# A Tutorial On R Package "QTLRel" (V0.2-10 or Later)

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#### 1 Introduction

Advanced intercross lines (AILs) are an ideal resource for fine-scale mapping of quantitative trait loci (QTL). Unlike populations such as F<sub>2</sub>, individuals in an AIL population are genetically unequally related. The unequal relatedness among individuals requires appropriate statistical models and computational tools. We have developed a computationally efficient R package 'QTLRel' for analysis of quantitative AIL data. Users can use the package to calculate condense identity coefficients (CIC) from large pedigrees, estimate variance-covariance component parameters, perform a genome-scan for QTL, assess genome-wide significance and plot results.

Though initially designed for analysis of AIL data, almost all of the major functions have a very general usage. For instance, the function scanOne can be used to perform a genome scan for nuclear family data where relatedness (or population structure) is a concern as long as relationship matrices are available.

To help users to understand and use the package 'QTLRel', we provide this tutorial with concrete examples that illustrate the process of data analyses using the package. However, this tutorial should not take the place of the R documentation; instead, it is better regarded as supplemental material.

Codes in this tutorial can be easily adapted for custom data.

## 2 Install and Load Package

We can install the package from within R using the command

> install.packages("QTLRel")

and load the package using the command

> library(QTLRel)

Package source and documentation are available on CRAN (here).

#### 3 Data Formats

In this tutorial, we'll use part of our real data in F<sub>8</sub>, including a pedigree 'pedF8', phenotype data 'pdatF8', genotype data 'gdatF8' and a genetic map 'gmapF8'. The sample size is 500.

The **pedigree** should be a data frame that includes individual ID 'id', father 'sire' and mother 'dam'. Other information such as sex and generation is optional. If given, 'sex' should be "M", "Male" or 1 for a male and "F", "Female" or 2 (other than 0 and 1) for a female,

and 'generation' can be numeric 0, 1, 2, ... or non-numeric "F0", "F1", "F2"... The special symbol 0 is reserved for unknown IDs. The following are five lines in the pedigree 'pedF8'.

#### > pedF8[137:141,]

		id	sex	${\tt generation}$	sire	dam	family
13	7 :	18411	M	F2	1	2	F1-25
13	8 :	18412	M	F2	1	2	F1-25
13	9 :	18871	F	F3	16164	18257	BrBDF2-01
14	0 :	18843	M	F3	16164	18257	BrBDF2-01
14	1 :	19418	F	F3	18417	18423	BrBDF2-02
>							

The **genotype data** should be a matrix or a data frame, with each row representing an observation and each column a marker locus. The row names (optional) can be individual IDs and the column names should be marker names. The following are five lines in the genotype data 'gdatF8'. Genotypes 1, 2 and 3 correspond to "AA", "AB" and "BB".

#### > gdatF8[1:5,1:5]

	rs6269442	rs13475701	rs13475706	rs3716083	rs3722996
31521	3	3	3	3	2
33424	3	3	3	3	3
31826	2	2	2	2	2
33609	1	1	2	2	1
33275	3	3	3	3	3
>					

The **phenotype data** 'pdatF8' has four columns: 'sex', 'age', 'bwt' and 'cage'. The variable 'bwt' is the body weight of an individual, 'age' is the age (in days) when body weight was measured, and 'cage' is the cage where the individual was bred. The following are five lines in the phenotype data 'pdatF8'. Note that there were a total of 163 cages.

#### > pdatF8[1:5,]

	sex	age	bwt	cage
31521	М	76	25.1	BDF8-57
33424	М	78	25.5	BDF8-144
31826	М	67	17.9	BDF8-63
33609	F	76	16.8	BDF8-132
33275	F	84	22.5	BDF8-119
>				

> length(unique(pdatF8\$cage))

```
[1] 163
>
```

>

The **genetic map** should be a data frame with colums 'snp', 'chr', 'dist', where 'snp' is the SNP (marker) name, 'chr' is the chromosome where the 'snp' is located, and 'dist' is the genetic distance in centi-Morgan (cM) on the chromosome. In 'gmapF8', the extra column 'phyPos' is the physical location (in bp; build 37) of the SNP on the chromosome.

```
> gmapF8[815:819,]
                         dist phyPos37
             snp chr
       rs6191324
815
                   19 54.6570 59396320
816
      rs30705190
                   19 57.7867 62267948
817
      rs13483712
                   X 4.3520
                              9129480
                   X 20.3860 33548080
818
      rs13483724
```

819 gnfX.023.543 X 23.1860 36252886

The above data can be downloaded from here. Suppose the data files are saved in a subfolder "data" under the R working directory. Then we can load the downloaded data using commands

```
> pedF8<- read.csv("data/pedF8.csv",header=TRUE,check.names=FALSE)
> pdatF8<- read.csv("data/pdatF8.csv",header=TRUE,check.names=FALSE)
> gdatF8<- read.csv("data/gdatF8.csv",header=TRUE,check.names=FALSE)
> gmapF8<- read.csv("data/gmapF8.csv",header=TRUE,check.names=FALSE)</pre>
```

These data sets are also saved in the workspace "QTLRelEx.RData" on the above download page, and can be loaded by command

```
> load("QTLRelEx.RData")
```

### 4 Statistical Models

Consider the following statistical model (see Cheng et al (2011) for the general model which this package can deal with)

$$y_i = \mathbf{x}_i' \mathbf{\beta} + x_i^* a^* + z_i^* d^* + u_i + \epsilon_i, \quad i = 1, 2, \dots, n$$
 (1)

where  $y_i$  is the trait value for the *i*-th individual,  $x_i$  represents covariates (e.g. sex) and  $\beta$  are the corresponding effects,  $x_i^*$  is 1, 0 or -1 if the genotype at the putative QTL is AA, Aa or aa and  $a^*$  is the additive effect of the putative QTL,  $z_i^*$  is 1 if the genotype at the

putative QTL is heterozygous or 0 if the genotype is homozygous and  $d^*$  is the dominance effect,  $u_i$  represents polygenic variation, and  $\epsilon_i$  denotes environmental effect. Assume that  $\epsilon_i \sim N(0, \sigma^2), i = 1, 2, \dots, n$  are independent, and  $\mathbf{u} = (u_1, u_2, \dots, u_n)' \sim N_n(\mathbf{0}, \mathbf{G})$  with  $\mathbf{G} = (g_{ij})$  and is independent of  $\boldsymbol{\epsilon} = (\epsilon_1, \epsilon_2, \dots, \epsilon_n)'$ . It is known that in general (e.g. Abney et al, 2000; Cheng et al, 2010)

$$g_{ij} = 2\Phi_{ij}\sigma_{a}^{2} + \Delta_{ij,7}\sigma_{d}^{2} + (4\Delta_{ij,1} + \Delta_{ij,3} + \Delta_{ij,5})Cov(a,d) + \Delta_{ij,1}\sigma_{h}^{2} + (\Delta_{ij,1} + \Delta_{ij,2} - f_{i}f_{j})\mu_{h}^{2} = g_{a,ij}\sigma_{a}^{2} + g_{d,ij}\sigma_{d}^{2} + g_{ad,ij}Cov(a,d) + g_{h,ij}\sigma_{h}^{2} + g_{m,ij}\mu_{h}^{2}$$
(2)

where  $\Phi_{ij}$  is the kinship coefficient between the *i*-th and *j*-th individuals,  $f_i$  is the inbreeding coefficient for the *i*-th individual, and  $\Delta_{ij}$ 's are condensed identity coefficients as defined in Lynch and Walsh (1998, pp.133) and can be calculated from the pedigree data.

## 5 Condensed Identity Coefficients

To fit model (1), we need identity coefficients in (2). Condensed identity coefficients (cic) can be calculated from the pedigree, using function cic.

Usage:

```
cic(ped,ids,inter,df=3,ask=FALSE,verbose=FALSE)
```

The 'ped' is the pedigree that should have at least three components 'id', 'sire' and 'dam'. Preferably, generation information is provided. Otherwise, the program will try to derive generation information based on the pedigree. The 'ids' specifies the IDs of the individuals for which to calculate the Jacquard condensed identity coefficients. If 'ids' is not specified, all individuals in the pedigree 'ped' will be considered. It is recommended that 'ids' be specified only for interested individuals because extra IDs may result in overwhelming computation.

Condensed identity coefficients can be derived from generalized kinship coefficients. Bottom-up and top-down are two strategies for calculating generalized kinship coefficients from a pedigree. The bottom-up approach starts from the target individuals and moves up till the founders. It requires minimal storage but the computational load can be approximately exponential in the number of generations. The bottom-up approach is not realistic if the number of generations is large and the number of individuals in each generation is not small. The top-down approach starts from founders and moves down to the target individuals. The computational load is approximately linear in the number of generations. However, the intermediate generalized kinship coefficients need to be stored, which may require extensive

storage if the number of individuals in a generation is large. This function allows for a hybrid of both bottom-up and top-down approaches by specifying intermediate generations via the argument 'inter' such that generations of too many members are passed over. If 'inter' is missing, the function will use 'df' to determine the "optimal" configuration of 'inter'. The default value of 'inter' is 3. If 'df = 0', then there will be no intermediate generations, which results in implementation of the bottom-up approach. If 'df' is large, then all generations will be used as intermediate generations, which calls for the top-down approach.

We can set 'ask = TRUE' to see if 'inter' produced by the program is feasible. If not, we can tune 'df' until it is satisfactory. Setting 'verbose = TRUE' will print out some messages to track the running process.

The output of cic is a matrix G of nine columns with G[,j] being the j-th Jacquard identity coefficients. Once we finish cic, we can run genMatrix to extract five genetic matrices, including additive genetic matrices "AA", dominance genetic matrix "DD" and other three that are described in Abney et al (2000), as well as inbreeding coefficients "ib".

Note: You may need the **administrative privilege** to run this function on systems such as Windows 7.

**Example 1**. Calculate condensed identity coefficients from a pedigree.

```
> # only interested individuals in F8
> id<- rownames(pdatF8)</pre>
> id[1:5]
[1] "31521" "33424" "31826" "33609" "33275"
> # check if phenotype and genotype data are from the same sample
> sum(!is.element(id,rownames(gdatF8)))
[1] 0
> sum(!is.element(rownames(gdatF8),id))
> sum(!is.element(id,pedF8$id))
[1] 0
> # running
> idcf<- cic(pedF8,ids=id,df=3,ask=TRUE,verbose=TRUE)</pre>
  Total free disk space needed: 577.0453 Mb...
  Carry-over number of individuals in each generation:
 F0
    F1
        F2
             F3 F4
                     F5
                         F6 F7 F8
  2
         72
             64
                 44
                     36
                          53 144 500
  Will go through generations: FO F1 F2 F4 F5 F6 F7 F8
Continue? Yes/No: Yes
```

```
This may take hours or days to finish!
  Please wait or press 'ctrl-c' to abort...
 F1 F2 F4 F5 F6 F7 F8
> # extract genetic matrices
> gmF8<- genMatrix(idcf); names(gmF8)</pre>
[1] "ib" "AA" "DD" "AD" "HH" "MH"
> dim(gmF8$AA)
[1] 500 500
> gmF8$AA[1:3,1:5]
          31521
                    33424
                               31826
                                         33609
                                                    33275
31521 1.3798828 0.7880859 0.3845215 0.7985840 0.7880859
33424 0.7880859 1.3793945 0.4257812 0.7678223 0.8813477
31826 0.3845215 0.4257812 1.0000000 0.3896484 0.4257812
```

Genetic matrices can also be estimated using genotypic data in the sense of identity-by-state (IBS). The function genMatrix has this capability in case there are only two founder strains or genotypes are only recorded as, say, "AA", "AB" and "BB" (QTLRel\_0.2.7 or later).

## 6 Variance Components

Once we have genetic matrices, we can proceed to estimate variance components that are needed in a genome scan. Estimating variance component parameters can be achieved via function estVC.

#### Usage:

```
estVC(y,x,v=vector("list",6),initpar,nit=25,
    method=c("Nelder-Mead","BFGS","CG","SANN"),
    control=list(),hessian=FALSE)
```

where 'y' is a numeric vector or a numeric matrix of one column that represents a phenotype, and 'x', if not missing, is a data frame or matrix that represents covariates. 'v' is a list of interested variance components (AA, DD, HH, AD, MH, EE,...), where "AA" and "DD" are respectively additive and dominance genetic matrices, "EE" is the residual matrix that is usually assumed to be an identity matrix, and "..." are other random components of interest. If a genetic component is not considered, it should be set to "NULL". If interested in Hessian matrix, we can set 'hessian = TRUE'. Here is an example.

Example 2. Estimate variance components.

```
> idx<- !is.na(pdatF8[,"bwt"])</pre>
> pdatTmp<- pdatF8[idx,] # remove missing values</pre>
> gdatTmp<- gdatF8[match(rownames(pdatTmp),rownames(gdatF8)),] # matching
>
> ii<- match(rownames(pdatTmp),rownames(gmF8$AA))</pre>
> vc<- estVC(y = pdatTmp[,"bwt"], x = pdatTmp[,c("sex","age")],</pre>
     v=list(AA=gmF8$AA[ii,ii],DD=gmF8$DD[ii,ii],HH=NULL,AD=NULL,MH=NULL,
            EE=diag(nrow(pdatTmp))))
> vc$value # likelihood
[1] -977.6066
> vc$par # parameter estimates
  (Intercept)
                                       sexM
                                                       age
                                                                       AA
 1.497242e+01 -5.951434e+00
                              0.000000e+00 1.040535e-01 1.863903e+00
           DD
 5.442390e+00 6.350363e-08
```

In the above example, the trait of interest is body weight 'bwt'. We include both additive and dominance genetic variance components but ignore the other three by setting HH=NULL, AD=NULL and MH=NULL. We also consider 'sex' and 'age' as covariates without questioning whether it makes sense to do so. Of course, we can test an effect as illustrated below.

**Example 3**. Include non-genetic variance components.

```
> ranMtr<- rem(~cage, data=pdatTmp) # matrice for random effect (cage)
> names(ranMtr)
[1] "cage"
> dim(ranMtr$cage)
[1] 496 496
> vc.cage<- estVC(y = pdatTmp[,"bwt"], x = pdatTmp[,c("sex","age")],
+ v=list(AA=gmF8$AA[ii,ii],DD=gmF8$DD[ii,ii],HH=NULL,AD=NULL,MH=NULL,
+ EE=diag(nrow(pdatTmp)), cage=ranMtr$cage))
> 2*(vc.cage$value - vc$value) # likelihood ratio test for cage effect
[1] 15.91948
```

The above example demonstrates how to include 'cage' as a non-genetic random effect, and also gives the likelihood ratio test statistic for 'cage' effect. However, we will ignore 'cage' effect as we move on.

# 7 Performing Genome Scans

Now we come to the point to perform a genome scan using function scanOne.

#### Usage:

```
scanOne(y, x, gdat, prdat=NULL, vc=NULL, intcovar=NULL,
numGeno=FALSE, test=c("None","F","Chisq"),
minorGenoFreq=0, rmv=TRUE)
```

The meanings of 'y' and 'x' are the same as in estVC. 'gdat' is genotype data in the format described previously. It is ignored if an object 'prdat' from genoProb is specified as an argument. 'vc' is an object from estVC or aicVC, or an estimated variance-covariance matrix induced by relatedness and environment. 'intcovar', if provided, specifies covariates that interact with QTL. Note that if we treat numerical coding of genotype as numerical value, we should set 'numGeno = TRUE'.

The function will return a list with at least the following components:

- 1) p: P-value at the snp (marker) if 'test' is "F" or "Chisq", or the log-likelihood ratio statistic at the SNP (marker) if 'test' is "None"
- 2) parameters: estimated parameters at all scanning loci, including additive effect 'a' and dominance effect 'd' if 'prdat' is not "NULL".

If the genome scan is based on genotype data 'gdat', then we need make sure there are not missing genotypes. Otherwise, we can call function genoImpute to impute missing genotypes.

#### Usage:

```
genoImpute(gdat,gmap,prd=NULL,step=Inf,gr=2,pos=NULL,
    method=c("Haldane","Kosambi"),na.str="NA",verbose=FALSE)
```

where 'gdat' is genotype data, 'gmap' is genetic map, and 'prd' is an object from genoProb if not "NULL". If we only impute missing genotypes at marker loci, we can specify 'step = Inf' (by default). The argument 'gr' indicates which generation is under consideration. The missing genotype is randomly assigned with a probability conditional on the genotypes of the flanking makers. Currently, this function only works for AIL. Here is an example.

**Example 4.** Genome scan without interactive covariates.

```
> sum(is.na(gdatTmp)) # number of missing genotypes
[1] 142
> gdatTmpImputed<- genoImpute(gdatTmp,gmapF8,gr=8,na.str=NA)
> # genome scan
```

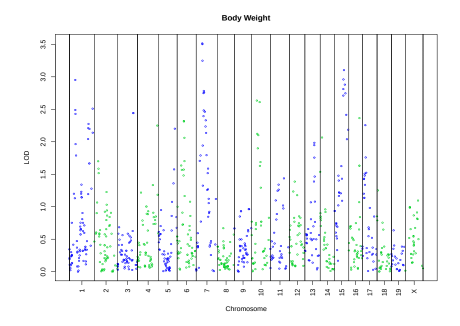


Figure 1 Sex and age are non-interactive covariates.

In the above example, we first call genoImpute to impute 142 missing genotypes and then perform a genome scan. The scan takes both sex and age as non-interactive covariates. We can also include interactive covariates as illustrated in the following example where age is a non-interactive covariate but sex is an interactive covariate. Figure 1 is the Manhattan plot of the mapping results.

**Example 5**. Genome scan with an interactive covariate.

Then QTL by sex interaction effect can be easily obtained as follows.

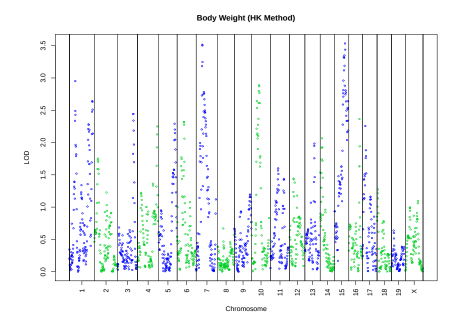


Figure 2 Sex and age are non-interactive covariates.

#### **Example 6**. Extract test statistics.

```
> # QTL by sex interaction
> lrtTmp<- lrt
> lrtTmp$p<- lrt.sex$p - lrt$p
>
> plot(lrtTmp,gmap=gmapF8,main="Body Weight: QTL by Sex Effect") # plotting
>
```

In case of AIL mapping populations, the package 'QTLRel' can also perform Haley-Knott interval mapping. This is achieved by first calling genoProb to calculate genotype probabilities and calling scanOne to scan the genome.

#### Usage:

where 'gdat' is genotype data whose entry should be 1, 2, 3 or 0, corresponding to "AA", "AB", "BB" or missing genotype, and 'gmap' is a genetic map. The argument 'step' defines scanning loci such that the "historical" genetic distance between any two adjacent loci for which probabilities are calculated is not larger than 'step'. If 'step = Inf' (or large enough),

probabilities will only be calculated at loci in both the columns of 'gdat' and the rows of 'gmap'. If 'step' is small, a large set of putative loci will be considered, including all loci defined by the columns of 'gdat and the rows of 'gmap'. The output is a list with the following components:

- 1) pr: a 3-D array with the first dimension equal to that of 'gdat', the second corresponding to three genotype and the third to the putative loci. The probabilities will be -1 if not imputable, which happens when the genotype data is missing at all loci on the chromosome.
- 2) chr: chromosome where the locus is located.
- 3) dist: genetic distance (in cM) of the locus from the first locus on the chromosome.
- 4) snp: SNP (marker) that the locus represents.

Again, it is currently only suitable for advanced intercross lines. Let's look at an example.

**Example 7**. (Haley-Knott method) Genome scan without interactive covariates.

```
> gdTmp<- gdatTmp
> gdTmp[is.na(gdTmp)]<- 0
> unique(c(as.matrix(gdTmp)))
[1] 3 2 1 0
> prDat<- genoProb(gdat=gdTmp,gmap=gmapF8,step=3,gr=8)
>
> # genome scan: Haley-Knott method
> lrtHK<- scanOne(y=pdatTmp[,"bwt"], x=pdatTmp[,c("sex","age")],
+ prd=prDat, vc=vc)
> plot(lrtHK,main="Body Weight (HK Method)") # plotting
```

In the above example, we define 'step = 3', which results in 1741 scanning loci including the 847 markers. Also note that we don't need genetic map information in plotting because the information is contained in the object "lrtHK". Figure 2 displays the mapping results. The following is another example using Haley-Knott interval mapping.

**Example 8.** (Haley-Knott method) Genome scan with sex being an interactive covariate.

```
> # genome scan: Haley-Knott method, interactive sex
> lrtHK.sex<- scanOne(y=pdatTmp[,"bwt"], x=pdatTmp[,c("age")],
+ prd=prDat, intcovar=pdatTmp[,c("sex")],vc=vc)</pre>
```

```
> plot(lrtHK.sex,main="Body Weight (HK Method)") # plotting
> 
> # Haley-Knott method, QTL by sex interaction
> lrtHKTmp<- lrtHK
> lrtHKTmp$p<- lrtHK.sex$p - lrtHK$p
> 
> plot(lrtHKTmp,main="Body Weight (HK Method): QTL by Sex Effect")
>
```

# 8 Empirical Thresholds

Empirical thresholds can be obtained by permutation. This simply repeats the genome scan with genotype data being shuffled among individuals. The function nullSim in the package can be called to estimate thresholds. However, we may write our own code to gain a better control. First, let's look at two permutation examples.

Example 9. Permutation test using marker genotype data.

```
> nn<- nrow(gdatTmpImputed) # sample size
> ntimes<- 1000 # number of simulations
> cvMtr<- NULL # matrix to save results of permuted data
> for(n in 1:ntimes){
        idx<- sample(1:nn,replace=FALSE) # permutation
        tmp<- scanOne(y=pdatTmp[,"bwt"],x=pdatTmp[,c("sex","age")],
            gdat=gdatTmpImputed[idx,], vc=vc)
+ cvMtr<- rbind(cvMtr,tmp$p)
+ cat(n,"/",ntimes,"\r") # track process
+ }
> 00 / 1000
```

In the above example, we shuffle the order of observations and store the shuffled order in variable "idx" so that "gdatTmpImputed[idx,]" is the permuted genotype data from "gdatTmpImputed". The code for genome scan is the same otherwise. The results are stored in the matrix "cvMtr". This process is repeated "ntimes = 1000" times. The 0.05 genome-wide significance threshold can be estimated by

```
> quantile(apply(cvMtr,1,max),0.95)/(2*log(10)) # in LOD
    95%
3.98763
```

Example 10. Permutation test in Haley-Knott interval mapping.

```
> # for HK method...
> cvMtrHK<- NULL
> for(n in 1:ntimes){
+    idx<- sample(1:nn,replace=FALSE)
+    prdTmp<- prDat
+    prdTmp$pr<- prdTmp$pr[idx,,]
+    tmp<- scanOne(y=pdatTmp[,"bwt"], x=pdatTmp[,c("sex","age")],
+    prd=prdTmp, vc=vc)
+    cvMtrHK<- rbind(cvMtrHK,tmp$p)
+    cat(n,"/",ntimes,"\r") # track process
+ }
> 00 / 1000
```

Since we have a pedigree here, we can perform gene dropping test. In the following example, we first call <code>genoSim</code>, which does gene dropping, to generate genotype data set "gdatTmp", and then perform the genome scan using "gdatTmp". This example can be easily adapted to the Haley-Knott version.

#### Example 11. Gene dropping test.

```
> pedR<- pedRecode(pedF8) # recode the pedigree
> ids<- rownames(gdatTmpImputed) # relevant individual IDs
> cvMtrGD<- NULL
> for(n in 1:ntimes){
        gdatTmp<- genoSim(pedR, gmapF8, ids=ids)
        tmp<- scanOne(y=pdatTmp[,"bwt"],x=pdatTmp[,c("sex","age")],
        gdat=gdatTmp, vc=vc)
+ cvMtrGD<- rbind(cvMtrGD,tmp$p)
+ cat(n,"/",ntimes,"\r")
+ }</pre>
```

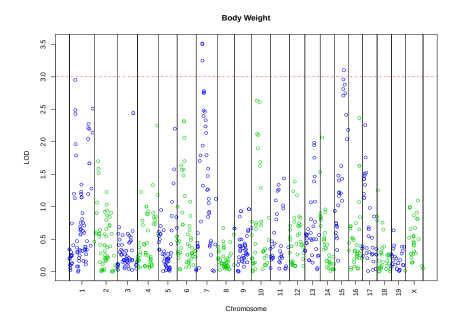


Figure 3 Plotting scanOne objects. Sex and age are non-interactive covariates.

# 9 Plotting

Object of scanOne can be plotted using R function plot, as illustrated in example 4 and 7. Here is one more example, in which we feed a threshold with 'cv' and tune the symbol size with 'cex'.

Example 12. Plotting scanOne objects (Figure 3).

```
> plot(lrt,cv=3,gmap=gmapF8,main="Body Weight (HK Method)",cex=1)
```

We can gain more control using function plotit. Here are two examples of calling function plotit. Users may check the documentation (here) for its usage.

**Example 13**. Plotting mapping results using function plotit (Figure 4).

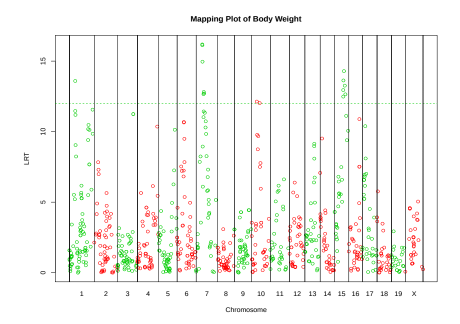


Figure 4 Plotting mapping results using function plotit. Sex and age are non-interactive covariates.

**Example 14**. Plotting mapping results by chromosome using function plotit (Figure 5).

# 10 Miscellaneous

In the package, there are a few other functions that may be useful.

- kinship calculates kinship coefficients from a pedigree.
- maic performs multiple QTL model section via AIC criterion. This function allows additive covariates but not interactive covariates.
- lodci calculates LOD support intervals.

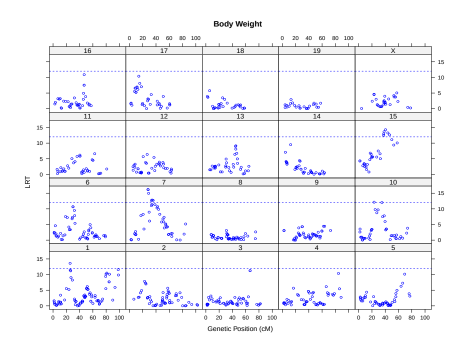


Figure 5 Plotting mapping results by chromosome using function plotit. Sex and age are non-interactive covariates.

Example 15. LOD support intervals based on Haley-Knott interval mapping.

• gls can estimate QTL effects and standard errors by fitting a multiple QTL model. The following example show how to fit a model including two putative QTL at the 674-th and 1365-th scanning loci.

Example 16. Estimated QTL effects and standard errors.

> # multiple QTL model estimates

```
> dtfTmp<- data.frame(</pre>
     y=pdatTmp[,"bwt"],
     age=pdatTmp[, "age"],
     sex=pdatTmp[,"sex"],
    a1=prDat$pr[,1,lc$index[1]]-prDat$pr[,3,lc$index[1]],
     d1=prDat$pr[,2,lc$index[1]],
+
     a2=prDat$pr[,1,lc$index[2]]-prDat$pr[,3,lc$index[2]],
+
     d2=prDat$pr[,2,1c$index[2]]
+ )
> est1<- gls(y~age+sex+a1+d1+a2+d2,data=dtfTmp,vc=vc)</pre>
> est2<- gls(y~age+sex*(a1+d1+a2+d2),data=dtfTmp,vc=vc)</pre>
> est1 # with age and sex being additive covariates
             Estimate Std. Error
                                                  Pr(>|t|)
                                     t value
Intercept 13.97465243 1.60575967
                                   8.7028294 4.977152e-17
age
           0.11225920 0.01591673
                                   7.0529061 6.014461e-12
sexF
          -5.94624248 0.14404054 -41.2817281 1.798428e-161
a1
          0.47577484 0.13217157
                                   3.5996761 3.511094e-04
           0.02658025 0.16582602
                                   0.1602900 8.727189e-01
d1
          -0.35522971 0.13403200 -2.6503350 8.301803e-03
a2
d2
           0.42040674 0.17596958
                                   2.3890876 1.726836e-02
> est2 # with sex being an interactive covariate
             Estimate Std. Error
                                                  Pr(>|t|)
                                      t value
Intercept 13.92886461 1.62037947
                                   8.59605103 1.142566e-16
                                   6.99647702 8.762561e-12
age
           0.11215587 0.01603033
sexF
          -5.82819712 0.27981590 -20.82868431 2.680314e-69
                                   2.88242786 4.121370e-03
           0.46295588 0.16061317
a1
          -0.01350022 0.21945982
                                  -0.06151568 9.509739e-01
d1
a2
          -0.29150868 0.17076744
                                  -1.70705070 8.845267e-02
d2
          0.54345166 0.23708915
                                   2.29218277 2.232256e-02
          0.03156263 0.21682315
                                   0.14556853 8.843225e-01
sexF.a1
sexF.d1
          0.07661133 0.30043387
                                   0.25500231 7.988294e-01
sexF.a2
         -0.14042193 0.22564183
                                  -0.62232222 5.340225e-01
sexF.d2
         -0.26528404 0.33162794 -0.79994479 4.241344e-01
>
```

• qtlVar calculates variance induced by QTL in a quantitative trait.

Example 17. QTL induced variation based on Haley-Knott interval mapping.

```
> ii<- match(rownames(pdatTmp),rownames(gmF8$AA))</pre>
> vcO<- estVC(y = pdatTmp[,"bwt"], v=list(AA=gmF8$AA[ii,ii],</pre>
     DD=gmF8$DD[ii,ii], HH=NULL, AD=NULL, MH=NULL,
     EE=diag(nrow(pdatTmp))))
> nb<- length(vc0$par) - sum(vc0$nn1)</pre>
> nr<- nrow(vc0$y)
> cov<- matrix(0,nrow=nr,ncol=nr)</pre>
> for(i in 1:vc0$nv)
     if(vc0$nnl[i]) cov<- cov + vc0$v[[i]]*vc0$par[nb+vc0$nn[i]]
> tv<- mean(diag(cov)) # total variation
> eff<- NULL # QTL effects
> for(n in 1:length(lrtHK$par)){
     eff<- rbind(eff,lrtHK$par[[n]][c("a","d")])</pre>
+ }
> eff<- data.frame(chr=lrtHK$chr,dist=lrtHK$dist,eff)</pre>
> qv<- qtlVar(eff,prDat$pr) # per QTL variation
> qv[lc$index]/tv*100 # per QTL heritability
[1] 0.9486277 0.9897365
```

The last command calculates variation percentage in body weight that are associated with two highest peaks in figure 2

• scanTwo calculates QTL-by-QTL interaction. Sometimes, people are interested in epistasis. The package provides this facility though it can be computationally expensive if the number of scanning loci is large. In the following example we calculate epistatic effect with sex and age being additive covariates and plot the results.

Example 18. Calculating epistasis and plotting.

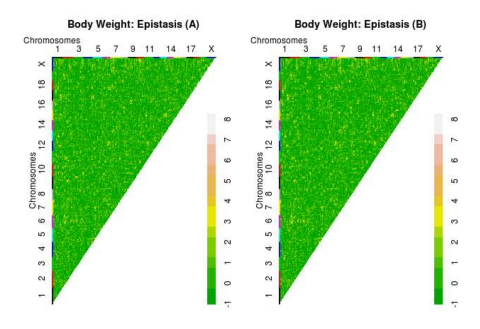


Figure 6 Plotting scanTwo objects. Sex and age are non-interactive covariates.

```
+ main="Body Weight: Epistasis (A)\n\n",xlab="",ylab="")
> plot(qqInt.2/(2*log(10)),
+ gmap=data.frame(snp=prDat$snp,chr=prDat$chr,dist=prDat$dist),
+ main="Body Weight: Epistasis (B)\n\n",xlab="",ylab="")
```

Note that we need a genetic map 'gmap' to plot the scanTwo objects. Refer to the R documentation for usage of scanTwo.

# 11 Citation

Package 'QTLRel' was built on the R and C code that implemented the methodology described in Cheng et al (2010). We ask users of the package to kindly cite the following papers:

Cheng R, Abney M, Palmer PP and Skol AD (2011). QTLRel: an R Package for Genomewide Association Studies in which Relatedness is a Concern. BMC Genet. 12: 66.

Cheng R, Lim JE, Samocha KE, Sokoloff G, Abney M, Skol AD and Palmer AA (2010). Genome-wide association studies and the problem of relatedness among advanced intercross lines and other highly recombinant populations. Genetics 185: 1033-1044.

We wish you success in your research!

## References

- Abney M, McPeek MS, Ober C (2000) Estimation of variance components of quantitative traits in inbred populations. Am J Hum Genet 141:629–650
- Cheng R, Lim JE, Samocha KE, Sokoloff G, Abney M, Skol AD, Palmer AA (2010) Genome-wide association studies and the problem of relatedness among advanced intercross lines and other highly recombinant populations. Genetics 185:1033–1044
- Cheng R, Abney M, Palmer AA, Skol AD (2011) Qtlrel: an R package for genome-wide association studies in which relatedness is a concern. BMC Genetics 12:66
- Lynch M, Walsh B (1998) Genetics and analysis of quantitative traits, vol 5. Sinauer Associates, Inc