Package 'Funmap2'

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${\sf R}$ topics documented:

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2 FM2.get.covariance

Description

Identifying or estimating the parameters of curve and covariance structure.

Usage

```
FM2.estimate.data(dat, curve.type=NULL, covar.type=NULL, pdf.file=NULL)
```

Arguments

dat	a data object returned by FM2.simulate or FM2.load.data
curve.type	string value indicating the specific curve type. optional values are listed in FM2.get.curve. The curve fitting is performed using the least-squares if 'auto' or NULL is assigned.
covar.type	string value indicating the specific type of covariance structure, optional values are listed in FM2.get.covariance. the MLE process identifies covariance matrix if 'auto' or NULL is assigned.
pdf.file	string value suggesting a PDF file name to illustrate the performance of curve fitting.

Value

A data object with the results of curve fitting and covariance identifying. This function updates or adds the estimation of curve fitting and covariance structure. FM2.load.data illustrates the structure of data object.

Examples

```
# data simulation using the default parameters
dat <- FM2.simulate();
dat;

# estimate the parameter of curve object and covariance structure
dat <- FM2.estimate.data(dat);
dat;</pre>
```

FM2.get.covariance Retriving covariance structure

Description

Retrive a covariance structure which characterizes the correlation between the measured phenotype.

Usage

```
FM2.get.covariance(covar.type)
```

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Arguments

covar.type string value indicating the type of covariance structure, such as "AR1", "SAD1", full list is described in the "details" section

Details

13 covariance structures are implemented in current version, including:

[1]	"AR1"	First-order Autoregressive
[2]	"SAD1"	First-order Structured Antedependence
[3]	"ARMA(1,1)"	First-order Autoregressive Moving Average
[4]	"ARH1"	Heterogeneous Autoregressive
[5]	"CS"	Compound Symmetry
[6]	"CSH"	Heterogeneous Compound Symmetry
[7]	"VS"	Variance Components
[8]	"SI"	Scaled Identity
[9]	"FA1"	Factor Analytic - First-order
[10]	"FAH1"	Heterogeneous Factor Analytic - First-order
[11]	"TOEP"	Toeplitz
[12]	"TOEPH"	Heterogeneous Toplitz
[13]	"HF"	Huynh-Feldt

The following summarize the parameters of covariance structure 'AR1'.

```
> x<-FM2.get.covariance("AR1");
> show(x);
    Class : fg.covariance.AR1
Covar.Type : AR1
Parameters : rho sigma2
```

Value

This functions returns a S4 object of covariance structure. You can use show or print command to check the summary information.

Examples

```
x<-FM2.get.covariance("SAD1");
x;
show(x);</pre>
```

FM2.get.curve

Retriving a curve object.

Description

Retrive a curve object.

FM2.get.curve

Usage

FM2.get.curve(curve.type)

Arguments

curve.type string value indicating the curve type, full list in the details.

Details

9 curves have been implemented in current version, including:

1) "Logistic"

$$g(t) = \frac{a}{1 + b * e^{-r*t}}$$

2) "Bi-Logistic"

$$g(t) = \frac{a1}{1 + b1 * e^{-r1 * t}} + \frac{a2}{1 + b2 * e^{-r2 * t}}$$

3) "Pharmacology"

$$g(t) = \frac{E_{max} * t}{Ec_{50} + t} + E_0$$

4) "Exponential"

$$q(t) = a * e^{-r * t}$$

5) "Bi-Exponential"

$$g(t) = a_1 * e^{-r_1 * t} + a_2 * e^{-r_2 * t}$$

6) "Power"

$$g(t) = a * t^b$$

7) "Legendre2", Legendre Polynomial(2nd-order)

$$g(t) = u_0 + u_1 * t + u_2 * (3 * t^2 - 1)/2$$

8) "Legendre3", Legendre Polynomial(3rd-order)

$$g(t) = u_0 + u_1 * t + u_2 * (2 * t^2 - 1)/2 + u_3 * (5 * t^3 - 3t)/2$$

9) "Legendre4", Legendre Polynomial(4th-order)

$$q(t) = u_0 + u_1 * t + u_2 * (2 * t^2 + 1)/2 + u_3 * (5 * t^3 - 3t)/2 + ...$$

The following introduces the summary infomation of a curve object.

```
> x<-FM2.get.curve("Logistic");
> show(x);
    Class : fg.curve.log
Curve Type : Logistic
Parameters : a b r
    Formula : y = a/(1+b*exp(-r*t))
```

Value

This function reurns a S4 object of curve. The structure is described in the details section. You can use show or print command to check the summary information.

Examples

```
curve <- FM2.get.curve("Logistic");
show(curve);</pre>
```

FM2.load.data

Loading data from the experiment files

Description

Load the experimental data from experiment files.

Usage

```
FM2.load.data( pheno.csv, time.csv, geno.csv, marker.csv, cross.type,
    curve.type = NULL,
    covar.type = NULL,
    pdf.file = NULL,
    log = FALSE )
```

pheno.csv	a CSV file of phenotypic traits . The format is described in the details section of FM2.load.data.
time.csv	a CSV file of measured time. The format is described in the details section of FM2.load.data.
geno.csv	a CSV file of genotype marker. The format is described in the details section of FM2.load.data.
marker.csv	a CSV file of marker definition. The format is described in the details section of ${\sf FM2.load.data}$
cross.type	string indicating the cross type, optional values are "F2", "BC" and "RIL".
curve.type	string indicating the curve type, optional values are "auto", "Logistic", "Exponential", "Power", "Legendre2",, the full list of curve is described in the details section of FM2.get.curve 'auto' or NULL force the function to do curve fitting by calling FM2.estimate.data
covar.type	string indicating the covariance type, including "auto", "AR1", "SAD1", "ARMA", "CS",, the full list of covariance structure is described in the details section of FM2.get.covariance. 'auto' or NULL force the function to estimate the covariance structure by calling FM2.estimate.data
pdf.file	string variable suggesting a PDF file name to illustrate the performance of curve fitting.
log	boolean variable indicating whether logarithm is applied to the phenotype data.

Details

The function returns a data object, which the structure is same as the simulation data, You can use print or str command to check the details in the data object. The following conents are exported by str command

```
List of 6
 $ obj.curve:Formal class 'fg.curve.log' [package "Funmap2"] with 2 slots
 $ obj.covar:Formal class 'fg.covariance.SAD1' [package "Funmap2"] with 3 slots
 $ obj.cross:List of 14
 $ obj.gen :List of 5
               : chr "../populus.BC.geno.csv"
  ..$ geno.csv
  ..$ marker.csv : chr "../populus.BC.marker.csv"
 ..$ marker.obj : NULL
  ..$ marker.table:'data.frame':
                                       275 obs. of 4 variables:
 ....$ Marker : Factor w/ 275 levels "A/15-620D",...: 227 251 122 123 124 244 186 243 272 61 ...
  ....$ Dist : num [1:275] 0 25.1 37 38.2 39.3 ...
  .. ..$ grp_idx: int [1:275] 1 1 1 1 1 1 1 1 1 1 ...
 ....$ Group : Factor w/ 22 levels "D1","D10","D11",..: 1 1 1 1 1 1 1 1 1 1 ...
  ..$ genos.matrix: int [1:78, 1:275] -1 -1 0 1 0 1 0 1 0 0 ...
  ....- attr(*, "dimnames")=List of 2
 ....$ : chr [1:78] "1" "10" "11" "12" ...
  ....$ : chr [1:275] "marker1" "marker2" "marker3" "marker4" ...
 $ obj.phe :List of 12
  ..$ pheno.csv : chr "../populus.BC.pheno.csv"
  ..$ time.csv
                  : NULL
                 : logi FALSE
 ..$ log
  ..$ sample.obs : int 78
  ..$ sample.times : int [1:11] 1 2 3 4 5 6 7 8 9 10 ...
 ..$ pheY
                 : num [1:78, 1:11] 1.3 2.1 1 1.7 1.2 1.1 0.9 1 1.8 2.4 ...
 ..$ pheX
                 : NULL
  ..$ pheT
                  : int [1:78, 1:11] 1 1 1 1 1 1 1 1 1 1 ...
  ..$ est.covar
                  :List of 2
  ..$ est.curve
                  :List of 7
  ..$ summary.curve:List of 5
  ....$ type : chr "Logistic"
  ....$ par : num [1:4] -6.401 32.937 5.131 0.455
  ....$ summary:'data.frame': 10 obs. of 9 variables:
 .. ... $ type: Factor w/ 10 levels "ABRK", "Bi-Exponential", ...: 8 3 1 9 4 2 10 5 6 7
  .. .. ..$ parm: num [1:10] 3 6 4 3 2 4 2 3 4 5
  .. .. ..$ AIC : num [1:10] 361 367 363 366 363 ...
  .. .. ..$ AICc: num [1:10] 5.66 5.75 5.69 5.72 5.68 ...
  ....$ BIC : num [1:10] 368 381 372 373 368 ...
  .....$ SSE : num [1:10] 7371 7370 7879 7766 ...
  .. .. ..$ MSE : num [1:10] 8.59 8.59 8.59 9.18 9.05 ...
  .....$ RMSE: num [1:10] 2.93 2.93 2.93 3.03 3.01 ...
  .....$ R2 : num [1:10] 0.00491 0.00491 0.00486 0.07421 0.05876 ...
  ..$ summary.covar:List of 4
              : chr "SAD1"
  .. ..$ type
 .. ..$ par
               : num [1:2] 1.055 -0.819
  ....$ summary: 'data.frame': 12 obs. of 4 variables:
 .....$ type: Factor w/ 12 levels "AR1", "ARH1", "ARMA(1,1)",...: 1 8 3 2 4 5 12 9 6 7 ...
```

```
.....$ L : num [1:12] -1249 -1132 -1227 -1190 -1850 ...
.....$ AIC : num [1:12] 2502 2268 2460 2404 3703 ...
.....$ BIC : num [1:12] 2506 2273 2468 2432 3708 ...
- attr(*, "class")= chr "FM2.dat"
```

The phenotype file, measured time file, genotype file and marker definition file must be a CSV file. The following sections illustrate the format of each data file.

1) The phenotype file.

The first column is individual ID and the rest columns are sample data for every measurement. It looks like the following file.

```
ID. 1st 2nd 3rd 4th 5th 6th 7th
1 2.9033 4.118 6.1495 7.8161 9.8379 12.963 14.918
2 4.3306 5.3783 7.0647 9.3624 11.439 NA 15.701
3 2.3997 4.052 5.5431 7.6933 9.8471 NA 12.849
4 3.3044 4.154 5.8924 7.7133 9.2144 10.945 NA
...
```

Please note missing data is coded as space or NA in all four data files.

2) **The measurement time file.** The first column is individual ID and the rest columns are sample data for each measurement. It looks like the following file.

```
ID. 1st 2nd 3rd 4th 5th 6th 7th
1 1 2 3 4 5 6 7
2 1 2 3 4 5 NA 7
3 1 2 3 4 5 NA 7
4 1 2 3 4 5 6 NA
...
```

3) **The genotype file.** The first column is individual ID and the rest columns are genotype data for each marker. An example is shown in the following table. Three genotypes (aa=0, Aa=1, AA=2) and missing data(coded as NA or -1) are valid maker values.

```
ID. marker1 marker2 marker3 marker4 marker5 marker6
1 1 1 0 1 NA 0
2 1 1 1 1 0 0
3 1 1 1 0 1 1
...
```

4) **The marker list file.** The first column is marker's ID, the rest columns are the marker's name, distance, group index, and group name for every marker. In the marker file, the distance field is a distance (in cM) in one chromosome or linkage group. The header row should be included.

```
id Marker Dist. Grp_idx Group
1 marker1 0 1 G1
2 marker2 20 1 G1
3 marker3 40 1 G1
```

Value

This function returns a S3 object with the class label of FM2.dat which structure is identical with the result from FM2.simulate.

```
obj.cross
                  the cross object.
                  the curve object.
obj.curve
obj.covar
                  the covariance structure(object).
                  the genotype data, including geno.csv, marker.csv, marker.obj, marker.table,
obj.gen
                  genos.matrix.
                  the phenotype list, each item is explained in below section.
obj.phe
obj.phe$pheno.csv
                  the phenotype file.
obj.phe$time.csv
                  the measured time file.
obj.phe$sample.obs
                  sample size
obj.phe$sample.times
                  the measure times
                  boolean value indicating whether log function is applied to the phenotype data
obj.phe$log
obj.phe$pheY
                  matrix, longitudinal phenotype traits
obj.phe$pheX
                  matrix, covariate data for all individuals.
obj.phe$pheT
                  matrix, measured times for all individuals.
obj.phe$est.covar
                  the estimation for covariance structure obtained from the call of FM2.estimate.data.
obj.phe$est.curve
                  the estimation for curve object obtained from the call of FM2.estimate.data.
obj.phe$summary.curve
                  the curve fitting results for all selected curves, if curve. type is NULL or 'auto',
                  all available curves are estimated.
sobj.phe$ummary.covar
                  the results of covariance estimation for all selected covariances, if covar. type
                  is NULL or 'auto', all available covariances are estimated.
```

Examples

FM2.permutation 9

Description

Execute permutation to get the cutoff value for significance levels p=0.05 and 0.01.

Usage

```
FM2.permutation( dat, res, grp.idx=NULL, options=list() )
```

Arguments

dat	a data object returned by FM2.simulate or FM2.load.data
res	a result object returned by FM2.qtlscan
grp.idx	a numeric vector indicating which chromosomes or groups get involved in the permutation test.
options	optional value for permutation control, including. debug : default=FALSE, indicating whether this function outputs the debug information. n.cores : default=1, the number of cpu cores for parallel computation. scan.step : default=1, an interval distance used to scan flanking marker, default is 1cm.

permu.loop: default=100, the count of permutation loop. **permu.filter.ratio**: default=1, indicating whether fast estimation algorithm on the basis of QTL filter is used or not. No any optimation for permutation with the default value(=1). If 0.01 is specified, permutation is performed on top 0.01

QTLs which are highly associated with phenotypic traits.

Details

If permutation count is greater than 100, more precise cutoff will be obtained. For example, 10000 times permutation can give the significance table which looks like the following table.

0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9
0.01	0.02	0.03	0.04	0.05	0.06	0.07	0.08	0.09
0.001	0.002	0.003	0.004	0.005	0.006	0.007	0.008	0.009
0.0001	0.0002	0.0003	0.0004	0.0005	0.0006	0.0007	0.0008	0.0009

If clusters or multiple CPU cores are available, the permutaion can use **snowfall** package to do parallel computation. In order to do that, the following is necessary.

- 1) **snowfall** is installed.
- 2) The cluster count should be specified in the options parameter

```
e.g. options=list(n.cores=10)
```

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Value

This function returns a new result object with the update of permutation results. The result objects is described in FM2.qtlscan.

Here we explain the updated results only in the item of obj.perm, a S3 object mainly including a p-value matrix(pv.table).

cross.type	the cross type
curve.type	the curve type
covar.type	the type of covariance structure
permu.loop	the permutation count
param	a list recording the paramters in this functrion calling, including permu.loop, permu.filter.ratio, scan.step, n.cores
full.res	matrix recording all the permutation results
pv.table	matrix which has two columns, the first column is significance leve and nd the second column is cutoff value.

Examples

```
dat <- FM2.simulate();
ret <- FM2.qtlscan(dat);
ret <- FM2.permutation(dat, ret, options=list(n.cores=10, permu.filter.ratio = 0.02, scan.step=2));
# only print the permutation part in the result object
show(ret$obj.permu);
# draw cutoff curve based on permutation results.
plot(ret$obj.permu, pdf.file="test.FM2.permu.pdf");</pre>
```

FM2.pipe

QTL mapping pipeline

Description

Perform standard pipeline for the experiment data based on Functional Mapping framework.

Usage

```
FM2.pipe( pheno.csv, time.csv, geno.csv, marker.csv, cross.type,
    curve.type = NULL,
    covar.type = NULL,
    model = "MLE",
    grp.idx = NULL,
    pdf.prefix = NULL,
    threshold = 0.05,
    threshold.type = "pvalue",
    options = list() )
```

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Arguments

pheno.csv	a CSV file of phenotypic traits . The format is described in the details section of ${\sf FM2.load.data}.$
time.csv	a CSV file of measure time. The format is described in the details section of $\ensuremath{FM2.load.data}.$
geno.csv	a CSV file of genotype marker. The format is described in the details section of $\ensuremath{FM2.load.data}.$
marker.csv	a CSV file of marker definition. The format is described in the details section of ${\sf FM2.load.data}$
cross.type	string value indicating the cross type, including "F2", "BC" and "RIL".
curve.type	string value indicating the curve type, including "auto", "logistic", "Exponential", "Power", "Legendre2",, the full list is described in the details section of FM2.get.curve.
covar.type	string value indicating the type of covariance structure, including "auto", "AR1", "SAD1", "ARMA", "CS",, the full list is described in the details section of FM2.get.covariance.
model	string value indicating the computation algorithm, currently only "MLE" is option.
grp.idx	a numeric vector indicating which chromosomes or groups get involved in the QTL scaning.
pdf.prefix	string value indicating the pefix name of pdf file exported by the pipeline
threshold	a numeric value indicating the criteria of signicant QTLs.
threshold.type	string value indicating the selection method of significant QTL, three optional values. 'pvalue', 'LR' and 'count'
options	optional values for the pipeline, see the following details.

Details

The options paramaters can slightly adjust the results and greatly speed up the computational process. the below explains all items in the options list:

- 1) **debug**, default=FALSE, indicating whether this function outputs the debug information.
- 2) **n.cores**, default=1, the cluster count or multiple cores for parallel permutation, used in FM2.qtlscan and FM2.permutation.
- 3) **scan.step**, default=2, an interval distance used to scan flanking marker, default is 1cm, used in FM2.qtlscan.
- 4) **peak.count**, default=5, a number determines how many significant QTLs will be selected, used in FM2.select.qtl.
- 5) **permu.loop**, default=100, the count of permutation loop, used in FM2.permutation.
- 6) **permu.filter.ratio**, default=1, indicating whether fast estimation algorithm on the basis of QTL filter is used or not in FM2.permutation. No any optimation for permutation under the default condition(=1). If 0.01 is specified, permutation is performed on top 0.01 QTLs which are highly associated with phenotypic traits.

FM2.pipe is a main pipeline of the Funmap2 package, it encapsulates a consecutive procedures, including:

1) Loading the phenotype, genotype and marker file, which is performed in the function FM2.load.data.

FM2.pipe

- 2) Summarize the data object returned by the function FM2.load.data
- 3) Performing the hypothesis tests on all chromosomes or specified chromosomes, which is performed in the function FM2.qtlscan.
- 4) Summarize the result object returned by the function FM2.qtlscan
- 5) Taking a long time to execute permutation parallelly or not, which is performed in the function FM2.permutation.
- 6) Selecting the significant QTLs according to the selection method and threshold, which is performed in the function FM2.select.qtl.
- 7) Outputing a PDF report which includes the summary of the data and QTL scaning results, which is performed in the function FM2.report

Value

The data object and the result object of hypothesis tests are returned in a list object,

data object with the S3 class label of "FM2.dat", the structure is decribed in the details section of FM2.load.data

ret result object with the S3 class label of "FM2.qtl.mle", the structure is decribed

in the details section of FM2.qtlscan

Examples

```
# Load the pre-installed data for the example
file.pheno.csv <- system.file("extdata", "populus.BC.pheno.csv", package="Funmap2")</pre>
file.geno.csv <- system.file("extdata","populus.BC.geno.csv", package="Funmap2")</pre>
file.marker.csv <- system.file("extdata", "populus.BC.marker.csv", package="Funmap2")</pre>
# Call the pipeline without permutation.
# Can't select QTL using pvalue due to the missing of permutation result.
r <- FM2.pipe( file.pheno.csv, NULL, file.geno.csv, file.marker.csv, "BC",</pre>
    curve.type="logistic",
    covar.type="auto",
    grp.idx = c(1:5),
    threshold = 3,
    threshold.type = "count",
    options=list(permu.loop=0) );
# Show the summary information of data object
show(r$dat);
# Show the summary information of result object
show(r$ret);
# Change the QTL criteria
r$ret <- FM2.select.qtl(r$ret, threshold = 40, threshold.type = "LR" );
show(r$ret);
# Make a report for the data analysis.
```

FM2.qtlscan

```
FM2.report("test.FM2.pipe.pdf", r$dat, r$ret );
```

FM2.qtlscan QTL scanning

Description

Perform QTL scaning for all QTLs to detect the significant ones based on the hypothesis test.

Usage

```
FM2.qtlscan( dat, model="MLE", grp.idx=NULL, options=list() )
```

Arguments

dat	a data object returned by FM2.simulate or FM2.load.data
model	string value indicateing which method will be used to test hypothesis, one optional value currently.
grp.idx	a numeric vector indicating which chromosomes or groups get involved in the QTL scaning.
options	optional list for QTL scanning, including: debug: default=FALSE, indicating whether this function outputs the debug information. n.cores: default=1, a number of cpu cores for parallel computation. scan.step: default=1, an interval distance used to scan flanking marker, default is 1cm. peak.count: default=5, a number indicating how many top(or significant) QTLs will be selected.

Details

This function returns a result object which can be inspected by the following method:

- 1) str command.
- 2) print command.
- 3) summary command.
- 4) plot command.

Hypothesis Test: For different genotypes, all parameters are identical.

The hypothesis testing scans every marker by the specified step (1cm). It maybe take a long time, so the Funmap2 package displays its progress after each chromosome (linkage group) has been calculated.

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After the QTL scanning, the package identifies the 5 top QTLs. At most one significant QTL is selected within each choromosome (group). The top QTLs are strongly displayed at the head of report.

Value

This function returns a S3 object with the class label of FM2.qtl.mle, including the following items:

	param	list recording the parameters for this function call.
	obj.phe	the phenotype data, copied from the data object (dat).
	obj.gen	the genotype data, copied from the data object (dat).
	obj.curve	the curve object, copied from the data object (dat).
	obj.covar	the covariance structure, copied from the data object (dat).
	obj.cross	the cross object, copied from the data object (dat).
	cross.type	the cross type, copied from the data object (dat).
	covar.type	the type of covariance structure, copied from the data object (dat).
	curve.type	the curve type, copied from the data object (dat).
	full.res	a matrix for all QTLs with the postion, likelihood ratio, curve parameters of different genes and covariance parameters.
	threshold.type	the selection method of significant QTLs, used in FM2.select.qtl
	threshold	the criteria of significant QTLs, used in FM2.select.qtl
qtl.peaks a numeric vector indicating the row index of significant QTLs in a This item is available after the calling of FM2.select.qtl		a numeric vector indicating the row index of significant QTLs in above matrix. This item is available after the calling of FM2.select.qtl $$
	obj.permu	the permutation result, obtained from the calling of FM2.permutation

Examples

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FM2	.report	
1 112	. 1 CDO1 L	

PDF report for data and result object

Description

Output a PDF report including the summary information and figures for the data and result object.

Usage

```
FM2.report( file.report.pdf, dat, res=NULL, options=list( debug=F ) )
```

Arguments

```
file.report.pdf
```

PDF file name.

dat a data object returned by FM2.simulate or FM2.load.data

res a result object returned by FM2.qtlscan

options option list including whether debug information is outputted.

Details

This function don't use the HaruPDF package anymore!!! It outputs the summary information and figures into a PDF file for the data object and the result object.

The following link is an example for FM2. report.

```
http://statgen.psu.edu/software/funmap/report_demo.pdf
```

Value

No return value.

Examples

```
dat <- FM2.simulate();
res <- FM2.qtlscan(dat);
FM2.report("test.FM.report.pdf", dat, ret);</pre>
```

FM2.select.qtl

Selecting significant QTLs

Description

Select significant QTLs according to the threshold and method.

Usage

```
FM2.select.qtl( res, threshold=0.05, threshold.type="pvalue" )
```

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Arguments

```
res a result object returned by FM2.qtlscan
threshold numeric value indicating the criteria of signicant QTLs.
threshold.type string value indicating the selection method of significant QTL, three optional values. 'pvalue', 'LR' and 'count'.
```

Details

Three methods can be used to show the significant QTLs:

- 1) use show or print command to check the significant QTLs;
- 2) use the plot command to show the significant QTLs and genetic curve at significant QTLs;
- 3) access the items of result object

```
e.g.
> res <- FM2.select.qtl(res, threshold=5, threshold.type="count")
> cat("The significant QTL list:\n");
> show( res$full.res[ res$qtl.peaks, 1:3] );
```

Value

A results with updated significant QTLs is returned.

Examples

FM2.simu.pipe

Pipeline for simulation test

Description

Demostrate the simulation test using the pipeline function.

FM2.simu.pipe

Usage

```
FM2.simu.pipe( cross.type = "BC", curve.type="Logistic", covar.type="AR1",
  simu.mrkdist = rep(20,10),
  simu.qtlpos = 95,
  simu.obs = 800,
  simu.times = 8,
  par.X = NULL,
  par0 = NULL,
  par1 = NULL,
  par2 = NULL,
  par.covar = NULL,
  phe.missing = 0.01,
  marker.missing = 0.01,
  threshold = 0.05,
  threshold.type = "pvalue",
  model = "MLE",
  pdf.prefix = NULL,
  options = list() )
```

Ī		
	cross.type	string value indicating the cross type, including "F2", "BC" and "RIL".
	curve.type	string value indicating the curve type, including "logistic", "Exponential", "Power" "Legendre2",, the full list is described in the details section of FM2.get.curve.
	covar.type	string value indicating the covariance type, including "AR1", "SAD1", "ARMA", "CS",, the full list is described in the details section of FM2.get.covariance.
	simu.mrkdist	numeric vector indicating the distance between the genomic marker.
	simu.qtlpos	numeric value indicating the significant QTL position.
	simu.obs	numeric value indicating the sample size.
	simu.times	numeric value indicating the measured times.
	par.X	numeric vector indicating covariate parameters.
	par0	numeric vector indicating curve parameters for gene QQ, default value is retrived from the curve object.
	par1	numeric vector indicating curve parameters for gene Qq, default value is retrived from the curve object.
	par2	numeric vector indicating curve parameters for gene qq, default value is retrived from the curve object.
	par.covar	numeric vector indicating covariance parameters, default value is retrived from the covariance structure.
	phe.missing	numeric value indicating the missing rate of phenotypic traits.
	marker.missing	numeric value indicating the missing rate of genomic markers.
	threshold	numeric value indicating the criteria of signicant QTLs.
	threshold.type	string value indicating the selection method of significant QTL, three optional values. 'pvalue', 'LR' and 'count'
	model	string value indicating the computation algorithm, currently only "MLE" is option.
	pdf.prefix	string value indicating the pefix name of pdf file exported by the pipeline
	options	optional list for the pipeline, see the details in FM2.pipe.

FM2.simulate

Details

The options paramater is described in FM2.pipe.

FM2.simu.pipe demostrates how to use the Funmap2 to do a simulation test, which includes the following steps:

- 1) Simulate a raw data object on the basis of the parameters.
- 2) Perform QTL scaning on all QTLs based on the hypothesis test.
- 3) Execute permutation to get a cutoff for significant QTLs.
- 4) Summarize all objects and plot all figures.
- 5) Export a PDF report including all summary information and figures.

Value

A list including the data object(dat) and the result object of QTL scaning (ret).

```
dat data object describled in FM2.simulate

ret result object of QTL scaning with permutation cutoff table describled in FM2.qtlscan
```

Examples

FM2.simulate

Data simulation

Description

Create a simulation data object for pipeline demonstration.

FM2.simulate

Usage

```
FM2.simulate( cross.type = "BC",
    curve.type = "Logistic",
    covar.type = "AR1",
    simu.mrkdist = rep(20,10),
    simu.qtlpos = 95,
    simu.obs = 800,
    simu.times = 8,
    par.X = NULL,
    par0 = NULL,
    par1 = NULL,
    par2 = NULL,
    par.covar = NULL,
    phe.missing = 0.01,
    marker.missing = 0.01,
    pdf.file = NULL )
```

Arguments

cross.type	string value indicating the cross type, including "F2", "BC" and "RIL".
curve.type	string value indicating the curve type, including "logistic", "Exponential", "Power", "Legendre2",, the curve list is described in the details section of FM2.get.curve.
covar.type	string value indicating the covariance type, including "AR1", "SAD1", "ARMA", "CS",, the covariance list is described in the details section of FM2.get.covariance.
simu.mrkdist	numeric vector indicating the distance between the genomic marker.
simu.qtlpos	numeric value indicating the significant QTL position.
simu.obs	numeric value indicating the sample size.
simu.times	numeric value indicating the measured times.
par.X	numeric vector indicating covariate parameters.
par0	numeric vector indicating curve parameters for gene QQ, default value is retrived from the curve object.
par1	numeric vector indicating curve parameters for gene Qq, default value is retrived from the curve object.
par2	numeric vector indicating curve parameters for gene qq, default value is retrived from the curve object.
par.covar	numeric vector indicating covariance parameters, default value is retrived from the covariance object.
phe.missing	numeric value indicating the missing rate of phenotypic traits.
marker.missing	numeric value indicating the missing rate of genomic markers.
pdf.file	string variable suggesting a PDF file name to illustrate the performance of curve fitting.

Details

The structure of simulation data is identical to experiment data object. The different points are listed below:

1) The items of pheno_file, geno_file and marker_file are made up by the Funmap2 package and will be used to assign the output filename as a filename prefix.

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2) In the genotype is coded by 1=Qq 2=QQ for backcross, 0=qq, 1=Qq, 2=QQ for F2 intercross and 0=qq, 2=QQ for RILs intercross.

Value

This function returns a S3 object with the class label of FM2.dat, which structure is same as the experiment data object obtained from the function FM2.load.data.

Examples

```
dat <- FM2.simulate("RIL", "Logistic", "SAD1", simu.obs=1000, simu.times = 7 );
#summarize the data information.
summary( dat );
plot(dat, pdf.file="test.FM2.simulate.pdf");
str(dat);</pre>
```

plot.FM2.dat

Plotting figures of data object

Description

Draw figures for a data object.

Usage

```
## S3 method for class 'FM2.dat'
plot( x, plot.type=NULL, pdf.file=NULL, ... )
```

Arguments

```
x a data object returned by FM2.simulate orFM2.load.data.

plot.type number, the plot type, 1 is for tiled curves and 2 is for overlapping curves.

pdf.file a pdf file name for the figure output, if no pdf file is specified, the plot command can output this figure in the R console.

... additional arguments affecting the plot produced.
```

Details

Two figures can be outputted to R console.

- 1) tiled curves for every individuals.
- 2) overlapping curves for every individuals

An example of this command is available in the following URL.

```
http://statgen.psu.edu/software/funmap/plot.data1.jpg.
http://statgen.psu.edu/software/funmap/plot.data2.jpg.
```

plot.FM2.qtl.mle 21

Examples

```
#check the codes in FM2.simulate() or FM2.load.data()
```

```
plot.FM2.qtl.mle
```

Plotting figures of QTL scanning

Description

Plot the figures based on the results of hypothesis test.

Usage

```
## S3 method for class 'FM2.qtl.mle'
plot(x, plot.type=NULL, pdf.file=NULL, ... )
```

Arguments

```
x a result object of hypothesis tests returned by FM2.qtlscan.

plot.type a number indicating which figure is plotted.

pdf.file a pdf file name for the figure output, if no pdf file is specified, the plot command can output this figure in the R console.

... additional arguments affecting the plot produced.
```

Details

The result object of QTL scaning can output three kinds of figure according to the parameter 'plot.type', including:

- 1) The LR profile for all chromosomes.
- 2) The LR profile for QTL postion.
- 3) The curve for QTL postion.

The examples can be viewed in the following url.

```
http://statgen.psu.edu/software/funmap/plot.t10-1.jpg.
http://statgen.psu.edu/software/funmap/plot.t10-2.jpg.
http://statgen.psu.edu/software/funmap/plot.t10-3.jpg.
```

Examples

```
#check the codes in FM2.qtlscan()
```

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```
plot.FM2.qtl.mle.perm Plotting figure of permutation result
```

Description

Draw a cutoff profile on the basis of the permutation result.

Usage

```
## S3 method for class 'FM2.qtl.mle.perm'
plot(x, pdf.file=NULL, ... )
```

Arguments

```
    an object of permutation result returned by FM2.permutation.
    a pdf file name for the figure output, if no pdf file is specified, the plot command can output this figure in the R console.
    additional arguments affecting the plot produced.
```

Details

This summary exports a figure based on the cutoff table in the permutation result. An example of this command is available in the following URL.

```
https://raw.githubusercontent.com/wzhy2000/Funmap2/master/img/plot.perm.jpg.
```

Examples

```
#check the example in the FM2.permutation()
```

```
summary.FM2.dat Summary of the data object
```

Description

Summarize information for the data object.

Usage

```
## S3 method for class 'FM2.dat'
summary( object, ... )
```

```
object a data object return by FM2.simulate or FM2.load.data ... additional arguments affecting the summary produced.
```

summary.FM2.qtl.mle 23

Details

The data object is described in FM2.load.data.

The following example demonstrates summary command for a data object.

```
Data set for FunMap model:
```

```
Date: 2010-03-19 03:49:36
Model: Logistic Curve
Cross: F2
Pheno. file: simu.pheno.LC.F2
Geno. file: simu.geno.LC.F2
Maker file: simu.marker.LC.F2
Sample size: 100
Sample times: 7
Marker count: 6
LC a: 19.83678
b: 8.96370
r: 0.47202
rho: 0.75430
sigma2: 0.58849
```

Value

No return values, only output the summary information on the R console.

Examples

```
#check the codes in FM2.simulate() or FM2.load.data()
```

```
summary.FM2.qtl.mle Summary of the result object.
```

Description

Summarize information for the QTL scanning results based on the hypothesis test.

Usage

```
## S3 method for class 'FM2.qtl.mle'
summary( object, ... )
```

```
object a result object returned by FM2.qtlscan which stores the results of hypothesis tests.additional arguments affecting the summary produced.
```

Details

The following sections demostrate the context of summary report. including:

- 1) Estimated parameters.
- 2) The significant QTL postions.

```
Hypothesis test 10:
   a2=a1 and b2=b1 and r2=r1
       Model: Logistic Curve
       Cross: Backcross
     QTL pos.: 50.1 (Group:8)
      QTL LR: 66.516
   QTL p-value: 0.000
Grwoth para(Qq): a2= 30.615, b2= 10.776, r2=
                                     0.538
Grwoth para(qq): a1= 23.707, b1= 9.449, r1=
                                     0.615
         rho: 0.953
      sigma2: 8.637
_____
No. Grp Pos. LR a1 b1
                                    r1
                                           a0
                                                   b0
                                                           r0
 1 8 50.100 66.516 30.615 10.776 0.538 23.707 9.449 0.615
 2 12 113.100 55.190 29.865 9.736 0.528 25.207 8.725 0.586
 3 13 12.000 50.963 29.518 9.723 0.526 24.926 8.906
                                                         0.602
 4 18 10.000 25.684 29.236 9.550 0.536 25.523 8.963
                                                         0.584
   1 151.300 24.162 25.998 8.520 0.575 28.801
                                                  9.672
                                                         0.536
```

Examples

```
#check the codes in FM2.qtlscan()
```

```
summary.FM2.qtl.mle.perm
Summary of permutation result
```

Description

Summarize the permutation result.

Usage

```
## S3 method for class 'FM2.qtl.mle.perm'
summary( object, ...)
```

```
object an object of permutation result returned by FM2.permutation.
... additional arguments affecting the summary produced.
```

Details

The summary command gives a table of cutoff values which starts at 90 If the p-value of x permutation count should be greater than 100/x.

The following gives an example of this summary command.

Permutation result:

Curve: Logistic Curve

Cross: BC Loop: 100

p-value	Cutoff
0.90000	0.76583
0.80000	1.45845
0.70000	2.21657
0.60000	3.09488
0.50000	4.16930
0.40000	5.52038
0.30000	7.33172
0.20000	9.97031
0.10000	14.86003
0.09000	15.63818
0.08000	16.50743
0.07000	17.51136
0.06000	18.68964
0.05000	20.09660
0.04000	21.77278
0.03000	23.96467
0.02000	26.98845
0.01000	32.80592

Examples

#check the example in the FM2.permutation()

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```