# Package 'growthfd'

January 20, 2022

Title Fitting FPCA-based growth curve model
Version 0.0.0.9000
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Description  This package provides a method for fiting an FPCA-based growth curve model described in the paper stated bellow. This research was funded by Technology Agency of the Czech Republic (Technologická agentura České republiky), grant number TL01000394.
Citation Kralik M., Klima O., Cuta M., Malina R., Koziel S., Polcerova L., Skultety-ova A., Spanel M., Kukla L., Zemcik P. Estimating Growth in Height from Limited Longitudinal Growth Data Using Full-Curves Training Dataset: A Comparison of Two Procedures of Curve Optimization-Functional Principal Component Analysis and SITAR. Children, 2021, vol. 8, n. 10., pages 934-955.
License LGPL (>= 3)
Encoding UTF-8
RoxygenNote 7.1.2
Imports minpack.lm, sitar (>= 1.2.0), fda, ggplot2, parallel, doParallel, foreach, flock, MASS
<b>Depends</b> R (>= 2.10)
LazyData true
Suggests rmarkdown, knitr
VignetteBuilder knitr
R topics documented:
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growthfd

Fit a FPCA Growth Curve Model to a population

#### **Description**

This function fits a model to the given measured data of a population.

#### Usage

```
growthfd(
  data,
  x,
  y,
  id,
  model,
  verbose = 1,
  bounds = "negative",
  filename = "",
  startFromId = NULL,
  parallel = F,
  scores.filename = "parallel.txt"
)
```

#### **Arguments**

data	Data frame containing age, height and id of individuals
X	Age at measured data points
у	Height at measured data points
id	Corresponding individual's id at measured data points
model	FPCA growth model to be fitted
verbose	Verbosity
bounds	Limitation of the interval for milestones estimation, 'negative' or 'inverse'
filename	File name for saving results after each individual
startFromId	Start the evaluation from this id
parallel	(Experimental) Parallel evaluation of the model fitting
scores.filename	

File name for continuous saving of the scores

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#### Value

List containing individuals id and model

#### **Examples**

```
filename <- system.file("extdata", "data.csv", package="growthfd", mustWork=TRUE)
csv <- read.csv(filename)
d <- data.frame('id'=as.factor(csv[,'id']), 'x'=csv[,'age'], 'y'=csv[,'height'])
fit<-growthfd(data=d, x=x, y=y, id=id, model=model.bgs.m)</pre>
```

growthfd.apv

Compute apv of model instance

#### **Description**

This function computes apv related to the certain instance of the model described by the given parameters.

#### Usage

```
growthfd.apv(model, par)
```

## **Arguments**

model FPCA growth model

par Params of the model, corresponding to some individual

#### Value

Age of maximum growth velocity

```
growthfd.ApvRegVelocity
```

Register a velocity curve at population apv

### **Description**

This function registers a curve corresponding to the supplied parameters onto the population apv.

#### Usage

```
growthfd.ApvRegVelocity(model, par)
```

#### **Arguments**

model FPCA growth model

par Params of the model, corresponding to the individual

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#### Value

Velocity at apv data frame

#### **Examples**

```
filename <- system.file("extdata", "data.csv", package="growthfd", mustWork=TRUE)
csv <- read.csv(filename)
d <- data.frame('id'=as.factor(csv[,'id']), 'x'=csv[,'age'], 'y'=csv[,'height'])
m <- d$id == 'John'
fit <- growthfd.fit(model.bgs.m, age=d$x[m], height=d$y[m])
data<-growthfd.ApvRegVelocity(model.bgs.m, fit$par)</pre>
```

growthfd.evaluate

Generate a Discrete Growth Curve

#### **Description**

This function evaluates a curve function for given ages. Depending on a degree of derivation, the function produces stature, velocity or acceleration curve.

#### Usage

```
growthfd.evaluate(x, par, model, deriv = 0)
```

## Arguments

X	Ages to be evaluated
par	Parameters of the model
model	FPCA growth model
deriv	Path to the input file

#### Value

Y-values of the evaluated curve

growthfd.fit	Fit a FPCA Growth Curve Model to measurements of a single individ-
	ual

#### **Description**

This function fits a model to the given measured data of a single individual.

#### Usage

```
growthfd.fit(model, age, height, nprint = 1)
```

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#### **Arguments**

model FPCA growth model to be fitted age Age at measured data points height Height at at measured data points nprint Verbosity

#### Value

An optimization result object

#### **Examples**

```
age <- c(6.9, 8.2, 10, 12.1)
height <- c(114, 122, 130, 141)
fit <- growthfd.fit(model.bgs.m, age=c(6.9, 8.2, 10, 12.1), height=c(114, 122, 130, 141))
x11()
growthfd.plot(model.bgs.m, fit$par)
points(age, height)
x11()
growthfd.plot(model.bgs.m, fit$par, from=0.5, deriv = 1)
x11()
growthfd.plot(model.bgs.m, fit$par, from=0.5, deriv = 2)</pre>
```

growthfd.plot

Plot a Growth Curve

#### **Description**

This function plots a stature, velocity or acceleration curve.

## Usage

```
growthfd.plot(model, par, deriv = 0, from = 0, to = 18)
```

## Arguments

model	FPCA growth model
par	Parameters of the model
deriv	Path to the input file
from	The lower age limit
to	The upper age limit

```
growthfd.plot.ApvRegVelocity

Plot a velocity curve registered at apv
```

#### **Description**

This function plots a velocity curve, registered at population (model) apv in comparison with the mean curve.

#### Usage

```
growthfd.plot.ApvRegVelocity(data)
```

#### **Arguments**

model

Data obtained using growthfd.ApvRegVelocity

#### Value

Velocity at apv plot

### **Examples**

```
filename <- system.file("extdata", "data.csv", package="growthfd", mustWork=TRUE)
csv <- read.csv(filename)
d <- data.frame('id'=as.factor(csv[,'id']), 'x'=csv[,'age'], 'y'=csv[,'height'])
m <- d$id == 'John'
fit <- growthfd.fit(model.bgs.m, age=d$x[m], height=d$y[m])
data<-growthfd.ApvRegVelocity(model.bgs.m, fit$par)
p<-growthfd.plot.ApvRegVelocity(data)
x11()
p</pre>
```

```
growthfd.plot.RegVelocities
```

Plot velocity boxplots registered on apv

## Description

This function plots boxplots in time of measurements, after registration of the individual on population apv.

## Usage

```
growthfd.plot.RegVelocities(model, par, ages, rndn = 100)
```

#### **Arguments**

model	Model
par	Parameters of the model fitted to the measurements
ages	Ages of measurements points
rndn	Count of random curves to be evaluated

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#### Value

GGPlot2 plot

growthfd.residuals

Compute residuals

## Description

This function computes residuals between measured stature data and data generated from the growth model.

## Usage

```
growthfd.residuals(x, y, par, model)
```

## Arguments

x Vector with input ages

y Vector with target height measurements

par Parameters of the model model FPCA growth model

#### Value

A vector of residuals

growthfd.std

Generate a Curve Function

## **Description**

This function generates a growth curve function based on given model and parameters, describing the growth phase and amplitude.

## Usage

```
growthfd.std(par, model)
```

## Arguments

par Phase and amplitude parameters

model FPCA growth model

#### Value

FDA function object

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model.bgs.f

FPCA model for girls

## **Description**

Model trained using 167 female individuals from Brno Growth Study (BGS).

## Usage

```
model.bgs.f
```

#### **Format**

An object of class list of length 3.

model.bgs.m

FPCA model for boys

## Description

Model trained using 167 male individuals from Brno Growth Study (BGS).

## Usage

```
model.bgs.m
```

## **Format**

An object of class list of length 3.

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