

Package ‘dQTG.seq’

March 22, 2023

Type Package

Title A BSA Software for Detecting All Types of QTLs in BC, DH, RIL and F2

Version 1.0.2

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Description The new (dQTG.seq1 and dQTG.seq2) and existing (SmoothLOD, G', deltaSNP and ED) bulked segregant analysis methods are used to identify various types of quantitative trait loci for complex traits via extreme phenotype individuals in bi-parental segregation populations (F2, backcross, doubled haploid and recombinant inbred line). The numbers of marker alleles in extreme low and high pools are used in existing methods to identify trait-related genes, while the numbers of marker alleles and genotypes in extreme low and high pools are used in the new methods to construct a new statistic Gw for identifying trait-related genes. dQTG-seq2 is feasible to identify extremely over-dominant and small-effect genes in F2. Li P, Li G, Zhang YW, Zuo JF, Liu JY, Zhang YM (2022, <[doi:10.1016/j.xplc.2022.100319](https://doi.org/10.1016/j.xplc.2022.100319)>).

Depends R (>= 3.5.0)

License GPL (>= 2)

Imports BB, data.table, doParallel, openxlsx, qtl, stringr, writexl, vroom, parallel, foreach

Encoding UTF-8

LazyData true

RoxygenNote 7.2.3

NeedsCompilation no

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Repository CRAN

Date/Publication 2023-03-22 09:10:06 UTC

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BSA	<i>BSA data</i>
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Description

BSA format of F2 dataset.

Usage

```
data(BSA)
```

Details

Input file for dQTG.seq function.

Dodata	<i>To perform method</i>
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Description

To perform method

Usage

```
Dodata(dir, calculatedata, chr, color1, CLO)
```

Arguments

dir	the path of the output
calculatedata	the inputdata
chr	chromosome
color1	the color of the chromosome
CLO	the numbers of CPU

Value

list

Examples

```
data(BSA)
dir<- tempdir()
data.calculatedata<-Readdata1(BSA)
Dodata(dir,calculatedata=data.calculatedata,chr="all",color1="blue",CL0=1)
```

dQTG.seq*Title The function of dQTG.seq*

Description

Title The function of dQTG.seq

Usage

```
dQTG.seq(dir, filegen, chr, color, CL0)
```

Arguments

dir	the path of the output
filegen	the input data
chr	the chromosome
color	the color
CL0	the numbers of CPU

Value

list

Examples

```
data(BSA)
dQTG.seq(dir=tempdir(),filegen=BSA,chr="all",color="blue",CL0=1)
```

Readdata1*Title readdata function*

Description

Title readdata function

Title readdata function

Usage`Readdata1(File)``Readdata1(File)`**Arguments**

File the input file

Value

list

list

Examples

```
data(BSA)
Readdata1(BSA)
data(BSA)
Readdata1(BSA)
```

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