# Package 'MOFA2'

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Type Package

Title Multi-Omics Factor Analysis v2

Version 1.5.2

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**Description** The MOFA2 package contains a collection of tools for training and analysing multiomic factor analysis (MOFA). MOFA is a probabilistic factor model that aims to identify principal axes of variation from data sets that can comprise multiple omic layers and/or groups of samples. Additional time or space information on the samples can be incorporated using the MEFISTO framework, which is part of MOFA2. Downstream analysis functions to inspect molecular features underlying each factor, vizualisation, imputation etc are available.

**Encoding** UTF-8

**Depends** R (>= 4.0)

Imports rhdf5, dplyr, tidyr, reshape2, pheatmap, ggplot2, methods, RColorBrewer, cowplot, ggrepel, reticulate, HDF5Array, grDevices, stats, magrittr, forcats, utils, corrplot, DelayedArray, Rtsne, uwot, basilisk, stringi

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BugReports https://github.com/bioFAM/MOFA2

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R topics documented:

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```

Function to add the MOFA representation onto a Seurat object

### **Description**

Function to add the MOFA latent representation to a Seurat object

# Usage

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```
add_mofa_factors_to_seurat(
  mofa_object,
  seurat_object,
  views = "all",
  factors = "all"
)
```

#### **Arguments**

mofa\_object a trained MOFA object. seurat\_object a Seurat object

views character vector with the view names, or numeric vector with view indexes.

Default is 'all'

factors character vector with the factor names, or numeric vector with the factor indexes.

Default is 'all'

#### **Details**

This function calls the CreateDimReducObject function from Seurat to store the MOFA factors.

#### Value

Returns a Seurat object with the 'reductions' slot filled with the MOFA factors. Also adds, if calculated, the UMAP/TSNE obtained with the MOFA factors.

```
# Generate a simulated data set
MOFAexample <- make_example_data()</pre>
```

calculate\_contribution\_scores

Calculate contribution scores for each view in each sample

### **Description**

This function calculates, \*for each sample\* how much each view contributes to its location in the latent manifold, what we call *contribution scores* 

#### Usage

```
calculate_contribution_scores(
  object,
  views = "all",
  groups = "all",
  factors = "all",
  scale = TRUE
)
```

### **Arguments**

object	a trained MOFA object.
views	character vector with the view names, or numeric vector with view indexes. Default is 'all'
groups	character vector with the group names, or numeric vector with group indexes. Default is 'all'
factors	character vector with the factor names, or numeric vector with the factor indexes. Default is 'all'
scale	logical indicating whether to scale the sample-wise variance explained values by the total amount of variance explained per view. This effectively normalises each view by its total variance explained. It is important when different amounts of variance is explained for each view (check with plot_variance_explained(, plot_total=TRUE))

#### **Details**

Contribution scores are calculated in three steps:

- Step 1 calculate variance explained for each cell i and each view m  $(R_{im})$ , using all factors
- Step 2(optional) scale values by the total variance explained for each view
- Step 3calculate contribution score  $(C_{im})$  for cell i and view m as:

$$C_{im} = \frac{R2_{im}}{\sum_{m} R2_{im}}$$

Note that contribution scores can be calculated using any number of data modalities, but it is easier to interpret when you specify two.

Please note that this functionality is still experimental, contact the authors if you have questions.

#### Value

adds the contribution scores to the metadata slot (samples\_metadata(MOFAobject)) and to the MOFAobject@cache slot

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
model <- calculate_contribution_scores(model)</pre>
```

calculate\_variance\_explained

Calculate variance explained by the model

# Description

This function takes a trained MOFA model as input and calculates the proportion of variance explained (i.e. the coefficient of determinations (R^2)) by the MOFA factors across the different views.

#### Usage

```
calculate_variance_explained(
  object,
  views = "all",
  groups = "all",
  factors = "all"
)
```

### **Arguments**

object a MOFA object.

views character vector with the view names, or numeric vector with view indexes.

Default is 'all'

groups character vector with the group names, or numeric vector with group indexes.

Default is 'all'

factors character vector with the factor names, or numeric vector with the factor indexes.

Default is 'all'

#### Value

a list with matrices with the amount of variation explained per factor and view.

#### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Calculate variance explained (R2)
r2 <- calculate_variance_explained(model)

# Plot variance explained values (view as x-axis, and factor as y-axis)
plot_variance_explained(model, x="view", y="factor")

# Plot variance explained values (view as x-axis, and group as y-axis)
plot_variance_explained(model, x="view", y="group")

# Plot variance explained values for factors 1 to 3
plot_variance_explained(model, x="view", y="group", factors=1:3)

# Scale R2 values
plot_variance_explained(model, max_r2 = 0.25)</pre>
```

calculate\_variance\_explained\_per\_sample

Calculate variance explained by the MOFA factors for each sample

# Description

This function takes a trained MOFA model as input and calculates, \*\*for each sample\*\* the proportion of variance explained (i.e. the coefficient of determinations (R^2)) by the MOFA factors across the different views.

#### Usage

```
calculate_variance_explained_per_sample(
  object,
  views = "all",
  groups = "all",
  factors = "all"
)
```

# Arguments

object	a MOFA object.
views	character vector with the view names, or numeric vector with view indexes. Default is 'all'
groups	character vector with the group names, or numeric vector with group indexes. Default is 'all'
factors	character vector with the factor names, or numeric vector with the factor indexes.

Default is 'all'

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#### Value

a list with matrices with the amount of variation explained per sample and view.

#### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Calculate variance explained (R2)
r2 <- calculate_variance_explained_per_sample(model)</pre>
```

cluster\_samples

K-means clustering on samples based on latent factors

#### **Description**

MOFA factors are continuous in nature but they can be used to predict discrete clusters of samples. The clustering can be performed in a single factor, which is equivalent to setting a manual threshold. More interestingly, it can be done using multiple factors, where multiple sources of variation are aggregated.

Importantly, this type of clustering is not weighted and does not take into account the different importance of the latent factors.

### Usage

```
cluster_samples(object, k, factors = "all", ...)
```

### **Arguments**

object a trained MOFA object.

k number of clusters (integer).

factors character vector with the factor name(s), or numeric vector with the index of the

factor(s) to use. Default is 'all'

... extra arguments passed to kmeans

#### **Details**

In some cases, due to model technicalities, samples can have missing values in the latent factor space. In such a case, these samples are currently ignored in the clustering procedure.

#### Value

output from kmeans function

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#### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Cluster samples in the factor space using factors 1 to 3 and K=2 clusters
clusters <- cluster_samples(model, k=2, factors=1:3)</pre>
```

compare\_elbo

Compare different trained MOFA objects in terms of the final value of the ELBO statistics and number of inferred factors

#### Description

Different objects of MOFA are compared in terms of the final value of the ELBO statistics. For model selection the model with the highest ELBO value is selected.

#### Usage

```
compare_elbo(models, log = FALSE, return_data = FALSE)
```

#### **Arguments**

models a list containing MOFA objects.

logical indicating whether to plot the log of the ELBO.

return\_data logical indicating whether to return a data.frame with the ELBO values per

model

#### Value

A ggplot object or the underlying data.frame if return\_data is TRUE

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model1 <- load_model(file)
model2 <- load_model(file)

# Compare ELBO between models
## Not run: compare_elbo(list(model1,model2))</pre>
```

compare\_factors

Plot the correlation of factors between different models

#### **Description**

Different MOFA objects are compared in terms of correlation between their factors.

#### Usage

```
compare_factors(models, ...)
```

#### **Arguments**

```
models a list with MOFA objects.
... extra arguments passed to pheatmap
```

#### **Details**

If assessing model robustness across trials, the output should look like a block diagonal matrix, suggesting that all factors are robustly detected in all model instances.

#### Value

Plots a heatmap of the Pearson correlation between latent factors across all input models.

# Examples

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model1 <- load_model(file)
model2 <- load_model(file)

# Compare factors between models
compare_factors(list(model1,model2))</pre>
```

```
correlate\_factors\_with\_covariates
```

Plot correlation of factors with external covariates

#### Description

Function to correlate factor values with external covariates.

# Usage

```
correlate_factors_with_covariates(
  object,
  covariates,
  factors = "all",
  groups = "all",
  abs = FALSE,
  plot = c("log_pval", "r"),
  alpha = 0.05,
  return_data = FALSE,
  transpose = FALSE,
  ...
)
```

# Arguments

object	a trained MOFA object.
covariates	<ul> <li>data.frame:a data.frame where the samples are stored in the rows and the covariates are stored in the columns. Use row names for sample names and column names for covariate names. Columns values must be numeric.</li> </ul>
	<ul> <li>character vector:character vector with names of columns that are present in the sample metadata (samples_metadata(model)</li> </ul>
factors	character vector with the factor name(s), or numeric vector with the index of the factor(s) to use. Default is 'all'.
groups	character vector with the groups names, or numeric vector with the indices of the groups of samples to use, or "all" to use samples from all groups.
abs	logical indicating whether to take the absolute value of the correlation coefficient (default is TRUE).
plot	character indicating whether to plot Pearson correlation coefficiens (plot="r") or log10 adjusted p-values (plot="log_pval").
alpha	p-value threshold
return_data	logical indicating whether to return the correlation results instead of plotting
transpose	logical indicating whether to transpose the plot
	extra arguments passed to corrplot (if plot=="r") or pheatmap (if plot=="log_pval").

# Value

A corrplot (if plot=="r") or  $plot=="log_pval"$ ) or the underlying data.frame if  $return\_data$  is TRUE

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covariates\_names

covariates\_names: set and retrieve covariate names

# **Description**

covariates\_names: set and retrieve covariate names

#### Usage

```
covariates_names(object)

covariates_names(object) <- value

## S4 method for signature 'MOFA'
covariates_names(object)

## S4 replacement method for signature 'MOFA, vector'
covariates_names(object) <- value</pre>
```

# Arguments

object a MOFA object.

value a character vector of covariate names

#### Value

character vector with the covariate names

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
covariates_names(model)</pre>
```

create\_mofa

create a MOFA object

# Description

Method to create a MOFA object. Depending on the input data format, this method calls one of the following functions:

```
• long data.frame: create_mofa_from_df
```

• List of matrices: create\_mofa\_from\_matrix

create\_mofa\_from\_df

- MultiAssayExperiment: create\_mofa\_from\_MultiAssayExperiment
- Seurat: create\_mofa\_from\_Seurat
- SingleCellExperiment: create\_mofa\_from\_SingleCellExperiment

Please read the documentation of the corresponding function for more details on your specific data format.

#### Usage

```
create_mofa(data, groups = NULL, extract_metadata = TRUE, ...)
```

#### **Arguments**

data one of the formats above

groups group information, only relevant when using the multi-group framework.

extract\_metadata

logical indicating whether to incorporate the sample metadata from the input object into the MOFA object (not relevant when the input is a list of matrices).

Default is TRUE.

further arguments that can be passed to the function depending on the inout data

format. See the dpcumentation of above functions for details.

#### Value

Returns an untrained MOFA object

### **Examples**

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")
# Load data (in long data.frame format)
load(file)
MOFAmodel <- create_mofa(dt)</pre>
```

create\_mofa\_from\_df

create a MOFA object from a data.frame object

#### **Description**

Method to create a MOFA object from a data.frame object

### Usage

```
create_mofa_from_df(df, extract_metadata = TRUE)
```

#### **Arguments**

df

data.frame object with at most 5 columns: sample, group, feature, view, value. The group column (optional) indicates the group of each sample when using the multi-group framework. The view column (optional) indicates the view of each feature when having multi-view data.

extract\_metadata

logical indicating whether to incorporate the extra columns as sample metadata into the MOFA object

#### Value

Returns an untrained MOFA object

#### **Examples**

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")
# Load data (in long data.frame format)
load(file)
MOFAmodel <- create_mofa_from_df(dt)</pre>
```

create\_mofa\_from\_matrix

create a MOFA object from a a list of matrices

### **Description**

Method to create a MOFA object from a list of matrices

#### Usage

```
create_mofa_from_matrix(data, groups = NULL)
```

#### **Arguments**

data A list of matrices, where each entry corresponds to one view. Samples are stored

in columns and features in rows. Missing values must be filled in prior to creat-

ing the MOFA object (see for example the CLL tutorial)

groups A character vector with group assignment for every sample. Default is NULL, no

group structure.

#### Value

Returns an untrained MOFA object

#### **Examples**

```
m <- make_example_data()
create_mofa_from_matrix(m$data)</pre>
```

### **Description**

Method to create a MOFA object from a MultiAssayExperiment object

#### Usage

```
create_mofa_from_MultiAssayExperiment(
  mae,
  groups = NULL,
  extract_metadata = FALSE
)
```

#### **Arguments**

mae a MultiAssayExperiment object

groups a string specifying column name of the colData to use it as a group variable. Al-

ternatively, a character vector with group assignment for every sample. Default

is NULL (no group structure).

extract\_metadata

logical indicating whether to incorporate the metadata from the MultiAssayEx-

periment object into the MOFA object

#### Value

Returns an untrained MOFA object

### **Description**

Method to create a MOFA object from a Seurat object

### Usage

```
create_mofa_from_Seurat(
    seurat,
    groups = NULL,
    assays = NULL,
    slot = "scale.data",
    features = NULL,
    extract_metadata = FALSE
)
```

# Arguments

seurat	Seurat object
groups	a string specifying column name of the samples metadata to use it as a group variable. Alternatively, a character vector with group assignment for every sample. Default is NULL (no group structure).
assays	assays to use, default is NULL, it fetched all assays available
slot	assay slot to be used (default is scale.data).
features	a list with vectors, which are used to subset features, with names corresponding to assays; a vector can be provided when only one assay is used
extract_metada	ta

logical indicating whether to incorporate the metadata from the Seurat object into the MOFA object

# Value

Returns an untrained MOFA object

```
create\_mofa\_from\_SingleCellExperiment\\ create\ a\ MOFA\ object\ from\ a\ SingleCellExperiment\ object
```

# Description

Method to create a MOFA object from a SingleCellExperiment object

### Usage

```
create_mofa_from_SingleCellExperiment(
    sce,
    groups = NULL,
    assay = "logcounts",
    extract_metadata = FALSE
)
```

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#### **Arguments**

sce SingleCellExperiment object

groups a string specifying column name of the colData to use it as a group variable. Al-

ternatively, a character vector with group assignment for every sample. Default

is NULL (no group structure).

assay assay to use, default is logcounts.

extract\_metadata

logical indicating whether to incorporate the metadata from the SingleCellEx-

periment object into the MOFA object

#### Value

Returns an untrained MOFA object

factors\_names

factors\_names: set and retrieve factor names

#### **Description**

factors\_names: set and retrieve factor names

### Usage

```
factors_names(object)
factors_names(object) <- value

## S4 method for signature 'MOFA'
factors_names(object)

## S4 replacement method for signature 'MOFA, vector'
factors_names(object) <- value</pre>
```

#### Arguments

object a MOFA object.

value a character vector of factor names

### Value

character vector with the factor names

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
factors_names(model)</pre>
```

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features\_metadata

features\_metadata: set and retrieve feature metadata

### **Description**

features\_metadata: set and retrieve feature metadata

#### Usage

```
features_metadata(object)
features_metadata(object) <- value

## S4 method for signature 'MOFA'
features_metadata(object)

## S4 replacement method for signature 'MOFA,data.frame'
features_metadata(object) <- value</pre>
```

# Arguments

object a MOFA object.

value data frame with feature information, it at least must contain the columns feature

and view

#### Value

a data frame with sample metadata

#### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
features_metadata(model)</pre>
```

features\_names

features\_names: set and retrieve feature names

#### **Description**

features\_names: set and retrieve feature names

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#### Usage

```
features_names(object)

features_names(object) <- value

## S4 method for signature 'MOFA'
features_names(object)

## S4 replacement method for signature 'MOFA,list'
features_names(object) <- value</pre>
```

### **Arguments**

object a MOFA object.

value list of character vectors with the feature names for every view

### Value

list of character vectors with the feature names for each view

#### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
features_names(model)</pre>
```

get\_covariates

Get sample covariates

# Description

Function to extract the covariates from a MOFA object using MEFISTO.

### Usage

```
get_covariates(
  object,
  covariates = "all",
  as.data.frame = FALSE,
  warped = FALSE
)
```

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### Arguments

object a MOFA object.

covariates character vector with the covariate name(s), or numeric vector with the covariate

index(es).

as.data.frame logical indicating whether to output the result as a long data frame, default is

FALSE.

warped logical indicating whether to extract the aligned covariates

#### Value

a matrix with dimensions (samples,covariates). If as.data.frame is TRUE, a long-formatted data frame with columns (sample,factor,value)

# **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
covariates <- get_covariates(model)</pre>
```

get\_data

Get data

# Description

Fetch the input data

### Usage

```
get_data(
  object,
  views = "all",
  groups = "all",
  features = "all",
  as.data.frame = FALSE,
  add_intercept = TRUE,
  denoise = FALSE,
  na.rm = TRUE
)
```

### **Arguments**

object a MOFA object.

views character vector with the view name(s), or numeric vector with the view in-

dex(es). Default is "all".

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groups	character vector with the group name(s), or numeric vector with the group index(es). Default is "all".
features	a *named* list of character vectors. Example: list("view1"=c("feature_1","feature_2"),
as.data.frame	logical indicating whether to return a long data frame instead of a list of matrices. Default is FALSE.
add_intercept	logical indicating whether to add feature intercepts to the data. Default is TRUE.
denoise	logical indicating whether to return the denoised data (i.e. the model predictions). Default is FALSE.
na.rm	remove NAs from the data.frame (only if as.data.frame is TRUE).

#### **Details**

By default this function returns a list where each element is a data matrix with dimensionality (D,N) where D is the number of features and N is the number of samples.

Alternatively, if as.data.frame is TRUE, the function returns a long-formatted data frame with columns (view,feature,sample,value). Missing values are not included in the the long data.frame format by default. To include them use the argument na.rm=FALSE.

#### Value

A list of data matrices with dimensionality (D,N) or a data.frame (if as.data.frame is TRUE)

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Fetch data
data <- get_data(model)

# Fetch a specific view
data <- get_data(model, views = "view_0")

# Fetch data in data.frame format instead of matrix format
data <- get_data(model, as.data.frame = TRUE)

# Fetch centered data (do not add the feature intercepts)
data <- get_data(model, as.data.frame = FALSE)

# Fetch denoised data (do not add the feature intercepts)
data <- get_data(model, denoise = TRUE)</pre>
```

#### Description

Function to obtain the default data options.

#### Usage

```
get_default_data_options(object)
```

#### **Arguments**

object

an untrained MOFA object

#### **Details**

This function provides a default set of data options that can be modified and passed to the MOFA object in the prepare\_mofa step (see example), i.e. after creating a MOFA object (using create\_mofa) and before starting the training (using run\_mofa) The data options are the following:

- scale\_views: logical indicating whether to scale views to have the same unit variance. As long as the scale differences between the views is not too high, this is not required. Default is FALSE.
- scale\_groups: logical indicating whether to scale groups to have the same unit variance. As long as the scale differences between the groups is not too high, this is not required. Default is FALSE.
- **use\_float32**: logical indicating whether use float32 instead of float64 arrays to increase speed and memory usage. Default is FALSE.

#### Value

Returns a list with the default data options.

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")
# Load data dt (in data.frame format)
load(file)
# Create the MOFA object
MOFAmodel <- create_mofa(dt)
# Load default data options</pre>
```

```
data_opts <- get_default_data_options(MOFAmodel)

# Edit some of the data options
data_opts$scale_views <- TRUE

# Prepare the MOFA object
MOFAmodel <- prepare_mofa(MOFAmodel, data_options = data_opts)</pre>
```

```
get_default_mefisto_options
```

Get default options for MEFISTO covariates

#### **Description**

Function to obtain the default options for the usage of MEFISTO covariates with MEFISTO

#### Usage

```
get_default_mefisto_options(object)
```

### **Arguments**

object

an untrained MOFA object

#### Details

The options are the following:

- scale\_cov: logical: Scale covariates?
- start\_opt: integer: First iteration to start the optimisation of GP hyperparameters
- **n\_grid**: integer: Number of points for the grid search in the optimisation of GP hyperparameters
- opt\_freq: integer: Frequency of optimisation of GP hyperparameters
- sparseGP: logical: Use sparse GPs to speed up the optimisation of the GP parameters?
- **frac\_inducing**: numeric between 0 and 1: Fraction of samples to use as inducing points (only relevant if sparseGP is TRUE)
- warping: logical: Activate warping functionality to align covariates between groups (requires a multi-group design)
- warping\_freq: numeric: frequency of the warping (only relevant if warping is TRUE)
- warping\_ref: A character specifying the reference group for warping (only relevant if warping is TRUE)
- warping\_open\_begin: logical: Warping: Allow for open beginning? (only relevant warping is TRUE)
- warping\_open\_end: logical: Warping: Allow for open end? (only relevant warping is TRUE)

- warping\_groups: Assignment of groups to classes used for alignment (advanced option). Needs to be a vector of length number of samples, e.g. a column of samples\_metadata, which needs to have the same value within each group. By default groups are used specified in 'create\_mofa'.
- model\_groups: logical: Model covariance structure across groups (for more than one group, otherwise FALSE)? If FALSE, we assume the same patterns in all groups.
- new\_values: Values for which to predict the factor values (for interpolation / extrapolation). This should be numeric matrix in the same format with covariate(s) in rows and new values in columns. Default is NULL, leading to no interpolation.

#### Value

Returns a list with default options for the MEFISTO covariate(s) functionality.

#### **Examples**

```
# generate example data
dd <- make_example_data(sample_cov = seq(0,1,length.out = 200), n_samples = 200,
n_factors = 4, n_features = 200, n_views = 4, lscales = c(0.5, 0.2, 0, 0))
# input data
data <- dd$data
# covariate matrix with samples in columns
time <- dd$sample_cov
rownames(time) <- "time"

# create mofa and set covariates
sm <- create_mofa(data = dd$data)
sm <- set_covariates(sm, covariates = time)

MEFISTO_opt <- get_default_mefisto_options(sm)</pre>
```

### Description

Function to obtain the default model options.

#### Usage

```
get_default_model_options(object)
```

### **Arguments**

object an untrained MOFA object

#### **Details**

This function provides a default set of model options that can be modified and passed to the MOFA object in the prepare\_mofa step (see example), i.e. after creating a MOFA object (using create\_mofa) and before starting the training (using run\_mofa) The model options are the following:

- **likelihoods**: character vector with data likelihoods per view: 'gaussian' for continuous data (Default for all views), 'bernoulli' for binary data and 'poisson' for count data.
- num\_factors: numeric value indicating the (initial) number of factors. Default is 15.
- **spikeslab\_factors**: logical indicating whether to use spike and slab sparsity on the factors (Default is FALSE)
- **spikeslab\_weights**: logical indicating whether to use spike and slab sparsity on the weights (Default is TRUE)
- ard\_factors: logical indicating whether to use ARD sparsity on the factors (Default is TRUE only if using multiple groups)
- **ard\_weights**: logical indicating whether to use ARD sparsity on the weights (Default is TRUE)

#### Value

Returns a list with the default model options.

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")

# Load data dt (in data.frame format)
load(file)

# Create the MOFA object
MOFAmodel <- create_mofa(dt)

# Load default model options
model_opts <- get_default_model_options(MOFAmodel)

# Edit some of the model options
model_opts$num_factors <- 10
model_opts$spikeslab_weights <- FALSE

# Prepare the MOFA object
MOFAmodel <- prepare_mofa(MOFAmodel, model_options = model_opts)</pre>
```

#### **Description**

Function to obtain the default options for stochastic variational inference.

### Usage

```
get_default_stochastic_options(object)
```

### **Arguments**

object an untrained MOFA

#### Details

This function provides a default set of stochastic inference options that can be modified and passed to the MOFA object in the prepare\_mofa step), i.e. after creating a MOFA object (using create\_mofa) and before starting the training (using run\_mofa) These options are only relevant when activating stochastic inference in training\_options (see example). The stochastic inference options are the following:

- batch\_size: numeric value indicating the batch size (as a fraction). Default is 0.5 (half of the data set).
- learning\_rate: numeric value indicating the learning rate. Default is 1.0
- forgetting\_rate: numeric indicating the forgetting rate. Default is 0.5
- start\_stochastic: integer indicating the first iteration to start stochastic inference Default is 1

#### Value

Returns a list with default options

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")
# Load data dt (in data.frame format)
load(file)
# Create the MOFA object
MOFAmodel <- create_mofa(dt)
# activate stochastic inference in training options
train_opts <- get_default_training_options(MOFAmodel)</pre>
```

```
train_opts$stochastic <- TRUE

# Load default stochastic options
stochastic_opts <- get_default_stochastic_options(MOFAmodel)

# Edit some of the stochastic options
stochastic_opts$learning_rate <- 0.75
stochastic_opts$batch_size <- 0.25

# Prepare the MOFA object
MOFAmodel <- prepare_mofa(MOFAmodel,
    training_options = train_opts,
    stochastic_options = stochastic_opts
)</pre>
```

### Description

Function to obtain the default training options.

# Usage

```
get_default_training_options(object)
```

#### Arguments

object an untrained MOFA

#### **Details**

This function provides a default set of training options that can be modified and passed to the MOFA object in the prepare\_mofa step (see example), i.e. after creating a MOFA object (using create\_mofa) and before starting the training (using run\_mofa) The training options are the following:

- maxiter: numeric value indicating the maximum number of iterations. Default is 1000. Convergence is assessed using the ELBO statistic.
- **drop\_factor\_threshold**: numeric indicating the threshold on fraction of variance explained to consider a factor inactive and drop it from the model. For example, a value of 0.01 implies that factors explaining less than 1% of variance (in each view) will be dropped. Default is -1 (no dropping of factors)
- **convergence\_mode**: character indicating the convergence criteria, either "fast", "medium" or "slow", corresponding to 0.0005%, 0.00005% or 0.000005% deltaELBO change.

28 get\_dimensions

- verbose: logical indicating whether to generate a verbose output.
- startELBO: integer indicating the first iteration to compute the ELBO (default is 1).
- freqELBO: integer indicating the first iteration to compute the ELBO (default is 1).
- **stochastic**: logical indicating whether to use stochastic variational inference (only required for very big data sets, default is FALSE).
- **gpu\_mode**: logical indicating whether to use GPUs (see details).
- seed: numeric indicating the seed for reproducibility (default is 42).

#### Value

Returns a list with default training options

### **Examples**

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")

# Load data dt (in data.frame format)
load(file)

# Create the MOFA object
MOFAmodel <- create_mofa(dt)

# Load default training options
train_opts <- get_default_training_options(MOFAmodel)

# Edit some of the training options
train_opts$convergence_mode <- "medium"
train_opts$startELBO <- 100
train_opts$seed <- 42

# Prepare the MOFA object
MOFAmodel <- prepare_mofa(MOFAmodel, training_options = train_opts)</pre>
```

 ${\tt get\_dimensions}$ 

Get dimensions

#### **Description**

Extract dimensionalities from the model.

#### Usage

```
get_dimensions(object)
```

#### **Arguments**

object

a MOFA object.

get\_elbo 29

#### **Details**

K indicates the number of factors, M indicates the number of views, D indicates the number of features (per view), N indicates the number of samples (per group) and C indicates the number of covariates.

### Value

list containing the dimensionalities of the model

# **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
dims <- get_dimensions(model)</pre>
```

get\_elbo

Get ELBO

### **Description**

Extract the value of the ELBO statistics after model training. This can be useful for model selection.

### Usage

```
get_elbo(object)
```

#### **Arguments**

object

a MOFA object.

#### **Details**

This can be useful for model selection.

### Value

Value of the ELBO

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
elbo <- get_elbo(model)</pre>
```

30 get\_expectations

ations Get expectations
-------------------------

#### **Description**

Function to extract the expectations from the (variational) posterior distributions of a trained MOFA object.

#### Usage

```
get_expectations(object, variable, as.data.frame = FALSE)
```

### **Arguments**

object a trained MOFA object.

variable variable name: 'Z' for factors and 'W' for weights.

as.data.frame logical indicating whether to output the result as a long data frame, default is

FALSE.

#### **Details**

Technical note: MOFA is a Bayesian model where each variable has a prior distribution and a posterior distribution. In particular, to achieve scalability we used the variational inference framework, thus true posterior distributions are replaced by approximated variational distributions. This function extracts the expectations of the variational distributions, which can be used as final point estimates to analyse the results of the model.

The priors and variational distributions of each variable are extensively described in the supplementary methods of the original paper.

#### Value

the output varies depending on the variable of interest:

- "Z"a matrix with dimensions (samples, factors). If as.data.frame is TRUE, a long-formatted data frame with columns (sample, factor, value)
- "W"a list of length (views) where each element is a matrix with dimensions (features, factors). If as.data.frame is TRUE, a long-formatted data frame with columns (view, feature, factor, value)

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
factors <- get_expectations(model, "Z")
weights <- get_expectations(model, "W")</pre>
```

get\_factors 31

get_factors	Get factors		

### **Description**

Extract the latent factors from the model.

### Usage

```
get_factors(
  object,
  groups = "all",
  factors = "all",
  scale = FALSE,
  as.data.frame = FALSE
)
```

# **Arguments**

object a trained MOFA object.

groups character vector with the group name(s), or numeric vector with the group index(es). Default is "all".

factors character vector with the factor name(s), or numeric vector with the factor index(es). Default is "all".

scale logical indicating whether to scale factor values.

as.data.frame logical indicating whether to return a long data frame instead of a matrix. De-

### Value

By default it returns the latent factor matrix of dimensionality (N,K), where N is number of samples and K is number of factors.

Alternatively, if as.data.frame is TRUE, returns a long-formatted data frame with columns (sample,factor,value).

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Fetch factors in matrix format (a list, one matrix per group)
factors <- get_factors(model)

# Concatenate groups
factors <- do.call("rbind",factors)</pre>
```

fault is FALSE.

32 get\_imputed\_data

```
# Fetch factors in data.frame format instead of matrix format
factors <- get_factors(model, as.data.frame = TRUE)</pre>
```

```
get_group_kernel
```

Get group covariance matrix

### **Description**

Extract the inferred group-group covariance matrix per factor

### Usage

```
get_group_kernel(object)
```

### Arguments

```
object
```

a MOFA object.

#### **Details**

This can be used only if covariates are passed to the MOFAobject upon creation and GP\_factors is set to True.

#### Value

A list of group-group correlation matrices per factor

```
get_imputed_data
```

Get imputed data

# Description

Function to get the imputed data. It requires the previous use of the impute method.

### Usage

```
get_imputed_data(
  object,
  views = "all",
  groups = "all",
  features = "all",
  as.data.frame = FALSE
)
```

#### **Arguments**

object	a trained MOFA object.
views	character vector with the view name(s), or numeric vector with the view index(es). Default is "all".
groups	character vector with the group $name(s)$ , or numeric vector with the group $index(es)$ . Default is "all".
features	list of character vectors with the feature names or list of numeric vectors with the feature indices. Default is "all".

logical indicating whether to return a long-formatted data frame instead of a list

as.data.frame of matrices. Default is FALSE.

#### **Details**

Data is imputed from the generative model of MOFA.

### Value

A list containing the imputed valued or a data.frame if as.data.frame is TRUE

### **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")</pre>
model <- load_model(file)</pre>
model <- impute(model)</pre>
imputed <- get_imputed_data(model)</pre>
```

```
get_interpolated_factors
```

Get interpolated factor values

# **Description**

Extract the interpolated factor values

### Usage

```
get_interpolated_factors(object, as.data.frame = FALSE, only_mean = FALSE)
```

### **Arguments**

```
a MOFA object
object
```

as.data.frame logical indicating whether to return data as a data.frame

logical indicating whether include only mean or also uncertainties only\_mean

34 get\_lengthscales

#### **Details**

This can be used only if covariates are passed to the object upon creation, GP\_factors is set to True and new covariates were passed for interpolation.

### Value

By default, a nested list containing for each group a list with a matrix with the interpolated factor values ("mean"), their variance ("variance") and the values of the covariate at which interpolation took place ("new\_values"). Alternatively, if as.data.frame is TRUE, returns a long-formatted data frame with columns containing the covariates and (factor, group, mean and variance).

get\_lengthscales

Get lengthscales

#### **Description**

Extract the inferred lengthscale for each factor after model training.

#### Usage

```
get_lengthscales(object)
```

### Arguments

object

a MOFA object.

#### **Details**

This can be used only if covariates are passed to the MOFAobject upon creation and GP\_factors is set to True.

#### Value

A numeric vector containing the lengthscale for each factor.

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
ls <- get_lengthscales(model)</pre>
```

get\_scales 35

get\_scales

Get scales

#### **Description**

Extract the inferred scale for each factor after model training.

#### Usage

```
get_scales(object)
```

#### **Arguments**

```
object
```

a MOFA object.

#### **Details**

This can be used only if covariates are passed to the MOFAobject upon creation and GP\_factors is set to True.

#### Value

A numeric vector containing the scale for each factor.

# **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
s <- get_scales(model)</pre>
```

```
get_variance_explained
```

Get variance explained values

### **Description**

Extract the latent factors from the model.

# Usage

```
get_variance_explained(
  object,
  groups = "all",
  views = "all",
  factors = "all",
  as.data.frame = FALSE
)
```

36 get\_weights

### **Arguments**

object	a trained MOFA object.
groups	character vector with the group name(s), or numeric vector with the group index(es). Default is "all".
views	character vector with the view name(s), or numeric vector with the view index(es). Default is "all".
factors	character vector with the factor $name(s)$ , or numeric vector with the factor index(es). Default is "all".
as.data.frame	logical indicating whether to return a long data frame instead of a matrix. Default is FALSE.

### Value

A list of data matrices with variance explained per group or a data.frame (if as.data.frame is TRUE)

# **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Fetch variance explained values (in matrix format)
r2 <- get_variance_explained(model)

# Fetch variance explained values (in data.frame format)
r2 <- get_variance_explained(model, as.data.frame = TRUE)</pre>
```

get\_weights

Get weights

### **Description**

Extract the weights from the model.

# Usage

```
get_weights(
  object,
  views = "all",
  factors = "all",
  abs = FALSE,
  scale = FALSE,
  as.data.frame = FALSE
)
```

groups\_names 37

### **Arguments**

object	a trained MOFA object.
views	character vector with the view name(s), or numeric vector with the view index(es). Default is "all".
factors	character vector with the factor name(s) or numeric vector with the factor index(es). Default is "all".
abs	logical indicating whether to take the absolute value of the weights.
scale	logical indicating whether to scale all weights from -1 to 1 (or from 0 to 1 if abs=TRUE).
as.data.frame	logical indicating whether to return a long data frame instead of a list of matrices. Default is FALSE.

## Value

By default it returns a list where each element is a loading matrix with dimensionality (D,K), where D is the number of features and K is the number of factors.

Alternatively, if as.data.frame is TRUE, returns a long-formatted data frame with columns (view,feature,factor,value).

## **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Fetch weights in matrix format (a list, one matrix per view)
weights <- get_weights(model)

# Fetch weights for factor 1 and 2 and view 1
weights <- get_weights(model, views = 1, factors = c(1,2))

# Fetch weights in data.frame format
weights <- get_weights(model, as.data.frame = TRUE)</pre>
```

groups\_names groups\_names: set and retrieve group names

## **Description**

groups\_names: set and retrieve group names

impute impute

### Usage

```
groups_names(object)
groups_names(object) <- value
## S4 method for signature 'MOFA'
groups_names(object)
## S4 replacement method for signature 'MOFA, character'
groups_names(object) <- value</pre>
```

### **Arguments**

object a MOFA object.

value character vector with the names for each group

#### Value

character vector with the names for each sample group

## **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
groups_names(model)
groups_names(model) <- c("my_group")</pre>
```

impute

Impute missing values from a fitted MOFA

## **Description**

This function uses the latent factors and the loadings to impute missing values.

### Usage

```
impute(
  object,
  views = "all",
  groups = "all",
  factors = "all",
  add_intercept = TRUE
)
```

interpolate\_factors 39

### Arguments

object	a MOFA object.
views	character vector with the view name(s), or numeric vector with view index(es).
groups	character vector with the group name(s), or numeric vector with group index(es).
factors	character vector with the factor names, or numeric vector with the factor index(es).

- response: gives mean for gaussian and poisson and probabilities for bernoulli.
- link: gives the linear predictions.
- **inRange**: rounds the fitted values from "terms" for integer-valued distributions to the next integer (default).

add\_intercept add feature intercepts to the imputation (default is TRUE).

#### **Details**

MOFA generates a denoised and condensed low-dimensional representation of the data that captures the main sources of heterogeneity of the data. This representation can be used to reconstruct the data, simply using the equation Y = WX. For more details read the supplementary methods of the manuscript.

Note that with impute you can only generate the point estimates (the means of the posterior distributions). If you want to add uncertainity estimates (the variance) you need to set impute=TRUE in the training options. See get\_default\_training\_options.

## Value

This method fills the imputed\_data slot by replacing the missing values in the input data with the model predictions.

# Examples

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Impute missing values in all data modalities
imputed_data <- impute(model, views = "all")

# Impute missing values in all data modalities using factors 1:3
imputed_data <- impute(model, views = "all", factors = 1:3)</pre>
```

interpolate\_factors

Interpolate factors in MEFISTO based on new covariate values

# Description

Function to interpolate factors in MEFISTO based on new covariate values.

40 load\_model

### Usage

```
interpolate_factors(object, new_values)
```

### **Arguments**

object a MOFA object trained with MEFISTO options and a covariate

new\_values a matrix containing the new covariate values to inter/extrapolate to. Should be

in the same format as the covariated used for training.

#### **Details**

This function requires the functional MEFISTO framework to be used in training. Use set\_covariates and specify mefisto\_options when preparing the training using prepare\_mofa. Currenlty, only the mean of the interpolation is provided from R.

#### Value

Returns the MOFA with interpolated factor values filled in the corresponding slot (interpolatedZ)

## **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
model <- interpolate_factors(model, new_values = seq(0,1.1,0.01))</pre>
```

load\_model

Load a trained MOFA

# Description

Method to load a trained MOFA

The training of mofa is done using a Python framework, and the model output is saved as an .hdf5 file, which has to be loaded in the R package.

### Usage

```
load_model(
    file,
    sort_factors = TRUE,
    on_disk = FALSE,
    load_data = TRUE,
    remove_outliers = FALSE,
    remove_inactive_factors = TRUE,
    verbose = FALSE,
    load_interpol_Z = FALSE
)
```

make\_example\_data 41

## Arguments

file an hdf5 file saved by the mofa Python framework

sort\_factors logical indicating whether factors should be sorted by variance explained (de-

fault is TRUE)

on\_disk logical indicating whether to work from memory (FALSE) or disk (TRUE).

This should be set to TRUE when the training data is so big that cannot fit into

memory.

On-disk operations are performed using the HDF5Array and DelayedArray frame-

work.

load\_data logical indicating whether to load the training data (default is TRUE, it can be

memory expensive)

remove\_outliers

logical indicating whether to mask outlier values.

remove\_inactive\_factors

logical indicating whether to remove inactive factors from the model.

verbose logical indicating whether to print verbose output (default is FALSE)

load\_interpol\_Z

(MEFISTO) logical indicating whether to load predictions for factor values based on latent processed (only relevant for models trained with covariates and Gaus-

sian processes, where prediction was enabled)

### Value

a MOFA model

### **Examples**

```
#' # Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)</pre>
```

make\_example\_data

Simulate a data set using the generative model of MOFA

### **Description**

Function to simulate an example multi-view multi-group data set according to the generative model of MOFA2.

42 make\_example\_data

# Usage

```
make_example_data(
  n_views = 3,
  n_features = 100,
  n_samples = 50,
  n_groups = 1,
  n_factors = 5,
  likelihood = "gaussian",
  lscales = 1,
  sample_cov = NULL,
  as.data.frame = FALSE
)
```

# Arguments

n_views	number of views
n_features	number of features in each view
n_samples	number of samples in each group
n_groups	number of groups
n_factors	number of factors
likelihood	likelihood for each view, one of "gaussian" (default), "bernoulli", "poisson", or a character vector of length $n\_views$
lscales	vector of lengthscales, needs to be of length $n_{\text{factors}}$ (default is $0$ - no smooth factors)
sample_cov	(only for use with MEFISTO) matrix of sample covariates for one group with covariates in rows and samples in columns or "equidistant" for sequential ordering, default is NULL (no smooth factors)
as.data.frame	return data and covariates as long dataframe

## Value

Returns a list containing the simulated data and simulation parameters.

```
# Generate a simulated data set
MOFAexample <- make_example_data()</pre>
```

MOFA 43

**MOFA** 

Class to store a mofa model

### **Description**

The MOFA is an S4 class used to store all relevant data to analyse a MOFA model

### **Slots**

data The input data

intercepts Feature intercepts

samples\_metadata Samples metadata

features\_metadata Features metadata.

imputed\_data The imputed data.

expectations expected values of the factors and the loadings.

dim\_red non-linear dimensionality reduction manifolds.

training\_stats model training statistics.

data\_options Data processing options.

training\_options Model training options.

stochastic\_options Stochastic variational inference options.

model\_options Model options.

mefisto\_options Options for the use of MEFISO

dimensions Dimensionalities of the model: M for the number of views, G for the number of groups, N for the number of samples (per group), C for the number of covariates per sample, D for the number of features (per view), K for the number of factors.

on\_disk Logical indicating whether data is loaded from disk.

cache Cache.

status Auxiliary variable indicating whether the model has been trained.

covariates optional slot to store sample covariate for training in MEFISTO

covariates\_warped optional slot to store warped sample covariate for training in MEFISTO

interpolated\_Z optional slot to store interpolated factor values (used only with MEFISTO)

plot\_ascii\_data

plot\_alignment

Plot covariate alignment acorss groups

## **Description**

Function to plot the alignment learnt by MEFISTO for the covariate values between different groups

### Usage

```
plot_alignment(object)
```

## Arguments

object

a MOFA object using MEFISTO with warping

#### **Details**

This function requires the functional MEFISTO framework to be used in training. Use set\_covariates and specify mefisto\_options when preparing the training using prepare\_mofa.

### Value

ggplot object showing the alignment

plot\_ascii\_data

Visualize the structure of the data in the terminal

# Description

A Fancy printing method

### Usage

```
plot_ascii_data(object, nonzero = FALSE)
```

# **Arguments**

object a MOFA object

nonzero a logical value specifying whether to calculate the fraction of non-zero values

(non-NA values by default)

### **Details**

This function is helpful to get an overview of the structure of the data as a text output

plot\_data\_heatmap 45

## Value

None

## **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_ascii_data(model)</pre>
```

plot\_data\_heatmap

Plot heatmap of relevant features

### **Description**

Function to plot a heatmap of the data for relevant features, typically the ones with high weights.

## Usage

```
plot_data_heatmap(
  object,
  factor,
  view = 1,
  groups = "all",
  features = 50,
  annotation_features = NULL,
  annotation_samples = NULL,
  transpose = FALSE,
  imputed = FALSE,
  denoise = FALSE,
  max.value = NULL,
  min.value = NULL,
  ...
)
```

### **Arguments**

object a MOFA object.

factor a string with the factor name, or an integer with the index of the factor.

view a string with the view name, or an integer with the index of the view. Default is

the first view.

groups groups to plot. Default is "all".

features if an integer (default), the total number of features to plot based on the absolute

value of the weights. If a character vector, a set of manually defined features.

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annotation\_features

annotation metadata for features (rows). Either a character vector specifying columns in the feature metadata, or a data.frame that will be passed to pheatmap as annotation\_col

annotation\_samples

annotation metadata for samples (columns). Either a character vector specifying columns in the sample metadata, or a data.frame that will be passed to pheatmap as annotation\_row

transpose logical indicating whether to transpose the heatmap. Default corresponds to

features as rows and samples as columns.

imputed logical indicating whether to plot the imputed data instead of the original data.

Default is FALSE.

denoise logical indicating whether to plot a denoised version of the data reconstructed

using the MOFA factors.

max.value numeric indicating the maximum value to display in the heatmap (i.e. the matrix

values will be capped at max.value).

min.value numeric indicating the minimum value to display in the heatmap (i.e. the matrix

values will be capped at min. value ). See predict. Default is FALSE.

... further arguments that can be passed to pheatmap

#### Details

One of the first steps for the annotation of a given factor is to visualise the corresponding weights, using for example plot\_weights or plot\_top\_weights.

However, one might also be interested in visualising the direct relationship between features and factors, rather than looking at "abstract" weights.

This function generates a heatmap for selected features, which should reveal the underlying pattern that is captured by the latent factor.

A similar function for doing scatterplots rather than heatmaps is plot\_data\_scatter.

#### Value

A pheatmap object

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_data_heatmap(model, factor = 1, show_rownames = FALSE, show_colnames = FALSE)</pre>
```

plot\_data\_overview 47

plot\_data\_overview Ove

Overview of the input data

#### **Description**

Function to do a tile plot showing the missing value structure of the input data

### Usage

```
plot_data_overview(
  object,
  covariate = 1,
  colors = NULL,
  show_covariate = FALSE,
  show_dimensions = TRUE
)
```

#### **Arguments**

object a MOFA object.

covariate (only for MEFISTO) specifies sample covariate to order samples by in the plot.

This should be a character or a numeric index giving the name or position of a column present in the covariates slot of the object. Default is the first sample

covariate in covariates slot. NULL does not order by covariate

colors a vector specifying the colors per view (see example for details).

show\_covariate (only for MEFISTO) boolean specifying whether to include the covariate in the

plot

show\_dimensions

logical indicating whether to plot the dimensions of the data (default is TRUE).

## Details

This function is helpful to get an overview of the structure of the data. It shows the model dimensionalities (number of samples, groups, views and features) and it indicates which measurements are missing.

#### Value

```
A ggplot object
```

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_data_overview(model)</pre>
```

48 plot\_data\_scatter

plot\_data\_scatter

Scatterplots of feature values against latent factors

### **Description**

Function to do a scatterplot of features against factor values.

## Usage

```
plot_data_scatter(
  object,
  factor = 1,
  view = 1,
  groups = "all",
  features = 10,
  sign = "all",
  color_by = "group",
  legend = TRUE,
  alpha = 1,
  shape_by = NULL,
  stroke = NULL,
  dot_size = 2.5,
  text_size = NULL,
  add_lm = TRUE,
  lm_per_group = TRUE,
  imputed = FALSE
)
```

### Arguments

object a MOFA object. factor string with the factor name, or an integer with the index of the factor. view string with the view name, or an integer with the index of the view. Default is the first view. groups groups to plot. Default is "all". if an integer (default), the total number of features to plot. If a character vector, features a set of manually-defined features. can be 'positive', 'negative' or 'all' (default) to show only positive, negative or sign all weights, respectively. specifies groups or values (either discrete or continuous) used to color the dots color\_by (samples). This can be either:

- the string "group": dots are coloured with respect to their predefined groups.
- a character giving the name of a feature that is present in the input data
- a character giving the same of a column in the sample metadata slot

plot\_data\_scatter 49

• a vector of the same length as the number of samples specifying the value

	for each sample.
	• a dataframe with two columns: "sample" and "color"
legend	logical indicating whether to add a legend
alpha	numeric indicating dot transparency (default is 1).
shape_by	specifies groups or values (only discrete) used to shape the dots (samples). This can be either:
	<ul> <li>the string "group": dots are shaped with respect to their predefined groups.</li> <li>a character giving the name of a feature that is present in the input data</li> <li>a character giving the same of a column in the sample metadata slot</li> <li>a vector of the same length as the number of samples specifying the value for each sample.</li> </ul>
	• a dataframe with two columns: "sample" and "shape"
stroke	numeric indicating the stroke size (the black border around the dots, default is NULL, infered automatically).
dot_size	numeric indicating dot size (default is 5).
text_size	numeric indicating text size (default is 5).
add_lm	logical indicating whether to add a linear regression line for each plot
lm_per_group	logical indicating whether to add a linear regression line separately for each

### **Details**

imputed

One of the first steps for the annotation of factors is to visualise the weights using plot\_weights or plot\_top\_weights. However, one might also be interested in visualising the direct relationship between features and factors, rather than looking at "abstract" weights.

logical indicating whether to include imputed measurements

A similar function for doing heatmaps rather than scatterplots is plot\_data\_heatmap.

#### Value

```
A ggplot object
```

group

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_data_scatter(model)</pre>
```

50 plot\_data\_vs\_cov

plot\_data\_vs\_cov

Scatterplots of feature values against sample covariates

# Description

Function to do a scatterplot of features against sample covariate values.

## Usage

```
plot_data_vs_cov(
 object,
  covariate = 1,
 warped = TRUE,
  factor = 1,
  view = 1,
  groups = "all",
  features = 10,
  sign = "all",
  color_by = "group",
  legend = TRUE,
  alpha = 1,
  shape_by = NULL,
  stroke = NULL,
  dot_size = 2.5,
  text_size = NULL,
  add_lm = FALSE,
  lm_per_group = FALSE,
  imputed = FALSE,
  return_data = FALSE
)
```

## Arguments

object	a MOFA object using MEFISTO.
covariate	string with the covariate name or a samples_metadata column, or an integer with the index of the covariate
warped	logical indicating whether to show the aligned covariate (default: TRUE), only relevant if warping has been used to align multiple sample groups
factor	string with the factor name, or an integer with the index of the factor to take top features from
view	string with the view name, or an integer with the index of the view. Default is the first view.
groups	groups to plot. Default is "all".
features	if an integer (default), the total number of features to plot (given by highest weights). If a character vector, a set of manually-defined features.

pioi_data_vs_cov	31

sign can be 'positive', 'negative' or 'all' (default) to show only features with highest positive, negative or all weights, respectively.

specifies groups or values (either discrete or continuous) used to color the dots (samples). This can be either:

• the string "group": dots are coloured with respect to their predefined groups.

**5** 1

- a character giving the name of a feature that is present in the input data
- a character giving the same of a column in the sample metadata slot
- a vector of the same length as the number of samples specifying the value for each sample.
- a dataframe with two columns: "sample" and "color"

legend logical indicating whether to add a legend

alpha numeric indicating dot transparency (default is 1).

shape\_by specifies groups or values (only discrete) used to shape the dots (samples). This can be either:

- the string "group": dots are shaped with respect to their predefined groups.
- a character giving the name of a feature that is present in the input data
- a character giving the same of a column in the sample metadata slot
- a vector of the same length as the number of samples specifying the value for each sample.
- a dataframe with two columns: "sample" and "shape"

stroke numeric indicating the stroke size (the black border around the dots, default is

NULL, inferred automatically).

dot\_size numeric indicating dot size (default is 5).

text\_size numeric indicating text size (default is 5).

add\_lm logical indicating whether to add a linear regression line for each plot

lm\_per\_group logical indicating whether to add a linear regression line separately for each

group

imputed logical indicating whether to include imputed measurements

return\_data logical indicating whether to return a data frame instead of a plot

### Details

mlat data via acri

color\_by

One of the first steps for the annotation of factors is to visualise the weights using plot\_weights or plot\_top\_weights and inspect the relationshio of the factor to the covariate(s) using plot\_factors\_vs\_cov. However, one might also be interested in visualising the direct relationship between features and covariate(s), rather than looking at "abstract" weights and possibly look at the interpolated and extrapolated values by setting imputed to True.

#### Value

Returns a ggplot2 object or the underlying dataframe if return\_data is set to TRUE.

52 plot\_dimred

### **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_data_vs_cov(model, factor = 3, features = 2)</pre>
```

plot\_dimred

Plot dimensionality reduction based on MOFA factors

# Description

Plot dimensionality reduction based on MOFA factors

# Usage

```
plot_dimred(
  object,
 method = c("UMAP", "TSNE"),
  groups = "all",
  show_missing = TRUE,
  color_by = NULL,
  shape_by = NULL,
  color_name = NULL,
  shape_name = NULL,
  label = FALSE,
  dot_size = 1.5,
  stroke = NULL,
  alpha_missing = 1,
  legend = TRUE,
  rasterize = FALSE,
  return_data = FALSE,
)
```

### **Arguments**

object	a trained MOFA object.
method	string indicating which method has been used for non-linear dimensionality reduction (either 'umap' or 'tsne')
groups	character vector with the groups names, or numeric vector with the indices of the groups of samples to use, or "all" to use samples from all groups.
show_missing	logical indicating whether to include samples for which shape_by or color_by is missing

plot\_dimred 53

color\_by specifies groups or values used to color the samples. This can be either: (1) a character giving the name of a feature present in the training data. (2) a character giving the same of a column present in the sample metadata. (3) a vector of the same length as the number of samples specifying discrete groups or continuous numeric values. shape\_by specifies groups or values used to shape the samples. This can be either: (1) a character giving the name of a feature present in the training data, (2) a character giving the same of a column present in the sample metadata. (3) a vector of the same length as the number of samples specifying discrete groups. color\_name name for color legend. shape\_name name for shape legend. logical indicating whether to label the medians of the clusters. Only if color\_by label is specified dot\_size numeric indicating dot size. stroke numeric indicating the stroke size (the black border around the dots, default is NULL, infered automatically). alpha\_missing numeric indicating dot transparency of missing data. legend logical indicating whether to add legend. rasterize logical indicating whether to rasterize plot using geom\_point\_rast

## **Details**

return\_data

This function plots dimensionality reduction projections that are stored in the dim\_red slot. Typically this contains UMAP or t-SNE projections computed using run\_tsne or run\_umap, respectively.

extra arguments passed to run\_umap or run\_tsne.

logical indicating whether to return the long data frame to plot instead of plotting

#### Value

Returns a ggplot2 object or a long data.frame (if return data is TRUE)

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Run UMAP
model <- run_umap(model)

# Plot UMAP
plot_dimred(model, method = "UMAP")

# Plot UMAP, colour by Factor 1 values
plot_dimred(model, method = "UMAP", color_by = "Factor1")</pre>
```

54 plot\_enrichment

```
# Plot UMAP, colour by the values of a specific feature
plot_dimred(model, method = "UMAP", color_by = "feature_0_view_0")
```

plot\_enrichment

Plot output of gene set Enrichment Analysis

# Description

Method to plot the results of the gene set Enrichment Analysisis

## Usage

```
plot_enrichment(
  enrichment.results,
  factor,
  alpha = 0.1,
  max.pathways = 25,
  text_size = 1,
  dot_size = 5
)
```

# Arguments

```
enrichment.results
```

output of run\_enrichment function

factor a string with the factor name or an integer with the factor index

alpha p.value threshold to filter out gene sets

max.pathways maximum number of enriched pathways to display

text\_size text size dot\_size dot size

## **Details**

it requires run\_enrichment to be run beforehand.

### Value

```
a ggplot2 object
```

```
plot_enrichment_detailed
```

Plot detailed output of the Feature Set Enrichment Analysis

## Description

Method to plot a detailed output of the Feature Set Enrichment Analysisis (FSEA).

Each row corresponds to a significant pathway, sorted by statistical significance, and each dot corresponds to a gene.

For each pathway, we display the top genes of the pathway sorted by the corresponding feature statistic (by default, the absolute value of the weight) The top genes with the highest statistic (max.genes argument) are displayed and labeled in black. The remaining genes are colored in grey.

# Usage

```
plot_enrichment_detailed(
  enrichment.results,
  factor,
  alpha = 0.1,
  max.genes = 5,
  max.pathways = 10,
  text_size = 3
)
```

### **Arguments**

```
enrichment.results
output of run_enrichment function

factor string with factor name or numeric with factor index
alpha p.value threshold to filter out feature sets

max.genes maximum number of genes to display, per pathway

max.pathways maximum number of enriched pathways to display

text_size size of the text to label the top genes
```

### Value

```
a ggplot2 object
```

56 plot\_factor

```
plot_enrichment_heatmap
```

Heatmap of Feature Set Enrichment Analysis results

# Description

This method generates a heatmap with the adjusted p.values that result from the feature set enrichment analysis. Rows are feature sets and columns are factors.

# Usage

```
plot_enrichment_heatmap(
  enrichment.results,
  alpha = 0.1,
  cap = 1e-50,
  log_scale = TRUE,
  ...
)
```

## Arguments

```
enrichment.results
output of run_enrichment function

alpha FDR threshold to filter out unsignificant feature sets which are not represented in the heatmap. Default is 0.10.

cap cap p-values below this threshold

log_scale logical indicating whether to plot the -log of the p.values.

... extra arguments to be passed to the pheatmap function
```

### Value

produces a heatmap

plot\_factor

Beeswarm plot of factor values

## **Description**

Beeswarm plot of the latent factor values.

plot\_factor 57

### Usage

```
plot_factor(
  object,
  factors = 1,
  groups = "all",
  group_by = "group",
  color_by = "group",
  shape_by = NULL,
  add_dots = TRUE,
  dot_size = 2,
  dot_alpha = 1,
  add_violin = FALSE,
  violin_alpha = 0.5,
  color_violin = TRUE,
  add_boxplot = FALSE,
  boxplot_alpha = 0.5,
  color_boxplot = TRUE,
  show_missing = TRUE,
  scale = FALSE,
  dodge = FALSE,
  color_name = "".
  shape_name = "",
  stroke = NULL,
  legend = TRUE,
  rasterize = FALSE
)
```

### **Arguments**

object

a trained MOFA object.

factors

character vector with the factor names, or numeric vector with the indices of the factors to use, or "all" to plot all factors.

groups

character vector with the groups names, or numeric vector with the indices of the groups of samples to use, or "all" to use samples from all groups.

group\_by

specifies grouping of samples:

- (default) the string "group": in this case, the plot will color samples with respect to their predefined groups.
- a character giving the name of a feature that is present in the input data
- a character giving the name of a column in the sample metadata slot
- a vector of the same length as the number of samples specifying the value for each sample.

color\_by

specifies color of samples. This can be either:

- (default) the string "group": in this case, the plot will color the dots with respect to their predefined groups.
- a character giving the name of a feature that is present in the input data
- a character giving the name of a column in the sample metadata slot

58 plot\_factor

• a vector of the same length as the number of samples specifying the value for each sample.

shape\_by specifies shape of samples. This can be either:

- (default) the string "group": in this case, the plot will shape the dots with respect to their predefined groups.
- a character giving the name of a feature that is present in the input data
- a character giving the name of a column in the sample metadata slot

• a vector of the same length as the number of samples specifying the value for each sample.

add\_dots logical indicating whether to add dots.

dot\_size numeric indicating dot size.

dot\_alpha numeric indicating dot transparency.

add\_violin logical indicating whether to add violin plots
violin\_alpha numeric indicating violin plot transparency.
color\_violin logical indicating whether to color violin plots.
add\_boxplot logical indicating whether to add box plots

boxplot\_alpha numeric indicating boxplot transparency.

color\_boxplot logical indicating whether to color box plots.

show\_missing logical indicating whether to remove samples for which shape\_by or color\_by

is missing.

scale logical indicating whether to scale factor values.

dodge logical indicating whether to dodge the dots (default is FALSE).

color\_name name for color legend (usually only used if color\_by is not a character itself).

shape\_name name for shape legend (usually only used if shape\_by is not a character itself).

stroke numeric indicating the stroke size (the black border around the dots).

legend logical indicating whether to add a legend to the plot (default is TRUE).

rasterize logical indicating whether to rasterize the plot (default is FALSE).

#### Details

One of the main steps for the annotation of factors is to visualise and color them using known covariates or phenotypic data.

This function generates a Beeswarm plot of the sample values in a given latent factor.

Similar functions are plot\_factors for doing scatter plots.

#### Value

Returns a ggplot2

plot\_factors 59

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Plot Factors 1 and 2 and colour by "group"
plot_factor(model, factors = c(1,2), color_by="group")

# Plot Factor 3 and colour by the value of a specific feature
plot_factor(model, factors = 3, color_by="feature_981_view_1")

# Add violin plots
plot_factor(model, factors = c(1,2), color_by="group", add_violin = TRUE)

# Scale factor values from -1 to 1
plot_factor(model, factors = c(1,2), scale = TRUE)</pre>
```

plot\_factors

Scatterplots of two factor values

## **Description**

Scatterplot of the values of two latent factors.

### Usage

```
plot_factors(
  object,
  factors = c(1, 2),
  groups = "all",
  show_missing = TRUE,
  scale = FALSE,
  color_by = NULL,
  shape_by = NULL,
  color_name = NULL,
  shape_name = NULL,
  dot_size = 2,
  alpha = 1,
  legend = TRUE,
  stroke = NULL,
  return_data = FALSE
)
```

# **Arguments**

object a trained MOFA object.

60 plot\_factors

factors a vector of length two with the factors to plot. Factors can be specified either as

a characters

groups character vector with the groups names, or numeric vector with the indices of

the groups of samples to use, or "all" to use samples from all groups.

show\_missing logical indicating whether to include samples for which shape\_by or color\_by

is missing

scale logical indicating whether to scale factor values.

color\_by specifies groups or values used to color the samples. This can be either: (1) a

character giving the name of a feature present in the training data. (2) a character giving the name of a column present in the sample metadata. (3) a vector of the name length as the number of samples specifying discrete groups or continuous

numeric values.

shape\_by specifies groups or values used to shape the samples. This can be either: (1) a

character giving the name of a feature present in the training data, (2) a character giving the name of a column present in the sample metadata. (3) a vector of the

same length as the number of samples specifying discrete groups.

color\_name name for color legend. shape\_name name for shape legend.

dot\_size numeric indicating dot size (default is 2).

alpha numeric indicating dot transparency (default is 1).

legend logical indicating whether to add legend.

stroke numeric indicating the stroke size (the black border around the dots, default is

NULL, infered automatically).

return\_data logical indicating whether to return the data frame to plot instead of plotting

#### **Details**

One of the first steps for the annotation of factors is to visualise and group/color them using known covariates such as phenotypic or clinical data. This method generates a single scatterplot for the combination of two latent factors. TO-FINISH... plot\_factors for doing Beeswarm plots for factors.

#### Value

Returns a ggplot2 object

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Scatterplot of factors 1 and 2
plot_factors(model, factors = c(1,2))
# Shape dots by a column in the metadata</pre>
```

plot\_factors\_vs\_cov 61

```
plot_factors(model, factors = c(1,2), shape_by="group")
# Scale factor values from -1 to 1
plot_factors(model, factors = c(1,2), scale = TRUE)
```

plot\_factors\_vs\_cov

Scatterplots of a factor's values againt the sample covariates

### **Description**

Scatterplots of a factor's values againt the sample covariates

### Usage

```
plot_factors_vs_cov(
  object,
  factors = "all",
  covariates = NULL,
 warped = TRUE,
  show_missing = TRUE,
  scale = FALSE,
  color_by = NULL,
  shape_by = NULL,
  color_name = NULL,
  shape_name = NULL,
  dot_size = 1.5,
  alpha = 1,
  stroke = NULL,
  legend = TRUE,
  rotate_x = FALSE,
  rotate_y = FALSE,
  return_data = FALSE,
  show_variance = FALSE
)
```

### **Arguments**

object a trained MOFA object using MEFISTO.

factors character or numeric specifying the factor(s) to plot, default is "all"

covariates specifies sample covariate(s) to plot against: (1) a character giving the name of

a column present in the sample covariates or sample metadata. (2) a character giving the name of a feature present in the training data. (3) a vector of the same length as the number of samples specifying continuous numeric values per

sample. Default is the first sample covariates in covariates slot

warped logical indicating whether to show the aligned covariate (default: TRUE), only

relevant if warping has been used to align multiple sample groups

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show\_missing (for 1-dim covariates) logical indicating whether to include samples for which shape\_by or color\_by is missing

scale logical indicating whether to scale factor values.

color\_by (for 1-dim covariates) specifies groups or values used to color the samples. This

can be either: (1) a character giving the name of a feature present in the training data. (2) a character giving the same of a column present in the sample metadata. (3) a vector of the same length as the number of samples specifying discrete

groups or continuous numeric values.

shape\_by (for 1-dim covariates) specifies groups or values used to shape the samples. This

can be either: (1) a character giving the name of a feature present in the training data, (2) a character giving the same of a column present in the sample metadata. (3) a vector of the same length as the number of samples specifying discrete

groups.

color\_name (for 1-dim covariates) name for color legend.
shape\_name (for 1-dim covariates) name for shape legend.

dot\_size (for 1-dim covariates) numeric indicating dot size.

alpha (for 1-dim covariates) numeric indicating dot transparency. stroke (for 1-dim covariates) numeric indicating the stroke size

legend (for 1-dim covariates) logical indicating whether to add legend.

rotate\_x (for spatial, 2-dim covariates) Rotate covariate on x-axis rotate\_y (for spatial, 2-dim covariates) Rotate covariate on y-axis

return\_data logical indicating whether to return the data frame to plot instead of plotting

show\_variance (for 1-dim covariates) logical indicating whether to show the marginal variance

of inferred factor values (only relevant for 1-dimensional covariates)

#### Details

To investigate the factors pattern along the covariates (such as time or a spatial coordinate) this function an be used to plot a scatterplot of the factor againt the values of each covariate

#### Value

Returns a ggplot2 object

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_factors_vs_cov(model)</pre>
```

plot\_factor\_cor 63

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Plot correlation matrix between latent factors

## **Description**

Function to plot the correlation matrix between the latent factors.

#### Usage

```
plot_factor_cor(object, method = "pearson", ...)
```

### **Arguments**

object a trained MOFA object.

method a character indicating the type of correlation coefficient to be computed: pearson

(default), kendall, or spearman.

... arguments passed to corrplot

#### **Details**

This method plots the correlation matrix between the latent factors.

The model encourages the factors to be uncorrelated, so this function usually yields a diagonal correlation matrix.

However, it is not a hard constraint such as in Principal Component Analysis and correlations between factors can occur, particularly with large number factors.

Generally, correlated factors are redundant and should be avoided, as they make interpretation harder. Therefore, if you have too many correlated factors we suggest you try reducing the number of factors.

### Value

Returns a symmetric matrix with the correlation coefficient between every pair of factors.

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Plot correlation between all factors
plot_factor_cor(model)</pre>
```

64 plot\_group\_kernel

plot	group	kernel

Heatmap plot showing the group-group correlations per factor

### **Description**

Heatmap plot showing the group-group correlations inferred by the model per factor

## Usage

```
plot_group_kernel(object, factors = "all", groups = "all", ...)
```

# Arguments

object	a trained MOFA object using MEFISTO.
factors	character vector with the factors names, or numeric vector indicating the indices of the factors to use
groups	character vector with the groups names, or numeric vector with the indices of the groups of samples to use, or "all" to use samples from all groups.
	additional parameters that can be passed to pheatmap

### **Details**

The heatmap gives insight into the clustering of the patterns that factors display along the covariate in each group. A correlation of 1 indicates that the module caputred by a factor shows identical patterns across groups, a correlation of zero that it shows distinct patterns, a negative correlation that the patterns go in opposite directions.

### Value

Returns a ggplot, gg object containing the heatmaps

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_group_kernel(model)</pre>
```

## **Description**

make a plot of interpolated covariates versus covariate

### Usage

```
plot_interpolation_vs_covariate(
  object,
  covariate = 1,
  factors = "all",
  only_mean = TRUE,
  show_observed = TRUE
)
```

### **Arguments**

```
object a trained MOFA object using MEFISTO.

covariate covariate to use for plotting
```

factors character or numeric specifying the factor(s) to plot, default is "all"

only\_mean show only mean or include uncertainties?

show\_observed include observed factor values as dots on the plot

### **Details**

to be filled

# Value

Returns a ggplot2 object

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
model <- interpolate_factors(model, new_values = seq(0,1.1,0.1))
plot_interpolation_vs_covariate(model, covariate = "time", factors = 1)</pre>
```

plot\_smoothness

plot_snaredness Barpiot snowing the snaredness per jactor	plot_sharedness	Barplot showing the sharedness per factor
---	-----------------	---

### **Description**

Barplot indicating a sharedness score (between 0 (non-shared) and 1 (shared)) per factor

## Usage

```
plot_sharedness(object, factors = "all", color = "#B8CF87")
```

## **Arguments**

object a trained MOFA object using MEFISTO.

factors character vector with the factors names, or numeric vector indicating the indices

of the factors to use

color for the shared part of the bar

#### **Details**

The sharedness score is calculated as the distance of the learnt group correlation matrix to the identity matrix in terms of the mean absolute distance on the off-diagonal elements.

## Value

Returns a ggplot2 object

plot_smoothness	Barplot showing the smoothness per factor	

## **Description**

Barplot indicating a smoothness score (between 0 (non-smooth) and 1 (smooth)) per factor

# Usage

```
plot_smoothness(object, factors = "all", color = "cadetblue")
```

## **Arguments**

object a trained MOFA object using MEFISTO.

factors character vector with the factors names, or numeric vector indicating the indices

of the factors to use

color for the smooth part of the bar

plot\_top\_weights 67

### **Details**

The smoothness score is given by the scale parameter for the underlying Gaussian process of each factor.

#### Value

Returns a ggplot2 object

### **Examples**

```
# Using an existing trained model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
smoothness_bars <- plot_smoothness(model)</pre>
```

plot\_top\_weights

Plot top weights

### **Description**

Plot top weights for a given factor and view.

### Usage

```
plot_top_weights(
  object,
  view = 1,
  factors = 1,
  nfeatures = 10,
  abs = TRUE,
  scale = TRUE,
  sign = "all"
)
```

## Arguments

object a trained MOFA object. a string with the view name, or an integer with the index of the view. view a character string with factors names, or an integer vector with factors indices. factors number of top features to display. Default is 10 nfeatures abs logical indicating whether to use the absolute value of the weights (Default is FALSE). logical indicating whether to scale all weights from -1 to 1 (or from 0 to 1 if scale abs=TRUE). Default is TRUE. sign can be 'positive', 'negative' or 'all' to show only positive, negative or all weights, respectively. Default is 'all'.

#### **Details**

An important step to annotate factors is to visualise the corresponding feature weights.

This function displays the top features with highest loading whereas the function plot\_top\_weights plots all weights for a given latent factor and view.

Importantly, the weights of the features within a view have relative values and they should not be interpreted in an absolute scale. Therefore, for interpretability purposes we always recommend to scale the weights with scale=TRUE.

### Value

Returns a ggplot2 object

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Plot top weights for Factors 1 and 2 and View 1
plot_top_weights(model, view = 1, factors = c(1,2))

# Do not take absolute value
plot_weights(model, abs = FALSE)</pre>
```

```
plot_variance_explained
```

Plot variance explained by the model

#### **Description**

plots the variance explained by the MOFA factors across different views and groups, as specified by the user. Consider using cowplot::plot\_grid(plotlist = ...) to combine the multiple plots that this function generates.

## Usage

```
plot_variance_explained(
  object,
  x = "view",
  y = "factor",
  split_by = NA,
  plot_total = FALSE,
  factors = "all",
  min_r2 = 0,
  max_r2 = NULL,
  legend = TRUE,
  use_cache = TRUE,
```

```
)
```

## **Arguments**

object	a MOFA object
X	character specifying the dimension for the x-axis ("view", "factor", or "group").
у	character specifying the dimension for the y-axis ("view", "factor", or "group").
split_by	character specifying the dimension to be faceted ("view", "factor", or "group").
plot_total	logical value to indicate if to plot the total variance explained (for the variable in the x-axis)
factors	character vector with a factor name(s), or numeric vector with the index(es) of the factor(s). Default is "all".
min_r2	minimum variance explained for the color scheme (default is 0).
max_r2	maximum variance explained for the color scheme.
legend	logical indicating whether to add a legend to the plot (default is TRUE).
use_cache	logical indicating whether to use cache (default is TRUE)
• • •	extra arguments to be passed to calculate_variance_explained

### Value

A list of ggplot objects (if plot\_total is TRUE) or a single ggplot object

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Calculate variance explained (R2)
r2 <- calculate_variance_explained(model)

# Plot variance explained values (view as x-axis, and factor as y-axis)
plot_variance_explained(model, x="view", y="factor")

# Plot variance explained values (view as x-axis, and group as y-axis)
plot_variance_explained(model, x="view", y="group")

# Plot variance explained values for factors 1 to 3
plot_variance_explained(model, x="view", y="group", factors=1:3)

# Scale R2 values
plot_variance_explained(model, max_r2=0.25)</pre>
```

```
plot_variance_explained_by_covariates
```

Plot variance explained by the smooth components of the model

# Description

This function plots the variance explained by the smooth components (Gaussian processes) underlying the factors in MEFISTO across different views and groups, as specified by the user.

## Usage

```
plot_variance_explained_by_covariates(
  object,
  factors = "all",
  x = "view",
  y = "factor",
  split_by = NA,
  min_r2 = 0,
  max_r2 = NULL,
  compare_total = FALSE,
  legend = TRUE
)
```

#### **Arguments**

object	a MOFA object
factors	character vector with a factor name(s), or numeric vector with the index(es) of the factor(s). Default is "all".
X	character specifying the dimension for the x-axis ("view", "factor", or "group").
У	character specifying the dimension for the y-axis ("view", "factor", or "group").
split_by	character specifying the dimension to be faceted ("view", "factor", or "group").
min_r2	minimum variance explained for the color scheme (default is 0).
max_r2	maximum variance explained for the color scheme.
compare_total	plot corresponding variance explained in total in addition
legend	logical indicating whether to add a legend to the plot (default is TRUE).

### **Details**

Note that this function requires the use of MEFISTO. To activate the functional MEFISTO framework, specify mefisto\_options when preparing the training using prepare\_mofa

#### Value

A list of ggplot objects (if compare\_total is TRUE) or a single ggplot object. Consider using cowplot::plot\_grid(plotlist = ...) to combine the multiple plots that this function generates.

### **Examples**

```
# load_model
file <- system.file("extdata", "MEFISTO_model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_variance_explained_by_covariates(model)

# compare to toal variance explained
plist <- plot_variance_explained_by_covariates(model, compare_total = TRUE)
cowplot::plot_grid(plotlist = plist)</pre>
```

plot\_variance\_explained\_per\_feature

Plot variance explained by the model for a set of features Returns a tile plot with a group on the X axis and a feature along the Y axis

# Description

Plot variance explained by the model for a set of features

Returns a tile plot with a group on the X axis and a feature along the Y axis

### Usage

```
plot_variance_explained_per_feature(
   object,
   view,
   features = 10,
   split_by_factor = FALSE,
   group_features_by = NULL,
   groups = "all",
   factors = "all",
   min_r2 = 0,
   max_r2 = NULL,
   legend = TRUE,
   return_data = FALSE,
   ...
)
```

### **Arguments**

```
object a MOFA object.
```

view a view name or index.

features a vector with indices or names for features from the respective view, or number

of top features to be fetched by their loadings across specified factors. "all" to

plot all features.

split\_by\_factor

logical indicating whether to split R2 per factor or plot R2 jointly

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```
group_features_by
column name of features metadata to group features by
groups a vector with indices or names for sample groups (default is all)
factors a vector with indices or names for factors (default is all)
min_r2 minimum variance explained for the color scheme (default is 0).
max_r2 maximum variance explained for the color scheme.
legend logical indicating whether to add a legend to the plot (default is TRUE).
return_data logical indicating whether to return the data frame to plot instead of plotting
extra arguments to be passed to calculate_variance_explained
```

#### Value

ggplot object

# Examples

```
# Using an existing trained model
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_variance_explained_per_feature(model, view = 1)</pre>
```

plot\_weights

Plot distribution of feature weights (weights)

### **Description**

An important step to annotate factors is to visualise the corresponding feature weights. This function plots all weights for a given latent factor and view, labeling the top ones. In contrast, the function plot\_top\_weights displays only the top features with highest loading.

## Usage

```
plot_weights(
 object,
  view = 1,
  factors = 1,
  nfeatures = 10,
  color_by = NULL,
  shape_by = NULL,
  abs = FALSE,
 manual = NULL,
  color_manual = NULL,
  scale = TRUE,
  dot_size = 1,
  text_size = 5,
 legend = TRUE,
  return_data = FALSE
)
```

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#### **Arguments**

object a MOFA object.

view a string with the view name, or an integer with the index of the view.

factors character vector with the factor name(s), or numeric vector with the index of the

factor(s).

nfeatures number of top features to label.

color\_by specifies groups or values (either discrete or continuous) used to color the dots

(features). This can be either:

• (default) the string "group": in this case, the plot will color the dots with respect to their predefined groups.

• a character giving the name of a feature that is present in the input data

• a character giving the same of a column in the features metadata slot

• a vector of the same length as the number of features specifying the value for each feature

• a dataframe with two columns: "feature" and "color"

shape\_by specifies groups or values (only discrete) used to shape the dots (features). This can be either:

• (default) the string "group": in this case, the plot will shape the dots with respect to their predefined groups.

• a character giving the name of a feature that is present in the input data

• a character giving the same of a column in the features metadata slot

• a vector of the same length as the number of features specifying the value for each feature

• a dataframe with two columns: "feature" and "shape"

abs logical indicating whether to take the absolute value of the weights.

manual A nested list of character vectors with features to be manually labelled (see the

example for details).

color\_manual a character vector with colors, one for each element of 'manual'

scale logical indicating whether to scale all weights from -1 to 1 (or from 0 to 1 if

abs=TRUE).

dot\_size numeric indicating the dot size.

text\_size numeric indicating the text size.

legend logical indicating whether to add legend.

return\_data logical indicating whether to return the data frame to plot instead of plotting

#### Value

A ggplot object or a data. frame if return\_data is TRUE

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Plot distribution of weights for Factor 1 and View 1
plot_weights(model, view = 1, factors = 1)

# Plot distribution of weights for Factors 1 to 3 and View 1
plot_weights(model, view = 1, factors = 1:3)

# Take the absolute value and highlight the top 10 features
plot_weights(model, view = 1, factors = 1, nfeatures = 10, abs = TRUE)

# Change size of dots and text
plot_weights(model, view = 1, factors = 1, text_size = 5, dot_size = 1)</pre>
```

### **Description**

Function to visualize the weights for a given set of factors in a given view.

This is useful to visualize the overall pattern of the weights but not to individually characterise the factors.

To inspect the weights of individual factors, use the functions plot\_weights and plot\_top\_weights

### Usage

```
plot_weights_heatmap(
  object,
  view = 1,
  features = "all",
  factors = "all",
  threshold = 0,
   ...
)
```

### **Arguments**

object a trained MOFA object.

view character vector with the view name(s), or numeric vector with the index of the

view(s) to use. Default is the first view.

features character vector with the feature name(s), or numeric vector with the index of

the feature(s) to use. Default is 'all'.

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factors character vector with the factor name(s), or numeric vector with the index of the

factor(s) to use. Default is 'all'.

threshold threshold on absolute weight values, so that weights with a magnitude below

this threshold (in all factors) are removed

... extra arguments passed to pheatmap.

#### Value

A pheatmap object

# **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_weights_heatmap(model)</pre>
```

```
plot_weights_scatter Scatterplots of weights
```

# **Description**

Scatterplot of the weights values for two factors

### Usage

```
plot_weights_scatter(
  object,
  factors,
  view = 1,
  color_by = NULL,
  shape_by = NULL,
  dot_size = 1,
  name_color = "",
  name_shape = "",
  show_missing = TRUE,
  abs = FALSE,
  scale = TRUE,
  legend = TRUE
```

# Arguments

object a trained MOFA object.

factors a vector of length two with the factors to plot. Factors can be specified either as

a characters using the factor names, or as numeric with the index of the factors

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view	character vector with the voiew name, or numeric vector with the index of the view to use. Default is the first view.
color_by	specifies groups or values used to color the features. This can be either
	<ul> <li>a character giving the same of a column in the feature metadata slot</li> <li>a vector specifying the value for each feature.</li> <li>a dataframe with two columns: "feature" and "color"</li> </ul>
shape_by	specifies groups or values used to shape the features. This can be either
	<ul> <li>a character giving the same of a column in the feature metadata slot</li> <li>a vector specifying the value for each feature.</li> <li>a dataframe with two columns: "feature" and "shape"</li> </ul>
dot_size	numeric indicating dot size.
name_color	name for color legend (usually only used if color_by is not a character itself)
name_shape	name for shape legend (usually only used if shape_by is not a character itself)
show_missing	logical indicating whether to include dots for which shape_by or color_by is missing
abs	logical indicating whether to take the absolute value of the weights.
scale	logical indicating whether to scale all weights from -1 to 1 (or from 0 to 1 if abs=TRUE).
legend	logical indicating whether to add a legend to the plot (default is TRUE).

### **Details**

One of the first steps for the annotation of factors is to visualise and group/color them using known covariates such as phenotypic or clinical data. This method generates a single scatterplot for the combination of two latent factors.

# Value

Returns a ggplot2 object

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
plot_weights_scatter(model, factors = 1:2)</pre>
```

predict 77

predict

Do predictions using a fitted MOFA

### **Description**

This function uses the latent factors and the weights to do data predictions.

### Usage

```
predict(
  object,
  views = "all",
  groups = "all",
  factors = "all",
  add_intercept = TRUE
)
```

# **Arguments**

object a MOFA object.

views character vector with the view name(s), or numeric vector with the view index(es). Default is "all".

groups character vector with the group name(s), or numeric vector with the group index(es). Default is "all".

factors character vector with the factor name(s) or numeric vector with the factor in-

...

dex(es). Default is "all".

add\_intercept add feature intercepts to the prediction (default is TRUE).

#### **Details**

MOFA generates a denoised and condensed low-dimensional representation of the data that captures the main sources of heterogeneity of the data. This representation can be used to reconstruct a denoised representation of the data, simply using the equation Y = WX. For more mathematical details read the supplementary methods of the manuscript.

# Value

Returns a list with the data reconstructed by the model predictions.

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Predict observations for all data modalities
predictions <- predict(model)</pre>
```

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prepare\_mofa

Prepare a MOFA for training

# Description

Function to prepare a MOFA object for training. It requires defining data, model and training options.

# Usage

```
prepare_mofa(
  object,
  data_options = NULL,
  model_options = NULL,
  training_options = NULL,
  stochastic_options = NULL,
  mefisto_options = NULL
)
```

#### **Arguments**

object an untrained MOFA data\_options list of data\_options (see get\_default\_data\_options details). If NULL, default options are used. model\_options list of model options (see get\_default\_model\_options for details). If NULL, default options are used. training\_options list of training options (see get\_default\_training\_options for details). If NULL, default options are used. stochastic\_options list of options for stochastic variational inference (see get\_default\_stochastic\_options for details). If NULL, default options are used. mefisto\_options list of options for mefisto (see get\_default\_mefisto\_options for details). If NULL, default options are used.

### **Details**

This function is called after creating a MOFA object (using create\_mofa) and before starting the training (using run\_mofa). Here, we can specify different options for the data (data\_options), the model (model\_options) and the training (training\_options, stochastic\_options). Take a look at the individual default options for an overview using the get\_default\_XXX\_options functions above.

#### Value

Returns an untrained MOFA with specified options filled in the corresponding slots

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### **Examples**

```
# Using an existing simulated data with two groups and two views
file <- system.file("extdata", "test_data.RData", package = "MOFA2")

# Load data dt (in data.frame format)
load(file)

# Create the MOFA object
MOFAmodel <- create_mofa(dt)

# Prepare MOFA object using default options
MOFAmodel <- prepare_mofa(MOFAmodel)

# Prepare MOFA object changing some of the default model options values
model_opts <- get_default_model_options(MOFAmodel)

model_opts$num_factors <- 10

MOFAmodel <- prepare_mofa(MOFAmodel, model_options = model_opts)</pre>
```

run\_enrichment

Run feature set Enrichment Analysis

# **Description**

Method to perform feature set enrichment analysis. Here we use a slightly modified version of the pcgse function.

#### Usage

```
run_enrichment(
  object,
  view,
  feature.sets,
  factors = "all",
  set.statistic = c("mean.diff", "rank.sum"),
  statistical.test = c("parametric", "cor.adj.parametric", "permutation"),
  sign = c("all", "positive", "negative"),
  min.size = 10,
  nperm = 1000,
  p.adj.method = "BH",
  alpha = 0.1,
  verbose = TRUE
)
```

#### **Arguments**

object a MOFA object.
view a character with

a character with the view name, or a numeric vector with the index of the view

to use.

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feature.sets data structure that holds feature set membership information. Must be a binary

membership matrix (rows are feature sets and columns are features). See details

below for some pre-built gene set matrices.

factors character vector with the factor names, or numeric vector with the index of the

factors for which to perform the enrichment.

set.statistic the set statistic computed from the feature statistics. Must be one of the follow-

ing: "mean.diff" (default) or "rank.sum".

statistical.test

the statistical test used to compute the significance of the feature set statistics under a competitive null hypothesis. Must be one of the following: "parametric"

(default), "cor.adj.parametric", "permutation".

sign use only "positive" or "negative" weights. Default is "all".

min. size Minimum size of a feature set (default is 10).

nperm number of permutations. Only relevant if statistical test is set to "permutation".

Default is 1000

p. adj. method Method to adjust p-values factor-wise for multiple testing. Can be any method

in p.adjust.methods(). Default uses Benjamini-Hochberg procedure.

alpha FDR threshold to generate lists of significant pathways. Default is 0.1

verbose boolean indicating whether to print messages on progress

#### **Details**

The aim of this function is to relate each factor to pre-defined biological pathways by performing a gene set enrichment analysis on the feature weights.

This function is particularly useful when a factor is difficult to characterise based only on the genes with the highest weight.

We provide a few pre-built gene set matrices in the MOFAdata package. See https://github.com/bioFAM/MOFAdata for details.

The function we implemented is based on the pcgse function with some modifications. Please read this paper https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4543476 for details on the math.

#### Value

a list with five elements:

**pval**: matrices with nominal p-values.

**pval.adj**: matrices with FDR-adjusted p-values.

feature.statistics:

matrices with the local (feature-wise) statistics.

**set.statistics**: matrices with the global (gene set-wise) statistics.

**sigPathways** list with significant pathways per factor.

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|--|

### **Description**

Function to train an untrained MOFA object.

# Usage

```
run_mofa(object, outfile = NULL, save_data = TRUE, use_basilisk = FALSE)
```

### **Arguments**

object an untrained MOFA object

outfile output file for the model (.hdf5 format). If NULL, a temporary file is created.

save\_data logical indicating whether to save the training data in the hdf5 file. This is useful

for some downstream analysis (mainly functions with the prefix plot\_data), but

it can take a lot of disk space.

use\_basilisk use basilisk to automatically install a conda environment with mofapy2 and

all dependencies? If FALSE (default), you should specify the right python binary when loading R with reticulate::use\_python(..., force=TRUE) or the right conda environment with reticulate::use\_condaenv(..., force=TRUE).

#### **Details**

This function is called once a MOFA object has been prepared (using prepare\_mofa) In this step the R package calls the mofapy2 Python package, where model training is performed. The interface with Python is done with the reticulate package. If you have several versions of Python installed and R is not detecting the correct one, you can change it using reticulate::use\_python when loading the R session. Alternatively, you can let us install mofapy2 for you using basilisk if you set use\_basilisk to TRUE

#### Value

```
a trained MOFA object
```

```
# Load data (in data.frame format)
file <- system.file("extdata", "test_data.RData", package = "MOFA2")
load(file)

# Create the MOFA object
MOFAmodel <- create_mofa(dt)

# Prepare the MOFA object with default options
MOFAmodel <- prepare_mofa(MOFAmodel)</pre>
```

run\_tsne

```
# Run the MOFA model
## Not run: MOFAmodel <- run_mofa(MOFAmodel, use_basilisk = TRUE)</pre>
```

run\_tsne

Run t-SNE on the MOFA factors

### Description

Run t-SNE on the MOFA factors

### Usage

```
run_tsne(object, factors = "all", groups = "all", ...)
```

#### **Arguments**

object a trained MOFA object.

factors character vector with the factor names, or numeric vector with the indices of the factors to use, or "all" to use all factors (default).

groups character vector with the groups names, or numeric vector with the indices of

the groups of samples to use, or "all" to use all groups (default).

... arguments passed to Rtsne

#### **Details**

This function calls Rtsne to calculate a TSNE representation from the MOFA factors. Subsequently, you can plot the TSNE representation with plot\_dimred or fetch the coordinates using plot\_dimred(..., method="TSNE", return\_data=TRUE). Remember to use set.seed before the function call to get reproducible results.

#### Value

Returns a MOFA object with the MOFA object@dim\_red slot filled with the t-SNE output

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Run
## Not run: model <- run_tsne(model, perplexity = 15)

# Plot
## Not run: model <- plot_dimred(model, method="TSNE")

# Fetch data
## Not run: tsne.df <- plot_dimred(model, method="TSNE", return_data=TRUE)</pre>
```

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run\_umap

Run UMAP on the MOFA factors

# **Description**

Run UMAP on the MOFA factors

### Usage

```
run_umap(
  object,
  factors = "all",
  groups = "all",
  n_neighbors = 30,
  min_dist = 0.3,
  metric = "cosine",
  ...
)
```

### **Arguments**

object a trained MOFA object. factors character vector with the factor names, or numeric vector with the indices of the factors to use, or "all" to use all factors (default). character vector with the groups names, or numeric vector with the indices of groups the groups of samples to use, or "all" to use all groups (default). n\_neighbors number of neighboring points used in local approximations of manifold structure. Larger values will result in more global structure being preserved at the loss of detailed local structure. In general this parameter should often be in the range 5 to 50. min\_dist This controls how tightly the embedding is allowed compress points together. Larger values ensure embedded points are more evenly distributed, while smaller values allow the algorithm to optimise more accurately with regard to local structure. Sensible values are in the range 0.01 to 0.5 metric choice of metric used to measure distance in the input space arguments passed to umap

#### **Details**

This function calls umap to calculate a UMAP representation from the MOFA factors For details on the hyperparameters of UMAP see the documentation of umap. Subsequently, you can plot the UMAP representation with plot\_dimred or fetch the coordinates using plot\_dimred(..., method="UMAP", return\_data=TRUE). Remember to use set.seed before the function call to get reproducible results.

84 samples\_metadata

### Value

Returns a MOFA object with the MOFAobject@dim\_red slot filled with the UMAP output

# **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Change hyperparameters passed to umap
## Not run: model <- run_umap(model, min_dist = 0.01, n_neighbors = 10)
# Plot
## Not run: model <- plot_dimred(model, method="UMAP")

# Fetch data
## Not run: umap.df <- plot_dimred(model, method="UMAP", return_data=TRUE)</pre>
```

samples\_metadata

samples\_metadata: retrieve sample metadata

# **Description**

samples metadata: retrieve sample metadata

### Usage

```
samples_metadata(object)
samples_metadata(object) <- value

## S4 method for signature 'MOFA'
samples_metadata(object)

## S4 replacement method for signature 'MOFA,data.frame'
samples_metadata(object) <- value</pre>
```

# Arguments

object a MOFA object.

value data frame with sample metadata, it must at least contain the columns sample

and group. The order of the rows must match the order of samples\_names(object)

#### Value

a data frame with sample metadata

samples\_names 85

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
samples_metadata(model)</pre>
```

samples\_names

samples\_names: set and retrieve sample names

# Description

samples\_names: set and retrieve sample names

# Usage

```
samples_names(object)
samples_names(object) <- value

## S4 method for signature 'MOFA'
samples_names(object)

## S4 replacement method for signature 'MOFA,list'
samples_names(object) <- value</pre>
```

# Arguments

object a MOFA object.

value list of character vectors with the sample names for every group

#### Value

list of character vectors with the sample names for each group

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
samples_names(model)</pre>
```

86 set\_covariates

select_model	Select a model from a list of trained MOFA objects based on the best
	ELBO value

# **Description**

Different objects of MOFA are compared in terms of the final value of the ELBO statistics and the model with the highest ELBO value is selected.

# Usage

```
select_model(models, plot = FALSE)
```

### **Arguments**

models a list containing MOFA objects.

plot boolean indicating whether to show a plot of the ELBO for each model instance

### Value

A MOFA object

set\_covariates Add covariates to a MOFA model

### **Description**

Function to add continuous covariate(s) to a MOFA object for training with MEFISTO

# Usage

```
set_covariates(object, covariates)
```

# **Arguments**

object an untrained MOFA

covariates Sample-covariates to be passed to the model. This can be either:

- a character, specifying columns already present in the samples\_metadata of the object
- a data.frame with columns "sample", "covariate", "value". Sample names need to match those present in the data
- a matrix with smaples in columns and covariate(s) in row(s)

Note that the covariate should be numeric and continuous.

subset\_factors 87

# **Details**

To activate the functional MEFISTO framework, specify mefisto\_options when preparing the training using prepare\_mofa

### Value

Returns an untrained MOFA with covariates filled in the corresponding slots

# **Examples**

```
#' # Simulate data
dd <- make_example_data(sample_cov = seq(0,1,length.out = 100), n_samples = 100, n_factors = 4)
# Create MOFA object
sm <- create_mofa(data = dd$data)
# Add a covariate
sm <- set_covariates(sm, covariates = dd$sample_cov)
sm</pre>
```

subset\_factors

Subset factors

# **Description**

Method to subset (or sort) factors

# Usage

```
subset_factors(object, factors, recalculate_variance_explained = TRUE)
```

# **Arguments**

object a MOFA object.

factors character vector with the factor names, or numeric vector with the index of the

factors.

recalculate\_variance\_explained

logical indicating whether to recalculate variance explained values. Default is

TRUE.

#### Value

A MOFA object

88 subset\_groups

### **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Subset factors 1 to 3
model <- subset_factors(model, factors = 1:3)</pre>
```

subset\_features

Subset features

### **Description**

Method to subset (or sort) features

### Usage

```
subset_features(object, view, features)
```

# **Arguments**

object a MOFA object.

view character vector with the view name or integer with the view index

features character vector with the sample names, numeric vector with the feature indices

or logical vector with the samples to be kept as TRUE.

# Value

A MOFA object

subset\_groups

Subset groups

### **Description**

Method to subset (or sort) groups

# Usage

```
subset_groups(object, groups)
```

# **Arguments**

object a MOFA object.

groups character vector with the groups names, numeric vector with the groups indices

or logical vector with the groups to be kept as TRUE.

subset\_samples 89

# Value

```
A MOFA object
```

# **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Subset the first group
model <- subset_groups(model, groups = 1)</pre>
```

subset\_samples

Subset samples

# **Description**

Method to subset (or sort) samples

### Usage

```
subset_samples(object, samples)
```

# **Arguments**

object a MOFA object.

samples character vector with the sample names or numeric vector with the sample in-

dices.

### Value

```
A MOFA object
```

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
# (TO-DO) Remove a specific sample from the model (an outlier)</pre>
```

90 summarise\_factors

subset\_views

Subset views

### **Description**

Method to subset (or sort) views

### Usage

```
subset_views(object, views)
```

# **Arguments**

object a MOFA object.

views character vector with the views names, numeric vector with the views indices,

or logical vector with the views to be kept as TRUE.

### Value

A MOFA object

# Examples

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)

# Subset the first view
model <- subset_views(model, views = 1)</pre>
```

 $summarise\_factors$ 

Summarise factor values using external groups

# Description

Function to summarise factor values using a discrete grouping of samples.

# Usage

```
summarise_factors(
  object,
  df,
  factors = "all",
  groups = "all",
  abs = FALSE,
  return_data = FALSE
)
```

views\_names 91

# Arguments

object	a trained MOFA object.
df	a data.frame with the columns "sample" and "level", where level is a factor with discrete group assignments for each sample.
factors	character vector with the factor name(s), or numeric vector with the index of the factor(s) to use. Default is 'all'.
groups	character vector with the groups names, or numeric vector with the indices of the groups of samples to use, or "all" to use samples from all groups.
abs	logical indicating whether to take the absolute value of the factors (default is $\ensuremath{FALSE}\xspace).$
return_data	logical indicating whether to return the fa instead of plotting

# Value

A ggplot object or a data. frame if return\_data is TRUE

views_names	views_names: set and retrieve view names

# Description

views\_names: set and retrieve view names

# Usage

```
views_names(object)
views_names(object) <- value

## S4 method for signature 'MOFA'
views_names(object)

## S4 replacement method for signature 'MOFA, character'
views_names(object) <- value</pre>
```

# Arguments

object a MOFA object.

value character vector with the names for each view

#### Value

character vector with the names for each view

92

# **Examples**

```
# Using an existing trained model on simulated data
file <- system.file("extdata", "model.hdf5", package = "MOFA2")
model <- load_model(file)
views_names(model)
views_names(model) <- c("viewA", "viewB")</pre>
```

%>%

Re-exporting the pipe operator See magrittr::Rpercent>Rpercent for details.

# Description

Re-exporting the pipe operator See magrittr::%>% for details.

# Usage

```
lhs %>% rhs
```

# **Arguments**

```
lhs see magrittr::%>%
rhs see magrittr::%>%
```

### Value

depending on lhs and rhs

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