Package 'mxfda'

February 19, 2025

Title A Functional Data Analysis Package for Spatial Single Cell Data

Version 0.2.2-1

Date 2025-02-19

Description Methods and tools for deriving spatial summary functions from single-cell imaging data and performing functional data analyses. Functions can be applied to other single-cell technologies such as spatial transcriptomics. Functional regression and functional principal component analysis methods

are in the 'refund' package <https:

//cran.r-project.org/package=refund> while calculation of the
spatial summary functions are from the 'spatstat' package https://spatstat.org/>.

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URL https://github.com/julia-wrobel/mxfda/,

http://juliawrobel.com/mxfda/

BugReports https://github.com/julia-wrobel/mxfda/issues/

Encoding UTF-8

RoxygenNote 7.3.2

- **Suggests** knitr, rmarkdown, testthat (>= 3.0.0), tidyverse, survival, ggpubr, spatialTIME, tibble, broom, refund.shiny, Seurat, SeuratObject
- **Imports** magrittr, rlang, tidyr, purrr, dplyr, ggplot2, lifecycle, methods, stats, refund (>= 0.1-35), reshape2, mgcv, spatstat.geom, spatstat.explore, SpatEntropy, SimDesign

Depends R (>= 2.10),

LazyData true

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Author Julia Wrobel [aut] (<https://orcid.org/0000-0001-6783-1421>), Alex Soupir [aut, cre] (<https://orcid.org/0000-0003-1251-9179>) Maintainer Alex Soupir <alex.soupir@moffitt.org> Repository CRAN Date/Publication 2025-02-19 21:10:02 UTC

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add_summary_function Add Summary Function

Description

Index

Sometimes other ways of calculating summary functions is wanted and is done in other packages, in this instance the data can be loaded into the mxFDA object.

bivariate

Usage

add_summary_function(mxFDAobject, summary_function_data, metric)

Arguments

mxFDAobject	object of class mxFDA	
summary_function_data		
	data frame with summary_key from $mxFDA$ object as key column for summary function	
metric	character vector with either 'uni' or 'bi' and 'k', 'l', or 'g'; e.g. 'uni g'	

Value

an updated mxFDA object with a derived value added. See make_mxfda() for more details.

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

bivariate bivariate

Description

Internal function called by extract_summary_functions to calculate a bivariate spatial summary function for a single image.

Usage

```
bivariate(
  mximg,
  markvar,
  mark1,
  mark2,
  r_vec,
  func = c(Kcross, Lcross, Gcross, entropy),
  edge_correction,
  empirical_CSR = FALSE,
  permutations = 1000
)
```

Arguments

Dataframe of cell-level multiplex imaging data for a single image. Should have variables x and y to denote x and y spatial locations of each cell.		
The name of the variable that denotes cell type(s) of interest. Character.		
Character string that denotes first cell type of interest.		
Character string that denotes second cell type of interest.		
Numeric vector of radii over which to evaluate spatial summary functions. Must begin at 0.		
Spatial summary function to calculate. Options are c(Kcross, Lcross, Gcross) which denote Ripley's K, Besag's L, and nearest neighbor G function, respectively, or entropy from Vu et al, 2023.		
edge_correction		
Character string that denotes the edge correction method for spatial summary function. For Kcross and Lcross choose one of c("border", "isotropic", "Ripley", "translate", "none"). For Gcross choose one of c("rs", "km", "han")		
logical to indicate whether to use the permutations to identify the sample-specific complete spatial randomness (CSR) estimation.		
integer for the number of permtuations to use if empirical_CSR is TRUE and exact CSR not calculable		

Details

[Stable]

Value

A data.frame containing:

r	the radius of values over which the spatial summary function is evaluated
sumfun	the values of the spatial summary function
csr	the values of the spatial summary function under complete spatial randomness
fundiff	sumfun - csr, positive values indicate clustering and negative values repulsion

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421. DOI: 10.1007/s11222-014-9485-x.

Vu, T., Seal, S., Ghosh, T., Ahmadian, M., Wrobel, J., & Ghosh, D. (2023). FunSpace: A functional and spatial analytic approach to cell imaging data using entropy measures. *PLOS Computational Biology*, 19(9), e1011490.

entropy

Creed, J. H., Wilson, C. M., Soupir, A. C., Colin-Leitzinger, C. M., Kimmel, G. J., Ospina, O. E., Chakiryan, N. H., Markowitz, J., Peres, L. C., Coghill, A., & Fridley, B. L. (2021). spatialTIME and iTIME: R package and Shiny application for visualization and analysis of immunofluorescence data. *Bioinformatics* (Oxford, England), 37(23), 4584–4586. https://doi.org/10.1093/bioinformatics/btab757

entropy

Entropy

Description

Entropy

Usage

entropy(df, r_vec, markvar)

Arguments

df	data frame with x and y columns, along with a column for point marks
r_vec	vector of length wanted for breaks (will be rescaled) with max value at max for measuring entropy
markvar	The name of the variable that denotes cell type(s) of interest. Character.

Details

[Experimental]

Value

data frame with entropy calculated for length(r_vec) bins within 0 to max(r_vec)

Author(s)

Thao Vu <thao.3.vu@cuanschutz.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Vu, T., Seal, S., Ghosh, T., Ahmadian, M., Wrobel, J., & Ghosh, D. (2023). FunSpace: A functional and spatial analytic approach to cell imaging data using entropy measures. *PLOS Computational Biology*, 19(9), e1011490.

Altieri, L., Cocchi, D., & Roli, G. (2018). A new approach to spatial entropy measures. *Environmental and ecological statistics*, 25, 95-110.

extract_entropy extract_entropy

Description

The extract_entropy() is used to compute spatial entropy at each distance interval for all cell types of interest. The goal is to capture the diversity in cellular composition, such as similar proportions across cell types or dominance of a single type, at a specific distance range. Additionally, spatial patterns, including clustered, independent, or regular, among cell types can also be acquired. In this example, we will look at the spatial heterogeneity across T cells, macrophages, and others. To focus on the local cell-to-cell interactions, we set the default maximum of the distance range (i.e., rmax) to be 400 microns. The default number of distance breaks/intervals is set to 50. Then, a sequence of distance breaks is generated by linearly decreasing from rmax to 0 on a log scale. At each distance range, partial spatial entropy and residual entropy are calculated as in Vu et al. (2023), Altieri et al. (2018). These spatial entropy functions can then be used as input functions for FPCA.

Usage

```
extract_entropy(mxFDAobject, markvar, marks, n_break = 50, rmax = 400)
```

Arguments

mxFDAobject	object of class mxFDA
markvar	The name of the variable that denotes cell type(s) of interest. Character.
marks	Character vector that denotes cell types of interest.
n_break	Total number of distance ranges/intervals of interest made from 0 to rmax for calculating entropy
rmax	Max distance between pairs of cells

Value

object of class mxFDA with a dataframe in the multivariate_summaries slot

extract_fpca_object Extract FPCA object

Description

Function that extracts the FPCA object created either by run_fpca() or run_mfpca() from the mxFDA object

Usage

extract_fpca_object(mxFDAobject, what)

Arguments

mxFDAobject	object of class mxFDA
what	what functional PCA data to extract, e.g. 'uni k'

Details

[Stable]

Output object can be visualized with refund.shiny::plot_shiny()

Value

fpca object created with run_fcm()

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

```
#load ovarian mxFDA object
data('ovarian_FDA')
```

#extract the fpca object
obj = extract_fpca_object(ovarian_FDA, "uni g fpca")

extract_fpca_scores Extract FPCA scores

Description

Extract FPCA scores

Usage

extract_fpca_scores(mxFDAobject, what)

Arguments

mxFDAobject	object of class mxFDA
what	what functional PCA data to extract, e.g. 'uni k'

Details

[Stable]

Value

fpca object

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

extract_model Extract Model

Description

Currently only extracts functional cox models not mixed functional cox models.

Usage

```
extract_model(mxFDAobject, metric, type, model_name)
```

Arguments

mxFDAobject	object of class mxFDA
metric	metric functional PCA data to extract, e.g. 'uni k'
type	one of "cox", "mcox", or "sofr" to specify the type of model to extract
model_name	character string of the model name to retrieve

Details

[Stable]

Value

fit functional model

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

extract_spatial_summary

Summarise spatial data in mxFDA object

Description

Summarise spatial data in mxFDA object

Usage

```
extract_spatial_summary(mxFDAobject, columns, grouping_columns = NULL)
```

Arguments

mxFDAobject	object of class mxFDA
columns	character vector for column heading for cells to summarise
grouping_colum	ns
	character vector of other columns to use as grouping, such as region classifica- tion column

Details

[Experimental]

Currently this function is experimental as it only handles data that has text in the columns. Eventually, will be able to handle any data inputs such as those from HALO where cells are designated as positive (1) or negative (0) for a cell phenotypes.

Value

data frame with percent of total points per spatial sample columns. If multiple levels are present in columns columns, multiple output columns will be provided.

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

```
#load data
data(lung_df)
#create data frames for `mxFDA` object
clinical = lung_df %>%
 dplyr::select(image_id, patient_id, patientImage_id, gender,
         age, survival_days, survival_status, stage) %>%
 dplyr::distinct()
#make small, just need to make sure it runs
spatial = lung_df %>%
 dplyr::select(-image_id, -gender, -age, -survival_days, -survival_status, -stage) %>%
 dplyr::filter(patientImage_id %in% clinical$patientImage_id[1:10])
#create `mxFDA` object
mxFDAobject = make_mxfda(metadata = clinical,
                         spatial = spatial,
                         subject_key = "patient_id",
                         sample_key = "patientImage_id")
#get markers
markers = colnames(mxFDAobject@Spatial) %>%
 grep("pheno", ., value = TRUE)
#extract summary
df = extract_spatial_summary(mxFDAobject, markers)
```

extract_summary_functions *Extract Summary Functions*

Description

Function to extract spatial summary functions from the Spatial slot of an mxFDA object

Usage

```
extract_summary_functions(
  mxFDAobject,
  r_vec = seq(0, 100, by = 10),
  extract_func = c(univariate, bivariate),
  summary_func = c(Kest, Lest, Gest),
  markvar,
  mark1,
  mark2 = NULL,
  edge_correction,
  empirical_CSR = FALSE,
  permutations = 1000
)
```

Arguments

mxFDAobject	object of class mxFDA	
r_vec	Numeric vector of radii over which to evaluate spatial summary functions. Must begin at 0.	
extract_func	Defaults to univariate, which calculates univariate spatial summary functions. Choose bivariate for bivariate spatial summary functions.	
summary_func	Spatial summary function to calculate. Options are c(Kest, Lest, Gest) which denote Ripley's K, Besag's L, and nearest neighbor G function, respectively.	
markvar	The name of the variable that denotes cell type(s) of interest. Character.	
mark1	Character string that denotes first cell type of interest.	
mark2	Character string that denotes second cell type of interest for calculating bivariate summary statistics. Not used when calculating univariate statistics.	
edge_correction		
	Character string that denotes the edge correction method for spatial summary function. For Kest and Lest choose one of c("border", "isotropic", "Ripley", "translate", "none"). For Gest choose one of c("rs", "km", "han")	
empirical_CSR	logical to indicate whether to use the permutations to identify the sample-specific complete spatial randomness (CSR) estimation. If there are not enough levels present in markvar column for permutations, the theoretical will be used.	
permutations	integer for the number of permtuations to use if empirical_CSR is TRUE and exact CSR not calculable	

Details

[Stable]

Complete spatial randomness (CSR) is the estimation or measure of a spatial summary function when the points or cells in a sample are randomly distributed, following no clustering or dispersion pattern. Some samples do have artifacts that may influence what CSR is under the distribution of points as they are found in the sample such as large regions of missing points or possibly in the case of tissue sections, necrotic tissue where cells are dead. Theoretical CSR requires points have

an equal chance of occurring anywhere in the sample that these artifacts violate, necessitating the need to estimate or calculate what this CSR would be for each sample independently. Previously Wilson et al. had demonstrated cases in which sample-specific CSR was important over the use of the theoretical in calculating how much the observed deviates from expected.

Value

an object of class mxFDA containing the corresponding spatial summary function slot filled. See make_mxfda() for object structure details.

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421. DOI: 10.1007/s11222-014-9485-x.

Wilson, C., Soupir, A. C., Thapa, R., Creed, J., Nguyen, J., Segura, C. M., Gerke, T., Schildkraut, J. M., Peres, L. C., & Fridley, B. L. (2022). Tumor immune cell clustering and its association with survival in African American women with ovarian cancer. PLoS computational biology, 18(3), e1009900. https://doi.org/10.1371/journal.pcbi.1009900

Creed, J. H., Wilson, C. M., Soupir, A. C., Colin-Leitzinger, C. M., Kimmel, G. J., Ospina, O. E., Chakiryan, N. H., Markowitz, J., Peres, L. C., Coghill, A., & Fridley, B. L. (2021). spatialTIME and iTIME: R package and Shiny application for visualization and analysis of immunofluorescence data. *Bioinformatics* (Oxford, England), 37(23), 4584–4586. https://doi.org/10.1093/bioinformatics/btab757

```
spatstat.explore::Kest()
spatstat.explore::Gest()
spatstat.explore::Lest()
spatstat.explore::Kcross()
spatstat.explore::Gcross()
spatstat.explore::Lcross()
```

Examples

```
#load ovarian FDA object
data('ovarian_FDA')
```

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Description

Function that transforms functional models from linear or additive functional cox models into afcmSurface or lfcmSurface objects to be plotted.

Usage

```
extract_surface(
  mxFDAobject,
  metric,
  model = NULL,
  r = "r",
  value = "fundiff",
  grid_length = 100,
  analysis_vars,
  p = 0.05,
  filter_cols = NULL
)
```

Arguments

mxFDAobject	object of class mxFDA with model model calculated wihtin
metric	spatial summary function to extract surface for
model	character string for the name of the model for metric data
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
grid_length	Length of grid on which to evaluate coefficient functions.
analysis_vars	Other variables used in modeling FCM fit.
р	numeric p-value used for predicting significant AFCM surface
filter_cols	a named vector of factors to filter summary functions to in c(Derived_Column = "Level_to_Filter") format

Value

a 4 element list of either class lfcmSurface or afcmSurface depending on the class of model

Surface	data.frame for term predictions for the surface of the metric * radius area
Prediction	data.frame for standard error of the terms for the above surface. AFCM models use the p to set the upper and lower standard errors of β_1

Metric	character of the spatial summary function used; helps keep track if running many models
P-value	a numeric value of the input p-value

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>
Alex Soupir <alex.soupir@moffitt.org>

I I

Examples

```
filter_spatial Filter Spatial data
```

Description

function to filter the spatial data slot of the mxFDA object.

Usage

```
filter_spatial(mxFDAobject, ..., based_on = "meta", force = FALSE)
```

Arguments

mxFDAobject	object of class mxFDA
	expressions that return a logical TRUE/FALSE value when evaluated on columns of the meta data slot. These expressions get passed to dplyr::filter() so must be compatible.
based_on	character for which data slot to use for filtering, either 'meta', or 'spatial'. Default to 'meta'.
force	logical whether or not to return empty spatial data if filtering results in 0 rows

lung_df

Value

object of class mxFDA with the spatial slot filtered. See make_mxfda() for more details on object

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

References

dplyr::filter()

Examples

```
#load ovarian mxFDA object
data(ovarian_FDA)
#filter ages greater than 50
```

ovarian_FDA_age50 = filter_spatial(ovarian_FDA, age >= 50, based_on = 'meta')

lung_df

Multiplex imaging data from a non-small cell lung cancer study.

Description

This data is adapted from the VectraPolarisData Bioconductor package. There are multiple ROIs for each patient. Data was filtered to include only the cells in the tumor compartment.

Usage

lung_df

Format

lung_df:

A data frame with 879,694 rows and 19 columns:

image_id Image id for a given patient

patient_id Unique patient id

age Patient age at time of cancer diagnosis

survival_days Survival time from diagnosis, in days

survival_status Censoring variable, 1 = death, 0 = censor

- **x** Cell x position
- y Cell y position ...

Source

https://bioconductor.org/packages/release/data/experiment/html/VectraPolarisData.
html

lung_FDA

Description

This data is adapted from the VectraPolarisData Bioconductor package. There are multiple ROIs for each patient.

Usage

lung_FDA

Format

lung_FDA:

An mxFDA object with augmented non-small cel lung cancer multiplex immunofluorescence data, and NN G(r) calculated:

Metadata information about the spatial samples with column sample_key column in both

Spatial cell-level information with x and y columns along with sample_key to link to Metadata

- subject_key column in Metadata that may have multiple sample_key values for each, akin to
 patient IDs
- sample_key column in both Metadata and Spatial that is a 1:1 with the samples (unique per sample)

univariate_summaries univariate summary slot with nearest neighbor G calculared

bivariate_summaries empty slot available for bivariate summaries

multiivariate_summaries empty slot available for multivariate summaries

functional_pca empty slot for functional PCA data of summaries

functional_cox empty slot for functional models

Details

Spatial summary functions of lung cancer multiplex imaging data.

This data is adapted from the VectraPolarisData Bioconductor package. Signal between the survival outcome and spatial summary functions has been augmented for teaching purposes. Spatial relationship is summarized using the nearest neighbor G function.

Includes only spatial samples that had 10 or more radii with calculable G function

Source

https://bioconductor.org/packages/release/data/experiment/html/VectraPolarisData.
html

make_mxfda

Description

Used to create an object of class mxFDA that can be used with the mxfda package for functional data analysis.

Usage

make_mxfda(metadata, spatial = NULL, subject_key, sample_key)

Arguments

metadata	metadata with columns subject_key and sample_key	
spatial	spatial information, either list or df, with column sample_key. Spatial can be empty if inputting data already derived. See add_summary_function() for more details.	
<pre>subject_key</pre>	column name in Metadata for subject ID	
sample_key	column linking Metadata to Spatial data	

Details

[Stable]

Value

S4 object of class mxFDA

Metadata	slot of class data.frame that contains sample and subject level information	
Spatial	slot of class data.frame that contains point level information within samples. An example would be cells belonging to $TMA\ cores$	
subject_key	slot of class character that corresponds to a column in the Metadata slot that groups samples at a subject level. An example would be " <i>patient_id</i> "	
sample_key	slot of class character that corresponds to a column both in the Metadata and Spatial slots that links samples to characteristics	
univariate_summaries		
	slot of class list where univariate summary functions calculated on Spatial would be stored	
bivariate_summaries		
	slot of class list where bivariate summary functions calculated on Spatial would be stored	
multiivariate_summaries		
	slot of class list where entropy summary functions calculated on $\ensuremath{Spatial}$ would be stored	

functional_pca	slot of class list where FPCA results are stored
functional_mpca	1
	slot of class list where MFPCA results are stored
functional_cox	slot of class list where functional cox model results are stored
functional_mcox	(
	slot of class list where mixed functional \cos model results are stored
scalar_on_function	
	slot of class list where functional models are fit to scalar responses

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

ovarian_FDA

Multiplex imaging data from an ovarian cancer tumor microarray

Description

This data is adapted from the VectraPolarisData Bioconductor package and comes from a tumormicroarray of tissue samples from 128 patients with ovarian cancer. There is one patient per subject.

Usage

ovarian_FDA

plot.afcmSurface

Format

ovarian_FDA:

An mxFDA object with augmented ovarian cancer multiplex immunofluorescence data, and NN G(r) calculated:

Metadata information about the spatial samples with column sample_key column in both

Spatial cell-level information with x and y columns along with sample_key to link to Metadata
subject_key column in Metadata that may have multiple sample_key values for each, akin to
patient IDs

sample_key column in both Metadata and Spatial that is a 1:1 with the samples (unique per sample)

univariate_summaries univariate summary slot with nearest neighbor G calculated

bivariate_summaries empty slot available for bivariate summaries

multiivariate_summaries empty slot available for multivariate summaries

functional_pca empty slot for functional PCA data of summaries

functional_cox empty slot for functional models

Details

Spatial summary functions of ovarian cancer multiplex imaging data.

This data is adapted from the VectraPolarisData Bioconductor package. Signal between the survival outcome and spatial summary functions has been augmented for teaching purposes. Spatial relationship is summarized using the nearest neighbor G function.

Source

https://bioconductor.org/packages/release/data/experiment/html/VectraPolarisData.
html

plot.afcmSurface Plot afcm object

Description

Plot afcm object

Usage

S3 method for class 'afcmSurface'
plot(x, ...)

Arguments

х	object of class afcmSurface to be plotted
	currently ignored

Value

object compatable with ggplot2

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

plot.lfcmSurface Plot lfcm surface

Description

Plot lfcm surface

Usage

S3 method for class 'lfcmSurface'
plot(x, ...)

Arguments

х	object of class lfcmSurface to be plotted
	currently ignored

Value

object compatable with ggplot2

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

plot.mxFDA

Description

Plot mxFDA object

Usage

```
## S3 method for class 'mxFDA'
plot(x, filter_cols = NULL, ...)
```

Arguments

х	object of class mxFDA to be plotted
filter_cols	column key to filter
	additional paramters including y, what, and sampleID to inform whats to be plotted

Details

[Stable]

If there are multiple metrics that are included in the derived table, an extra parameter filter_cols in the format of c(Derived_Column = "Level_to_Filter") will return curves from the Derived_Column with the level_to_Filter

When plotting mFPCA objects, additional arguments level1 and level2 help indicate which FPCA from level 1 and level 2 to plot

Value

object of class ggplot compatible the ggplot2 aesthetics

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

Examples

```
#plot fpca
plot(ovarian_FDA, what = 'uni g fpca', pc_choice = 1)
```

plot.sofr Plot sofr object

Description

Plot sofr object

Usage

S3 method for class 'sofr'
plot(x, ...)

Arguments

х	object of class sofr to be plotted
	currently ignored

Value

object compatable with ggplot2

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>
Alex Soupir <alex.soupir@moffitt.org>

plot_fpc

Create plot of mean +/- scaled eigenfunctions from FPCA

Description

Produces a ggplot with mean plus or minus two standard deviations of a selected FPC.

Usage

plot_fpc(obj, pc_choice)

Arguments

obj	fpca object to be plotted.
pc_choice	FPC to be plotted.

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plot_mfpc

Details

[Superseded]

Value

object of class ggplot

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

plot_mfpc

Create plot of mean +/- scaled eigenfunctions from FPCA

Description

Produces a ggplot with mean plus or minus two standard deviations of a selected FPC.

Usage

plot_mfpc(obj, pc_choice_level1, pc_choice_level2)

Arguments

obj fpca object to be plotted. pc_choice_level1, pc_choice_level2 FPC to be plotted.

Details

[Superseded]

Value

list of objects of class ggplot

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

run_fcm

Description

Fit a functional Cox regression model.

Usage

```
run_fcm(
    mxFDAobject,
    model_name,
    formula,
    event = "event",
    metric = "uni k",
    r = "r",
    value = "fundiff",
    afcm = FALSE,
    smooth = FALSE,
    filter_cols = NULL,
    ...,
    knots = NULL
)
```

Arguments

mxFDAobject	Dataframe of spatial summary functions from multiplex imaging data, in long format. Can be estimated using the function extract_summary_functions or provided separately.
model_name	character string to give the fit model in the functional cox slot
formula	Formula to be fed to mgcv in the form of survival_time ~ $x1 + x2$. Does not contain functional predictor. Character valued. Data must contain censoring variable called "event".
event	character string for the column in Metadata that contains $1\!/\!0$ for the survival event
metric	name of calculated spatial metric to use
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
afcm	If TRUE, runs additive functional Cox model. If FALSE, runs linear functional cox model. Defaults to linear functional cox model.
smooth	Option to smooth data using FPCA. Defaults to FALSE.
filter_cols	a named vector of factors to filter summary functions to in c(Derived_Column = "Level_to_Filter") format

run_fpca

•••	Optional other arguments to be passed to fpca.face
knots	Number of knots for defining spline basis.

Details

[Stable]

Value

A list which is a linear or additive functional Cox model fit. See mgcv::gam for more details.

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>
Alex Soupir <alex.soupir@moffitt.org>

Examples

run_fpca

run_fpca

Description

This is a wrapper for the function fpca. face from the refund package. EXPAND

Usage

```
run_fpca(
    mxFDAobject,
    metric = "uni k",
    r = "r",
    value = "fundiff",
    knots = NULL,
    analysis_vars = NULL,
    lightweight = FALSE,
    filter_cols = NULL,
    ...
)
```

Arguments

mxFDAobject	$object \ of \ class \ mxFDA \ created \ by \ make_mxfda \ with \ metrics \ derived \ with \ extract_summary_functions$
metric	name of calculated spatial metric to use
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
knots	Number of knots for defining spline basis. Defaults to the number of measure- ments per function divided by 2.
analysis_vars	Optional list of variables to be retained for downstream analysis.
lightweight	Default is FALSE. If TRUE, removes Y and Yhat from returned FPCA object. A good option to select for large datasets.
filter_cols	a named vector of factors to filter summary functions to in c(Derived_Column = "Level_to_Filter") format
	Optional other arguments to be passed to fpca.face

Details

[Stable]

The filter_cols parameter is useful when the summary function was input by the user using add_summary_function() and the multiple marks were assessed; a column called "Markers" with tumor infiltrating lymphocytes as well as cytotoxic T cells. This parameter allows for filtering down to include only one or the other.

Value

A mxFDA object with the functional_pca slot filled for the respective spatial summary function containing:

mxfundata	The original dataframe of spatial summary functions, with scores from FPCA
	appended for downstream modeling
fpc_object	A list of class "fpca" with elements described in the documentation for refund::fpca.face

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421. DOI: 10.1007/s11222-014-9485-x.

run_mfcm

Examples

run_mfcm

Run function Cox models for data with multiple samples per subject

Description

Fit a functional Cox regression model when there are multiple functions per subject, which arise from multiple samples per subject. It is not necessary for all subjects to have the same number of samples. The function first performs a multilevel functional principal components analysis (MF-PCA) decomposition to the spatial summary function. Then, the average curve for each subject is used in a functional Cox model (FCM). Variation around each subject's mean is captured by calculating the standard deviation of the level 2 scores from MFPCA, then including this as a scalar variable in the FCM called "level2_score_sd".

Usage

```
run_mfcm(
  mxFDAobject,
  model_name,
  formula,
  event = "event",
  metric = "uni k",
  r = "r",
  value = "fundiff",
  afcm = FALSE,
  filter_cols = NULL,
  pve = 0.99,
  ...,
  knots = NULL
)
```

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Arguments

mxFDAobject Dataframe of spatial summary functions from multiplex imaging data, in long format. Can be estimated using the function extract_summary_functions or provided separately. character string to give the fit model in the functional cox slot

formula	Formula to be fed to mgcv in the form of survival_time ~ $x1 + x2$. Does not contain functional predictor. Character valued. Data must contain censoring variable called "event".
event	character string for the column in Metadata that contains $1\!/\!0$ for the survival event
metric	name of calculated spatial metric to use
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
afcm	If TRUE, runs additive functional Cox model. If FALSE, runs linear functional cox model. Defaults to linear functional cox model.
filter_cols	a named vector of factors to filter summary functions to in c(Derived_Column = "Level_to_Filter") format
pve	Proportion of variance explained by multilevel functional principal components analysis in mfpca step
	Optional other arguments to be passed to fpca.face
knots	Number of knots for defining spline basis.

Details

[Stable]

Value

A list which is a linear or additive functional Cox model fit. See mgcv::gam for more details.

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>
Alex Soupir <alex.soupir@moffitt.org>

Examples

run_mfpca

run_fpca

Description

This is a wrapper for the function mfpca.face from the refund package. EXPAND

Usage

```
run_mfpca(
    mxFDAobject,
    metric = "uni k",
    r = "r",
    value = "fundiff",
    knots = NULL,
    lightweight = FALSE,
    ...
)
```

Arguments

mxFDAobject	$object \ of \ class \ mxFDA \ created \ by \ make_mxfda() \ with \ metrics \ derived \ with \ extract_summary_functions \ and \ an$
metric	name of calculated spatial metric to use
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
knots	Number of knots for defining spline basis.Defaults to the number of measure- ments per function divided by 2.
lightweight	Default is FALSE. If TRUE, removes Y and Yhat from returned mFPCA object. A good option to select for large datasets.
	Optional other arguments to be passed to mfpca.face

Details

[Stable]

Value

A mxFDA object with the functional_mpca slot for the respective spatial summary function containing:

mxfundata	The original dataframe of spatial summary functions, with scores from FPCA
	appended for downstream modeling
fpc_object	A list of class "fpca" with elements described in the documentation for refund::fpca.face

 run_sofr

Author(s)

unknown <first.last@domain.extension>

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421. DOI: 10.1007/s11222-014-9485-x.

Examples

#load data
data(lung_FDA)

#run mixed fpca
lung_FDA = run_mfpca(lung_FDA, metric = 'uni g')

run_sofr

Run Scalar on Function Regression

Description

Fit a scalar-on-function regression model. Uses refund::pfr under the hood for computations, and stores results in the mxfda object.

Usage

```
run_sofr(
  mxFDAobject,
  model_name,
  formula,
  family = "gaussian",
  metric = "uni k",
  r = "r",
  value = "fundiff",
  smooth = FALSE,
  filter_cols = NULL,
  ...,
  knots = NULL
)
```

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run_sofr

Arguments

mxFDAobject	Dataframe of spatial summary functions from multiplex imaging data, in long format. Can be estimated using the function extract_summary_functions or provided separately.
model_name	character string to give the fit model
formula	Formula to be fed to mgcv in the form of outcome ~ $x1 + x2$. Does not contain functional predictor. Character valued.
family	Exponential family distribution to be passed to mgcv::gam. Defaults to "gaussian". Select "binomial" for binary outcome.
metric	Name of calculated spatial metric to use
r	Character string, the name of the variable that identifies the function domain (usually a radius for spatial summary functions). Default is "r".
value	Character string, the name of the variable that identifies the spatial summary function values. Default is "fundiff".
smooth	Option to smooth data using FPCA. Defaults to FALSE.
filter_cols	a named vector of factors to filter summary functions to in c(Derived_Column = "Level_to_Filter") format
	Optional other arguments to be passed to fpca.face
knots	Number of knots for defining spline basis.

Details

[Stable]

Value

A list which is a linear or additive functional Cox model fit. See mgcv::gam for more details.

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

Examples

#load ovarian mxFDA object
data('ovarian_FDA')

```
# run scalar on function regression model with a binary outcome (stage)
# also known as functional logistic regression
```

summary.mxFDA

Summary method for object of class mxFDA

Description

Summary method for object of class mxFDA

Usage

S3 method for class 'mxFDA'
summary(object, ...)

Arguments

object	object of class mxFDA
	unused currently

Details

[Stable]

Value

summary of object to the R console

Author(s)

Alex Soupir <alex.soupir@moffitt.org>

univariate

Description

Internal function called by extract_summary_functions() to calculate a univariate spatial summary function for a single image.

Usage

```
univariate(
  mximg,
  markvar,
  mark1,
  mark2,
  r_vec,
  func = c(Kest, Lest, Gest),
  edge_correction,
  empirical_CSR = FALSE,
  permutations = 1000
)
```

Arguments

mximg	Dataframe of cell-level multiplex imaging data for a single image. Should have variables x and y to denote x and y spatial locations of each cell.
markvar	The name of the variable that denotes cell type(s) of interest. Character.
mark1	dummy filler, unused
mark2	dummy filler, unused
r_vec	Numeric vector of radii over which to evaluate spatial summary functions. Must begin at 0.
func	Spatial summary function to calculate. Options are c(Kest, Lest, Gest) which denote Ripley's K, Besag's L, and nearest neighbor G function, respectively.
edge_correction	
	Character string that denotes the edge correction method for spatial summary function. For Kest and Lest choose one of c("border", "isotropic", "Ripley", "translate", "none"). For Gest choose one of c("rs", "km", "han")
empirical_CSR	logical to indicate whether to use the permutations to identify the sample-specific complete spatial randomness (CSR) estimation.
permutations	integer for the number of permtuations to use if empirical_CSR is TRUE and exact CSR not calculable

Details

[Stable]

Value

A data.frame containing:

r	the radius of values over which the spatial summary function is evaluated
sumfun	the values of the spatial summary function
csr	the values of the spatial summary function under complete spatial randomness
fundiff	sumfun - csr, positive values indicate clustering and negative values repulsion

Author(s)

Julia Wrobel <julia.wrobel@emory.edu>

Alex Soupir <alex.soupir@moffitt.org>

References

Creed, J. H., Wilson, C. M., Soupir, A. C., Colin-Leitzinger, C. M., Kimmel, G. J., Ospina, O. E., Chakiryan, N. H., Markowitz, J., Peres, L. C., Coghill, A., & Fridley, B. L. (2021). spatialTIME and iTIME: R package and Shiny application for visualization and analysis of immunofluorescence data. *Bioinformatics* (Oxford, England), 37(23), 4584–4586. https://doi.org/10.1093/bioinformatics/btab757

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