

# Package ‘REBET’

April 24, 2025

**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

**git\_url** <https://git.bioconductor.org/packages/REBET>

**git\_branch** RELEASE\_3\_21

**git\_last\_commit** ce98409

**git\_last\_commit\_date** 2025-04-15

**Repository** Bioconductor 3.21

**Date/Publication** 2025-04-23

**Author** Bill Wheeler [cre],  
Bin Zhu [aut],  
Lisa Mirabello [ctb],  
Nilanjan Chatterjee [ctb]

**Maintainer** Bill Wheeler <[wheelerb@imsweb.com](mailto:wheelerb@imsweb.com)>

## Contents

data . . . . .	2
REBET . . . . .	2
rebet . . . . .	3

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

---

data	<i>Data for the example</i>
------	-----------------------------

---

### 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
```

---

REBET	<i>The REBET package</i>
-------	--------------------------

---

### 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>

---

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

---

**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](#)