

# Package ‘REBET’

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**Title** The subREgion-based BurdEn Test (REBET)

**Version** 1.26.0

**Date** 2018-08-13

**Description** There is an increasing focus to investigate the association between rare variants and diseases. The REBET package implements the subREgion-based BurdEn Test which is a powerful burden test that simultaneously identifies susceptibility loci and sub-regions.

**Imports** stats, utils

**Depends** ASSET

**Suggests** RUnit, BiocGenerics

**License** GPL-2

**biocViews** Software, VariantAnnotation, SNP

**RoxygenNote** 6.0.1

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## Contents

|                 |   |
|-----------------|---|
| data . . . . .  | 2 |
| REBET . . . . . | 2 |
| rebet . . . . . | 3 |

|              |          |
|--------------|----------|
| <b>Index</b> | <b>5</b> |
|--------------|----------|

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|------|-----------------------------|
| data | <i>Data for the example</i> |
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### Description

Data for the example.

### Details

The data contains a binary phenotype vector response, a genotype matrix genotypes consisting of 20 rare-variant SNPs, and the sub-region annotation vector subRegions for the [rebet](#) example.

### See Also

[rebet](#)

### Examples

```
data(data, package="REBET")

# Display some of the data
table(response)
dim(genotypes)
subRegions
```

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|       |                          |
|-------|--------------------------|
| REBET | <i>The REBET package</i> |
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### Description

An R package for the subREgion-based BurdEn Test (REBET).

### Details

In rare-variant association studies, aggregating rare and/or low frequency variants, may increase statistical power for detection of the underlying susceptibility gene or region. However, it is unclear which variants, or class of them, in a gene contribute most to the association. This subregion-based burden test (REBET) simultaneously selects susceptibility genes and identifies important underlying sub-regions. The sub-regions are predefined based on shared common biologic characteristics, such as the protein domain or possible functional impact. Based on a subset-based approach considering local correlations between combinations of test statistics of sub-regions, REBET is able to properly control the type I error rate while adjusting for multiple comparisons in a computationally efficient manner. See the reference for the details of this test. The main function in this package is [rebet](#), which performs the REBET test.

**Author(s)**

Bin Zhu <bin.zhu@nih.gov>, Lisa Mirabello and Nilanjan Chatterjee

**References**

Zhu, B., Mirabello, L., Chatterjee, N. (2018) A Subregion-based Burden Test for Simultaneous Identification of Susceptibility Loci and Sub-regions within Genetic Epidemiology. In press. <https://doi.org/10.1002/gepi.221>

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|       |                                                |
|-------|------------------------------------------------|
| rebet | <i>The subREgion-based BurdEn Test (REBET)</i> |
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**Description**

A Subregion-based Burden Test for Simultaneous Identification of Susceptibility Loci and Sub-regions within

**Usage**

```
rebet(response, genotypes, subRegions, responseType=NULL,
      covariates=NULL, shape1=1, shape2=1, saveMem=FALSE)
```

**Arguments**

|              |                                                                                                                                                                                                |
|--------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| response     | Numerical vector of phenotypes. A binary phenotype must be coded as 0 and 1.                                                                                                                   |
| genotypes    | Matrix of genotypes with each column as a locus.                                                                                                                                               |
| subRegions   | Sub-region annotation vector with length equal to the number of columns of genotypes. In the returned object, these regions will appear as <code>paste("Region_", subRegions, sep="")</code> . |
| responseType | NULL, "continuous" or "binary". If NULL, then "continuous" or "binary" will be chosen based on Y. The default is NULL.                                                                         |
| covariates   | NULL or matrix of covariates. The default is NULL.                                                                                                                                             |
| shape1       | The shape1 parameter in the beta distribution. The default is 1.                                                                                                                               |
| shape2       | The shape2 parameter in the beta distribution. The default is 1.                                                                                                                               |
| saveMem      | TRUE or FALSE to conserve memory (see details). The default is FALSE.                                                                                                                          |

**Details**

See the reference for details of this method.

Missing values in any of `response`, `genotypes` or `covariates` will be removed before the analysis. Setting `saveMem` to TRUE will allow for the analysis of a much larger number of subjects, but will take more time to compute. When `saveMem` is FALSE, there needs to be enough memory available to hold two or three  $N \times N$  matrices, where  $N$  is the number of subjects.

This function calls the `h.traits` function in the [ASSET](#) package.

**Value**

The object returned from `h.traits` in the `ASSET` package.

**Author(s)**

Bin Zhu <bin.zhu@nih.gov>, Lisa Mirabello and Nilanjan Chatterjee

**References**

Zhu, B., Mirabello, L., Chatterjee, N. (2018) A Subregion-based Burden Test for Simultaneous Identification of Susceptibility Loci and Sub-regions within Genetic Epidemiology. In press. <https://doi.org/10.1002/gepi.221>

**Examples**

```
data(data, package="REBET")

res <- rebet(response, genotypes, subRegions)
h.summary(res)
```

# Index

\* **SNP, gene, rare variant**

rebet, [3](#)

\* **data**

data, [2](#)

\* **package**

REBET, [2](#)

ASSET, [3](#), [4](#)

data, [2](#)

genotypes (data), [2](#)

h.traits, [3](#), [4](#)

REBET, [2](#)

rebet, [2](#), [3](#)

response (data), [2](#)

subRegions (data), [2](#)