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NΑ
  4: 14 120.04434
                     7176.222
                                                           [C7H6NO] +
                                                                               H3N
                                                                                     305
#>
                                   FALSE
                                               C7H6NO
   5: 19 122.07222
                     2246.000
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NA
#> 6: 21 135.08004
                     1565.556
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NA
#> 7: 23 137.07039
                     5348.667
                                    TRUE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NΑ
#> 8: 24 137.09572 2026.889
                                   FALSE
                                                 <NA>
                                                                < NA >
                                                                              <NA>
                                                                                      NA
#> 9: 26 138.09116 12356.667
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NA
#> 10: 27 139.07503 5020.667
                                   FALSE
                                                 <NA>
                                                                <NA>
                                                                              <NA>
                                                                                      NA
```

More advanced examples for these functions are shown below.

```
as.data.table(formulas, OM = TRUE)

as.data.table(compounds, fGroups = fGroups) # add feature group information
as.data.table(compounds, fragments = TRUE) # include information of all annotated

index = 1, groupName = "M120_R268_30",

MSPeakLists = mslists, onlyAnnotated = TRUE) # only include annotated

index = 1, groupName = "M120_R268_30",

index = 1, groupName = "M120_R268_30",

MSPeakLists = mslists, formulas = formulas) # include formula

index = annotations
```

5.2 Filtering

During a non-target workflow it is not uncommon that some kind of data-cleanup is necessary. Datasets are often highly complex, which makes separating data of interest from the rest highly important. Furthermore, general cleanup typically improves the quality of the dataset, for instance by removing low scoring annotation results or features that are unlikely to be 'correct' (e.g. noise or present in blanks). For this reason patRoon supports many different filters that easily clean data produced during the workflow in a highly customizable way.

All major workflow objects (e.g. featureGroups, compounds, components etc.) support filtering operations by the filter() generic. This function takes the object to be filtered as first argument and any remaining arguments describe the desired filter options. The filter() generic function then returns the modified object back. Some examples are shown below.

```
# remove low intensity (<500) features
features <- filter(features, absMinIntensity = 500)

# remove features with intensities lower than 5 times the blank
fGroups <- filter(fGroups, blankThreshold = 5)

# only retain compounds with >1 explained MS/MS peaks
compounds <- filter(compounds, minExplainedPeaks = 1)</pre>
```

The following sections will provide a more detailed overview of available data filters.

NOTE Some other R packages (notably dplyr) also provide a filter() generic function. To use the filter() function from different packages you may need to explicitly specify which one to use in your script. This can be done by prefixing it with the package name, e.g. patRoon::filter(...), dplyr::filter(...) etc.

5.2.1 Features

There are many filters available for feature data:

Filter	Classes	Remarks
absMinIntensity,	features,	Minimum intensity
relMinIntensity	featureGroups	

Filter	Classes	Remarks
preAbsMinIntensity,	featureGroups	Minimum intensity prior to other filtering
${\tt preRelMinIntensity}$		(see below)
retentionRange, mzRange,	features,	Filter by feature properties
mzDefectRange,	featureGroups	
${\tt chromWidthRange}$		
${\tt absMinAnalyses},$	featureGroups	Minimum feature abundance in all analyses
relMinAnalyses		
${\tt absMinReplicates},$	featureGroups	Minimum feature abundance in different
relMinReplicates		replicates
${\tt absMinFeatures},$	featureGroups	Only keep analyses with at least this amount
relMinFeatures		of features
$\verb"absMinReplicateAbundance",$	featureGroups	Minimum feature abundance in a replicate
${\tt relMinReplicateAbundance}$		group
${\tt maxReplicateIntRSD}$	featureGroups	Maximum relative standard deviation of
		feature intensities in a replicate group.
blankThreshold	featureGroups	Minimum intensity factor above blank
		intensity
rGroups	featureGroups	Only keep (features of) these replicate groups
results	featureGroups	Only keep feature groups with
		formula/compound annotations or
		componentization results

Application of filters to feature data is important for (environmental) non-target analysis. Especially blank and replicate filters (i.e. blankThreshold and absMinReplicateAbundance/relMinReplicateAbundance) are important filters and are highly recommended to always apply for cleaning up your dataset.

All filters are available for feature group data, whereas only a subset is available for feature objects. The main reason is that other filters need grouping of features between analyses. Regardless, in patRoon filtering feature data is less important, and typically only needed when the number of features are extremely large and direct grouping is undesired.

From the table above you can notice that many filters concern both absolute and relative data (i.e. as prefixed with abs and rel). When a relative filter is used the value is scaled between θ and 1. For instance:

```
# remove features not present in at least half of the analyses within a replicate group
fGroups <- filter(fGroups, relMinReplicateAbundance = 0.5)</pre>
```

An advantage of relative filters is that you will not have to worry about the data size involved. For instance, in the above example the filter always takes half of the number of analyses within a replicate group, even when replicate groups have different number of analyses.

Note that multiple filters can be specified at once. Especially for feature group data the order of filtering may impact the final results, this is explained further in the reference manual (i.e. ?`feature-filtering`). Some examples are shown below.

```
maxReplicateIntRSD = 0.75, # remove features with intensity RSD in
                  → replicates >75%
                  blankThreshold = 5, # remove features <5x intensity of (average) blank
                  \rightarrow intensity
                  removeBlanks = TRUE) # remove blank analyses from object afterwards
# filter by feature properties
fGroups <- filter(mzDefectRange = c(0.8, 0.9),
                  chromWidthRange = c(6, 120))
# remove features not present in at least 3 analyses
fGroups <- filter(fGroups, absMinAnalyses = 3)
# remove features not present in at least 20% of all replicate groups
fGroups <- filter(fGroups, relMinReplicates = 0.2)
# only keep data present in replicate groups "repl1" and "repl2"
# all other features and analyses will be removed
fGroups <- filter(fGroups, rGroups = c("repl1", "repl2"))
# only keep feature groups with compound annotations
fGroups <- filter(fGroups, results = compounds)</pre>
# only keep feature groups with formula or compound annotations
fGroups <- filter(fGroups, results = list(formulas, compounds))
```

5.2.2 Suspect screening

Several additional filters are available for feature groups obtained with screenSuspects():

Filter	Classes	Remarks
onlyHits	featureGroupsScreenOmgly retain feature groups assigned to one or	
		more suspects.
selectHitsBy	featureGroupsScree	en Some et the feature group that matches best with
		a suspect (in case there are multiple).
selectBestFGroups	featureGroupsScree	en Singlect the suspect that matches best with a
		feature group (in case there are multiple).
maxLevel, maxFormRank,	featureGroupsScreen Ongly retain suspect hits with	
maxCompRank	identification/annotation ranks below	
_		threshold.
minAnnSimForm, minAnnSimComp,	featureGroupsScreenPhymove suspect hits with annotation similarity	
minAnnSimBoth	•	scores below this value.
absMinFragMatches,	featureGroupsScreen Onely keep suspect hits with a minimum	
relMinFragMatches	•	(relative) number of fragment matches from the
		suspect list.

NOTE: most filters only remove suspect hit results. Set onlyHits=TRUE to also remove any feature groups that end up without suspect hits.

The selectHitsBy and selectBestFGroups filters are useful to remove duplicate hits (one suspect assigned to multiple feature groups assigned to the same suspect, respectively). The former selects based on either best identification level (selectHitsBy="level") or highest mean intensity

(selectHitsBy="intensity"). The selectBestFGroups can only be TRUE/FALSE and always selects by best identification level.

Some examples are shown below.

```
# only keep feature groups assigned to at least one suspect
fGroupsSusp <- filter(fGroupsSusp, onlyHits = TRUE)
# remove duplicate suspect to feature group matches and keep the best
fGroupsSusp <- filter(fGroupsSusp, selectHitsBy = "level")
# remove suspect hits with ID levels >3 and make sure no feature groups
# are present without suspect hits afterwards
fGroupsSusp <- filter(fGroupsSusp, maxLevel = 3, onlyHits = TRUE)</pre>
```

5.2.3 Annotation

There are various filters available for handling annotation data:

Filter	Classes	Remarks
absMSIntThr, absMSMSIntThr, relMSIntThr, relMSIntThr	MSPeakLists	Minimum intensity of mass peaks
topMSPeaks, topMSMSPeaks	MSPeakLists	Only keep most intense mass peaks
withMSMS	MSPeakLists	Only keep results with MS/MS data
minMSMSPeaks	MSPeakLists	Only keep an MS/MS peak list if it contains a minimum number of peaks (excluding the precursor peak)
annotatedBy	MSPeakLists	Only keep MS/MS peaks that have formula or compound annotations
minExplainedPeaks	formulas, compounds	Minimum number of annotated mass peaks
elements, fragElements, lossElements	formulas, compounds	Restrain elemental composition
topMost	formulas, compounds	Only keep highest ranked candidates
minScore, minFragScore, minFormulaScore	compounds	Minimum compound scorings
scoreLimits	formulas, compounds	Minimum/Maximum scorings
OM	formulas, compounds	Only keep candidates with likely elemental composition found in organic matter

Several intensity related filters are available to clean-up MS peak list data. For instance, the topMSPeaks/topMSMSPeaks filters provide a simple way to remove noisy data by only retaining a defined number of most intense mass peaks. Note that none of these filters will remove the precursor mass peak of the feature itself.

The filters applicable to formula and compound annotation generally concern minimal scoring or chemical properties. The former is useful to remove unlikely candidates, whereas the second is useful to focus on certain study specific chemical properties (e.g. known neutral losses).

Common examples are shown below.

```
# intensity filtering
mslists <- filter(mslists,</pre>
```

```
absMSIntThr = 500, # minimum MS mass peak intensity of 500
                  relMSMSIntThr = 0.1) # minimum MS/MS mass peak intensity of 10%
# only retain 10 most intens mass peaks
# (feature mass is always retained)
mslists <- filter(mslists, topMSPeaks = 10)</pre>
# remove MS/MS peaks without compound annotations
mslists <- filter(mslists, annotatedBy = compounds)</pre>
# remove MS/MS peaks not annotated by either a formula or compound candidate
mslists <- filter(mslists, annotatedBy = list(formulas, compounds))</pre>
# only keep formulae with 1-10 sulphur or phosphorus elements
formulas <- filter(formulas, elements = c("S1-10", "P1-10"))</pre>
# only keep candidates with MS/MS fragments that contain 1-10 carbons and 0-2 oxygens
formulas <- filter(formulas, fragElements = "C1-1000-2")</pre>
# only keep candidates with CO2 neutral loss
formulas <- filter(formulas, lossElements = "CO2")</pre>
# only keep the 15 highest ranked candidates with at least 1 annotated MS/MS peak
compounds <- filter(compounds, minExplainedPeaks = 1, topMost = 15)
# minimum in-silico score
compounds <- filter(compounds, minFragScore = 10)</pre>
# candidate should be referenced in at least 1 patent
# (only works if database lists number of patents, e.q. PubChem)
compounds <- filter(compounds,</pre>
                    scoreLimits = list(numberPatents = c(1, Inf))
```

NOTE As of patRoon 2.0 MS peak lists are **not** re-generated after a filtering operation (unless the **reAverage** parameter is explicitly set to TRUE). The reason for this change is that re-averaging invalidates any formula/compound annotation data (e.g. used for plotting and reporting) that were generated prior to the filter operation.

5.2.4 Components

Finally several filters are available for components:

Filter	Remarks
size adducts, isotopes rtIncrement, mzIncrement	Minimum component size Filter features by adduct/istopes annotation Filter homologs by retention/mz increment range

Note that these filters are only applied if the components contain the data the filter works on. For instance, filtering by adducts will *not* affect components obtained from homologous series.

As before, some typical examples are shown below.

NOTE As mentioned before, components are still in a relative young development phase and results should always be verified!

5.2.5 Negation

All filters support *negation*: if enabled all specified filters will be executed in an opposite manner. Negation may not be so commonly used, but allows greater flexibility which is sometimes needed for advanced filtering steps. Furthermore, it is also useful to specifically isolate the data that otherwise would have been removed. Some examples are shown below.

```
# keep all features/analyses _not_ present from replicate groups "repl1" and "repl2"
fGroups <- filter(fGroups, rGroups = c("repl1", "repl2"), negate = TRUE)

# only retain features with a mass defect outside 0.8-0.9
fGroups <- filter(mzDefectRange = c(0.8, 0.9), negate = TRUE)

# remove duplicate suspect hits and only keep the _worst_ hit
fGroupsSusp <- filter(fGroupsSusp, selectHitsBy = "level", negate = TRUE)

# remove candidates with CO2 neutral loss
formulas <- filter(formulas, lossElements = "CO2", negate = TRUE)

# select 15 worst ranked candidates
compounds <- filter(compounds, topMost = 15, negate = TRUE)

# only keep components with <5 features
componInt <- filter(componInt, minSize = 5, negate = TRUE)</pre>
```

5.3 Subsetting

The previous section discussed the filter() generic function to perform various data cleaning operations. A more generic way to select data is by *subsetting*: here you can manually specify which parts of an object

should be retained. Subsetting is supported for all workflow objects and is performed by the R subset operator ("["]). This operator either subsets by one or two arguments, which are referred to as the i and j arguments.

Class	Argument i	Argument j	Remarks
features featureGroups MSPeakLists	analyses analyses analyses	feature groups feature groups	peak lists for feature groups will be re-averaged when subset on analyses (by default)
formulas compounds components	feature groups feature groups components	feature groups	

For objects that support two-dimensional subsetting (e.g. featureGroups, MSPeakLists), either the i or j argument is optional. Furthermore, unlike subsetting a data.frame, the position of i and j does not change when only one argument is specified:

```
df[1, 1] # subset data.frame by first row/column
df[1] # subset by first column
df[1, ] # subset by first row

fGroups[1, 1] # subset by first analysis/feature group
fGroups[, 1] # subset by first feature group (i.e. column)
fGroups[1] # subset by first analysis (i.e. row)
```

The subset operator allows three types of input:

- A logical vector: elements are selected if corresponding values are TRUE.
- A numeric vector: select elements by numeric index.
- A character vector: select elements by their name.

When a logical vector is used as input it will be re-cycled if necessary. For instance, the following will select by the first, third, fifth, etc. analysis.

```
fGroups[c(TRUE, FALSE)]
```

In order to select by a character you will need to know the names for each element. These can, for instance, be obtained by the groupNames() (feature group names), analyses() (analysis names) and names() (names for components or feature groups for featureGroups objects) generic functions.

Some more examples of common subsetting operations are shown below.

```
# select first three analyses
fList[1:3]

# select first three analyses and first 500 feature groups
fGroups[1:3, 1:500]

# select all feature groups from first component
fGroupsNT <- fGroups[, componNT[[1]]$group]</pre>
```

```
# only keep feature groups with formula annotation results
fGroupsForms <- fGroups[, groupNames(formulas)]

# only keep feature groups with either formula or compound annotation results
fGroupsAnn <- fGroups[, union(groupNames(formulas), groupNames(compounds))]

# select first 15 components
components[1:15]

# select by name
components[c("CMP1", "CMP5")]

# only retain feature groups in components for which compound annotations are
# available
components[, groupNames(compounds)]</pre>
```

In addition, feature groups can also be subset by given replicate groups or annotation/componentization results (similar to filter()). Similarly, suspect screening results can also be subset by given suspect names.

```
# equal as filter(fGroups, rGroups = ...)
fGroups[rGroups = c("repl1", "repl2")]
# equal as filter(fGroups, results = ...)
fGroups[results = compounds]
# only keep feature groups assigned to given suspects
fGroupsSusp[suspects = c("1H-benzotriazole", "2-Hydroxyquinoline")]
```

NOTE As of patRoon 2.0 MS peak lists are **not** re-generated after a subsetting operation (unless the reAverage parameter is explicity set to TRUE). The reason for this change is that re-averaging invalidates any formula/compound annotation data (e.g. used for plotting and reporting) that were generated prior to the subset operation.

5.3.1 Prioritization workflow

An important use case of subsetting is prioritization of data. For instance, after statistical analysis only certain feature groups are deemed relevant for the rest of the workflow. A common prioritization workflow is illustrated below:



During the first step the workflow object is converted to a suitable format, most often using the as.data.frame() function. The converted data is then used as input for the prioritization strategy. Finally, these results are then used to select the data of interest in the original object.

A very simplified example of such a process is shown below.

```
featTab <- as.data.frame(fGroups, average = TRUE)

# prioritization: sort by (averaged) intensity of the "sample" replicate group
# (from high to low) and then obtain the feature group identifiers of the top 5.</pre>
```

```
featTab <- featTab[order(featTab$standard, decreasing = TRUE), ]
groupsOfInterest <- featTab$group[1:5]

# subset the original data
fGroups <- fGroups[, groupsOfInterest]

# fGroups now only contains the feature groups for which intensity values in the
# "sample" replicate group were in the top 5</pre>
```

5.4 Deleting data

The delete() generic function can be used to manually delete workflow data. This function is used internally within patRoon to implement filtering and subsetting operations, but may also be useful for advanced data processing.

Like the subset operator this function accepts a i and j parameter to specify which data should be operated on:

Class	Argument i	Argument j
features	analysis	feature index
featureGroups	analysis	feature group
formulas, compounds	feature group	candidate index
components	component	feature group

If i or j is not specified (NULL) then data is removed for the complete selection. Some examples are shown below:

```
# delete 2nd feature in analysis-1
fList <- delete(fList, i = "analysis-1", j = 2)</pre>
# delete first ten features in all analyses
fList <- delete(fList, i = NULL, j = 1:10)</pre>
# completely remove third/fourth analyses from feature groups
fGroups <- delete(fGroups, i = 3:4)
# delete specific feature group
fGroups <- delete(fGroups, j = "M120 R268 30")
# delete range of feature groups
fGroups \leftarrow delete(fGroups, j = 500:750)
# remove all results for a feature group
formulas <- delete(formulas, i = "M120 R268 30")
# remove top candidate for all feature groups
compounds <- delete(compounds, j = 1)</pre>
# remove a component
components <- delete(components, i = "CMP1")</pre>
# remove specific feature group from a component
components <- delete(components, i = "CMP1", j = "M120_R268_30")</pre>
# remove specific feature group from all components
components <- delete(components, j = "M120_R268_30")</pre>
```

The j parameter can also be a function: in this case it is called repeatedly on parts of the data to select what should be deleted. How the function is called and what it should return depends on the workflow data class:

Class	Called on every	First argument	Second argument	Return value
features	analysis	data.table with features	analysis name	Features indices (as integer or logical)
featureGroups	feature group	vector with group intensities	feature group name	The analyses of the features to remove (as character, integer, logical)
formulas, compounds	feature group	data.table with annotations	feature group name	Candidate indices (rows)
components	component	data.table with the component	component name	The feature groups (as character, integer)

Some examples for this:

```
# remove features with intensities below 5000
fList <- delete(fList, j = function(f, ...) f$intensity <= 5E3)

# same, but for features in all feature groups from specific analyses
fGroups <- delete(i = 1:3, j = function(g, ...) g <= 5E3)

# remove formula candidates with high relative mass deviation
formulas <- delete(formulas, j = function(ft, ...) ft$error > 5)
```

5.5 Unique and overlapping features

Often an analysis batch is composed of different sample groups, such as different treatments, influent/effluent etc. In such scenarios it may be highly interesting to evaluate uniqueness or overlap between these samples. Furthermore, extracting overlapping or unique features is a simple but effective prioritization strategy.

The overlap() and unique() functions can be used to extract overlapping and unique features between replicate groups, respectively. Both functions return a subset of the given featureGroups object. An overview of their arguments is given below.

Argument	Function(s)	Remarks
which	unique(), overlap()	The replicate groups to compare.
relativeTo	unique()	Only return unique features compared to these replicate groups
		(NULL for all). Replicate groups in which are ignored.
outer	unique()	If TRUE then only return features which are also unique among
		the compared replicates groups.
exclusive	overlap	Only keep features that only overlap between the compared
		replicate groups.

Some examples:

In addition, several plotting functions are discussed in the visualization section that visualize overlap and uniqueness of features.

5.6 MS similarity

The *spectral similarity* is used to compare spectra from different features. For this purpose the **spectrumSimilarity** function can be used. This function operates on MS peak lists, and accepts the following function arguments:

Argument	Remarks
MSPeakLists	The MS peak lists object from which peak lists data should be taken.
${\tt groupName1},$	The name(s) of the first and second feature group(s) to compare
groupName2	
analysis1, analysis2	The analysis names of the data to be compared. Set this when feature data
	(instead of feature group data) should be compared.
MSLevel	The MS level: 1 or 2 for MS and MS/MS, respectively.
specSimParams	Parameters that define how similarities are calculated.
NAToZero	If TRUE then NA values are converted to zeros. NA values are reported if a comparison cannot be made because of missing peak list data.
	comparison cannot be made because of missing peak list data.

The specSimParams argument defines the parameters for similarity calculations. It is a list, and the default values are obtained with the getDefSpecSimParams() function:

```
getDefSpecSimParams()
```

```
#> $method
#> [1] "cosine"
#>
#> $removePrecursor
#> [1] FALSE
#>
#> $mzWeight
#> [1] 0
#>
```

```
#> $intWeight
#> [1] 1
#>
#> $absMzDev
#> [1] 0.005
#>
#> $relMinIntensity
#> [1] 0.05
#>
#> $minPeaks
#> [1] 1
#>
#> $shift
#> [1] "none"
#>
#> $setCombineMethod
#> [1] "mean"
```

The method field describes the calculation measure: this is either "cosine" or "jaccard".

The shift field is primarily useful when comparing MS/MS data and defines if and how a spectral shift should be performed prior to similarity calculation:

- "none": The default, no shifting is performed.
- "precursor" The mass difference between the precursor mass of both spectra (*i.e.* the feature mass) is first calculated. This difference is then subtracted from each of the mass peaks of the second spectrum. This shifting increases similarity if the MS fragmentation process itself occurs similarly (*i.e.* if both features show similar neutral losses).
- "both" This combines both shifting methods: first peaks are aligned that have the same mass, then the precursor strategy is applied for the remaining mass peaks. This shifting method yields higher similarities if either fragment masses or neutral losses are similar.

To override a default setting, simply pass it as an argument to getDefSpecSimParams:

```
getDefSpecSimParams(shift = "both")
```

For more details on the various similarity calculation parameters see the reference manual (?getDefSpecSimParams). Some examples are shown below:

#> [1] 0.08589848

#> [1] 0.1111111

#> [1] 0.08589848

The spectrumSimilarity function can also be used to calculate *multiple* similarities. Simply specify multiple feature group names for the groupNameX parameters. Alternatively, if you want to compare the same set of feature groups with each other pass their names only as the groupName1 parameter:

```
# compare two pairs
spectrumSimilarity(mslists,
                  groupName1 = c("M120_R268_30", "M137_R249_53"),
                  groupName2 = c("M146_R309_68", "M192_R355_191"),
                  MSLevel = 2, specSimParams = getDefSpecSimParams(shift = "both"))
#>
               M146 R309 68 M192 R355 191
#> M120_R268_30
                   0.520052
                               0.08589848
                               0.03372542
#> M137_R249_53
                   0.197720
# compare all
spectrumSimilarity(mslists, groupName1 = groupNames(mslists),
                  MSLevel = 2, specSimParams = getDefSpecSimParams(shift = "both"))
#>
                M120_R268_30 M137_R249_53 M146_R309_68 M192_R355_191
#> M120 R268 30
                  1.00000000 0.20406381
                                            0.52005204
                                                          0.08589848
#> M137_R249_53
                  0.20406381 1.00000000
                                            0.19772004
                                                          0.03372542
                  0.52005204 0.19772004
#> M146_R309_68
                                            1.00000000
                                                          0.08524785
#> M192_R355_191
                  0.08589848 0.03372542
                                            0.08524785
                                                          1.00000000
```

5.7 Visualization

5.7.1 Features and annatation data

Several generic functions are available to visualize feature and annotation data:

Generic	Classes	Remarks
plot()	featureGroups,	Scatter plot for retention and m/z values
	${\tt featureGroupsComparison}$	
<pre>plotInt()</pre>	featureGroups	Intensity profiles across analyses
<pre>plotChroms()</pre>	featureGroups, components	Plot extracted ion chromatograms (EICs)

Generic	Classes	Remarks
plotSpectrum	(MSPeakLists, formulas, compounds, components	Plots (annotated) spectra
<pre>plotStructur plotScores() plotGraph()</pre>	e Compounds formulas, compounds componentsNT	Draws candidate structures Barplot for candidate scoring Draws interactive graphs of linked homologous series

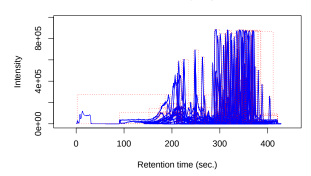
The most common plotting functions are plotChroms(), which plots chromatographic data for features, and plotSpectrum(), which will plot (annotated) spectra. An overview of their most important function arguments are shown below.

Argument	Generic	Remarks
rtWindow	plotChroms()	Extra time (in s) +/- retention limits of plotted
		features (useful to zoom out)
retMin	<pre>plotChroms()</pre>	If TRUE plot retention times in minutes
topMost	<pre>plotChroms()</pre>	Only draw this amount of highest intensity
		features in each group.
topMostByRGroup	<pre>plotChroms()</pre>	If TRUE then the topMost parameter specifies the
		top most intense features in each replicate group
		to draw (e.g. $topMost=1$ would draw the most
		intense feature for each replicate group).
${\tt showPeakArea}, {\tt showFGroupRect}$	<pre>plotChroms()</pre>	Fill peak areas / draw rectangles around feature
		groups?
title	${\tt plotChroms()},$	Override plot title
	<pre>plotSpectrum()</pre>	
colourBy	<pre>plotChroms()</pre>	Colour individual feature groups ("fGroups") or
		replicate groups ("rGroups"). By default nothing
		is coloured ("none")
showLegend	plotChroms()	Display a legend? (only if colourBy!="none")
onlyPresent	<pre>plotChroms()</pre>	Only plot EICs for analyses where a feature was
		detected? Setting to FALSE is useful to inspect if
	()	a feature was 'missed.'
xlim, ylim	plotChroms(),	Override x/y axis ranges, i.e. to manually set
	plotSpectrum()	plotting range.
groupName, analysis,	<pre>plotSpectrum()</pre>	What to plot. See examples below.
precursor, index	7 . (2	TITL 1 A DAG DAG DAG
MSLevel	<pre>plotSpectrum()</pre>	Whether to plot an MS or MS/MS spectrum
	7 (0)	(only MSPeakLists)
formulas	<pre>plotSpectrum()</pre>	Whether formula annotation should be added
	1 - + 0 + ()	(only compounds)
plotStruct	<pre>plotSpectrum()</pre>	Whether the structure should be added to the
	-1 -+C+ ()	plot (only compounds)
mincex	<pre>plotSpectrum()</pre>	Minimum annotation font size (only
		formulas/compounds)

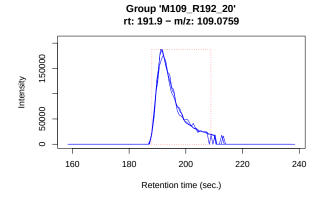
Note that we can use subsetting to select which feature data we want to plot, e.g.

plotChroms(fGroups[1:2]) # only plot EICs from first and second analyses.

142 feature groups

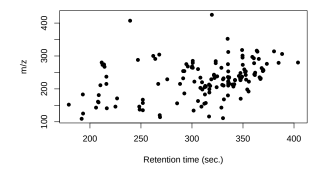


plotChroms(fGroups[, 1]) # only plot all features of first group



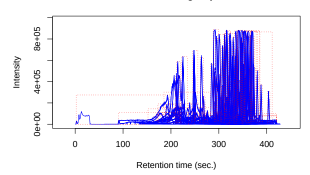
The plotStructure() function will draw a chemical structure for a compound candidate. In addition, this function can draw the maximum common substructure (MCS) of multiple candidates in order to assess common structural features.

Some other common and less common plotting operations are shown below.

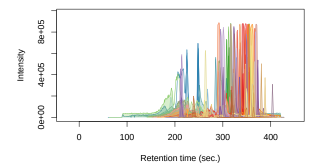


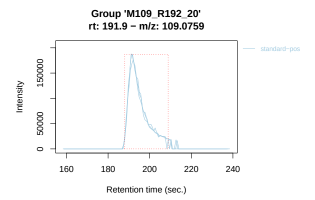
plotChroms(fGroups) # plot EICs for all features

142 feature groups

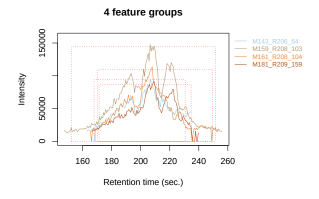


142 feature groups

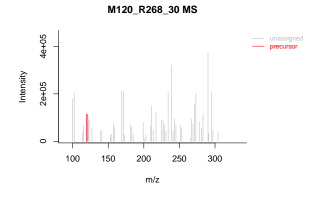




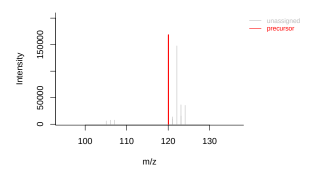
plotChroms(components, index = 7, fGroups = fGroups) # EICs from a component

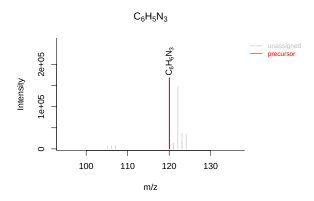


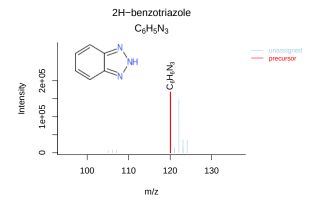
 $\verb|plotSpectrum(mslists, "M120_R268_30")| \textit{# non-annotated MS spectrum}|$

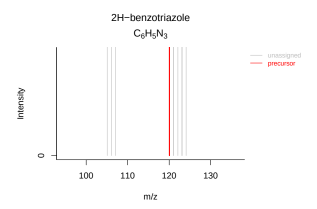


M120_R268_30 MSMS

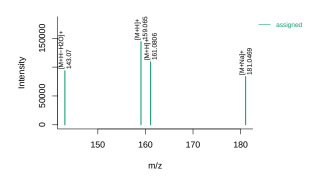




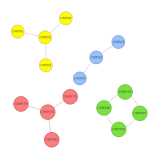




```
plotSpectrum(components, index = 7) # component spectrum
```



Inspect homologous series
plotGraph(componNT)



5.7.2 Overlapping and unique data

There are three functions that can be used to visualize overlap and uniqueness between data:

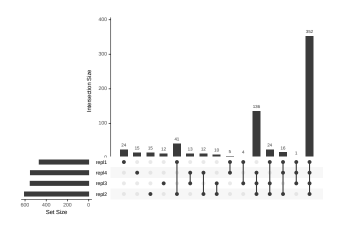
Generic	Classes
plotVenn	featureGroups, featureGroupsComparison, formulas, compounds
${ t plotUpSet}$	${\tt feature Groups, feature Groups Comparison, formulas, compounds}$
plotChord	${\tt feature Groups Comparison}$

The most simple comparison plot is a Venn diagram (i.e. plotVenn()). This function is especially useful for two or three-way comparisons. More complex comparisons are better visualized with UpSet diagrams (i.e. plotUpSet()). Finally, chord diagrams (i.e. plotChord()) provide visually pleasing diagrams to assess overlap between data.

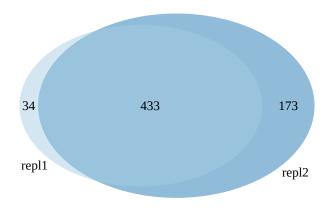
These functions can either be used to compare feature data or different objects of the same type. The former is typically used to compare overlap or uniqueness between features in different replicate groups, whereas comparison between objects is useful to visualize differences in algorithmic output. Besides visualization, note that both operations can also be performed to modify or combine objects (see unique and overlapping features and algorithm consensus).

As usual, some examples are shown below.

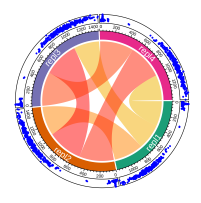
plotUpSet(fGroups) # compare replicate groups



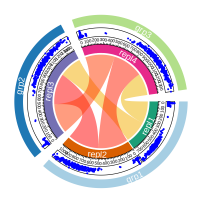
plotVenn(fGroups, which = c("repl1", "repl2")) # compare some replicate groups

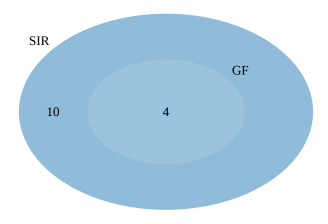


plotChord(fGroups, average = TRUE) # overlap between replicate groups



```
# compare with custom made groups
plotChord(fGroups, average = TRUE,
    outer = c(repl1 = "grp1", repl2 = "grp1", repl3 = "grp2", repl4 = "grp3"))
```

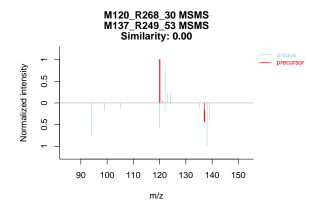


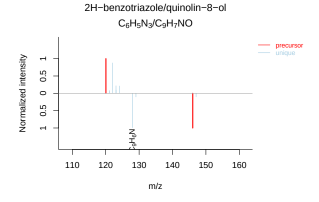


5.7.3 MS similarity

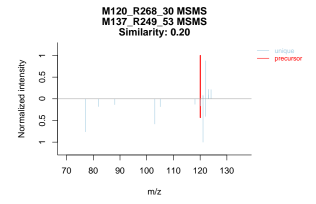
The plotSpectrum function is also useful to visually compare (annotated) spectra. This works for MSPeakLists, formulas and compounds object data.

```
plotSpectrum(mslists, groupName = c("M120_R268_30", "M137_R249_53"), MSLevel = 2)
```





The specSimParams argument, which was discussed in MS similarity, can be used to configure the similarity calculation:

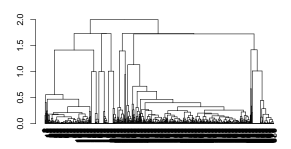


5.7.4 Hierarchical clustering results

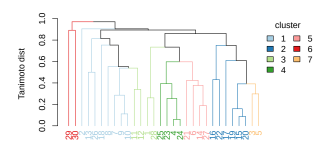
In patRoon hierarchical clustering is used for some componentization algorithms and to cluster candidate compounds with similar chemical structure (see compound clustering). The functions below can be used to visualize their results.

Generic	Classes	Remarks
plot()	All	Plots a dendrogram
<pre>plotInt()</pre>	${\tt componentsIntClust}$	Plots normalized intensity profiles in a cluster
plotHeatMap()	${\tt componentsIntClust}$	Plots an heatmap
plotSilhouettes()componentsClust		Plot silhouette information to determine the
		cluster amount
<pre>plotStructure()</pre>	compoundsCluster	Plots the maximum common substructure
		(MCS) of a cluster

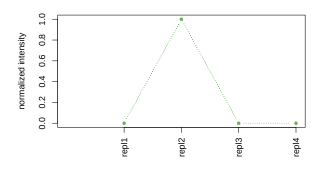
plot(componInt) # dendrogram



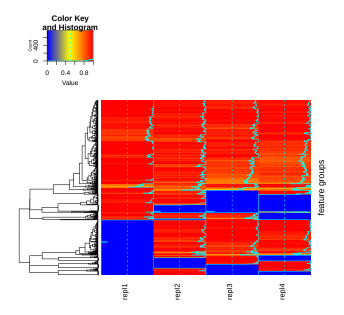
plot(compsClust, groupName = "M120_R268_30") # dendrogram for clustered compounds

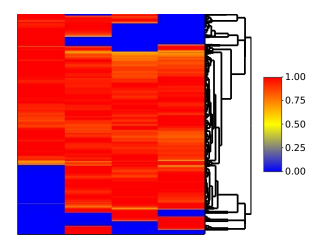


plotInt(componInt, index = 4) # intensities of 4th cluster

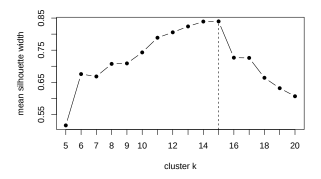


plotHeatMap(componInt) # plot heatmap





plotSilhouettes(componInt, 5:20) # plot silhouettes (e.g. to obtain ideal cluster amount)



5.7.5 Generating EICs in DataAnalysis

If you have Bruker data and the DataAnalysis software installed, you can automatically add EIC data in a DataAnalysis session. The addDAEIC() will do this for a single m/z in one analysis, whereas the addAllDAEICs() function adds EICs for all features in a featureGroups object.

5.8 Interactively explore and review data

The checkFeatures and checkComponents functions start a graphical user interface (GUI) which allows you to interactively explore and review feature and components data, respectively.

```
checkFeatures(fGroups) # inspect features and feature groups
checkComponents(componCAM, fGroups) # inspect components
```

Both functions allow you to easily explore the data in an interactive way. Furthermore, these functions allow you to remove unwanted data. This is useful to remove for example features that are actually noise and feature groups that shouldn't be in the same component. To remove an unwanted feature, feature group or components, simply uncheck its 'keep' checkbox. The next step is to save the selections you made. A *check session* is a file that stores which data should be removed. Once the session file is saved the filter function can be used to actually remove the data:

```
fGroupsF <- filter(fGroups, checkFeaturesSession = TRUE)
componCAMF <- filter(componCAM, checkComponentsSession = TRUE)</pre>
```

If you saved the session and you re-launch the GUI it will restore the selections made earlier. The clearSession argument can be used to fully clear a session before starting the GUI, hence, all the data will be restored to their 'keep state.'

```
checkFeatures(fGroups, clearSession = TRUE) # start GUI with fresh session
```

It is also possible to use multiple different sessions. This is especially useful if you do not want to overwrite previous session data or want to inspect different objects. In this case the session file name should be specified:

```
checkFeatures(fGroups, "mysession.yml")
fGroupsF <- filter(fGroups, checkFeaturesSession = "mysession.yml")</pre>
```

The default session names are "checked-features.yml" and "checked-components.yml" for feature and component data, respectively.

The extension of session file names is .yml since the YAML file format is used. An advantage of this format is that it is easily readable and editable with a text editor.

Note that the session data is tied to the feature group names of your data. This means that, for instance, when you re-group your feature data after changing some parameters, the session data you prepared earlier cannot be used anymore. Since probably quite some manual work went into creating the session file, a special function is available to import a session that was made for previous data. This function tries its best to guess the new feature group name based on similarity of their retention times and m/z values.

```
checkFeatures(fGroups) # do manual inspection

fGroups <- groupFeatures(fList, ...) # re-group with different parameters

importCheckFeaturesSession("checked-features.yml", "checked-features-new.yml", fGroups)

checkFeatures(fGroups, session = "checked-features-new.yml") # inspect new data</pre>
```

Take care to monitor the messages that importCheckFeaturesSession may output, as it may be possible that some 'old' feature groups are not found or are matched by multiple candidates of the new dataset.

Some additional parameters exist to the functions described in this section. As usualy check the reference manual for more details (e.g. ?checkFeatures).

NOTE Although the GUI tools described here allow you to easily filter out results, it is highly recommended to first prioritize your data to avoid doing a lot of unneeded manual work.

5.9 Reporting

The previous sections showed various functionalies to inspect and visualize results. An easy and automated way to do this automatically is by using the *reporting* functionality of patRoon. The following three reporting functions are available:

- reportCSV(): exports workflow data to comma-separated value (csv) files
- reportPDF(): generates simple reports by plotting workflow data in portable document files (PDFs)
- reportHTML(): generates interactive and easily explorable reports

There are many different arguments available to configure the reporting process. Some common arguments are listed below; for a complete listing see the reference manual (e.g. ?reporting).

Argument	Functions	Remarks
fGroups, formulas, compounds, formulas, components, compsCluster	All	Objects to plot. Only fGroups is mandatory.
MSPeakLists	<pre>reportPDF(), reportHTML()</pre>	The MSPeakLists object that was used to generate annotation data. Only needs to be specified if formulas or compounds are reported.
path	All	Directory path where report files will be stored ("report" by default).
formulasTopMost, compoundsTopMost EICOnlyPresent	<pre>reportPDF(), reportHTML() reportPDF(), reportHTML()</pre>	Report no more than this amount of highest ranked candidates. Only plot an EIC for an analysis if a feature was detected.
selfContained	reportHTML()	Outputs to a single and self contained .html file. Handy to share reports, but not recommended for large amounts of data.

Which data will be reported is fully configurable. The only workflow object that must be specified are the feature groups (i.e. with the fGroups argument), all other data (e.g. compounds, components) are optional. This means that reporting can be performed at every stage during the workflow, which, for instance, can be useful to quickly inspect results when testing out various settings to generate workflow data.

When formula or compound results are reported with reportPDF() or reportHTML() then only the top ranked candidates are considered. This limitation is often necessary as reporting many candidates will take considerable time. By default the top 5 for each feature group are reported, however, this number can be changed with the formulasTopMost and compoundsTopMost arguments.

Some typical examples:

6 Sets workflows

In LC-HRMS screening workflows it is typically desired to be able to detect a broad range of chemicals. For this reason, the samples are often measured twice: with positive and negative ionization. Most data processing steps are only suitable for data with the same polarity, for instance, due to the fact that the m/z values in mass spectra are inherently different (e.g. [M+H]+ vs [M-H]-) and MS/MS fragmentation occurs differently. As a result, the screening workflow has to be done twice, which generally requires more time and complicates comparing and interpretation of the complete (positive and negative) dataset.

In patRoon version 2.0 the sets workflow is introduced. This allows you to perform a single non-target screening workflow from different sets of analyses files. Most commonly, each set represents a polarity, hence, there is a positive and negative set. However, more than two sets are supported, and other distinctions between sets are also possible, for instance, samples that were measured with different MS/MS techniques. Another important advantage of the sets workflow is that MS/MS data from different sets can be combined to provide more comprehensive annotations of features. The most important limitation is that (currently) the chromatographic method that was used when analyzing the samples from each set needs to be equal, since retention times are used to group features among the sets.

Performing a sets workflow usually only requires small modifications compared to a 'regular' patRoon workflow. This chapter outlines how to perform such workflows and how to use its unique functionality for data processing. It is assumed that the reader is already familiar with performing 'regular' workflows, which were discussed in the previous chapters.

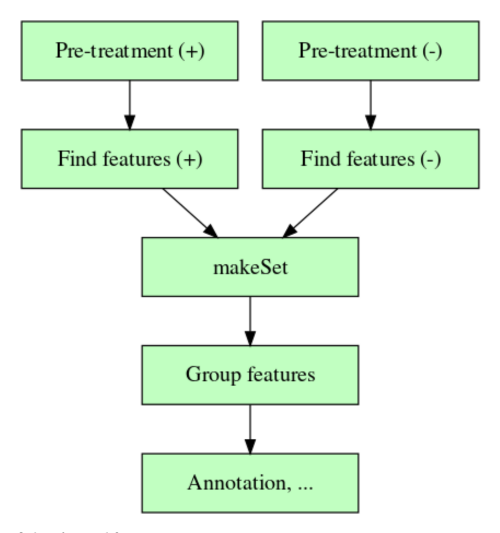
6.1 Initiating a sets workflow

A sets workflow is not much different than a 'regular' (or non-sets) workflow. For instance, consider the following workflow:

This example uses the example data from patRoonData to obtain a feature group dataset, which is cleaned-up afterwards. Then, feature groups are annotated and all the results are reported.

Converting this to a sets workflow:

This workflow will do all the steps for positive and negative data.



Only a few modifications were necessary:

- The analysis information is obtained for positive and negative data (i.e. per set)
- Features are found for each set separately.
- makeSet is used to combine the feature data
- There is no need to specify the adduct anymore in the annotation steps.

NOTE The analysis names for the analysis information must be *unique* for each row, even among sets. Furthermore, replicate groups should not contain analyses from different sets.

The key principle to make sets workflows work is performed by $\mathtt{makeSet}$. This method function takes different features objects (or featureGroups, discussed later) to combine the feature data across sets. During this step features are neutralized: the feature m/z data is converted to neutral feature masses. This step ensures that when features are grouped with $\mathtt{groupFeatures}$, its algorithms are able to find the same feature among different sets, even when different MS ionization modes were used during acquisition. However, please note that (currently) no additional chromatographic alignment steps between sets are performed. For this reason, the chromatographic methodology that is used to acquire the data must be the same for all sets.

The feature neutralization step relies on adduct data. In the example above, it is simply assumed that all features measured with positive mode are protonated (M+H) species, and all negative features are deprotonated (M-H). It is also possible to use adduct annotations for neutralization; this is discussed later.

NOTE The newProject tool can be used to easily generate a sets workflow. Simply select "both" for the *Ionization* option.

6.2 Generating sets workflow data

As was shown in the previous section, the generation of workflow data with a sets workflow largely follows that as what was discussed in the previous chapters. The same generator functions are used:

Workflow step	Function	Output S4 class
Grouping features Suspect screening MS peak lists Formula annotation Compound annotation Componentization	<pre>groupFeatures() screenSuspects() generateMSPeakLists() generateFormulas() generateCompounds() generateComponents()</pre>	featureGroupsSet featureGroupsScreeningSet MSPeakListsSet formulasSet compoundsSet algorithm dependent

(the data pre-treatment and feature finding steps have been omitted as they are not specific to sets workflows).

While the same function generics are used to generate data, the class of the output objects differ (e.g. formulasSet instead of formulas). However, since all these classes *inherit* from their non-sets workflow counterparts, using the workflow data in a sets workflow is nearly identical to what was discussed in the previous chapters (further discussed in the next section).

As discussed before, an important step is the neutralization of features. Other workflow steps also have internal mechanics to deal with data from different sets:

Workflow step	Handling of set data
Finding/Grouping features	Neutralization of m/z values
Suspect screening	Merging results from screening performed for each set
Componentization	Algorithm dependent (discussed below)
MS peak lists	MS data is obtained and stored per set. The final peak lists are
	combined (not averaged)
Formula/Compound annotation	Annotation is performed for each set separately and used to generate a final consensus

In most cases the algorithms of the workflow steps are first performed for each set, and this data is then merged. To illustrate the importance of this, consider these examples

- A suspect screening with a suspect list that contains known MS/MS fragments
- Annotation where MS/MS fragments are used to predict the chemical formula
- Componentization in order to establish adduct assignments for the features

In all cases data is used that is highly dependent on the MS method (eg polarity) that was used to acquire the sample data. Nevertheless, all the steps needed to obtain and combine set data are performed automatically in the background, and are therefore largely invisible.

NOTE Because feature groups in sets workflows always have adduct annotations, it is never required to specify the adduct or ionization mode when generating annotations, components or do suspect screening (*i.e.* the adduct/ionization arguments should not be specified).

6.2.1 Componentization

When the componentization algorithms related to adduct/isotope annotations (e.g. CAMERA, RAMClustR and cliqueMS) and nontarget are used, then componentization occurs per set and the final object (a componentsSet or componentsNTSet) contains all the components together. Since these algorithms are highly dependent upon MS data polarity, no attempt is made to merge components from different sets.

The other componentization algorithms work on the complete data. For more details, see the reference manual (?generateComponents).

6.2.2 Formula and compound annotation

For formula and compound annotation, the data generated for each set is combined to generate a set consensus. The annotation tables are merged, scores are averaged and candidates are re-ranked. More details can be found in the reference manual (e.g. ?generateCompounds). In addition, it possible to only keep candidates that exist in a minimum number of sets. For this, the setThreshold and setThresholdAnn argument can be used:

```
# candidate must be present in all sets
formulas <- generateFormulas(fGroups, mslists, "genform", setThreshold = 1)
# candidate must be present in all sets with annotation data
compounds <- generateCompounds(fGroups, mslists, "metfrag", setThresholdAnn = 1)</pre>
```

In the first example, a formula candidate for a feature group is only kept if it was found for all of the sets. In the second example, a compound candidate is only kept if it was present in all of the sets with annotation data available. The following examples of a common positive/negative sets workflow illustrate the differences:

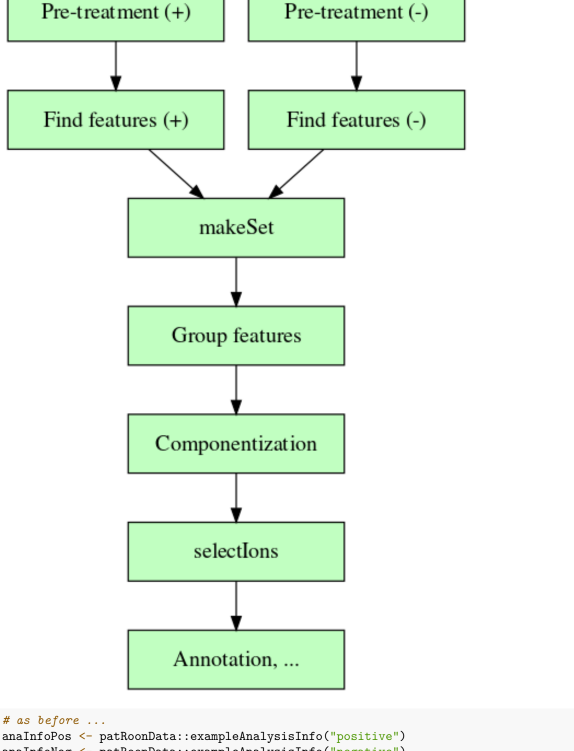
Candidate	annotations	candidate present	setThreshold=1	setThresholdAnn=1
#1	+, -	+, -	Keep	Keep
#2	+, -	+	Remove	Remove
#3	+	+	Remove	Keep

For more information refer to the reference manual (e.g. ?generateCompounds).

6.3 Selecting adducts to improve grouping

The selectIons() and adduct() functions discussed before can also improve sets workflows. This is because the adduct annotations can be used to improve feature neutralization, which in turn will improve grouping features between positive and negative ionization data. Once adduct annotations are set the features will be re-neutralized and re-grouped.

A typical workflow with selectIons looks like this:



```
anaInfoPos <- patRoonData::exampleAnalysisInfo("positive")
anaInfoNeg <- patRoonData::exampleAnalysisInfo("negative")
fListPos <- findFeatures(anaInfoPos, "openms")
fListNeg <- findFeatures(anaInfoNeg, "openms")

fGroupsPos <- groupFeatures(fListPos, "openms")
fGroupsNeg <- groupFeatures(fListNeg, "openms")</pre>
```

The first part of the workflow is exactly the same as was introduced in the beginning of this chapter. Furthermore, note that for sets workflows, selectIons needs a preferential adduct for each set.

The adducts function can also be used to obtain and modify adduct annotations. For sets workflows, these functions operate per set:

```
adducts(fGroups, set = "positive")[1:5]
adducts(fGroups, set = "positive")[4] <- "[M+K]+"</pre>
```

If you want to modify annotations for multiple sets, it is best to delay the re-gouping step:

```
adducts(fGroups, set = "positive", reGroup = FALSE)[4] <- "[M+K]+"
adducts(fGroups, set = "negative", reGroup = TRUE)[10] <- "[M-H20]-"</pre>
```

Setting reGroup=FALSE will not perform any re-neutralization and re-grouping, which preserves feature group names and safes processing time. However, it is **crucial** that the re-grouping step is eventually performed at the end.

6.4 Processing data

All data objects that are generated during a sets workflow *inherit* from the classes from a 'regular' workflow. This means that, with some minor exceptions, *all* of the data processing functionality discussed in the previous chapter (e.g. subsetting, inspection, filtering, plotting, reporting) is also applicable to a sets workflow. For instance, the as.data.table() method can be used for general inspection:

```
as.data.table(compounds)[1:5, c("group", "score", "compoundName", "set")]
```

```
#> group score compoundName set

#> 1: M198_R317_272 4.025217 3-(4-chlorophenyl)-1,1-dimethylurea positive,negative

#> 2: M198_R317_272 1.946688 3-(3-chlorophenyl)-1,1-dimethylurea positive,negative

#> 3: M198_R317_272 1.745251 1-(4-chlorophenyl)-3-ethylurea positive,negative

#> 4: M198_R317_272 1.838637 3-amino-5-chloro-N,N-dimethylbenzamide positive,negative

#> 5: M198_R317_272 2.366149 5-[[(2R)-azetidin-2-yl]methoxy]-2-chloropyridine positive,negative
```

In addition, some the data processing functionality contains additional functionality for a sets workflow:

```
# only keep feature groups that have positive data
fGroupsPos <- fGroups[, sets = "positive"]
# only keep feature groups that have feature data for all sets
fGroupsF <- filter(fGroups, relMinSets = 1)

# only keep feature groups with features present in both polarities
fGroupsPosNeg <- overlap(fGroups, which = c("positive", "negative"), sets = TRUE)
# only keep feature groups with features that are present only in positive mode
fGroupsOnlyPos <- unique(fGroups, which = "positive", sets = TRUE)</pre>
```

And plotting:



The reference manual for the workflow objects contains specific notes applicable to sets workflows (?featureGroups, ?compounds etc).

6.5 Advanced

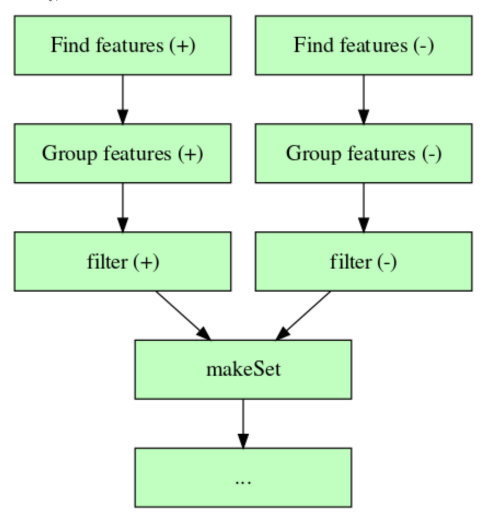
6.5.1 Initiating a sets workflow from feature groups

The makeSet function can also be used to initiate a sets workflow from feature groups:

In this case makeSet combines the positive and negative (un-grouped) features, neutralizes them and regroups them all together (with the algorithm specified by groupAlgo).

While this option involves some extra steps, an advantage is that allows processing the feature data before they are combined, e.g.:

Visually, this workflow looks like this:



Of course, any other processing steps on the feature groups data such as subsetting and visually checking features are also possible before the sets workflow is initiated. Furthermore, it is also possible to perform adduct annotations prior to grouping, which is an alternative way to improve neutralization to what was discussed before.

6.5.2 Inspecting and converting set objects

The following generic functions may be used to inspect or convert data from sets workflows:

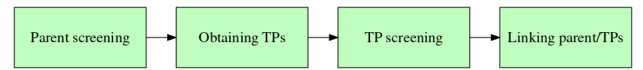
Generic	Purpose	Notes
sets	Return the names of the sets in this object.	
setObjects	Obtain the raw data objects that were used to construct this object.	Not available for features and feature groups.
unset	Converts this object to a regular workflow object.	The set argument must be given to specify which of the set data is to be converted. This function will restore the original m/z values of features.

These methods are heavily used internally, but rarely needed otherwise. More details can be found in the reference manual.

7 Transformation product screening

This chapter describes the various functionality for screening of transformation products (TPs), which are introduced since patRoon 2.0. Screening for TPs, i.e. chemicals that are formed from a parent chemical by e.g. chemical or biological processes, has broad applications. For this reason, the TP screening related functionality is designed to be flexible, thus allowing one to use a workflow that is best suited for a particular study.

Regardless, the TP screening workflow in patRoon can be roughly summarized as follows:



- Parent screening During this step a common patRoon workflow is used to screen for the parent chemicals of interest. This could be a full non-target analysis with compound annotation or a relative simple suspect or target screening.
- **Obtaining TPs** Data is obtained of potential TPs for the parents of interest. The TPs may originate from a library or predicted *in-silico*. Note that in some workflows this step is omitted (discussed later).
- TP screening A suspect screening is performed to find the TPs in the analysis data.
- Linking parents and TPs In the step the parent features are linked with the TP features. Several post-processing functionality exists to improve and prioritize the data.

The next sections will outline more details on these steps are performed and configured. The last section in this chapter outlines several example workflows.

NOTE The new Project tool can be used to easily generate a workflow with transformation product screening.

7.1 Obtaining transformation product data

The generateTPs function is used to obtain TPs for a particular set of parents. Like other workflow generator functions (findFeatures, generateCompounds), several algorithms are available that do the actual work.

Algorithm	Usage	Remarks
BioTransformer	<pre>generateTPs(algorithm = "biotransformer",)</pre>	Predicts TPs with full structural information
Library	<pre>generateTPs(algorithm = "library",)</pre>	Obtains transformation products from a library (PubChem transformations or custom)
Metabolic logic	<pre>generateTPs(algorithm = "logic",)</pre>	Uses pre-defined logic to predict TPs based on common elemental differences (e.g. hydroxylation, demethylation). Based on Schollee et al. (2015).

The biotransformer and library algorithms provide full structural information of the TPs (e.g. formula, SMILES, predicted Log P). However, these algorithms also depend on the full chemical structure of the parent compound. Hence, these algorithms are typically suitable when parents are known in advance or were found by a suspect screening. On the other hand, metabolic logic only requires the feature mass, and this simplicity allows it to predict TPs for all features. This algorithm is most suitable for full non-target analysis, however, extra care must be taken to rule out false positives.

An overview of common arguments for TP generation is listed below.

Argument	Algorithm(s)	Remarks
parents	biotransformer, library	The input parents. See section below.
fGroups	logic	The input feature groups to calculate TPs for.
type	biotransformer	The prediction type: "env", "ecbased", "cyp450", "phaseII", "hgut", "superbio", "allHuman". See BioTransformer for more details.
TPLibrary/transformations adduct	library/logic logic	Custom TP library/transformation rules. The assumed adduct of the parents (e.g. "[M+H]+"). Not needed when adduct annotations are available.

7.1.1 Parent input

The input parent structures for the biotransformer and library algorithms must be one the following:

- A suspect list (follows the same format as suspect screening)
- A feature groups object with screening results (e.g. obtained with screenSuspects, see suspect screening)
- A compounds object obtained with compound annotation

In the former two cases the parent information is taken from the suspect list or from the hits in a suspect screening worklow, respectively. The last case is more suitable for when the parents are not completely known. In this case, the candidate structures from a compound annotation are used as input to obtain TPs. Since *all* the candidates are used, it is highly recommend to filter the object in advance, for instance, with

the topMost filter. For library, the parent input is optional: if no parents are specified then TP data for all parents in the database is used.

For the logic algorithm TPs are predicted directly for feature groups. Since this algorithm can only perform very basic validity checks, it is strongly recommended to first prioritize the feature group data.

Some typical examples:

7.1.2 Processing data

Similar to other workflow data, several generic functions are available to inspect the data:

Generic	Classes	Remarks
length()	All	Returns the total number of transformation products
names()	All	Returns the names of the parents
<pre>parents()</pre>	All	Returns a table with information about the parents
<pre>products()</pre>	All	Returns a list with for each parent a table with TPs
as.data.table(),	All	Convert all the object information into a
as.data.frame		data.table/data.frame
"[[" / " $$$ " operators	All	Extract TP information for a specified parent

Some examples:

TPs[["DEET"]][, ...cols]

```
#> name formula InChIKey
#> 1: (-)-bisoprolol C18H31NO4 VHYCDWMUTMEGQY-KRWDZBQOSA-N
#> 2: (-)-Epigallocatechin C15H14O7 XMOCLSLCDHWDHP-IUODEOHRSA-N
#> 3: (-)-Epigallocatechin gallate C22H18O11 WMBWREPUVVBILR-WIYYLYMNSA-N
#> 4: (-)-ethosuximide C7H11NO2 HAPOVYFOVVWLRS-SSDOTTSWSA-N
#> 5: (-)-etodolac C17H21NO3 NNYBQONXHNTVIJ-QGZVFWFLSA-N
```

```
#> name formula InChIKey
#> 1: DEET-TP-N,N-diethyl-m-hydroxymethylbenzamide C12H17NO2 FRZJZRVZZNTMAW-UHFFFAOYSA-N
```

```
#> 2: DEET-TP-N-Ethyl-N-(2-hydroxyethyl)-3-methylbenzamide C12H17NO2 KVTUZBGZTRABBQ-UHFFFAOYSA-N
#> 3:
                                      DEET-TP-acetaldehyde
                                                                C2H4O IKHGUXGNUITLKF-UHFFFAOYSA-N
#> 4:
                               DEET-TP-N-ethyl-m-toluamide C10H13NO FPINATACRXASTP-UHFFFAOYSA-N
TPs[[2]][, ..cols]
#>
                                                                       name
                                                                              formula
#> 1: (-)-Epigallocatechin-TP-(-)-Epigallocatechin, 3p-hydroxy-glucuronide C21H22O13 CQDATFYRGVZXLF-ZVH
as.data.table(TPs)[1:5, 1:3]
                            parent
#>
                                                                  transformation
                                             O-deisopropylation / Human Phase I
#> 1:
                    (-)-bisoprolol
```

#> 3: (-)-Epigallocatechin gallate Aromatic-OH glucuronidation / Human Phase II (-)-Epigallocatechin ga
#> 4: (-)-Epigallocatechin gallate Aromatic-OH glucuronidation / Human Phase II (-)-Epigallocatechin ga

Aliphatic Hydroxylation / Human Phase I

(-)-Epi

(-)-eth

In addition, the following generic functions are available to modify or convert the object data:

(-)-ethosuximide

#> 5:

Generic	Classes	Remarks
"[" operator	All	Subset this object on given parents
filter	transformationProductsBT Filters this object	
${\tt convertToSuspects}$	All	Generates a suspect list of all TPs (and optionally
		parents) that is suitable for screenSuspects
${\tt convertToMFDB}$	transformationProductsBT,Generates a MetFrag database for all TPs (and	
	${\tt transformationProductsLib pation} {\tt nally parents})$	

(-)-Epigallocatechin Aromatic-OH glucuronidation / Human Phase II

```
TPs2 <- TPs[1:10] # only keep results for first ten parents
# remove transformation products that are isomers to their parent or sibling TPs
# may simplify data as these are often difficult to identify
TPsF <- filter(TPs, removeParentIsomers = TRUE, removeTPIsomers = TRUE)</pre>
# remove duplicate transformation products from each parent
# these can occur if different pathways yield the same TPs
TPsF <- filter(TPs, removeDuplicates = TRUE)</pre>
# only keep TPs that have a structural similarity to their parent of >= 0.5
TPsF <- filter(TPs, minSimilarity = 0.5)
# do a suspect screening for all TPs and their parents
suspects <- convertToSuspects(TPs, includeParents = TRUE)</pre>
fGroupsScr <- screenSuspects(fGroups, suspects, onlyHits = TRUE)</pre>
# use the TP data for a specialized MetFrag database
convertToMFDB(TPs, "TP-database.csv", includeParents = FALSE)
compoundsTPs <- generateCompounds(fGroups, mslists, "metfrag", database = "csv",
                                   extraOpts = list(LocalDatabasePath =

¬ "TP-database.csv"))
```

The convertToSuspects function is always part of a workflow with biotransformer or library TPs. This is discussed further in the next section. The convertToMFDB function is especially handy with biotransformer workflows, as it allows generating a compound database for TPs that may not be available in other databases. This is further demonstrated in the first example.

7.1.3 Custom libraries and transformations

By default the library and logic algorithms use data that is installed with patRoon (based on PubChem transformations and Schollee et al. (2015), respectively). However, it is also possible to use custom data.

To use a custom TP library a simple data.frame is needed with the names, SMILES and optionally log P values for the parents and TPs. The log P values are used for prediction of the retention time direction of a TP compared to its parent, as is discussed further in the next section. The following small library has two TPs for benzotriazole and one for DEET:

```
#> parent_name parent_SMILES TP_name TP_SMILES
#> 1 1H-Benzotriazole C1=CC2=NNN=C2C=C1 1-Methylbenzotriazole C1=CC2=CC2C=C2N=N1
#> 2 1H-Benzotriazole C1=CC2=NNN=C2C=C1 1-Hydroxybenzotriazole C1=CC=C2C(=C1)N=NN2O
#> 3 DEET CCN(CC)C(=0)C1=CC=CC(=C1)C N-ethyl-m-toluamide CCNC(=0)C1=CC=CC(=C1)C
```

To use this library, simply pass it to the TPLibrary argument:

```
TPs <- generateTPs("library", TPLibrary = myTPLib)</pre>
```

Similarly, for logic a table with custom transformation rules can be specified for TP calculations:

```
#> transformation add sub retDir
#> 1 hydroxylation 0 -1
#> 2 demethylation CH2 -1
```

The add and sub columns are used to denote the elements that are added or subtracted by the reaction. These are used to calculate mass differences between parents and TPs. The retDir column is used to indicate the retention time direction of the parent compared to the TP: -1 (elutes before parent), 1 (elutes after parent) or 0 (similar or unknown). The next section describes how this data can be used to filter TPs. The custom rules can be used by passing them to the transformations argument:

7.2 Linking parent and transformation product features

This section discusses one of the most important steps in a TP screening workflow, which is to link feature groups of parents with those of candidate transformation products. During this step, *components* are made, where each component consist of one or more feature groups of detected TPs for a particular parent. Note that componentization was already introduced before, but for very different algorithms. However, the data format for TP componentization is highly similar. After componentization, several filters are available to clean and prioritize the data. These can even allow workflows without obtaining potential TPs in advance, which is discussed in the last subsection.

7.2.1 Componentization

Like other algorithms, the generateComponents generic function is used to generate TP components, by setting the algorithm parameter to "tp".

The following arguments are of importance:

Argument	Remarks
fGroups	The input feature groups for the parents
fGroupsTPs	The input feature groups for the TPs
ignoreParents	Set to TRUE to ignore feature groups in fGroupsTPs that also occur in fGroups
TPs	The input transformation products, ie as generated by generateTPs()
${ t MSPeakLists},$	Annotation objects used for similarity calculation between the parent and its TPs
formulas,	
compounds	
minRTDiff	The minimum retention time difference (seconds) of a TP for it to be considered to elute differently than its parent.

7.2.1.1 Feature group input The fGroups, fGroupsTPs and ignoreParents arguments are used by the componentization algorithm to identify which feature groups can be considered as parents and which as TPs. Three scenarios are possible:

- 1. fGroups=fGroupsTPs and ignoreParents=FALSE: in this case no distinction is made, and all feature groups are considered a parent or TP (default if fGroupsTPs is not specified).
- 2. fGroups and fGroupsTPs contain different subsets of the *same* featureGroups object and ignoreParents=FALSE: only the feature groups in fGroups/fGroupsTPs are considered as parents/TPs.
- 3. As above, but with ignoreParents=TRUE: the same distinction is made as above, but any feature groups in fGroupsTPs are ignored if also present in fGroups.

The first scenario is often used if it is unknown which feature groups may be parents or which are TPs. Furthermore, this scenario may also be used if the dataset is sufficiently simple, for instance, because a suspect screening with the results from convertToSuspects (discussed in the previous section) would reliably discriminate between parents and TPs. A workflow with the first scenario is demonstrated in the second example.

In all other cases it is recommended to use either the second or third scenario, since making a prior distinction between parent and TP feature groups greatly simplifies the dataset and reduces false positives. A relative simple example where this can be used is when there are two sample groups: before and after treatment.

In this example, only those feature groups present in the "before" replicate group are considered as parents, and those in "after" may be considered as a TP. Since it is likely that there will be some overlap in feature groups between both sample groups, the <code>ignoreParents</code> flag can be used to not consider any of the overlap for TP assignments:

More sophisticates ways are of course possible to provide an upfront distinction between parent/TP feature groups. In the fourth example a workflow is demonstrated where fold changes are used.

NOTE The feature groups specified for fGroups/fGroupsTPs must always originate from the same featureGroups object.

For the library and biotransformer algorithms it is mandatory that a suspect screening of parents and TPs is performed prior to componentization. This is necessary for the componentization algorithm to map the feature groups that belong to a particular parent or TP. To do so, the convertToSuspects function is used to prepare the suspect list:

If a parent screening was already performed in advance, for instance when the input parents to generateTPs are screening results, the screening results for parents and TPs can also be combined. The second example demonstrates this.

Note that in the case a parent suspect is matched to multiple feature groups, a component is made for each match. Similarly, if multiple feature groups match to a TP suspect, all of them will be incorporated in the component.

When TPs were generated with the logic algorithm a suspect screening must also be carried out in advance. However, in this case it is not necessary to include the parents (since each parent equals a feature group no mapping is necessary). The onlyHits variable to screenSuspects must not be set in order to keep the parents.

```
# only screen for TPs
suspects <- convertToSuspects(TPs, includeParents = FALSE)
# but keep all other feature groups as these may be parents
fGroupsScr <- screenSuspects(fGroups, suspects, onlyHits = FALSE)
# do the componentization...</pre>
```

7.2.1.2 Annotation similarity calculation If additional annotation data for parents and TPs is given to the componentization algorithm, it will be used to calculate various similarity properties. Often, the chemical structure for a transformation product is similar to that of its parent. Hence, there is a good chance that a parent and its TPs also share similar MS/MS data.

Firstly, if MS peak lists are provided, then the spectrum similarity is calculated between each parent and its potential TP candidates. This is performed with all the three different alignment shifts (see the spectrum similarity section for more details).

In case formulas and/or compounds objects are specified, then a parent/TP comparison is made by counting the number of fragments and neutral losses that they share (by using the formula annotations). This property is mainly used for non-target workflows where the identity for a parent and TP is not yet well established. For this reason, fragments and neutral losses reported for all candidates for the parent/TP feature group are considered. Hence, it is highly recommend to pre-treat the annotation objects, for instance, with the topMost filter. If both formulas and compounds are given the results are pooled. Note that each unique fragment/neutral loss is only counted once, thus multiple formula/compound candidates with the same annotations will not skew the results.

7.2.2 Processing data

The output of TP componentization is an object of the componentsTPs class. This *derives* from the 'regular' components class, therefore, all the data processing functionality described before (extraction, subsetting, filtering etc) are also valid for TP components.

Several additional filters are available to prioritize the data:

Filter	Remarks
retDirMatch	If TRUE only keep TPs with an expected chromatographic retention direction compared to the parent.
<pre>minSpecSim, minSpecPrec, minSpecSimBoth</pre>	The minimum spectrum similarity between the parent and TP. Calculated with no, "precursor" and "both" alignment shifting (see spectrum similarity).
minFragMatches, minNLMatches formulas	Minimum number of formula fragment/neutral loss matches between parent and TP (discussed in previous section). A formulas object used to further verify candidate TPs that were generated by the logic algorithm.

The retDirMatch filter compares the expected and observed retention time direction of a TP in order to decide if it should be kept. The direction is a value of either -1 (TP elutes before parent), +1 (TP elutes after parent) or 0 (TP elutes very close to the parent or its direction is unknown). The directions are taken from the generated transformation products. For the library and biotransformer algorithms the log P values are compared of a TP and its parent. Here, it is assumed that lower log P values result in earlier elution (i.e. typical with reversed phase LC). For the logic algorithm the retention time direction is taken from the transformation rules table. Note that specifying a large enough value for the minRTDiff argument to generateComponents is important to ensure that some tolerance exists while comparing retention time directions of parent and TPs. This filter does nothing if either the observed or expected direction is zero.

When TPs data was generated with the logic algorithm it is recommended to use the formulas filter. This filter uses formula annotations to verify that (1) a parent feature group contains the elements that are subtracted during the transformation and (2) the TP feature group contains the elements that were added during the transformation. Since the 'right' candidate formula is most likely not yet known, this filter looks at all candidates. Therefore, it is recommended to filter the formulas object, for instance, with the topMost filter.

7.2.3 Omitting transformation product input

The TPs argument to generateComponents can also be omitted. In this case every feature group of fGroupTPs is considered to be a potential TP for the potential parents specified for fGroups. An advantage is that the screening workflow is not limited to any known TPs or transformations. However, such a workflow has high demands on prioritiation steps before and after the componentization to rule out the many false positives that may occur.

When no transformation data is supplied it is crucial to make a prior distinction between parent and TP feature groups. Afterwards, the MS/MS spectral and other annotation similarity filters mentioned in the previous section may be a powerful way to further prioritize data.

The fourth example demonstrates such a workflow.

7.2.4 Reporting TP components

The TP components can be reported with the reportHTML function. This is done by setting the components function argument (i.e. equally to all other component types). The results will be displayed with a customized format that allows easy exploring of each parent with its TPs.

```
reportHTML(fGroups, components = componTP)
```

7.3 Example workflows

The next subsections demonstrate several approaches to perform a TP screening workflow with patRoon. In all examples it is assumed that feature groups were already obtained (with the findFeatures and groupFeatures functions) and stored in the fGroups variable.

The workflows with patRoon are designed to be flexible, and the examples here are primarily meant to implement your own workflow. Furthermore, some of the techniques used in the examples can also be combined. For instance, the Fold change classification and MS/MS similarity filters applied in the fourth example could also be applied to any of the other examples.

7.3.1 Screen predicted TPs for targets

The first example is a simple workflow where TPs are predicted for a set of given parents with BioTransformer and subsequently screened. A MetFrag compound database is generated and used for annotation.

7.3.2 Screening TPs from a library for suspects

In this example TPs of interest are obtained for the parents that surfaced from of a suspect screening. The steps of this workflow are:

- 1. Suspect screening parents.
- 2. Obtain TPs for the suspect hits from a library.
- 3. A second suspect screening is performed for TPs and the original parent screening results are amended. Note that the parent data is needed for componentization.
- 4. Both parents and TPs are annotated using a database generated from their chemical structures.
- 5. Some prioritization is performed by
 - a. Only keeping candidate structures for which in-silico fragmentation resulted in at least one annotated MS/MS peak.
 - b. Only keeping suspect hits with an estimated identification level of 3 or better.
- 6. The TP components are made and only feature groups with parent/TP assignments are kept.
- 7. All results are reported.

```
# step 1
fGroupsScr <- screenSuspects(fGroups, patRoonData::suspectsPos, adduct = "[M+H]+")
# step 2
TPs <- generateTPs("library", parents = fGroupsScr)</pre>
# step 3
suspects <- convertToSuspects(TPs)</pre>
fGroupsScr <- screenSuspects(fGroupsScr, suspects, adduct = "[M+H]+", onlyHits = TRUE,

→ amend = TRUE)

# step 4
mslistsScr <- generateMSPeakLists(fGroupsScr, "mzr")</pre>
convertToMFDB(TPs, "TP-database.csv", includeParents = TRUE)
compoundsScr <- generateCompounds(fGroupsScr, mslistsScr, "metfrag", adduct = "[M+H]+",</pre>

    database = "csv",

                                   extraOpts = list(LocalDatabasePath =

¬ "TP-database.csv"))

# step 5a
compoundsScr <- filter(compoundsScr, minExplainedPeaks = 1)</pre>
# step 5b
fGroupsScrAnn <- annotateSuspects(fGroupsScr, MSPeakLists = mslistsScr,
                                   compounds = compoundsScr)
fGroupsScrAnn <- filter(fGroupsScrAnn, maxLevel = 3, onlyHits = TRUE)
# step 6
componTP <- generateComponents(fGroupsScrAnn, "tp", TPs = TPs, MSPeakLists = mslistsScr,</pre>
                                compounds = compoundsScr)
fGroupsScrAnn <- fGroupsScrAnn[results = componTP]</pre>
# step 7
reportHTML(fGroupsScrAnn, MSPeakLists = mslistsScr, compounds = compoundsScr,
           components = componTP)
```

7.3.3 Non-target screening of predicted TPs

This example uses metabolic logic to calculate possible TPs for all feature groups from a complete non-target screening. This example demonstrates how a workflow can be performed when little is known about the identity of the parents. The steps of this workflow are:

- 1. Formula annotations are performed for all feature groups.
- 2. These results are then limited to the top 5 candidates, and only feature groups with annotations are kept.
- $3. \ \,$ The TPs are calculated for all remaining feature groups.
- 4. A suspect screening is performed to find the TPs. Unlike the previous example feature groups without hits are kept (discussed here).
- 5. The components are generated
- 6. The components are filtered:
 - a. The TPs must follow an expected retention time direction
 - b. The parent/TPs should have at least one candidate formula that fits with the transformation.
- 7. Only feature groups are kept with parent/TP assignments and all results are reported.

```
mslists <- generateMSPeakLists(fGroups, "mzr")</pre>
formulas <- generateFormulas(fGroups, mslists, "genform", adduct = "[M+H]+")
formulas <- filter(formulas, topMost = 5)</pre>
fGroups <- fGroups[results = formulas]</pre>
# step 3
TPs <- generateTPs("logic", fGroups = fGroups, adduct = "[M+H]+")
# step 4
suspects <- convertToSuspects(TPs)</pre>
fGroupsScr <- screenSuspects(fGroups, suspects, adduct = "[M+H]+", onlyHits = FALSE)
# step 5
componTP <- generateComponents(fGroupsScr, "tp", TPs = TPs, MSPeakLists = mslists,</pre>

    formulas = formulas)

# step 6
componTP <- filter(componTP, retDirMatch = TRUE, formulas = formulas)</pre>
# step 7
fGroupsScr <- fGroupsScr[results = componTP]</pre>
reportHTML(fGroupsScr, MSPeakLists = mslists, formulas = formulas, components = componTP)
```

7.3.4 Non-target screening of TPs by annotation similarities

This example shows a workflow where no TP data from a prediction or library is used. Instead, this workflow relies on statistics and MS/MS data to find feature groups which may potentially have a parent - TP relationship. The workflow is similar to that of the previous example. The steps of this workflow are:

- 1. Fold changes (FC) between two sample groups are calculated to classify which feature groups are decreasing (i.e. parents) or increasing (i.e. TPs).
- 2. Feature groups without classification are removed.

- 3. Formula annotations are performed like the previous example.
- 4. The componentization is performed and the FC classifications are used to specify which feature groups are to be considered parents or TPs.
- 5. Only TPs are kept that show a high MS/MS spectral similarity and share at least one fragment with their parent.
- 6. Only feature groups are kept with parent/TP assignments and all results are reported.

```
# step 1
tab <- as.data.table(fGroups, FCParams = getFCParams(c("before", "after")))</pre>
groupsParents <- tab[classification == "decrease"]$group</pre>
groupsTPs <- tab[classification == "increase"]$group</pre>
# step 2
fGroups <- fGroups[, union(groupsParents, groupsTPs)]</pre>
# step 3
mslists <- generateMSPeakLists(fGroups, "mzr")</pre>
formulas <- generateFormulas(fGroups, mslists, "genform", adduct = "[M+H]+")</pre>
formulas <- filter(formulas, topMost = 5)</pre>
fGroups (- fGroups[results = formulas]
# step 4
componTP <- generateComponents(algorithm = "tp",</pre>
                                 fGroups = fGroups[, groupsParents],
                                 fGroupsTPs = fGroups[, groupsTPs],
                                 MSPeakLists = mslists, formulas = formulas)
# step 5
componTP <- filter(componTP, minSpecSimBoth = 0.75, minFragMatches = 1)</pre>
fGroups <- fGroups[results = componTP]</pre>
reportHTML(fGroups, MSPeakLists = mslists, formulas = formulas, components = componTP)
```

8 Advanced usage

8.1 Adducts

When generating formulae and compound annotations and some other functionalities it is required to specify the adduct species. Behind the scenes, different algorithms typically use different formats. For instance, in order to specify a protonated species...

- GenForm either accepts "M+H" and "+H"
- MetFrag either accepts the numeric code 1 or "[M+H]+"
- SIRIUS accepts "[M+H]+"

In addition, most algorithms only accept a limited set of possible adducts, which do not necessarily all overlap with each other. The GenFormAdducts() and MetFragAdducts() functions list the possible adducts for GenForm and MetFrag, respectively.

In order to simplify the situation patRoon internally uses its own format and converts it automatically to the algorithm specific format when necessary. Furthermore, during conversion it checks if the specified adduct

format is actually allowed by the algorithm. Adducts in patRoon are stored in the adduct S4 class. Objects from this class specify which elements are added and/or subtracted, the final charge and the number of molecules present in the adduct (e.g. 2 for a dimer).

```
adduct(add = "H") # [M+H]+
adduct(sub = "H", charge = -1) # [M-H]-
adduct(add = "K", sub = "H2", charge = -1) # [M+K-H2]-
adduct(add = "H3", charge = 3) # [M+H3]3+
adduct(add = "H", molMult = 2) # [2M+H]+
```

A more easy way to generate adduct objects is by using the as.adduct() function:

```
as.adduct("[M+H]+")
as.adduct("[M+H2]2+")
as.adduct("[2M+H]+")
as.adduct("[M-H]-")
as.adduct("+H", format = "genform")
as.adduct(1, isPositive = TRUE, format = "metfrag")
```

In fact, the adduct argument to workflow functions such as generateFormulas() and generateCompounds() is automatically converted to an adduct class with the as.adduct() function if necessary:

```
formulas <- generateFormulas(..., adduct = adduct(sub = "H", charge = -1))
formulas <- generateFormulas(..., adduct = "[M-H]-") # same as above</pre>
```

More details can be found in the reference manual (?adduct and ?`adduct-utils`).

8.2 Feature parameter optimization

Many different parameters exist that may affect the output quality of feature finding and grouping. To avoid time consuming manual experimentation, functionality is provided to largely automate the optimization process. The methodology, which uses design of experiments (DoE), is based on the excellent Isotopologue Parameter Optimization (IPO) R package. The functionality of this package is directly integrated in patRoon. Some functionality was added or changed, the most important being support for other feature finding and grouping algorithms besides XCMS and basic optimization support for qualitative parameters. Nevertheless, the core optimization algorithms are largely untouched.

This section will introduce the most important concepts and functionality. Please see the reference manual for more information (e.g. ?`feature-optimization`).

NOTE The SIRIUS and SAFD algorithms are currently not (yet) supported.

8.2.1 Parameter sets

Before starting an optimization experiment we have to define *parameter sets*. These sets contain the parameters and (initial) numeric ranges that should be tested. A parameter set is defined as a regular list, and can be easily constructed with the <code>generateFeatureOptPSet()</code> and <code>generateFGroupsOptPSet()</code> functions (for feature finding and feature grouping, respectively).

```
pSet <- generateFeatureOptPSet("openms") # default test set for OpenMS

pSet <- generateFeatureOptPSet("openms", chromSNR = c(5, 10)) # add parameter

# of course manually making a list is also possible (e.g. if you don't want to test the

→ default parameters)

pSet <- list(noiseThrInt = c(1000, 5000))
```

When optimizing with XCMS or KPIC2 a few things have to be considered. First of all, when using the XCMS3 interface (i.e. algorithm="xcms3") the underlying method that should be used for finding and grouping features and retention alignment should be set. In case these are not set default methods will be used.

In addition, when optimizing feature grouping (both XCMS interfaces and KPIC2) we need to set the grouping and retention alignment parameters in two different nested lists: these are groupArgs/retcorArgs (algorithm="xcms"), groupParams/retAlignParams (algorithm="xcms3") or groupArgs/alignArgs (algorithm="kpic2").

```
pSetFG <- list(groupParams = list(bw = c(20, 30))) # xcms3
pSetFG <- list(retcorArgs = list(gapInit = c(0, 7))) # xcms
pSetFG <- list(groupArgs = list(mz_weight = c(0.3, 0.9))) # kpic2</pre>
```

When a parameter set has been defined it should be used as input for the optimizeFeatureFinding() or optimizeFeatureGrouping() functions.

Similar to findFeatures(), the first argument to optimizeFeatureFinding() should be the analysis information. Note that it is not uncommon to perform the optimization with only a subset of (representative) analyses (i.e. to reduce processing time).

```
ft0pt <- optimizeFeatureFinding(anaInfo[1:2, ], "openms", pSet) # only use first two \rightarrow analyses
```

From the parameter set a design of experiment will be automatically created. Obviously, the more parameters are specified, the longer such an experiment will take. After an experiment has finished, the optimization algorithm will start a new experiment where numeric ranges for each parameter are increased or decreased in order to more accurately find optimum values. Hence, the numeric ranges specified in the parameter set are only *initial* ranges, and will be changed in subsequent experiments. After each experiment iteration the results will be evaluated and a new experiment will be started as long as better results were obtained during the last experiment (although there is a hard limit defined by the maxIterations argument).

For some parameters it is recommended or even necessary to set hard limits on the numeric ranges that are allowed to be tested. For instance, setting a minimum feature intensity threshold is highly recommended

to avoid excessive processing time and potentially suboptimal results due to excessive amounts of resulting features. Configuring absolute parameter ranges is done by setting the paramRanges argument.

Depending on the used algorithm, several default absolute limits are imposed. These may be obtained with the getDefFeaturesOptParamRanges() and getDefFGroupsOptParamRanges() functions.

The common operation is to optimize numeric parameters. However, parameters that are not numeric (i.e. *qualitative*) need a different approach. In this case you will need to define multiple parameter sets, where each set defines a different qualitative value.

In the above example there are two parameter sets: both define the numeric chromFWHM parameter, whereas the qualitative isotopeFilteringModel parameter has a different value for each. Note that we had to specify the chromFWHM twice, this can be remediated by using the templateParams argument:

As its name suggests, the templateParams argument serves as a template parameter set, and its values are essentially combined with each given parameter set. Note that current support for optimizing qualitative parameters is relatively basic and time consuming. This is because tests are essentially repeated for each parameter set (e.g. in the above example the chromFWHM parameter is optimized twice, each time with a different value for isotopeFilteringModel).

8.2.2 Processing optmization results

The results of an optimization process are stored in objects from the S4 optimizationResult class. Several methods are defined to inspect and visualize these results.

The optimizedParameters() function is used to inspect the best parameter settings. Similarly, the optimizedObject() function can be used to obtain the object that was created with these settings (i.e. a features or featureGroups object).

```
optimizedParameters(ftOpt) # best settings for whole experiment
```

```
#> $chromFWHM
#> [1] 3.75
#>
```

```
#> $mzPPM
#> [1] 13.5
#>
#> $minFWHM
#> [1] 1.4
#>
#> $maxFWHM
#> [1] 24
optimizedObject(ftOpt) # features object with best settings for whole experiment
#> A featuresOpenMS object
#> Hierarchy:
#> features
       |-- featuresOpenMS
#> ---
#> Object size (indication): 595 kB
#> Algorithm: openms
#> Total feature count: 3681
#> Average feature count/analysis: 3681
#> Analyses: solvent-pos-1 (1 total)
#> Replicate groups: solvent-pos (1 total)
#> Replicate groups used as blank: solvent-pos (1 total)
Results can also be obtained for specific parameter sets/iterations.
optimizedParameters(ftOpt, 1) # best settings for first parameter set
#> $chromFWHM
#> [1] 3.75
#> $mzPPM
#> [1] 13.5
#>
#> $minFWHM
#> [1] 1.4
#>
#> $maxFWHM
#> [1] 24
optimizedParameters(ftOpt, 1, 1) # best settings for first parameter set and experiment
\hookrightarrow iteration
#> $chromFWHM
#> [1] 5
#>
#> $mzPPM
#> [1] 10
#>
#> $minFWHM
#> [1] 1
```

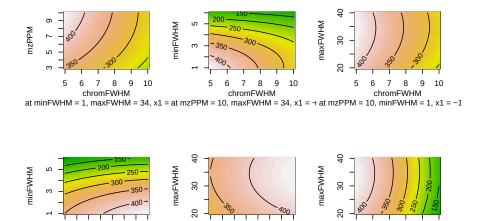
```
#>
#> $maxFWHM
#> [1] 34
```

optimizedObject(ftOpt, 1) # features object with best settings for first parameter set

The plot() function can be used to visualize optimization results. This function will plot graphs for results of all tested parameter pairs. The graphs can be contour, image or perspective plots (as specified by the type argument).

```
plot(ftOpt, paramSet = 1, DoEIteration = 1) # contour plots for first param

→ set/experiment
```



3 4 5 6 7 8 9

t chromFWHM = 5, maxFWHM = 34, x2 at chromFWHM = 5, minFWHM = 1, x2 = at chromFWHM = 5, mzPPM = 10, x3 = -

7 8

plot(ftOpt, paramSet = 1, DoEIteration = 1, type = "persp") # pretty perspective plots

minFWHM







at minFWHM = 1, maxFWHM = 34, x1 = at mzPPM = 10, maxFWHM = 34, x1 = \Rightarrow at mzPPM = 10, minFWHM = 10, x1 = \Rightarrow at mzPPM = 10, minFWHM = 10, x1 = \Rightarrow at mzPPM = 10, minFWHM = 10, x1 = \Rightarrow at mzPPM = 10, minFWHM = 10, x1 = \Rightarrow at mzPPM = 10, minFWHM = 10, x1 = \Rightarrow at mzPPM = 10, x1 = \Rightarrow







t chromEWHM = 5, maxEWHM = 34, x2 at chromEWHM = 5, minEWHM = 1, x2 = at chromEWHM = 5, mzPPM = 10, x3 = -

Please refer to the reference manual for more methods to inspect optimization results (e.g. ?optimizationResult).

8.3 Chromatographic peak qualities

The algorithms used by findFeatures detect chromatographic peaks automatically to find the features. However, it is common that not all detected features have 'proper' chromatographic peaks, and some features could be just noise. The MetaClean R package supports various quality measures for chromatographic peaks. The quality measures include Gaussian fit, symmetry, sharpness and others. In addition, MetaClean averages all feature data for each feature group and adds a few additional group specific quality measures (e.g. retention time consistency). Please see Chetnik, Petrick, and Pandey (2020) for more details. The calculations are integrated into patRoon, and are easily performed with the calculatePeakQualities() generic function.

```
fList <- calculatePeakQualities(fList) # calculate for all features
fGroups <- calculatePeakQualities(fGroups) # calculate for all features and groups
```

Most often the featureGroups method is only used, unless you want to filter features (discussed below) prior to grouping.

An extension in patRoon is that the qualities are used to calculate *peak scores*. The score for each quality measure is calculated by normalizing and scaling the values into a 0-1 range, where zero is the worst and one the best. Note that most scores are relative, hence, the values should only be used to compare features among each other. Finally, a totalScore is calculated which sums all individual scores and serves as a rough overall score indicator for a feature (group).

The qualities and scores are easily obtained with the as.data.table() function.

```
# (limit rows/columns for clarity)
as.data.table(fList)[1:5, 26:30]
```

```
GaussianSimilarityScore SharpnessScore TPASRScore ZigZagScore totalScore
#> 1:
                    0.6314046
                                3.443351e-02 0.9956949
                                                           0.9103221
                                                                       6.302180
                    0.9633994
                                9.900530e-10 0.9944988
#> 2:
                                                           0.3565674
                                                                       6.513205
                    0.3613087
#> 3:
                                7.565147e-10 0.8006569
                                                           0.9999449
                                                                       5.651379
```

```
#> 5: 0.3676623 1.000000e+00 0.9907657 0.8435805 5.825267

# the qualities argument is necessary to include the scores.

# valid values are: "quality", "score" or "both"
as.data.table(fGroups, qualities = "both")[1:5, 25:29]
```

8.600747e-03 0.9405262

0.9637153

5.892201

```
{\tt TPASRScore}\ {\tt ZigZagScore}\ {\tt ElutionShiftScore}\ {\tt RetentionTimeCorrelationScore}\ {\tt totalScore}
#>
#> 1: 0.7305554
                   0.9962254
                                      0.8421657
                                                                      0.9937370
                                                                                  7.930701
#> 2: 0.0000000
                  0.9744541
                                      0.9960804
                                                                                  5.962788
                                                                      0.6808412
#> 3: 0.6140008 0.9171568
                                      0.9015949
                                                                      0.9683740
                                                                                  7.471384
#> 4: 0.8227904
                  0.8907734
                                      0.9403958
                                                                      0.9948720
                                                                                  8.450124
#> 5: 0.9848653
                  0.8667116
                                      0.5754979
                                                                      0.9978621
                                                                                  8.739507
```

The feature quality values can also be reviewed interactively with reports generated with reportHTML (see Reporting) and with checkFeatures (see here). The filter function can be used filter out low scoring features and feature groups:

8.3.1 Applying machine learning with MetaClean

0.9151027

#> 4:

An important feature of MetaClean is to use the quality measures to train a machine learning model to automatically recognize 'good' and 'bad' features. patRoon provides a few extensions to simplify training and using a model. Furthermore, while MetaClean was primarily designed to work with XCMS, the extensions of patRoon allow the usage of data from all the algorithms supported by patRoon.

The getMCTrainData function can be used to convert data from a feature check session to training data that can be used by MetaClean. This allows you to use interactively select good/bad peaks. The workflow looks like this:

```
# untick the 'keep' checkbox for all 'bad' feature groups
checkFeatures(fGroupsTrain, "train_session.yml")

# get train data. This gives comparable data as MetaClean::getPeakQualityMetrics()
trainData <- getMcTrainData(fGroupsTrain, "train_session.yml")

# use train data with MetaClean with MetaClean::runCrossValidation(),
# MetaClean::getEvaluationMeasures(), MetaClean::trainClassifier() etc
# --> see the MetaClean vignette for details
```

Once you have created a model with MetaClean it can be used with the predictCheckFeaturesSession() function:

```
predictCheckFeaturesSession(fGroups, "model_session.yml", model)
```

This will generate another *check session file*: all the feature groups that are considered good will be with a 'keep' state, the others without. As described elsewhere, the checkFeatures function is used to review the results from a session and the filter function can be used to remove unwanted feature groups. Note that calculatePeakQualitites() must be called before getMCTrainData/predictCheckFeaturesSession can be used.

NOTE MetaClean only predicts at the feature group level. Thus, only the kept feature groups from a *feature check session* will be used for training, and any indivual features that were marked as removed will be ignored.

8.4 Exporting and converting feature data

The feature group data obtained during the workflow can be exported to various formats with the export() generic function. There are currently three formats supported: "brukerpa" (Bruker ProfileAnalysis), "brukertasq" (Bruker TASQ) and "mzmine" (mzMine). The former exports a 'bucket table' which can be loaded in ProfileAnalysis, the second and third export a target list that can be processed with TASQ and mzMine, respectively.

The getXCMSSet() function converts a features or featureGroups object to an xcmsSet object which can be used for further processing with xcms. Similarly, the getXCMSnExp() function can be used for conversion to an XCMS3 style XCMSnExp object, and the getPICSet() function can be used to convert features to KPIC2 data.

Some examples for these functions are shown below.

8.5 Algorithm consensus

With patRoon you have the option to choose between several algorithms for most workflow steps. Each algorithm is typically characterized by its efficiency, robustness, and may be optimized towards certain data properties. Comparing their output is therefore advantegeous in order to design an optimum workflow. The consensus() generic function will compare different results from different algorithms and returns a consensus, which may be based on minimal overlap, uniqueness or simply a combination of all results from involved objects. The output from the consensus() function is of similar type as the input types and is

therefore compatible to any 'regular' further data processing operations (e.g. input for other workflow steps or plotting). Note that a consensus can also be made from objects generated by the same algorithm, for instance, to compare or combine results obtained with different parameters (e.g. different databases used for compound annotation).

The consensus() generic is defined for most workflow objects. Some of its common function arguments are listed below.

Argument Classes	Remarks
obj, All	Two or more objects (of the same type) that should be compared to generate the consensus.
compThreshoddmpounds, formulas,	The minimum overlap (relative/absolute) for a
relAbundanceeatureGroupsComparison	result (feature, candidate) to be kept.
absAbundance,	
formThreshold	
uniqueFrom compounds, formulas,	Only keep <i>unique</i> results from specified objects.
featureGroupsComparison	
uniqueOutercompounds, formulas,	Should be combined with uniqueFrom. If TRUE
featureGroupsComparison	then only results are kept which are <i>also</i> unique between the objects specified with uniqueFrom.

Note that current support for generating a consensus between components objects is very simplistic; here results are not compared, but the consensus simply consists a combination of all the components from each object.

Generating a consensus for feature groups involves first generating a featureGroupsComparison object. This step is necessary since (small) deviations between retention times and/or mass values reported by different feature finding/grouping algorithms complicates a direct comparison. The comparison objects are made by the comparison() function, and its results can be visualized by the plotting functions discussed in the previous chapter.

Some examples are shown below

8.6 Compound clustering

When large databases such as PubChem or ChemSpider are used for compound annotation, it is common to find *many* candidate structures for even a single feature. While choosing the right scoring settings can significantly improve their ranking, it is still very much possible that many candidates of potential interest remain. In this situation it might help to perform *compound clustering*. During this process, all candidates

for a feature are clustered hierarchically on basis of similar chemical structure. From the resulting cluster the maximum common substructure (MCS) can be derived, which represents the largest possible substructure that 'fit' in all candidates. By visual inspection of the MCS it may be possible to identify likely common structural properties of a feature.

In order to perform compound clustering the makeHCluster() generic function should be used. This function heavily relies on chemical fingerprinting functionality provided by rcdk.

```
compounds <- generateCompounds(...) # get our compounds
compsClust <- makeHCluster(compounds)</pre>
```

This function accepts several arguments to fine tune the clustering process:

- method: the clustering method (e.g. "complete" (default), "ward.D2"), see ?hclust for options
- fpType: finger printing type ("extended" by default), see ?get.fingerprint
- fpSimMethod: similarity method for generating the distance method ("tanimoto" by default), see ?fp.sim.matrix

For all arguments see the reference manual (?makeHClust).

The resulting object is of type compoundsCluster. Several methods are defined to modify and inspect these results:

```
# plot MCS of first cluster from candidates of M109_R116_61
plotStructure(compsClust, groupName = "M109_R116_61", 1)

# plot dendrogram
plot(compsClust, groupName = "M109_R116_61")

# re-assign clusters for a feature group
compsClust <- treeCut(compsClust, k = 5, groupName = "M109_R116_61")

# ditto, but automatic cluster determination
compsClust <- treeCutDynamic(compsClust, minModuleSize = 3, groupName = "M109_R116_61")</pre>
```

For a complete overview see the reference manual (?compoundsCluster).

8.7 Basic quantitative and regression analysis

While patRoon is currently primarily focused on qualitative analyses, some basic quantitative analysis can also be performed, for instance, to estimate concentrations of features. In fact, other types of data that may be useful for regression analysis can be set such as sample dilution factor or sampling time. The latter may, for instance, be used to isolate features with increasing or decreasing intensity. Regardless of what kind of regression analysis is performed, here we simply refer the values to be calculated as concentrations. In order to use this functionality, an extra column (conc) should be added to the analysis information, for instance:

For analyses with known concentrations (e.g. standards) the conc column should be set; for all others the value should be set to NA.

The as.data.table() function (or as.data.frame()) can then be used to calculate regression data and estimate concentrations:

```
# use areas for quantitation and make sure that feature data is reported
# (otherwise no concentrations are calculated)
# (only relevant columns are shown for clarity)
as.data.table(fGroups, areas = TRUE, features = TRUE, regression = TRUE)
```

```
#>
                group conc
                                  RSQ intercept slope
                                                         conc_reg
     1: M109_R192_20
#>
                        1 0.71446367 193338.67
                                                -4928
                                                        1.3649892
#>
     2: M109 R192 20
                         2 0.71446367 193338.67
                                                -4928 1.2700216
#>
        M109 R192 20
                         3 0.71446367 193338.67
                                                -4928 3.3649892
#>
        M111_R330_23
                        1 0.08902714 85338.67
                                                  -370 -0.8468468
        M111_R330_23
                         2 0.08902714 85338.67
                                                  -370 5.6936937
#>
#>
#> 422: M407_R239_672
                         2 0.99560719 210036.00 -11734 2.0767002
#> 423: M407_R239_672
                         3 0.99560719 210036.00 -11734
                                                       2.9616499
#> 424: M425_R319_676
                        1 0.46488086 193198.67
                                                 10896
                                                        1.6194322
#> 425: M425_R319_676
                         2 0.46488086 193198.67
                                                 10896
                                                       0.7611356
#> 426: M425_R319_676
                         3 0.46488086 193198.67
                                                10896
                                                       3.6194322
```

Calculated concentrations are stored in the conc_reg column, alongside while other regression data (i.e. RSQ, intercept, slope columns). To perform basic trend analysis the RSQ (i.e. R squared) can be used:

```
fGroupsTab <- as.data.table(fGroups, areas = TRUE, features = FALSE, regression = TRUE)
# subset fGroups with reasonable correlation
increasingFGroups <- fGroups[, fGroupsTab[RSQ >= 0.8, group]]
```

8.8 Fold changes

A specific statistical way to prioritize feature data is by Fold changes (FC). This is a relative simple method to quickly identify (significant) changes between two sample groups. A typical use case is to compare the feature intensities before and after an experiment.

To perform FC calculations we first need to specify its parameters. This is best achieved with the getFCParams() function:

```
getFCParams(c("before", "after"))
```

```
#> $rGroups
#> [1] "before" "after"
#>
#> $thresholdFC
#> [1] 0.25
#>
#> $thresholdPV
#> [1] 0.05
#>
#> $zeroMethod
```

```
#> [1] "add"
#>
#> $zeroValue
#> [1] 0.01
#>
#> $PVTestFunc
#> function (x, y)
#> t.test(x, y, paired = TRUE)$p.value
#> <bytecode: 0x55dbe71a2710>
#> <environment: 0x55dbd5466898>
#>
#> $PVAdjFunc
#> function (pv)
#> p.adjust(pv, "BH")
#> <bytecode: 0x55dbe71a2a20>
#> <environment: 0x55dbd5466898>
```

In this example we generate a list with parameters in order to make a comparison between two replicate groups: before and after. Several advanced parameters are available to tweak the calculation process. These are explained in the reference manual (?featureGroups).

The as.data.table function for feature groups is used to perform the FC calculations.

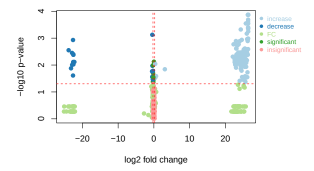
```
#>
                group
                                FC
                                        FC_log
                                                              PV_log classification
#>
           M99_R14_1 8.837494e-01 -0.17829070 0.22523941 0.64735561 insignificant
     1:
            M99 R4 2 8.500464e-01 -0.23438649 0.77848844 0.10874783 insignificant
#>
     2:
           M100 R7 3 8.009186e-01 -0.32027248 0.80475149 0.09433821
                                                                                 FC
#>
     3:
#>
           M100_R5_4 4.140000e+06 21.98119934 0.53420412 0.27229277
                                                                                 FC
     4:
#>
     5:
           M100_R28_5 9.594972e-01 -0.05964952 0.97571237 0.01067819
                                                                      insignificant
#> 676: M425_R319_676 2.149907e+07 24.35777069 0.00983421 2.00726053
                                                                           increase
#> 677: M427_R10_677 1.059937e+00 0.08397893 0.37317466 0.42808786
                                                                      insignificant
#> 678: M427_R319_678 7.776800e+06 22.89074521 0.53420412 0.27229277
                                                                                 FC
#> 679: M432_R383_679 9.816400e+06 23.22676261 0.34750809 0.45903508
                                                                                 FC
#> 680: M433_R10_680 1.132909e+00 0.18003240 0.29525423 0.52980387 insignificant
```

The classification column allows you to easily identify if and how a feature changes between the two sample groups. This can also be used to prioritize feature groups:

```
tab <- as.data.table(fGroups, FCParams = myFCParams)
# only keep feature groups that significantly increase or decrease
fGroupsChanged <- fGroups[, tab[classification %in% c("increase", "decrease")]$group]</pre>
```

The plotVolcano function can be used to visually the FC data:

```
plotVolcano(fGroups, myFCParams)
```



8.9 Caching

In patRoon lengthy processing operations such as finding features and generating annotation data is *cached*. This means that when you run such a calculation again (without changing any parameters), the data is simply loaded from the cache data instead of re-generating it. This in turn is very useful, for instance, if you have closed your R session and want to continue with data processing at a later stage.

The cache data is stored in a sqlite database file. This file is stored by default under the name cache.sqlite in the current working directory (for this reason it is very important to always restore your working directory!). However, the name and location can be changed by setting a global package option:

```
options(patRoon.cache.fileName = "~/myCacheFile.sqlite")
```

For instance, this might be useful if you want to use a shared cache file between projects.

After a while you may see that your cache file can get quite large. This is especially true when testing different parameters to optimize your workflow. Furthermore, you may want to clear the cache after you have updated patRoon and want to make sure that the latest code is used to generate the data. At any point you can simply remove the cache file. A more fine tuned approach which doesn't wipe all your cached data is by using the clearCache() function. With this function you can selectively remove parts of the cache file. The function has two arguments: what, which specifies what should be removed, and file which specifies the path to the cache file. The latter only needs to be specified if you want to manage a different cache file.

In order to figure what is in the cache you can run clearCache() without any arguments:

clearCache()

```
#> Please specify which cache you want to remove. Available are:
#> - EICData (3 rows)
#> - MSPeakListsAvg (4 rows)
#> - MSPeakListsMzR (97 rows)
#> - MSPeakListsSetAvg (2 rows)
#> - annPeakListSet (1 rows)
#> - annotateSuspects (1 rows)
#> - calculatePeakQualities (3 rows)
#> - componentsCAMERA (1 rows)
#> - componentsNontarget (1 rows)
#> - compoundsCluster (1 rows)
#> - compoundsMetFrag (30 rows)
```

```
#> - featAnnUnset (2 rows)
#> - featureGroupsOpenMS (5 rows)
#> - featuresOpenMS (69 rows)
#> - filterFGroups_blank (3 rows)
#> - filterFGroups_intensity (9 rows)
#> - filterFGroups minReplicates (85 rows)
#> - filterFGroups replicateAbundance (6 rows)
#> - filterFGroups_replicate_group (18 rows)
#> - filterFGroups retention (2 rows)
#> - filterMSPeakLists (4 rows)
#> - formulasFGroupConsensus (2 rows)
#> - formulasGenForm (89 rows)
#> - formulasSIRIUS (5 rows)
#> - loadIntensities (69 rows)
#> - mzREIC (3403 rows)
#> - reportPlots (334 rows)
#> - screenSuspects (4 rows)
#> - screenSuspectsPrepList (3 rows)
#> - specData (12 rows)
#> - all (removes complete cache database)
```

Using this output you can re-run the function again, for instance:

```
clearCache("featuresOpenMS")
clearCache(c("featureGroupsOpenMS", "formulasGenForm")) # clear multiple
clearCache("OpenMS") # clear all with OpenMS in name (ie partial matched)
clearCache("all") # same as simply removing the file
```

8.10 Parallelization

Some steps in the non-target screening workflow are inherently computationally intensive. To reduce computational times patRoon is able to perform *parallelization* for most of the important functionality. This is especially useful if you have a modern system with multiple CPU cores and sufficient RAM.

For various technical reasons several parallelization techniques are used, these can be categorized as parallelization of R functions and multiprocessing. The next sections describe both parallelization approaches in order to let you optimize the workflow.

8.10.1 Parellization of R functions

Several functions of patRoon support parallelization.

Function	Purpose	Remarks
findFeatures	Obtain feature data	Only envipick and kpic2 algorithms
generateComponents optimizeFeatureFinding, optimizeFeatureGrouping	Generate components Optimize feature finding/grouping parameters	Only cliquems algorithm. Discussed here.
calculatePeakQualities	Calculate feature (group) qualities	Discussed here.

The parallelization is achieved with the future and future.apply R packages. To enable parallelization of these functions the parallel argument must be set to TRUE and the future framework must be properly configured in advance. For example:

```
# setup three workers to run in parallel
future::plan("multisession", workers = 3)

# find features with enviPick in parallel
fList <- findFeatures(anaInfo, "envipick", parallel = TRUE)</pre>
```

It is important to properly configure the right future plan. Please see the documentation of the future package for more details.

8.10.2 Multiprocessing

patRoon relies on several external (command-line) tools to generate workflow data. These commands may be executed in *parallel* to reduce computational times ('multiprocessing'). The table below outlines the tools that are executed in parallel.

Tool	Used by	Notes
msConvert	convertMSFiles(algorithm="pwiz",)	
FileConverter	<pre>convertMSFiles(algorithm="openms",)</pre>	
${\tt Feature Finder Metabo}$	<pre>findFeatures(algorithm="openms",)</pre>	
julia	<pre>findFeatures(algorithm="safd",)</pre>	
SIRIUS	<pre>findFeatures(algorithm="sirius",)</pre>	
MetaboliteAdductDecl	h geger ateComponents(algorithm="openms",	
)	
GenForm	<pre>generateFormulas(agorithm="genform",)</pre>	
SIRIUS	<pre>generateFormulas(agorithm="sirius",),</pre>	Only if splitBatches=TRUE
	<pre>generateCompounds(agorithm="sirius",)</pre>	
MetFrag	<pre>generateCompounds(agorithm="metfrag",)</pre>	
pngquant	reportHTML()	Only if $optimizePng=TRUE$
BioTransformer	<pre>generateTPs(algorithm =</pre>	Disabled by default (see
	"biotransformer")	<pre>?generateTPs for details).</pre>

Multiprocessing is either performed by executing processes in the background with the processx R package (classic interface) or by futures, which were introduced in the previous section. An overview of the characteristics of both parallelization techniques is shown below.

classic	future
requires little or no configuration works with all tools only supports parallelization on the local computer	configuration needed to setup doesn't work with pngquant and slower with GenForm allows both local and cluster computing

Which method is used is controlled by the patRoon.MP.method package option. Note that reportHTML() will always use the classic method for pngquant.

8.10.2.1 Classic multiprocessing interface The classic interface is the 'original' method implemented in patRoon, and is therefore well tested and optimized. It is easier to setup, works well with all tools, and is therefore the default method. It is enabled as follows:

```
options(patRoon.MP.method = "classic")
```

The number of parallel processes is configured through the patRoon.MP.maxProcs option. By default it is set to the number of available CPU cores, which results usually in the best performance. However, you may want to lower this, for instance, to keep your computer more responsive while processing or limit the RAM used by the data processing workflow.

```
options(patRoon.MP.maxProcs = 2) # do not execute more than two tools in parallel.
```

This will change the parallelization for the complete workflow. However, it may be desirable to change this for only a part the workflow. This is easily achieved with the withOpt() function.

```
# do not execute more than two tools in parallel.
options(patRoon.MP.maxProcs = 2)

# ... but execute up to four GenForm processes
withOpt(MP.maxProcs = 4, {
   formulas <- generateFormulas(fGroups, "genform", ...)
})</pre>
```

The withOpt function will temporarily change the given option(s) while executing a given code block and restore it afterwards (it is very similar to the with_options() function from the withr R package). Furthermore, notice how withOpt() does not require you to prefix the option names with patRoon..

8.10.2.2 Multiprocessing with futures The primary goal of the "future" method is to allow parallel processing on one or more external computers. Since it uses the future R package, many approaches are supported, such as local parallelization (similar to the classic method), cluster computing via multiple networked computers and more advanced HPC approaches such as slurm via the future.batchtools R package. This parallelization method can be activated as follows:

```
options(patRoon.MP.method = "future")

# set a future plan

# example 1: start a local cluster with four nodes
future::plan("cluster", workers = 4)

# example 2: start a networked cluster with four nodes on PC with hostname "otherpc"
future::plan("cluster", workers = rep("otherpc", 4))
```

Please see the documentation of the respective packages (e.g. future and future.batchtools) for more details on how to configure the workers.

The withOpt() function introduced in the previous subsection can also be used to temporarily switch between parallelization approaches, for instance:

```
# default to future parallelization
options(patRoon.MP.method = "future")
future::plan("cluster", workers = 4)

# ... do workflow

# do classic parallelization for GenForm
withOpt(MP.method = "classic", {
    formulas <- generateFormulas(fGroups, "genform", ...)
})

# .. do more workflow</pre>
```

8.10.2.3 Logging Most tools that are executed in parallel will log their output to text files. These files may contain valuable information, for instance, when an error occurred. By default, the logfiles are stored in the log directory placed in the current working directory. However, you can change this location by setting the patRoon.MP.logPath option. If you set this option to FALSE then no logging occurs.

8.10.3 Notes when using parallelization with futures

Some important notes when using the future parallelization method:

- GenForm currently performs less optimal with future multiprocessing to the classic approach. Nevertheless, it may still be interesting to use the future method to move the computations to another system to free up resources on your local system.
- Behind the scenes the future.apply package is used to schedule the tools to be executed. The patRoon.MP.futureSched option sets the value for the future.scheduling argument to the future_lapply() function, and therefore allows you to tweak the scheduling.
- Make sure that patRoon and in case of multiprocessing that the tool to be executed (MetFrag, SIRIUS etc.) are exactly the *same* version on all computing hosts.
- Make sure that patRoon is properly configured on all hosts, e.g. set the patRoon.path.XXX options to ensure all tools can be found.
- For MetFrag annotation: if a local database such as PubChemLite is used, it must be present on each computing node as well. Furthermore, the local computer (even if not used for the computations) also must have this file present. Like the previous point, make sure that the patRoon.path.XXX options are set properly.
- If you encounter errors then it may be handy to switch to future::plan("sequential") and see if it works or you get more descriptive error messages.
- In order to restart the nodes, for instance after re-configuring patRoon, updating R packages etc, simply re-execute future::plan(...).
- Setting the future.debug package option to TRUE may give you more insight what is happening to find problems.

9 References

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