

Generating Marker Summary Reports Using the *GeneticsBase*

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1 Introduction

This document demonstrates how to use the *GeneticsBase* (version 0.6.0) package to generate marker summary tables *for studies with a small number of markers*. It is written as a step-by-step tutorial. For additional details on each of the R functions utilized, please see the individual help pages

Note: The textual displays described here are not suitable for large numbers of markers. They are intended for reviewing detailed information on a small number of markers, such as those in candidate gene studies, or a small set of markers achieving a 'quality' or 'significance' cutoff from a larger set.

2 Example

2.1 Prepare phenotype data

The first step is to prepare the phenotype data. It may be in the form of a SAS dataset, SAS export file, comma-delimited text file (CSV), tab-delimited text file (TSV), or Microsoft Excel spreadsheet file (XLS). It should have one row per observation and one column per variable, and must contain a subject identifier variable that can be used to match observations with the corresponding genotype data.

2.2 Prepare genotype data

You also need to store the genetic call data in a file that can be read into R. GeneticsBase packge accepts genotype data in a variety of formats:

- standard pedigree (ped) format.

a2m	apoe					
50103	5010004	5090005	5090004	2	2	1
2	3	4				
50103	5010005	5090005	5090004	2	2	1
1	3	4				
50105	5010049	5090021	5090022	2	2	1
1	4	4				
50105	5010070	5090020	5090019	1	2	1
1	3	4				

- hapmap format : The hapmap .ped format is a variant of the standard pedigree format. A portion of the first two lines of the hapmap file for chromosome 1 are shown below:

```
rs2298011 rs1320571 rs11721 rs4018608 rs6685064 rs604618 ....
1347 14 0 0 1 1 1 3 3 3 1 1 2 2 3 3 3 2 2 3 3 2 .....
```

- Pfizer format: First few lines of an example file in Pfizer's data format are shown below:

```
Locus,Gene Marker,Locus Start,Project,Protocol,Sample ID,Donor ID,Genotype
A1A,C1556G,-1243,P234,103,1028022.1,1028022,G/G
A1B,T127A,20141,P234,103,1028022.1,1028022,A/T
A1B,T5094A,102358,P234,103,1028022.1,1028022,A/T
A1A,C1556G,-1243,P234,103,1035130.1,1035130,G/G
```

- Perlegen format: A portion of first two lines of data in the Perlegen format are shown below:

```
snp_id genotype
753527 rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr.....
752848 hhahaararhhahrrhhahaharhaahhahrhhahhnha.....
```

2.3 Load the genotype and phenotype data

In **GeneticsBase**, both genotype and phenotype data is loaded by a single function *readgenes*. This function *readGenes* has four primary arguments: *gfile*, *gformat*, *file*, and *pformat*. Arguments *gfile* and *pfile* are the names of files containing the genotype and phenotype data (respectively), and arguments *gformat* and *pformat* are the corresponding file formats for the genotype and phenotype data.

Various types of data can be loaded by *readGenes* function. Example files are provided in **data** subdirectory of the **GeneticsBase** package. To access these execute

```
> library(GeneticsBase)
> setwd(file.path(.path.package("GeneticsBase"), "data"))
```

Supported file formats include:

- fbat .ped format

The Alzheimer's example dataset is stored in the Fbat variant of the .ped Pedigree Format. As it does not include phenotype data, we only use the genotype filename and file type arguments:

```
> ALZH <- readGenes(gfile = "ALZH.ped", gformat = "fbat.ped")
```

The CAMP example dataset is from from the 'Childhood Asthma Management Program (CAMP)' and includes both genotype and phenotype information. It can be loaded by:

```
> CAMP <- readGenes(gfile = "CAMP.ped", gformat = "fbat.ped", pfile = "CAMPZ.phe",
+ pformat = "fbat.phe")
```

- HAPMAP .ped format

A subset of the data for the International HapMap project is available in the hapmap example data set. This file can be loaded via:

```
> hapmapchr1 <- readGenes(gfile = "hapmapchr1.ped", gformat = "hapmap")
```

- Pfizer genetics data format

```
> PfizerExample <- readGenes.pfizer("PfizerExample.txt", format = "Listing")
• Perlegen data format

> PerlegenExample <- readGenes("PerlegenExample.txt", gformat = "perlegen")
```

2.4 Reviewing the data

For the purpose of this example, we are going to use CAMP data set, which can be loaded manually as shown in the previous sub-section, or via

```
> library(GeneticsBase)
> data(CAMP)

Reading 8 markers and 2011 subjects from `CAMP.ped' ...
generating 'geneSet' object...

Successfully read the pedigree file `CAMP.ped'.

Number of Markers: 8
Number of Subjects: 2011
Number of Families: 651

Reading 12 vars from `CAMPZ.phe' ... Done.

Number of Phenotype Variables: 12
Number of Observations : 2011
```

Now you can see a brief summary of the data that was loaded by simply entering the name of the object on a line by itself:

```
> CAMP

geneSet object
-----

Number of Markers: 8
Number of Observations: 2011

Sample variables: family, pid, father, mother, sex, affected, zposfevp, zposfvcp, zlog

Genetic data:

  1.1900 1.1667 1.978 2.1391 2.1988 2.109      649.1837 650.1736 650.1908
m709 1/1    1/1    1/1    1/1    1/1    1/1    ... 1/1    1/1    1/1
m654 1/1    1/1    1/1    1/1    2/1    1/1    ... 2/1    <NA>   1/1
m47  1/1    1/2    1/2    1/2    2/2    1/2    ... 2/2    1/1    1/2
p46  2/2    2/2    2/2    2/2    1/2    2/2    ... 1/2    2/2    2/2
p79  2/2    2/1    2/1    2/1    1/1    2/1    ... 1/1    <NA>   2/1
p252 2/2    1/2    1/2    1/2    <NA>   ... 1/2    2/2    1/2
p491 1/1    1/1    1/1    1/1    1/1    1/1    ... 1/1    <NA>   1/1
p523 1/1    1/2    1/2    1/2    1/2    1/2    ... 1/2    1/1    1/2
```

```

650.1675 651.568 651.1725
m709 1/1      1/1      1/1
m654 1/1      2/2      2/1
m47  1/2      2/2      2/2
p46  1/2      1/1      1/2
p79  2/1      1/1      1/1
p252 1/2     2/2      1/2
p491 1/1      1/1      1/1
p523 1/2     1/1      1/2

```

Warning messages:

```

1: geneSet Object has 121 observations. Only first and last 6 displayed
in: .local(object, ...)

```

The phenotype data can be extracted from the CAMP data object using the `sampleInfo` command:

```

> pdata <- sampleInfo(CAMP)
> summary(pdata)

```

family	pid	father	mother
Min. : 1.0	Min. : 1.0	Min. : 0.0	Min. : 0.0
1st Qu.:165.0	1st Qu.: 503.5	1st Qu.: 0.0	1st Qu.: 0.0
Median :327.0	Median :1006.0	Median : 0.0	Median : 0.0
Mean :326.9	Mean :1006.0	Mean : 340.9	Mean : 367.7
3rd Qu.:489.0	3rd Qu.:1508.5	3rd Qu.: 521.0	3rd Qu.: 637.5
Max. :651.0	Max. :2011.0	Max. :2009.0	Max. :2010.0

sex	affected	zposfevp	zposfvcp
Min. :1.000	Min. :0.0000	Min. :-3.234e+00	Min. :-2.880e+00
1st Qu.:1.000	1st Qu.:0.0000	1st Qu.:-6.790e-01	1st Qu.:-6.250e-01
Median :1.000	Median :0.0000	Median : 1.000e-03	Median :-1.600e-02
Mean :1.453	Mean :0.7041	Mean : 1.431e-05	Mean :-3.433e-05
3rd Qu.:2.000	3rd Qu.:2.0000	3rd Qu.: 6.275e-01	3rd Qu.: 6.005e-01
Max. :2.000	Max. :2.0000	Max. : 4.021e+00	Max. : 4.041e+00

...

2.5 Generate the tables

We can generate a variety of summary tables on our genetics data.

- Allele information

```
> alleleSummary(CAMP)
```

Gene	Marker	Position	Group	Allele	Count	Freq	CI-Lower	CI-Upper
ALL	m709	?	ALL	1	3904	0.998	0.996	0.999
			ALL	2	8	0.002	0.001	0.004
m654	?		ALL	1	2491	0.638	0.623	0.654
			ALL	2	1411	0.362	0.346	0.377

m47	?	ALL	1	1417	0.369	0.354	0.384
		ALL	2	2427	0.631	0.616	0.646
p46	?	ALL	1	1557	0.401	0.385	0.416
		ALL	2	2329	0.599	0.584	0.615
p79	?	ALL	1	2407	0.626	0.611	0.642

...

- Genotype information

```
> genotypeSummary(CAMP)
```

Gene	Marker	Position	Group	Genotype	Count	Freq	CI-Lower	CI-Upper	Expected
?	m709	?	ALL	1/1	1948	0.996	0.993	0.998	1948.008
				1/2	8	0.004	0.002	0.007	7.984
				2/2	0	0.000			0.008
				NA	55				
?	m654	?	ALL	1/1	826	0.423	0.401	0.445	795.115
				1/2	839	0.430	0.408	0.452	900.769
				2/2	286	0.147	0.131	0.162	255.115
				NA	60				
?	m47	?	ALL	1/1	276	0.144	0.128	0.159	261.172
				1/2	865	0.450	0.428	0.472	894.656
				2/2	781	0.406	0.384	0.428	766.172
				NA	55				

...

- Marker information
- Linkage disequilibrium, text view

```
> ld <- LD(CAMP)
```

```
> ld
```

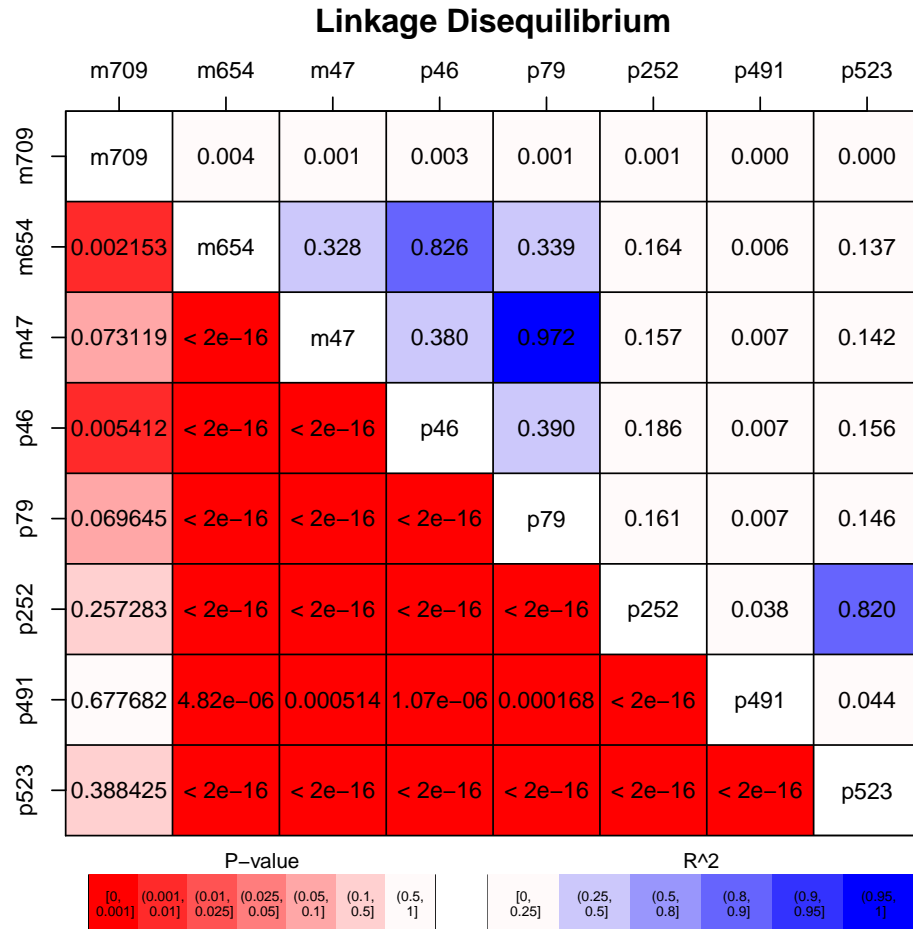
```
-----
Pairwise LD
-----
```

	m709	m654	m47	p46	p79	p252	p491	p523
m709 D	0.001	-0.001	0.001	-0.001	0.000	0.000	0.000	0.000
m709 D'	1.000	0.997	1.000	0.998	0.972	1.000	0.858	
m709 Corr.	0.061	-0.035	0.056	-0.035	-0.025	-0.005	-0.019	
m709 R^2	0.004	0.001	0.003	0.001	0.001	0.000	0.000	
LD X^2	9.000	3.000	7.000	3.000	1.000	0.000	0.000	
P-value	0.00215	0.0731	0.00541	0.0696	0.257	0.678	0.388	
m709 LOD	2.044	0.697	1.680	0.715	0.279	0.038	0.162	
m709 n	2011	2011	2011	2011	2011	2011	2011	
m654 D		-0.133	0.214	-0.135	-0.081	-0.004	-0.071	

...

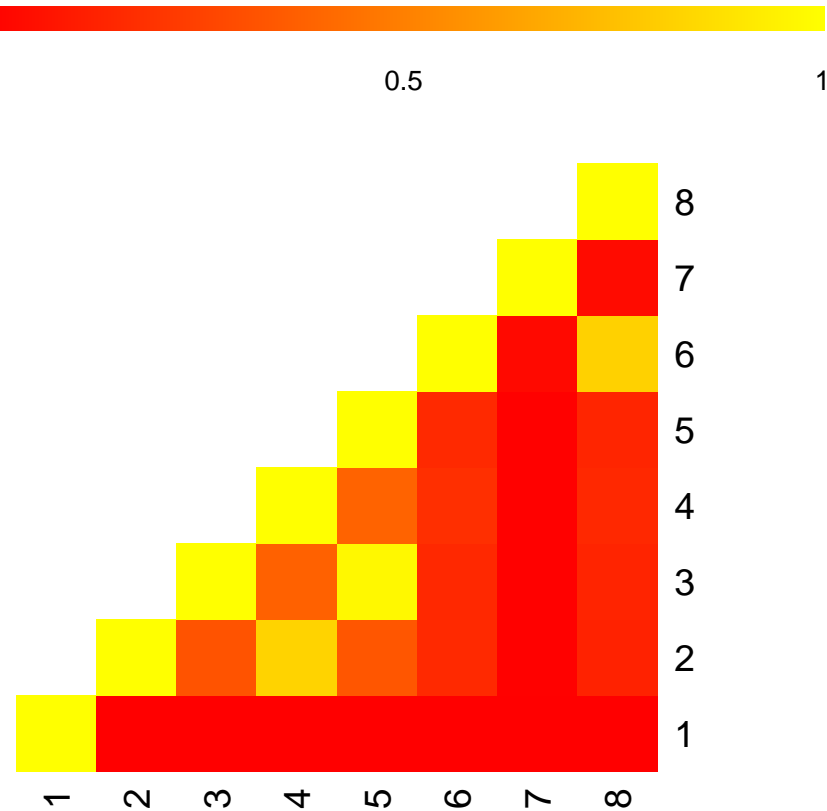
- Linkage disequilibrium, matrix plot

> `plot(ld)`



- Linkage disequilibrium, graphical view using LDView

```
> r2 <- t(ld@"R^2")
> diag(r2) <- 1
> LDView(r2, labRow = markerNames(CAMP))
```



3 Generating tables for inclusion in reports

To make it simple to include the summary tables in written reports, they can be written to files in a variety of formats, including plain text, html, and LaTeX.

3.1 Plain text files

```
> aS <- alleleSummary(CAMP)
> txt(aS, file = "CAMP_alleleSummary.txt")
```

3.2 LaTeX files

```
> aS <- alleleSummary(CAMP)
> latex(aS)
```

	Gene	Marker	Position	Group	Allele	Count	Freq	CI-Lower	CI-Upper
1	ALL	m709	?	ALL	1	3904	0.998	0.996	0.999
2				ALL	2	8	0.002	0.001	0.004
3									
4		m654	?	ALL	1	2491	0.638	0.623	0.654
5				ALL	2	1411	0.362	0.346	0.377
6									
7		m47	?	ALL	1	1417	0.369	0.354	0.384
8				ALL	2	2427	0.631	0.616	0.646
9									
10		p46	?	ALL	1	1557	0.401	0.385	0.416
11				ALL	2	2329	0.599	0.584	0.615
12									
13		p79	?	ALL	1	2407	0.626	0.611	0.642
14				ALL	2	1435	0.374	0.358	0.389
15									
16		p252	?	ALL	1	811	0.222	0.209	0.236
17				ALL	2	2835	0.778	0.764	0.791
18									
19		p491	?	ALL	1	3865	0.989	0.986	0.992
20				ALL	2	43	0.011	0.008	0.014
21									
22		p523	?	ALL	1	3148	0.804	0.791	0.816
23				ALL	2	768	0.196	0.184	0.209
24									

```
> gs <- genotypeSummary(CAMP[-2, ])
```

```
> latex(gs)
```

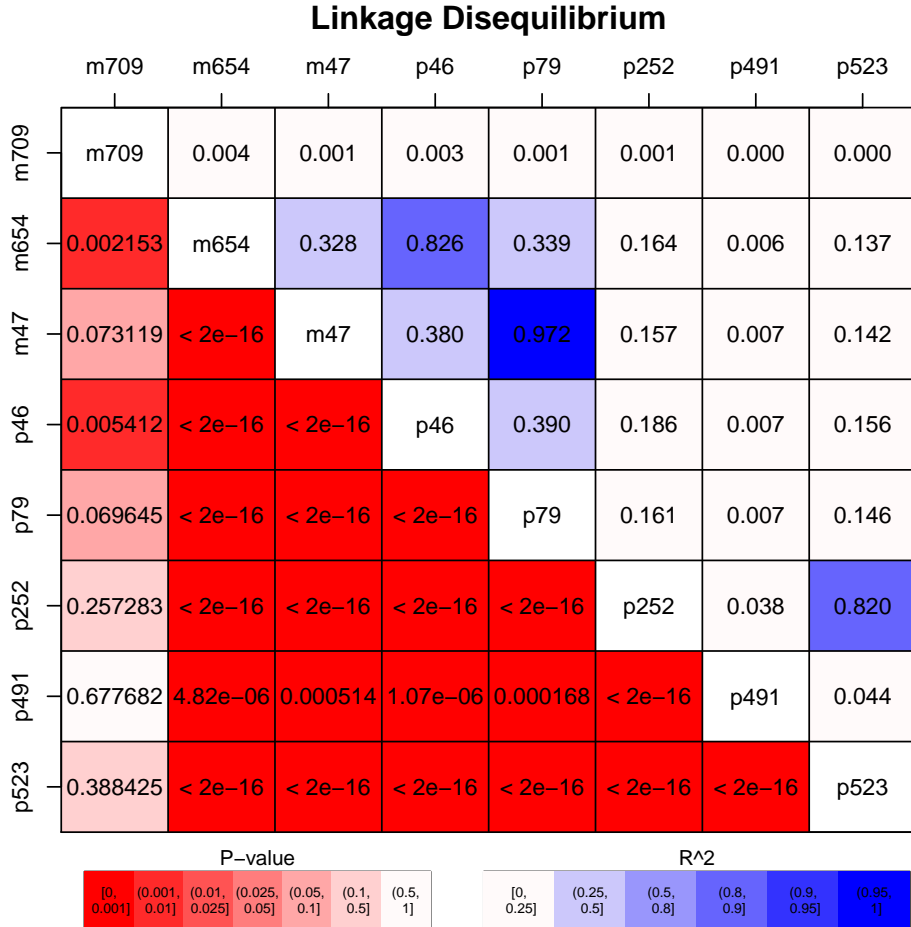
	Gene	Marker	Position	Group	Genotype	Count	Freq	CI-Lower	CI-Upper	Expected	Obs-Exp	HWE X\^2	P-value
1	?	m709	?	ALL	1/1	1948	0.996	0.993	0.998	1948.008	-0.008	0.008	1
2					1/2	8	0.004	0.002	0.007	7.984	0.016		
3					2/2	0	0.000			0.008	-0.008		
4					NA	55							
5													
6	?	m47	?	ALL	1/1	276	0.144	0.128	0.159	261.172	14.828	2.112	0.142
7					1/2	865	0.450	0.428	0.472	894.656	-29.656		
8					2/2	781	0.406	0.384	0.428	766.172	14.828		
9					NA	89							
10	?	p46	?	ALL	1/1	328	0.169	0.152	0.186	311.921	16.079	2.308	0.129
11					1/2	901	0.464	0.442	0.486	933.158	-32.158		
12					2/2	714	0.367	0.346	0.389	697.921	16.079		
13					NA	68							
14													
15	?	p79	?	ALL	1/1	774	0.403	0.381	0.425	753.989	20.011	3.807	0.053
16					1/2	859	0.447	0.425	0.470	899.023	-40.023		
17					2/2	288	0.150	0.134	0.166	267.989	20.011		
18					NA	90							
19													
20	?	p252	?	ALL	1/1	107	0.059	0.048	0.070	90.198	16.802	5.177	0.0228
21					1/2	597	0.327	0.306	0.349	630.605	-33.605		
22					2/2	1119	0.614	0.591	0.636	1102.198	16.802		
23					NA	188							
24													
25	?	p491	?	ALL	1/1	1911	0.978	0.971	0.984	1911.237	-0.237	0.242	1
26					1/2	43	0.022	0.016	0.029	42.527	0.473		
27					2/2	0	0.000			0.237	-0.237		
28					NA	57							
29													
30	?	p523	?	ALL	1/1	1274	0.651	0.630	0.672	1265.309	8.691	1.552	0.223
31					1/2	600	0.306	0.286	0.327	617.381	-17.381		
32					2/2	84	0.043	0.034	0.052	75.309	8.691		
33					NA	53							
34													
35													

```
> ld <- LD(CAMP)
```

```
> latex(ld)
```

	m709	m654	m47	p46	p79	p252	p491	p523
m709 D	0.001	-0.001	0.001	-0.001	0.000	0.000	0.000	0.000
m709 D'	1.000	0.997	1.000	0.998	0.972	1.000	0.858	
m709 Corr.	0.061	-0.035	0.056	-0.035	-0.025	-0.005	-0.019	
m709 \$R^2\$	0.004	0.001	0.003	0.001	0.001	0.000	0.000	
LD \$\chi^2\$	9	3	7	3	1	0	0	
P-value	0.002153	0.073119	0.005412	0.069645	0.257283	0.677682	0.388425	
m709 LOD	2.044	0.697	1.680	0.715	0.279	0.038	0.162	
m709 n	2011	2011	2011	2011	2011	2011	2011	
m654 D		-0.133	0.214	-0.135	-0.081	-0.004	-0.071	
m654 D'		0.992	0.978	0.997	0.994	1.000	0.994	
m654 Corr.		-0.572	0.909	-0.582	-0.405	-0.080	-0.370	
m654 \$R^2\$		0.328	0.826	0.339	0.164	0.006	0.137	
LD \$\chi^2\$		965	2727	1040	468	20	416	
P-value		< 2e-16	< 2e-16	< 2e-16	< 2e-16	4.82e-06	< 2e-16	
m654 LOD		209.756	592.178	226.003	101.725	4.540	90.541	
m654 n		2011	2011	2011	2011	2011	2011	
m47 D			-0.146	0.230	-0.080	-0.004	-0.072	
m47 D'			0.991	0.991	0.966	0.998	0.987	
m47 Corr.			-0.617	0.986	-0.396	-0.082	-0.376	
m47 \$R^2\$			0.380	0.972	0.157	0.007	0.142	
LD \$\chi^2\$			1138	3462	413	12	395	
P-value			< 2e-16	< 2e-16	< 2e-16	0.000514	< 2e-16	
m47 LOD			247.311	751.833	89.707	2.620	85.947	
m47 n			2011	2011	2011	2011	2011	
p46 D				-0.148	-0.088	-0.004	-0.077	
p46 D'				0.993	0.980	1.000	0.983	
p46 Corr.				-0.624	-0.432	-0.086	-0.395	
p46 \$R^2\$				0.390	0.186	0.007	0.156	
LD \$\chi^2\$				1185	498	23	433	
P-value				< 2e-16	< 2e-16	1.07e-06	< 2e-16	
p46 LOD				257.333	108.232	5.169	94.059	
p46 n				2011	2011	2011	2011	
p79 D					-0.081	-0.004	-0.073	
p79 D'					0.972	0.999	1.000	
p79 Corr.					-0.401	-0.082	-0.382	
p79 \$R^2\$					0.161	0.007	0.146	
LD \$\chi^2\$					432	14	430	
P-value					< 2e-16	0.000168	< 2e-16	
p79 LOD					93.840	3.073	93.524	
p79 n					2011	2011	2011	
p252 D						0.008	0.148	
p252 D'						1.000	0.990	
p252 Corr.						0.196	0.905	
p252 \$R^2\$						0.038	0.820	
LD \$\chi^2\$						77	2088	
P-value						< 2e-16	< 2e-16	
p252 LOD						16.742	453.461	
p252 n						2011	2011	
p491 D							0.009	
p491 D'							1.000	
p491 Corr.							0.210	
p491 \$R^2\$							0.044	
LD \$\chi^2\$							88	
P-value							< 2e-16	
p491 LOD							19.116	
p491 n							2011	
p523 D								
p523 D'								
p523 Corr.								
p523 \$R^2\$								
LD \$\chi^2\$								
P-value								
p523 LOD								
p523 n								

```
> plot(1d)
```



3.3 HTML files

3.4 Graphics files

As usual, plots can be generated in any format R supports.

We can also output everything all at once to a set of files, encoded as plain text (`format="print"`), `html` (`format="html"`), or `LaTeX` (`format="latex"`):

```
> PGtables(CAMP, filename = "CAMP", sep = "_", format = "html")

Creating CAMP_alleleSummary.html ...
Creating CAMP_genotypeSummary.html ...
Creating CAMP_LD.html ...
Creating CAMP_LD.pdf ...
Done.
```

which creates a set of html and a PDF files in the current directory.

Figure 1: HTML allele summary table

	Gene	Marker	Position	Group	Allele	Count	Freq	CI-Lower	CI-Upper
1	ALL	m709	?	ALL	1	2534	0.998	0.997	1.000
2				ALL	2	4	0.002	0.000	0.003
3				ALL	NA	0			
4									
5		m654	?	ALL	1	1630	0.647	0.629	0.666
6				ALL	2	888	0.353	0.334	0.371
7				ALL	NA	0			
8									
9		m47	?	ALL	1	924	0.371	0.352	0.390
10				ALL	2	1564	0.629	0.610	0.648
11				ALL	NA	0			
12									
13		p46	?	ALL	1	990	0.395	0.376	0.414
14				ALL	2	1516	0.605	0.586	0.624
15				ALL	NA	0			
16									
17		p79	?	ALL	1	1556	0.625	0.607	0.644
18				ALL	2	932	0.375	0.356	0.393
19				ALL	NA	0			
20									
21		p252	?	ALL	1	546	0.231	0.214	0.247
22				ALL	2	1822	0.769	0.753	0.786
23				ALL	NA	0			
24									
25		p491	?	ALL	1	2499	0.989	0.985	0.993
26				ALL	2	27	0.011	0.007	0.015
27				ALL	NA	0			
28									
29		p523	?	ALL	1	2031	0.799	0.783	0.814
30				ALL	2	511	0.201	0.186	0.217
31				ALL	NA	0			
32									

Confidence intervals width is 95%, computed using the exact quantiles for the binomial distribution.

Figure 2: HTML genotype summary table

Mozilla Firefox

file:///localhost/Users/warnes/src/r-genetics/GeneticsBase, ▾

Gene	Marker	Position	Group	Genotype	Count	Freq	CI-Lower	CI-Upper	Expected	Obs-Exp	HWE X^2	P-value
1	?	m709	?	ALL	1/1	1265	0.997	0.994	0.999	1265.003	-0.003	0.003
2					1/2	4	0.003	0.001	0.006	3.994	0.006	
3					2/2	0	0.000			0.003	-0.003	
4					NA	0						
5												
6	?	m654	?	ALL	1/1	536	0.426	0.399	0.453	527.581	8.419	1.080
7					1/2	558	0.443	0.416	0.471	574.837	-16.837	
8					2/2	165	0.131	0.113	0.150	156.581	8.419	
9					NA	0						
10												
11	?	m47	?	ALL	1/1	171	0.137	0.119	0.157	171.579	-0.579	0.005
12					1/2	582	0.468	0.441	0.496	580.842	1.158	
13					2/2	491	0.395	0.367	0.422	491.579	-0.579	
14					NA	0						
15												
16	?	p46	?	ALL	1/1	197	0.157	0.137	0.178	195.551	1.449	0.029
17					1/2	596	0.476	0.448	0.504	598.899	-2.899	
18					2/2	460	0.367	0.341	0.394	458.551	1.449	
19					NA	0						
20												
21	?	p79	?	ALL	1/1	488	0.392	0.365	0.420	486.563	1.437	0.030
22					1/2	580	0.466	0.439	0.494	582.875	-2.875	
23					2/2	176	0.141	0.122	0.161	174.563	1.437	
24					NA	0						
25												
26	?	p252	?	ALL	1/1	68	0.057	0.045	0.071	62.947	5.053	0.685
27					1/2	410	0.346	0.319	0.373	420.106	-10.106	
28					2/2	706	0.596	0.568	0.624	700.947	5.053	
29					NA	0						
30												
31	?	p491	?	ALL	1/1	1236	0.979	0.970	0.987	1236.144	-0.144	0.147
32					1/2	27	0.021	0.013	0.030	26.711	0.289	
33					2/2	0	0.000			0.144	-0.144	
34					NA	0						

Done Adblock Now: Partly Cloudy, 53° F Fri: 52° F Sat: 52° F

Figure 3: HTML linkage disequilibrium table

Mozilla Firefox

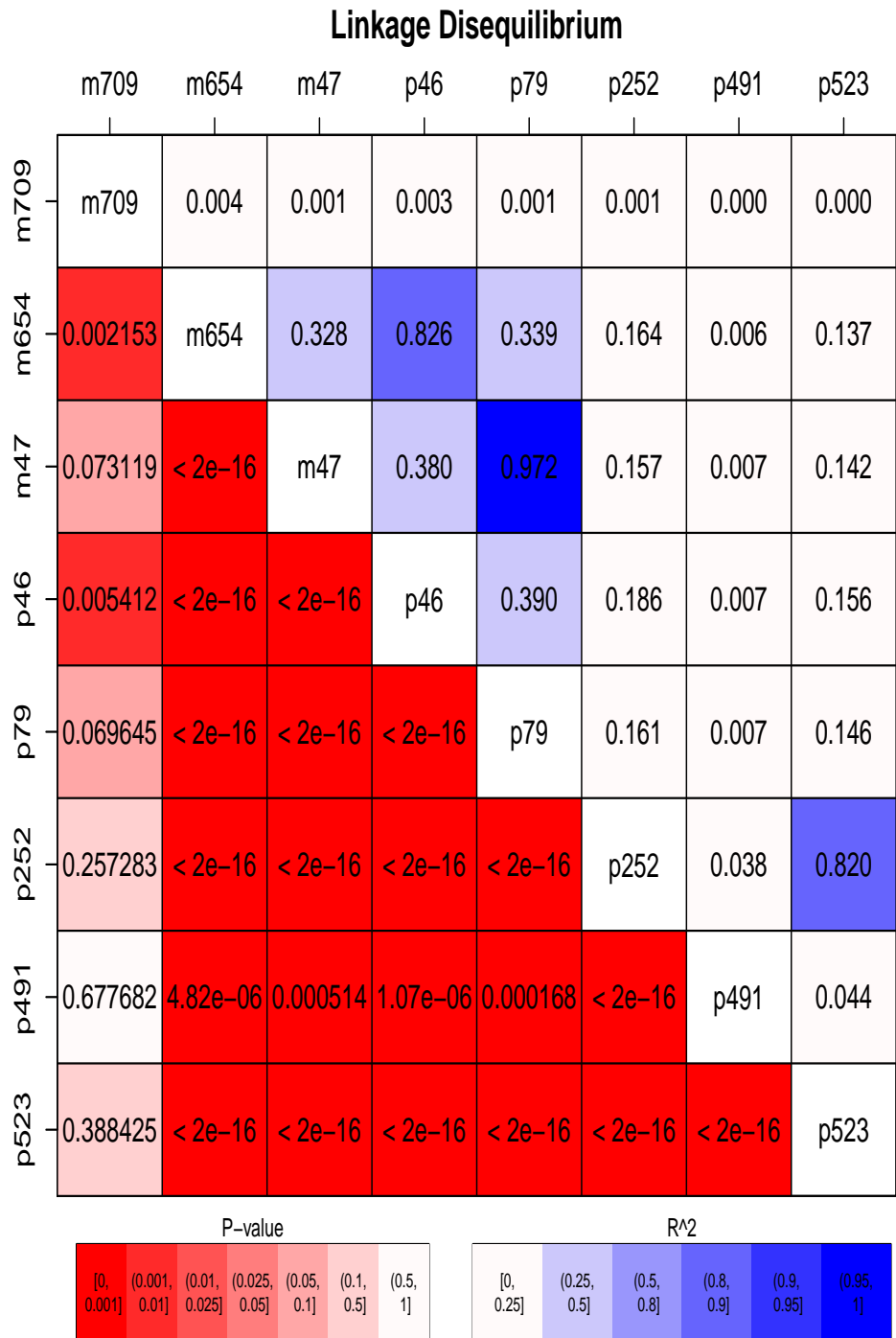
file:///localhost/Users/warnes/src/r-genetics/GeneticsBase, ▾

html to pdf

	m709	m654	m47	p46	p79	p252	p491	p523
m709 D	0.001	-0.001	0.001	-0.001	0.000	0.000	0.000	0.000
m709 D'	1.000	0.999	0.999	0.999	0.047	1.000	0.151	
m709 Corr.	0.055	-0.031	0.050	-0.031	0.004	-0.004	0.012	
m709 R^2	0.003	0.001	0.002	0.001	0.000	0.000	0.000	
LD X^2	4	2	3	2	0	0	0	
P-value	0.039830	0.127371	0.069309	0.125046	0.895257	0.767100	0.702909	
m709 LOD	0.917	0.505	0.716	0.511	0.004	0.019	0.032	
m709 n	1303	1303	1303	1303	1303	1303	1303	
m654 D		-0.130	0.212	-0.132	-0.082	-0.004	-0.070	
m654 D'		0.996	0.978	0.996	0.991	1.000	0.990	
m654 Corr.		-0.563	0.905	-0.570	-0.405	-0.077	-0.366	
m654 R^2		0.317	0.818	0.325	0.164	0.006	0.134	
LD X^2		588	1698	619	300	12	255	
P-value		< 2e-16	< 2e-16	< 2e-16	< 2e-16	0.000391	< 2e-16	
m654 LOD		127.716	368.786	134.516	65.332	2.730	55.408	
m654 n		1303	1303	1303	1303	1303	1303	
m47 D			-0.143	0.231	-0.083	-0.004	-0.075	
m47 D'			0.992	0.991	0.966	0.998	1.000	
m47 Corr.			-0.608	0.985	-0.407	-0.081	-0.388	
m47 R^2			0.370	0.970	0.166	0.007	0.150	
LD X^2			699	2213	268	7	273	
P-value			< 2e-16	< 2e-16	< 2e-16	0.007251	< 2e-16	
m47 LOD			151.847	480.561	58.250	1.566	59.344	
m47 n			1303	1303	1303	1303	1303	
p46 D				-0.145	-0.090	-0.004	-0.078	
p46 D'				0.993	0.985	1.000	0.991	
p46 Corr.				-0.615	-0.439	-0.085	-0.400	
p46 R^2				0.379	0.192	0.007	0.160	
LD X^2				719	335	14	282	
P-value				< 2e-16	< 2e-16	0.000113	< 2e-16	
p46 LOD				156.342	72.789	3.237	61.413	
p46 n				1303	1303	1303	1303	
n79 D					-0.083	-0.004	-0.075	

Done Adblock Now: Partly Cloudy, 53° F Fri: 52° F Sat: 52° F

Figure 4: Linkage disequilibrium plot



4 Subsetting by Group

The `alleleSummary` and `genotypeSummary` functions also allow you to create tables which show the summary information separated out by a grouping variable, which must be discrete “factor” variables (such as `Sex`).

To accomplish this, add the argument `by=Sex` to the function call. For example:

```
> alleleSummary(CAMP, by = "sex")
```

Gene	Marker	Position	Group	Allele	Count	Freq	CI-Lower	CI-Upper
ALL	m709	?	1	1	3904	0.998	0.996	0.999
				2	8	0.002	0.001	0.004
	m654	?	1	1	3904	0.998	0.996	0.999
				2	8	0.002	0.001	0.004
m47	m654	?	1	1	2491	0.638	0.623	0.654
				2	1411	0.362	0.346	0.377
	p46	?	1	1	2491	0.638	0.623	0.654
				2	1411	0.362	0.346	0.377
p46	m47	?	1	1	1417	0.369	0.354	0.384
				2	2427	0.631	0.616	0.646
	p79	?	1	1	1417	0.369	0.354	0.384
				2	2427	0.631	0.616	0.646
p79	p46	?	1	1	1557	0.401	0.385	0.416
				2	2329	0.599	0.584	0.615
	p79	?	1	1	1557	0.401	0.385	0.416
				2	2329	0.599	0.584	0.615
p252	p79	?	1	1	2407	0.626	0.611	0.642
				2	1435	0.374	0.358	0.389
	p252	?	1	1	2407	0.626	0.611	0.642
				2	1435	0.374	0.358	0.389
p252	p252	?	1	1	811	0.222	0.209	0.236
				2	2835	0.778	0.764	0.791
	p491	?	1	1	811	0.222	0.209	0.236
				2	2835	0.778	0.764	0.791
p491	p523	?	1	1	3865	0.989	0.986	0.992
				2	43	0.011	0.008	0.014
	p523	?	1	1	3865	0.989	0.986	0.992
				2	43	0.011	0.008	0.014
p523		?	1	1	3148	0.804	0.791	0.816

1	2	768	0.196	0.184	0.209
2	1	3148	0.804	0.791	0.816
2	2	768	0.196	0.184	0.209

Footer:

Confidence intervals width is 95%, computed using
the exact quantiles for the binomial
distribution.

This will display a table within a separate block within each marker for each level of the variable **Sex**.
To control whether the summary table for entire data in addition to individual factor levels, add `includeOverall=TRUE` or `includeOverall=FALSE` (the default) as appropriate.

A Example R script

```
> library(GeneticsBase)
> data(CAMP)
> PGtables(CAMP, filename = "test", format = "html")
> PGtables(CAMP, filename = "test", format = "latex")
```

References

- Warnes GR. “The Genetics Package: Utilities for handling genetic data” *R News*, Volume 3, Issue 1, June 2003.
- Warnes GR. “genetics”, a package for handling marker-based genetic data within the open-source statistical package “R”. The package includes function to compute allele frequencies, use genetic markers in statistical models, estimate disequilibrium, and test for departure from Hardy-Weinberg equilibrium.
<http://cran.us.r-project.org/src/contrib/PACKAGES.html#genetics>, 2002-