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Abstract

DAPAR (Differential Analysis of Protein Abundance with R) and ProStaR (Proteomics and Statistics with R) are two Bioconductor packages that contain the necessary functions to analyze proteomics data (DAPAR), as well as the corresponding graphical user interfaces (ProStaR). This document guides the practitioner through the use of DAPAR (R command lines) and ProStaR (click-button interface, so that no programming skill is required).

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1 Introduction

DAPAR and *ProStaR* are a series of software dedicated to the processing of proteomics data. More precisely, they are devoted to the analysis of quantitative datasets produced in bottom-up discovery proteomics with a LC-MS/MS pipe-line (Liquid Chromatography and Tandey Mass spectrometry). The experiment package *DAPARdata* is the companion package for *ProStaR* and *DAPAR*. It contains many datasets that can be used as example.

DAPAR (Differential Analysis of Protein Abundance with R) is an R package that contains all the necessary functions to:

- Import/export a quantitative dataset. Here, a quantitative dataset denotes a table where each protein is represented by a line and and each replicate is represented by a column; each cell of the table contains the abundance of a given protein in a given sample; the replicates are clustered into different conditions (or groups), and the purpose of the analysis is to isolate the few proteins the abundance of which significantly differ between the conditions (or groups).
- Compute and display meaningful statistics regarding the quantitative dataset.
- Perform the various processing steps of a complete quantitative analysis:

 (i) filtering and data cleaning;
 (ii) cross-replicate normalization;
 (iii) missing value imputation;
 (iv) aggregation of peptide intensities into protein intensities;
 (v) statistical tests and false discovery rate computation;
 (vi) Gene Ontology (GO) analysis (grouping and category enrichment).

This package can be used on its own; or as a complement to the numerous Bioconductor packages (https://www.bioconductor.org/) it is compliant with; or through the *ProStaR* interface. *ProStaR* (Proteomics and Statistics with R) is a web-interface based on Shiny (http://shiny.rstudio.com/) that provides Graphical User Interfaces (GUI) to all the *DAPAR* functionalities, so as to guide any practitioner that is not comfortable with R programming through the complete quantitative analysis process.

In *DAPAR*, a serie of functions may be called from two types of variables: a dataframe that contains quantitative data and an object of class MSnSet. The first case has been developped to make *DAPAR* easier to use (in command line) for users who do not want to work with MSnSet files and to be compliant in the future with the Proline Suite (http://proline.profiproteomics.fr/).

2 Installation

There are 3 ways to use DAPAR:

- The first one is to use *DAPAR* alone, through command lines or scripts. To do so, the user simply has to install *DAPAR* on his/her own workstation, as instructed in Section 2.2;
- The second one is to use DAPAR along with its graphical interface ProStaR, and to have them running on the user's station (referred to as stand-alone install). In such case, it is necessary to install DAPAR first, as instructed in Section 2.2, and ProStaR then, as instructed in Section 2.1.1;
- In the case where several *ProStaR* users who are not confortable with R (programming or installing), it is best to have a single version of *DAPAR* and *ProStaR* running on a shiny server installed on a Unix/Linux server. The users will use *ProStaR* through a web browser, exactly if it were locally installed, yet, a single install has to be administrated. In that case, *DAPAR* has to be classically installed (Section 2.2), while on the other hand, the installation of *ProStaR* is slightly different on a server (Section 2.1.2).

For a stand-alone use, both *DAPAR* and *ProStaR* can run on any operating system (Unix/Linux, Mac OS X and Windows) as long as R is installed. In any case (stand-alone or server), a recent version of R (\geq 3.4) is needed.

2.1 ProStaR (with DAPAR)

ProStaR can be run in two differents ways: standalone or server. The prerequested packages described above have to be installed on the server if the user run a shiny-server to distribute *ProStaR* or on a local machine if *ProStaR* is run locally.

2.1.1 Stand-alone version

To run the stand-alone version, it is necessary to install the package in a directory where the user have read/write permissions. If the user have administrator privileges, then in a R console, enter:

```
> source("http://www.bioconductor.org/biocLite.R")
```

```
> biocLite("Prostar")
```

This step will automatically install the following packages: *shiny*, *rhandsontable*, *DAPARdata*, *DAPAR*, *data.table*, *DT*, *shinyjs*, *openxlsx*, *shinyAce*, *highcharter*, *rhandsontable*, *htmlwidgets*, *webshot*, *R.utils*.

Once the package is installed, to launch *ProStaR*, then enter:

```
> library(Prostar)
```

> Prostar()

A new window of the default web browser opens.

2.1.2 Server version

This version uses a Shiny Server (https://github.com/rstudio/shiny-server). It is a server program that makes Shiny applications available over the web. Please follow installation instructions if you do not have a server yet.

Once a Shiny server is available in your network, the first step is to install *Prostar* as described in Section 2.1.1 in order to have the dependencies installed.

Then, execute the following line in order to get the install directory of Prostar:

> installed.packages()["Prostar","LibPath"]

The result of this command is now referred as */path_to_Prostar*.

Change the owner of the shiny-server directory and log as shiny # sudo chown shiny /srv/shiny-server # sudo su shiny

Create a directory named *ProStaR* in the Shiny Server directory with the user shiny as owner and then copy the Prostar files.

Create the directory for the shiny application # mkdir /srv/shiny-server/Prostar Copy the ProstarApp directory within the shiny-server directory # sudo cp -r /path_to_Prostar/ProstarApp/ /srv/shiny-server/Prostar Change the owner of the shiny-server directory # sudo chown -R shiny /srv/shiny-server Give the following permissions to the www directory # chmod 755 /srv/shiny-server/Prostar/www Check if the configuration file of shiny-server is correct. For more details, please visit http://rstudio.github.io/shiny-server/latest/.

Now, the application should be available via a web browser at http://servername:port/Prostar.

2.2 DAPAR (alone)

To install the package DAPAR from the source file with administrator rights, start R and enter:

```
> source("http://www.bioconductor.org/biocLite.R")
```

```
> biocLite("DAPAR")
```

This step will automatically install the following packages:

- From CRAN: RColorBrewer, Cairo, png, lattice, reshape2, tmvtnorm, norm, ggplot2, imputeLCMD, gplots, openxlsx, knitr, cp4p, doParallel, foreach, scales, stats, grDevices, vioplot, sm, graphics, utils, parallel, openxlsx, Matrix, highcharter, dplyr, tidyr, imp4p, readxl, Ime4,graph
- From Bioconductor: *MSnbase*, *DAPARdata*, *preprocessCore*, *impute*, *limma*, *pcaMethods*, *clusterProfiler*, *AnnotationDbi*, *siggenes*

2.3 DAPARdata

The installation of the package *DAPARdata* follows the classical way for Bioconductor packages. In a R console, enter:

```
> source("http://www.bioconductor.org/biocLite.R")
```

```
> biocLite("DAPARdata")
```

3 Navigating through the ProStaR interface

3.1 Overview of the interface

As illustrated on Fig. 1, *ProStaR* proposes a classical Graphical User Interface (GUI) to visualize and interact with the data. On the top, a navbar menu helps navigating through the various *DAPAR* functionalities and running them. It is divided into five submenus:

Prostar	Dataset manager +	Descriptive statistics	Data processing +	Help	Dataset versions
					None •
S. Wieczorek 'DAPAR & Pr Bioinformatic	R and ProStaR softwar , F. Combes, C. Lazar, Q. oStaR: software to perfor s 33(1), 135–136, 2017 /10.1093/bioinformatics/b	Gial-Gianetto, L. Gatto, A m statistical analyses in q			и. Ferro, C. Bruley and T. Burger.
Nore specific t is compose • Prostar	ProStaR form a software ally it is designed to proc d of two distinct R packa (version 1.8.5); the web i (version 1.9.1); a collect	ess relative quantitative d ges : based graphical user inter	ata from discovery exp face to DAPAR	riments.	proteomics.
DAPAR includ Here is a brie • Descri • Filterin • Cross • Missin • Aggre • Differe	des wrappers to numerou f overview of the available ptive statistics are availa g options allows pruning replicate normalization, g values imputation with pation from peptide to pr	s other R packages, eithe functionalities: ible, for exploration and v the protein or peptide list so as to make the quanti different methods, deper- tein intensity values; udes null hypothesis sign	r available on CRAN or isualization of the quan according to various c tative values comparab nding on the nature of t ificance testing as well	itative data itative data iteria (miss e between ne missing as multiple	aset; sing values, contaminants, reverse sequences); n the different analyzed samples; y values; b testing correction (for false discovery rate estimatif
For more deta	alls, please refer to the "H	elp" tab.			

Figure 1: Default screen of ProStaR

- **ProStar**: The welcome page, depicted on Fig. 1.
- Dataset manager: It contains the tools to import and export datasets;
- Descriptive statistics: It provides different visualization tools that are helpful to understand the dataset, and to picture the influence of the various processing;
- Data processing: This is the heart of ProStaR, where all the DAPAR functionalities can be accessed to;
- **Help**: A serie of informations about the software, associated references, etc.

On the right hand side of the navbar menu, a dropdown menu referred to as "Dataset versions" makes it possible to navigate back through the history of the processing. Its use is detailed in Section 3.6

3.2 Dataset manager

The "Dataset manager" allows the user to open, import or export quantitative datasets. *ProStaR* and *DAPAR* use the MSnSet format which is part of the package *MSnbase*: It is either possible to load existing MSnSet files (see Section 3.2.1) or to import text (-tabulated) and Excel files (see Section 3.2.2). The "Demo mode" allows to load the datasets of the package *DAPARdata* (see Section 3.2.5).

3.2.1 Open MSnSet

The user can upload a dataset that is already formated as an MSnSet file, by clicking on "Open MSnSet File" (see Fig. 2). This action opens a popup window, so as to let the user choose the appropriate file. Once the file is uploaded, a short summary of the dataset is shown, which includes the number of samples, the number of proteins in the dataset, the percentage of missing values and the number of lines which only contain missing values.

Once done, the menu of "Dataset versions" is updated to "Original - peptide"

Prostar	Dataset manager -	<u>III</u> De	scriptive statistics	Data processing -	Help	Dataset versions of Exp1_R2_pept
						Original - peptide 👻
Open a MS	Snset file		Quick over	view of the d	ataset	
Browse	Exp1_R2_pept.MSn Upload complete	Iset	There are 14 Percentage	samples in your data. .048 peptides in your da of missing values: 18.82 nes with only NA values	2 %	
			As your dataset co to the differential a		, you should	I impute them prior to proceed
				ontains lines with no va ed to the analysis of th		hould remove them with the filter

Figure 2: Open a MSnSet file

or "Original - protein" whether the file contains quantitative information about peptides or proteins (see Section 3.6). All the plots in the "Descriptive statistics" submenu (see Section 3.3) become accessible and all the widgets to interact with *ProStaR* are preloaded.

Command line: It is possible to open an MSnSet dataset directly in command line (*i.e.* without *ProStaR* interface), using function readRDS().

The user can find examples of MSnSet file in the 'extdata' directory of *DA*-*PARdata*. The user can find this directory by entering:

> installed.packages()["DAPARdata","LibPath"]

3.2.2 Import

Alternatively, the user can create a quantitative dataset in the MSnSet format, on the basis of TSV (Tab Separated Values) or Excel files (in format xls or xlsx) that contain the results of a proteomics analysis. To do so, one has to click on

"Convert data to MSnSet". Then, the right panel splits into 5 tabs that guide the user through the procedure to create the MSnSet object. Let us describe the import format first, and the import procedure then.

Import data

Data are imported through a text file (.txt) formatted as a column separated file, with tabulations (*i.e.* "\t" character) as column separator. The first line of the text file contains the column names. A minimum of 4 columns with quantitative values are necessary: As ProStaR is made for label-free discovery proteomics, it is required to have a minima 2 conditions (or groups, or labels) to compare; moreover, 2 replicates per conditions are necessary, so that it is possible to compute a condition-wise variance. Regarding the quantitative values, the decimal separator is ".", and the intensities may be either log-transformed or not. It is also advised to have an additional columns that contains IDs (that is that gives a unique name to each line of the dataset). If such a unique ID does not appear in the dataset, one will be automatically generated.

Of course, it is possible to have more columns with quantitative values; as well as additional columns with other information. The latter ones will be considered as metadata when imported.

It is recommended to avoid special characters such as " \sharp ", "@", "\$", "%", etc. that are automatically removed. Similarly spaces in column names are replaced by ".".

If the dataset describes proteins, each line should correspond to one and only one protein. If the dataset describes peptides, each line should correspond to one and only one peptide. In addition, it is necessary to have a column that lists (separated by a ";") all the parent proteins of each peptide, as this piece of information is mandatory for the aggregation. It is also necessary that these parent proteins are describe by a unique ID (so as to avoid confusion between the proteins).

As an example, please find at the following address a dataset abstract that can be inspired from: http://www.bioconductor.org/packages/release/bioc/readmes/Prostar/README.

Import procedure

Several panels guide the user through the different steps of the procedure:

Select file: Select the CSV/txt or Excel (*.xls, *.xlsx) file to import (see Fig. 3). The file must contain a table where each line corresponds to a peptide or protein, except the first one which must contain the names of the colums. If the user chooses an Excel file, a dropdown menu appears and ask the user to select the spreadsheet containing the data.

As it appears in Fig. 3, some options allows for specifying if data are related to peptides or proteins, if the abundance values are already log-transformed or not ¹, and also if 0 and NaN values should be replaced by **NA**.

v to create a MSn	Sot file from a tabulated t			none 🔻
	Set file from a tabulateu-t	ext file.		
2 - Data Id	3 - Exp. and feat. data	4 - Samples metadata	5 - Convert	
sv, .tsv, .xis, .xisx	files)			
xp1_R2_pept.txt				
Upload complete				
protein dataget	2			
r protein dataset	?			
et	?			
et et				
et et ready log-transfo				
et et				
et et ready log-transfo				
et et ready log-transfo	ormed ?			
et et ready log-transfo unchanged)	ormed ?			
		sv, .tsv, .xls, .xlsx files) xp1_R2_pept.txt	sv, .tsv, .xls, .xlsx files) xp1_R2_pept.txt	sv, .tsv, .xls, .xlsx files) xp1_R2_pept.txt

¹In *DAPAR*, the analysis is always conducted on logtransformed data. They may have previously been transformed, but if not, then DAPAR automatically performs the transformation. The user should not try applying any DAPAR processing on data that are not log-transformed, for the result would be dubious.

Figure 3: Importing an Excel file, tab 1

Data ID: This step is to set the column that corresponds to the unique ID of peptides or proteins. The user has two options: let *ProStaR* creates such an ID by itself or choose among the columns available in the data file. If choosing the second option, then a drop-down menu appears and provides the list of the column names. A column corresponding to the unique ID of the peptides or proteins should be selected (see Fig. 4). If the column contains non unique IDs, a warning alerts the user and suggests him to choose another column.

These steps allow to create a MSnSet file from a tabulated-text file. 1 - Select file 2 - Data Id 3 - Exp. and feat. data 4 - Samples metadata 5 - Convert Please select among the columns ofyour data the one that corresponds to a unique ID of the peptides . If you choose the automatic ID, Prostar will build an index.	Prostar	Dataset manager -	Lel Descriptive statistics	Data processing -	Help	Dataset versions
Please select among the columns ofyour data the one that corresponds to a unique ID of the peptides . If you choose the automatic ID, Prostar will build an index. Auto ID	These steps allow	to create a MSnSet file f	rom a tabulated-text file.			
If you choose the automatic ID, Prostar will build an index. Auto ID	1 - Select file	2 - Data Id 3 - Ex	p. and feat. data 4 - San	nples metadata 5 -	Convert	
Auto ID	Please select amor	ng the columns ofyour d	ata the one that corresponds	to a unique ID of the pe	ptides .	
	-	automatic ID, Prostar	will build an index.			
S User ID						
	 User ID 					
	id		•			
id 👻						

Figure 4: Importing a CSV file, tab 2

Exp. and Feat. data: In the "Quantitative data" list, select (one after the other) the columns that correspond to the quantitative data. Each time the user selects an item in the list, it is moved up to the field above (see Fig. 5).

If an item is selected by mistake, it can be removed by pressing on the SUPPR key.

Please note that the decimal separator should be ".".



Figure 5: Importing a CSV file, tab 3

Sample metadata: In this tab, the user fills the informations related to the samples, *i.e.* the experimental design. The colum named *Experiment* is filled by default with the name of the different samples. The user fills the other columns: *Label* corresponds to the conditions of the experiment that will be compared during the differential analysis; *Bio.Rep, Tech.rep* and *Analyt.Rep.* correspond respectively to the biological, technical and analytical replicates (Fig. 6). The column Label is mandatory (for the subsequent differential analysis), the other ones are optional.

For the case of a peptide dataset, and in order to be able to agregate the peptides intensities on proteins ones : there should be a column indicating, for each peptide, the protein/s the peptide belongs to. If a peptide belongs to more than one protein, proteins names should be separated by ";".

	Prostar Da	taset manager	- Ind Des	criptive statistics	Data process	ing - Help	Dataset versions
se	steps allow to cre	ate a MSnSet fi	ile from a tabul	ated-text file.			
- 8	Select file 2 -	Data Id 3 -	- Exp. and feat	data 4 - Sa	mples metadata	5 - Convert	
	Exper	ment	Label	Bio.Rep	Tech.Rep	Analyt.Rep	
1	Intensity.D.R1	ment	D	1	Tech.Rep	Analyt.Rep	
1 2 3		ment		Bio.Rep 1 2 3	Tech.Rep	Analyt.Rep	
	Intensity.D.R1 Intensity.D.R2	ment	D	1 2	Tech.Rep	Analyt.Rep	
3	Intensity.D.R1 Intensity.D.R2 Intensity.D.R3	ment	D D D	1 2	Tech.Rep	Analyt.Rep	

Figure 6: Importing a CSV file, tab 4

Convert: Finally, enter the name of the MSnSet to be created (Fig. 7) and click on "Convert data". The data are converted and automatically loaded in *ProStaR*. The name of the file appears in the upper right hand side of the screen, as a title to the drop-down menu "Dataset version of <dataset_name>".

Prostar	Dataset manager 👻	In Descriptive statistics	Data processing +	Help	Dataset versions of UPS_test
These steps allow t	o create a MSnSet file fr 2 - Data Id 3 - Exi		les metadata 5 - Ci	onvert	Original - peptide 🔻
Enter the name of		, and reat. data 4 - Samp	ies metauata 5 - G	Unvert	
Convert data The conversion vizualize your o		aset has been automati	ically loaded in me	emory. No	ow, you can switch to the Descriptive statistics panel to

Figure 7: Importing a CSV file, tab 5

Command line: In *DAPAR*, the function to create an MSnSet from a CSV file is createMSnSet().

3.2.3 Export dataset to a file

Once an MSnSet has been created, it is possible to save it as a MSnSet binary object (so that next time, it is not necessary to create it, and a simple uploads makes it, as described in Section 3.2.1). It is also possible to export it as an Excel spreadsheet (in xlsx format). To do so, one simply goes on the corresponding tab and select the appropriate option (Fig. 8).

					Exp1_R2_pept Original - peptide -
Export to file	Generate report (E	Beta)			Original - peptide +
hoose the exp	ort format of the datas	et and choose a name.			
ile format					
		•			
MSnset					
MSnset					
MSnset					
	nns you want to keep a		f any column is specifie	d, all meta	idata in your dataset will be exporte
	nns you want to keep a		f any column is specifie	d, all meta	idata in your dataset will be exporte
	nns you want to keep a		f any column is specifie	d, all meta	idata in your dataset will be exporte
	nns you want to keep a		f any column is specifie	d, all meta	data in your dataset will be exporte
	nns you want to keep a		f any column is specifie	d, all meta	idata in your dataset will be exporte
	nns you want to keep a		f any column is specifie	d, all meta	idata in your dataset will be exporte
elect the colun		as metadata. By default, i	f any column is specifie	d, all meta	idata in your dataset will be exporte
elect the colun	nns you want to keep a	as metadata. By default, i	f any column is specifie	d, all meta	udata in your dataset will be exporte
elect the colun	of the files to be crea	as metadata. By default, i	f any column is specifie	d, all meta	udata in your dataset will be exporte

Figure 8: Exporting to an Excel file

Command line: When working exclusively with *DAPAR*, the functions are writeMSnSetToExcel() (to export in Excel format) and saveRDS() (to export in MSnSet format).

The user can download the plots showed in *Prostar* by right-clicking on the plot. A contextual menu appears and let the user choose either "Save image as" or "Copy image". In the latter case, he/she has to paste the image in appropriate software.

3.2.4 Export plots as report (Beta)

In addition to the exported datasets (see previous section), the user can generate and download a report, dynamically created on demand (e.g. at the end of an analysis), through the interface presented on Fig. 9.

The left hand side of the interface shows a list of checkboxes (one for each dataset created during the analysis). By default, all the datasets are selected. The user chooses which results to include in the report. Even if no analysis has been performed (e.g. right after the upload of a dataset), the plots of the Descriptive Statistics panel are available.

The right hand side of the interface allows the user to choose the size and resolution of the images and the format of the report: PDF, HTML or DOC file.

After selecting the desired options, one clicks on the button "Generate a report". Then, ProStaR rebuilds all the required images and includes them in the report file. During this step, the button "Download" remains disabled. Only when the report is ready, the "Download" buttons is enabled.

Prostar Dataset manager - Descr	iptive statistics Data processing - Help	Exp1_R	versions of 2_pept
Export to file Generate report (Beta)		GOAna	alysis - protei n
hoose the datasets to export	Choose the size of images (PNC)	Choose the report document format
Original - peptide	1200 * 800	-	PDF O HTML O Word
Filtered - peptide Normalized - peptide	Choose the resolution		Enter the name of the report
Imputed - peptide	150	-	
Aggregated - protein			
DiffAnalysis.Limma - protein GOAnalysis - protein			
GOAnalysis - protein			

Figure 9: Exporting analysis as a report

Please note this report generation functionality is still in Beta version, so that some bugs may remain. In the future, and depending on the users' expectations, the report will be completed:

 So far, the R commands run for the report generation does not appear in the log console (see 3.2.6);

- The texts accompanying the figures is minimal;
- The report cannot be customized.

In the future, we hope to improve it, as well as to generate additional ready-touse files for publication (Material & Methods sketch, R script for reproducibility, etc.)

3.2.5 Demo mode

In order to facilitate first steps with Prostar, the "demo mode" menu allows the user to access the datasets contained in the package *DAPARdata*. When the user chooses one of those datasets, it is automatically loaded in *Prostar*. It can be used to easily test the various functionalities of *Prostar* (Fig. 10).

Prostar Data	iset manager 👻	Descriptive statistics Data p	rocessing - Help Dataset versions of Exp1_R2_pept
			Original - peptide 👻
Choose a demo o	dataset	Quick overvie	w of the dataset
Exp1_R2_pept		There are 6 samp	
Show PDF doc		 Percentage of mis 	septides in your data. sing values: 18.82 % vith only NA values = 715
Load demo data	aset	Exp1_R2_pept.pdf	1/3 Č 🛓
			of 'Exp1_R2_pept.Rd'
			October 6, 2016
			Exp1_R2_pept Exp1_R2_pept dataset
			Description
			This dataset is the final outcome of a quantitative mass spectrometry-based protomic analysis of two samples containing different concentrations of 48 human proteim (RPS1 standard from Sigma-Aldrich) within a constant year background (see Giai Gianetto et al. (2016) for details). It contains the abundance values of the different human and yeast peptidise identified and quantified in these two conditions. The two conditions represent the measured abundances of peptides when respectively 5 finol and 10 from 0 (UPS1 human proteins were maked with the yeast extra telefore mass spectrometry analyses. This results in a concentration ratio of 2. Three technical replicates were acquired for each condition.
			The dataset is either available as a CSV file (see inst/extdatafExpl_R2_pept.txt), or as a MSnSet structure (Expl_R2_pept). In the latter case, the quantitative data are those of the raw intensities.
			Usage
			<pre>data(Exp1_R2_pept)</pre>
			Format
			An object of class MSnSet related to peptide quantification. It contains 6 samples divided into two conditions (10fmol and 5fmol) and 14048 peptides.
			The data frame exprs(Exp1_R2_pept) contains six columns that are the quantitation of peptides for the six replicates.
			The data frame fData(Exp1_R2_pept) contains the meta data about the peptides.
			The data frame pData(Exp1_R2_pept) contains the experimental design and gives few informations about the samples.

Figure 10: Loadind a demo dataset

A checkbox allows the user to show the PDF file in the interface; by default, it is unchecked.

3.2.6 Session log

Each time the user validates a processing step (by clicking on the "Save < *the_step*>" button, see Section 3.4), the entire related information (such as the method name and its parameters) is added to the table shown in the "Session log" tab (see Figure 11). Hence, this table is a history of how the data were processed during the session. Let us note that, if a dataset is processed, then saved and reloaded in a new session, the session log is naturally empty. To have a complete view on the previous processing applied to a given dataset, please refer to Section 3.3.2).

Moreover, in the "R source code" tab, the user has access to the R commands (from DAPAR) that have been used to process its dataset (see Figure 12). This code may be copy and paste in a R script to automate the analysis of a dataset.

3.3 Descriptive statistics

Several plots (one plot per tab) are proposed to help the user to have a quick and as complete as possible overview of his/her dataset. This menu is an essential element for the user to check that each processing step indeed gave the expected result.

3.3.1 Missing value summary

The barplot on the left represents the number of missing values in each sample. The different colors correspond to the different conditions (or label). The histogram in the middle displays the distribution of missing values; the red bin counts the peptide or protein lines that only contains missing values (Fig. 13). The barplot on the right shows the distribution of missing values per condition.

Pr	ostar I	Dataset manager 👻		Descriptive statistics	Dat	ta processing - H		Dataset versions of Exp1_R2_pept			
								Normalized - peptide			
	session	R source code (E	eta)								
mow	15 - en	itries							Search:		
	Date		0	Dataset	0	History					
1	Mon Oct	16 09:10:58 2017		Normalized - peptide		Current dataset has o	change	d. Now, it is Normalized - peptide			
2	Mon Oct	16 09:10:56 2017		Normalized - peptide		Normalization : data r	normal	ized with the method Quantile Cen	tering		
3	Mon Oct	16 09:10:38 2017		Filtered - peptide		Current dataset has o	change	d. Now, it is Filtered - peptide			
4	Mon Oct	16 09:10:37 2017		Filtered - peptide		Filtering :Filtered with	whole	Matrix (threshold = 1). Contamina	nts deleted Rev	erse deleted	
5	Mon Oct	16 09:06:12 2017		Original - peptide		Current dataset has o	change	d. Now, it is Original - peptide			
6	Mon Oct	16 09:06:12 2017		Original - peptide		Open : file opened					
7											

Figure 11: Example of the log of a session in ProStaR

Command line: In *DAPAR*, the functions for these three plots are:

- for the dataframe parameter: mvPerLinesHisto_HC(), mvHisto_HC() and mvPerLinesHistoPerCondition_HC(),
- for an object of class MSnSet: wrapper.mvPerLinesHisto_HC(), wrap per.mvHisto_HC() and wrapper.mvPerLinesHistoPerCondition_HC().

3.3.2 Data explorer

This panel allows viewing the content of the MSnSet structure. It is made of four tables, which can be displayed one at a time thanks to the radio button on the left menu. The first one, named "Quantitative data" contains quantitative values (see Fig. 14). The missing values are represented by empty cells.

The second tab is named "Protein metadata" or "Peptide metadata". It contains the metadata of the proteins (see Fig. 15).

The third tab is named "Replicate metadata". The information displayed here is the one entered by the user during the import step (see Fig. 16).

The last tab, named "Dataset history" contains the log of the previous processing. Contrarily to the "Session log" panel (see Section 3.2.6), the information here does not relate to the session, and is saved from a session to the next one.

Command line: The *DAPAR* functions to get the three first tables are in fact those from the *MSnbase* package: exprs() (Quantitative data), fData() (Analyte metadata) and pData() (Replicate metadata). Similarly, the "Dataset history" information is also accessible. In fact, it is stored in the slot (process ingData) of the current MSnSet object. In a R console, if obj is the current dataset, it can be accessed by entering:

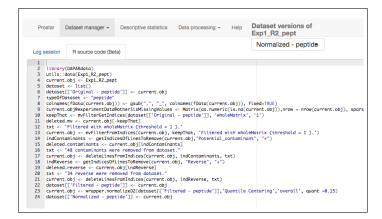


Figure 12: Example of the R code generated during a session in ProStaR

Prosta	ir Dataset manager +	Descriptive statistics	Data processing +	Help	Exp1	set versions of _R2_pept
Overview	v Miss. values Data	explorer Corr. matrix	Heatmap Box	kplot Violinplot Densityplot	CV distr.	ginal - peptide 👻
These barp	lots display the distribution of	f missing values in the dat	aset.			
	#[non-NA values] b	oy replicate =		#[lines] with X NA values	=	#[lines] with X NA values (condition-wise)
3000 —			1500			12.5k
2500		2 495	1250			
2500 -			1250			104
2000 —			1000			
1500 -			750		715	7.5k
1500 -			750			54
1000 —			- 500			
			250			2.5k
			230			
500						

Figure 13: Histrograms for the overview of the missing values

Prostar Dataset manager - L	M. Descriptive statistics	Data process	ing + Help			Dataset versions of Exp1_R2_pept		
Overview Miss. values Data e	corr. matrix	Heatmap	Boxplot Violinplot	Densityplot	Original - p	peptide -		
	Show 10 ¢ en	tries				Search:		
Choose the tab to display	Inte	nsity.D.R1 🗄	Intensity.D.R2 0	Intensity.D.R3	Intensity.E.R1 0	Intensity.E.R2 0	Intensity.E.R3	
 Quantitative data 	0	25.278	24.996	24.487	25.502	25.025	24.69	
Peptides metadata Replicate metadata	1	24.422	24.291	24.17	24.69	24.623	24.65	
 Dataset history 	2	24.527	24.411	24.376	24.822	24.585	24.75	
	3	24.088	23.866	24.335	24.532	24.674	24.76	
Show full length intensities	4	24.607	24.836	24.414	24.455	24.608	24.92	
	5					22.18	21.93	
	6	24.684	24.302	24.255	24.937	24.614	24.58	
	7	27.113	27.18	27.27	27.38	27.351	27.38	
	8	21.839	21.811	23.116	21.859	22.157		
	9	31.092	31.104	31.195	31.208	31.213	31.29	
	Showing 1 to 10 c	f 14,048 entries			Previous 1 2	3 4 5	1405 Next	

Figure 14: View of quantitative data in the MSnSet dataset

> getProcessingInfo(obj)

			Exp1_R2_pept	
Overview Miss. values Data exp	plorer Corr. matrix Heatmap Boxplot	Violinplot Densityplo	original - peptide -	
	Show 10 ¢ entries		Search:	-
Choose the tab to display	Sequence 0	Length Missed.cleava	ges 🕴 Mass 🌾 Proteins	
 Quantitative data 	0 AAAAQDEITGDGTTTVVCIVGEIIR	25	0 2559.285 sp[P39079]TCPZ_YEAST	
 Peptides metadata Replicate metadata 	1 AAADAISDIEIK	12	0 1215.6347 sp[P09938]RIR2_YEAST	
O Dataset history	2 AAADAISDIEIKDSK	15	1 1545.7886 sp[P09938]RIR2_YEAST	
	3 AAAEGVANIHIDEATGEMVSK	21	0 2112.0157 sp[P15180[SYKC_YEAST	
Show full length intensities	4 AAAEYEKGEYETAISTINDAVEQGR	25	1 2714.2671 sp[P15705 STI1_YEAST	
	5 AAAISIIGTAVQNNIDSQNNFMK	23	0 2419.2166 sp[P38260]FES1_YEAST	
	6 AAAPGIQIVAGEGFQSPIEDR	21	0 2125.0804 sp[Q04697]GSF2_YEAST	
	7 AAAPTVVFIDEIDSIAK	17	0 1758.9404 sp P25694 CDC48_YEAST	
	8 AACIVQNGIATWFPIAVTK	19	0 2059.0925 sp[P33892]GCN1_YEAST	
	9 AADAIIK	8	0 813.496 sp[P00925]ENO2_YEAST;sp[P00924]ENO1_YE	EAS
	Showing 1 to 10 of 14,048 entries		Previous 1 2 3 4 5 1405	N

Figure 15: View of feature meta-data in the MSnSet dataset

Prostar Dataset manager - Li	Descriptive statistics	Data processing - Help			Dataset versions of Exp1_R2_pept	of
					Original - peptide	-
Overview Miss. values Data ex	Show 15 ¢ entries	Heatmap Boxplot	Violinplot Dens	ityplot CV distr.		Search:
Choose the tab to display		Experiment	÷ Label	Bio.Rep	Tech.Rep	Analyt.Rep
Quantitative data	Intensity.D.R1	Intensity.D.R1	D	1	1	1
 Peptides metadata Replicate metadata 	Intensity.D.R2	Intensity.D.R2	D	2	2	2
 Dataset history 	Intensity.D.R3	Intensity.D.R3	D	3	3	3
	Intensity.E.R1	Intensity.E.R1	Е	1	4	4
Show full length intensities	Intensity.E.R2	Intensity.E.R2	Е	2	5	5
	Intensity.E.R3	Intensity.E.R3	E	3	6	6
	Showing 1 to 6 of 6 er	tries				Previous 1 N

Figure 16: View of samples meta-data in the MSnSet dataset

3.3.3 Correlation matrix

In this tab, it is possible to visualize the extent to which the replicates correlate or not (see Fig. 17). The contrast in the matrix may be changed by modifying the color gradient.



Figure 17: Correlation matrix for the quantitative data

Command line: In *DAPAR*, the corresponding function are:

- for the dataframe parameter: corrMatrixD_HC(),
- for an object of class MSnSet: wrapper.corrMatrixD_HC().

3.3.4 Heatmap

A heatmap is drawn with the associated dendrogram (see Fig. 18). The colors represent the intensities: red for high intensities and green for low intensities. White color is reserved for missing values. The dendrogram shows the hierarchical classification of the samples. This classification can be tuned by two parameters:

- Distance: the parameter *method* of the function stats::dist. The default value is 'euclidean'
- **Linkage**: the parameter *method* of the function **stats::hclust**. The default value is *'complete'*



Figure 18: Heatmap and dendrogram for the quantitative data

Command line: In *DAPAR*, the corresponding function are:

- for the dataframe parameter: heatmapD(),
- for an object of class MSnSet: wrapper.heatmapD().

3.3.5 Boxplot

The protein distribution by replicates is summarized with boxplots (see Fig. 19). The user can change the legend of the samples (X-axis) by checking items in the checkboxes group. The colors of the boxes correspond to the different conditions (column **Label** in the table of *Samples Meta Data*).

Command line: In *DAPAR*, the corresponding functions are:

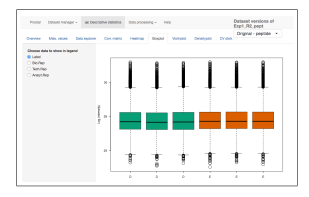


Figure 19: Boxplot for the quantitative data

- for the dataframe parameter: boxPlotD(),
- for an object of class MSnSet: wrapper.boxPlotD().

3.3.6 Violin plot

The protein distribution by replicates is summarized with violin plots (see Fig. 20). The user can change the legend of the samples (X-axis) by checking items in the checkboxes group. The colors of the boxes correspond to the different conditions (column **Label** in the table of *Samples Meta Data*).

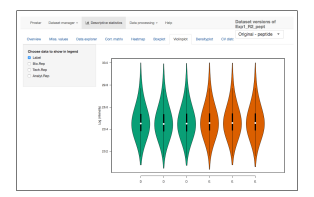


Figure 20: Violin plot for the quantitative data

Command line: In *DAPAR*, the corresponding functions are:

- for the dataframe parameter: violinPlotD(),
- for an object of class MSnSet: wrapper.violinPlotD().

3.3.7 CV distribution

This plot shows the distribution of the CV of the log-intensity of proteins for each condition (see Fig. 21). For better visualization, the user can zoom in by click-and-drag (see Fig. 22).

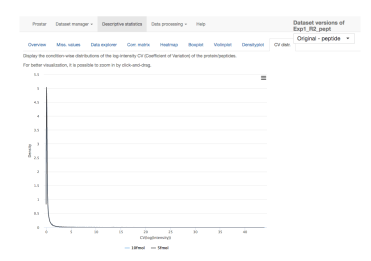
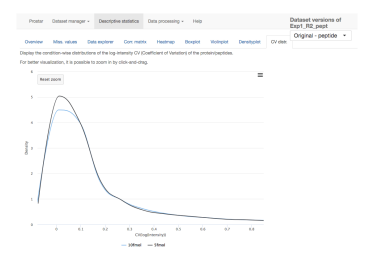


Figure 21: CV distribution for the quantitative data





Command line: In *DAPAR*, the corresponding function is

- for the dataframe parameter: CVDistD_HC(),
- for an object of class MSnSet: wrapper.CVDistD_HC().

3.3.8 Density plot

This plots shows the distribution of the log-intensity of proteins for each condition (see Fig. 23).

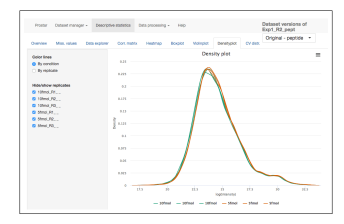


Figure 23: Densityplot the quantitative data

Two options are available to custom the plot:

- Plot to show which defines how to color the replicates: one color for each condition (value "By condition") or one color per replicate (value "By replicate"). By default, the data are colored by condition.
- Select data to show Select the replicates to display. By default, all the replicates are showed.

Command line: In *DAPAR*, the corresponding functions are:

- for the dataframe parameter: densityPlotD_HC(),
- for an object of class MSnSet: wrapper.densityPlotD_HC().

3.4 Data processing

The "Data processing" menu contains the 5 predefined steps of a quantitative analysis. They are designed to be used in a specific order:

- 1. Filtering
- 2. Normalization
- 3. Missing values imputation
- 4. Aggregation
- 5. Differential analysis

6. Gene Ontology analysis

For each step, several algorithms or parameters are available, and they are toroughly detailled in the sequel of this section.

During each of these steps, it is possible to test several options, and to observe the influence of the processing in the descriptive statistics menu (see Section 3.3), which is dynamically updated.

Finally, once the ultimate tuning is chosen for a given step, it is advised to save the processing. By doing so, another dataset appears in the "Dataset versions" list (see Section 3.6). Thus, it is possible to go back to any previous step of the analysis if necessary, without starting back the analysis from scratch.

3.4.1 Filtering

In this step, the user may decide to delete several peptides or proteins according to several criteria: If the amount of missing values is too important to expect confident processing (Tab 1); or if they are identified as reverse sequences (for target-decoy approaches) or contaminants (Tab 2).

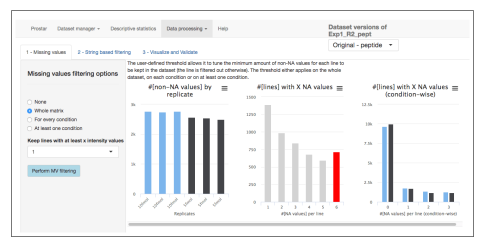


Figure 24: Interface of the filtering tool - 1

To filter the missing values (first tab called "Missing values"), the choice of the lines to be deleted is made by different options (see Fig. 24):

- **None**: No filtering, the quantitative data is left unchanged. This is the default option;
- Whole Matrix: The lines (across all conditions) in the quantitative dataset which contain less non-missing value than a user-defined threshold are deleted;

							R2_pe						
- Missing values 2 - String based f	iltering 3 - Visualize and Valid	ate				Origi	inal - pe	eptide	•				
String based filtering options													Ξ
• • • •	Quantitative data (13873 lines)											99.3	35%
Filter contaminants													
Choose column													
Potential_contaminant	Contaminants (49 lines)	0.35%											
Choose prefix	containing (15 miles)	0.55%											
+													
Filter reverse	Reverse (42 lines)	0.3%											
Choose column													
Reverse -		0	10	20	30	40	50 Pource	60 entage	70	80	90	100	11
Choose prefix	String-based filt	ering	don	е									
+		-											

Figure 25: Interface of the filtering tool - 2

Prostar Dataset manager - D	escriptive st	atistics	Data processing	✓ Help		Exp1_R2	versions of _pept		
- Missing values 2 - String based	filtering	3 - Vis	ualize and Validate			Original	- peptide 🔻		
	Show	15 \$	entries					Search:	
Filtered data display		$\mathbf{ID} \ \diamondsuit$	Intensity.D.R1	Intensity.D.R2	Inter	nsity.D.R3	Intensity.E.R1	Intensity.E.R2 \u00e9	Intensity.E.
Choose the data to display	5	5						22.18	21.932
 Quantitative data 	8	8	21.839	21.811	23.11	6	21.859	22.157	
Meta data	12	12					23.338	22.731	
Choose the type of filtered data Deleted on missing values	13	13		24.567	24.36	2	24.229		24.225
Deleted contaminants	20	20		22.79	22.10	1		23.112	
Deleted reverse	21	21							
	22	22	22.245		21.80	1			
Show full length intensities	23	23	22.240	22.372	21.00				
-				22.312					
Save filtered dataset	24	24							23.152
	29	29	25.894	24.454	23.52	2	25.283	23.529	
	35	35	22.783	22.346			22.405	21.4	21.218
	38	38		22.933	22.48	5	23.343	23.52	23.519
	39	39							
	42	42							
	43	43	22.293	22.998			23.6	23.54	23.259
	Show	ing 1 to	15 of 5,205 entries			Previous	1 2 3	4 5	347 Next
		-	the data, validate th	- 6u					

Figure 26: Interface of the filtering tool - 3

- For every condition: The lines for which each condition contain less non-missing value than a user-defined threshold are deleted;
- At least one condition: The lines for which at least one condition contain less non-missing value than a user-defined threshold are deleted;

The user can visualize the effect of filter options without changing the current dataset by clicking on "Perform filtering". If the filtering does not produce the expected effect, the user can test another one. To do so, one simply has to choose another method in the list and click again on "Perform filtering". The plots are automatically updated. This action does not modify the dataset but offers a preview of the filtered data. The user can visualize as many times he/she wants several filtering options.

Afterwards, the user can choose to remove contaminants and reverses in the second tab called "String based filtering". To do so, he/she selects the appropriate columns of the metadata listed in the dropdown menus. Then, he/she specifies in each of these columns the prefix chain of characters that identifies the analytes to filter. Note : the button "Perform string-based filtering" is disabled until all the fields are complete.

Remark: If he/she has no idea of the prefixes, he/she can switch to the Data Explorer in the Descriptive Statistics menu, so as to visualize the corresponding metadata.

Once the choices are made, the user click on "Perform string-based filtering" to remove corresponding lines. Then, the barplot beside shows the proportion of quantitative data, contaminants and reverses that were filtered out.

Once the filtering is appropriately tuned, the user goes to the last tab (called "Visualize and Validate") (see Fig. 26), to visualize the set of analytes that have been previously filtered. On the left panel, one chooses among the lines filtered on missing values, contaminants or reverse; Then, the corresponding data table is diplayed on the right panel. Finally, one clicks on "Save filtered dataset" so as to validate the user's choice and to apply it to the dataset. The information related to the type of filtering as well as to the chosen options appears in the Session log tab (see Section 3.2.6). A new dataset is created; it becomes the new current dataset and its name appears in the **Dataset versions** menu at the top of the screen. All plots and tables available in *ProStaR* are automatically updated.

Command line: In *DAPAR*, the function to filter missing values is mvFilter(). The other types of filters corresponds to classical data structure manipulation with R.

3.4.2 Normalization

The next step is to normalize the replicates so as to have more accurate comparisons. *ProStaR* offers a number of different normalization routines that are described below. In order to visualize the data after normalization, three plots are displayed: a boxplot, a plot that displays the differences between data before and after the normalization and a densityplot (see Fig. 27). The first and the third plots are the same as the one showed in **Descriptive Statistics**, thus they have the same options (see Sections 3.3.5 and 3.3.8).

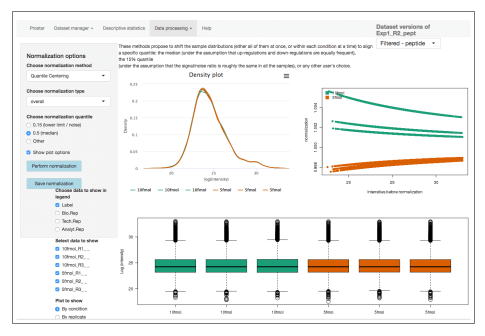


Figure 27: Interface of the normalization tool

If no normalization is necessary, it is possible to skip this step. If the user wants to compare the influence of several normalization methods, it is possible to select them in a row, and to alternate between this menu and the "Descriptive statistics" one. It is possible to go back to the original dataset by selecting "None". Several methods are implemented:

- **Global alignment** These methods propose normalizations of important magnitude that should be cautiously used:
 - sum by columns operates on the original scale (not the log2 one) and propose to normalize each abundance by the total abundance of the sample (so as to focus on the analyte proportions among each sample).
 - quantile alignment proposes to align the quantiles of all the replicates as described in [ref1]; practically it amounts to replace abundances by order statistics.

- **Quantile centering** These methods propose to shift the sample distributions (either all of them at once, or within each condition at a time) to align a specific quantile: the median (under the assumption that up-regulations and down-regulations are equally frequent), the 15 quantile (under the assumption that the signal/noise ratio is roughly the same in all the samples), or any other user's choice. Two parameters are available:
 - Normalization type: the centering can operate over the entire dataset (value "overall") or over each condition (value "within conditions"),
 - Value of quantile (in %): 0.15 (lower limit/noise), 0.5 (median) or Other (In that case, the user can choose its own quantile value).
- **Mean centering** These methods propose to shift the sample distributions (either all of them at once, or within each condition at a time) to align their means. It is also possible to force unit variance (or not). Two parameters are available:
 - Normalization type: the centering can operate over the entire dataset (value "overall") or over each condition (value "within conditions"),
 - **Include variance reduction:** Let the user choose if he/she wants to rescale the dataset to have unitary variance.

Each time the user selects a method, the explanation above is displayed. The user can visualize the effect of a normalization method without changing the current dataset. If the normalization does not produce the expected effect, the user can test another one. To do so, one simply has to choose another method in the list and click on "Perform normalization". The plots are automatically updated. This action does not modify the dataset but offers a preview of the normalized quantitative data. The user can visualize as many times he/she wants several normalization methods. Once he finds the correct one, he/she validates his/her choice by clicking on "Save normalization". Then, a new "normalized" dataset is created and loaded in memory. The method of normalization that has been used is added to the Session log tab (see section 3.2.6). It becomes the new current dataset and the name "Normalized" appears in "Dataset versions". All plots and tables in other menus are automatically updated.

Command line: In *DAPAR*, the functions for the plot "before-after normalization" are:

- for the dataframe parameter: compareNormalizationD(),
- for an object of class MSnSet: wrapper.compareNormalizationD() .

The corresponding functions for the normalization are:

- for the dataframe parameter: normalizeD(),
- for an object of class MSnSet: wrapper.normalizeD().

3.4.3 Imputation

Two plots are available in order to help the user choose the right imputation method for his dataset (see Fig. 28).

The scatter plot on the left hand side displays the proteins in a space spanned by the mean abundance (x axis) and the number of missing values (y axis). Note that for each protein, as many points (of different colors) as conditions are displayed, for each condition is processed independently of the others. As a result, the maximum value on the y axis is given by the number of replicates in a condition (depending on the filtering step). Let us note that the points have been slightly jittered on the y axis to enhance a better visualization.

This plot indicates how the missing values are distributed over the range of intensity: if there are lots of missing values in the low intensity region (indicating a censoring mechanism produced the missing values) or if they are uniformly distributed.

The heatmap on the right hand side clusters the proteins according to their distribution of missing values across the conditions. Each line of the map depicts a protein. On the contrary, the columns do not depicts the replicates anymore, as the abundance values have been reordered so as to cluster the missing values together. Similarly, the proteins have been reordered, so as to cluster the proteins that have a similar amount of missing values distributed in the same way over the conditions. Each line is colored so as to depicts the mean abundance value within each condition. This heatmap is also helpful to decide what is the main origin of missing values (random missingness or censoring of the low intensities).

The user can choose one of the several available imputation methods, depending on the type of missing values:

If the missing values are due to a mixture of censorship and of randomness mechanisms, it is advised to use the functions of the package *imp4p*. This package is a collection of proteomic-specific multiple imputation methods that operate on peptide-level datasets and which propose to impute each missing value according to its nature (censored or random). Two parameters are available: the number of iterations (the more iterations, the more accurate the results, yet the more time-consuming) and a check-

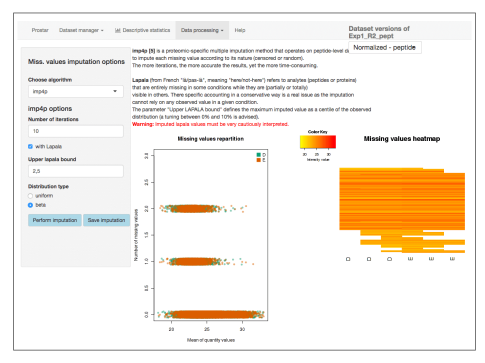


Figure 28: Interface of the imputation of missing values tool

box to let the user choose if he/she wants to impute the LAPALA. This term coined from French "là/pas-là"(meaning "here/not-here") refers to analytes (peptides or proteins) that are entirely missing in some conditions while they are (partially or totally) visible in others. Their specific accounting in a conservative way is a real issue as the imputation cannot rely on any observed value in a given condition. The parameter "Upper LAPALA bound" defines the maximum imputed value as a centile of the observed distribution (a tuning between 0% and 10% is advised).

• As an alternative, it is possible to rely on several basics methods from the state of the art to perform imputation: KNN (K Nearest Neighbors) from *impute* package, MLE (Maximum Likelihood Estimation) from *norm* package, or deterministic quantile imputation. In the latter, the user tunes a specific quantile, which is computed replicate-wise, so as to provide an imputation value for each replicate. Possibly, it is possible to multiply these values by a user-tuned factor (for instance, one may use $q(5\%) \times 0.9$ instead of q(1%)).

To date, we advise to use IMP4P for peptide-level imputation and other methods for protein-level imputation.

The user can visualize the effect of an imputation method without changing the current dataset. If the imputation does not produce the expected effect, the user can test another one. To do so, one simply has to choose another method in the

list and click on "Perform imputation". The plots are automatically updated. This action does not modify the dataset but offers a preview of the imputed quantitative data. The user can visualize as many times he/she wants several imputation methods. Once he finds the correct one, he/she validates his/her choice by clicking on "Save imputation". Then, a new "imputed" dataset is created and loaded in memory. The method of imputation used is added to the Session log tab (see Section 3.2.6). This new dataset becomes the new current dataset and the name "Imputed" appears in "Dataset versions". All plots and tables in other menus are automatically updated.

Command line: In *DAPAR*, the function used to impute the missing values is mvImputation(). The two aforementioned plots are obtained with respectively:

- for the dataframe parameter: mvTypePlot() and mvImage(),
- for an object of class MSnSet: wrapper.mvTypePlot() and wrapper.mvImage().

3.4.4 Aggregation

When working on a protein dataset, this step should be bypassed. On the other hand, when working on peptide datasets, one may want to conduct the differential analysis at protein level, for proteins are the biological units of interest. To do so, it is necessary to estimate the abundance of the proteins on the basis of those of the peptides. This is what the Aggregation step is made for.

First, the user chooses the "protein id" of the dataset, i.e. the column in the metadata, that contains the IDs of all the parent proteins for each peptide. Two barplots show up (Fig. 29). They provide the distribution of proteins according to their number of peptides (either all of them, or only those which are specific to a single protein). These statistics are helpful to visualize the adjacency matrix of the peptide-protein graph, that is sometime rather big.

Second, a checkbox is used to indicate whether the user wants the shared peptides to be accounted for during the aggregation process.

Third, the aggregation method itself ust be chosen:

- Sum: that is the sum of the peptide intensities,
- Mean: the mean of the peptide intensities,
- Sum on top n: that is the sum over the N peptides with the highest median intensities - in this case, the additional parameter N must be tuned.

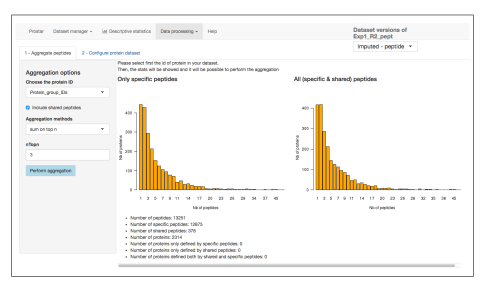


Figure 29: Interface of the agregation tool - 1

Prostar Dataset ma	anager - Descriptive statistics	Data processing -	Help -	ataset versions of p1_R2_pept
1 - Aggregate peptides	2 - Configure protein dataset		I	mputed - peptide 👻
	eta-data (related to proteins) that hav ains the protein ID if you wish to perfo		new protein	dataset.
Save aggregation				

Figure 30: Interface of the agregation tool - 2

On the next tab, the user selects the columns of the peptide dataset that are of interest to be kept in the metadata of the protein dataset (e.g. the sequence of the peptides). The effect of this action is to compile, for a given parent-protein, the information of all of its child-peptides, and to store them in a dedicated column. Once done, one validates the user's choice by clicking on "Save aggregation". Then, a new "aggregated" dataset is created and loaded in memory. The aggregation method that was finally used is recorded in the Session log tab (see section 3.2.6). This new dataset becomes the new current dataset and the name "Aggregated" appears in "Dataset versions". All plots and tables in other menus are automatically updated. As the new dataset is a protein one, the "Aggregation" menu has been disabled. Thus, the interface automatically switches to the "Descriptive Statistics" menu in order to let the user check the results of the aggregation step.

The aggregation being more computationaly demanding than other processing steps, the current version of *ProStaR* does not provide the same flexibility regarding the parameter tuning. Here, it is necessary to save the aggregation result first, then, check the results in the "descriptive statistics", and possibly to go back to the imputed dataset with the "Dataset versions" dropdown menu to test another aggregation tuning. Contrarily to other processing steps, it is not possible to visualize on-the-fly the consequences of the parameter tuning, and to save it afterwards. We are currently working on improving this issue for the next versions of *ProStaR*.

Naturally, the output of this step is not a peptide dataset anymore, but a protein dataset. As a result, all the plots available in *ProStaR* are deeply modified. For instance, the barplots summarising the peptide-protein graphs disapear because they have become meaningless.

Command line: In *DAPAR*, the function used to compute the adjacency matrix peptides-proteins is **BuildAdjacencyMatrix()** and the one used to agregate the peptides into proteins is **AggregatePeptides()**. The aforementioned plot is obtained with the functions **GraphPepProt()**.

3.4.5 Differential analysis

This step cannot be conducted if the dataset still contains some missing values: They must be imputed before.

The differential analysis is divided into four steps, each impersonated by a different tab:

- Volcano plot,
- *p*-value calibration,
- FDR,
- Validate & save.

Volcano plot (see Fig. 31): It is a scatter plot where each analyte is represented by 2 coordinates, namely a p-value on the Y-axis (more precisely $-\log 10(p$ -value)) and a fold change (FC) on the X-axis. Regarding the computation of the p-values, two tests are available in *DAPAR*, depending on the user's choice: the Welch t-test (from package *stats*) and the moderated t-test (from package *limma*). As an option, it is possible to redefined the sets of conditions that are tested one against the other. Then, the p-values are computed and a volcanoplot is displayed. Finally, the user can tune a threshold on the FC. It allows discriminating some analytes for which the difference of expression between the condition is not important enough to be biologically relevant.

This is an interactive plot which reacts to mouse's events :

- When the user puts the pointer of his mouse over a point of the plot, a tooltip window appears and shows some informations about that point. He can select the items to show in the Select widget where the different choices correspond to the columns of the feature meta-data table. The tooltip window is automatically updated,
- When the user clicks on a point, a table is displayed above the volcanoplot. It shows the values of intensities for all the samples related to the selected point. The cells colored in blue indicate that the corresponding value was a missing value in the original dataset and has been imputed,
- The user can click and draw a rectangle on the plot to zoom in. By clicking on the button named "Reset zoom", the user can return to the entire plot.

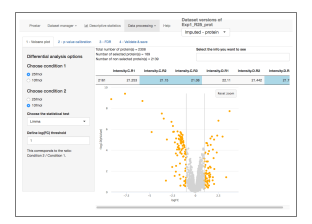


Figure 31: Volcanoplot of the differential analysis tool - 1

p-value calibration (see Fig. 32): In this tab, the fonctionalities of *CP4P* have been wrapped. Future versions of *ProStaR* will propose a more refined integration. To date, we redirect the reader to the *CP4P* tutorial: https://sites.google.com/site/thomasburgerswebpage/download/tutorial-CP4P-4.pdf?attredirects= 0.

FDR (see Fig. 33): This tab also displays the volcano plot. A threshold along the *p*-value axis can be tuned by the user, so as to discriminate the differentially abondant proteins (which are highlighted). A horizontal straight line is drawn to visualize the threshold. The corresponding FDR is computed. The user can adjust the thresholds in order to select the maximum of proteins by minimizing the FDR.

Command line: In *DAPAR*, the function used to compute the FDR is diffAna ComputeFDR().

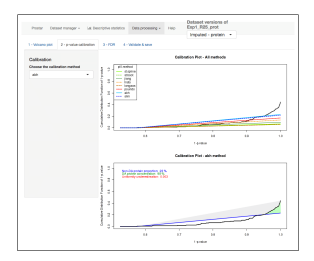






Figure 33: p-Value threshold of the differential analysis tool - 3

Validate & save (see Fig. 34): A table shows the results of the statistical test (see Fig. 34): the value of -log10(p-value) and the Fold Change (*i.e.* the log2 of the ratio of the mean values per condition). Finally, it is advised to save the results by clicking on "Save diff analysis". Then, a new "DiffAnalysis" dataset is created and loaded in memory. This dataset is the same as the previous one, except that three columns have been added in the "Quantitative data" table: "-log10(p-value)", "Fold Change" and "Significant". The two first contain the coordinates of the proteins on the volcano plot, and the third one contains a boolean value indicating whether each protein is differentially abondant or not. As with the other processing steps, the information related to the user's choices is added to the "Session log" tab (see section 3.2.6) of this new dataset. It becomes the new current dataset and its name, "DiffAnalysis.<test>" (where <test> indicates the test performed), appears in "Dataset versions". All plots

and tables in other menus are automatically updated. Note that it is possible to keep stored in memory different "DiffAnalysis" datasets: one for each type of $\langle test \rangle$.

Prostar Dataset manager - Int De	scriptive 3 - F		4 - Validate & save	ing - Help	Exp1_R25_prot DiffAnalysis.Limma - protein				
	Show	10 ;	entries			Search:			
Save diff analysis		id	Å	logFC 🍦		P.Value 🕴		Sign	ificant
	10	10	-	1.50192465004003	0.0000076856	9647037205			
	11	11	-	1.63084289353196	0.0001773	0370606419			
	12	12	-	1.78856451062824	0.000038253	6871337211			
	13	13	-	1.73373175682144	0.000071943	2309892085			
	14	14	-	1.41582407003764	0.0000094145	0556717109			
	15	15		2.12242770429974	0.00021154	5774617801			
	16	16	-	1.78238193213426	0.000090754	4460088037			
	17	17		1.51192780533265	0.000016058	9994651451			
	18	18	-	1.26164572409617	0.00039351	1686388655			
	19	19	-	1.57455400751554	0.000009123	7713955135			
	Showir	ng 1 to	10 of 95 entries		Previous 1 2	3 4 5	ō	10	Nex

Figure 34: Table of the results of statistical test in the differential analysis tool

Command line: The *DAPAR* functions for the Welch *t*-test and moderated *t*-test are diffAnaWelch() and diffAnaLimma(), respectively. These functions return a data.frame which contains 2 columns: the p-values and the Fold Change of the test. These columns can be added to the current MSnSet object imputed_dataset (as explained earlier) with the function diffAnaSave():

> res <- diffAnaLimma(imputed_dataset, condition1, condition2)
> obj <- diffAnaSave(imputed_dataset, res, "limma", condition1, condition2)</pre>

Moreover, diffAnaSave() adds the aforementioned third column named "Significant" to the MSnSet object. Two optional arguments allows the user defining the thresholds on the *p*-values and on the Fold Change, so has to be more or less stringent on the number of proteins called "Significant".

3.4.6 Gene Ontology analysis (Beta)

The Gene Ontology (GO, www.geneontology.org) is a controlled vocabulary for annotating three biological aspects of gene products. This ontology is made of three parts : Molecular Function (MF), Biological Process (BP) and Cellular Component (CC).

GO analysis is the last step proposed in the "Data processing" menu. It aims to provide the user with a "global view" of what is impacted (in a biological point of view) in his/her experiment, by showing which GO terms are represented (GO classification tab), or over-represented compared to a reference (GO Enrichment tab).

Prostar relies on the package *clusterProfiler* to perform both GO Classification and GO Enrichment. We propose a GO analysis interface with four separated tabs (see Fig. 35):

- GO Setup
- GO classification
- Go Enrichement
- Save GO analysis

Prostar Dataset manager +	Descriptive s	itatistics I	Data processing 👻	Help	Dataset versions of Exp1_R2_pept	
GO Setup GO Classification	30 Enrichme	ent Save	GO analysis		Aggregated - protein	
General setup Source of protein ID Select a column in dataset Choose a file	Thes	e proteins are	eins have not been listed in the table be		2314).	
Select column which contains prote	in ID (Show	v 10 ¢ ent	ries		Search:	
Leading_razor_protein		protid 🔅	nb.pep.used.Inte	nsity.D.R1 🔅	nb.pep.used.Intensity.D.R2 \u00e1	nb.pep.used.Intensity.D.R3 n
Id From	40	40		13	13	13
UNIPROT	15	15		6	6	6
Genome Wide Annotation	54	54		3	3	3
Yeast (org.Sc.sgd.db)	28	28		26	26	26
Ontology	30	30		15	15	15
Molecular Function (MF)	50	50		10	10	10
	18	18		1	1	1
Map proteins IDs	32	32		13	13	13
	35	35		10	10	10
	34	34		3	3	3

Figure 35: Input parameters for GO analysis (GO Setup tab)

The left-hand side of the **GO Setup** tab allows it to set the input parameters, namely:

- Source of protein ID: user indicates either a column in the current dataset or chooses a file (1 ID per line).
- Id From: the type of ID supplied (UNIPROT by default).
- Genome Wide Annotation: the organism to consider for the analysis.
- Ontology: the level of the ontology to work with.

Once these parameters filled, clicking on "Map proteins IDs" launches the mapping of the IDs onto the GO categories of the annotation package. Then, on the right-hand side of the panel, the proportion of proteins that cannot be mapped onto the annotation package is indicated (this informative ouput does not interrupt the process, unless no protein maps).

Command line: The R function to map protein IDs on the annotation package is **bitr()** (function of the Bioconductor package *clusterProfiler*). In *DAPAR*, the call to the **bitr** is integrated inside the group_GO() and enrich_GO() functions. The mapping is done implicitly and the user does not have to perform the mapping.

Next step is to perform either GO Classification or GO Enrichment (or both).

In the **GO Classification** tab (see Fig. 36), one has to indicate which level(s) of the ontology to consider. Then clicking on the "Perform GO grouping" button launches the analysis (function groupGO() of the *clusterProfiler* package). The graphics shows the most represented GO categories for a user-defined ontology at (a) user-defined level(s).

Command line: The *DAPAR()* function to perform the GO classification is group_GO(). It returns a 'groupGOResult' instance. The plot as seen in the GO-Classification tab of ProStaR (see Fig. 36) can be generated with the barplot() function.

```
> ggo <- group_GO(data, idFrom="UNIPROT", orgdb=org.Hs.eg.db, ont="MF", level=2)
> barplotGroupGO_HC(ggo)
```

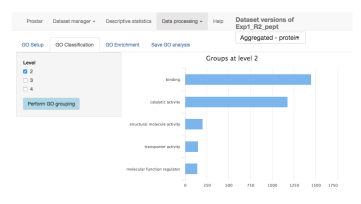


Figure 36: Tab to perform the GO classification

The **GO Enrichment** tab (see Fig. 37) allows it to know which GO categories are significantly enriched in the users list, compared to a chosen reference ('background' or 'universe'). This background can either be :

(i) the entire organism (in this case, the totality of the proteins identified with an "ENTREZGENE" ID in the annotation package specified in the GO Setup tab constitutes the background), or

(ii) the entire dataset loaded in ProStaR (e.g. all the proteins IDs of the 'Leading_razor_protein' column of the dataset, as illustrated on Fig. 35), or

(iii) a custom IDs list provided in a separate file by the user (one ID per line).

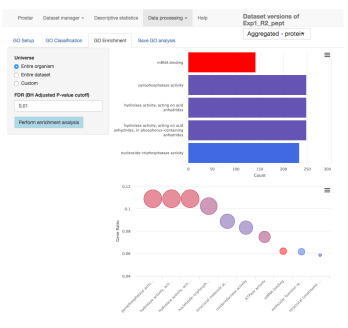


Figure 37: GO enrichment tab

The enrichment tab calls the groupGO() function of the *clusterProfiler* package. This function performs a significance test for each category, followed by a multiple test correction at a user-defined level. Concretely, this level is tuned thanks to the "FDR (BH Adjusted *P*-value cutoff)" field. Analysis is launched by clicking the "Perform enrichment analysis" button.

Once the analysis has been performed, the result is displayed via two graphics on the right-hand side of the panel (see Fig. 37). The first one (top) is a barplot showing the five most significant categories. The length of each bar represents the number of proteins within the corresponding category. The second one (bottom) is a dotplot ranked by decreasing *GeneRatio*, which reads:

 $GeneRatio = \frac{\#(Genes \text{ of the input list in this category})}{\#(Total number of Genes in the category)}.$

Command line: The *DAPAR()* function to perform the GO Enrichment is en rich_GO(). It returns an 'enrichResult' instance. The plots as seen in the GO Enrichment tab of ProStaR (see Fig. 37) can be generated with the barplot() and dotplot() functions. The universe argument allows to indicate the list of IDs considered as the reference (or background) to which the IDs input list of the user will be compared. To consider the whole organism IDs as a reference, one can extract it from the annotation package with the *DAPAR* function univ_AnnotDbPkg(). (see code below)

```
> univ.Hs<-univ_AnnotDbPkg(org.Hs.eg.db)</pre>
```

```
> ego<-group_GO(data, idFrom="UNIPROT", orgdb=org.Hs.eg.db, ont="MF",</pre>
```

```
+ universe=univ.Hs, pval=0.05, pAdjustMethod="BH")
```

```
> barplotEnrichGO_HC(ego)
```

```
> scatterplotEnrichG0_HC(ego)
```

The last tab is the **Save GO** analysis one. It allows saving the results: GO classification, GO enrichment, or both (see Fig. 38). Then, a new GOAnalysis dataset is created and loaded in memory.

As usual in ProStaR, it is possible to export this new dataset via the Dataset manager menu , either in MSnSet or in Excel format.

Prostar	Dataset manager 👻	Descriptive statistics	Data processing -	Help	Dataset versions of Exp1_R2_pept
GO Setup	GO Classification	GO Enrichment	ave GO analysis		Aggregated - protein
Choose wit	hich GO analysis to sav	re			
 Enrichm 					
 Both 					
Save ana	lysis				

Figure 38: Saving the GO analysis

Command line: In *DAPAR*, the function to save the results of the GO analysis in the MSnSet dataset is GOAnalysisSave().

Nota Bene: The GO Analysis functionalities are in beta version and improvements will be considered in the future, such as for instance:

- authorizing a larger range of input format ID;
- handling external mapping file between ID and GO ontology (or even a custom ontology), so as to be able to work on other organisms than those for which a Bioconductor annotation package exists.

Any additional suggestion is welcome.

3.5 Help

The Help screen offers various information:

- **The MSnSet format**. On this screen, there is a link to an article about the MSnSet format in order to explain its architecture to the user,
- Refs. The references associated and/or related to the packages DAPAR and ProStaR.

3.6 Versions of dataset

This major element of the Dataset manager is not in the corresponding menu, but on the contrary is detached on the right hand side of the navbar. The reason is, it is convenient to have a constant view on it. It is a drop-down menu that lists the different versions of dataset of interest, i.e. the restauration points that were progressively saved along the quantitative analysis.

Basically, each time the modifications of the current dataset are saved, the new dataset does not overwrite the previous one. On the contrary, the different versions are stored in memory. Thus, *ProStaR* keeps a history of all processing performed on a dataset. Concretely, right after creating or uploading a dataset, only a single dataset is available: it is named "Original (peptide)" or "Original (protein)" depending on the data being related to peptides or proteins. This information is registered in the MSnSet file (the slot "typeOfData" of experimentData(object)). After the filtering step, if the user saves his/her results, another dataset becomes available, named "Filtered (peptide)" or "Filtered (protein)". Similarly, after the saving of the normalization, of the imputation of missing values, of the aggregation into proteins and of the differential analysis, a new dataset is created and stored. Each time a new dataset is created, it is by default the one on which the processing goes on. However, the previous one is accessible through the "Dataset versions" drop-down menu.

At any time, the name of the current dataset and the type of data are displayed. If the user needs to return to a previous dataset (for example, the current dataset is "Imputed" and the user wants to return to "Filtered"), he/she chooses it in the select field. The dataset is then automatically loaded in memory and becomes the current one; the new dataset becomes the new current one. Naturally, all the plots that are displayed throughout the various panels of *ProStaR* are dynamically updated without any action from the user.

Remarks:

- Let us note that if the user saves the current step (let us say the imputation step), then goes back to a previous step (say the normalization step) and start working on this older dataset (for instance, by performing another imputation) and then saves it, the new version of the processing overwrites the previous version (the older imputation is lost and only the newest one is stored in memory): in fact, only a single version of the dataset can be saved for a given processing step.
- For a refined analysis regarding the influence of a processing step, it also possible to switch from an older to a newer dataset (that has been saved before) with the "Dataset versions" drop-down menu, and to observe the variations in the "Descriptive statistics" menu.

4 Bugs

Both packages *DAPAR* and *Prostar* are under active development. Despite our attention bugs may remain. To signal any, as well as typos, suggestions, etc. or even to ask a question, please contact us by email. Please join to the message as much information as possible, a reproducible example and the output of sessionInfo().

Here follow some error messages that the user may encounter and the tip to work around (please note that this section will be enriched with your feedbacks):

5 Session information

- R version 3.5.1 Patched (2018-07-12 r74967), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Running under: Ubuntu 16.04.5 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.7-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.7-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Loaded via a namespace (and not attached): BiocStyle 2.8.2, Rcpp 0.12.18, backports 1.1.2, compiler 3.5.1, digest 0.6.16, evaluate 0.11, htmltools 0.3.6, knitr 1.20, magrittr 1.5, rmarkdown 1.10, rprojroot 1.3-2, stringi 1.2.4, stringr 1.3.1, tools 3.5.1, yaml 2.2.0