

# Users' Guide

## MORE

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## Contents

<b>1</b>	<b>Introduction</b>	<b>3</b>
<b>2</b>	<b>Getting started</b>	<b>4</b>
<b>3</b>	<b>Input data</b>	<b>4</b>
<b>4</b>	<b>Generating the regression models with MORE</b>	<b>6</b>
4.1	Arguments for more() function . . . . .	6
4.2	Recommendations for users . . . . .	10
4.3	more output . . . . .	10
4.4	Running an example . . . . .	14
<b>5</b>	<b>Retrieving significant regulations from MORE results</b>	<b>17</b>
5.1	RegulationPerCondition input parameters . . . . .	17
5.2	Interpreting RegulationPerCondition output with an example . . . . .	17
5.3	RegulationInCondition . . . . .	18
<b>6</b>	<b>Plotting MORE results</b>	<b>19</b>
6.1	summaryPlot function . . . . .	19
6.2	FilterRegulationPerCondition . . . . .	20
6.3	differentialRegPlot function . . . . .	20
6.4	plotMORE function . . . . .	21
6.5	Plots for PLS models . . . . .	26
6.6	networkMORE function . . . . .	27
<b>7</b>	<b>Functional Enrichment analysis</b>	<b>31</b>

7.1	Over Representation Analysis . . . . .	31
7.2	Gene Set Enrichment Analysis . . . . .	32
<b>8</b>	<b>How to cite MORE package</b>	<b>33</b>

# 1 Introduction

One of the most common questions to be addressed when performing a multi-omics experiment is how the levels of given biological entities are being regulated by other biological entities under certain conditions. An example of this type of study would be understanding the regulatory mechanisms behind the changes in gene expression.

Potential regulators of a given gene such as miRNAs, transcription factors (TF), methylation sites, etc., can be either retrieved or predicted from public databases or obtained by a combination of experimental and computational procedures. However, a methodology for selecting the specific regulators of a particular biological system studied under certain conditions is required. This is the goal of the MORE (Multi-Omics REgulation) method: modeling a target omic (such as gene expression, protein levels, etc.) as a function of experimental variables, such as diseases or treatments, and the potential regulators of the target features. The idea is to obtain specific candidate regulators for the biological system under study by applying regression models, as Multiple Linear Regression models (MLR) or Partial Least Squares (PLS) regression. MORE facilitates the application of MLR or PLS to multi-omic data.

MORE requires several data inputs: a target omic abundance data matrix, regulatory omic data matrix/matrices, conditions of considered samples, and potential associations between target omic features and regulators (optional). With this input data, MORE generates the initial model equation, which is different for each target omic feature because each one of them has different potential regulators. MORE admits numerical omic data (continuous or discrete) or binary data.

It is recommended to fit MORE models only to target omic features that present significant changes in any of the conditions studied, for example, when considering Gene Expression data as target omic to differentially expressed genes (DEGs). DEGs can be selected with the standard procedures depending on the conditions, but DEGs selection is not included in the MORE algorithm and must be done by the user.

This idea can be extended to potential regulators since regulators that do not change are not good candidates to regulate the target omic. Even so, MORE has several functionalities to filter regulators with missing values or low variation, treat highly correlated regulators, and perform efficient variable selection.

MORE package also includes a function to retrieve the significant regulations and the magnitude of the regulatory effect under each condition considered and an additional function to graphically investigate the relationship between target omic features and regulators.

## 2 Getting started

The MORE method is available as an R package from <https://github.com/BiostatOmics/MORE.git>. As for other packages in GitHub, it can be installed from R with the following instructions:

```
> install.packages("devtools")
> devtools::install_github("BiostatOmics/MORE")
```

## 3 Input data

This section describes the main data files required by MORE to generate the regression models.

**Target data** Expression/abundance values for features of a target omic (e.g. gene expression data), in rows, under each condition, in columns. MORE accepts either a **matrix** or a **data frame**. In MORE it should be provided to **targetData** argument. See an example below:

```
> head(TestData$GeneExpressionDE)
   Sample_1 Sample_2 Sample_3 Sample_4 Sample_5 Sample_6 Sample_7 Sample_8 Sample_9
ENSMUSG00000000078 16.35068 16.43845 16.08969 16.18012 15.98894 16.24338 16.14683 16.20514 15.53671
ENSMUSG0000056999 14.04936 15.00476 15.39196 15.38862 15.48909 15.74698 16.15151 15.97799 13.66564
ENSMUSG0000024873 11.70604 11.92110 12.88689 12.96273 12.75602 12.67522 12.99590 13.11888 10.93298
ENSMUSG0000015461 16.08382 16.20221 16.27489 16.29576 16.03688 15.71148 15.45131 14.66903 14.17544
ENSMUSG0000058135 13.02257 15.52288 16.35661 16.42303 16.38726 16.13189 15.76151 15.01974 13.93261
ENSMUSG0000038208 13.52974 13.64620 13.55908 13.39410 13.36335 13.34064 13.23755 13.32766 13.15812
   Sample_10 Sample_11 Sample_12 Sample_13 Sample_14 Sample_15 Sample_16
ENSMUSG00000000078 15.27647 15.355672 15.005449 15.36413 15.16874 15.65070 15.38943
ENSMUSG0000056999 14.99625 15.436961 15.854549 15.76431 15.81027 15.65091 15.53761
ENSMUSG0000024873 11.92766 11.730738 11.649180 11.99320 12.10198 12.74034 12.69285
ENSMUSG0000015461 12.50220 9.709291 3.501127 11.07505 13.25271 14.35546 14.98628
ENSMUSG0000058135 15.07768 15.749751 15.709056 15.87783 15.57376 15.74364 14.80781
ENSMUSG0000038208 13.23678 12.969414 13.338755 13.46027 13.00960 12.95872 12.75443
```

**Condition** Matrix or data frame containing the conditions of the samples, such as treatments, diseases, strains, dose of a drug, etc. The rows of the object must be the same as the columns in **targetData** and in the same order, as shown below. In MORE it should be provided to **condition** argument

```
> TestData$edesign
   Group
Sample_1 "Control"
Sample_2 "Control"
Sample_3 "Control"
Sample_4 "Control"
Sample_5 "Control"
Sample_6 "Control"
Sample_7 "Control"
Sample_8 "Control"
Sample_9 "Disease"
Sample_10 "Disease"
Sample_11 "Disease"
Sample_12 "Disease"
Sample_13 "Disease"
Sample_14 "Disease"
Sample_15 "Disease"
Sample_16 "Disease"
```

**Regulatory omic data** This object must be a list where each element is a matrix or data frame containing the data for each “regulatory” omic (miRNA expression, transcription factor expression, etc.), with a structure similar to target data: regulators in rows and conditions in columns (the columns must be the same as in target data and in the same order). In MORE it should be provided to **regulatoryData** argument. See the example below (TestData\$data.omics\$‘miRNA-seq’).

```
> head(TestData$data.omics$`miRNA-seq`)
   Sample_1 Sample_2 Sample_3 Sample_4 Sample_5 Sample_6 Sample_7 Sample_8 Sample_9
mmu-miR-125a-5p    7100    7229   10066    8868   12649   16321   14760   15164   1655
mmu-miR-141-5p      19      49     40     107     176     183     168     343      58
mmu-miR-145a-3p  117329   85793   73670   57547   43926   21426   11769      2   11593
mmu-miR-148b-3p  125403  136517  137890  190417  150718  217291  203966  203880  108489
mmu-miR-150-5p     1256    1098     707    1025     599     253     190      0   1289
mmu-miR-152-3p     344     445     322     529     655    1082    1645    1441      63
   Sample_10 Sample_11 Sample_12 Sample_13 Sample_14 Sample_15 Sample_16
mmu-miR-125a-5p     166       8     562    2818    4997    7528    9159
mmu-miR-141-5p       8       2       9      13     117     121     283
mmu-miR-145a-3p    1539     125    7075   19253   32778   61536   106987
mmu-miR-148b-3p  116911  102194   94160  126998  150972  171944  205767
mmu-miR-150-5p      844     617     517     374     370     164      0
mmu-miR-152-3p      17       0     121      94     270     665     687
```

**Associations** For each regulatory omic, associations between potential regulators and target features can be optionally provided to indicate which are the potential regulators to be incorporated into the initial equation of the regression model. The association objects must be data frames and stored in a single **list** (attached below the example of miRNA-seq, TestData\$associations\$‘miRNA-seq’). The names of the elements of this list must be the names of the list of regulatory omic data and must be in the same order. In MORE it should be provided to **associations** argument.

If the user wants to consider all regulators of an omic as potential regulators they must set to NULL the object of this omic in the **associations** list. Moreover, if the user does not provide the list of **associations**, all regulators of all omics in **regulatoryData** will be considered potential regulators for all target features. However, this option can be very time-consuming. By default, NULL.

```
> head(TestData$associations$`miRNA-seq`)
   Gene          ID
1 ENSMUSG00000024873 mmu-miR-335-3p
2 ENSMUSG00000024873 mmu-miR-1912-3p
3 ENSMUSG00000024873 mmu-miR-615-5p
4 ENSMUSG00000024873 mmu-miR-322-5p
5 ENSMUSG00000024873 mmu-miR-1894-3p
6 ENSMUSG00000024873 mmu-miR-7082-3p
```

## 4 Generating the regression models with MORE

The **more** function in MORE adjusts a regression model for each target feature (gene, protein, metabolite, etc.) in the **targetData** object to determine which regulators and conditions (treated vs. non-treated, control vs disease, etc.) have a significant effect on the response variable inferred by advanced variable selection methods. MORE can apply more conventional regression models such as Multiple Linear Regression models (MLR) in combination with Elastic Net (EN) or Iterative Sparse Group Lasso (ISGL) regularization regression methods if the user selects 'MLR' as the method to apply. It can also use a dimensionality reduction technique such as Partial Least Squares (PLS) if the selected method is 'PLS1' or 'PLS2'. These are the arguments the function accepts, described in detail in Section 4.1.

```
more(targetData, regulatoryData, associations, omicType = NULL,
      condition = NULL, clinic = NULL, clinicType = NULL,
      minVariation = 0, percNA = 0.2, scaleType = 'auto',
      epsilon = 0.00001, interactions = TRUE, varSel = 'EN',
      alfaEN = NULL, correlation = 0.7, groupingISGL = 'MF_0.7',
      alfa = 0.05, vip = 0.8, method = 'MLR', parallel = FALSE,
      seed = 123)
```

### 4.1 Arguments for **more()** function

**targetData** Matrix or data frame containing data from a target omic with its features in rows and samples in columns. The row names must be the target omic features IDs; e.g. when gene expression is considered as the targetData, gene IDs should be the row names.

**regulatoryData** List where each element corresponds to a different omic data type (miRNAs, transcription factors, methylation, etc.). The names of this list will be the omics, and each element of the list is a matrix or data frame with omic regulators in rows and samples in columns. *Attention!* We clarify that the user can not use as regulatory omic the data considered as **targetData**. For considering those co-expression networks see *Recommendations for users* section.

**associations** List where each element corresponds to a different omic data type (miRNAs, transcription factors, methylation, etc.). The names of the elements of the list will be the omics (in the same order as in **regulatoryData**). Each element is a data frame with two columns (optionally three) describing the potential interactions between target omic features and potential regulators for that omic. The first column must contain the regulators, the second the target features IDs, and an additional column can be added to describe the type of interaction (for example, in methylation data, if a CpG site is located in the promoter region of the gene, in the first exon, or any other information). Optionally, the user can set the **associations** data frame

of an omic equal to NULL if they want to consider all the regulators of that omic as potential regulators for all the target features. They can even set **associations** to NULL if they want to consider all regulators of all omics in **regulatoryData** as potential regulators to all target features. Even if it can be done, we do not recommend the user to do it as it can be very time-consuming.

**omicType** Vector with as many elements as the number of omics, indicating whether the omic values are numeric (0) or binary (1). When NULL is indicated, **MORE** will estimate which type of omics are provided and display them on the screen. If a single value is provided, the type for all the omics is set to that value. By default, NULL. If the estimated type of omics are incorrect, the user must halt the process and manually specify the correct omic type.

**condition** Data frame or matrix describing the condition to which samples belong. Rows must be the samples, that is, the columns in the **targetData**, and columns must be the conditions to be included in the model, such as disease, treatment, etc.

**clinic** Data frame or matrix containing clinical variables values where rows must represent samples and columns variables.

**clinicType** Vector with as many elements as the number of clinical variables, indicating whether the variables values are numeric (0) or categorical/binary (1). When NULL is indicated, **MORE** will estimate which type of variables are provided and display them on the screen. If a single value is provided, the type for all the variables is set to that value. By default, NULL. If the estimated type of variables are incorrect, the user must halt the process and manually specify the clinic type.

**minVariation** Vector with as many elements as the number of omics (names of this vector will be the omics), indicating the minimum change in the standard deviation that a regulator must show across conditions in order not to be considered as having low variation and be removed from the regression models, for numerical regulators. For binary regulators, the minimum change in the proportion a regulator must show across conditions. When a single value is given, the minimum change will be considered the same for all omics and equal to this value. The user has the option to set this argument to NA if they do not want to provide a value but want to filter more than constant regulators across conditions. In this case, the value will be calculated as the 10% of the maximum observed variability across conditions for continuous regulators and as the 10% of the maximum observed proportion difference across conditions for binary regulators. Additionally, the user can combine both functionalities; indeed, the user has the option to provide a vector containing the minimum change in the standard deviation for some omics and NA others. By default, this argument is set to 0. If the selected threshold is excessively restrictive, resulting in the removal of all regulators, the model will fail to run and a warning message will be shown.

**percNA** Maximum percentage of missing values allowed in regulatoryData to construct the models. Only used in PLS models since MLR models do not allow missing values. By default, 0.2.

**scaleType** Type of scaling to be applied when adjusting a model, if scaling is requested. It can be: 'none', 'auto', 'softBlock', or 'hardBlock'. The first applies the autoscaling method; so that scales each variable independently. The second and the third apply block-scaling (soft and hard block-scaling respectively), where each regulatory omic is considered a block.

**epsilon** A threshold for the convergence tolerance in the MLR model. By default, 0.00001.

**interactions** If TRUE (default), **MORE** allows for interactions between each regulator and the conditions, or phenotypes under study.

**varSel** Variable selection method to apply. The options are different for MLR or PLS. Four options:

**EN** Applies a Multiple Linear Regression (MLR) model with ElasticNet (EN) regularization.

**ISGL** Applies a Multiple Linear Regression (MLR) model with Iterative Sparse Group Lasso (ISGL) regularization.

**Jack** Applies the Jack-Knife resampling technique to compute the significance of the coefficients in Partial Least Squares (PLS) models.

**Perm** Applies a resampling technique to compute the significance of the coefficients in Partial Least Squares (PLS) models in which the response variable is permuted 100 times to obtain the distribution of the coefficients and compute then their associated p-value.

**alfaEN** ElasticNet mixing parameter ( $\alpha$ ). By default, NULL. These are the values that can be passed to this argument:

**NULL**  $\alpha$  parameter will be automatically optimized by cross-validation. For computational efficiency, only values ranging from 0 to 1 in increments of 0.1 will be tested.

**Value between 0 and 1** ElasticNet is applied with this  $\alpha$  being the combination between ridge and lasso penalization.

**Vector of  $\alpha$ 's** ElasticNet will be applied for each of the  $\alpha$  values provided in the vector, and the one that yields the best cross-validation performance will be selected.

**Value 0** A ridge penalty will be applied.

**Value 1** A lasso penalty will be applied.

The shrinkage parameter ( $\lambda$ ) will be optimized by cross-validation in all cases.

**correlation** Correlation threshold (in absolute value) for the Multicollinearity Filter (MF) to decide which regulators are correlated, in which case, a/some representative of the group of correlated regulators is chosen to enter the model. By default, 0.7.

**groupingISGL** Option to create the groups for the Iterative Sparse Group Lasso regularization. By default, 'MF\_0.7'. There are two options:

**MF\_X** Takes the same approach as in the multicollinearity filter. Grouping the variables which correlate more than X. E.g: MF\_0.7 groups variables with correlation higher than 0.7. By default, MF\_0.7.

**PCA\_X** Extracts as many PCA components to explain X percent of the variability of the data and assigns the variables to the component in which they had the highest loadings. E.g: PCA\_0.8 extracts the minimum number of components required to explain at least 80% of the variability and groups variables according to the component they obtained the highest loading.

**alfa** Significance level to consider a regulator as significant in PLS models. By default, 0.05.

**vip** The Variable Importance in Projection, VIP, score threshold to apply together with the **alfa** threshold to take a variable as significant; both requirements should be met to take a variable as significant. By default, 0.8.

**method** Regression model to be applied: 'MLR' for Multiple Linear Regression, 'PLS1' for a different Partial Least Squares model to apply for each of the target features, or 'PLS2' for a single Partial Least Squares model for all target features (same model for all target features). The user must be aware that in the 'PLS2' model, the **association** list will not be considered and must be set to NULL.

**parallel** If FALSE, **MORE** will be run sequentially. If TRUE, **MORE** will be run using parallelization with as many cores as the available ones minus one so the system is not overcharged. If the user wants to specify how many cores they want to use, they can also provide the number of cores to use in this parameter. Parallelization is only implemented for MLR with ElasticNet regularization and PLS1 methods. By default, FALSE.

**seed** Sets the seed to guaranty reproducibility of the results for the methods that require random sampling of the observations (i.e., all MLR approaches and PLS with Perm variable selection). By default, 123.

## 4.2 Recommendations for users

1. If the omic considered in **targetData** is to be considered as a regulatory omic, the user **MUST** provide an association matrix in which each target feature is explicitly excluded as a potential regulator of itself.
2. When running the PLS2 methodology in **MORE**, we recommend running it in a cluster rather than on the desktop due to its memory consumption. More or less 25000 regulators require more than 30GB of memory to create the model. The memory usage grows exponentially according to the number of regulators and target features included in the model. The number of phenotypes or conditions also affects.
3. When selecting PLS2 methodology and considering high-dimensional datasets **targetData** or **regulatoryData** datasets, we recommend using '*Perm*' methodology for p-value computation (**varSel='Perm'**) because of memory consumption efficiency.
4. If the user still faces problems due to memory when running PLS2 models, we recommend running separate models of **MORE**: One for each condition by setting interactions = FALSE. To create the differential network, the user would only have to merge the columns of the considered conditions in the **RegulationPerCondition** matrices.

## 4.3 more output

The object returned by **more** function varies depending on the selected method.

### 4.3.1 more output for MLR

The object returned by the **more** when fitting a MLR model (with elasticnet or ISGL penalization indifferently) is a list that contains the following elements:

**ResultsPerTargetF** is a list with as many elements as target features in the **targetData** object. For each target feature, there is a list containing the following information:

**Y** Data frame with the response variable values for that target feature (y), the values fitted by the model (fitted.y), and the residuals of the model (residuals).

**X** Data frame with all the predictors included in the final model.

**coefficients** Matrix with the estimated coefficients for the regulators selected as relevant by the EN or the ISGL regularization methods.

**allRegulators** Data frame with all the initial potential regulators in rows and the following information in columns: target feature, regulator, omic, area (the third optional column in associations), filter (if the regulator has been filtered out of the model, this column indicates the reason), and Rel (1 if the regulator is

considered relevant and 0 if not). Regarding the filter column, several values are possible:

- *MissingValue*: If the regulator has been filtered out of the study because it has missing values.
- *LowVariation*: If the regulator has been filtered out of the study because it has lower variability than the threshold set by the user in **minVariation** parameter.
- *Model*: When the regulator is included in the initial equation model.
- *omic\_mcX\_X\_X*: For example, *TF\_mc1\_1\_R*. This notation is related to highly correlated regulators and how they are treated to avoid the multicollinearity problem. Only present for MLR models when EN regularization is selected. Following the *TF\_mc1\_1\_R* example, two or more regulators, which potentially regulate the target feature, are highly correlated (in absolute value). In such cases, one is chosen as the representative and indicated with *\_R*. The rest of the regulators considered if they are directly highly correlated to it are labeled with *\_P*, which means that they are positively correlated with the representative and with *\_N* if negatively correlated. Once a representative is taken for them, if there are still highly correlated regulators, the process is repeated and indicated in the *\_1* of the example. An additional row is then added to this table, with the regulator *TF\_mc1\_1\_R* and the filter label being *Model*, since only this representative is considered in the model. When there are several groups of correlated regulators for the same omic, it is indicated with *\_mc1*, *\_mc2*, etc.

**relevantRegulators** A character vector containing the relevant regulators.

**GlobalSummary** List that contains the following elements:

**GoodnessOfFit** Matrix that collects the R-squared value (which for MLRs is defined as the percentage of deviance explained by the model), the adjusted R-squared value, the Root Mean Square Error (RMSE), the Normalized Root Mean Square Error (NRMSE) and the number of relevant regulators for all the target features that had at least a relevant regulator.

**ReguPerTargetF** Matrix containing, for each omic and target feature, the number of initial regulators, the number of regulators included in the initial model, and the number of relevant regulators.

**TargetFNOmodel** List of target features for which the final MLR model with the EN or ISGL regularization could not be obtained. There are four possible reasons for that, and they are indicated: “Too many missing values”, “-Inf/Inf values”, “No regulators left after NA/LowVar filtering” and “No regulators selected after variable selection”.

**TargetFNOregu** List of target features for which there were no initial regulators, only generated in case any target feature was under this condition.

**GlobalRegulators** Vector containing the top 25% regulators, ranked by the number of target features they relevantly regulate, with a minimum of 10 regulated target features.

**HubTargetF** Vector containing the top 25% target features, ranked by the number of relevant regulators acting on them, as inferred by MORE models. At least they have to be regulated by a minimum of 10 regulators.

**Arguments** List with the arguments used to generate the model: conditions, minimum degrees of freedom in the residuals, significance level, etc.

#### 4.3.2 more output for PLS

The object returned by the **more** when fitting a PLS (PLS1 or PLS2 indifferently) model is a list that contains the following elements:

**ResultsPerTargetF** is a list with as many elements as target features in the **targetData** object. For each target feature, there is a list containing the following information:

**Y** Data frame with the response variable values for that target feature (**y**), the values fitted by the model (**fitted.y**), and the residuals of the model (**residuals**).

**X** Data frame with all the predictors included in the final model.

**coefficients** Matrix with the estimated coefficients for the regulators selected as significant by the selected p-value computation method (**varSel**) and whose Variable Importance in Projection (**vip**) has been higher than the threshold.

**allRegulators** Data frame with all the initial potential regulators in rows and the following information in columns: target feature, regulator, omic, area (the third optional column in associations), filter (if the regulator has been filtered out of the model, this column indicates the reason), and Sig (1 if the regulator is considered significant and 0 if not). Regarding the filter column, several values are possible:

- *MissingValue*: If the regulator has been filtered out of the study because it has missing values.
- *LowVariation*: If the regulator has been filtered out of the study because it has lower variability than the threshold set by the user in **minVariation** parameter.
- *Model*: When the regulator is included in the initial equation model.

**significantRegulators** A character vector containing the significant regulators.

**GlobalSummary** List that contains the following elements:

**GoodnessOfFit** Matrix that collects the R-squaredY value (the R squared of the response variable), the Q-squared (the goodness of prediction), the square root of the mean error between the actual and the predicted responses (RMSE), the Normalized Root Mean Square Error (NRMSE) and the number of significant regulators for all the target features that had at least a significant regulator.

**ReguPerTargetF** Matrix containing, for each omic and target feature, the number of initial regulators, the number of regulators included in the initial model, and the number of significant regulators.

**TargetFNOmodel** List of target features for which the final PLS model could not be obtained. There are three possible reasons for that, and they are indicated: "Too many missing values", "-Inf/Inf values", and "No regulators left after NA/LowVar filtering".

**TargetFNOregu** List of target features for which there were no initial regulators, only generated in case any target feature was under this condition.

**GlobalRegulators** Vector containing the top 25% regulators, ranked by the number of target features they significantly regulate, with a minimum of 10 regulated target features.

**HubTargetF** Vector containing the top 25% target features, ranked by the number of significant regulators acting on them, as inferred by MORE models. At least they have to be regulated by a minimum of 10 regulators.

**Arguments** List with the arguments used to generate the model: conditions, significance level, etc.

#### 4.3.3 more summary

Making use of the output object returned by **more**, in both cases in MLR and PLS models, the user can ask for a summary of the results obtained by:

```
summary(object, plot.more=FALSE)
```

This summary takes two arguments as input:

**object** MORE object obtained from applying **more** function, indifferent to the method that has been used ('MLR', 'PLS1' or 'PLS2').

**plot.more** If TRUE, the top 10 global regulators will be plotted against the target features they regulate. By default, FALSE. This option can be very time-consuming if global regulators regulate more than 50 target features and is therefore not recommended unless the user knows that only a small number of target features are involved. Instead, it is recommended to first use the summary function with the

`plot.more` parameter set to FALSE to verify that there are no more than 50 target feature-regulator associations. For visualizing specific regulatory relationships, the use of the `plotMORE` function is recommended.

The summary function will print the following information on the screen:

1. Number of genes for which a model was computed.
2. Number of genes that did not have initial regulators.
3. Number of genes for which the final model could not be obtained.
4. The mean of relevant/significant regulators of the genes.
5. Top 10 hub genes and the number of relevant/significant regulators for each.
6. Top 10 global regulators and the number of genes they regulate. They will have to regulate at least 10 genes to be considered global regulators.
7. If required with `plot.more = TRUE`, the plots of the global regulators against the genes they regulate.

## 4.4 Running an example

An example of the execution of `more` function with the 'MLR' option is shown below using the data file `TestData.RData`, which is included in the package. This dataset is a reduced version of a database whose full version is available [here](#).

In this file, the `targetData` matrix corresponds to the gene expression data obtained by RNA-seq (**GeneExpressionDE**), and there is a list with four matrices of regulators in the `data.omics` object (all values are normalized):

**miRNA-seq** miRNA expression data.

**ChIP-seq** chromatin immunoprecipitation sequencing data.

**TF** TF expression data.

The experimental design matrix (**edesign**) contains the information about the experimental groups corresponding to the 16 samples available in this dataset.

We can run the following `more` code to obtain the regression models for our genes, in this case MLR model with ElasticNet variable selection:

```

> set.seed(123)
> SimMLR = more(targetData = TestData$GeneExpressionDE ,
  regulatoryData = TestData$data.omics ,
  associations = TestData$associations ,
  omicType = c(1,0,0) , condition = TestData$edesign ,
  minVariation = 0 , interactions = TRUE ,
  alfaEN = NULL , varSel = 'EN' , correlation = 0.7 ,
  method = 'MLR')

```

The following code shows the estimated coefficients of some relevant regulators in the final MLR model computed by **more** for the gene ENSMUSG00000024873:

	coefficient
(Intercept)	1.229948e+01
mmu-miR-16-1-3p	-7.696939e-04
mmu-miR-1905	3.594030e-02
mmu-miR-301a-5p	9.645046e-03
mmu-miR-1249-5p	1.596555e-03
mmu-miR-7084-3p	4.651677e-02
mmu-miR-92a-2-5p	4.347362e-02
mmu-miR-761	5.235964e-03
mmu-miR-7081-5p	-1.688525e-01
mmu-miR-1912-5p	-3.021328e-02
mmu-miR-7086-5p	6.904835e-02
mmu-miR-665-3p	-3.685801e-02
mmu-miR-191-3p	2.342803e-02
mmu-miR-301b-5p	-1.273607e-02
mmu-miR-879-5p	5.323761e-02
miRNA-seq_mc2_1_R	8.295371e-03
miRNA-seq_mc4_1_R	-9.639108e-02
miRNA-seq_mc5_1_R	3.556168e-01
Group_Disease:mmu-miR-322-5p	-2.589291e-04
Group_Disease:mmu-miR-16-1-3p	-4.848563e-05
Group_Disease:mmu-miR-1912-5p	-4.479328e-02
Group_Disease:mmu-miR-181d-3p	-5.101563e-02

The **allRegulators** table reports, for each gene, its regulators together with their associated omic, area (if provided), applied filter, and if the regulator is considered relevant or not. In this example (see the **filter** column), within the miRNA-seq omic, the regulator mmu-miR-335-3p was selected as a representative regulator (R) for being correlated to other regulators. In the filter column, the correlations are indicated using (R) for the representative, (P), and (N) for positive and negative correlation with the representative, respectively. In the same column, Model means that the regulator was included in the model by itself. Finally, the **Rel** column indicates whether a regulator is considered relevant in the final model, returning 1 for relevant regulators and 0 otherwise.

For instance, the regulator mmu-miR-186-5p is labeled as miRNA – seq\_mc5\_1\_N in the **filter** column, indicating that it is negatively correlated with its representative regulator

miRNA – seq\_mc5\_1\_R. Although mmu-miR-186-5p did not enter the regression model directly, it is still considered relevant because its representative regulator was selected as relevant, as could be seen in the latest rows.

A similar situation occurs for mmu-miR-1224-3p, whose filter entry miRNA – seq\_mc8\_P indicates a positive correlation with its corresponding representative regulator (miRNA – seq\_mc8\_R). In this case, the regulator belongs to a different correlation group (mc8) than the previous example (mc5), illustrating how distinct correlated regulator groups are handled independently.

```
> head(SimMLR$ResultsPerTargetF$ENSMUSG00000024873$allRegulators)
  targetF      regulator      omic area      filter Rel
mmu-miR-335-3p ENSMUSG00000024873 mmu-miR-335-3p miRNA-seq      miRNA-seq_mc1_R  0
mmu-miR-1912-3p ENSMUSG00000024873 mmu-miR-1912-3p miRNA-seq      Model    0
mmu-miR-615-5p ENSMUSG00000024873 mmu-miR-615-5p miRNA-seq      Model    0
mmu-miR-322-5p ENSMUSG00000024873 mmu-miR-322-5p miRNA-seq      Model    1
mmu-miR-1894-3p ENSMUSG00000024873 mmu-miR-1894-3p miRNA-seq      Model    0
mmu-miR-7082-3p ENSMUSG00000024873 mmu-miR-7082-3p miRNA-seq      Model    0
> tail(SimMLR$ResultsPerTargetF$ENSMUSG00000024873$allRegulators,15)
  targetF      regulator      omic area      filter Rel
mmu-miR-1224-3p ENSMUSG00000024873 mmu-miR-1224-3p miRNA-seq      miRNA-seq_mc8_P  0
mmu-miR-3109-3p ENSMUSG00000024873 mmu-miR-3109-3p miRNA-seq      miRNA-seq_mc7_N  0
mmu-miR-301b-5p ENSMUSG00000024873 mmu-miR-301b-5p miRNA-seq      Model    1
mmu-miR-291a-5p ENSMUSG00000024873 mmu-miR-291a-5p miRNA-seq      miRNA-seq_mc5_1_N 1
mmu-miR-879-5p ENSMUSG00000024873 mmu-miR-879-5p miRNA-seq      Model    1
mmu-miR-449a-5p ENSMUSG00000024873 mmu-miR-449a-5p miRNA-seq      miRNA-seq_mc4_1_N 1
mmu-miR-1929-3p ENSMUSG00000024873 mmu-miR-1929-3p miRNA-seq      Model    0
miRNA-seq_mc1_R ENSMUSG00000024873 miRNA-seq_mc1_R miRNA-seq      Model    0
miRNA-seq_mc2_1_R ENSMUSG00000024873 miRNA-seq_mc2_1_R miRNA-seq      Model    1
miRNA-seq_mc3_R ENSMUSG00000024873 miRNA-seq_mc3_R miRNA-seq      Model    0
miRNA-seq_mc4_1_R ENSMUSG00000024873 miRNA-seq_mc4_1_R miRNA-seq      Model    1
miRNA-seq_mc5_1_R ENSMUSG00000024873 miRNA-seq_mc5_1_R miRNA-seq      Model    1
miRNA-seq_mc6_R ENSMUSG00000024873 miRNA-seq_mc6_R miRNA-seq      Model    0
miRNA-seq_mc7_R ENSMUSG00000024873 miRNA-seq_mc7_R miRNA-seq      Model    0
miRNA-seq_mc8_R ENSMUSG00000024873 miRNA-seq_mc8_R miRNA-seq      Model    0
```

Finally, we can ask for a brief summary of the created model calling the summary function:

```
> summary(SimMLR)
A model was computed for 20 target features.
0 target features had no initial regulators.
For 1 target features, the final MLR model could not be obtained.
Target features presented a mean of 32.73684 relevant regulators.
These are the top 1 hub target features and the number of relevant regulators for each:
ENSMUSG0000049624 ENSMUSG0000012535 ENSMUSG0000038208 ENSMUSG0000021583 ENSMUSG0000016018 ENSMUSG0000041995 ENSMUSG0000050439 ENSMUSG0000033985
  91      69      68      60      55      44      28      28
ENSMUSG0000024873 ENSMUSG0000036932
  27      25
There were not global regulators (regulators that regulate more than 10 target features).
```

## 5 Retrieving significant regulations from MORE results

The function **RegulationPerCondition** is applied to the **more** output. It returns a summary table containing all the relevant/significant regulations, that is, all the pairs target feature-regulator considered relevant/significant in MORE models (depending if a MLR or a PLS model was applied). Moreover, it provides the regression coefficient that relates the target feature and the regulator for each experimental condition after testing if this coefficient is relevant/significant or not.

```
RegulationPerCondition(output, filterR2 = 0)
```

### 5.1 RegulationPerCondition input parameters

**output** Object containing the output of **more** function.

**filterR2** Filters out target features with lower  $R^2$  than the specified threshold. By default, 0 (no filter is applied).

### 5.2 Interpreting RegulationPerCondition output with an example

Following the previous example, we can run the **RegulationPerCondition** function.

```
> myresults = RegulationPerCondition(SimMLR)
```

The output is the following table, where some pairs target feature-regulator are shown:

```
> head(myresults)
   targetF      regulator      omic area representative Group_Control Group_Disease
1 ENSMUSG00000000078 mmu-miR-139-3p miRNA-seq           -0.002225 -0.002225
2 ENSMUSG00000000078          Mef2d       TF            0.000000 -0.771200
3 ENSMUSG00000000078          Rxra       TF            0.000000 -0.004460
4 ENSMUSG00000024873 mmu-miR-16-1-3p miRNA-seq           -0.001215 -0.001291
5 ENSMUSG00000024873 mmu-miR-1905 miRNA-seq            0.056730  0.056730
6 ENSMUSG00000024873 mmu-miR-301a-5p miRNA-seq            0.015220  0.015220
> tail(myresults)
   targetF      regulator      omic area representative Group_Control Group_Disease
526 ENSMUSG00000036932      Nfe2l2       TF           Nfe2l2      0.000 -0.2076
527 ENSMUSG00000036932        Rara       TF           Nfe2l2      0.000  0.2076
528 ENSMUSG00000036932       Runx1       TF           Nfe2l2      0.000  0.2076
529 ENSMUSG00000036932       Srebf2       TF           Zfp787     -0.405 -0.5161
530 ENSMUSG00000036932        Tcf3       TF           Nfe2l2      0.000  0.2076
531 ENSMUSG00000036932       Zfp787       TF           Zfp787     -0.405 -0.5161
```

This table shows all the relevant regulations for all target features. The **representative** column indicates if the regulator was chosen as the representative of a correlated group of regulators or, otherwise, which regulator was taken as the representative of the group. When no information is provided in this column, it means that the regulator was not part of a correlated group of regulators. Regulators correlated positively with the representative will have the same coefficients (same sign) as the representative, while negatively correlated regulators will have the same coefficients as the representative but with the opposite sign.

The final columns correspond to the regression coefficients of each regulator for each experimental group. In this case, the experimental design matrix (**edesign**) contained two conditions, so the column `Group_Control` corresponds to the first condition, and `Group_Disease` corresponds to the second one. These are the conclusions we can draw from the coefficients:

- If two experimental groups have the same coefficients, it means that the regulator has the same effect on the target feature in both groups. E.g. regulators *mmu-miR-139-3p*, *mmu-miR-1905* and *mmu-miR-301a-5p* of the example shown.
- If one of the coefficients is 0, it means that the regulator does not have an effect on the target feature under this experimental condition. E.g. regulators *Mef2d*, *Rxra*, *Nfe2l2*, *Rara*, *Runx1* and *Tcf3* of the example shown.
- Experimental groups with different non-zero coefficients indicate that the regulator influences the target feature in all these experimental groups, but the magnitude of the effect is not the same for all these groups. E.g., in the illustrated case, this behavior is observed for the regulators *mmu-miR-16-1-3p*, *Srebf2*, and *Zfp787*.

The last two regulators are of particular interest: *Zfp787* was identified as representative of *Srebf2*. As a result, *Srebf2* did not enter the regression model directly and instead inherits the coefficient values estimated for its representative.

### 5.3 RegulationInCondition

In case the user is interested in a particular biological condition, to summarize the information, MORE includes the function **RegulationInCondition** which takes as input the output of **RegulationPerCondition** and the condition in which they are interested. It returns a list containing the hub genes, global regulators, and regulators with their coefficients specific to that condition.

```
RegulationInCondition(output_regpcond, cond)
```

Following the previous example, we can run the **RegulationInCondition** function for example for the *Disease* condition.

```
> myres_dis = RegulationInCondition(myresults, 'Disease')
```

The results contain a compact version of **RegulationPerCondition** only with those regulators that present a relevant/significant regulation in the specified condition, the global regulators and the hub genes in that condition.

As the dataset used contains only 20 target features, no global regulator was found (difficult to regulate at least 10 of the considered target features). To see a better example about this dataset, please look at the tutorial also available on [Github](#) which uses the extended version of the dataset.

```

> myres_dis$GlobalRegulators
character(0)
> myres_dis$HubTargetF
[1] "ENSMUSG00000012535" "ENSMUSG0000038208" "ENSMUSG0000021583" "ENSMUSG0000016018" "ENSMUSG00000041995" "ENSMUSG00000050439" "ENSMUSG00000033985"
[8] "ENSMUSG0000024873" "ENSMUSG0000036932" "ENSMUSG00000058135"
> head(myres_dis$RegulationInCondition)
  targetF regulator      omic Group Disease
1 ENSMUSG0000000078 mmu-miR-139-3p mirNA-seq -0.002225
2 ENSMUSG0000000078 Mef2d       TF   -0.771200
4 ENSMUSG0000024873 mmu-miR-16-1-3p mirNA-seq -0.001291
5 ENSMUSG0000024873 mmu-miR-1905 mirNA-seq  0.056730
6 ENSMUSG0000024873 mmu-miR-301a-5p mirNA-seq  0.015220
7 ENSMUSG0000024873 mmu-miR-1249-5p mirNA-seq  0.002520

```

## 6 Plotting MORE results

The MORE package includes several plots for the interpretation of the results.

### 6.1 summaryPlot function

The function **summaryPlot** graphically represents the summary of the relationship between target features and regulators found when creating the models. It creates two types of summary plots depending on user specifications.

```
summaryPlot(output, output_regpcond, filterR2 = 0, byTargetF = TRUE)
```

#### 6.1.1 summaryPlot input parameters

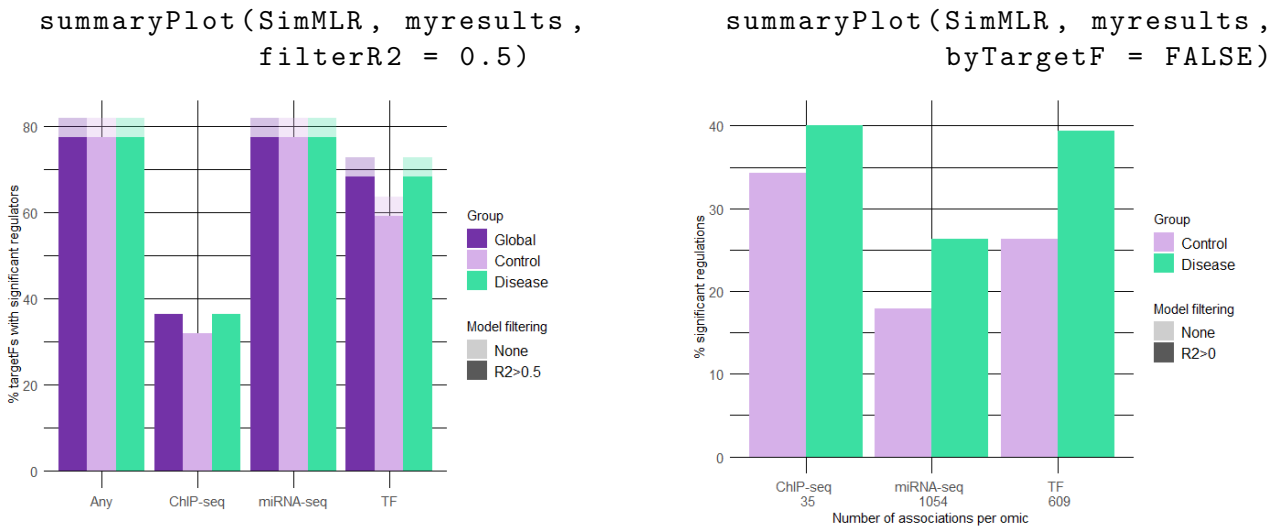
**output** Object generated by the function **more**.

**outputRegpcond** Object generated by the function **RegulationPerCondition** when applied to a **more** object.

**filterR2** Highlights the results for the genes that showed a R2 above the one indicated in this parameter. By default, 0.

**byTargetF** If TRUE (default), the function plots the percentage of target features with significant regulators globally and per omic. If FALSE, it plots the percentage of significant regulations per omic.

As an example we introduce the summary plots generated for the model previously created with **more**:



If the user needs an explanation of the presented plots, please look at the tutorial also available on [Github](#) which explains them.

## 6.2 FilterRegulationPerCondition

The MORE package includes the function **FilterRegulationPerCondition** to filter out the genes with low  $R^2$  from the **RegulationPerCondition** matrix. The function filters out genes with  $R^2$  below the one specified by the user. Thanks to **summaryPlot** the user can be aware of the percentage of genes that will be filtered out when setting a specific threshold.

```
FilterRegulationPerCondition(output, outputRegpcond, filterR2=0)
```

This function takes two arguments as input:

**output** Output object of running **more** function.

**outputRegpcond** Output object of running **RegulationPerCondition** function.

**filterR2** By default, 0. Filters out the results for the genes that showed a  $R^2$  below the one indicated.

## 6.3 differentialRegPlot function

The MORE function **differentialRegPlot** graphically visualizes the differential regulators across experimental conditions and analyzed omics. The function creates an upset plot in which the proportions of the barplots show the proportion of the identified regulators

across omic types for each combination of the experimental conditions. Numbers within bars indicate the absolute counts of regulators for each omic in each specific combination.

```
differentialRegPlot(output, outputRegPcond)
```

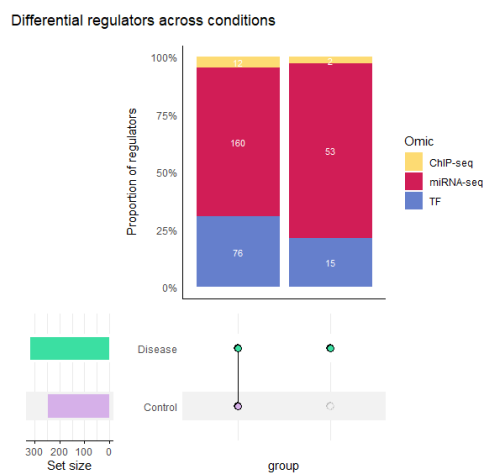
This function takes two arguments as input:

**output** Output object of running **more** function.

**outputRegPcond** Output object of running **RegulationPerCondition** function.

If we run the code for the example we were analyzing, it generates the following plot:

```
differentialRegPlot(SimMLR, myresults)
```



## 6.4 plotMORE function

The **plotMORE** function represents the relationship between the target features and regulators graphically: for a given pair target feature-regulator, to explore the regulators of a given target feature, or to analyze which target features are regulated by a specific regulator.

```
plotMORE(output, targetF, regulator = NULL, simplify = FALSE,
         reguValues = NULL, plotPerOmic = FALSE, targetF.col = 1,
         regu.col = NULL, order = TRUE, xlab = "", cont.var = NULL,
         cond2plot = NULL)
```

### 6.4.1 plotMORE input parameters

**output** Object generated by the function **more**.

**targetF** ID of the target feature to be plotted.

**regulator** ID of the regulator to be plotted. If NULL (default value), all the regulators of the target feature are plotted.

**simplify** If TRUE, a boxplot (if the regulator is binary) or a Scatterplot (otherwise) is plotted to represent the relationship between the target feature and the regulator provided to the function. If FALSE (default), the target feature and the regulator profiles will be plotted. When many samples are provided to create the models, it is hard to differentiate something in this second option.

**reguValues** Vector containing the values of a regulator that the user can optionally provide. If NULL (default value), these values are taken from **output** as long as they are available.

**plotPerOmic** If TRUE, all the relevant regulators of the given target feature and the same omic are plotted in the same graph. If FALSE (default value), each regulator is plotted in a separate plot.

**targetF.col** Color to plot the target feature. By default, biostatomics colors.

**regu.col** Color to plot the regulator. If NULL (default), a color will be assigned by the function, that will be different for each regulatory omic.

**order** If TRUE (default), the values in X-axis are ordered according to target feature expression.

**xlab** Label for the X-axis.

**cont.var** Vector with length equal to the number of observations in data, which optionally may contain the values of the numerical variable (e.g. time) to be plotted on the X-axis. By default, NULL. It plots a range for each observation in which the observation could take values taking into account the numerical variable introduced.

**cond2plot** Vector or factor indicating the experimental group of each value to represent. If NULL (default), the labels are taken from the experimental design matrix.

**smooth** If TRUE (default), smoothing is applied via splines to the actual targetF and regulator values so that profiles can be more easily seen. The smoothed profiles would be displayed in bright colors while the real values are displayed with some degradation.

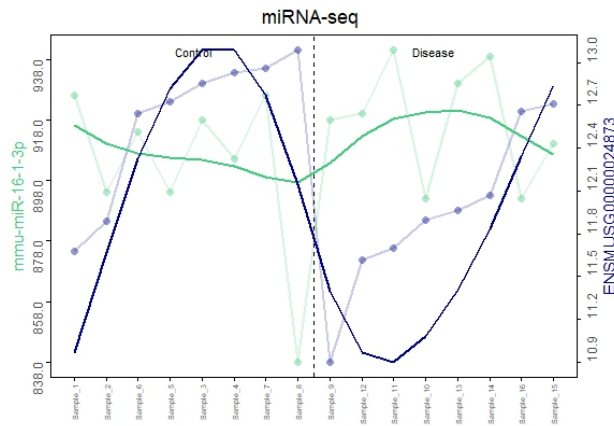
**size** Size of the X-axis labels. By default, 3.

**breakby** Range in which values should be displayed in Y-axis for regulators and target features, respectively. By default, c(0.3,0.3).

## 6.4.2 Interpretation of MORE plots

Following the previous example, the MORE graphic below represents the expression profile of a given gene (ENSMUSG00000024873) and the values for a relevant regulator of this gene (miRNA regulator, mmu-miR-16-1-3p). It can be generated with the following code:

```
> plotMORE(output = SimMLR, targetF = "ENSMUSG00000024873",
           regulator = "mmu-miR-16-1-3p", plotPerOmic = FALSE,
           targetF.col = "blue4", order = TRUE,
           regu.col = "seagreen3", size = 5, breakby = c(20,0.3))
```



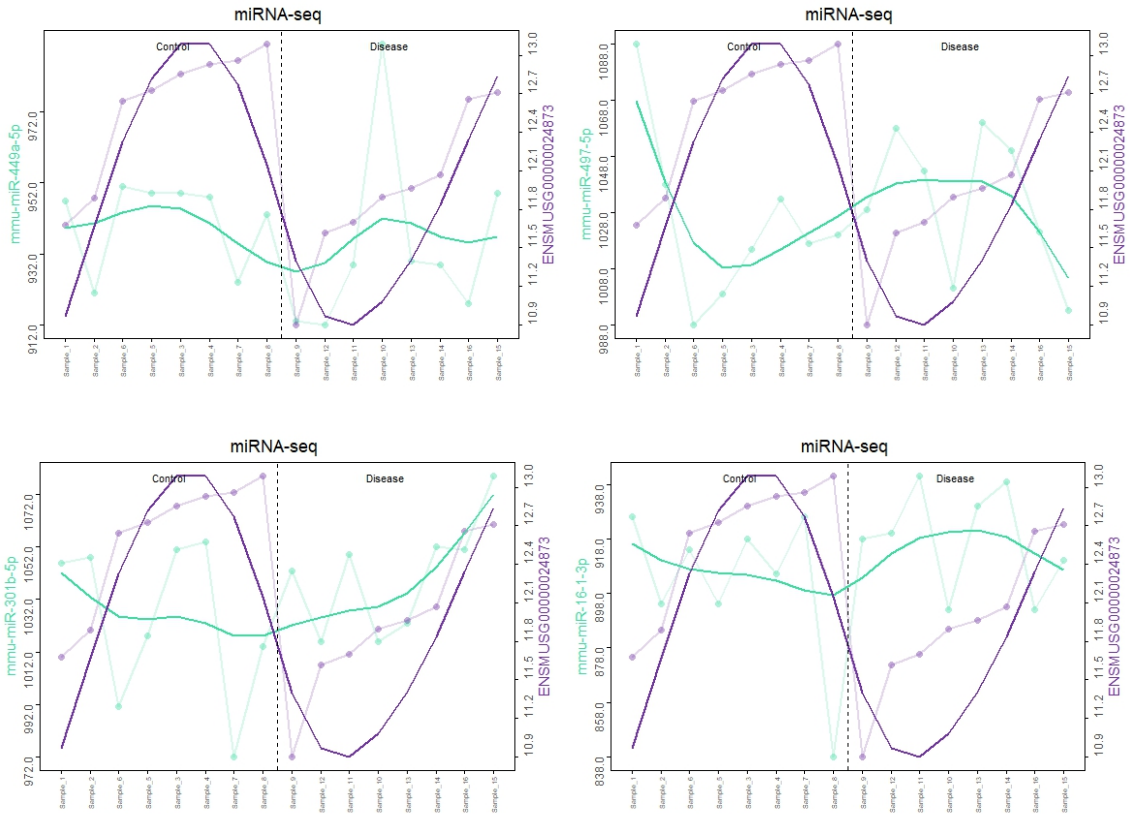
The X-axis is divided into two conditions (Control or Disease), and within each condition, the observations are displayed. The right Y-axis shows the expression values for the gene (plotted in blue), while the left Y-axis indicates the values for the regulator (plotted in green). The gradient-colored points represent the observed (raw) values of the target feature and the regulator, while the solid-colored line shows the smoothed curve fitted to these values.

If we set the regulator argument to NULL, all the relevant regulators of gene ENSMUSG-00000024873 will be plotted (27 regulators). Only 4 will be presented:

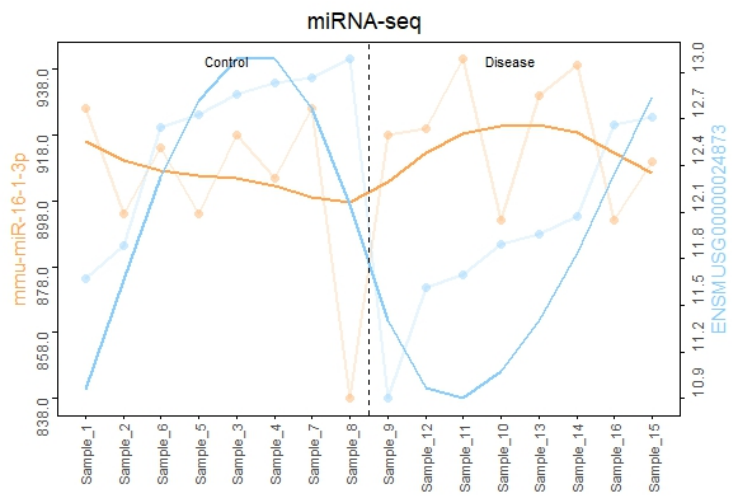
```
> plotMORE(output = SimMLR, targetF = "ENSMUSG00000024873",
           regulator = NULL, order = TRUE, plotPerOmic = FALSE,
           size = 5, breakby = c(20,0.3))
```

The title of each plot indicates the omic represented in that plot and the area. The values for relevant regulators are plotted in different colors according to the omic. The values for the gene are plotted in sea green, as indicated in the previous code.

If we want to plot all the genes that are considered to be relevantly regulated by a given regulator (e.g. mmu-miR-16-1-3p), we must set the gene argument to NULL as follows. In this case, the miRNA regulates one gene: ENSMUSG00000024873.

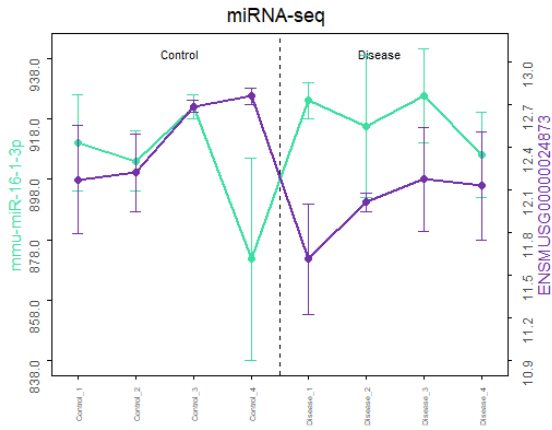


```
> plotMORE(output = SimMLR, targetF = NULL,
           regulator = "mmu-miR-16-1-3p", plotPerOmic = FALSE,
           order = FALSE, targetF.col = "skyblue1",
           regu.col = "tan1", size = 8, breakby=c(20,0.3))
```



Additionally, if the user knows that the samples belong to different time points, they can provide this information in **cont.var** vector. A 'confidence interval' will be plotted for each of the time points to which the samples belong to. The code and the resulting graph, where the gene is plotted in purple, and the regulator is plotted in green, can be found below.

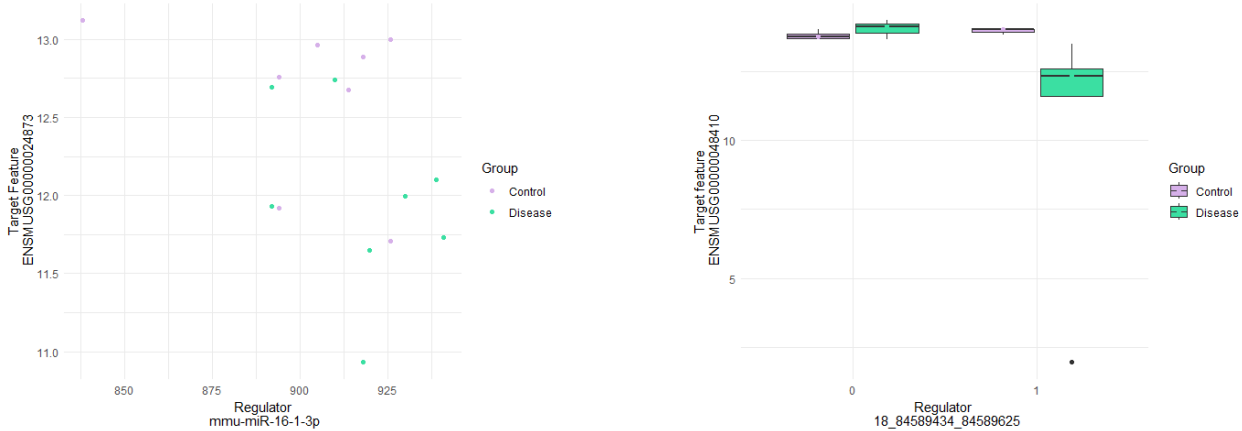
```
> plotMORE(output = SimMLR, targetF = "ENSMUSG00000024873",
           regulator = "mmu-miR-16-1-3p", plotPerOmic = FALSE,
           cont.var = c(1,2,3,4,1,2,3,4,1,2,3,4,1,2,3,4), size = 5,
           breakby=c(20,0.3))
```



In this example, we considered that there are four different time points and two different replicates for each. For that reason, in **cont.var** we specify the time point to which each of the samples belong to.

In this case, as only 16 samples are plotted the profiles can clearly be seen. However, in cases where there are many observations (let say more than 300), we encourage the user to apply the **simplify** parameter equal to TRUE to see better the relationships between a gene and a regulator. An example for binary regulator and continuous regulator are shown:

```
> plotMORE(output = SimMLR, targetF = "ENSMUSG00000024873",
           regulator = "mmu-miR-16-1-3p", simplify =TRUE)
> plotMORE(output = SimMLR, targetF = "ENSMUSG00000048410",
           regulator = "18_84589434_84589625", simplify =TRUE)
```



## 6.5 Plots for PLS models

Models created by MLR regressions in MORE apply a previous multi-collinearity filter, and users can see in the table generated by **RegulationPerCondition** the regulators that co-regulate the target feature expression. This multi-collinearity filter is not applied in PLS models as these models benefit from multi-collinearity. In order to analyze the co-regulators and the groups to which they are related, in MORE, we introduce two functions:

### 6.5.1 plotWeight function

The **plotWeight** function in MORE plots the weighting star of the regulators identified as significant in PLS models.

```
plotWeight(output, targetF, axe1 = 1, axe2 = 2)
```

**output** Object generated by the function **more**. It must be generated either by PLS1 or PLS2 models.

**targetF** ID of the target feature to be plotted.

**axe1** Component to plot in the X-axis. Default, 1.

**axe2** Component to plot in the X-axis. Default, 2.

### 6.5.2 plotScores function

The **plotScores** function in MORE plots the scores of the samples under the PLS model generated by the regulators identified as significant.

```
plotScores(output, targetF, axe1 = 1, axe2 = 2)
```

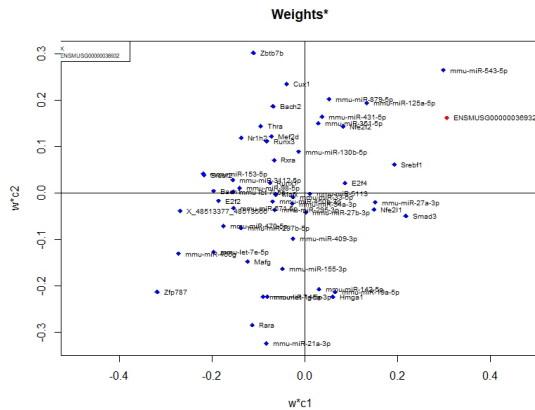
**output** Object generated by the function **more**. It must be generated either by PLS1 or PLS2 models.

**targetF** ID of the target feature to be plotted.

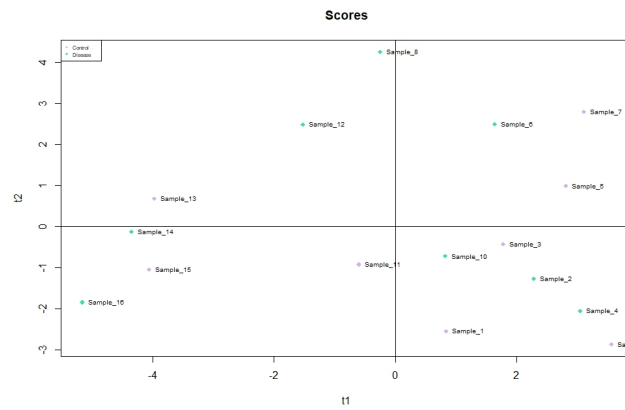
**axe1** Component to plot in the X-axis. Default, 1.

**axe2** Component to plot in the X-axis. Default, 2.

```
plotWeight(SimPLS ,
targetF= "ENSMUSG00000036932")
```



```
plotScores(SimPLS ,
targetF= "ENSMUSG00000036932")
```



If the user needs an explanation of the presented plots, please look at the tutorial also available on [Github](#) which explains them.

To generate the examples below, we run a PLS model with the PLS1 method and Jack-Knife variable selection using the rest of the default parameters of MORE for PLS models and the same input data we used to create the SimMLR model.

## 6.6 networkMORE function

MORE package includes the function **networkMORE** to graphically represent the regulatory networks obtained with the **more** function.

```
networkMORE(outputRegpcond , cytoscape = TRUE , group1 = NULL ,
group2 = NULL , pc = 0 , pathway = NULL ,
annotation = NULL , save = FALSE)
```

### 6.6.1 networkMORE input parameters

**outputRegpcond** Object generated by the application of **RegulationPerCondition** function to a **more** object.

**cytoscape** If TRUE (default), the function plots the network in *Cytoscape*. For that, it is necessary to install *RCy3* package and to maintain *Cytoscape* software opened

while running the function. If FALSE, the network is plotted in R using *igraph* package and saves it so that the user can introduce it later to *Cytoscape*. This option is not recommended for plotting huge networks, as the visualization is complex.

**group1** Name of the group to take as a reference in the creation of differential networks. If it is not provided, the networks of all groups will be plotted. If it is provided without providing *group2* only the specific network for this group would be created. By default, NULL.

**group2** Name of the group to compare to the reference in the differential network creation. Differential networks are computed by subtracting the regulator coefficients obtained in this condition from those of the reference condition. If it is not provided, the networks of all groups will be plotted. By default, NULL.

**pc** Value between 0 and 1 for the proportion of significant/relevant regulators to be plotted in the network. When having networks with many nodes, users can decide to only plot the regulators with the highest coefficients in the models (in absolute value). By default, 0, which means that all significant/relevant regulators will be plotted.

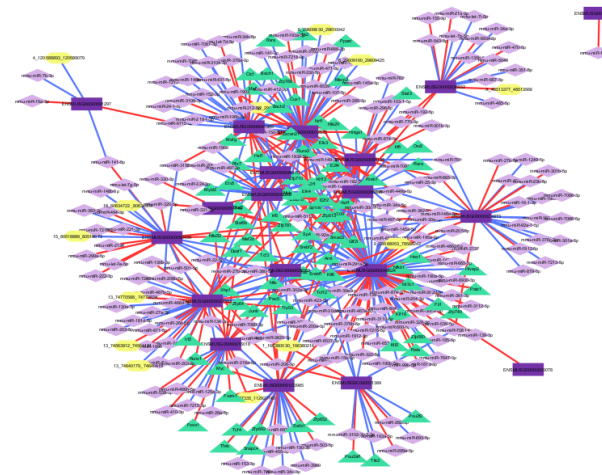
**pathway** If provided, the function will print the regulatory network only for the target features involved in the specified pathway instead of the entire regulatory network. The provided pathway name must have the same terminology as in the annotation matrix. By default, NULL.

**annotation** Annotation matrix with target features in the first column, pathway ID in the second and pathway name in the third. Only necessary when a specific pathway has to be plotted. By default, NULL.

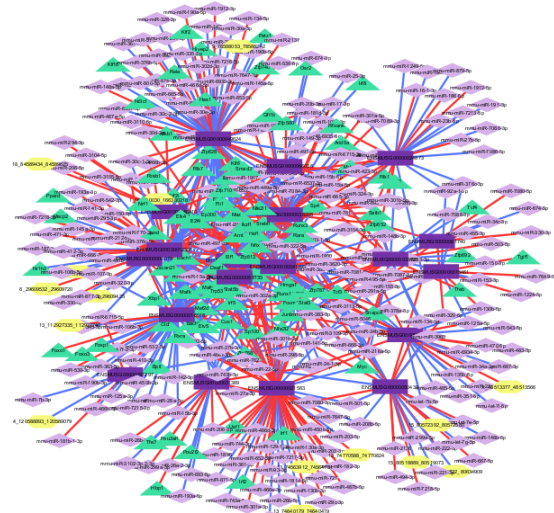
**save** If TRUE, only when *cytoscape* is set to FALSE, networks are saved in *gml* extension. By default, FALSE.

Following previous example, networks can be plotted by:

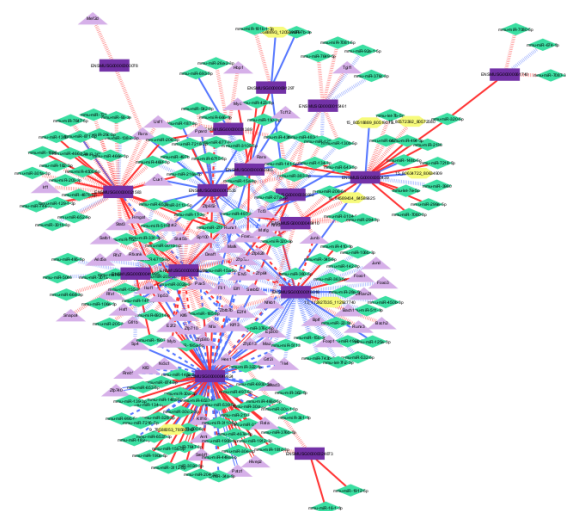
```
networkMORE(myresults, cytoscape = TRUE)
networkMORE(myresults, cytoscape = TRUE, group1 = 'Control',
            group2 = 'Disease')
```



Regulatory network in Control



Regulatory network in Disease



Differential regulatory network  
Disease vs. Control

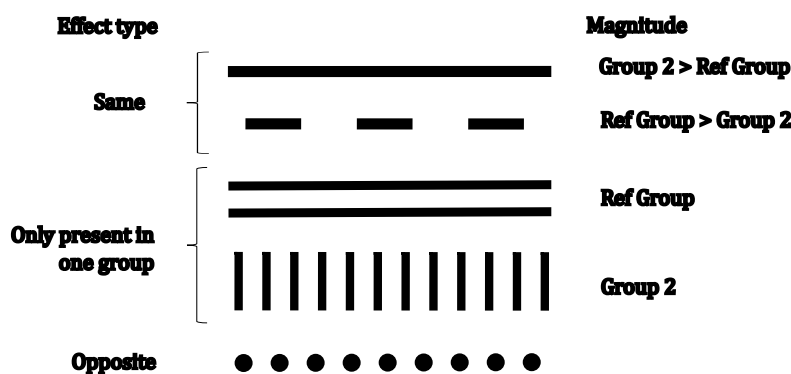
**NOTE 1:** By default, regulators repressing the target feature expression will be connected by red edges and regulators activating the target feature expression will be connected by blue edges.

**NOTE 2:** By default, response variable (Gene Expression, protein expression...), transcription factors (TFs) and microRNAs (miRNAs) will be plotted with rectangle, triangle and diamond shapes, respectively. Considering there is still no consensus in the literature on the shapes of all omics, we applied the ones by default to the rest of the omics. Users can modify these default options.

**NOTE 3:** Considering that we want to compare two groups, group1 (reference) and group2, in the differential network creation the different line styles will represent different scenarios.

- When both groups present same type of effect (activator or repressor):
  - When effect of reference is bigger than compared group, dashed lines will be used.
  - When the compared group presents bigger effect than reference, solid lines will be used.
- When one of the groups does not present a regulation but the other does:
  - When reference group is the only one which shows a regulation it will be represented by double parallel lines.
  - When the compared group is the only one which shows a regulation it will be represented by vertical lines.
- When compared groups present opposite type of effect, dotted lines will be used to represent it.

We summarize in the figure below the line types used in MORE differential networks depending on the effect types of the groups and the magnitude of them.



**NOTE 4:** When creating large networks, the R–Cytoscape connection may take a noticeable amount of time, as several layers and visual properties are applied during the connection process. In some cases, this step may take from several seconds up to a few minutes. Additionally, for networks with a very large number of nodes, the graphical rendering itself may also require several minutes. If no network visualization appears within approximately two/three minutes, we recommend stopping the connection and instead saving the networks to disk and loading them directly into Cytoscape. This approach allows users to define and customize their own network styles more efficiently. If the user wishes to apply MORE’s default style, we recommend first generating a small network induced by a subset of the **RegulationPerCondition** output and running **networkMORE** for a single condition. This will load MORE’s default style into Cytoscape, which will then be automatically applied to subsequently loaded networks.

## 7 Functional Enrichment analysis

Apart from creating networks, the MORE package also includes functionalities to perform different types of functional enrichment analyses.

### 7.1 Over Representation Analysis

The Over Representation Analysis (ORA) methodology determines if a particular biological function or pathway is significantly enriched in a target set of genes compared to a reference set. In MORE, ORA is implemented through the **oraMORE** function, which allows enrichment analysis on different biologically meaningful target set of genes derived from MORE models. In MORE, we can choose three options for the target set of genes: (1) the hub target features, (2) the target features regulated by a given global regulator in a specific condition, or (3) the target features regulated by regulators of a specific omic type. In the second option, MORE will compute as many ORAs as global regulators.

```
oraMORE(output, outputRegincond, byHubs = TRUE, byOmic = NULL,  
annotation, alpha = 0.05, p.adjust.method = "fdr")
```

#### 7.1.1 oraMORE input parameters

**output** Object generated by the function **more**.

**outputRegincond** Object generated by the function **RegulationInCondition**.

**byHubs** Indicates whether to perform the ORA for the Hub target features, TRUE, or for the target features regulated by the global regulators, FALSE. By default, TRUE.

**byOmic** If provided, it performs the ORA to the regulators of the specified omic (it must

follow the same nomenclature that in regulatoryData). Incompatible with other methodologies specified in byHubs parameter. By default, NULL.

**annotation** Annotation matrix with target features in the first column, pathway ID in the second and pathway name in the third. Only necessary when a specific pathway has to be plotted. By default, NULL.

**alpha** p-value cutoff to consider. By default, 0.05.

**p.adjust.method** p-value adjustment method to consider. By default, 'fdr'.

### 7.1.2 ORA use cases in MORE

As introduced before, the **oraMORE** function can be applied to different target gene sets derived from MORE models, depending on the biological question of interest. The following sections describe each use case and the corresponding parameter settings.

#### ORA on hub target features

```
oraMORE(output, outputRegincond, byHubs = TRUE, byOmic = NULL,  
annotation, alpha = 0.05, p.adjust.method = "fdr")
```

#### ORA on target features regulated by a given global regulator

```
oraMORE(output, outputRegincond, byHubs = FALSE, byOmic = NULL,  
annotation, alpha = 0.05, p.adjust.method = "fdr")
```

#### ORA on target features regulated by regulators of a specific omic type

```
oraMORE(output, outputRegincond, byHubs = FALSE, byOmic = 'X',  
annotation, alpha = 0.05, p.adjust.method = "fdr")
```

where 'X' indicates the omic of interest, which must be specified using the same name as in the **regulatoryData** list.

Following previous example, ORA on hub target features can be performed by:

```
load('annotation.Rdata')  
oraMORE(SimMLR, myres_dis, annotation)
```

To run this example an annotation matrix is loaded, but this annotation matrix is not incorporated in MORE, is up to the user to load their own annotation matrix.

## 7.2 Gene Set Enrichment Analysis

Gene Set Enrichment Analysis (GSEA) evaluates whether a set of genes shows statistically significant differences between two experimental conditions. In MORE we center the analysis of the differences in the number of regulators of the genes in the experimental

conditions. **gseaMORE** function performs GSEA analysis to MORE output making use of *clusterProfiler* package, so that all plots included in that package could be applied to the output object of running that function.

```
gseaMORE(outputRegincond, outputRegincond2 = NULL, annotation,
alpha = 0.05, p.adjust.method = "fdr")
```

### 7.2.1 gseaMORE input parameters

**outputRegincond** Object generated by the function **RegulationInCondition**.

**outputRegincond2** Object generated by the function **RegulationInCondition** for another group different from the previously considered. By default, NULL. If NULL, the analysis will be centered only in the study group, so the results will apply only to that group and will not mean the statistical differences between groups. If provided **MORE** computes and internal score comparing the number of significant regulators in both groups.

**annotation** Annotation matrix with target features in the first column, pathway ID in the second and pathway name in the third. Only necessary when a specific pathway has to be plotted. By default, NULL.

**alpha** p-value cutoff to consider. By default, 0.05.

**p.adjust.method** p-value adjustment method to consider. By default, 'fdr'.

Following previous example, GSEA can be performed by:

```
myres_cont = RegulationInCondition(myresults, 'Control')
gseaMORE(myres_cont, myres_dis, annotation,
alpha = 0.05, p.adjust.method = "fdr")
```

**NOTE 5:** In the case where no condition is provided, if the user wants to use functions that require **RegulationInCondition** as input, instead, it must provide the output of **RegulationPerCondition**.

## 8 How to cite MORE package

Aguerralde-Martin M, Clemente-Císcar M, Conesa A, Tarazona S. MORE interpretable multi-omic regulatory networks to characterise phenotypes. Brief Bioinform. 2025; 26(3): bbaf270. doi:10.1093/bib/bbaf270.

Aguerralde-Martin, Maider; Clemente-Císcar, Mónica; Conesa, Ana; Tarazona, Sonia. (2023). MORE: Multi-Omics REgulation. R package version 1.0.