

# Package: glmmSeq (via r-universe)

September 18, 2024

**Title** General Linear Mixed Models for Gene-Level Differential Expression

**Version** 0.5.5

**Description** Using mixed effects models to analyse longitudinal gene expression can highlight differences between sample groups over time. The most widely used differential gene expression tools are unable to fit linear mixed effect models, and are less optimal for analysing longitudinal data. This package provides negative binomial and Gaussian mixed effects models to fit gene expression and other biological data across repeated samples. This is particularly useful for investigating changes in RNA-Sequencing gene expression between groups of individuals over time, as described in: Rivellesse, F., Surace, A. E., Goldmann, K., Sciacca, E., Cubuk, C., Giorli, G., ... Lewis, M. J., & Pitzalis, C. (2022) Nature medicine <[doi:10.1038/s41591-022-01789-0](https://doi.org/10.1038/s41591-022-01789-0)>.

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<https://github.com/myles-lewis/glmmSeq>

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**Repository** <https://myles-lewis.r-universe.dev>

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fcPlot	<i>Plotly or ggplot fold change plots</i>
--------	---

---

## Description

Plotly or ggplot fold change plots

## Usage

```
fcPlot(
  object,
  x1var,
  x2var,
  x1Values = NULL,
  x2Values = NULL,
  pCutoff = 0.01,
  labels = c(),
  useAdjusted = FALSE,
  plotCutoff = 1,
  graphics = "ggplot",
  fontSize = 12,
  labelFontSize = 4,
  colours = c("grey", "goldenrod1", "red", "blue"),
```

```

    verbose = FALSE,
    ...
  )

```

### Arguments

object	A glmmSeq object created by <code>glmmSeq::glmmSeq()</code> .
x1var	The name of the first (inner) x parameter
x2var	The name of the second (outer) x parameter
x1Values	Timepoints or categories in x1var used to calculate fold change. If NULL the first two levels in x1var are used.
x2Values	Categories in x2var to be compared on x and y axis.
pCutoff	The significance cut-off for colour-coding (default = 0.01)
labels	Row names or indices to label on plot
useAdjusted	whether to use adjusted p-values (must have q-values in object). Default = FALSE
plotCutoff	Which probes to include on plot by significance cut-off (default = 1, for all markers)
graphics	Graphics system to use: "ggplot" or "plotly"
fontSize	Font size
labelFontSize	Font size for labels
colours	Vector of colours to use for significance groups
verbose	Whether to print statistics
...	Other parameters to pass to plotly or ggplot

### Value

Returns a plot for fold change between x1Values in one x2Value subset on x axis and fold change in the other x2Value on the y axis.

### Examples

```

data(PEAC_minimal_load)

disp <- apply(tpm, 1, function(x) {
  (var(x, na.rm = TRUE)-mean(x, na.rm = TRUE))/(mean(x, na.rm = TRUE)**2)
})

glmmFit <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  countdata = tpm[1:5, ],
  metadata = metadata,
  dispersion = disp,
  verbose = FALSE)

fcPlot(object = glmmFit,
  x1var = "Timepoint",

```

```
x2var = "EULAR_6m",
x2Values = c("Good", "Non-response"),
pCutoff = 0.05,
useAdjusted = FALSE,
plotCutoff = 1,
graphics = "plotly")
```

---

ggmodelPlot

*Mixed model effects plot using ggplot2*


---

## Description

Plot to show differences between groups and over time using ggplot2.

## Usage

```
ggmodelPlot(
  object,
  geneName = NULL,
  x1var = NULL,
  x2var = NULL,
  x2shift = NULL,
  xlab = NULL,
  ylab = geneName,
  plab = NULL,
  title = geneName,
  logTransform = is(object, "GlmSeq"),
  shapes = 19,
  colours = "grey60",
  lineColours = "grey60",
  markerSize = 1,
  fontSize = 12,
  alpha = 0.7,
  x2Offset = 5,
  addPoints = TRUE,
  addModel = TRUE,
  modelSize = 4,
  modelColours = "blue",
  modelLineSize = 1,
  modelLineColours = modelColours,
  addBox = FALSE,
  ...
)
```

## Arguments

**object** A `glmmSeq`/`lmmSeq` object created by `glmmSeq::glmmSeq()` or `glmmSeq::lmmSeq()`

geneName	The gene/row name to be plotted
x1var	The name of the first (inner) x parameter, typically 'time'. This is anticipated to have different values when matched by ID.
x2var	The name of an optional second (outer) x parameter, which should be a factor.
x2shift	Amount to shift along x axis for each level of x2var. By default the function will arrange each level of x2var side by side.
xlab	Title for the x axis
ylab	Title for the y axis
plab	Optional character vector of labels for p-values. These must align with column names in object@stats\$pvals.
title	Plot title. If NULL gene name is used
logTransform	Whether to perform a log10 transform on the y axis
shapes	The marker shapes (default=19)
colours	The marker colours as vector
lineColours	The line colours (default='grey60') as vector
markerSize	Size of markers (default=1)
fontSize	Plot font size
alpha	Line and marker opacity (default=0.7)
x2offset	Vertical adjustment to secondary x-axis labels (default=5)
addPoints	Whether to add underlying data points (default=TRUE)
addModel	Whether to add the fit model with markers (default=TRUE)
modelSize	Size of model points (default=4)
modelColours	Colour of model fit markers (default="blue") as vector
modellineSize	Size of model points (default=1) as vector
modellineColours	Colour of model fit lines
addBox	Logical whether to add boxplots for mean and IQR
...	Other parameters to pass to <code>ggplot2::theme()</code> .

**Value**

Returns a paired plot for matched samples.

**Examples**

```
data(PEAC_minimal_load)

disp <- apply(tpm, 1, function(x){
  (var(x, na.rm=TRUE)-mean(x, na.rm=TRUE))/(mean(x, na.rm=TRUE)**2)
})

MS4A1glmm <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  countdata = tpm['MS4A1', , drop = FALSE],
```

```

        metadata = metadata,
        dispersion = disp,
        verbose = FALSE)

ggmodelPlot(object = MS4A1glmm,
            geneName = 'MS4A1',
            x1var = 'Timepoint',
            x2var = 'EULAR_6m',
            colours = c('skyblue', 'goldenrod1', 'mediumvioletred'))

```

---

glmmQvals

*Glmm Sequencing qvalues*


---

## Description

Add qvalue columns to the glmmSeq dataframe

## Usage

```
glmmQvals(object, cutoff = 0.05, verbose = TRUE)
```

## Arguments

object	A glmmSeq/lmmSeq object created by <code>glmmSeq::glmmSeq()</code> .
cutoff	Prints a table showing the number of probes considered significant by the pvalue cut-off (default=0.05)
verbose	Logical whether to print the number of significant probes (default=TRUE)

## Value

Returns a GlmmSeq object with results for gene-wise general linear mixed models with adjusted p-values using the qvalue function

## Examples

```

data(PEAC_minimal_load)
disp <- apply(tpm, 1, function(x) {
  (var(x, na.rm=TRUE)-mean(x, na.rm = TRUE))/(mean(x, na.rm = TRUE)**2)
})
MS4A1glmm <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
                  countdata = tpm[1:5, ],
                  metadata = metadata,
                  dispersion = disp[1:5],
                  verbose=FALSE)
MS4A1glmm <- glmmQvals(MS4A1glmm)

```

---

`glmmRefit`*Refit mixed effects model*

---

## Description

Based on a 'GlmmSeq' or 'lmmSeq' class result object, this function attempts to refit an identical model for a specific gene based on the data and fitting parameters stored in the results object and refitting using either `lme4::glmer()` for GlmmSeq objects or `lmer()` for lmmSeq objects. The fitted model can then be passed on to other packages such as `emmeans` to look at estimated marginal means for the model.

## Usage

```
glmmRefit(object, gene, ...)  
  
## S3 method for class 'lmmSeq'  
glmmRefit(object, gene, formula = object@formula, ...)  
  
## S3 method for class 'GlmmSeq'  
glmmRefit(  
  object,  
  gene,  
  formula = object@formula,  
  control = object@info$control,  
  family = NULL,  
  ...  
)
```

## Arguments

<code>object</code>	A fitted results object of class <code>GlmmSeq</code> or <code>lmmSeq</code>
<code>gene</code>	A character value specifying a single gene to extract a fitted model for
<code>...</code>	Optional arguments passed to either <code>lme4::glmer()</code> or <code>lme4::lmer()</code>
<code>formula</code>	Optional formula to use when refitting model
<code>control</code>	Optional control parameters, see <code>lme4::lmerControl()</code> or <code>lme4::glmerControl()</code>
<code>family</code>	Optional GLM family when refitting GLMM using <code>lme4::glmer()</code> or <code>glmmTMB()</code>

## Value

Fitted model of class `lmerMod` in the case of LMM, or `glmerMod` or `glmmTMB` for a GLMM dependent on the original method.

**Examples**

```

data(PEAC_minimal_load)
disp <- apply(tpm, 1, function(x) {
  (var(x, na.rm = TRUE)-mean(x, na.rm = TRUE))/(mean(x, na.rm = TRUE)**2)
})
glmmtest <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  countdata = tpm[1:2, ],
  metadata = metadata,
  dispersion = disp,
  verbose = FALSE)

# show summary for single gene
summary(glmmtest, "MS4A1")

# refit a single model using lme4::glmer()
fit <- glmmRefit(glmmtest, "MS4A1")

# refit model with reduced formula
fit2 <- glmmRefit(glmmtest, "MS4A1",
  formula = count ~ Timepoint + EULAR_6m + (1 | PATID))

# LRT
anova(fit, fit2)

```

---

glmmSeq

*GLMM with negative binomial distribution for sequencing count data*


---

**Description**

Fits many generalised linear mixed effects models (GLMM) with negative binomial distribution for analysis of overdispersed count data with random effects. Designed for longitudinal analysis of RNA-Sequencing count data.

**Usage**

```

glmmSeq(
  modelFormula,
  countdata,
  metadata,
  id = NULL,
  dispersion = NA,
  sizeFactors = NULL,
  reduced = NULL,
  modelData = NULL,
  designMatrix = NULL,
  method = c("lme4", "glmmTMB"),
  control = NULL,

```



```

    family = nbinom2,
    cores = 1,
    removeSingles = FALSE,
    zeroCount = 0.125,
    verbose = TRUE,
    returnList = FALSE,
    progress = FALSE,
    ...
)

```

## Arguments

modelFormula	the model formula. This must be of the form " $\sim \dots$ " where the structure is assumed to be "counts $\sim \dots$ ". The formula must include a random effects term. For more information on formula structure for random effects see <a href="#">lme4::glmer()</a>
countdata	the sequencing count data matrix with genes in rows and samples in columns
metadata	a dataframe of sample information with variables in columns and samples in rows
id	Optional. Used to specify the column in metadata which contains the sample IDs to be used in repeated samples for random effects. If not specified, the function defaults to using the variable after the " " in the random effects term in the formula.
dispersion	a numeric vector of gene dispersion. Not required for glmmTMB models, as these determine and fit dispersion for each gene.
sizeFactors	size factors (default = NULL). If provided the glmer offset is set to log(sizeFactors). For more information see " <a href="#">lme4::glmer()</a> "
reduced	Optional reduced model formula. If this is chosen, a likelihood ratio test is used to calculate p-values instead of the default Wald type 2 Chi-squared test.
modelData	Optional dataframe. Default is generated by call to <code>expand.grid</code> using levels of variables in the formula. Used to calculate model predictions (estimated means & 95% CI) for plotting via <a href="#">modelPlot</a> . It can therefore be used to add/remove points in <a href="#">modelPlot</a> .
designMatrix	custom design matrix, used only for prediction
method	Specifies which package to use for fitting GLMM models. Either "lme4" or "glmmTMB" depending on whether to use <a href="#">lme4::glmer</a> or <a href="#">glmmTMB::glmmTMB</a> to fit GLMM models.
control	the glmer optimizer control. If method = "lme4" default is <code>glmerControl(optimizer = "bobyqa")</code> . If method = "glmmTMB" default is <code>glmmTMBControl()</code>
family	Only used with glmmTMB models. Default is <code>nbinom2</code> . See <a href="#">glmmTMB::nbinom2</a>
cores	number of cores to use. Default = 1.
removeSingles	whether to remove individuals without repeated measures (default = FALSE)
zeroCount	numerical value to offset zeroes for the purpose of log (default = 0.125)
verbose	Logical whether to display messaging (default = TRUE)

<code>returnList</code>	Logical whether to return results as a list or <code>glmmSeq</code> object (default = FALSE). Useful for debugging.
<code>progress</code>	Logical whether to display a progress bar
<code>...</code>	Other parameters to pass to <code>lme4::glmer()</code>

### Details

This function is a wrapper for `lme4::glmer()`. By default, p-values for each model term are computed using Wald type 2 Chi-squared test as per `car::Anova()`. The underlying code for this has been optimised for speed. However, if a reduced model formula is specified by setting `reduced`, then a likelihood ratio test is performed instead using `stats::anova`. This will double computation time since two GLMM have to be fitted.

Parallelisation is provided using `parallel::mclapply` on Unix/Mac or `parallel::parLapply` on PC.

Setting `method = "glmmTMB"` enables an alternative method of fitting GLMM using the `glmmTMB` package. This gives access to a variety of alternative GLM family functions. Note, `glmmTMB` negative binomial models are substantially slower to fit than `glmer` models with known dispersion due to the extra time taken by `glmmTMB` to optimise the dispersion parameter.

The `id` argument is usually optional. By default the `id` column in the metadata is determined as the term after the bar in the random effects term of the model. Note that `id` is not passed to `glmer` or `glmmTMB`. It is only really used to remove singletons from the countdata matrix and metadata dataframe. The `id` is also stored in the output from `glmmSeq` and used by plotting function `modelPlot()`. However, due to its flexible nature, in theory `glmmSeq` should allow for more than one random effect term, although this has not been tested formally. In this case, it is probably prudent to specify a value for `id`.

### Value

Returns an S4 class `GlmmSeq` object with results for gene-wise general linear mixed models. A list of results is returned if `returnList` is TRUE which is useful for debugging. If all genes return errors from `glmer`, then an error message is shown and a character vector containing error messages for all genes is returned.

### See Also

[lme4::glmer](#) [lme4::glmerControl](#) [glmmTMB::glmmTMB](#) [glmmTMB::nbinom2](#) [glmmTMB::glmmTMBControl](#) [car::Anova](#)

### Examples

```
data(PEAC_minimal_load)
disp <- apply(tpm, 1, function(x) {
  (var(x, na.rm = TRUE)-mean(x, na.rm = TRUE))/(mean(x, na.rm = TRUE)**2)
})
MS4A1glmm <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
                    countdata = tpm[1:2, ],
                    metadata = metadata,
                    dispersion = disp,
                    verbose = FALSE)
names(attributes(MS4A1glmm))
```

---

GlmSeq-class	<i>An S4 class to define the glmmSeq output</i>
--------------	---

---

**Description**

An S4 class to define the glmmSeq output

**Slots**

info List including the matched call, dispersions, offset, designMatrix

formula The model formula

stats Statistics from fitted models

predict Predicted values

reduced Optional reduced formula for LRT

countdata The input expression data with count data in rows

metadata The input metadata

modelData Model data for predictions

optInfo Information on whether the model was singular or converged

errors Any errors

vars List of variables stored from the original call, including the id variable (by default automatically identified from the random effect term in the model) and removeSingles argument

---

lmmSeq	<i>Linear mixed models for data matrix</i>
--------	--

---

**Description**

Fits many linear mixed effects models for analysis of gaussian data with random effects, with parallelisation and optimisation for speed. It is suitable for longitudinal analysis of high dimensional data. Wald type 2 Chi-squared test is used to calculate p-values.

**Usage**

```
ImmSeq(
  modelFormula,
  maindata,
  metadata,
  id = NULL,
  offset = NULL,
  test.stat = c("Wald", "F", "LRT"),
  reduced = NULL,
  modelData = NULL,
  designMatrix = NULL,
  control = lmerControl(),
  cores = 1,
  removeSingles = FALSE,
  verbose = TRUE,
  returnList = FALSE,
  progress = FALSE,
  ...
)
```

**Arguments**

modelFormula	the model formula. This must be of the form " $\sim \dots$ " where the structure is assumed to be "gene $\sim \dots$ ". The formula must include a random effects term. See formula structure for random effects in <a href="#">lme4::lmer()</a>
maindata	data matrix with genes in rows and samples in columns
metadata	a dataframe of sample information with variables in columns and samples in rows
id	Optional. Used to specify the column in metadata which contains the sample IDs to be used in repeated samples for random effects. If not specified, the function defaults to using the variable after the " " in the random effects term in the formula.
offset	Vector containing model offsets (default = NULL). If provided the <code>lmer()</code> offset is set to <code>offset</code> . See <a href="#">lme4::lmer()</a>
test.stat	Character value specifying test statistic. Current options are "Wald" for type 2 Wald Chi square test using code derived and modified from <a href="#">car::Anova</a> to improve speed for matrix tests. Or "F" for conditional F tests using Saiterthwaite's method of approximated Df. This uses <a href="#">lmerTest::lmer</a> and is somewhat slower.
reduced	Optional reduced model formula. If this is chosen, a likelihood ratio test is used to calculate p-values instead of the default Wald type 2 Chi-squared test.
modelData	Optional dataframe. Default is generated by call to <code>expand.grid</code> using levels of variables in the formula. Used to calculate model predictions (estimated means & 95% CI) for plotting via <a href="#">modelPlot</a> . It can therefore be used to add/remove points in <a href="#">modelPlot</a> .
designMatrix	Optional custom design matrix generated by call to <code>model.matrix</code> using <code>modelData</code> and <code>FEformula</code> . Used to calculate model predictions for plotting.

<code>control</code>	the lmer optimizer control (default = <code>lmerControl()</code> ). See <code>lme4::lmerControl()</code> .
<code>cores</code>	number of cores to use for parallelisation. Default = 1.
<code>removeSingles</code>	whether to remove individuals with no repeated measures (default = FALSE)
<code>verbose</code>	Logical whether to display messaging (default = TRUE)
<code>returnList</code>	Logical whether to return results as a list or ImmSeq object (default = FALSE). Helpful for debugging.
<code>progress</code>	Logical whether to display a progress bar
<code>...</code>	Other parameters passed to <code>lmerTest::lmer()</code> . Only available if <code>test.stat = "F"</code> .

## Details

By default, p-values for each model term are computed using Wald type 2 Chi-squared test as per `car::Anova()`. The underlying code for this has been optimised for speed. However, if a reduced model formula is specified by setting `reduced`, then a likelihood ratio test (LRT) is performed instead using `anova`. This will double computation time since two LMM have to be fitted for each gene. For LRT, models being compared are optimised by maximum likelihood and not REML (`REML=FALSE`).

Two key methods are used to speed up computation above and beyond simple parallelisation. The first is to speed up `lme4::lmer()` by calling `lme4::lFormula()` once at the start and then updating the `lFormula` output with new data. The 2nd speed up is through optimised code for repeated type 2 Wald Chi-squared tests (original code was derived from `car::Anova`). For example, elements such as the hypothesis matrices are generated only once to reduce unnecessarily repetitive computation, and the generation of p-values from Chi-squared values is vectorised and performed at the end. F-tests using the `lmerTest` package have not been optimised and are therefore slower.

Parallelisation is performed using `parallel::mclapply` on unix/mac and `parallel::parLapply` on windows. Progress bars use `pbmclapply::pbmclapply` on unix/mac and `pbapply::pbapply` on windows.

The `id` argument is usually optional. By default the `id` column in the metadata is determined as the term after the bar in the random effects term of the model. Note that `id` is not passed to `lmer`. It is only really used to remove singletons from the `maindata` matrix and `metadata` dataframe. The `id` is also stored in the output from `ImmSeq` and used by plotting function `modelPlot()`. However, due to its flexible nature, in theory `ImmSeq` should allow for more than one random effect term, although this has not been tested formally. In this case, it is probably prudent to specify a value for `id`.

## Value

Returns an S4 class `ImmSeq` object with results for gene-wise linear mixed models; or a list of results if `returnList` is TRUE, which is useful for debugging. If all genes return errors from `lmer`, then an error message is shown and a character vector containing error messages for all genes is returned.

## Examples

```
data(PEAC_minimal_load)
logtpm <- log2(tpm + 1)
lmmtest <- ImmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  maindata = logtpm[1:2, ],
  metadata = metadata,
```

```

                                verbose = FALSE)
names(attributes(lmmtest))

```

---

lmmSeq-class	<i>An S4 class to define the lmmSeq output</i>
--------------	--

---

### Description

An S4 class to define the lmmSeq output

### Slots

info List including matched call, offset, designMatrix  
 formula The model formula  
 stats Statistics from fitted models  
 predict Predicted values  
 reduced Optional reduced formula for LRT  
 maindata The input expression data with variables in rows  
 metadata The input metadata  
 modelData Model data for predictions  
 optInfo Information on whether the model was singular or converged  
 errors Any errors  
 vars List of variables stored from the original call

---

maPlot	<i>MA plots</i>
--------	-----------------

---

### Description

MA plots

### Usage

```

maPlot(
  object,
  x1var,
  x2var,
  x1Values = NULL,
  x2Values = NULL,
  pCutoff = 0.01,
  plotCutoff = 1,
  zeroCountCutoff = 50,

```

```

    colours = c("grey", "midnightblue", "mediumvioletred", "goldenrod"),
    labels = c(),
    fontSize = 12,
    labelFontSize = 4,
    useAdjusted = FALSE,
    graphics = "ggplot",
    verbose = FALSE
  )

```

### Arguments

object	A glmmSeq object created by <code>glmmSeq::glmmSeq()</code> .
x1var	The name of the first (inner) x parameter
x2var	The name of the second (outer) x parameter
x1Values	Timepoints or categories in x1var to be used to calculate fold change. If NULL the first two levels in x1var are used.
x2Values	Categories in x2var to be compared on x and y axis.
pCutoff	The significance cut-off for colour-coding (default=0.01)
plotCutoff	Which probes to include by significance cut-off (default=1 for all markers)
zeroCountCutoff	Which probes to include by minimum counts cut-off (default=50)
colours	Vector of colours to use for significance groups
labels	Row names or indices to label on plot
fontSize	Font size
labelFontSize	Font size for labels
useAdjusted	whether to use adjusted p-values (must have q-values in object)
graphics	Either "ggplot" or "plotly"
verbose	Whether to print statistics

### Value

List of three plots. One plot for each x2Value and one combined figure

### Examples

```

data(PEAC_minimal_load)

disp <- apply(tpm, 1, function(x){
  (var(x, na.rm=TRUE)-mean(x, na.rm=TRUE))/(mean(x, na.rm=TRUE)**2)
})

resultTable <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  countdata = tpm[1:5, ],
  metadata = metadata,
  dispersion = disp)

```

```
plots <- maPlot(resultTable,
                x1var='Timepoint',
                x2var='EULAR_6m',
                x2Values=c('Good', 'Non-response'),
                graphics="plotly")

plots$combined
```

---

metadata	<i>Minimal metadata from PEAC</i>
----------	-----------------------------------

---

### Description

Minimal metadata for paired longitudinal response analysis.

### Usage

```
metadata
```

### Format

A data frame

**PATID** Id for matching patients

**Timepoint** timepoints

**EULAR\_6m** response data

---

modelPlot	<i>Mixed model effects plot</i>
-----------	---------------------------------

---

### Description

Plot to show differences between groups over time using base graphics.

### Usage

```
modelPlot(
  object,
  geneName = NULL,
  x1var = NULL,
  x2var = NULL,
  x2shift = NULL,
  xlab = NA,
  ylab = geneName,
  plab = NULL,
  title = geneName,
```



```

logTransform = is(object, "GlmSeq"),
shapes = 21,
colours = "grey60",
lineColours = "grey60",
markerSize = 0.5,
fontSize = NULL,
alpha = 0.7,
addModel = TRUE,
addPoints = TRUE,
modelSize = 2,
modelColours = "royalblue",
modelLineSize = 1,
modelLineColours = modelColours,
errorBarLwd = 2.5,
errorBarLength = 0.05,
...
)

```

### Arguments

object	A glmmSeq/lmmSeq object created by <code>glmmSeq::glmmSeq()</code> or <code>glmmSeq::lmmSeq()</code>
geneName	The gene/row name to be plotted
x1var	The name of the first (inner) x parameter, typically 'time'. This is anticipated to have different values when matched by ID.
x2var	The name of an optional second (outer) x parameter, which should be a factor.
x2shift	Amount to shift along x axis for each level of x2var. By default the function will arrange each level of x2var side by side. Lower values of x2shift or x2shift = 0 can be used to overlap plots similar to 'dodge' or stagger them.
xlab	Title for the x axis
ylab	Title for the y axis
plab	Optional character vector of labels for p-values. These must align with column names in <code>object@stats\$pvals</code> .
title	Plot title. If NULL gene name is used
logTransform	Whether to perform a log10 transform on the y axis
shapes	The marker shapes (default=19)
colours	The marker colours (default='red') as vector or named vector
lineColours	The line colours (default='grey60') as vector or named vector
markerSize	Size of markers (default=2)
fontSize	Plot font size
alpha	Line and marker opacity (default=0.7)
addModel	Whether to add the fit model with markers (default=TRUE)
addPoints	Whether to add underlying data points (default=TRUE)
modelSize	Size of model points (default=2)

**modelColours**    Colour of model fit markers (default="black") as vector or named vector  
**modelLineSize**    Size of model points (default=1) as vector or named vector  
**modelLineColours**  
                     Colour of model fit lines.  
**errorBarLwd**      Line width of error bars  
**errorBarLength**    Head width of error bars  
**...**              Other parameters to pass to `graphics::plot()`

**Value**

Returns a paired plot for matched samples

**Examples**

```

data(PEAC_minimal_load)

disp <- apply(tpm, 1, function(x){
  (var(x, na.rm=TRUE)-mean(x, na.rm=TRUE))/(mean(x, na.rm=TRUE)**2)
})

MS4A1glmm <- glmmSeq(~ Timepoint * EULAR_6m + (1 | PATID),
  countdata = tpm[1:2, ],
  metadata = metadata,
  dispersion = disp)

modelPlot(object=MS4A1glmm,
  geneName = 'MS4A1',
  x1var = 'Timepoint',
  x2var='EULAR_6m')

```

---

summary.lmmSeq

*Summarise a 'glmmSeq'/'lmmSeq' object*


---

**Description**

Summarise results from `glmmSeq` or `lmmSeq` analysis

**Usage**

```

## S3 method for class 'lmmSeq'
summary(object, gene = NULL, digits = max(3L, getOption("digits") - 3L), ...)

## S3 method for class 'GlmmSeq'
summary(object, gene = NULL, ...)

```

**Arguments**

object	an object of class "GlimmSeq" or "lmmSeq"
gene	an optional character value specifying a single gene whose results are summarised
digits	integer, used for number formatting
...	arguments to be passed to other methods

**Value**

If gene=NULL a dataframe of results for all genes is returned. Otherwise the output of GLMM or LMM model results for a single gene including coefficients, test statistics, p-values is printed and the dataframe for all genes is returned invisibly.

**See Also**

[glimmSeq\(\)](#), [lmmSeq\(\)](#)

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tpm	<i>TPM count data from PEAC</i>
-----	---------------------------------

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**Description**

Transcripts Per Million (TPM) count data for PEAC synovial biopsies.

**Usage**

tpm

**Format**

An object of class `matrix` (inherits from `array`) with 50 rows and 123 columns.

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