

# Package ‘denovolyzeR’

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**Title** Statistical Analyses of De Novo Genetic Variants

**Version** 0.2.0

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**Description** An integrated toolset for the analysis of de novo (sporadic) genetic sequence variants. denovolyzeR implements a mutational model that estimates the probability of a de novo genetic variant arising in each human gene, from which one can infer the expected number of de novo variants in a given population size. Observed variant frequencies can then be compared against expectation in a Poisson framework. denovolyzeR provides a suite of functions to implement these analyses for the interpretation of de novo variation in human disease.

**Depends** R (>= 3.1.0)

**Imports** dplyr (>= 0.3), reshape2 (>= 1.4)

**License** GPL-3

**LazyData** true

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**URL** <http://denovolyzeR.org>

**BugReports** <http://github.com/jamesware/denovolyzeR/issues>

**RoxygenNote** 5.0.1

**NeedsCompilation** no

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**Repository** CRAN

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autismDeNovos	<i>de novo variants found in 1,078 autism trios</i>
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### Description

de novo variants found in 1,078 autism trios, published in Nature Genetics(<http://www.nature.com/doifinder/10.1038/ng.3050>)

### Format

A data frame with 1096 obs of 2 variables:

**gene** Gene symbol of gene containing de novo variant

**class** Functional class of variant: "syn" = synonymous, "mis" = missense, "non" = nonsense, "splice" = canonical splice site, "frameshift" = frameshift indel

### References

<http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4222185/>

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denovolyze	<i>Evaluates burden of de novo variation against expectation</i>
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### Description

Determines whether the test population carry more *de novo* variants than expected. Variants may be grouped by variant class (e.g. are there more LOF variants than expected, across the whole dataset?), or by gene (are there more variants of a given class in SCN2A?).

**Usage**

```
denovolyze(genes, classes, nsamples, groupBy = "class",
  includeGenes = "all", includeClasses = c("syn", "mis", "misD", "non",
  "stoploss", "startgain", "splice", "frameshift", "lof", "prot", "protD",
  "all"), geneId = "geneName", signifP = 3, roundExpected = 1,
  probTable = NULL, misD = NULL)
```

```
denovolyzeByClass(genes, classes, nsamples, groupBy = "class",
  includeGenes = "all", includeClasses = c("syn", "mis", "lof", "prot",
  "all"), geneId = "geneName", signifP = 3, roundExpected = 1,
  probTable = NULL)
```

```
denovolyzeByGene(genes, classes, nsamples, groupBy = "gene",
  includeGenes = "all", includeClasses = c("lof", "prot"),
  geneId = "geneName", signifP = 3, roundExpected = 1, probTable = NULL)
```

**Arguments**

genes	A vector of genes containing de novo variants.
classes	A vector of classes of de novo variants. Standard supported classes are "syn" (synonymous), "mis" (missense), "non" (nonsense), "splice" (splice), "frameshift" (frameshift) and "lof" (loss of function = non + splice + frameshift). Additional classes that are supported by the code, but are not included in the built-in probability tables, are "stoploss", "startloss", "misD" (damaging missense). These labels may be used for user-supplied probability tables. If "misD" is present, then "mis" (in the input) implies non-damaging missense.
nsamples	Number of individuals considered in de novo analysis.
groupBy	Results can be tabulated by "gene", or by variant "class"
includeGenes	Genes to include in analysis. "all" or a vector of gene names.
includeClasses	Determines which variant classes are tabulated in output. In addition to the input classes, summaries can be produced for "prot" (protein-altering = mis + lof), "all", and "protD" (protein damaging = misD + lof, only available if misD included in user-specified probability table). If "misD" is present, then "mis" will return statistics for all missense. Non-damaging missense are not analysed separately.
geneId	Gene identifier used. One of "hgncID", "hgncSymbol", "enstID", "ensgID" or "geneName" (default, equals <code>ensembl "external_gene_name"</code> )
signifP	Number of significant figures used to round p-values in output.
roundExpected	Number of decimal places used to round expected burdens in output.
probTable	Probability table. A user-defined table of probabilities can be provided here, to replace the probability table included in the package.
misD	If the user-specified probability table contains probabilities for a sub-category of missense variants (e.g. predicted to be damaging by an in silico algorithm), this column should be called misD, or the alternative name should be specified here.

**Details**

Analyses can be restricted to a subset of genes, and/or a subset of variant classes  
See vignette("denovolyzeR\_intro") for more information.

**Value**

Returns a data frame

**Functions**

- denovolyzeByClass: denovolyzeByClass
- denovolyzeByGene: denovolyzeByGene

**Examples**

```
### denovolyze

denovolyze(genes=autismDeNovos$gene,
           classes=autismDeNovos$class,
           nsamples=1078)

### denovolyzeByClass

denovolyzeByClass(genes=autismDeNovos$gene,
                  classes=autismDeNovos$class,
                  nsamples=1078)

# this convenience function is identical to:

denovolyze(genes=autismDeNovos$gene,
           classes=autismDeNovos$class,
           nsamples=1078,
           groupBy="class",
           includeClasses=c("syn", "mis", "lof", "prot", "all"),
           includeGenes="all"
           )

### denovolyzeByGene

denovolyzeByGene(genes=autismDeNovos$gene,
                 classes=autismDeNovos$class,
                 nsamples=1078)

# this is identical to:

denovolyze(genes=autismDeNovos$gene,
           classes=autismDeNovos$class,
           nsamples=1078,
           groupBy="gene",
           includeClasses=c("lof", "prot"),
```

```
includeGenes="all"
)
```

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denolyzeMultiHits *Determine significance of genes with multiple de novos*

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### Description

Are there more genes containing >1 *de novos* than expected?

### Usage

```
denolyzeMultiHits(genes, classes, nsamples, nperms = 100,
  includeGenes = "all", includeClasses = c("syn", "mis", "lof", "prot",
  "all"), nVars = "actual", geneId = "geneName", probTable = NULL,
  misD = NULL, signifP = 3, roundExpected = 1)
```

### Arguments

genes	A vector of genes containing de novo variants.
classes	A vector of classes of de novo variants. Standard supported classes are "syn" (synonymous), "mis" (missense), "non" (nonsense), "splice" (splice), "frameshift" (frameshift) and "lof" (loss of function = non + splice + frameshift). Additional classes that are supported by the code, but are not included in the built-in probability tables, are "stoploss", "startloss", "misD" (damaging missense). These labels may be used for user-supplied probability tables. If "misD" is present, then "mis" (in the input) implies non-damaging missense.
nsamples	Number of individuals considered in de novo analysis.
nperms	Number of permutations
includeGenes	Genes to include in analysis. "all" or a vector of gene names.
includeClasses	Determines which variant classes are tabulated in output. In addition to the input classes, summaries can be produced for "prot" (protein-altering = mis + lof), "all", and "protD" (protein damaging = misD + lof, only available if misD included in user-specified probability table). If "misD" is present, then "mis" will return statistics for all missense. Non-damaging missense are not analysed separately.
nVars	Select whether expected number of multihits is determined by "expected" total number of variants, or "actual" total. Actual (default) is more conservative.
geneId	Gene identifier used. One of "hgncID", "hgncSymbol", "enstID", "ensgID" or "geneName" (default, equals ensembl "external_gene_name")
probTable	Probability table. A user-defined table of probabilities can be provided here, to replace the probability table included in the package.

misD	If the user-specified probability table contains probabilities for a sub-category of missense variants (e.g. predicted to be damaging by an in silico algorithm), this column should be called misD, or the alternative name should be specified here.
signifP	Number of significant figures used to round p-values in output.
roundExpected	Number of decimal places used to round expected burdens in output.

### Details

See vignette (denovostats\_intro) for more information.

### Value

Returns a data.frame

### Examples

```
denolyzeMultiHits(genes=autismDeNovos$gene,  
                  classes=autismDeNovos$class,  
                  nsamples=1078)
```

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denolyzeR

*A package for the analysis of de novo sequencing variants*

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### Description

A package for the analysis of *de novo* sequencing variants

### Author(s)

James Ware <j.ware@imperial.ac.uk>

### References

<http://github.com/jamesware/denolyzeR>

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fmrpGenes

*FMRP genes*


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### Description

837 genes found to interact with the fragile X mental retardation protein (FMRP)

### Format

A vector of gene symbols

### References

<http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4222185/>

<http://dx.doi.org/10.1016/j.cell.2011.06.013>

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parseInput

*Checks input for errors*


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### Description

An internal function to check inputs

### Usage

```
parseInput(genes = genes, classes = classes, nsamples = nsamples,
  groupBy = groupBy, includeGenes = includeGenes,
  includeClasses = includeClasses, geneId = geneId, signifP = signifP,
  roundExpected = roundExpected, probTable = NULL)
```

### Arguments

genes	A vector of genes containing de novo variants.
classes	A vector of classes of de novo variants. Standard supported classes are "syn" (synonymous), "mis" (missense), "non" (nonsense), "splice" (splice), "frameshift" (frameshift) and "lof" (loss of function = non + splice + frameshift). Additional classes that are supported by the code, but are not included in the built-in probability tables, are "stoploss", "startloss", "misD" (damaging missense). These labels may be used for user-supplied probability tables. If "misD" is present, then "mis" (in the input) implies non-damaging missense.
nsamples	Number of individuals considered in de novo analysis.
groupBy	Results can be tabulated by "gene", or by variant "class"
includeGenes	Genes to include in analysis. "all" or a vector of gene names.

includeClasses	Determines which variant classes are tabulated in output. In addition to the input classes, summaries can be produced for "prot" (protein-altering = mis + lof), "all", and "protD" (protein damaging = misD + lof, only available if misD included in user-specified probability table). If "misD" is present, then "mis" will return statistics for all missense. Non-damaging missense are not analysed separately.
geneId	Gene identifier used. One of "hgncID", "hgncSymbol", "enstID", "ensgID" or "geneName" (default, equals ensembl "external_gene_name")
signifP	Number of significant figures used to round p-values in output.
roundExpected	Number of decimal places used to round expected burdens in output.
probTable	Probability table. A user-defined table of probabilities can be provided here, to replace the probability table included in the package.

**Value**

warning or error if any invalid input, else assigns variables back to parent function

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PermuteMultiHits	<i>Permutes x variants across a genelist, and counts genes with multiple hits</i>
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**Description**

An internal function called by denovolyzeMultiHits

**Usage**

```
PermuteMultiHits(x, y, nperms = 100, class = "lof", geneId = "geneName",
  includeGenes = "all", probTable = pDNM)
```

**Arguments**

x	Total number of de novo variants observed in dataset
y	Number of genes with >1 de novo variant (of class "class") in the population
nperms	Number permutations
class	In c("lof", "mis", "syn", "prot")
geneId	Gene identifier used. One of "hgncID", "hgncSymbol", "enstID", "ensgID" or "geneName" (default, equals ensembl "external_gene_name")
includeGenes	Genes to include in analysis. "all" or a vector of gene names.
probTable	Probability table. A user-defined table of probabilities can be provided here, to replace the probability table included in the package.

**Value**

Returns a named vector of 5 values



**See Also**[denovolyzeMultiHits](#)

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viewProbabilityTable *Displays underlying de novo probability tables*

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**Description**

Tabulates probability of *de novo* variant for each protein-coding variant class, for each gene. Values are probability of a *de novo* variant per chromosome per generation. i.e. expected number of de novos for a given gene/class =  $p * 2 * nsamples$ .

**Usage**

```
viewProbabilityTable(format = "wide")
```

**Arguments**

format                    option to display table in wide format (default; one line per gene), or long format

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