

Package ‘gammaFuncModel’

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Type Package

Title Non-Linear Mixed Effects Model Based on the Gamma Function Form

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Description Identifies biomarkers that exhibit differential response dynamics by time across groups and estimates kinetic properties of biomarkers.

License GPL

RoxygenNote 7.2.3

Imports Rdpack, nlme, dplyr, cubature, rootSolve, patchwork, scales, ggplot2, future.apply, gridExtra, rlang, grDevices, stats, grid

NeedsCompilation no

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calculate_AUC	<i>Function that produces Area Under the Curve(AUC) property for a single individual in a particular group, for a specific metabolite</i>
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Description

Function that produces Area Under the Curve(AUC) property for a single individual in a particular group, for a specific metabolite

Usage

```
calculate_AUC(f, upperbound)
```

Arguments

f	function that returns the prediction of a metabolite concentration, for a single individual in a particular group
upperbound	Numeric value that serves as the upperbound of integration

Value

AUC for this metabolite, in a particular group for a single individual

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(cubature)
require(dplyr)
require(nlme)
modify.df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = NA
)
for (i in 1:10) {
  for (d in 1:3) {
    C0 <- runif(1, 10, 15) # initial concentration
```

```

k <- runif(1, 0.1, 0.3) # decay rate constant
modify.df$Concentration[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d] <-
  C0 * exp(-k * modify.df$Time[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d])
}
}
covariates <- c("ID", "Diet", "Age", "BMI")
model <- gammaFunction(
  modify.df,
  covariates,
  time_grp_inter = FALSE,
  return_ml_model = FALSE, include_grp = TRUE
)[[1]]
test_data <- modify.df %>%
  filter(Diet == 1 & ID == "04") %>%
  select(-c("Concentration", "ID", "Diet"))
f_dat = modify.df %>%
  filter(Diet == 1 & ID == "04") %>%
  select(-Concentration)
f <- generate_f_function(
  data = f_dat,
  model = model,
  grp_var = 1,
  grp_name = "Diet",
  ID = "04",
  ref = 1
)
AUC <- calculate_AUC(f, 9)
AUCInf <- calculate_AUC(f, Inf)

```

calculate_Cmax	<i>Function that produces Cmax property for a single individual in a particular group, for a specific metabolite</i>
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Description

Function that produces Cmax property for a single individual in a particular group, for a specific metabolite

Usage

```
calculate_Cmax(data, model, grp_var, ID, grp_name = "Diet", Tmax)
```

Arguments

data	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other individual characteristics covariates (excluding other forms of 'Time') Note: Data must be complete (No missing values).
model	Fitted model for the metabolite in question

grp_var	Value of the grouping variable
ID	Subject ID
grp_name	Name of the grouping variable. Default is 'Diet'
Tmax	for this metabolite, in a particular group for a single individual

Value

Cmax for this metabolite, in a particular group for a single individual

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(rootSolve)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = round(runif(270, 5, 15), 2)
)
covariates <- c("ID", "Diet", "Age", "BMI")
model <- gammaFunction(
  df,
  covariates,
  time_grp_inter = FALSE,
  return_ml_model = FALSE,
  include_grp = TRUE
)[[1]]
test_data = df %>% filter(Diet == 1 & ID == "02") %>% select(-c("Concentration", "ID", "Diet"))
Tmax <- calculate_Tmax(data = test_data, model, grp_var = 1, ID = "02", grp_name = 'Diet', ref = 1)
Cmax <- calculate_Cmax(data = test_data, model, grp_var = 1, ID = "02", grp_name = "Diet", Tmax)
```

calculate_half_life *Function that produces Half-life property for a single individual in a particular group, for a specific metabolite*

Description

Function that produces Half-life property for a single individual in a particular group, for a specific metabolite

Usage

```
calculate_half_life(f, Tmax, Cmax)
```

Arguments

f	function that returns the prediction of a metabolite concentration, for a single individual in a particular group
Tmax	Tmax property of a metabolite, for a single individual in a particular group
Cmax	Cmax property of a metabolite, for a single individual in a particular group

Value

Half-life for this metabolite, in a particular group for a single individual

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(rootSolve)
require(dplyr)
require(nlme)

modify.df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = NA
)
for (i in 1:10) {
  for (d in 1:3) {
    C0 <- runif(1, 10, 15) # initial concentration
    k <- runif(1, 0.1, 0.3) # decay rate constant
    modify.df$Concentration[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d] <-
      C0 * exp(-k * modify.df$Time[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d])
  }
}
covariates <- c("ID", "Diet", "Age", "BMI")
```

```

model <- gammaFunction(
  modify.df,
  covariates,
  time_grp_inter = FALSE,
  return_ml_model = FALSE,
  include_grp = TRUE
)[[1]]
test_data = modify.df %>%
  filter(Diet == 1 & ID == "03") %>%
  select(-c("Concentration", "ID", "Diet"))
Tmax <- calculate_Tmax(data = test_data, model, grp_var = 1, ID = "03", grp_name = 'Diet', ref = 1)
Cmax <- calculate_Cmax(data = test_data, model, grp_var = 1, ID = "03", grp_name = "Diet", Tmax)
f_dat = modify.df %>% filter(Diet == 1 & ID == "03") %>% select(-Concentration)
f <- generate_f_function(
  data = f_dat,
  model = model,
  grp_var = 1,
  grp_name = "Diet",
  ID = "03",
  ref = 1)
half_life <- calculate_half_life(f, Tmax, Cmax)

```

calculate_Tmax	<i>Function that produces Tmax property for a single individual in a particular group, for a specific metabolite</i>
----------------	----------------------------------------------------------------------------------------------------------------------

Description

Function that produces Tmax property for a single individual in a particular group, for a specific metabolite

Usage

```
calculate_Tmax(data, model, grp_var, ID, grp_name = "Diet", ref = 1)
```

Arguments

data	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other individual characteristics covariates(excluding other forms of 'Time') Note: Data must be complete (No missing values);
model	Fitted model for the metabolite in question
grp_var	Value of the grouping variable
ID	Subject ID
grp_name	Name of the grouping variable. Default is 'Diet'
ref	numeric or character. The reference level for the grouping variable, as a factor

Value

Tmax for this metabolite, in a particular group for a single individual

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(rootSolve)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = round(runif(270, 5, 15), 2)
)
covariates <- c("ID", "Diet", "Age", "BMI")
model <- gammaFunction(
  df,
  covariates,
  time_grp_inter = FALSE,
  return_ml_model = FALSE,
  include_grp = TRUE
)[[1]]
test_data = df %>% filter(Diet == 1 & ID == "01") %>% select(-c("Concentration", "ID", "Diet"))
Tmax <- calculate_Tmax(data = test_data, model, grp_var = 1, ID = "01", grp_name = 'Diet', ref = 1)
```

diffGrpResponse

Function that produces a summary table for coefficient estimates, their p-values and LRT p-values for every metabolite in the dataframe

Description

Function that produces a summary table for coefficient estimates, their p-values and LRT p-values for every metabolite in the dataframe

Usage

```
diffGrpResponse(
  df,
  met_vec,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp = "Diet",
  random_formula = ~1 + Time | ID/Diet,
  correlation_formula = corSymm(form = ~Time | ID/Diet),
  weights = varIdent(form = ~1 | Time)
)
```

Arguments

df	Data frame containing columns Group(numeric or character); ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.
met_vec	Vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
time_terms	Vector that contains all additional form of the covariate 'Time' (including the 'Time' covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp	Grouping variable;
random_formula	Random effects formula for the model, nested effects of Diet within ID (could also add random slope for 'Time');
correlation_formula	Correlation formula. Default is autogressive but can accommodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity

Value

Data frame that contains the coefficient estimates, their corresponding p-values; LRT p-values for Time-Group interactions (for every 'Time' term);LRT p-values for Group and Time-Group interactions (for every 'Time' term); as well as the fitted models for each metabolite

References

- Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>
- Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```

require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
covariates <- c("ID", "Diet", "Age", "BMI")
result <- diffGrpResponse(df, metvar, covariates)[[1]]
summary(result)

```

diffGrpResponse_parallel

Parallelized version of diffGrpResponse()

Description

Parallelized version of diffGrpResponse()

Usage

```

diffGrpResponse_parallel(
  df,
  met_vec,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp = "Diet",
  random_formula = ~1 + Time | ID/Diet,
  correlation_formula = corSymm(form = ~Time | ID/Diet),
  weights = varIdent(form = ~1 | Time)
)

```

Arguments

df Data frame containing columns Group(numeric or character); ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations
 Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.

met_vec	Vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
time_terms	Vector that contains all additional form of the covariate "Time" (including the "Time" covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp	Grouping variable;
random_formula	Random effects formula for the model, nested effects of Diet within ID (could also add random slope for "Time");
correlation_formula	Correlation formula. Default is autogressive but can accommodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity

Value

Data frame that contains the coefficient estimates, their corresponding p-values; LRT p-values for Time-Group interactions (for every "Time" term); LRT p-values for Group and Time-Group interactions (for every "Time" term); as well as the fitted models for each metabolite

Note

This function uses parallel processing via the 'future.apply' package. To enable parallel execution, runs the following before calling this function:

```
library(future.apply) plan(multisession, workers = parallel::detectCores() - 1)
```

You only need to set the plan once per session.

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
```

```

colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
covariates <- c("ID", "Diet", "Age", "BMI")
library(future.apply)
plan(multisession, workers = parallel::detectCores() - 1)
result <- diffGrpResponse(df, metvar, covariates)[[1]]
summary(result)

```

gammaFunction	<i>Implementation of the novel non-linear mixed-effects model based on gamma function form with nested covariance structure where random effects are specified for each Diet level within each subject (ID), capturing within-subject correlation across dietary conditions. to identify metabolites that responds to time differentially across dietary groups</i>
---------------	---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Description

Implementation of the novel non-linear mixed-effects model based on gamma function form with nested covariance structure where random effects are specified for each Diet level within each subject (ID), capturing within-subject correlation across dietary conditions. to identify metabolites that responds to time differentially across dietary groups

Usage

```

gammaFunction(
  data,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp = "Diet",
  random_formula = ~1 + Time | ID/Diet,
  correlation_formula = corSymm(form = ~Time | ID/Diet),
  weights = varIdent(form = ~1 | Time),
  time_grp_inter = TRUE,
  return_ml_model = FALSE,
  include_grp
)

```

Arguments

data	Data frame that contains the 'ID' column along with all covariates as well as concentration column, named 'Concentration', for a single metabolite Note: All non-concentration columns must be complete (No missing values); the concentration column can have missing values in the forms of either numeric 0 or 'NA'.
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates

time_terms	Vector that contains all additional form of the covariate 'Time' (including the 'Time' covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp	Grouping variable;
random_formula	Random effects formula for the model, nested effects of Diet within ID (could also add random slope for 'Time');
correlation_formula	Correlation formula. Default is autorrgressive but can accomodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity
time_grp_inter	Boolean value that indicates if the model should include interactions terms of 'time_terms' with 'Group';
return_ml_model	Boolean value that indicates if the model should fit "ML" model as well as "REML" model(default)
include_grp	boolean value to indicate whether or not 'grp' should be included in the model construction

Value

mixed effects models for a single metabolites: one with REML, the other with ML

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pineiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = round(runif(270, 5, 15), 2)
)
covariates <- c("ID", "Diet", "Age", "BMI")
model <- gammaFunction(
  df,
  covariates,
  random_formula = ~ 1 | ID/Diet,
  correlation_formula = corAR1(form = ~ Time | ID/Diet),
```

```

weights = NULL,
include_grp = TRUE)[[1]]
summary(model)

```

generatePlot

Function that generate plots for metabolite models

Description

Function that generate plots for metabolite models

Usage

```

generatePlot(
  graph,
  df,
  met_vec,
  covariates,
  grp = "Diet",
  models,
  save_path = NULL
)

```

Arguments

graph	character string, 'None' by default. If not 'None, in addition to returning models, produces pdf file of graphs based on the specific value of 'graph'.
df	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations; Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.
met_vec	the vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates;
grp	is the grouping variable;
models	a list of fitted non-linear mixed effects metabolite models
save_path	location (file path, not directory) where the pdf file will be saved (must end in '.pdf'); default is NULL, i.e. pdf is saved to a temporary location

Value

A pdf file for fitted concentration curves that is saved to a user provided file location; otherwise saved to a temporary location

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
require(patchwork)
require(scales)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
covariates <- c("ID", "Diet", "Age", "BMI")
mods <- generate_models(df = df, met_vec = metvar, covariates = covariates, graph = 'None')
generatePlot(
  graph = "individual_separated",
  df = df,
  met_vec = metvar,
  covariates = covariates,
  grp = "Diet",
  models = mods,
  save_path = NULL
)
```

generate_f_function *Function produce predictions from the model*

Description

Function produce predictions from the model

Usage

```
generate_f_function(data, model, grp_var, grp_name = "Diet", ID, ref = 1)
```

Arguments

data	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other individual characteristics covariates (excluding other forms of 'Time') Note: Data must be complete (No missing values).
model	Fitted model for the metabolite in question
grp_var	Value of the grouping variable
grp_name	Name of the grouping variable. Default is 'Diet'
ID	Subject ID
ref	reference group

Value

f function that produces the prediction from this model for a specific individual in a specific group

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
modify.df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1),
  Concentration = NA
)
for (i in 1:10) {
  for (d in 1:3) {
    C0 <- runif(1, 10, 15) # initial concentration
    k <- runif(1, 0.1, 0.3) # decay rate constant
    modify.df$Concentration[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d] <-
      C0 * exp(-k * modify.df$Time[modify.df$ID == sprintf("%02d", i) & modify.df$Diet == d])
  }
}
covariates <- c("ID", "Diet", "Age", "BMI")
model <- gammaFunction(
  modify.df,
  covariates,
  time_grp_inter = FALSE,
  return_ml_model = FALSE,
  include_grp = TRUE
)[[1]]
test_data = modify.df %>%
  filter(Diet == 1 & ID == "04") %>%
  select(-c("Concentration", "ID", "Diet"))
f_dat = modify.df %>% filter(Diet == 1 & ID == "04") %>% select(-Concentration)
f <- generate_f_function(
  data = f_dat,
  model = model,
```

```

    grp_var = 1,
    grp_name = "Diet",
    ID = "04",
    ref = 1
  )

```

generate_models

Function that produces a fitted gamma model for each metabolite

Description

Function that produces a fitted gamma model for each metabolite

Usage

```

generate_models(
  df,
  met_vec,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp_name = "Diet",
  random_formula = ~1 + Time | ID/Diet,
  correlation_formula = corSymm(form = ~Time | ID/Diet),
  weights = varIdent(form = ~1 | Time),
  graph = "None",
  save_path = NULL
)

```

Arguments

df	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.
met_vec	the vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
time_terms	is the vector that contains all additional form of the covariate 'Time' (including the 'Time' covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp_name	is the grouping variable;
random_formula	is the random effects formula for the model, nested effects of Diet within ID (could also add random slope for 'Time');

correlation_formula	is the correlation formula. Default is autogressive but can accommodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity;
graph	character string, 'None' by default. If not 'None, in addition to returning models, produces pdf file of graphs based on the specific value of 'graph'.
save_path	location where the pdf file will be saved; default is NULL, i.e. pdf is saved to a temporary location

Value

List that contains fitted models for each metabolite and a pdf file for fitted concentration curves.

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
covariates <- c("ID", "Diet", "Age", "BMI")
mods <- generate_models(
  df = df,
  met_vec = metvar,
  covariates = covariates,
  graph = 'None',
  save_path = NULL)
```

grpResp2Time	<i>Function that produces a summary table for coefficient estimates, their p-values and LRT p-values for every metabolite in the dataframe, for a single Group</i>
--------------	--------------------------------------------------------------------------------------------------------------------------------------------------------------------

Description

Function that produces a summary table for coefficient estimates, their p-values and LRT p-values for every metabolite in the dataframe, for a single Group

Usage

```
grpResp2Time(
  df,
  met_vec,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp = "Diet",
  random_formula = ~1 | ID,
  correlation_formula = corAR1(form = ~Time | ID),
  weights = NULL
)
```

Arguments

df	Data frame containing information for a single group, containing columns grp; ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.
met_vec	Vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
time_terms	Vector that contains all additional form of the covariate 'Time' (including the 'Time' covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp	Grouping variable (should be a single valued column);
random_formula	Random effects formula for the model, within ID (could also add random slope for 'Time');
correlation_formula	Correlation formula. Default is autogressive but can accommodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity

Value

Data frame that contains the coefficient estimates, their corresponding p-values as well as LRT p-values for 'Time' terms

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
df_single_diet <- subset(df, Diet == 1)
covariates <- c("ID", "Diet", "Age", "BMI")
result_SD <- grpResp2Time(df_single_diet, metvar, covariates)[[1]]
summary(result_SD)
```

grpResp2Time_parallel *Vectorized version of grpRes2Time()*

Description

Vectorized version of grpRes2Time()

Usage

```
grpResp2Time_parallel(
  df,
  met_vec,
  covariates,
  time_terms = c("Time", "log(Time)"),
  grp = "Diet",
```

```

random_formula = ~1 | ID,
correlation_formula = corAR1(form = ~Time | ID),
weights = NULL
)

```

Arguments

df	Data frame containing information for a single group, containing columns grp; ID(subject ID: character); Time(positive: numeric); other Time terms (numeric); other individual characteristics covariates; as well columns of metabolite concentrations Note: All non-concentration columns must be complete (No missing values); concentration columns can have missing values in the forms of either numeric 0 or 'NA'.
met_vec	Vector of metabolite names
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
time_terms	Vector that contains all additional form of the covariate "Time" (including the "Time" covariate), and must contain 'log(Time)', other forms also include I(Time^2) and I(Time^3);
grp	Grouping variable (should be a single valued column);
random_formula	Random effects formula for the model, within ID (could also add random slope for 'Time');
correlation_formula	Correlation formula. Default is autoregressive but can accommodate other forms such as unstructured covariance or exponential covariance;
weights	specify a variance function that models heteroscedasticity

Value

Data frame that contains the coefficient estimates, their corresponding p-values as well as LRT p-values for 'Time' terms

Note

This function uses parallel processing via the 'future.apply' package. To enable parallel execution, runs the following before calling this function:

```
library(future.apply) plan(multisession, workers = parallel::detectCores() - 1)
```

You only need to set the plan once per session.

References

- Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>
- Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```

require(gammaFuncModel)
require(dplyr)
require(nlme)
df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
concentration_data <- replicate(10, round(runif(270, 5, 15), 2))
colnames(concentration_data) <- metvar[1:10]
df <- cbind(df, as.data.frame(concentration_data))
df_single_diet <- subset(df, Diet == 1)
covariates <- c("ID", "Diet", "Age", "BMI")
library(future.apply)
plan(multisession, workers = parallel::detectCores() - 1)
result_SD <- grpResp2Time_parallel(df_single_diet, metvar, covariates)[[1]]
summary(result_SD)

```

pk_calculation	<i>Function that returns a data frame for Tmax, Cmax, half-life, AUC and AUCInf for metabolites</i>
----------------	-----------------------------------------------------------------------------------------------------

Description

Function that returns a data frame for Tmax, Cmax, half-life, AUC and AUCInf for metabolites

Usage

```
pk_calculation(df, met_vec, models, grp_name = "Diet", covariates, ref = 1)
```

Arguments

df	Data frame containing columns Group(factor); ID(subject ID: character); Time(positive: numeric); other individual characteristics covariates (excluding other forms of 'Time') Note: Data must be complete (No missing values).
met_vec	Vector of metabolite names
models	Fitted models for all metabolites of interest
grp_name	Name of the grouping variable
covariates	Vector containing the names of the "ID" covariate, grouping covariate and other covariates excluding any "Time" covariates
ref	reference level for the grouping variable. could be numeric or character

Value

Data frame with the pharmacokinetic properties of each metabolite

References

Wickham, H. (2022). dplyr: A Grammar of Data Manipulation. R package version 1.0.10. Available at: <https://CRAN.R-project.org/package=dplyr>

Pinheiro, J. C., & Bates, D. M. (2022). nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-153. Available at: <https://CRAN.R-project.org/package=nlme>

Examples

```
require(gammaFuncModel)
require(dplyr)

df <- data.frame(
  ID = rep(sprintf("%02d", 1:10), each = 9 * 3),
  Time = rep(rep(1:9, each = 3), 10),
  Diet = as.factor(rep(1:3, times = 9 * 10)),
  Age = rep(sample(20:70, 10, replace = TRUE), each = 9 * 3),
  BMI = round(rep(runif(10, 18.5, 35), each = 9 * 3), 1)
)
metvar <- paste0("met", 1:10)
n_rows <- nrow(df)
concentration_data <- sapply(1:10, function(m) {
  shape <- runif(1, 2, 5)
  scale <- runif(1, 1, 3)
  rgamma(n_rows, shape = shape, scale = scale)
})
colnames(concentration_data) <- metvar
df <- cbind(df, as.data.frame(concentration_data))
covariates <- c("ID", "Diet", "Age", "BMI")
mods <- generate_models(df = df, met_vec = metvar, covariates = covariates, graph = 'None')
result <- pk_calculation(
  df = df,
  met_vec = metvar,
  models = mods,
  grp_name = "Diet",
  covariates = covariates
)
```

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