# **Haplo Stats**

(version 1.6.0)

# Statistical Methods for Haplotypes When Linkage Phase is Ambiguous

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# Contents

1	$\mathbf{Intr}$	oduction	4							
	1.1	Updates	4							
	1.2	Operating System and Installation	4							
	1.3	R Basics	4							
	ъ.		_							
2		a Setup	5							
	2.1	Example Data	5							
	2.2	Creating a Genotype Matrix	6							
	2.3	Preview Missing Data: summaryGeno	6							
	2.4	Random Numbers and Setting Seed	8							
3	Han	plotype Frequency Estimation: haplo.em	8							
•	3.1	Algorithm	8							
	3.2	Example Usage	8							
	3.3	Summary Method	9							
	3.4	Control Parameters for haplo.em	10							
	3.5	Haplotype Frequencies by Group Subsets	11							
	0.0	Traplotype Troquencies by Group Subsection 1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.								
4	Pow	ver and Sample Size for Haplotype Association Studies	<b>12</b>							
	4.1	Quantitative Traits: haplo.power.qt	12							
	4.2	Case-Control Studies: haplo.power.cc	14							
5	Цор	plotype Score Tests: haplo.score	15							
J	5.1	Quantitative Trait Analysis	15 15							
	$5.1 \\ 5.2$	Binary Trait Analysis	16							
	5.2 $5.3$	v	17							
		Ordinal Trait Analysis	18							
	5.4	Haplotype Scores, Adjusted for Covariates								
	5.5	Plots and Haplotype Labels	19							
	5.6	Skipping Rare Haplotypes	21							
	5.7	Score Statistic Dependencies: the eps.svd parameter	21							
	5.8	Haplotype Model Effect	22							
	5.9	Simulation p-values	22							
6	Reg	Regression Models: haplo.glm 24								
	6.1	New and Updated Methods for haplo.glm	24							
	6.2	Preparing the data frame for haplo.glm	24							
	6.3	Rare Haplotypes	25							
	6.4	Regression for a Quantitative Trait	25							
		Fitting Haplotype x Covariate Interactions	27							
	6.6	Regression for a Binomial Trait	28							
	0.0	6.6.1 Caution on Rare Haplotypes with Binomial Response	29							
	6.7	Control Parameters	29							
		6.7.1 Controlling Genetic Models: haplo.effect	30							
		6.7.2 Selecting the Baseline Haplotype	31							
		6.7.3 Rank of Information Matrix and eps.svd (NEW)	32							
		6.7.4 Rare Haplotypes and haplo.min.info	35							

7	Met	shods for haplo.glm (NEW)	36				
	7.1	fitted.values	36				
	7.2	residuals	36				
	7.3	vcov					
	7.4	anova and Model Comparison	37				
8	Exte	ended Applications	39				
	8.1	Combine Score and Group Results: haplo.score.merge	39				
	8.2	Case-Control Haplotype Analysis: haplo.cc	39				
	8.3	Score Tests on Sub-Haplotypes: haplo.score.slide	42				
		8.3.1 Plot Results from haplo.score.slide	42				
	8.4	Scanning Haplotypes Within a Fixed-Width Window: haplo.scan	44				
	8.5	Sequential Haplotype Scan Methods: seqhap	45				
		8.5.1 Plot Results from seqhap					
	8.6	Creating Haplotype Effect Columns: haplo.design					
9	Lice	ense and Warranty	51				
10	10 Acknowledgements						
$\mathbf{A}$	Counting Haplotype Pairs When Marker Phenotypes Have Missing Alleles 55						

# 1 Introduction

Haplo Stats is a suite of R routines for the analysis of indirectly measured haplotypes. The statistical methods assume that all subjects are unrelated and that haplotypes are ambiguous (due to unknown linkage phase of the genetic markers), while also allowing for missing alleles.

The user-level functions in Haplo Stats are:

- haplo.em: for the estimation of haplotype frequencies and posterior probabilities of haplotype pairs for each subject, conditional on the observed marker data
- haplo.glm: generalized linear models for the regression of a trait on haplotypes, with the option of including covariates and interactions
- haplo.score: score statistics to test associations between haplotypes and a variety of traits, including binary, ordinal, quantitative, and Poisson
- haplo.score.slide: haplo.score computed on sub-haplotypes of a larger region
- sequentially scan markers in enlarging a haplotype for association with a trait
- haplo.cc: run a combined analysis for haplotype frequencies, scores, and regression results for a case-control study
- haplo.power.qt/haplo.power.cc: power or sample size calculatins for quantitative or binary trait
- haplo.scan: search for a trait locus for all sizes of sub-haplotypes within a fixed maximum window width for all markers in a region
- haplo.design: create a design matrix for haplotype effects

This manual explains the basic and advanced usage of these routines, with guidelines for running the analyses and interpreting results. We provide many of these details in the function help pages, which are accessed within an R session using help(haplo.em), for example. We also provide brief examples in the help files, which can be run in the R session with example(haplo.em).

#### 1.1 Updates

The last major update to Haplo Stats included updates to haplo.glm in section 6 and new methods written for it that resemble glm class methods. These methods include residuals, fitted.values, vcov, and anova, and they are detailed in section 7. For full history of updates see the NEWS file, or type news(package="haplo.stats") in the R command prompt.

#### 1.2 Operating System and Installation

Haplo Stats version 1.6.0 is written for R (version 3.0.1). It has been uploaded to the Comprehensive R Archive Network (CRAN), and is made available on various operating systems through CRAN. Package installation within R is made simple from within R using *install.packages("haplo.stats")*, but other procedures for installing R packages can be found at the R project website (http://www.r-project.org).

#### 1.3 R Basics

For users who are new to the R environment, we demonstrate some basic concepts. In the following example we create a vector of character alleles and use the *table* function to get allele counts. We first show how to save the results of *table*(snp) into an R session variable, *tab*. We show that *tab* is an object of the *table* class, and use the print and summary methods that are defined for *table* objects. Note that when you enter just *tab* or *table*(snp) at the prompt, the print method is invoked.

```
R > snp < -c("A", "T", "T", "A", "A", "T", "T")
R> tab <- table(snp)</pre>
R> tab
snp
A T
3 4
R> class(tab)
[1] "table"
R> print.table(tab)
snp
A T
3 4
R> summary(tab)
Number of cases in table: 7
Number of factors: 1
R>
```

The routines in *haplo.stats* are computationally intensive and return lots of information in the returned object. Therefore, we assign classes to the returned objects and provide various methods for each of them.

# 2 Data Setup

We first show some typical steps when you first load a package and look for details on a function of interest. In the sample code below, we load *haplo.stats*, check which functions are available in the package, view a help file, and run the example that is within the help file.

```
R> # load the library, load and preview at demo dataset
R> library(haplo.stats)
R> ls(name="package:haplo.stats")
R> help(haplo.em)
R> example(haplo.em)
```

#### 2.1 Example Data

The haplo.stats package contains three example data sets. The primary data set used in this manual is named (hla.demo), which contains 11 loci from the HLA region on chromosome 6, with covariates, qualitative, and quantitative responses. Within /haplo.stats/data/hla.demo.tab the data is stored in tab-delimited format. Typically data stored in this format can be read in using read.table(). Since the data is provided in the package, we load the data in R using data() and view the names of the columns. Then to make the columns of hla.demo accessible without typing it each time, we attach it to the current session.

```
R> # load and preview demo dataset stored in ~/haplo.stats/data/hla.demo.tab
R> data(hla.demo)
R> names(hla.demo)
```

```
[1] "resp"
                                                       "DPB.a1"
                                                                   "DPB.a2"
                                                                                "DPA.a1"
                  "resp.cat"
                             "male"
                                          "age"
     "DPA.a2"
                                                       "DMB.a2"
                                                                   "TAP1.a1"
                                                                                "TAP1.a2"
[8]
                 "DMA.a1"
                              "DMA.a2"
                                          "DMB.a1"
[15] "TAP2.a1"
                 "TAP2.a2"
                              "DQB.a1"
                                          "DQB.a2"
                                                       "DQA.a1"
                                                                   "DQA.a2"
                                                                                "DRB.a1"
[22] "DRB.a2"
                 "B.a1"
                              "B.a2"
                                          "A.a1"
                                                       "A.a2"
```

R> # attach hla.demo to make columns available in the session R> attach(hla.demo)

The column names of *hla.demo* are shown above. They are defined as follows:

- resp: quantitative antibody response to measles vaccination
- resp.cat: a factor with levels "low", "normal", "high", for categorical antibody response
- male: gender code with 1="male", 0="female"
- age: age (in months) at immunization

The remaining columns are genotypes for 11 HLA loci, with a prefix name (e.g., "DQB") and a suffix for each of two alleles (".a1" and ".a2"). The variables in *hla.demo* can be accessed by typing *hla.demo*\$ before their names, such as *hla.demo*\$resp. Alternatively, it is easier for these examples to attach *hla.demo*, (as shown above with attach()) so the variables can be accessed by simply typing their names.

# 2.2 Creating a Genotype Matrix

Many of the functions require a matrix of genotypes, denoted below as geno. This matrix is arranged such that each locus has a pair of adjacent columns of alleles, and the order of columns corresponds to the order of loci on a chromosome. If there are K loci, then the number of columns of geno is 2K. Rows represent the alleles for each subject. For example, if there are three loci, in the order A-B-C, then the 6 columns of geno would be arranged as A.a1, A.a2, B.a1, B.a2, C.a1, C.a2. For illustration, three of the loci in hla.demo will be used to demonstrate some of the functions. Create a separate data frame for 3 of the loci, and call this geno. Then create a vector of labels for the loci.

```
R> geno <- hla.demo[,c(17,18,21:24)]
R> label <-c("DQB","DRB","B")
```

The *hla.demo* data already had alleles in two columns for each locus. For many SNP datasets, the data is in a one column format, giving the count of the minor allele. To assist in converting this format to two columns, a function named *geno1to2* has been added to the package. See its help file for more details.

#### 2.3 Preview Missing Data: summaryGeno

Before performing a haplotype analysis, the user will want to assess missing genotype data to determine the completeness of the data. If many genotypes are missing, the functions may take a long time to compute results, or even run out of memory. For these reasons, the user may want to remove some of the subjects with a lot of missing data. This step can be guided by using the *summaryGeno* function, which checks for missing allele information and counts the number of potential haplotype pairs that are consistent with the observed data (see the Appendix for a description of this counting scheme).

The codes for missing alleles are defined by the parameter *miss.val*, a vector to define all possible missing value codes. Below, the result is saved in *geno.desc*, which is a data frame, so individual rows may be printed. Here we show the results for subjects 1-10, 80-85, and 135-140, some of which have missing alleles.

```
R> geno.desc <- summaryGeno(geno, miss.val=c(0,NA))
R> print(geno.desc[c(1:10,80:85,135:140),])
```

missing0	missing1	missing2	N.enum.rows
3	0	0	4
3	0	0	4
3	0	0	4
3	0	0	2
3	0	0	4
3	0	0	2
3	0	0	4
3	0	0	2
3	0	0	2
3	0	0	1
3	0	0	4
2	0	1	1800
3	0	0	2
3	0	0	1
3	0	0	2
3	0	0	4
3	0	0	4
3	0	0	2
1	0	2	129600
3	0	0	4
3	0	0	4
3	0	0	4
	3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	3 0 3 0 3 0 3 0 3 0 3 0 3 0 3 0 3 0 3 0	3       0       0         3       0       0

The columns with 'loc miss-' illustrate the number of loci missing either 0, 1, or 2 alleles, and the last column, num\_enum\_rows, illustrates the number of haplotype pairs that are consistent with the observed data. In the example above, subjects indexed by rows 81 and 137 have missing alleles. Subject #81 has one locus missing two alleles, while subject #137 has two loci missing two alleles. As indicated by num\_enum\_rows, subject #81 has 1,800 potential haplotype pairs, while subject #137 has nearly 130,000.

The 130,000 haplotype pairs is considered a large number, but haplo.em, haplo.score, and haplo.glm complete in roughly 3-6 minutes (depending on system limits or control parameter settings). If person #137 were removed, the methods would take less than half that time. It is preferred to keep people if they provide information to the analysis, given that run time and memory usage are not too much of a burden.

When a person has no genotype information, they do not provide information to any of the methods in haplo.stats. Furthermore, they cause a much longer run time. Below, using the table function on the third column of geno.desc, we can tabulate how many people are missing two alleles at any at of the three loci. If there were people missing two alleles at all three loci, they should be removed. The second command below shows how to make an index of which people to remove from hla.demo because they are missing all their alleles.

```
R> # find if there are any people missing all alleles
R> table(geno.desc[,3])

0  1  2
218  1  1

R> ## create an index of people missing all alleles
R> miss.all <- which(geno.desc[,3]==3)
R> # use index to subset hla.demo
R> hla.demo.updated <- hla.demo[-miss.all,]</pre>
```

# 2.4 Random Numbers and Setting Seed

Random numbers are used in several of the functions (e.g., to determine random starting frequencies within haplo.em, and to compute permutation p-values in haplo.score). To reproduce calculations involving random numbers, we use set.seed() before any function that uses random numbers. Section 6.7.2 shows one example of setting the seed for haplo.glm. We illustrate setting the seed below.

```
R> # this is how to set the seed for reproducing results where haplo.em is R> # involved, and also if simulations are run. In practice, don't reset seed. R> seed <- c(17, 53, 1, 40, 37, 0, 62, 56, 5, 52, 12, 1)
R> set.seed(seed)
```

# 3 Haplotype Frequency Estimation: haplo.em

# 3.1 Algorithm

For genetic markers measured on unrelated subjects, with linkage phase unknown, haplo.em computes maximum likelihood estimates of haplotype probabilities. Because there may be more than one pair of haplotypes that are consistent with the observed marker phenotypes, posterior probabilities of haplotype pairs for each subject are also computed. Unlike the usual EM which attempts to enumerate all possible haplotype pairs before iterating over the EM steps, our progressive insertion algorithm progressively inserts batches of loci into haplotypes of growing lengths, runs the EM steps, trims off pairs of haplotypes per subject when the posterior probability of the pair is below a specified threshold, and then continues these insertion, EM, and trimming steps until all loci are inserted into the haplotype. The user can choose the batch size. If the batch size is chosen to be all loci, and the threshold for trimming is set to 0, then this reduces to the usual EM algorithm. The basis of this progressive insertion algorithm is from the "snphap" software by David Clayton[1]. Although some of the features and control parameters of haplo.em are modeled after snphap, there are substantial differences, such as extension to allow for more than two alleles per locus, and some other nuances on how the algorithm is implemented.

#### 3.2 Example Usage

We use haplo.em on geno for the 3 loci defined above and save the result in an object named save.em, which has the haplo.em class. The print method would normally print all 178 haplotypes from save.em, but to keep the results short for this manual, we give a quick glance of the output by using the option nlines=10, which prints only the first 10 haplotypes of the full results. The nlines parameter has been employed in some of Haplo Stats' print methods for when there are many haplotypes. In practice, it is best to exclude this parameter so that the default will print all results.

```
R> save.em <- haplo.em(geno=geno, locus.label=label, miss.val=c(0,NA))
R> names(save.em)
```

```
[1] "lnlike"
                                       "1r"
                      "lnlike.noLD"
                                                         "df.lr"
                                                                          "hap.prob"
[6] "hap.prob.noLD" "converge"
                                       "locus.label"
                                                         "indx.subj"
                                                                          "subj.id"
[11] "post"
                                       "hap2code"
                                                         "haplotype"
                                                                          "nreps"
                      "hap1code"
[16] "rows.rem"
                      "max.pairs"
                                       "control"
```

R> print(save.em, nlines=10)

\_\_\_\_\_

```
Haplotypes
```

-----

DQB DRB B hap.freq

```
1
    21
                0.00232
2
    21
          2
            7
                0.00227
3
    21
         2 18
                0.00227
4
    21
         3 8
                0.10408
5
         3 18
                0.00229
6
    21
          3 35
                0.00570
7
                0.00378
    21
          3 44
8
    21
          3 45
                0.00227
9
    21
          3 49
                0.00227
10
          3 57
                0.00227
```

\_\_\_\_\_\_

#### Details

\_\_\_\_\_\_

```
lnlike = -1847.675
lr stat for no LD = 632.5085 , df = 125 , p-val = 0
```

# **Explanation of Results**

The print methods shows the haplotypes and their estimated frequencies, followed by the final log-likelihood statistic and the lr stat for no LD, which is the likelihood ratio test statistic contrasting the lnlike for the estimated haplotype frequencies versus the lnlike under the null assuming that alleles from all loci are in linkage equilibrium. We note that the trimming by the progressive insertion algorithm can invalidate the lr stat and the degrees of freedom (df).

#### 3.3 Summary Method

The summary method for a haplo.em object on save.em shows the list of haplotypes per subject, and their posterior probabilities:

R> summary(save.em, nlines=7)

		1	1	1
1	1	58	78	1.00000
2	2	13	143	0.12532
3	2	17	138	0.87468
4	3	25	168	1.00000
5	4	13	39	0.28621
6	4	17	38	0.71379
7	5	55	94	1.00000

\_\_\_\_\_\_

Number of haplotype pairs: max vs used

\_\_\_\_\_\_

```
2
           1
                     3
                        84 135
         18
                0
                     0
                          0
                              0
1
2
         50
                4
                     0
                          0
                              0
4
        116
              29
                          0
                              0
                     1
                0
1800
           0
                     0
                          1
                              0
129600
           0
                0
                          0
                              1
```

# **Explanation of Results**

The first part of the summary output lists the subject id (row number of input geno matrix), the codes for the haplotypes of each pair, and the posterior probabilities of the haplotype pairs. The second part gives a table of the maximum number of pairs of haplotypes per subject, versus the number of pairs used in the final posterior probabilities. The haplotype codes remove the clutter of illustrating all the alleles of the haplotypes, but may not be as informative as the actual haplotypes themselves. To see the actual haplotypes, use the show.haplo=TRUE option, as in the following example.

R> # show full haplotypes, instead of codes
R> summary(save.em, show.haplo=TRUE, nlines=7)

\_\_\_\_\_

Subjects: Haplotype Codes and Posterior Probabilities								
	subj.id	hap1.DQB	hap1.DRB	hap1.B	hap2.DQB	hap2.DRB	hap2.B	posterior
58	1	31	11	61	32	4	62	1.00000
13	2	21	7	7	62	2	44	0.12532
17	2	21	7	44	62	2	7	0.87468
25	3	31	1	27	63	13	62	1.00000
13.1	4	21	7	7	31	7	44	0.28621
17.1	4	21	7	44	31	7	7	0.71379
55	5	31	11	51	42	8	55	1.00000

Number of haplotype pairs: max vs used

\_\_\_\_\_\_

X	1	2	3	84	135
1	18	0	0	0	0
2	50	4	0	0	0
4	116	29	1	0	0
1800	0	0	0	1	0
129600	0	0	0	0	1

#### 3.4 Control Parameters for haplo.em

A set of control parameters can be passed together to haplo.em as the "control" argument. This is a list of parameters that control the EM algorithm based on progressive insertion of loci. The default values are set by a function called haplo.em.control (see help(haplo.em.control) for a complete description). Although the user can accept the default values, there are times when control parameters may need to be adjusted.

These parameters are defined as:

- insert.batch.size: Number of loci to be inserted in a single batch.
- min.posterior: Minimum posterior probability of haplotype pair, conditional on observed marker genotypes. Posteriors below this minimum value will have their pair of haplotypes "trimmed" off the list of possible pairs.
- max.iter: Maximum number of iterations allowed for the EM algorithm before it stops and prints an error.
- n.try: Number of times to try to maximize the *lnlike* by the EM algorithm. The first try will use, as initial starting values for the posteriors, either equal values or uniform random variables, as determined by *random.start*. All subsequent tries will use uniform random values as initial starting values for the posterior probabilities.

• max.haps.limit: Maximum number of haplotypes for the input genotypes. Within haplo.em, the first step is to try to allocate the sum of the result of geno.count.pairs(), if that exceeds max.haps.limit, start by allocating max.haps.limit. If that is exceeded in the progressive-insertions steps, the C function doubles the memory until it can longer request more.

One reason to adjust control parameters is for finding the global maximum of the log-likelihood. It can be difficult in particular for small sample sizes and many possible haplotypes. Different maximizations of the log-likelihood may result in different results from haplo.em, haplo.score, or haplo.glm when rerunning the analyses. The algorithm uses multiple attempts to maximize the log-likelihood, starting each attempt with random starting values. To increase the chance of finding the global maximum of the log-likelihood, the user can increase the number of attempts (n.try), increase the batch size (insert.batch.size), or decrease the trimming threshold for posterior probabilities (min.posterior).

Another reason to adjust control parameters is when the algorithm runs out of memory because there are too many haplotypes. If max.haps.limit is exceeded when a batch of markers is added, the algorithm requests twice as much memory until it runs out. One option is to set max.haps.limit to a different value, either to make haplo.em request more memory initially, or to request more memory in smaller chunks. Another solution is to make the algorithm trim the number of haplotypes more aggressively by decreasing insert.batch.size or increasing min.posterior. Any changes to these parameters should be made with caution, and not drastically different from the default values. For instance, the default for min.posterior used to be 1e-7, and in some rare circumstances with many markers in only moderate linkage disequilibrium, some subjects had all their possible haplotype pairs trimmed. The default is now set at 1e-9, and we recommend not increasing min.posterior much greater than 1e-7.

The example below gives the command for increasing the number of tries to 20, and the batch size to 2, since not much more can be done for three markers.

# 3.5 Haplotype Frequencies by Group Subsets

To compute the haplotype frequencies for each level of a grouping variable, use the function haplo.group. The following example illustrates the use of a binomial response based on resp.cat, y.bin, that splits the subjects into two groups.

```
R> ## run haplo.em on sub-groups
R> ## create ordinal and binary variables
R> y.bin <- 1*(resp.cat=="low")
R> group.bin <- haplo.group(y.bin, geno, locus.label=label, miss.val=0)
R> print(group.bin, nlines=15)

Counts per Grouping Variable Value

group
0 1
157 63

Haplotype Frequencies By Group
```

```
21
             8 0.00232 0.00335
1
                                      NA
2
    21
        10
             8 0.00181 0.00318
                                      NA
             8 0.00274
3
    21
                                      NA
4
    21
          2 18 0.00227 0.00318
                                      NA
5
    21
             7 0.00227 0.00318
                                      NA
6
          3 18 0.00229 0.00637
    21
                                      NA
7
    21
          3 35 0.00570 0.00639
                                      NA
8
    21
          3 44 0.00378 0.00333 0.01587
9
    21
          3 45 0.00227
                             NA
                                      NA
10
    21
          3 49 0.00227
                             NA
                                      NA
11
    21
          3 57 0.00227
                             NA
                                      NA
    21
12
          3 70 0.00227
                             NA 0.00000
13
    21
             8 0.10408 0.06974 0.19048
          4 62 0.00455 0.00637
14
    21
15
    21
          7 13 0.01072
                             NA 0.02381
```

### Explanation of Results

The group.bin object can be very large, depending on the number of possible haplotypes, so only a portion of the output is illustrated above (limited again by *nlines*). The first section gives a short summary of how many subjects appear in each of the groups. The second section is a table with the following columns:

- The first column gives row numbers.
- The next columns (3 in this example) illustrate the alleles of the haplotypes.
- Total are the estimated haplotype frequencies for the entire data set.
- The last columns are the estimated haplotype frequencies for the subjects in the levels of the group variable (y.bin=0 and y.bin=1 in this example). Note that some haplotype frequencies have an NA, which appears when the haplotypes do not occur in the subgroups.

# 4 Power and Sample Size for Haplotype Association Studies

It is known that using haplotypes has greater power than single-markers to detect genetic association in some circumstances. There is little guidance, however, in determining sample size and power under different circumstances, some of which include: marker type, dominance, and effect size. The *haplo.stats* package now includes functions to calculate sample size and power for haplotype association studies, which is flexible to handle these multiple circumstances.

Based on work in Schaid 2005[2], we can take a set of haplotypes with their population frequencies, assign a risk to a subset of the haplotypes, then determine either the sample size to achieve a stated power, or the power for a stated sample size. Sample size and power can be calculated for either quantitative traits or case-control studies.

#### **4.1 Quantitative Traits:** haplo.power.qt

We assume that quantitative traits will be modeled by a linear regression. Some well-known tests for association between haplotypes and the trait include score statistics[3] and an F-test[4]. For both types of tests, power depends on the amount of variance in the trait that is explained by haplotypes, or a multiple correlation coefficient,  $R^2$ . Rather than specifying the haplotype coefficients directly, we calculate the vector of coefficients based on an  $R^2$  value.

In the example below, we load an example set of haplotypes that contain 5 markers, and specify the indices of the at-risk haplotypes; in this case, whichever haplotype has allele 1 at the 2nd and 3rd markers.

We set the first haplotype (most common) as the baseline. With these values we calculate the vector of coefficients for haplotype effects from find.haplo.beta.qt using an  $R^2 = 0.01$ . Next, we use haplo.power.qt to calculate the sample size for the set of haplotypes and their coefficients, type-I error (alpha) set to 0.05, power at 80%, and the same mean and variance used to get haplotype coefficients. Then we use the sample size needed for 80% power for un-phased haplotypes (2,826) to get the power for both phased and un-phased haplotypes.

```
R> # load a set of haplotypes (hap-1 from Schaid 2005)
R>
R> data(hapPower.demo)
R> #### an example using save.em hla markers may go like this.
R> # keep <- which(save.em$hap.prob > .004) # get an index of non-rare haps
R> # hfreq <- save.em$hap.prob[keep]</pre>
R> # hmat <- save.em$haplotype[keep,]</pre>
R> # hrisk <- which(hmat[,1]==31 & hmat[,2]==11) # contains 3 haps with freq=.01
R> # hbase <- 4 # 4th hap has mas freq of .103
R> ####
R>
R> ## separate the haplotype matrix and the frequencies
R> hmat <- hapPower.demo[,-6]</pre>
R> hfreq <- hapPower.demo[,6]</pre>
R> ## Define risk haplotypes as those with "1" allele at loc2 and loc3
R> hrisk <- which(hmat$loc.2==1 & hmat$loc.3==1)</pre>
R> # define index for baseline haplotype
R> hbase <- 1
R> hbeta.list <- find.haplo.beta.qt(haplo=hmat, haplo.freq=hfreq, base.index=hbase,
                             haplo.risk=hrisk, r2=.01, y.mu=0, y.var=1)
R> hbeta.list
$r2
[1] 0.01
$beta
$base.index
[1] 1
$haplo.risk
[1] 8 10 16 17
R> ss.qt <- haplo.power.qt(hmat, hfreq, hbase, hbeta.list$beta,
+
                     y.mu=0, y.var=1, alpha=.05, power=.80)
R> ss.qt
$ss.phased.haplo
[1] 2091
$ss.unphased.haplo
[1] 2826
```

# 4.2 Case-Control Studies: haplo.power.cc

The steps to compute sample size and power for case-control studies is similar to the steps for quantitative traits. If we assume a log-additive model for haplotype effects, the haplotype coefficients can be specified first as odds ratios (OR), and then converted to logistic regression coefficients according to log(OR).

In the example below, we assume the same baseline and risk haplotypes defined in section 4.1, give the risk haplotypes an odds ratio of 1.50, and specify a population disease prevalence of 10%. We also assume cases make up 50% (case.frac) of the studyś subjects. We first compute the sample size for this scenario for Type-I error (alpha) at 0.05 and 80% power, and then compute power for the sample size required for un-phased haplotypes (4,566).

```
$power.phased.haplo
[1] 0.8
$power.unphased.haplo
[1] 0.8
R> # Compute power for given sample size
R> power.cc <- haplo.power.cc(hmat, hfreq, hbase, hbeta.cc, case.frac=.5, prevalence=.1,
                 alpha=.05, sample.size=4566)
R> power.cc
$ss.phased.haplo
[1] 4566
$ss.unphased.haplo
「1] 4566
$power.phased.haplo
[1] 0.9206568
$power.unphased.haplo
[1] 0.8000695
```

# 5 Haplotype Score Tests: haplo.score

The haplo.score routine is used to compute score statistics to test association between haplotypes and a wide variety of traits, including binary, ordinal, quantitative, and Poisson. This function provides several different global and haplotype-specific tests for association and allows for adjustment for non-genetic covariates. Haplotype effects can be specified as additive, dominant, or recessive. This method also has an option to compute permutation p-values, which may be needed for sparse data when distribution assumptions may not be met. Details on the background and theory of the score statistics can be found in Schaid et al.[3].

#### 5.1 Quantitative Trait Analysis

First, we assess a haplotype association with a quantitative trait in hla.demo called resp. To tell haplo.score the trait is quantitative, specify the parameter trait.type="gaussian" (a reminder that a gaussian distribution is assumed for the error terms). The other arguments, all set to default values, are explained in the help file. Note that rare haplotypes can result in unstable variance estimates, and hence unreliable test statistics for rare haplotypes. We restrict the analysis to get scores for haplotypes with a minimum sample count using min.count=5. For more explanation on handling rare haplotypes, see section 5.6. Below is an example of running haplo.score with a quantitative trait, then viewing the results using the print method for the haplo.score class. (again, output shortened by nlines).

\_\_\_\_\_

Haplotype Effect Model: additive

Global Score Statistics

global-stat = 30.6353, df = 18, p-val = 0.03171

#### Haplotype-specific Scores

DQB DRB B Hap-Freq Hap-Score p-val [1,] 21 3 8 0.10408 -2.39631 0.01656 [2,] 31 44 0.02849 -2.24273 0.02491 [3,] 51 44 0.01731 -0.99357 0.32043 1 [4,] 63 13 44 0.01606 -0.84453 0.39837 [5,] 63 2 7 0.01333 -0.50736 0.6119 [6,] 32 4 60 0.0306 -0.46606 0.64118 [7,] 21 44 0.02332 -0.41942 0.67491 7 [8,] 62 2 44 0.01367 -0.26221 0.79316 [9,] 62 2 18 0.01545 -0.21493 0.82982 [10,] 51 1 27 0.01505 0.01539 0.98772

# **Explanation of Results**

First, the model effect chosen by haplo.effect is printed across the top. The section Global Score Statistics shows results for testing an overall association between haplotypes and the response. The global-stat has an asymptotic  $\chi^2$  distribution, with degrees of freedom (df) and p-value as indicated. Next, Haplotype-specific scores are given in a table format. The column descriptions are as follows:

- The first column gives row numbers.
- The next columns (3 in this example) illustrate the alleles of the haplotypes.
- Hap-Freq is the estimated frequency of the haplotype in the pool of all subjects.
- Hap-Score is the score for the haplotype, the results are sorted by this value. Note, the score statistic should not be interpreted as a measure of the haplotype effect.
- p-val is the asymptotic  $\chi_1^2$  p-value, calculated from the square of the score statistic.

#### 5.2 Binary Trait Analysis

Let us assume that "low" responders are of primary interest, so we create a binary trait that has values of 1 when resp.cat is "low", and 0 otherwise. Then in haplo.score specify the parameter trait.type="binomial".

```
Haplotype Effect Model: additive
                             Global Score Statistics
global-stat = 33.70125, df = 18, p-val = 0.01371
                           Haplotype-specific Scores
     DQB DRB B Hap-Freq Hap-Score p-val
[1,] 62 2
           7 0.05098 -2.19387 0.02824
 [2,] 51 1
            35 0.03018 -1.58421 0.11315
 [3,] 63 13 7 0.01655 -1.56008 0.11874
 [4,] 21 7 7 0.01246 -1.47495 0.14023
 [5,] 32 4 7 0.01678 -1.00091 0.31687
 [6,] 32 4 62 0.02349 -0.6799
                               0.49657
 [7,] 51 1
            27 0.01505 -0.66509 0.50599
 [8,] 31 11 35 0.01754 -0.5838 0.55936
 [9,] 31 11 51 0.01137 -0.43721 0.66196
[10,] 51 1
           44 0.01731 0.00826 0.99341
5.3
     Ordinal Trait Analysis
To create an ordinal trait, here we convert resp.cat (described above) to numeric values, y.ord (with levels
1, 2, 3). For haplo.score, use y.ord as the response variable, and set the parameter trait.type = "ordinal".
R> # scores w/ ordinal trait
R> y.ord <- as.numeric(resp.cat)</pre>
R> score.ord <- haplo.score(y.ord, geno, trait.type="ordinal",
                    x.adj = NA, min.count=5,
                    locus.label=label,
                    miss.val=0, simulate=FALSE)
R> print(score.ord, nlines=7)
                         Haplotype Effect Model: additive
                             Global Score Statistics
global-stat = 15.23209, df = 18, p-val = 0.64597
______
                           Haplotype-specific Scores
------
```

DQB DRB B Hap-Freq Hap-Score p-val

```
[1,] 32
            62 0.02349 -2.17133 0.02991
[2,] 21
        3
            8 0.10408 -1.34661 0.17811
[3,] 32
            7 0.01678
                       -1.09487
                                 0.27357
[4,] 62
        2
              0.05098
                       -0.96874
                                 0.33268
            7
[5,] 21
        7
            44 0.02332
                        -0.83747
                                 0.40233
[6,] 63
           7
              0.01655
        13
                       -0.80787 0.41917
            7 0.01246 -0.63316 0.52663
[7,] 21
```

#### Warning for Ordinal Traits

When analyzing an ordinal trait with adjustment for covariates (using the x.adj option), the software requires the rms package, distributed by Frank Harrell [5]. If the user does not have these packages installed, then it will not be possible to use the x.adj option. However, the unadjusted scores for an ordinal trait (using the default option x.adj=NA) do not require these packageses. Check the list of your local packages in the list shown from entering library() in your prompt.

#### 5.4 Haplotype Scores, Adjusted for Covariates

To adjust for covariates in *haplo.score*, first set up a matrix of covariates from the example data. For example, use a column for male (1 if male; 0 if female), and a second column for age. Then pass the matrix to *haplo.score* using parameter x.adj. The results change slightly in this example.

```
DQB DRB B Hap-Freq Hap-Score p-val
[1,] 21
         3
             8 0.10408 -2.4097
[2,] 31
             44 0.02849 -2.25293 0.02426
         4
[3,] 51
             44 0.01731
                         -0.98763
         1
                                   0.32333
[4,] 63
         13
            44 0.01606 -0.83952 0.40118
[5,] 63
         2
             7 0.01333
                         -0.48483
                                   0.6278
[6,] 32
         4
             60 0.0306
                          -0.46476
                                   0.64211
[7,] 21
         7
             44 0.02332
                         -0.41249
                                   0.67998
[8,] 62
         2
             44 0.01367
                         -0.26443
                                   0.79145
[9,] 62
         2
             18 0.01545
                         -0.20425
                                   0.83816
[10,] 51
             27 0.01505 0.02243
                                   0.9821
```

# 5.5 Plots and Haplotype Labels

A convenient way to view results from haplo.score is a plot of the haplotype frequencies (Hap-Freq) versus the haplotype score statistics (Hap-Score). This plot, and the syntax for creating it, are shown in Figure 1.

Some points on the plot may be of interest. To identify individual points on the plot, use locator.haplo(score.gaus), which is similar to locator(). Use the mouse to select points on the plot. After points are chosen, click on the middle mouse button, and the points are labeled with their haplotype labels. Note, in constructing Figure 1, we had to define which points to label, and then assign labels in the same way as done within the locator.haplo function.

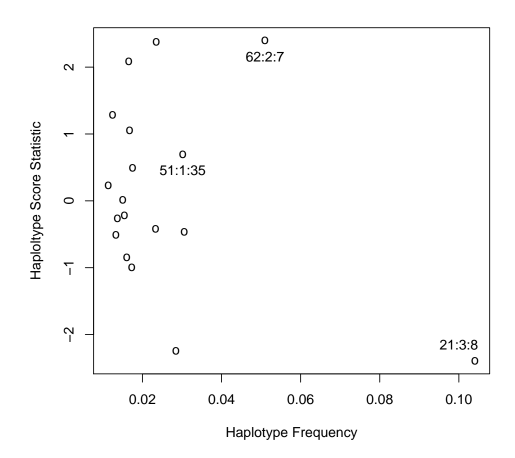


Figure 1: Haplotype Statistics: Score vs. Frequency, Quantitative Response

### 5.6 Skipping Rare Haplotypes

For the haplo.score, the skip.haplo and min.count parameters control which rare haplotypes are pooled into a common group. The min.count parameter is a recent addition to haplo.score, yet it does the same task as skip.haplo and is the same idea as haplo.min.count used in haplo.glm.control for haplo.glm. As a guideline, you may wish to set min.count to calculate scores for haplotypes with expected haplotype counts of 5 or greater in the sample. We concentrate on this expected count because it adjusts to the size of the input data. If N is the number of subjects and f the haplotype frequency, then the expected haplotype count is  $count = 2 \times N \times f$ . Alternatively, you can choose  $skip.haplo = \frac{count}{2 \times N}$ .

In the following example we try a different cut-off than before, min.count=10, which corresponds to skip.haplo of  $10 \div (2 \times 220) = .045$ . In the output, see that the global statistic, degrees of freedom, and p-value change because of the fewer haplotypes, while the haplotype-specific scores do not change.

```
R> # increase skip.haplo, expected hap counts = 10
R> score.gaus.min10 <- haplo.score(resp, geno, trait.type="gaussian",
                            x.adj = NA, min.count=10,
                            locus.label=label, simulate=FALSE)
R> print(score.gaus.min10)
                             Haplotype Effect Model: additive
                                 Global Score Statistics
global-stat = 20.66451, df = 7, p-val = 0.0043
                               Haplotype-specific Scores
     DQB DRB B Hap-Freq Hap-Score p-val
[1,] 21
            8 0.10408 -2.39631 0.01656
[2,] 31
            44 0.02849 -2.24273 0.02491
[3,] 32
            60 0.0306
                         -0.46606 0.64118
[4,] 21
        7
            44 0.02332 -0.41942 0.67491
[5,] 51
            35 0.03018 0.69696
                                   0.48583
[6,] 32
        4
            62 0.02349
                        2.37619
                                   0.01749
[7,] 62
            7 0.05098 2.39795
                                  0.01649
```

#### 5.7 Score Statistic Dependencies: the eps.svd parameter

The global score test is calculated using the vector of scores and the generalized inverse of their variance-covariance matrix, performed by the *Ginv* function. This function determines the rank of the variance matrix by its singular value decomposition, and an epsilon value is used as the cut-off for small singular values. If all of the haplotypes in the sample are scored, then there is dependence between them and the variance matrix is not of full rank. However, it is more often the case that one or more rare haplotypes are not scored because of low frequency. It is not clear how strong the dependencies are between the remaining score statistics, and likewise, there is disparity in calculating the rank of the variance matrix. For these instances we give the user control over the epsilon parameter for *haplo.score* with *eps.svd*.

We have seen instances where the global score test had a very significant p-value, but none of the haplotype-specific scores showed strong association. In such instances, we found the default epsilon value

in Ginv was incorrectly considering the variance matrix as having full rank, and the misleading global score test was corrected with a larger epsilon for Ginv.

# 5.8 Haplotype Model Effect

haplo.score allows non-additive effects for scoring haplotypes. The possible effects for haplotypes are additive, dominant, and recessive. Under recessive effects, fewer haplotypes may be scored, because subjects are required to be homozygous for haplotypes. Furthermore, there would have to be min.count such persons in the sample to have the recessive effect scored. Therefore, a recessive model should only be used on samples with common haplotypes. In the example below with the gaussian response, set the haplotype effect to dominant using parameter haplo.effect = "dominant". Notice the results change slightly compared to the score.gaus.add results above.

```
R> # score w/gaussian, dominant effect
R>
R> score.gaus.dom <- haplo.score(resp, geno, trait.type="gaussian",
                          x.adj=NA, min.count=5,
                          haplo.effect="dominant", locus.label=label,
                          simulate=FALSE)
R> print(score.gaus.dom, nlines=10)
                             Haplotype Effect Model: dominant
                                 Global Score Statistics
global-stat = 29.56133, df = 18, p-val = 0.04194
                               Haplotype-specific Scores
     DQB DRB B Hap-Freq Hap-Score p-val
 [1,] 21
         3
              8 0.10408 -2.23872 0.02517
 [2,] 31
              44 0.02849 -2.13233 0.03298
 [3,] 51
              44 0.01731 -0.99357
                                   0.32043
         1
 [4,] 63
         13
             44 0.01606
                         -0.84453
                                    0.39837
 [5,] 63
         2
             7 0.01333 -0.50736 0.6119
 [6,] 32
              60 0.0306
                          -0.46606 0.64118
 [7,] 21
         7
              44 0.02332 -0.41942
                                   0.67491
 [8,] 62
         2
              44 0.01367
                          -0.26221
                                    0.79316
 [9,] 62
         2
              18 0.01545
                         -0.21493
                                    0.82982
[10,] 51
              27 0.01505
                         0.01539
                                    0.98772
```

#### 5.9 Simulation p-values

When simulate=TRUE, haplo.score gives simulated p-values. Simulated haplotype score statistics are the re-calculated score statistics from a permuted re-ordering of the trait and covariates and the original ordering of the genotype matrix. The simulated p-value for the global score statistic (Global sim. p-val) is the number of times the simulated global score statistic exceeds the observed, divided by the total number of simulations.

Likewise, simulated p-value for the maximum score statistic (Max-stat sim. p-val) is the number of times the simulated maximum haplotype score statistic exceeds the observed maximum score statistic, divided by the total number of simulations. The maximum score statistic is the maximum of the square of the haplotype-specific score statistics, which has an unknown distribution, so its significance can only be given by the simulated p-value. Intuitively, if only one or two haplotypes are associated with the trait, the maximum score statistic should have greater power to detect association than the global statistic.

The score.sim.control function manages control parameters for simulations. The haplo.score function employs the simulation p-value precision criteria of Besag and Clifford[6]. These criteria ensure that the simulated p-values for both the global and the maximum score statistics are precise for small p-values. The algorithm performs a user-defined minimum number of permutations (min.sim) to guarantee sufficient precision for the simulated p-values for score statistics of individual haplotypes. Permutations beyond this minimum are then conducted until the sample standard errors for simulated p-values for both the global-stat and max-stat score statistics are less than a threshold (p.threshold \* p-value). The default value for p.threshold= $\frac{1}{4}$  provides a two-sided 95% confidence interval for the p-value with a width that is approximately as wide as the p-value itself. Effectively, simulations are more precise for smaller p-values. The following example illustrates computation of simulation p-values with min.sim=1000.

```
R> # simulations when binary response
R> score.bin.sim <- haplo.score(y.bin, geno, trait.type="binomial",
               x.adj = NA, locus.label=label, min.count=5,
               simulate=TRUE, sim.control = score.sim.control() )
R> print(score.bin.sim)
                            Haplotype Effect Model: additive
                                Global Score Statistics
global-stat = 33.70125, df = 18, p-val = 0.01371
                           Global Simulation p-value Results
Global sim. p-val = 0.0095
Max-Stat sim. p-val = 0.00563
Number of Simulations, Global: 2842, Max-Stat: 2842
                               Haplotype-specific Scores
     DQB DRB B Hap-Freq Hap-Score p-val
 [1,] 62
         2
             7
                0.05098 -2.19387 0.02824 0.02991
 [2,] 51
         1
             35 0.03018 -1.58421 0.11315 0.13863
 [3,] 63
         13
             7 0.01655 -1.56008 0.11874 0.19177
 [4,] 21
         7
             7 0.01246 -1.47495 0.14023 0.15588
 [5,] 32
         4
             7 0.01678 -1.00091 0.31687 0.25123
```

0.49657 0.47467

[6,] 32

[7,] 51 1

62 0.02349 -0.6799

27 0.01505 -0.66509 0.50599 0.63089

```
[8,] 31
              35 0.01754
                           -0.5838
                                      0.55936 0.6506
          11
 [9,] 31
                           -0.43721
          11
              51 0.01137
                                      0.66196 0.91872
[10,] 51
          1
              44 0.01731
                           0.00826
                                      0.99341 1
[11,] 32
              60 0.0306
                           0.03181
                                      0.97462 0.95074
          4
[12,] 62
          2
              44 0.01367
                           0.16582
                                      0.8683 0.91872
[13,] 63
                           0.22059
          13
              44 0.01606
                                      0.82541 0.7266
[14,] 63
                                      0.76555 0.89163
          2
                 0.01333
                           0.2982
[15,] 62
          2
              18 0.01545
                           0.78854
                                      0.43038 0.6608
[16,] 21
          7
              44 0.02332
                           0.84562
                                      0.39776 0.39796
[17,] 31
          4
              44 0.02849
                           2.50767
                                      0.01215 0.01161
[18,] 21
                 0.10408
                           3.77763
                                      0.00016 0.00035
```

# 6 Regression Models: haplo.glm

The haplo.glm function computes the regression of a trait on haplotypes, and possibly other covariates and their interactions with haplotypes. We currently support the gaussian, binomial, and Poisson families of traits with their canonical link functions. The effects of haplotypes on the link function can be modeled as either additive, dominant (heterozygotes and homozygotes for a particular haplotype assumed to have equivalent effects), or recessive (homozygotes of a particular haplotype considered to have an alternative effect on the trait). The basis of the algorithm is a two-step iteration process; the posterior probabilities of pairs of haplotypes per subject are used as weights to update the regression coefficients, and the regression coefficients are used to update the haplotype posterior probabilities. See Lake et al.[7] for details.

# 6.1 New and Updated Methods for haplo.glm

We initially wrote haplo.glm with a focus on creating a basic print method for results. We have now refined the haplo.glm class to look and act as much like a glm class object as possible with methods defined specifically for the haplo.glm class. We provide print and summary methods that make use of the corresponding methods for glm and then add extra information for the haplotypes and their frequencies. Furthermore, we have defined for the haplo.glm class some of the standard methods for regression fits, including residuals, fitted.values, vcov, and anova. We describe the challenges that haplotype regression presents with these methods in section 7.

#### **6.2** Preparing the data frame for haplo.glm

A critical distinction between haplo.glm and all other functions in Haplo Stats is that the definition of the regression model follows the S/R formula standard (see lm or glm). So, a data.frame must be defined, and this object must contain the trait and other optional covariates, plus a special kind of genotype matrix (geno.glm for this example) that contains the genotypes of the marker loci. We require the genotype matrix to be prepared using setupGeno(), which handles character, numeric, or factor alleles, and keeps the columns of the genotype matrix as a single unit when inserting into (and extracting from) a data.frame. The setupGeno function recodes all missing genotype value codes given by miss.val to NA, and also recodes alleles to integer values. The original allele codes are preserved within an attribute of geno.glm, and are utilized within haplo.glm. The returned object has class model.matrix, and it can be included in a data.frame to be used in haplo.glm. In the example below we prepare a genotype matrix, geno.glm, and create a data.frame object, glm.data, for use in haplo.glm.

```
R> # set up data for haplo.glm, include geno.glm,
R> # covariates age and male, and responses resp and y.bin
R> geno <- hla.demo[,c(17,18,21:24)]
R> geno.glm <- setupGeno(geno, miss.val=c(0,NA), locus.label=label)
R> attributes(geno.glm)
```

```
$dim
[1] 220
         6
$dimnames
$dimnames[[1]]
NULL
$dimnames[[2]]
[1] "DQB.a1" "DQB.a2" "DRB.a1" "DRB.a2" "B.a1"
                                                  "B.a2"
$class
[1] "model.matrix"
$unique.alleles
$unique.alleles[[1]]
 [1] "21" "31" "32" "33" "42" "51" "52" "53" "61" "62" "63" "64"
$unique.alleles[[2]]
 [1] "1" "2" "3" "4"
                        "7" "8" "9" "10" "11" "13" "14"
$unique.alleles[[3]]
 [1] "7" "8" "13" "14" "18" "27" "35" "37" "38" "39" "41" "42" "44" "45" "46" "47" "48"
[18] "49" "50" "51" "52" "55" "56" "57" "58" "60" "61" "62" "63" "70"
R> y.bin <- 1*(resp.cat=="low")</pre>
R> glm.data <- data.frame(geno.glm, age=age, male=male, y=resp, y.bin=y.bin)
```

#### 6.3 Rare Haplotypes

The issue of deciding which haplotypes to use for association is critical in *haplo.glm*. By default it will model a rare haplotype effect so that the effects of other haplotypes are in reference to the baseline effect of the one common happlotype. The rules for choosing haplotypes to be modeled in *haplo.glm* are similar to the rules in *haplo.score*: by a minimum frequency or a minimum expected count in the sample.

Two control parameters in haplo.glm.control may be used to control this setting: haplo.freq.min may be set to a selected minimum haplotype frequency, and haplo.min.count may be set to select the cut-off for minimum expected haplotype count in the sample. The default minimum frequency cut-off in haplo.glm is set to 0.01. More discussion on rare haplotypes takes place in section 6.7.4.

#### 6.4 Regression for a Quantitative Trait

The following illustrates how to fit a regression of a quantitative trait y on the haplotypes estimated from the geno.glm matrix, and the covariate male. For na.action, we use na.geno.keep, which keeps a subject with missing values in the genotype matrix if they are not missing all alleles, but removes subjects with missing values (NA) in either the response or covariate.

```
Call:
```

```
haplo.glm(formula = y ~ male + geno.glm, family = gaussian, data = glm.data,
    na.action = "na.geno.keep", locus.label = label, control = haplo.glm.control(haplo.freq.min = 0.02)
    x = TRUE)
```

#### Deviance Residuals:

```
Min 1Q Median 3Q Max
-2.46945 -0.92052 -0.06533 0.94874 2.37199
```

#### Coefficients:

	coef	se	t.stat	pval
(Intercept)	1.06436	0.34283	3.10464	0.002
male	0.09735	0.15521	0.62723	0.531
geno.glm.17	0.28022	0.43549	0.64346	0.521
geno.glm.34	-0.31713	0.34342	-0.92342	0.357
geno.glm.77	0.22167	0.36126	0.61360	0.540
geno.glm.78	1.14144	0.38382	2.97390	0.003
geno.glm.100	0.55557	0.36427	1.52517	0.129
geno.glm.138	0.98229	0.30329	3.23875	0.001
<pre>geno.glm.rare</pre>	0.39765	0.18191	2.18591	0.030

(Dispersion parameter for gaussian family taken to be 1.269581)

Null deviance: 297.01 on 219 degrees of freedom Residual deviance: 267.88 on 211 degrees of freedom

AIC: 687.65

Number of Fisher Scoring iterations: 268

#### Haplotypes:

```
DQB DRB B hap.freq
geno.glm.17
               21
                    7 44
                          0.02291
geno.glm.34
                          0.02858
               31
                    4 44
geno.glm.77
               32
                    4 60
                          0.03022
geno.glm.78
               32
                    4 62 0.02390
geno.glm.100
               51
                    1 35
                          0.03008
geno.glm.138
               62
                    2
                       7
                          0.05023
geno.glm.rare
                          0.71000
                       *
haplo.base
               21
                    3 8 0.10409
```

#### **Explanation of Results**

 is the default) is labeled as haplo.base; more information on the baseline may be found in section 6.7.2.

#### 6.5 Fitting Haplotype x Covariate Interactions

Interactions are fit by the standard S-language model syntax, using a '\*' in the model formula to indicate main effects and interactions. Some other formula constructs are not supported, so use the formula parameter with caution. Below is an example of modeling the interaction of *male* and the haplotypes. Because more terms will be estimated in this case, we limit how many haplotypes will be included by increasing *haplo.min.count* to 10.

```
R> # glm fit haplotypes with covariate interaction
R> fit.inter <- haplo.glm(formula = y ~ male * geno.glm,</pre>
                    family = gaussian, data=glm.data,
                    na.action="na.geno.keep",
+
                    locus.label = label,
                    control = haplo.glm.control(haplo.min.count = 10))
R> summary(fit.inter)
Call:
haplo.glm(formula = y ~ male * geno.glm, family = gaussian, data = glm.data,
   na.action = "na.geno.keep", locus.label = label, control = haplo.glm.control(haplo.min.count = 10))
Deviance Residuals:
                     Median
               1Q
                                            Max
-2.23387 -0.90661 -0.05953
                              0.96140
                                        2.48859
Coefficients:
                      coef
                                     t.stat pval
                                 se
(Intercept)
                   0.97536 0.52268 1.86607 0.063
male
                   0.25806 0.67351 0.38315 0.702
geno.glm.17
                   0.14443 0.54544 0.26479 0.791
geno.glm.34
                  -0.17161 0.66773 -0.25700 0.797
geno.glm.77
                   0.80523  0.64951  1.23975  0.216
geno.glm.78
                   0.49557 0.56574 0.87596 0.382
                   0.52310  0.48067  1.08828  0.278
geno.glm.100
geno.glm.138
                   1.15704 0.42325 2.73371 0.007
geno.glm.rare
                   0.45547 0.28721 1.58587 0.114
male:geno.glm.17
                   0.50872 0.87531 0.58119 0.562
male:geno.glm.34
                  male:geno.glm.77
                  -0.90084 0.79114 -1.13865 0.256
male:geno.glm.78
                   1.26376 0.77131 1.63846 0.103
male:geno.glm.100
                            0.77470 0.06549 0.948
                   0.05074
                            0.61903 - 0.72027 0.472
male:geno.glm.138 -0.44587
male:geno.glm.rare -0.09787 0.37197 -0.26312 0.793
(Dispersion parameter for gaussian family taken to be 1.27362)
   Null deviance: 297.01 on 219 degrees of freedom
Residual deviance: 259.82 on 204 degrees of freedom
AIC: 694.93
```

Number of Fisher Scoring iterations: 120

#### Haplotypes:

```
DQB DRB B hap.freq
geno.glm.17
              21
                   7 44 0.02346
geno.glm.34
              31
                   4 44 0.02845
geno.glm.77
              32
                   4 60 0.03060
geno.glm.78
              32
                   4 62 0.02413
geno.glm.100
              51
                   1 35 0.03013
geno.glm.138
              62
                   2 7 0.05049
geno.glm.rare
              *
                   * * 0.70863
              21
haplo.base
                   3 8 0.10410
```

#### **Explanation of Results**

The listed results are as explained under section 6.4. The main difference is that the interaction coefficients are labeled as a concatenation of the covariate (*male* in this example) and the name of the haplotype, as described above. In addition, estimates may differ because the model has changed.

#### 6.6 Regression for a Binomial Trait

Next we illustrate the fitting of a binomial trait with the same genotype matrix and covariate.

```
R> # gender and haplotypes fit on binary response,
R> # return model matrix
R> fit.bin <- haplo.glm(y.bin ~ male + geno.glm, family = binomial,
              data=glm.data, na.action = "na.geno.keep",
              locus.label=label,
               control = haplo.glm.control(haplo.min.count=10))
R> summary(fit.bin)
haplo.glm(formula = y.bin ~ male + geno.glm, family = binomial,
   data = glm.data, na.action = "na.geno.keep", locus.label = label,
    control = haplo.glm.control(haplo.min.count = 10))
Deviance Residuals:
   Min
             1Q
                 Median
                               30
                                       Max
-1.5559 -0.7996 -0.6473 1.0591
                                    2.4348
Coefficients:
                coef
                          se t.stat pval
(Intercept)
              1.5457 0.6547 2.3610 0.019
male
             -0.4802 0.3308 -1.4518 0.148
geno.glm.17
             -0.7227
                      0.8011 -0.9022 0.368
geno.glm.34
              0.3641 0.6798 0.5356 0.593
geno.glm.77
             -0.9884 0.7328 -1.3489 0.179
geno.glm.78
             -1.4093 0.8543 -1.6496 0.101
geno.glm.100 -2.5907 1.1278 -2.2971 0.023
geno.glm.138 -2.7156 0.8524 -3.1860 0.002
geno.glm.rare -1.2610 0.3537 -3.5647 0.000
```

(Dispersion parameter for binomial family taken to be 1)

```
Null deviance: 263.50 on 219 degrees of freedom Residual deviance: 233.46 on 211 degrees of freedom
```

AIC: 251.11

Number of Fisher Scoring iterations: 61

#### Haplotypes:

	DQB	DRB	В	hap.freq
geno.glm.17	21	7	44	0.02303
geno.glm.34	31	4	44	0.02843
geno.glm.77	32	4	60	0.03057
geno.glm.78	32	4	62	0.02354
geno.glm.100	51	1	35	0.02977
geno.glm.138	62	2	7	0.05181
<pre>geno.glm.rare</pre>	*	*	*	0.70880
haplo.base	21	3	8	0.10405

#### **Explanation of Results**

The underlying methods for haplo.glm are based on a prospective likelihood. Normally, this type of likelihood works well for case-control studies with standard covariates. For ambiguous haplotypes, however, one needs to be careful when interpreting the results from fitting haplo.glm to case-control data. Because cases are over-sampled, relative to the population prevalence (or incidence, for incident cases), haplotypes associated with disease will be over-represented in the case sample, and so estimates of haplotype frequencies will be biased. Positively associated haplotypes will have haplotype frequency estimates that are higher than the population haplotype frequency. To avoid this problem, one can weight each subject. The weights for the cases should be the population prevalence, and the weights for controls should be 1 (assuming the disease is rare in the population, and controls are representative of the general population). See Stram et al.[8] for background on using weights, and see the help file for haplo.glm for how to implement weights.

The estimated regression coefficients for case-control studies can be biased by either a large amount of haplotype ambiguity and mis-specified weights, or by departures from Hardy-Weinberg Equilibrium of the haplotypes in the pool of cases and controls. Generally, the bias is small, but tends to be towards the null of no association. See Stram et al. [8] and Epstein and Satten [9] for further details.

#### 6.6.1 Caution on Rare Haplotypes with Binomial Response

If a rare haplotype occurs only in cases or only in controls, the fitted values would go to 0 or 1, where R would issue a warning. Also, the coefficient estimate for that haplotype would go to positive or negative infinity, If the default haplo.min.count=5 were used above, this warning would appear. To keep this from occuring in other model fits, increase the minimum count or minimum frequency.

#### 6.7 Control Parameters

Additional parameters are handled using *control*, which is a list of parameters providing additional functionality in *haplo.glm*. This list is set up by the function *haplo.glm.control*. See the help file (*help(haplo.glm.control*)) for a full list of control parameters, with details of their usage. Some of the options are described here.

#### 6.7.1 Controlling Genetic Models: haplo.effect

The haplo effect control parameter for haplo glm instructs whether the haplotype effects are fit as additive, dominant, or recessive. That is, haplo effect determines whether the covariate (x) coding of haplotypes follows the values in Table 1 for each effect type. Heterozygous means a subject has one copy of a particular haplotype, and homozygous means a subject has two copies of a particular haplotype.

Table 1: Coding haplotype covariates in a model matrix

Hap - Pair	additive	dominant	recessive
Heterozygous	1	1	0
Homozygous	2	1	1

Note that in a recessive model, the haplotype effects are estimated only from subjects who are homozygous for a haplotype. Some of the haplotypes which meet the *haplo.freq.min* and *haplo.count.min* cut-offs may occur as homozygous in only a few of the subjects. As stated in 5.8, recessive models should be used when the region has multiple common haplotypes.

The default haplo effect is additive, whereas the example below illustrates the fit of a dominant effect of haplotypes for the gaussian trait with the gender covariate.

```
R> # control dominant effect of haplotypes (haplo.effect)
R> # by using haplo.glm.control
R> fit.dom <- haplo.glm(y ~ male + geno.glm, family = gaussian,</pre>
         data = glm.data, na.action = "na.geno.keep",
         locus.label = label,
         control = haplo.glm.control(haplo.effect='dominant',
         haplo.min.count=8))
R> summary(fit.dom)
Call:
haplo.glm(formula = y ~ male + geno.glm, family = gaussian, data = glm.data,
    na.action = "na.geno.keep", locus.label = label, control = haplo.glm.control(haplo.effect = "dominar
        haplo.min.count = 8))
Deviance Residuals:
     Min
                1Q
                      Median
                                     3Q
                                              Max
-2.48099 -1.01196
                     0.01035
                               1.00557
                                          2.48801
```

#### Coefficients:

	coef	se	t.stat	pval
(Intercept)	1.64935	0.37350	4.41593	0.000
male	0.07969	0.15726	0.50673	0.613
geno.glm.17	-0.06035	0.42317	-0.14262	0.887
geno.glm.34	-0.66499	0.36392	-1.82731	0.069
geno.glm.77	-0.07339	0.34665	-0.21171	0.833
geno.glm.78	0.85369	0.36421	2.34394	0.020
geno.glm.100	0.24697	0.34561	0.71458	0.476
geno.glm.138	0.67295	0.28163	2.38944	0.018
geno.glm.rare	0.11195	0.34006	0.32922	0.742

```
(Dispersion parameter for gaussian family taken to be 1.300586)
```

Null deviance: 297.01 on 219 degrees of freedom Residual deviance: 274.42 on 211 degrees of freedom

AIC: 692.96

Number of Fisher Scoring iterations: 91

#### Haplotypes:

```
DQB DRB B hap.freq
geno.glm.17
              21
                    7 44 0.02297
                    4 44 0.02855
geno.glm.34
               31
geno.glm.77
               32
                    4 60 0.03019
geno.glm.78
               32
                    4 62
                         0.02391
geno.glm.100
               51
                    1 35 0.03003
geno.glm.138
               62
                    2 7
                         0.05023
geno.glm.rare
                *
                         0.71003
haplo.base
               21
                    3 8 0.10408
```

#### 6.7.2 Selecting the Baseline Haplotype

The haplotype chosen for the baseline in the model is the one with the highest frequency. Sometimes the most frequent haplotype may be an at-risk haplotype, and so the measure of its effect is desired. To specify a more appropriate haplotype as the baseline in the binomial example, choose from the list of other common haplotypes, fit.bin\$haplo.common. To specify an alternative baseline, such as haplotype 77, use the control parameter haplo.base and haplotype code, as in the example below.

```
R> # control baseline selection, perform the same exact run as fit.bin,
R> # but different baseline by using haplo.base chosen from haplo.common
R> fit.bin$haplo.common
    17 34 77 78 100 138
R> fit.bin$haplo.freq.init[fit.bin$haplo.common]
[1] 0.02332031 0.02848720 0.03060053 0.02349463 0.03018431 0.05097906
R> fit.bin.base77 <- haplo.glm(y.bin ~ male + geno.glm, family = binomial,
         data = glm.data, na.action = "na.geno.keep",
         locus.label = label,
+
         control = haplo.glm.control(haplo.base=77,
         haplo.min.count=8))
R> summary(fit.bin.base77)
Call:
haplo.glm(formula = y.bin ~ male + geno.glm, family = binomial,
    data = glm.data, na.action = "na.geno.keep", locus.label = label,
    control = haplo.glm.control(haplo.base = 77, haplo.min.count = 8))
Deviance Residuals:
   Min
              1Q
                   Median
                                3Q
                                        Max
```

```
-1.5559 -0.7996 -0.6473 1.0591 2.4348
```

#### Coefficients:

```
coef
                               t.stat pval
(Intercept)
              -0.4311
                       1.3586 -0.3173 0.751
              -0.4802 0.3308 -1.4518 0.148
male
geno.glm.4
               0.9884
                       0.7328
                              1.3489 0.179
geno.glm.17
               0.2657
                       1.0254
                               0.2591 0.796
geno.glm.34
               1.3525
                       0.9223
                              1.4665 0.144
geno.glm.78
              -0.4209
                       1.0430 -0.4035 0.687
geno.glm.100
              -1.6023
                       1.3007 -1.2319 0.219
geno.glm.138
              -1.7273
                       1.0321 -1.6736 0.096
geno.glm.rare -0.2726
                      0.6834 -0.3989 0.690
```

(Dispersion parameter for binomial family taken to be 1)

```
Null deviance: 263.50 on 219 degrees of freedom Residual deviance: 233.46 on 211 degrees of freedom
```

AIC: 251.11

Number of Fisher Scoring iterations: 61

#### Haplotypes:

	DQB	DRB	В	hap.freq
geno.glm.4	21	3	8	0.10405
geno.glm.17	21	7	44	0.02303
geno.glm.34	31	4	44	0.02843
geno.glm.78	32	4	62	0.02354
geno.glm.100	51	1	35	0.02977
geno.glm.138	62	2	7	0.05181
<pre>geno.glm.rare</pre>	*	*	*	0.70880
haplo.base	32	4	60	0.03057

#### **Explanation of Results**

The above model has the same haplotypes as fit.bin, except haplotype 4, the old baseline, now has an effect estimate while haplotype 77 is the new baseline. Due to randomness in the starting values of the haplotype frequency estimation, different runs of haplo.glm may result in a different set of haplotypes meeting the minimum counts requirement for being modeled. Therefore, once you have arrived at a suitable model, and you wish to modify it by changing baseline and/or effects, you can make results consistent by controlling the randomness using set.seed, as described in section 2.4. In this document, we use the same seed before making fit.bin and fit.bin.base77.

#### 6.7.3 Rank of Information Matrix and eps.svd (NEW)

Similar to recent additions to haplo.score in section 5.7, we give the user control over the epsilon parameter determining the number of singular values when determining the rank of the information matrix in haplo.glm. Finding the generalized inverse of this matrix can be problematic when either the response variable or a covariate has a large variance and is not scaled before passed to haplo.glm. The rank of the information matrix is determined by the number of non-zero singular values a small cutoff, epsilon. When the singular values for the coefficients are on a larger numeric scale than those for the haplotype frequencies, the generalized

inverse may incorrectly determine the information matrix is not of full rank. Therefore, we allow the user to specify the epsilon as *eps.svd* in the control parameters for *haplo.glm*. A simpler fix, which we strongly suggest, is for the user to pre-scale any continuous responses or covariates with a large variance.

Here we demonstrate what happens when we increase the variance of a gaussian response by 2500. We see that the coefficients are all highly significant and the rank of the information matrix is much smaller than the scaled gaussian fit.

```
R> glm.data$ybig <- glm.data$y*50</pre>
R> fit.gausbig <- haplo.glm(formula = ybig ~ male + geno.glm, family = gaussian,
            data = glm.data, na.action = "na.geno.keep", locus.label = label,
            control = haplo.glm.control(haplo.freq.min = 0.02), x = TRUE)
R> summary(fit.gausbig)
Call:
haplo.glm(formula = ybig ~ male + geno.glm, family = gaussian,
    data = glm.data, na.action = "na.geno.keep", locus.label = label,
    control = haplo.glm.control(haplo.freq.min = 0.02), x = TRUE)
Deviance Residuals:
    Min
                1Q
                      Median
                                    3Q
                                              Max
-123.472
           -46.026
                      -3.267
                                47.450
                                         118.550
Coefficients:
                  coef
                             se
                                  t.stat pval
(Intercept)
                                 30.6849 0.000
               53.2180
                         1.7343
                4.8675
                         6.0042
                                  0.8107 0.418
male
geno.glm.17
                         0.2579 54.3195 0.000
               14.0111
                         1.0033 -15.8044 0.000
geno.glm.34
              -15.8563
geno.glm.77
               11.0835
                         0.9978 11.1078 0.000
geno.glm.78
               57.0720
                         0.3855 148.0436 0.000
geno.glm.100
               27.7784
                         0.3228
                                 86.0564 0.000
geno.glm.138
               49.1143
                         1.0334
                                 47.5256 0.000
               19.8824
geno.glm.rare
                         3.4583
                                  5.7493 0.000
(Dispersion parameter for gaussian family taken to be 3173.952)
```

Null deviance: 742530 on 219 degrees of freedom Residual deviance: 669704 on 211 degrees of freedom

AIC: 2408.9

Number of Fisher Scoring iterations: 268

#### Haplotypes:

DQB DRB B hap.freq geno.glm.17 21 7 44 0.02291 4 44 0.02858 geno.glm.34 31 geno.glm.77 32 4 60 0.03022 geno.glm.78 32 4 62 0.02390 geno.glm.100 51 1 35 0.03008 geno.glm.138 62 2 7 0.05023 geno.glm.rare \* 0.71000 haplo.base 21 3 8 0.10409

```
R> fit.gausbig$rank.info
[1] 175
R> fit.gaus$rank.info
[1] 182
  Now we set a smaller value for the eps.svd control parameter and find the fit matches the original Gaussian
fit.
R> fit.gausbig.eps <- haplo.glm(formula = ybig ~ male + geno.glm, family = gaussian,
            data = glm.data, na.action = "na.geno.keep", locus.label = label,
+
            control = haplo.glm.control(eps.svd=1e-10, haplo.freq.min = 0.02), x = TRUE)
R> summary(fit.gausbig.eps)
Call:
haplo.glm(formula = ybig ~ male + geno.glm, family = gaussian,
    data = glm.data, na.action = "na.geno.keep", locus.label = label,
    control = haplo.glm.control(eps.svd = 1e-10, haplo.freq.min = 0.02),
    x = TRUE
```

#### Deviance Residuals:

Min 1Q Median 3Q Max -123.472 -46.026 -3.267 47.450 118.550

#### Coefficients:

	coef	se	t.stat	pval
(Intercept)	53.2180	17.1414	3.1046	0.002
male	4.8675	7.7603	0.6272	0.531
geno.glm.17	14.0111	21.7745	0.6435	0.521
geno.glm.34	-15.8563	17.1712	-0.9234	0.357
geno.glm.77	11.0835	18.0631	0.6136	0.540
geno.glm.78	57.0720	19.1910	2.9739	0.003
geno.glm.100	27.7784	18.2133	1.5252	0.129
geno.glm.138	49.1143	15.1646	3.2387	0.001
<pre>geno.glm.rare</pre>	19.8824	9.0957	2.1859	0.030

(Dispersion parameter for gaussian family taken to be 3173.952)

Null deviance: 742530 on 219 degrees of freedom Residual deviance: 669704 on 211 degrees of freedom

AIC: 2408.9

Number of Fisher Scoring iterations: 268

#### Haplotypes:

DQB DRB B hap.freq geno.glm.17 21 7 44 0.02291 geno.glm.34 31 4 44 0.02858 geno.glm.77 32 4 60 0.03022 geno.glm.78 32 4 62 0.02390

```
geno.glm.100 51 1 35 0.03008
geno.glm.138 62 2 7 0.05023
geno.glm.rare * * * 0.71000
haplo.base 21 3 8 0.10409
```

R> fit.gausbig.eps\$rank.info

[1] 182

#### 6.7.4 Rare Haplotypes and haplo.min.info

Another notable control parameter is the minimum frequency for a rare haplotype to be included in the calculations for standard error (se) of the coefficients, or *haplo.min.info*. The default value is 0.001, which means that haplotypes with frequency less than that will be part of the rare haplotype coefficient estimate, but it will not be used in the standard error calculation.

The following example demonstrates a possible result when dealing with the rare haplotype effect. We show with the hla genotype data one consequence for when this occurs. However, we make it happen by setting haplo.freq.min equal to haplo.min.info, which we advise strongly against in your analyses.

```
R> ## set haplo.freq.min and haplo.min.info to same value to show how the
R> ## rare coefficient may be modeled but standard error estimate is not
R> ## calculated because all haps are below haplo.min.info
R> ## warning expected
R> fit.bin.rare02 <- haplo.glm(y.bin ~ geno.glm, family = binomial,</pre>
         data = glm.data, na.action = "na.geno.keep", locus.label = label,
         control = haplo.glm.control(haplo.freq.min=.02, haplo.min.info=.02))
R> summary(fit.bin.rare02)
Call:
haplo.glm(formula = y.bin ~ geno.glm, family = binomial, data = glm.data,
   na.action = "na.geno.keep", locus.label = label, control = haplo.glm.control(haplo.freq.min = 0.02,
        haplo.min.info = 0.02))
Deviance Residuals:
   Min
              1Q
                   Median
                                3Q
                                        Max
-1.4558 -0.7238 -0.7238
                            1.0382
                                     2.3083
Coefficients:
                 coef
                           se
                              t.stat pval
(Intercept)
               1.2409
                       1.3238
                              0.9374 0.350
```

```
geno.glm.17   -0.6068   1.5630   -0.3882   0.698
geno.glm.34   0.3189   1.2678   0.2516   0.802
geno.glm.77   -1.0719   1.3666   -0.7844   0.434
```

geno.glm.78 -1.3593 4.3659 -0.3113 0.756 geno.glm.100 -2.3984 2.0878 -1.1487 0.252

geno.glm.138 -2.6096 1.5043 -1.7347 0.084

geno.glm.rare -1.2233 NA NA NA

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 263.50 on 219 degrees of freedom Residual deviance: 235.59 on 212 degrees of freedom

AIC: 251.24

Number of Fisher Scoring iterations: 71

#### Haplotypes:

	DQB	DRB	В	hap.freq
geno.glm.17	21	7	44	0.02302
geno.glm.34	31	4	44	0.02841
geno.glm.77	32	4	60	0.03058
geno.glm.78	32	4	62	0.02353
geno.glm.100	51	1	35	0.02980
geno.glm.138	62	2	7	0.05190
<pre>geno.glm.rare</pre>	*	*	*	0.70872
haplo.base	21	3	8	0.10403

# **Explanation of Results**

The above results show the standard error for the rare haplotype coefficient is "NaN", or "Not a Number" in R, which is a consequence of having most, or all, of the rare haplotypes discarded for the standard error estimate. In other datasets there may be only a few haplotypes between *haplo.min.info* and *haplo.freq.min*, and may yield misleading results for the rare haplotype coefficient. For this reason, we recommend that any inference made on the rare haplotypes be made with caution, if at all.

# 7 Methods for haplo.glm (NEW)

The latest updates to haplo.stats is our work to make haplo.glm to act similar to a glm object with methods to compare and assess model fits. In this section we describe the challenges and caveats of defining these methods for a haplo.glm object and show how to use them.

#### 7.1 fitted.values

A challenge when defining methods for haplo.glm is that we account for the ambiguity in a person's haplotype pair. To handle this in the glm framework, the response and non-haplotype covariates are expanded for each person with a posterior probability of the haplotype given their genotype as a weight. The returned object from haplo.glm looks somewhat like a regular glm, but the model matrix, response, and thus the fitted values, are all expanded. Users who want to work with the expanded versions of those items are welcome to access them from the returned object.

We now provide a method to get the fitted values for each person, fitted.haplo.glm. These collapsed fitted values are calculated by a weighted sum of the expanded fitted values for each person where the weights are the posterior probabilities of the person's expanded haplotype pairs.

#### 7.2 residuals

The residuals within the *haplo.glm* object are also expanded for the haplotype pairs for subjects. We provide residuals.haplo.glm to get the collapsed deviance, pearson, working, and response residuals for each person. Because we have not implemented a predict method for *haplo.glm*, the method does not calculate partial residuals.

#### 7.3 vcov

We provide vcov.haplo.glm as a method to get the variance-covariance matrix of model parameters in the haplo.glm object. Unlike the standard glm object, this matrix is computed and retained in the returned

object. We do this because the model parameters are the model coefficients and the haplotype frequencies, and it is computationally-intensive to compute.

We show how to get the variance matrix for all the parameters and for only the model coefficients.

```
R> varmat <- vcov(fit.gaus)</pre>
R> dim(varmat)
[1] 182 182
R> varmat <- vcov(fit.gaus, freq=FALSE)</pre>
R> dim(varmat)
[1] 9 9
R> print(varmat, digits=2)
               (Intercept)
                                male geno.glm.17 geno.glm.34 geno.glm.77 geno.glm.78
                                          -0.0674
                                                       -0.0544
                                                                   -0.0526
(Intercept)
                     0.118 -0.01513
                                                                               -0.05123
male
                    -0.015
                           0.02409
                                           0.0065
                                                       -0.0022
                                                                   -0.0038
                                                                                0.00082
                           0.00646
                    -0.067
                                          0.1897
                                                       0.0335
                                                                    0.0206
geno.glm.17
                                                                                0.03217
geno.glm.34
                    -0.054 -0.00218
                                          0.0335
                                                       0.1179
                                                                    0.0226
                                                                                0.02753
geno.glm.77
                    -0.053 -0.00375
                                          0.0206
                                                       0.0226
                                                                    0.1305
                                                                                0.02602
geno.glm.78
                    -0.051
                            0.00082
                                          0.0322
                                                       0.0275
                                                                    0.0260
                                                                                0.14732
geno.glm.100
                    -0.059
                            0.00674
                                          0.0370
                                                       0.0298
                                                                    0.0204
                                                                                0.02853
geno.glm.138
                    -0.059
                            0.00362
                                          0.0307
                                                       0.0254
                                                                    0.0256
                                                                                0.02195
geno.glm.rare
                    -0.058
                            0.00142
                                          0.0320
                                                       0.0283
                                                                    0.0278
                                                                                0.02478
               geno.glm.100 geno.glm.138 geno.glm.rare
(Intercept)
                    -0.0595
                                  -0.0587
                                                 -0.0583
male
                     0.0067
                                   0.0036
                                                  0.0014
geno.glm.17
                     0.0370
                                   0.0307
                                                  0.0320
geno.glm.34
                     0.0298
                                   0.0254
                                                  0.0283
geno.glm.77
                     0.0204
                                   0.0256
                                                  0.0278
geno.glm.78
                     0.0285
                                   0.0220
                                                  0.0248
geno.glm.100
                     0.1327
                                   0.0214
                                                  0.0281
                                                  0.0288
geno.glm.138
                     0.0214
                                   0.0920
geno.glm.rare
                     0.0281
                                   0.0288
                                                  0.0331
```

# 7.4 anova and Model Comparison

We use the anova.glm method as a framework for anova.haplo.glm to allow comparisons of model fits. We limit the model comparisons to multiple nested model fits, which requires that each model to be compared is either a haplo.glm or glm fitted object. We eliminate the functionality of testing sub-models of a single fit because removal of a single covariate would require re-fitting of the reduced model to get updated coefficient and haplotype frequency estimates with a maximized log-likelihood. We decided to simplify the usage and require that all models to be compared are fully fitted.

As with the *anova.glm* method, it is difficult to check for truly nested models, so we pass the responsibility on to the user. We discuss some of the requirements.

One type of two-model comparison is between models with haplotypes (expanded subjects) and a reduced model without haplotypes. We check for the same sample size in these models by comparing the collapsed sample size from a haplo.glm fit to the sample size from the glm fit, which we remind users is only a loose check of model comparability.

The other comparison of two models in *anova.haplo.glm* is to compare two models that contain the same genotypes, and inherently the same haplotypes. This is more tricky because a subject may not have the same

expanded set of possible haplotype pairs across two fits of haplo.glm unless the same seed is set before both fits. Even if a seed is the same, the other effects that are different between the two models will affect the haplotype frequency estimates, and may still result in a different expansion of haplotype pairs per subject. Our check of the collapsed sample size for the two models still applies with the same pitfalls, but a better assurance of model comparability is to use the same seed.

In the haplo.glm fit we provide the likelihood ratio test of the null model against the full model, which is the most appropriate test available for haplo.glm objects, but it is difficult to compare the log-likelihood across two haplo.glm fits. Therefore, we remain consistent with glm model comparison [10], and use the difference in deviance to compare models. Furthermore, we restrict the asymptotic test for model comparison to be the  $\chi^2$  test for goodness of fit.

Below we show how to get the LRT from the fit.gaus result, then show how to compare some of the nested models fit above, including a regular glm fit of  $y \sim male$ . The anova method requires the nested model to be given first, and any anova with a haplo.glm object should explicitly call anova.haplo.glm.

```
R> fit.gaus$1rt
$1rt
[1] 22.34062
$df
[1] 8
R> glmfit.gaus <- glm(y~male, family="gaussian", data=glm.data)
R> anova.haplo.glm(glmfit.gaus, fit.gaus)
Analysis of Deviance Table
Model 1: y ~ male
Model 2: y ~ male + geno.glm
  Resid. Df Resid. Dev Df Deviance Pr(>Chi)
1
        218
                297.00
2
        211
                267.88 7
                            29.114 0.001752 **
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
R> anova.haplo.glm(fit.gaus, fit.inter)
Analysis of Deviance Table
Model 1: y ~ male + geno.glm
Model 2: y ~ male * geno.glm
  Resid. Df Resid. Dev Df Deviance Pr(>Chi)
        211
                267.88
1
                259.82 7
        204
                            8.0631
                                     0.5017
R> anova.haplo.glm(glmfit.gaus, fit.gaus, fit.inter)
Analysis of Deviance Table
Model 1: y ~ male
Model 2: y ~ male + geno.glm
Model 3: y ~ male * geno.glm
  Resid. Df Resid. Dev Df Deviance Pr(>Chi)
```

```
1 218 297.00

2 211 267.88 7 29.1137 0.001804 **

3 204 259.82 7 8.0631 0.501696

---
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
```

# 8 Extended Applications

The following functions are designed to wrap the functionality of the major functions in Haplo Stats into other useful applications.

# 8.1 Combine Score and Group Results: haplo.score.merge

When analyzing a qualitative trait, such as binary, it can be helpful to align the results from haplo.score with haplo.group. To do so, use the function haplo.score.merge, as illustrated in the following example:

```
R> # merge haplo.score and haplo.group results
R> merge.bin <- haplo.score.merge(score.bin, group.bin)
R> print(merge.bin, nlines=10)
```

------

Haplotype Scores, p-values, and Frequencies By Group

```
DQB DRB
            B Hap.Score
                           p.val Hap.Freq y.bin.0 y.bin.1
    62
         2
            7
1
               -2.19387 0.02824
                                  0.05098 0.06789 0.01587
2
    51
         1 35
               -1.58421 0.11315
                                  0.03018 0.03754 0.00907
3
    63
        13
            7
               -1.56008 0.11874
                                  0.01655 0.02176
4
    21
         7
            7
               -1.47495 0.14023
                                  0.01246 0.01969
                                                        NA
5
    32
         4
            7
               -1.00091 0.31687
                                  0.01678 0.02628 0.00794
6
    32
         4 62
               -0.67990 0.49657
                                  0.02349 0.01911
7
    51
         1 27
               -0.66509 0.50599
                                  0.01505 0.01855 0.00907
8
    31
        11 35
               -0.58380 0.55936
                                  0.01754 0.01982 0.01587
9
    31
        11 51
               -0.43721 0.66196
                                  0.01137 0.01321
                0.00826 0.99341
                                  0.01731 0.01595 0.00000
10
    51
```

# Explanation of Results

The first column is a row index, the next columns (3 in this example) illustrate the haplotype, the Hap.Score column is the score statistic and p.val the corresponding  $\chi^2$  p-value. Hap.Freq is the haplotype frequency for the total sample, and the remaining columns are the estimated haplotype frequencies for each of the group levels (y.bin) in this example). The default print method only prints results for haplotypes appearing in the haplo.score output. To view all haplotypes, use the print option all.haps=TRUE, which prints all haplotypes from the haplo.group output. The output is ordered by the score statistic, but the order.by parameter can specify ordering by haplotypes or by haplotype frequencyies.

## 8.2 Case-Control Haplotype Analysis: haplo.cc

We provide haplo.cc to run and combine the results of haplo.score, haplo.group, and haplo.glm for case-control data. The function performs a score test and a glm on the same haplotypes. The parameters that determine which haplotypes are used are haplo.min.count and haplo.freq.min, which are set in the control parameter, as done for haplo.glm.

Below we run haplo.cc setting the minimum haplotype frequency at 0.02. The print results are shown, in addition to the names of the objects stored in the cc.hla result.

```
R> # demo haplo.cc where haplo.min.count is specified
R> # use geno, and this function prepares it for haplo.glm
R> y.bin <- 1*(hla.demo$resp.cat=="low")</pre>
R> cc.hla <- haplo.cc(y=y.bin, geno=geno, locus.label = label,
              control=haplo.glm.control(haplo.freq.min=.02))
R> print(cc.hla, nlines=25, digits=2)
______
                     Global Score Statistics
______
global-stat = 29, df = 8, p-val = 0.00029
______
                   Counts for Cases and Controls
______
control
       case
  157
        63
        Haplotype Scores, p-values, Hap-Frequencies (hf), and Odds Ratios
                           (95% CI)
______
  DQB DRB B Hap-Score p-val pool.hf control.hf case.hf glm.eff OR.lower
                                                     OR.
147 62 2 7
           -2.103 0.03546 0.0490
                             0.0679 0.0159
                                        Eff
                                              0.0138 0.072
     1 35
98
           -1.583 0.11344 0.0302
                             0.0376 0.0089
                                          Eff
                                              0.0097 0.086
   51
                             0.0263 0.0079
78
   32 4 7
           -1.393 0.16349 0.0227
                                          Eff
                                              0.0047 0.058
77
   32 4 62
          -0.496 0.62001 0.0212
                                              0.0517 0.281
                             0.0191
                                   NA
                                          Eff
```

76 32 4 60 0.028 0.97762 0.0307 0.0315 0.0238 Eff 0.0763 0.318 21 7 44 1.069 0.28516 0.0217 0.0175 0.0476 Eff 0.1253 0.661 16 31 4 44 2.516 0.01186 0.0285 0.0150 0.0635 0.3425 1.318 52 Eff 21 3 8 3.776 0.00016 0.1042 0.0693 0.1905 NA 1.000 11 Base 21 1 8 NA 0.0023 0.0033 R 0.1443 0.290 1 NANA2 21 10 8 NA NA 0.0023 0.0032 NA R 0.1443 0.290 3 21 2 18 NANA 0.0023 0.0032 NAR 0.1443 0.290 21 2 7 NA 0.0023 4 NA0.0032 0.1443 0.290 NAR NA 0.0046 5 21 3 18 NA0.0067 NAR 0.1443 0.290 6 21 3 35 NANA 0.0057 0.0065 NAR 0.1443 0.290 7 21 3 44 NANA 0.0036 0.0033 0.0159 R 0.1443 0.290 8 21 3 49 NANA 0.0023 NANAR 0.1443 0.290 9 21 3 57 NANA 0.0024 NANAR 0.1443 0.290 21 3 70 10 NANA 0.0023 NANAR 0.1443 0.290 NA 0.0045 12 21 4 62 0.0064 0.1443 0.290 NANAR NA 0.0108 7 13 13 21 NANA 0.0238 R 0.1443 0.290 14 21 7 18 NANA 0.0025 NANAR 0.1443 0.290 15 21 7 35 NANA 0.0024 NA 0.0079 0.1443 0.290 R 7 45 NA 0.0023 17 21 R 0.1443 0.290 NA0.0032 NA7 50 NA 0.0045 0.0032 0.0079 R 18 21 NA0.1443 0.290 NA NA 0.0023 19 21 7 57 0.0064 NA R 0.1443 0.290

OR.upper

147 0.38

98 0.76

```
0.72
78
77
         1.53
76
         1.32
16
         3.48
52
         5.07
11
           NA
         0.58
1
2
         0.58
3
         0.58
4
         0.58
5
         0.58
6
         0.58
7
         0.58
8
         0.58
9
         0.58
10
         0.58
12
         0.58
13
         0.58
14
         0.58
15
         0.58
17
         0.58
18
         0.58
         0.58
19
R> names(cc.hla)
[1] "cc.df"
                     "group.count"
                                      "score.lst"
                                                       "fit.lst"
                                                                         "ci.prob"
[6] "exclude.subj"
```

#### **Explanation of Results**

First, from the names function we see that cc.hla also contains score.lst and fit.lst, which are the haplo.score and haplo.glm objects, respectively. For the printed results of haplo.cc, first are the global statistics from haplo.score, followed by cell counts for cases and controls. The last portion of the output is a data frame containing combined results for individual haplotypes:

- **Hap-Score:** haplotype score statistic
- p-val: haplotype score statistic p-value
- sim p-val: (if simulations performed) simulated p-value for the haplotype score statistic
- pool.hf: haplotype frequency for the pooled sample
- control.hf: haplotype frequencies for the control sample only
- case.hf: haplotype frequencies for the case sample only
- **glm.eff:** one of three ways the haplotype appeared in the glm model: *Eff:* modeled as an effect; *Base*: part of the baseline; and *R*: a rare haplotype, included in the effect of pooled rare haplotypes
- OR.lower: Odds Ratio confidence interval lower limit
- OR: Odds Ratio for each effect in the model
- OR.upper: Odds Ratio confidence interval upper limit

Significance levels are indicated by the p-values for the score statistics, and the odds ratio (OR) confidence intervals for the haplotype effects. Note that the Odds Ratios are effect sizes of haplotypes, assuming haplotype effects are multiplicative. Since this last table has many columns, lines are wrapped in the output in this manual. You can align wrapped lines by the haplotype code which appears on the far left. Alternatively, instruct the print function to only print digits significant digits, and set the width settings for output in your session using the options() function.

#### 8.3 Score Tests on Sub-Haplotypes: haplo.score.slide

To evaluate the association of sub-haplotypes (subsets of alleles from the full haplotype) with a trait, the user can evaluate a "window" of alleles by *haplo.score*, and slide this window across the entire haplotype. This procedure is implemented by the function *haplo.score.slide*. To illustrate this method, we use all 11 loci in the demo data, *hla.demo*.

First, make the geno matrix and the locus labels for the 11 loci. Then use *haplo.score.slide* for a window of 3 loci (*n.slide=3*), which will slide along the haplotype for all 9 contiguous subsets of size 3, using the previously defined gaussian trait resp.

```
R> # haplo.score on 11 loci, slide on 3 consecutive loci at a time
R > geno.11 \leftarrow hla.demo[,-c(1:4)]
R> label.11 <- c("DPB","DPA","DMA","DMB","TAP1","TAP2","DQB","DQA","DRB","B","A")
R> score.slide.gaus <- haplo.score.slide(hla.demo$resp, geno.11, trait.type =
                   "gaussian", n.slide=3, min.count=5, locus.label=label.11)
R> print(score.slide.gaus)
  start.loc score.global.p global.p.sim max.p.sim
1
          1
                    0.21550
                                                  NA
          2
2
                    0.09366
                                        NA
                                                  NA
3
          3
                    0.39042
                                        NA
                                                  NA
4
          4
                    0.48771
                                        NA
                                                  NA
          5
5
                    0.13747
                                        NA
                                                  NA
6
          6
                    0.14925
                                        NA
                                                  NA
7
          7
                    0.11001
                                        NA
                                                  NA
8
          8
                    0.00996
                                        NA
                                                  NA
          9
                    0.04255
                                        NA
                                                  NA
```

#### **Explanation of Results**

The first column is the row index of the nine calls to haplo.score, the second column is the number of the starting locus of the sub-haplotype, the third column is the global score statistic p-value for each call. The last two columns are the simulated p-values for the global and maximum score statistics, respectively. If you specify simulate = TRUE in the function call, the simulated p-values would be present.

#### **8.3.1** Plot Results from haplo.score.slide

The results from haplo.score.slide can be easily viewed in a plot shown in Figure 2 below. The x-axis has tick marks for each locus, and the y-axis is the  $-log_{10}(pval)$ . To select which p-value to plot, use the parameter pval, with choices "global", "global.sim", and "max.sim" corresponding to p-values described above. If the simulated p-values were not computed, the default is to plot the global p-values. For each p-value, a horizontal line is drawn at the height of  $-log_{10}(pval)$  across the loci over which it was calculated. For example, the p-value score.global.p = 0.009963 for loci 8-10 is plotted as a horizontal line at y = 2.002 spanning the  $8^{th}$ , and  $10^{th}$  x-axis tick marks.

R> # plot global p-values for sub-haplotypes from haplo.score.slide R> plot.haplo.score.slide(score.slide.gaus)

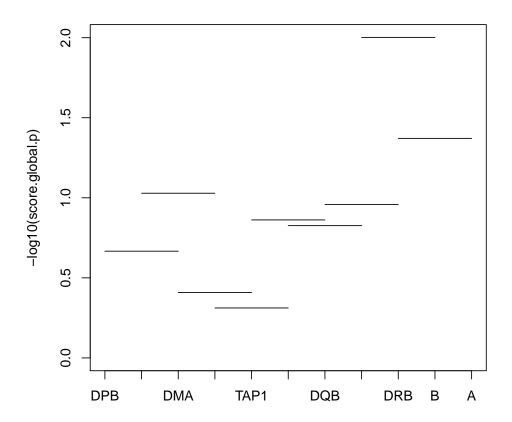


Figure 2: Global p-values for sub-haplotypes; Gaussian Response

# 8.4 Scanning Haplotypes Within a Fixed-Width Window: haplo.scan

Another method to search for a candidate locus within a genome region is haplo.scan, an implementation of the method proposed in Cheng et al. 2005 [11]. This method searches for a region for which the haplotypes have the strongest association with a binary trait by sliding a window of fixed width over each marker locus, and then scans over all haplotype lengths within each window. This latter step, scanning over all possible haplotype lengths within a window, distinguishes haplo.scan from haplo.score.slide (which considers only the maximum haplotype length within a window). To account for unknown linkage phase, the function haplo.em is called prior to scanning, to create a list of haplotype pairs and posterior probabilities. To illustrate the scanning window, consider a 10-locus dataset. When placing a window of width 3 over locus 5, the possible haplotype lengths that contain locus 5 are three (loci 3-4-5, 4-5-6, and 5-6-7), two (loci 4-5 and 5-6) and one (locus 5). For each of these loci subsets a score statistic is computed, which is based on the difference between the mean vector of haplotype counts for cases and that for controls. The maximum of these score statistics, over all possible haplotype lengths within a window, is the locus-specific test statistic, or the locus scan statistic. The global test statistic is the maximum over all computed score statistics. To compute p-values, the case/control status is randomly permuted. Below we run haplo.scan on the 11-locus HLA dataset with a binary response and a window width of 3, but first we use the results of summary Geno to choose subjects with less than 50,000 haplotype pairs to speed calculations with all 11 polymorphic loci with many missing alleles.

```
R > geno.11 \leftarrow hla.demo[,-c(1:4)]
R> y.bin <- 1*(hla.demo$resp.cat=="low")
R> hla.summary <- summaryGeno(geno.11, miss.val=c(0,NA))
R> # track those subjects with too many possible haplotype pairs ( > 50,000)
R> many.haps <- (1:length(y.bin))[hla.summary[,4] > 50000]
R> # For speed, or even just so it will finish, make y.bin and geno.scan
R> # for genotypes that don't have too many ambigous haplotypes
R> geno.scan <- geno.11[-many.haps,]</pre>
R> y.scan <- y.bin[-many.haps]</pre>
R> # scan haplotypes for regions within width of 3 for each locus.
R> # test statistic measures difference in haplotype counts in cases and controls
R> # p-values are simulated for each locus and the maximum statistic,
R> # we do 100 simuations here, should use default settings for analysis
R>
R> scan.hla <- haplo.scan(y.scan, geno.scan, width=3,
          sim.control=score.sim.control(min.sim=100, max.sim=100),
          em.control=haplo.em.control())
R> print(scan.hla)
  Call:
haplo.scan(y = y.scan, geno = geno.scan,
   width = 3, em.control = haplo.em.control(),
    sim.control = score.sim.control(min.sim = 100,
        max.sim = 100)
                         Locus Scan-statistic Simulated P-values
          loc-1 loc-2 loc-3 loc-4 loc-5 loc-6 loc-7 loc-8 loc-9 loc-10 loc-11
                                            0 0
sim.p-val
                    0
                          0
                                0
                                      0
                                                        0
                                                               0
```

Max-Stat Simulated Global p-value: 0
Number of Simulations: 100

### **Explanation of Results**

In the output we report the simulated p-values for each locus test statistic. Additionally, we report the loci (or locus) which provided the maximum observed test statistic, and the Max-Stat Simulated Global p-value is the simulated p-value for that maximum statistic. We print the number of simulations, because they are performed until p-value precision criteria are met, as described in section 5.9. We would typically allow simulations to run under default parameters rather than limiting to 100 by the control parameters.

#### 8.5 Sequential Haplotype Scan Methods: seghap

Another approach for choosing loci for haplotype associations is seqhap, as described in Yu and Schaid, 2007 [12]. The seqhap method performs three tests for association of a binary trait over a set of bi-allelic loci. When evaluating each locus, loci close to it are added in a sequential manner based on the Mantel-Haenszel test [13]. For each marker locus, three tests are provided:

- single locus, the traditional single-locus  $\chi_1^2$  test of association,
- sequential haplotype, based on a haplotype test for sequentially chosen loci,
- sequential sum, based on the sum of a series of conditional  $\chi^2$  statistics.

All three tests are assessed for significance with permutation p-values, in addition to the asymptotic p-value. The point-wise p-value for a statistic at a locus is the fraction of times that the statistic for the permuted data is larger than that for the observed data. The regional p-value is the chance of observing a permuted test statistic, maximized over a region, that is greater than that for the observed data.

Similar to the permutation p-values in *haplo.score* as described in section 5.9, permutations are performed until a precision threshold is reached for the regional p-values. A minimum and maximum number of permutations specified in the *sim.control* parameter list ensure a certain accuracy is met for every simulation p-value, yet having a limit to avoid infinite run-time.

Below is an example of using seqhap on data with case-control response for a chromosome region. First set up the binary response, y, with 0=control, 1=case, then a genotype matrix with two columns per locus, and a vector of chromosome positions. The genotype data is available in the seqhap.dat dataset while the chromosome positions are in seqhap.pos. The following example runs seqhap with default settings for permutations and threshold parameters.

```
_______
                            Single-locus Chi-square Test
_____
Regional permuted P-value based on single-locus test is 0.13191
      chi.stat perm.point.p asym.point.p
loc-1 1.22062 0.27729741
                           0.26924
loc-2 1.35462 0.23245526
                             0.24447

      loc-3
      5.20288
      0.02010859
      0.02255

      loc-4
      3.36348
      0.05972250
      0.06666

      loc-5
      3.55263
      0.06153227
      0.05945

loc-6 0.39263 0.53026342
                             0.53092
loc-7 5.54913 0.01829881
                              0.01849
loc-8 3.74740 0.05469535

      loc-8
      3.74740
      0.05469535
      0.05289

      loc-9
      0.03602
      0.85682687
      0.84947

      loc-10
      1.99552
      0.17313493
      0.15777

______
                                  Sequential Scan
_____
Loci Combined in Sequential Analysis
seq-loc-1 1
seq-loc-2 2 3 4 5
seq-loc-3 3 4 5
seq-loc-4 4 3
seq-loc-5 5
seq-loc-6 6 7
seq-loc-7 7
seq-loc-8 8
seq-loc-9 9
seq-loc-10 10
______
                             Sequential Haplotype Test
______
Regional permuted P-value based on sequential haplotype test is 0.016489
          hap.stat df perm.point.p asym.point.p
seq-loc-1 1.22062 1 0.310878745 0.26924
seq-loc-2 24.16488 12 0.027950935
                                    0.01932
seg-loc-3 19.78808 6 0.005228232
                                    0.00302
seq-loc-4 14.95765 3 0.003016288
                                    0.00185
seq-loc-5 3.55263 1 0.096722300
                                    0.05945
seq-loc-6 5.45723 2 0.114216771
                                    0.06531
seq-loc-7 5.54913 1 0.038608486
                                    0.01849

      seq-loc-8
      3.74740
      1
      0.103961392
      0.05289

      seq-loc-9
      0.03602
      1
      0.867886588
      0.84947
```

0.15777

seq-loc-10 1.99552 1 0.219384677

\_\_\_\_\_\_

Regional pe	ermuted P-	-val	lue based on	sequential sum	test is	0.0032174
	<pre>sum.stat</pre>	df	<pre>perm.point.p</pre>	asym.point.p		
seq-loc-1	1.22062	1	0.3108787452	0.26924		
seq-loc-2	21.15360	4	0.0008043435	0.00030		
seq-loc-3	18.65769	3	0.0008043435	0.00032		
seq-loc-4	14.61897	2	0.0020108586	0.00067		
seq-loc-5	3.55263	1	0.1033581339	0.05945		
seq-loc-6	5.43826	2	0.1150211140	0.06593		
seq-loc-7	5.54913	1	0.0386084858	0.01849		
seq-loc-8	3.74740	1	0.1041624774	0.05289		
seq-loc-9	0.03602	1	0.8678865876	0.84947		
seq-loc-10	1.99552	1	0.2193846773	0.15777		

#### **Explanation of Results**

The output above first shows n.sim, the number of permutations needed for precision on the regional p-values. Next, in the printed results, the first section (Single-locus Chi-square Test) shows a table with columns for single-locus tests. The table includes test statistics, permuted p-values, and asymptotic p-values based on a  $\chi_1^2$  distribution. The second section (Sequential Scan) shows which loci are combined for association. In this example, the table shows the first locus is not combined with other loci, whereas the second locus is combined with loci 3, 4, and 5. The third section (Sequential Haplotype Test), shows the test statistics for the sequential haplotype method with degrees of freedom and permuted and asymptotic p-values. The fourth section (Sequential Sum Test) shows similar information for the sequential sum tests.

#### **8.5.1** Plot Results from seqhap

The results from seqhap can be viewed in a useful plot shown in Figure 3. The plot is similar to the plot for haplo.score.slide results, with the x-axis having tick marks for all loci and the y-axis is the -log10() of p-value for the tests performed. For the sequential result for each locus, a horizontal line at the height of -log10(p-value) is drawn across the loci combined. The start locus is indicated by a filled triangle and other loci combined with the start locus are indicated by an asterisk or circle. The choices for pval include "hap" (sequential haplotype asymptotic p-value), "hap.sim" (sequential haplotype simulated p-value), "sum" (sequential sum asymptotic p-value), and "sum.sim" (sequential sum simulated p-value). The other parameter option is single, indicating whether to plot a line for the single-locus tests.

R> # plot global p-values for sub-haplotypes from haplo.score.slide R> plot(seqhap.out, pval="hap", single=TRUE, las=2)

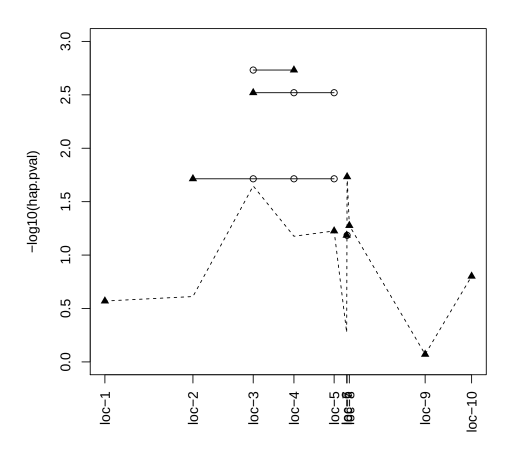


Figure 3: Plot p-values for sequential haplotype scan and single-locus tests

## 8.6 Creating Haplotype Effect Columns: haplo.design

In some instances, the desired model for haplotype effects is not possible with the methods given in haplo.glm. Examples include modeling just one haplotype effect, or modeling an interaction of haplotypes from different chromosomes, or analyzing censored data. To circumvent these limitations, we provide a function called haplo.design, which will set up an expected haplotype design matrix from a haplo.em object, to create columns that can be used to model haplotype effects in other modeling functions.

The function haplo.design first creates a design marix for all pairs of haplotypes over all subjects, and then uses the posterior probabilities to create a weighted average contribution for each subject, so that the number of rows of the final design matrix is equal to the number of subjects. This is sometimes called the expectation-substitution method, as proposed by Zaykin et al. 2002 [4], and using this haplotype design matrix in a regression model is asymptotically equivalent to the score statistics from haplo.score (Xie and Stram 2005 [14]). Although this provides much flexibility, by using the design matrix in any type of regression model, the estimated regression parameters can be biased toward zero (see Lin and Zeng, 2006 [15] for concerns about the expectation-substitution method).

In the first example below, using default parameters, the returned data frame contains a column for each haplotype that meets a minimum count in the sample *min.count*. The columns are named by the code they are assigned in *haplo.em*.

```
R> # create a matrix of haplotype effect columns from haplo.em result
R> hap.effect.frame <- haplo.design(save.em)</pre>
R> names(hap.effect.frame)
 [1] "hap.4"
                "hap.13"
                           "hap.17"
                                      "hap.34"
                                                 "hap.50"
                                                            "hap.55"
                                                                       "hap.69"
 [9] "hap.78"
                "hap.99"
                           "hap.100" "hap.102" "hap.138" "hap.140" "hap.143" "hap.155"
[17] "hap.162"
                "hap.165"
R> hap.effect.frame[1:10,1:8]
                       hap.17 hap.34 hap.50 hap.55 hap.69 hap.77
   hap.4
             hap.13
       0 0.000000 0.0000000
                                     0
                                                    0
1
                                            0
                                                            0
                                                                   0
2
       0 0.1253234 0.8746766
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
3
       0 0.0000000 0.0000000
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
4
       0 0.2862131 0.7137869
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
5
       0 0.000000 0.0000000
                                     0
                                            0
                                                            0
                                                                   0
                                                    1
6
       1 0.0000000 1.0000000
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
7
       0 0.000000 0.0000000
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
8
       0 0.0000000 0.0000000
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
9
       0 0.0000000 0.0000000
                                     0
                                            0
                                                    0
                                                            0
                                                                   0
10
       0 0.0000000 0.0000000
                                            0
                                                    0
                                                            0
                                                                   0
```

Additionally, haplo.design gives the user flexibility to make a more specific design matrix with the following parameters:

- hapcodes: codes assigned in the haplo.em object, the only haplotypes to be made into effects
- haplo.effect: the coding of haplotypes as additive, dominant, or recessive
- haplo.base: code for the baseline haplotype
- min.count: minimum haplotype count

This second example below creates columns for specific haplotype codes that were most interesting in score.gaus.add, haplotypes with alleles 21-3-8 and 62-2-7, corresponding to codes 4 and 138 in haplo.em, respectively. Assume we want to test their individual effects when they are coded with haplo.effect="dominant".

```
R> # create haplotype effect cols for haps 4 and 138
R> hap4.hap138.frame <- haplo.design(save.em, hapcodes=c(4,138),
                                 haplo.effect="dominant")
R> hap4.hap138.frame[1:10,]
  hap.4 hap.138
      0 0.0000000
1
2
      0 0.8746766
3
      0.0000000
4
      0.0000000
5
      0 0.0000000
6
      1 0.0000000
7
      0 1.0000000
8
      0 0.0000000
      0 0.1358696
9
10
      0 0.0000000
R> dat.glm <- data.frame(resp, male, age,
                       hap.4=hap4.hap138.frame$hap.4,
+
                      hap.138=hap4.hap138.frame$hap.138)
R> glm.hap4.hap138 <- glm(resp ~ male + age + hap.4 + hap.138,
                       family="gaussian", data=dat.glm)
R> summary(glm.hap4.hap138)
Call:
glm(formula = resp ~ male + age + hap.4 + hap.138, family = "gaussian",
   data = dat.glm)
Deviance Residuals:
               1Q
                    Median
                                  3Q
                                           Max
-2.32614 -1.07489 -0.06559
                                       2.39044
                            1.04483
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 1.913834 0.229577
                               8.336 9.11e-15 ***
           0.048588 0.155290
                               0.313
                                         0.7547
male
           -0.002651
                      0.011695 -0.227
                                         0.8209
age
           -0.405530 0.195857 -2.071
                                         0.0396 *
hap.4
           hap.138
Signif. codes: 0 âĂŸ***âĂŹ 0.001 âĂŸ**âĂŹ 0.01 âĂŸ*âĂŹ 0.05 âĂŸ.âĂŹ 0.1 âĂŸ âĂŹ 1
(Dispersion parameter for gaussian family taken to be 1.318277)
   Null deviance: 297.01 on 219 degrees of freedom
Residual deviance: 283.43 on 215 degrees of freedom
AIC: 692.07
Number of Fisher Scoring iterations: 2
```

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# **Appendix**

# A Counting Haplotype Pairs When Marker Phenotypes Have Missing Alleles

The following describes the process for counting the number of haplotype pairs that are consistent with a subject's observed marker phenotypes, allowing for some loci with missing data. Note that we refer to marker phenotypes, but our algorithm is oriented towards typical markers that have a one-to-one correspondence with their genotypes. We first describe how to count when none of the loci have missing alleles, and then generalize to allow loci to have either one or two missing alleles. When there are no missing alleles, note that homozygous loci are not ambiguous with respect to the underlying haplotypes, because at these loci the underlying haplotypes will not differ if we interchange alleles between haplotypes. In contrast, heterozygous loci are ambiguous, because we do not know the haplotype origin of the distinguishable alleles (i.e., unknown linkage phase). However, if there is only one heterozygous locus, then it doesn't matter if we interchange alleles, because the pair of haplotypes will be the same. In this situation, if parental origin of alleles were known, then interchanging alleles would switch parental origin of haplotypes, but not the composition of the haplotypes. Hence, ambiguity arises only when there are at least two heterozygous loci. For each heterozygous locus beyond the first one, the number of possible haplotypes increases by a factor of 2, because we interchange the two alleles at each heterozygous locus to create all possible pairs of haplotypes. Hence, the number of possible haplotype pairs can be expressed as  $2^x$ , where x = H - 1, if H (the number of heterozygous loci) is at least 2, otherwise x = 0.

Now consider a locus with missing alleles. The possible alleles at a given locus are considered to be those that are actually observed in the data. Let  $a_i$  denote the number of distinguishable alleles at the locus. To count the number of underlying haplotypes that are consistent with the observed and missing marker data, we need to enumerate all possible genotypes for the loci with missing data, and consider whether the imputed genotypes are heterozygous or homozygous.

To develop our method, first consider how to count the number of genotypes at a locus, say the  $i^{th}$  locus, when either one or two alleles are missing. This locus could have either a homozygous or heterozygous genotype, and both possibilities must be considered for our counting method. If the locus is considered as homozygous, and there is one allele missing, then there is only one possible genotype; if there are two alleles missing, then there are  $a_i$  possible genotypes. A function to perform this counting for homozygous loci is denoted  $f(a_i)$ . If the locus is considered as heterozygous, and there is one allele missing, then there are  $a_i - 1$  possible genotypes; if there are two alleles missing, then there are  $\frac{a_i(a_i-1)}{2}$  possible genotypes. A function to perform this counting for heterozygous loci is denoted  $g(a_i)$  These functions and counts are summarized in Table A.1.

**Table A.1:** Factors for when a locus having missing allele(s) is counted as homozygous(f()) or heterozygous(g())

Number of	Homozygous	Heterozygous
missing alleles	function $f(a_i)$	function $g(a_i)$
1	1	$a_i - 1$
2	$a_i$	$\frac{a_i(a_i-1)}{2}$

Now, to use these genotype counting functions to determine the number of possible haplotype pairs, first consider a simple case where only one locus, say the  $i^{th}$  locus, has two missing alleles. Suppose that the phenotype has H heterozygous loci (H is the count of heterozygous loci among those without missing data). We consider whether the locus with missing data is either homozygous or heterozygous, to give the count of possible haplotype pairs as

$$a_i 2^x + \left[ \frac{a_i (a_i - 1)}{2} \right] 2^{x+1} \tag{1}$$

where again x = H - 1 if H is at least 2, otherwise x = 0. This special case can be represented by our more general genotype counting functions as

$$f(a_i) 2^x + g(a_i) 2^{x+1} (2)$$

When multiple loci have missing data, we need to sum over all possible combinations of heterozygous and homozygous genotypes for the incomplete loci. The rows of Table A.2 below present these combinations for up to m=3 loci with missing data. Note that as the number of heterozygous loci increases (across the columns of Table A.2), so too does the exponent of 2. To calculate the total number of pairs of haplotypes, given observed and possibly missing genotypes, we need to sum the terms in Table A.2 across the appropriate row. For example, with m=3, there are eight terms to sum over. The general formulation for this counting method can be expressed as

$$TotalPairs = \sum_{j=0}^{m} \sum_{combo} C(combo, j)$$
(3)

where combo is a particular pattern of heterozygous and homozygous loci among the loci with missing values (e.g., for m = 3, one combination is the first locus heterozygous and the  $2^{nd}$  and  $3^{rd}$  third as homozygous), and C(combo, j) is the corresponding count for this pattern when there are i loci that are heterozygous (e.g., for m = 3 and j = 1, as illustrated in Table A.2).

**Table A.2:** Genotype counting terms when m loci have missing alleles, grouped by number of heterozygous loci (out of m)

$\lfloor m \rfloor$	j = 0  of  m	j = 1  of  m	j = 2 of m	j = 3  of  m
0	$2^x$			
1	$f(a_1)2^x$	$g(a_1)2^{x+1}$		
2	$f(a_1)f(a_2)2^x$	$g(a_1)f(a_2)2^{x+1}$	$g(a_1)g(a_2)2^{x+1}$	
		$f(a_1)g(a_2)2^{x+1}$		
3	$\int f(a_1)f(a_2)f(a_3)2^x$	$g(a_1)f(a_2)f(a_3)2^{x+1}$	$g(a_1)g(a_2)f(a_3)2^{x+2}$	$g(a_1)g(a_2)g(a_3)2^{x+2}$
		$f(a_1)g(a_2)f(a_3)2^{x+1}$	$g(a_1)f(a_2)g(a_3)2^{x+2}$	
		$f(a_1)f(a_2)g(a_3)2^{x+1}$	$f(a_1)g(a_2)g(a_3)2^{x+2}$	

# References

- [1] Clayton, David. Personal web page, software list. <a href="http://www-gene.cimr.cam.ac.uk/clayton/software/">http://www-gene.cimr.cam.ac.uk/clayton/software/</a>. Accessed April 1, 2004.
- [2] Schaid DJ. Power and Sample Size for Testing Associations of Haplotypes with Complex Traits. *Annals of Human Genetics* 2005;70:116-130.
- [3] Schaid DJ, Rowland CM, Tines DE, Jacobson RM, Poland GA. Score tests for association between traits and haplotypes when linkage phase is ambiguous. Am J Hum Genet 2002;70:425-34.
- [4] Zaykin DV, Westfall PH, Young SS, Karnoub MA, Wagner MJ, Ehm MG. Testing Association of Statistically Inferred Haplotypes with Discreet and Continuous Traits in Samples of Unrelated Individuals. Human Heredity 2002;53:79-91.
- [5] Harrell, FE. Regression Modeling Strategies. New York: Springer-Verlag; 2001.
- [6] Besag J, Clifford P. Sequential Monte Carlo p-Values. Biometrika 1991;78:301-304.
- [7] Lake S, Lyon H, Silverman E, Weiss S, Laird N, Schaid D. Estimation and tests of haplotype-environment interaction when linkage phase is ambiguous. *Human Heredity* 2003;55:56-65.
- [8] Stram D, Pearce C, Bretsky P, Freedman M, Hirschhorn J, Altshuler D, Kolonel L, Henderson B, Thomas D. Modeling and E-M estimation of haplotype-specific relative risks from genotype data for case-control study of unrelated individuals. *Hum Hered* 2003;55:179-190.
- [9] Epstein MP, Satten GA. Inference on haplotype effects in case-control studies using unphased genotype data. Am J Hum Genet 2003;73:1316-1329.
- [10] McCullagh P, Nelder JA. Generalized Linear Models, Second Edition. Boca Raton, FL: Chapman and Hall. 1989:35-36.
- [11] Cheng R, Ma JZ, Elston RC, Li MD. Fine Mapping Functional Sites or Regions from Case-Control Data Using Haplotypes of Multiple Linked SNPs. *Annals of Human Genetics* 2005;69: 102-112.
- [12] Yu Z, Schaid DJ. Sequential haplotype scan methods for association analysis. Gen Epi 2007;31:553-564.
- [13] Mantel N, Haenszel W. Statistical aspects of the analysis of data from retrospective studies of disease. J Nat Cancer Inst 1959;22:719-48.
- [14] Xie R, Stram DO. Asymptotic equivalence between two score tests for haplotype-specific risk in general linear models. *Gen Epi* 2005;29:166-170.
- [15] Lin DY, Zeng D. Likelihood-based inference on haplotype effects in genetic association studies. J Am Stat Assoc 2006;101:473.