

Package ‘mlr3fda’

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Title Extending 'mlr3' to Functional Data Analysis

Version 0.5.0

Description Extends the 'mlr3' ecosystem to functional analysis by adding support for irregular and regular functional data as defined in the 'tf' package. The package provides 'PipeOps' for preprocessing functional columns and for extracting scalar features, thereby allowing standard machine learning algorithms to be applied afterwards. Available operations include simple functional features such as the mean or maximum, smoothing, interpolation, flattening, and functional 'PCA'.

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URL <https://mlr3fda.mlr-org.com>, <https://github.com/mlr-org/mlr3fda>

BugReports <https://github.com/mlr-org/mlr3fda/issues>

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'PipeOpFDASmooth.R' 'PipeOpFDATsfeatures.R'
'PipeOpFDAWavelets.R' 'PipeOpFDAZoom.R' 'PipeOpFPCA.R'
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mlr3fda-package

mlr3fda: Extending 'mlr3' to Functional Data Analysis

Description

Extends the 'mlr3' ecosystem to functional analysis by adding support for irregular and regular functional data as defined in the 'tf' package. The package provides 'PipeOps' for preprocessing functional columns and for extracting scalar features, thereby allowing standard machine learning algorithms to be applied afterwards. Available operations include simple functional features such as the mean or maximum, smoothing, interpolation, flattening, and functional 'PCA'.

Data types

To extend mlr3 to functional data, two data types from the tf package are added:

- `tfd_irreg` - Irregular functional data, i.e. the functions are observed for potentially different inputs for each observation.
- `tfd_reg` - Regular functional data, i.e. the functions are observed for the same input for each individual.

Lang M, Binder M, Richter J, Schratz P, Pfisterer F, Coors S, Au Q, Casalicchio G, Kotthoff L, Bischl B (2019). “mlr3: A modern object-oriented machine learning framework in R.” *Journal of Open Source Software*. doi:10.21105/joss.01903. <https://joss.theoj.org/papers/10.21105/joss.01903>.

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See Also

Useful links:

- <https://mlr3fda.mlr-org.com>
- <https://github.com/mlr-org/mlr3fda>
- Report bugs at <https://github.com/mlr-org/mlr3fda/issues>

mlr_pipeops_fda.bsignal

B-spline Feature Extraction

Description

This PipeOp extracts features from functional data using B-spline basis functions. The extracted features are B-spline coefficients that represent the functional data in the B-spline basis space. For more details, see `FDbboost::bsignal()`, which is called internally.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `inS :: character(1)`
Type of effect in the covariate index: one of "smooth", "linear", "constant". Default is "smooth".
- `knots :: numeric()`
Either the number of interior knots or a vector of their positions.
- `boundary.knots :: numeric(2)`
Boundary points at which to anchor the B-spline basis. Lower and upper boundary points for the spline basis. Defaults to the range of the data.
- `degree :: integer(1)`
The degree of the regression spline. Default is 3L.
- `differences :: integer(1)`
Order of difference penalty. Default is 1L.
- `df :: numeric(1)`
Trace of the hat matrix, controlling smoothness. Default is 4.
- `lambda :: any`
Smoothing parameter of the penalty term.
- `center :: logical(1)`
Reparameterize the unpenalized part to zero-mean? Default is FALSE.
- `cyclic :: logical(1)`
If true the fitted coefficient function coincides at the boundaries.
- `Z :: any`
Custom transformation matrix for the spline design.
- `penalty :: character(1)`
The penalty type: "ps" (P-spline) or "pss" (shrinkage). Default is "ps".
- `check.ident :: logical(1)`
Use checks for identifiability of the effect. Default is FALSE.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDABsignal
```

Methods

Public methods:

- `PipeOpFDABsignal$new()`
- `PipeOpFDABsignal$clone()`

`PipeOpFDABsignal$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDABsignal$new(id = "fda.bsignal", param_vals = list())
```

Arguments:

id (character(1))

Identifier of resulting object, default "fda.bsignal".

param_vals (named list())

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default list().

PipeOpFDABsignal\$clone(): The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDABsignal$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_bsignal = po("fda.bsignal")
task_bsignal = po_bsignal$train(list(task))[[1L]]
task_bsignal$data()
```

mlr_pipeops_fda.cor *Cross-Correlation of Functional Data*

Description

Calculates the cross-correlation between two functional vectors using `tf::tf_crosscor()`. Note that it only operates on regular data and that the cross-correlation assumes that each column has the same domain.

To apply this PipeOp to irregular data, convert it to a regular grid first using `PipeOpFDAInterpol`. If you need to change the domain of the columns, use `PipeOpFDAScaleRange`.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `arg::numeric()`
Grid to use for the cross-correlation.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDACor
```

Methods

Public methods:

- [PipeOpFDACor\\$new\(\)](#)
- [PipeOpFDACor\\$clone\(\)](#)

`PipeOpFDACor$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDACor$new(id = "fda.cor", param_vals = list())
```

Arguments:

`id` (character(1))

Identifier of resulting object, default "fda.cor".

`param_vals` (named list())

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default list().

`PipeOpFDACor$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDACor$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
set.seed(1234L)
dt = data.table(y = 1:100, x1 = tf::tf_rgp(100L), x2 = tf::tf_rgp(100L))
task = as_task_regr(dt, target = "y")
po_cor = po("fda.cor")
task_cor = po_cor$train(list(task))[[1L]]
task_cor
```

mlr_pipeops_fda.extract

Extract Simple Features from Functional Columns

Description

This is the class that extracts simple features from functional columns. Note that it only operates on values that were actually observed and does not interpolate.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `drop :: logical(1)`
Whether to drop the original functional features and only keep the extracted features. Note that this does not remove the features from the backend, but only from the active column role feature. Initial value is `TRUE`.
- `features :: list() | character()`
A list of features to extract. Each element can be either a function or a string. If the element is a function it requires the following arguments: `arg` and `value` and returns a numeric. For string elements, the following predefined features are available: "mean", "max", "min", "slope", "median", "var". Initial value is `c("mean", "max", "min", "slope", "median", "var")`.
- `left :: numeric()`
The left boundary of the window. Initial value is `-Inf`. The window is specified such that all values `>=left` and `<=right` are kept for the computations.
- `right :: numeric()`
The right boundary of the window. Initial value is `Inf`.

Naming

The new names generally append a `_feature` to the corresponding column name. However this can lead to name clashes with existing columns. This is solved as follows: If a column was called "x" and the feature is "mean", the corresponding new column will be called "x_mean". In case of duplicates, unique names are obtained using `make.unique()` and a warning is given.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAExtract
```

Methods

Public methods:

- `PipeOpFDAExtract$new()`
- `PipeOpFDAExtract$clone()`

`PipeOpFDAExtract$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAExtract$new(id = "fda.extract", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.extract".

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDAExtract$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAExtract$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_fmean = po("fda.extract", features = "mean")
task_fmean = po_fmean$train(list(task))[[1L]]

# add more than one feature
pop = po("fda.extract", features = c("mean", "median", "var"))
task_features = pop$train(list(task))[[1L]]

# add a custom feature
po_custom = po("fda.extract",
  features = list(mean = function(arg, value) mean(value, na.rm = TRUE))
)
task_custom = po_custom$train(list(task))[[1L]]
task_custom
```

```
mlr_pipeops_fda.flatten
```

Flatten Functional Columns

Description

Convert regular functional features (e.g. all individuals are observed at the same time-points) to new columns, one for each input value to the function.

Parameters

The parameters are the parameters inherited from [PipeOpTaskPreprocSimple](#).

Naming

The new names generally append `_1`, `_2`, ... to the corresponding column name. However this can lead to name clashes with existing columns. This is solved as follows: If a column was called "x", the corresponding new columns will be called "x_1", "x_2", etc. In case of duplicates, unique names are obtained using `make.unique()` and a warning is given.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAFlatten
```

Methods**Public methods:**

- [PipeOpFDAFlatten\\$new\(\)](#)
- [PipeOpFDAFlatten\\$clone\(\)](#)

`PipeOpFDAFlatten$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAFlatten$new(id = "fda.flatten", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.flatten".

`param_vals` (named `list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDAFlatten$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAFlatten$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
pop = po("fda.flatten")
task_flat = pop$train(list(task))[[1L]]
```

mlr_pipeops_fda.fourier

Fast Fourier Transform Features

Description

This `PipeOp` extracts Fourier coefficients from functional columns. For more details, see [stats::fft\(\)](#), which is called internally. Only the one-sided spectrum is returned since the input is real-valued (Oppenheim and Schaffer, 2010).

Parameters

The parameters are the parameters inherited from [PipeOpTaskPreprocSimple](#), as well as the following parameters:

- `type` :: `character(1)`
Which feature to extract from the Fourier coefficients. "amplitude" returns the magnitude. "phase" returns the phase shift in degrees (values in [-180, 180]). Initial value is "phase".

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAFourier
```

Methods**Public methods:**

- `PipeOpFDAFourier$new()`
- `PipeOpFDAFourier$clone()`

`PipeOpFDAFourier$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAFourier$new(id = "fda.fourier", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.fourier".

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDAFourier$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAFourier$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

References

Oppenheim, V A, Schafer, W R (2010). *Discrete-Time Signal Processing*, 3rd edition. Pearson. ISBN 978-0131988422.

Examples

```
task = tsk("fuel")
po_fourier = po("fda.fourier")
task_fourier = po_fourier$train(list(task))[[1L]]
task_fourier$data()
```

mlr_pipeops_fda.fpca *Functional Principal Component Analysis*

Description

This PipeOp applies a functional principal component analysis (FPCA) to functional columns and then extracts the principal components as features. This is done using a (truncated) weighted SVD. To apply this PipeOp to irregular data, convert it to a regular grid first using [PipeOpFDAInterpol](#). For more details, see `tf::tfb_fpc()`, which is called internally.

Parameters

The parameters are the parameters inherited from [PipeOpTaskPreproc](#), as well as the following parameters:

- `pve :: numeric(1)`
The percentage of variance explained that should be retained. Default is 0.995.
- `n_components :: integer(1)`
The number of principal components to extract. This parameter is initialized to Inf.

Naming

The new names generally append a `_pc_{number}` to the corresponding column name. If a column was called "x" and there are three principal components, the corresponding new columns will be called "x_pc_1", "x_pc_2", "x_pc_3".

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> PipeOpFPCA
```

Methods

Public methods:

- [PipeOpFPCA\\$new\(\)](#)
- [PipeOpFPCA\\$clone\(\)](#)

`PipeOpFPCA$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFPCA$new(id = "fda.fpca", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.fpca".

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

PipeOpFPCA\$clone(): The objects of this class are cloneable with this method.

Usage:

```
PipeOpFPCA$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_fpca = po("fda.fpca", n_components = 3L)
task_fpca = po_fpca$train(list(task))[[1L]]
task_fpca$data()
```

```
mlr_pipeops_fda.interpol
```

Interpolate Functional Columns

Description

Interpolate functional features (e.g. all individuals are observed at different time-points) to a common grid. This is useful if you want to compare functional features across observations. The interpolation is done using the `tf` package. See `tfd()` for details.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `grid::character(1) | numeric()`
The grid to use for interpolation. If `grid` is numeric, it must be a sequence of values to use for the grid or a single value that specifies the number of points to use for the grid, requires `left` and `right` to be specified in the latter case. If `grid` is a character, it must be one of:
 - `"union"`: This option creates a grid based on the union of all argument points from the provided functional features. This means that if the argument points across features are $\setminus(t_1, t_2, \dots, t_n)$, then the grid will be the combined unique set of these points. This option is generally used when the argument points vary across observations and a common grid is needed for comparison or further analysis.
 - `"intersect"`: Creates a grid using the intersection of all argument points of a feature. This grid includes only those points that are common across all functional features, facilitating direct comparison on a shared set of points.
 - `"minmax"`: Generates a grid within the range of the maximum of the minimum argument points to the minimum of the maximum argument points across features. This bounded grid encapsulates the argument point range common to all features. Note: For regular functional data this has no effect as all argument points are the same. Initial value is `"union"`.

- `method` :: character(1)
One of:
 - "linear": applies linear interpolation without extrapolation (see `tf::tf_approx_linear()`).
 - "spline": applies cubic spline interpolation (see `tf::tf_approx_spline()`).
 - "fill_extend": applies linear interpolation with constant extrapolation (see `tf::tf_approx_fill_extend()`).
 - "locf": applies "last observation carried forward" interpolation (see `tf::tf_approx_locf()`).
 - "nocb": applies "next observation carried backward" interpolation (see `tf::tf_approx_nocb()`).
 Default is "linear".
- `left` :: numeric()
The left boundary of the window. The window is specified such that all values \geq left and \leq right are kept for the computations.
- `right` :: numeric()
The right boundary of the window.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAInterpol
```

Methods

Public methods:

- `PipeOpFDAInterpol$new()`
- `PipeOpFDAInterpol$clone()`

`PipeOpFDAInterpol$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAInterpol$new(id = "fda.interpol", param_vals = list())
```

Arguments:

`id` (character(1))

Identifier of resulting object, default "fda.interpol".

`param_vals` (named list())

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default list().

`PipeOpFDAInterpol$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAInterpol$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
pop = po("fda.interpol")
task_interpol = pop$train(list(task))[[1L]]
task_interpol$data()
```

mlr_pipeops_fda.random_effect

Extract Random Effects from Functional Columns

Description

This is the class that extracts random effects, specifically random intercepts and random slopes, from functional columns. This PipeOp fits a linear mixed model, specifically a random intercept and random slope model, using the `lme4::lmer()` function. The target variable is the value of the functional feature which is regressed on the functional feature's argument while subject id determines the grouping structure. After model estimation, the random effects are extracted and assigned to the correct id.

Parameters

The parameters are the parameters inherited from [PipeOpTaskPreprocSimple](#).

Naming

The new names append `_random_intercept` and `_random_slope` to the corresponding column name of the functional feature.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDARandomEffect
```

Methods

Public methods:

- [PipeOpFDARandomEffect\\$new\(\)](#)
- [PipeOpFDARandomEffect\\$clone\(\)](#)

`PipeOpFDARandomEffect$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDARandomEffect$new(id = "fda.random_effect", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default `"fda.random_effect"`.

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDARandomEffect$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDARandomEffect$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("dti")
po_fre = po("fda.random_effect")
task_fre = po_fre$train(list(task))[[1L]]
```

```
mlr_pipeops_fda.register
```

Register (Align) Functional Columns

Description

Aligns functional features by estimating warping functions against a template using `tf::tf_register()`. Registration reduces phase (horizontal) variability while preserving amplitude (vertical) variability, which is useful when curves share a common shape but differ in the timing of their features.

During training, a template is learned for each functional column (either estimated iteratively from the data or supplied via `args`). The template is stored as part of the `$state` and reused at predict time so that new observations are aligned to the same reference as the training data.

Supported methods are "srvf", "cc", and "affine". The "landmark" method from `tf::tf_register()` is not supported because it requires per-observation landmark positions for both training and prediction data, which does not fit a stateful preprocessing step. For landmark registration, use `tf::tf_register()` directly and feed the aligned data into the task.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreproc`, as well as the following parameters:

- `method` :: character(1)
Registration method. One of:
 - "srvf": elastic registration using the Square Root Velocity Framework via `fdasrvf::time_warping()`. Requires regular grids and the **fdasrvf** package. Default template is the Karcher mean.
 - "cc": continuous-criterion registration with monotone spline warps. Requires regular grids. Default template is the arithmetic mean.
 - "affine": affine (shift and/or scale) registration with warps of the form $h(t) = a \cdot t + b$. Supports regular and irregular grids. Default template is the arithmetic mean.
 Default is "srvf".
- `args` :: named list()
Method-specific arguments passed to `tf::tf_register()` via `...`. See the help page of `tf::tf_estimate_warps()` for valid arguments (e.g. `lambda` / `penalty_method` for "srvf"; `nbasis`, `lambda`, `crit`, `conv`, `iterlim` for "cc"; `type`, `shift_range`, `scale_range` for "affine"). An optional template entry is honored at training time and stored in the state.
- `max_iter` :: integer(1)
Maximum number of Procrustes-style template refinement iterations during training. Default is 3. Ignored at predict time because the stored template is used directly.
- `tol` :: numeric(1)
Convergence tolerance for template refinement during training. Default is 0.01.

State

`$state$templates` contains the learned template (as a length-1 `tf` vector) for each functional column.

Super classes

`mlr3pipelines::PipeOp` -> `mlr3pipelines::PipeOpTaskPreproc` -> `PipeOpFDARegister`

Methods**Public methods:**

- `PipeOpFDARegister$new()`
- `PipeOpFDARegister$clone()`

`PipeOpFDARegister$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDARegister$new(id = "fda.register", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default `"fda.register"`.

`param_vals` (named `list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDARegister$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDARegister$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
set.seed(1)
task = tsk("fuel")
po_reg = po("fda.register", method = "affine", args = list(type = "shift_scale"))
task_reg = po_reg$train(list(task))[[1L]]
task_reg$data(cols = c("NIR", "UVVIS"))
```

mlr_pipeops_fda.scalerange

Linearly Transform the Domain of Functional Data

Description

Linearly transform the domain of functional data so they are between lower and upper. The formula for this is $x' = offset + x * scale$, where $scale$ is $(upper - lower) / (max(x) - min(x))$ and $offset$ is $-min(x) * scale + lower$. The same transformation is applied during training and prediction.

Parameters

The parameters are the parameters inherited from [PipeOpTaskPreproc](#), as well as the following parameters:

- `lower :: numeric(1)`
Target value of smallest item of input data. Initialized to 0.
- `upper :: numeric(1)`
Target value of greatest item of input data. Initialized to 1.

Super classes

`mlr3pipelines::PipeOp` -> `mlr3pipelines::PipeOpTaskPreproc` -> `PipeOpFDAScaleRange`

Methods

Public methods:

- `PipeOpFDAScaleRange$new()`
- `PipeOpFDAScaleRange$clone()`

`PipeOpFDAScaleRange$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAScaleRange$new(id = "fda.scalerange", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.scalerange".

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDAScaleRange$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAScaleRange$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_scale = po("fda.scalerange", lower = -1, upper = 1)
task_scale = po_scale$train(list(task))[[1L]]
task_scale$data()
```

mlr_pipeops_fda.smooth

Smooth Functional Columns

Description

Smooths functional data using `tf::tf_smooth()`. This preprocessing operator is similar to `PipeOpFDAInterpol`, however it does not interpolate to unobserved x-values, but rather smooths the observed values.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `method` :: character(1)
One of:
 - "lowess": locally weighted scatterplot smoothing (default)
 - "rollmean": rolling mean
 - "rollmedian": rolling median
 - "savgol": Savitzky-Golay filtering
 All methods but "lowess" ignore non-equidistant arg values.
- `args` :: named list()
List of named arguments that is passed to `tf_smooth()`. See the help page of `tf_smooth()` for default values.
- `verbose` :: logical(1)
Whether to print messages during the transformation. Is initialized to FALSE.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDASmooth
```

Methods**Public methods:**

- `PipeOpFDASmooth$new()`
- `PipeOpFDASmooth$clone()`

`PipeOpFDASmooth$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDASmooth$new(id = "fda.smooth", param_vals = list())
```

Arguments:

```
id (character(1))
```

Identifier of resulting object, default "fda.smooth".

```
param_vals (named list())
```

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default list().

`PipeOpFDASmooth$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDASmooth$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_smooth = po("fda.smooth", method = "rollmean", args = list(k = 5))
task_smooth = po_smooth$train(list(task))[[1L]]
task_smooth
task_smooth$data(cols = c("NIR", "UVVIS"))
```

```
mlr_pipeops_fda.tsfeats
```

Time Series Feature Extraction

Description

This PipeOp extracts time series features from functional columns.

For more details, see `tsfeatures::tsfeatures()`, which is called internally.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `features::character()`
Function names which return numeric vectors of features. All features returned by these functions must be named if they return more than one feature. Default is `c("frequency", "stl_features", "entropy", "acf_features")`.
- `scale::logical(1)`
If TRUE, data is scaled to mean 0 and sd 1 before features are computed. Default is TRUE.
- `trim::logical(1)`
If TRUE, data is trimmed by `trim_amount` before features are computed. Values larger than `trim_amount` in absolute value are set to NA. Default is FALSE.

- `trim_amount` :: `numeric(1)`
Default level of trimming. Default is 0.1.
- `parallel` :: `logical(1)`
If TRUE, the features are computed in parallel. Default is FALSE.
- `multiprocess` :: `any`
The function from the future package to use for parallel processing. Default is `future::multisession()`.
- `na.action` :: `any`
A function to handle missing values. Default is `stats::na.pass()`.

Naming

The new names generally append a `_{feature}` to the corresponding column name. If a column was called "x" and the feature is "trend", the corresponding new column will be called "x_trend".

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDATsfeatures
```

Methods

Public methods:

- `PipeOpFDATsfeatures$new()`
- `PipeOpFDATsfeatures$clone()`

`PipeOpFDATsfeatures$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDATsfeatures$new(id = "fda.tsfeats", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.tsfeats".

`param_vals` (`named list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDATsfeatures$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDATsfeatures$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_tsfeats = po("fda.tsfeats")
task_tsfeats = po_tsfeats$train(list(task))[[1L]]
task_tsfeats$data()
```

mlr_pipeops_fda.wavelets

Discrete Wavelet Transform Features

Description

This PipeOp extracts discrete wavelet transform coefficients from functional columns. For more details, see `wavelets::dwt()`, which is called internally.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `filter` :: `character(1) | numeric() | wavelets::wt.filter()`
Specifies which filter should be used. Must be either `wavelets::wt.filter()` object, an even numeric vector or a string. In case of a string must be one of "d"|"la"|"b1"|"c" followed by an even number for the level of the filter. The level of the filter needs to be smaller or equal than the time-series length. For more information and acceptable filters see `help(wt.filter)`. Default is "la8".
- `n.levels` :: `integer(1)`
An integer specifying the level of the decomposition.
- `boundary` :: `character(1)`
Boundary to be used. "periodic" assumes circular time series, for "reflection" the series is extended to twice its length. Default is "periodic".
- `fast` :: `logical(1)`
Should the pyramid algorithm be calculated with an internal C function? Default is TRUE.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAWavelets
```

Methods

Public methods:

- `PipeOpFDAWavelets$new()`
- `PipeOpFDAWavelets$clone()`

`PipeOpFDAWavelets$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAWavelets$new(id = "fda.wavelets", param_vals = list())
```

Arguments:

`id` (`character(1)`)

Identifier of resulting object, default "fda.wavelets".

`param_vals` (named `list()`)

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default `list()`.

`PipeOpFDAWavelets$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAWavelets$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
po_wavelets = po("fda.wavelets")
task_wavelets = po_wavelets$train(list(task))[[1L]]
task_wavelets$data()
```

mlr_pipeops_fda.zoom *Zoom In/Out on Functional Columns*

Description

Zoom in or out on functional features by restricting their domain to a specified window. This operation extracts a subset of each function by defining new lower and upper boundaries, effectively cropping the functional data to focus on a specific region of interest. Calls `tf::tf_zoom()` from package `tf`.

Parameters

The parameters are the parameters inherited from `PipeOpTaskPreprocSimple`, as well as the following parameters:

- `begin` :: `numeric()`
The lower limit of the domain. Can be a single value applied to all functional columns, or a numeric of length equal to the number of observations. The window includes all values where argument `>= begin`. If not specified, defaults to the lower limit of each function's domain.
- `end` :: `numeric()`
The upper limit of the domain.

Super classes

```
mlr3pipelines::PipeOp -> mlr3pipelines::PipeOpTaskPreproc -> mlr3pipelines::PipeOpTaskPreprocSimple
-> PipeOpFDAZoom
```

Methods

Public methods:

- [PipeOpFDAZoom\\$new\(\)](#)
- [PipeOpFDAZoom\\$clone\(\)](#)

`PipeOpFDAZoom$new()`: Initializes a new instance of this Class.

Usage:

```
PipeOpFDAZoom$new(id = "fda.zoom", param_vals = list())
```

Arguments:

`id` (character(1))

Identifier of resulting object, default "fda.zoom".

`param_vals` (named list())

List of hyperparameter settings, overwriting the hyperparameter settings that would otherwise be set during construction. Default list().

`PipeOpFDAZoom$clone()`: The objects of this class are cloneable with this method.

Usage:

```
PipeOpFDAZoom$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Examples

```
task = tsk("fuel")
pop = po("fda.zoom", begin = 50, end = 100)
task_zoom = pop$train(list(task))[[1L]]
task_zoom$data()
```

 mlr_tasks_dti

Diffusion Tensor Imaging (DTI) Regression Task

Description

This dataset contains two functional covariates and three scalar covariates. The goal is to predict the PASAT score. `pasat` represents the PASAT score at each visit. `subject_id` represents the subject ID. `cca` represents the fractional anisotropy tract profiles from the corpus callosum. `sex` indicates subject's sex. `rcst` represents the fractional anisotropy tract profiles from the right corticospinal tract. Rows containing NAs are removed.

This is a subset of the full dataset, which is contained in the package `refund`.

Format

`R6::R6Class` inheriting from `mlr3::TaskRegr`.

Dictionary

This `mlr3::Task` can be instantiated via the dictionary `mlr3::mlr_tasks` or with the associated sugar function `mlr3::tsk()`:

```
mlr_tasks$get("dti")
tsk("dti")
```

Meta Information

- Task type: “regr”
- Dimensions: 340x4
- Properties: “groups”
- Has Missings: FALSE
- Target: “pasat”
- Features: “cca”, “rest”, “sex”

References

Goldsmith, Jeff, Bobb, Jennifer, Crainiceanu, M C, Caffo, Brian, Reich, Daniel (2011). “Penalized functional regression.” *Journal of Computational and Graphical Statistics*, **20**(4), 830–851.

Brain dataset courtesy of Gordon Kindlmann at the Scientific Computing and Imaging Institute, University of Utah, and Andrew Alexander, W. M. Keck Laboratory for Functional Brain Imaging and Behavior, University of Wisconsin-Madison.

See Also

- Chapter in the `mlr3book`: https://mlr3book.mlr-org.com/chapters/chapter2/data_and_basic_modeling.html
- Package `mlr3data` for more toy tasks.
- Package `mlr3oml` for downloading tasks from <https://www.openml.org>.
- Package `mlr3viz` for some generic visualizations.
- Dictionary of Tasks: `mlr3::mlr_tasks`
- `as.data.table(mlr_tasks)` for a table of available Tasks in the running session (depending on the loaded packages).
- `mlr3fselect` and `mlr3filters` for feature selection and feature filtering.
- Extension packages for additional task types:
 - Unsupervised clustering: `mlr3cluster`
 - Probabilistic supervised regression and survival analysis: <https://mlr3proba.mlr-org.com/>.

Other Task: `mlr_tasks_fuel`, `mlr_tasks_phoneme`

mlr_tasks_fuel	<i>Fuel Regression Task</i>
----------------	-----------------------------

Description

This dataset contains two functional covariates and one scalar covariate. The goal is to predict the heat value of some fuel based on the ultraviolet radiation spectrum and infrared ray radiation and one scalar column called h2o.

This is a subset of the full dataset, which is contained in the package FDboost.

Format

R6::R6Class inheriting from mlr3::TaskRegr.

Dictionary

This mlr3::Task can be instantiated via the dictionary mlr3::mlr_tasks or with the associated sugar function mlr3::tsk():

```
mlr_tasks$get("fuel")
tsk("fuel")
```

Meta Information

- Task type: “regr”
- Dimensions: 129x4
- Properties: -
- Has Missings: FALSE
- Target: “heatan”
- Features: “NIR”, “UVVIS”, “h2o”

References

Brockhaus, Sarah, Scheipl, Fabian, Hothorn, Torsten, Greven, Sonja (2015). “The functional linear array model.” *Statistical Modelling*, **15**(3), 279–300.

See Also

- Chapter in the mlr3book: https://mlr3book.mlr-org.com/chapters/chapter2/data_and_basic_modeling.html
- Package **mlr3data** for more toy tasks.
- Package **mlr3oml** for downloading tasks from <https://www.openml.org>.
- Package **mlr3viz** for some generic visualizations.
- Dictionary of Tasks: mlr3::mlr_tasks

- `as.data.table(mlr_tasks)` for a table of available **Tasks** in the running session (depending on the loaded packages).
- **mlr3fselect** and **mlr3filters** for feature selection and feature filtering.
- Extension packages for additional task types:
 - Unsupervised clustering: **mlr3cluster**
 - Probabilistic supervised regression and survival analysis: <https://mlr3proba.mlr-org.com/>.

Other Task: `mlr_tasks_dti`, `mlr_tasks_phoneme`

`mlr_tasks_phoneme` *Phoneme Classification Task*

Description

The task contains a single functional covariate and 5 equally big classes (aa, ao, dcl, iy, sh). The aim is to predict the class of the phoneme from the functional feature, which is a log-periodogram. This is a subset of the full dataset, which is contained in the package `fda.usc`.

Format

`R6::R6Class` inheriting from `mlr3::TaskClassif`.

Dictionary

This `mlr3::Task` can be instantiated via the dictionary `mlr3::mlr_tasks` or with the associated sugar function `mlr3::tsk()`:

```
mlr_tasks$get("phoneme")
tsk("phoneme")
```

Meta Information

- Task type: “classif”
- Dimensions: 250x2
- Properties: “multiclass”
- Has Missings: FALSE
- Target: “class”
- Features: “X”

References

Ferraty, Frédéric, Vieu, Philippe (2003). “Curves discrimination: a nonparametric functional approach.” *Computational Statistics & Data Analysis*, **44**(1-2), 161–173.

See Also

- Chapter in the **mlr3book**: https://mlr3book.mlr-org.com/chapters/chapter2/data_and_basic_modeling.html
- Package **mlr3data** for more toy tasks.
- Package **mlr3oml** for downloading tasks from <https://www.openml.org>.
- Package **mlr3viz** for some generic visualizations.
- **Dictionary of Tasks**: `mlr3::mlr_tasks`
- `as.data.table(mlr_tasks)` for a table of available **Tasks** in the running session (depending on the loaded packages).
- **mlr3fselect** and **mlr3filters** for feature selection and feature filtering.
- Extension packages for additional task types:
 - Unsupervised clustering: **mlr3cluster**
 - Probabilistic supervised regression and survival analysis: <https://mlr3proba.mlr-org.com/>.

Other Task: `mlr_tasks_dti`, `mlr_tasks_fuel`

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