# $\begin{array}{c} {\rm PLINK}~({\rm v1.04}) \\ {\rm A~whole\text{-}genome~association~toolset} \end{array}$

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# Contents

1	$\operatorname{Get}$	ting started with PLINK	L
	1.1	Citing PLINK	1
	1.2	Reporting problems, bugs and questions	1
	1.3	Download	2
	1.4	Development version source code	2
	1.5	General installation notes	3
	1.6	Windows/MS-DOS notes	3
	1.7	UNIX/Linux notes	4
	1.8	Source code compilation	4
	1.9	Running PLINK from the command line	ô
	1.10	Viewing PLINK output files	7
<b>2</b>	Bas	ic usage / data formats	3
	2.1	Running PLINK	3
	2.2	PED files	9
		2.2.1 Different PED file formats: missing fields	1
	2.3	MAP files	2
		2.3.1 Chromosome codes	2
		2.3.2 Allele codes	3
	2.4	Transposed filesets	4
	2.5	Long-format filesets	4
	2.6	Binary PED files	6
	2.7	Alternate phenotype files	7
		2.7.1 Creating a new binary phenotype automatically	3
	2.8	Covariate files	3
	2.9	Cluster files	9
	2.10	Set files	9
3	Dat	a management tools 2	1
	3.1	Recode and reorder a sample	1
		3.1.1 Transposed genotype files	2
		3.1.2 Additive and dominance components	2
		3.1.3 Listing by genotype	4
	3.2	Write SNP list files	4
	3.3	Update SNP positions	5
	3.4	Write covariate files	5
	3.5	Write cluster files	
	3.6	Flip DNA strand for SNPs	
	3.7	Using LD to identify incorrect strand assignment in a subset of the sample	
	3.8	Merge two filesets	

	3.9	Merge multiple filesets	29
	3.10	Extract a subset of SNPs: command line options	30
		3.10.1 Based on a single chromosome (chr)	30
		3.10.2 Based on a range of SNPs (from andto)	30
		3.10.3 Based on single SNP (and window) (snp andwindow)	30
		3.10.4 Based on multiple SNPs and ranges (snps)	30
		3.10.5 Based on physical position (from-kb, etc)	31
		3.10.6 Based on a set file (gene)	31
	3.11	Extract a subset of SNPs: file-list options	31
			32
			32
	3.14	Extract a subset of individuals	34
	3.15	Remove a subset of individuals	34
			34
	3.17	Create a SET file based on a list of ranges	35
		<u> </u>	
4	Sun	<b>v</b>	36
	4.1		36
	4.2	Obligatory missing genotypes	37
	4.3		39
	4.4		40
	4.5	1 11	40
	4.6	v 0 1	42
	4.7	1 0	43
	4.8	Linkage disequilibrium based SNP pruning	43
	4.9		44
	4.10	Sex check	45
	4.11	Pedigree errors	45
_	T1		47
5	inci		
			47
	F 1		47
	5.1		47
	5.2		48
	5.3		48
	5.4	V 0 1	48
	5.5	Mendel error rate	49
6	Pon	pulation stratification	50
Ū	6.1		50
	6.2		52
	6.3	<u> </u>	53
	6.4		56
	6.5	v	57
	6.6	<b>~</b> .	57
	0.0	Outlier detection diagnostics	01
7	IBS	/IBD estimation	<b>5</b> 9
	7.1	·	59
	7.2	Inbreeding coefficients	60
	7.2 7.3	· · · · · · · · · · · · · · · · · · ·	60 61
		Runs of homozygosity	

	7.4.2	Remove very closely related individuals
	7.4.3	Prune the set of SNPs
	$7.4.4 \\ 7.4.5$	Detecting shared segments (extended, shared haplotypes)
8 A		on analysis
		case/control association test
_		
		's Exact test (allelic association)
		nate / full model association tests
		fied analyses
		g for heterogeneous association
		ing's T(2) multilocus association test
8	.7 Quant	itative trait association
8	.8 Genot	ype means for quantitative traits
8	.9 Quant	itative trait interaction (GxE)
8	.10 Linear	and logistic models
	8.10.1	Basic usage
		Covariates and interactions
		Flexibly specifying the model
		Flexibly specifying joint tests
		Multicollinearity
8		used tests
		tment for multiple testing: Bonferroni, Sidak, FDR, etc
_		
	amily-ba	sed association analysis
		y-based association (TDT)
	-	TDT
		t of origin analysis
		I: family-based association for disease traits
9	.5 QFAM	I: family-based association tests for quantitative traits
10 F	Permutati	ion procedures
	10.0.1	Conceptual overview of permutation procedures
	10.0.2	Label-swapping and gene-dropping
		Adaptive and max(T) permutation
		Computational issues
1		(adaptive) permutation procedure
		ive permutation parameters
	-	[] permutation
	,	dropping permutation
1		
		Basic within family QTDT
		Discordant sibling test
		parenTDT/parenQTDT
		Standard association for singleton, unrelated individuals
1	U.5 Withii	n-cluster permutation
11 N	/ ////////////////////////////////////	ker haplotype tests
		ication of haplotypes to be estimated
		mputed lists of multimarker tests
		ating haplotype frequencies
		g for haplotype-based case/control and quantitative trait association
		type-based TDT association test

	11.6	Imputing multimarker haplotypes	100
	11.7	Tabulating individuals' haplotype phases	101
12	Con	ditional haplotype-based association testing	L02
	12.1	Basic usage for conditional haplotype-based testing	103
	12.2	Specifying the type of test	105
		12.2.1 Testing a specific haplotype	105
		12.2.2 Testing whether SNPs have independent effects	106
		12.2.3 Omnibus test controlling for X	109
	12.3	General specification of haplotype groupings	110
		12.3.1 Manually specifying haplotypes	110
		12.3.2 Manually specifying SNPs	111
	12.4	Covariates and additional SNPs	111
	12.5	General setting of linear constraints	112
13	Pro	xy association	113
		Proxy association: basic usage	113
		13.1.1 Heuristic for selection of proxy SNPs	
		13.1.2 Specifying the type of association test	
	13.2	Refining a single SNP association	
		Automating for multiple references SNPs	
		Providing some degree of robustness to non-random genotyping failure	
11	CNII		
14		P imputation and association testing Basic steps for using PLINK imputation functions	122
	14.1	14.1.1 Strand issues	
	149		
		Combined imputation and association analysis of case/control data	
	14.5	Modifying options for basic imputation/association testing	
	111	14.3.1 Parameters modifying selection of proxies	
		Imputing discrete genotype calls	
	14.5	Verbose output options	120
<b>15</b>		1 91	L <b>2</b> 8
		Basic usage for LD-based clumping	
	15.2	Verbose report	
		15.2.1 Annotation by SNP details and genomic co-ordinates	
		Combining multiple result files (potentially from different SNP panels)	
	15.4	Selecting the single best proxy	132
16	Gen	e reporting tool	<b>134</b>
	16.1	Basic usage	134
	16.2	Other options	135
17	Epis	stasis	136
	_		136
		17.1.1 A faster epistasis option	138
	17.2	Case-only epistasis	
		Gene-based tests of epistasis	

18	R plugin functions	139
	18.1 Basic usage for R plug-ins	. 139
	18.2 Defining the R plug-in function	. 140
	18.3 Example of debugging an R plug-in	. 140
	18.4 Setting up the Rserve package	. 143
19	SNP annotation database lookup	144
	19.1 Basic usage for SNP lookup function	. 144
	19.2 Gene-based SNP lookup	. 146
	19.3 Description of the annotation information	. 147
20	SNP simulation routine	148
	20.1 Basic usage	. 148
	20.2 Resimulating a sample from the same population	. 149
<b>21</b>	SNP scoring routine	151
	21.1 Basic usage	. 151
20	Dana armanian (CNIV) data	150
22	Rare copy number variant (CNV) data 22.1 Basic support for segmental CNV data	153
	22.2 Creating MAP files for CNV data	
	22.3 Loading CNV data files	
	22.4 Filtering of CNV data based on CNV type	
	22.5 Filtering of CNV data based on genomic location	
	22.5.1 Defining overlap for partially overlapping CNVs and regions	
	22.5.2 Filtering by chromosomal co-ordinates	
	22.6 Filtering of CNV data based on frequency	
	22.6.1 Alternative frequency filtering specification	
	22.6.2 Miscellaneous commands frequency filtering commands	
	22.7 Association analysis of segmental CNV data	
	22.8 Association mapping with segmental CNV data	
	22.9 Writing new CNV lists	
	22.9.1 Creating UCSC browser CNV tracks	
	22.10Listing intersected genes and regions	
	22.11Reporting sets of overlapping segmental CNVs	
	22.12Illustration of the different CNV frequency filtering commands	
	22.12.1 Region-based, or locus-based, frequency filtering (default)	
	22.12.2 Alternative frequency filtering approach	
	22.12.3 In summary	. 171
<b>23</b>	Common copy number polymorphism (CNP) data	172
	23.1 Format for common CNVs (generic variant format)	
	23.2 Association models for combined SNP and common CNV data	. 174
24	Resources available for download	175
	24.1 The Phase 2 HapMap as a PLINK fileset	
	24.2 Teaching materials and example dataset	. 175
	24.3 Multimarker test lists	. 176
	24.4 Gene sets	. 177
	24.5 Gene range lists	. 177

25	6 Miscellaneous	178
	25.1 Output modifiers	. 178
	25.2 Analyses with different species	. 178
	25.3 Matrix of pairwise LD (genotype correlation)	
	25.3.1 Filtering the output	
	25.3.2 Obtaining LD values for a specific SNP versus all others	
	25.3.3 Obtaining a matrix of LD values	
	25.3.4 Haplotype-based LD calculations	
	25.4 Known issues	
26	5 FAQ and Hints	181
	26.1 Can I convert my binary PED fileset back into a standard PED/MAP fileset?	. 181
	26.2 To speed up input of a large fileset	
	26.3 Why are no indidividuals included in the analysis?	. 182
	26.4 Why are my results different from an analysis using program X?	
	26.5 How large a file can PLINK handle?	. 182
	26.6 Why does my linear/logistic regression output have all NA's?	. 183
	26.7 What kind of computer do I need to run PLINK?	. 183
	26.8 Can I analyse multiple phenotypes in a single run (e.g. for gene expression datasets)?	. 184
	26.9 How does PLINK handle the X chromosome in association tests?	. 184
	26.10Can/why can't gPLINK perform a particular PLINK command?	. 184
	26.11When I include covariates withlinear orlogistic, what do the p-values mean?	. 185
$\mathbf{A}$	Reference Tables	186
	A.1 Options	. 186
	A.2 Output files (alphabetical listing)	

# Chapter 1

# Getting started with PLINK

This page contains some important information on learning to use PLINK and how to handle any problems you encounter.

We suggest that after downloading PLINK you first try the tutorial. This will familiarize you with the basic PLINK commands.

# 1.1 Citing PLINK

If you use PLINK in any published work, please cite both the software (as an electronic resource/URL) and the manuscript describing the methods.

Package: PLINK (including version number)

Author: Shaun Purcell

URL: http://pngu.mgh.harvard.edu/purcell/plink/
Purcell S, Neale B, Todd-Brown K, Thomas L, Ferreira MAR,

Bender D, Maller J, Sklar P, de Bakker PIW, Daly MJ & Sham PC (2007)
PLINK: a toolset for whole-genome association and population-based

linkage analysis. American Journal of Human Genetics, 81.

# 1.2 Reporting problems, bugs and questions

If you have any problems with PLINK or would like to report a bug, please follow these steps:

#### PLEASE READ THIS SECTION BEFORE E-MAILING!

When an analysis does not report the results you expect, or when PLINK seemingly gives different answers to previous versions or to other software packages, or the last time you ran it, etc, please feel me to e-mail me

plink AT chgr DOT mgh DOT harvard DOT edu

**but** also please consider the following before doing so:

- Please first check the Frequently Asked Questions list to see if your question has already been answered
- Please check the LOG file, it often contains important information. For example, did it filter out some individuals based on genotyping rate or missing phenotype/sex information which you were not expecting?
- Please check the format of your data: is it plain text? does each file have the correct number of rows, etc. Are the missing value codes appropriate?

- Please recheck the web-documentation: sometimes the syntax of an option may change.
- If the above steps do not resolve your problem, then please e-mail me plink AT chgr dot mgh dot harvard dot edu (this is different from the mailing list i.e. your e-mail will only be sent to me, not the whole list). The more specific your e-mail, the easier it will be for me to diagnose any problem or error. Please include:
  - The whole LOG file(s)
  - The type of machine you were using
  - Ideally, please try to make some reduced dataset that replicates the problem that you are able to send to me in a ZIP file, so that I will be able to recreate the problem; any data sent to me for these purposes will be immediately deleted after I have resolved the problem.

**HINT** The more of the above steps you follow, the more likely you are to receive a timely, useful response! If you haven't heard within a week or so, please feel free to send a reminder e-mail...

**IMPORTANT** I am willing and able to advise on the use of specific features implemented in PLINK: to diagnose whether they are working as intended and to give a generic description of a procedure or method, if it is unclear from the web documentation. I'm afraid I will not necessarily be able to give specific advice on any one particular dataset, why you should use one method over another, what it all means, etc...

This page contains some important information regarding how to set up and use PLINK. Individuals familiar with using command line programs can probably skip most of this page.

#### 1.3 Download

**PLINK** is now available for free download. Below are links to ZIP files containing binaries compilied on various platforms as well as the C/C++ source code. Linux/Unix users should download the source code and compile (see notes below).

These downloads also contain a version of gPLINK, an (optional) GUI for PLINK. Please see these pages for instructions on use of gPLINK.

**Remember** This release is considered a *stable* release, although please remember that, just like any other non-trivial computer program, we cannot guarantee that it does not contain bugs...

```
        Platform File Version

        Linux (x86.64)
        plink-1.04-x86.64.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-x86.64.zip
        v1.04

        Linux (x86.64)
        plink-1.04-i686.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-i686.zip
        v1.04

        MS-DOS
        plink-1.04-dos.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-acs.zip
        v1.04

        Apple Mac (PPC)
        plink-1.04-mac.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-acs.zip
        v1.04

        Apple Mac (Intel)
        plink-1.04-mac.intel.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-acc.zip
        v1.04

        C/C++ source (.zip)
        plink-1.04-src.zip http://pngu.mgh.harvard.edu/~purcell/dist/plink-1.04-src.zip
        v1.04
```

One more thing... If you download PLINK please either join the very low-volume e-mail list (link from Introduction page) or drop an e-mail to plink AT chgr dot mgh dot harvard dot edu letting me know you've downloaded a copy.

For old versions of PLINK please visit the archive.

# 1.4 Development version source code

You can download the very latest development source code in this ZIP file http://pngu.mgh.harvard.edu/~purcell/dist/plink-latest.zip. This is really, strongly not recommended for most users. The code posted here could change on a daily basis and is not versioned.

The principle reason for including the source code here is to allow access for specific users to specific, new features. These features are described here.

#### 1.5 General installation notes

The PLINK executable file should be placed in either the current working directory or somewhere in the command path. This means that typing

```
plink
or
./plink
```

at the command line prompt will run PLINK, no matter which current directory you happen to be in. PLINK is a command line program – clicking on an icon with the mouse will get you nowhere.

Below, on this page, is a general overview of how to use the command line to run PLINK. The next sections give details about how to install PLINK on different platforms.

# 1.6 Windows/MS-DOS notes

Unzipping the downloaded ZIP file should reveal a single executable program plink.exe. The Windows/MS-DOS version of PLINK is also a command line program, and is run by typing

```
plink options...
```

not by clicking on the icon with the mouse. Open a DOS windows by selecting "Command Prompt" from the start menu, or entering "command" or "cmd" in the "Run..." option of the start menu.

The folders c:\windows\ or c:\winnt\ are typically in the path, so these are good places to copy the file plink.exe to. You can copy the plink.exe file using Windows, as you would copy-and-paste any file (e.g. using the right-button menu or the keyboard shortcuts control-C (paste) and control-V (paste).

Alternatively, if you know that you will only ever run PLINK on files in a single folder, then you can paste plink.exe into that folder, e.g. C:\work\genetics\. The disadvantage of this approach is that PLINK will not be available from the command line if you are in a folder other than this one.

Once you have copied plink.exe to the correct location, you can test whether or not PLINK is available (i.e. in your command path) by simply typing

#### plink

at the command line. You should see something like the following message:

Microsoft Windows XP [Version 5.1.2600]

ERROR: No file [ plink.ped ] exists.

Options in effect:

```
(C) Copyright 1985-2001 Microsoft Corp.
C:\>plink
0------
               - 1
                   v0.991
                              27/Jul/2006
       PI.TNK!
|-----|
  (C) 2006 Shaun Purcell, GNU General Public License, v2
     http://pngu.mgh.harvard.edu/purcell/plink/
@------@
Web-based version check ( --noweb to skip )
Connecting to web... OK, v0.991 is current
*** Pre-Release Testing Version ***
Writing this text to log file [ plink.log ]
Analysis started: Fri Jul 28 10:07:57 2006
```

Do not worry about this *error message* – normally you would specify your own PED/MAP file names to analyse (i.e. the default input filename is plink.ped).

Please ask your system administrator for help if you do not understand this.

HINT In MS-DOS, you can to increase the width of the window to avoid output lines wrapping around and being hard to read. To do this under Windows XP DOS: right click on the top title/menu bar of the window and select Properties / Layout / Window Size / Width – increse the width value to a larger value (e.g. 120, or as large as possible without the window getting too big to fit on your screen!).

## 1.7 UNIX/Linux notes

If you are not familiar with the concept of the path variable, ask your system administrator to help. In a UNIX/Linux environment, this would mean either copying the PLINK executable to a folder such as

```
/usr/local/bin/
```

or

~bin/

assuming these directories exist and are in the path. To see which directories are in the path, typing

\$PATH

at the command prompt will often work. To create a directory, say called bin in your home directory and add it to the path, try

```
mkdir /bin
export PATH=$PATH:/bin/
```

although this will depend on which shell you are using. Some shells do not include the current directory in the path: in this case, you might need to prefix all PLINK commands with the characters ./, e.g.

```
./plink --file mydata --assoc
```

# 1.8 Source code compilation

PLINK is also distributed as C/C++ source code, which you can compile for your particular system using any standard C/C++ compile. Download the .zip or .tar.gz files and perform the following steps:

```
or
unzip plink-0.99s-src.zip
```

or use a graphical tool such as WinZip to extract the contents of the archive. This should create a directory called

```
plink-0.99s-src
```

(the exact version number might be different, of course). On the command line, move to that directory and simply type make :

```
cd plink-0.99s
```

You will need a C/C++ compiler installed on your system for the next step. Linux distributions will include gcc/g++ by default. Ask your system administrator about installing a C/C++ compiler if you do not have one already (Windows, MS-DOS users).

**Hint** PLINK has not been exhaustively tested on different compilers. We sugest you use a recent download of MinGW for Windows, or at least gcc 4.1.

WARNING We suggest using the most recent stable release of the compiler available on your platform to avoid compilation problems. For most platforms this means gcc 4.2 as of writing this. Some issues with specific older compiler and specific platforms have been detected, e.g. gcc 3.3.3 on a SGI Altix 3700 system.

Use a standard text editor such as emacs, pico or WordPad to edit the Makefile to suit your particular platform: the top of the Makefile should look like this:

```
______
#
#
   Makefile for PLINK
#
#
   Supported platforms
#
       Unix / Linux
                                 LINUX
#
       Windows
                                 WIN
#
       Mac
                                 MAC
#
                                 SOLARIS
       Solaris
#
#
   Compilation options
#
       R plugins
                                 WITH_R_PLUGINS
#
       Web-based version check
                                 WITH_WEBCHECK
#
       Ensure 32-bit binary
                                 FORCE_32BIT
#
# Set this variable to either UNIX, MAC or WIN
SYS = UNIX
# Leave blank after "=" to disable; put "= 1" to enable
WITH_R_PLUGINS = 1
WITH_WEBCHECK = 1
FORCE_32BIT =
# Put C++ compiler here; Windows has it's own specific version
CXX_UNIX = g++
CXX_WIN = c:\bin\mingw\bin\mingw32-g++.exe
# Do not edit below this line
```

The steps to edit this:

- Change the SYS variable to your platform, e.g. WIN for Windows
- For the next three options, put either a 1 or leave blank to turn on or off these options, respectively.
  - WITH\_R\_PLUGINS This enables support for R plugins using Rserve as described here. Currently
    this only works for Unix-based machines.
  - If you want to disable the web-based version check option (not recommended) or if compilation fails with this on, you might try removing the 1
  - When compiling on a 64-bit machine, this option can force (when set) a 32 bit binary (assumes all necessary libraries, etc) are in place

• Edit the CXX\_\* variable to point to the C/C++ compiler you wish to use

To pass any extra commands to the compiler (e.g. location of libraries, etc), you can of course edit CXX\_FLAGS (even though it is below the do not edit line...)

You should then just type

make

and PLINK should (hopefully) start compiling. You should use GNU version, which is sometimes called gmake on some platforms (e.g. FreeBSD). It is also possible that you have installed make but it is not in your path and/or your version of make.exe is called something slightly different, in which case use the full path, e.g. change the following to suit your system:

```
c:\mingw\bin\mingw32-make
```

**NOTE** Often problems in compilation will reflect system-specific / compiler-specific problems: unfortunately, we are not able to give detailed advice on how to do this. If things do not work and you are unsure, you will need to enlist the help of your systems/IT department.

You should see something like the following output (abbreviated)

```
g++ -03 -I. -DUNIX -static -c plink.cpp
g++ -03 -I. -DUNIX -static -c options.cpp
g++ -03 -I. -DUNIX -static -c input.cpp
...
g++ -03 -static -o plink plink.o options.o input.o binput.o
helper.o genome.o snpfilter.o indfilter.o locus.o multi.o
regress.o crandom.o cluster.o output.o informative.o affpair.o
assoc.o bins.o epi.o phase.o trio.o sharing.o genepi.o sets.o
perm.o mh.o genedrop.o gxe.o merge.o hotel.o multiple.o
```

After a minute or so, this will have created an executable binary file called plink (or plink.exe for Windows/MSDOS users).

# 1.9 Running PLINK from the command line

A typical session might involve running several commands, e.g. to produce summary statistics on missing data, to exclude some SNPs based on these results, to run an association analysis. Each command involves a separate instantiation of plink – note that PLINK does not remember any parameter settings between different runs or store any other information. In otherwords, if you want to perform two association tests with different PED files, but only including SNPs that are above a certain minor allele frequency in both runs, you would use the following:

```
plink --ped file1.ped --map file1.map --maf 0.05 --assoc plink --ped file2.ped --map file2.map --maf 0.05 --assoc In otherwords, the following sequence would not work: plink --ped file1.ped --map file1.map --maf 0.05 plink --ped file1.ped --map file1.map --assoc MAF returns to default 0.01 plink --ped file2.ped --map file2.map --assoc As above
```

## 1.10 Viewing PLINK output files

UPDATE We are developing the tool gPLINK to integrate PLINK with Haploview http://www.broad.mit.edu/mpg/haploview/. Haploview 4.0 provides a number of features for viewing, filtering and plotting PLINK results files. This is intended to supplant the methods suggested below.

All the output files that PLINK generates are plain-text, space-delimited files. Most files will have the same number of fields per line and will have the field names in the first line, facilitating use of a spreadsheet or statistics package to view and process the results.

For small results files, simply printing the files to the terminal or viewing in a text-editor should work well. In Windows/MS-DOS use the type command, e.g.

type mydata.assoc

to view a results file. Alternatively, you can call up WordPad from the command line as follows:

write mydata.assoc

If you are using a Unix/Linux system, then commands such as cat, more or less can be used to display the results; alternatively text-editors such as pico, emacs or vi.

Of course, Unix/Linux users also have available the entire range of text-processing tools (grep, gawk, perl, sort, head, etc) and shell-scripting tools, as well as powerful text-editors (emacs, etc) that are ideal for processing very large result files. Another alternative is to use a statistics package such as the R package www.r-project.org which will provide powerful visualisation tools also.

Windows/MS-DOS users have fewer options for handling very large results files: For moderate size files (e.g. up to 50K SNPs), you could use Excel. For larger files, you can either install cygwin http://www.cygwin.com/ to provide a Linux-like environment, or use a statistics package such as the R package www.r-project.org.

**Personal opinion...** Although a MS-DOS version of PLINK is supported, we would, in general, advise any any researchers planning on performing many large-scale analyses to look into adopting a Linux environment, if they are not already using this.

# Chapter 2

# Basic usage / data formats

PLINK is a command line program written in C/C++. All commands involve typing plink at the command prompt (e.g. DOS window or Unix terminal) followed by a number of options (all starting with --option) to specify the data files / methods to be used. All results are written to files with various extensions. The name of the file is by default plink.ext where .ext will change depending on the content of the file. Often these files will be large: using a package such as R is suggested for visualising and tabulating output. The majority of output files are in a standard plain text 'rectangular' format, with one header row and a fixed number of columns per line. A complete list of all options and output file types is given in the reference section

# 2.1 Running PLINK

PLINK is a command-line program: clicking on an icon will get you nowhere: please consult these notes on downloading and installing PLINK. Open up a command prompt or terminal window and perform all analyses by typing commands as described below.

```
plink --file mydata
```

where we expect two files: in this case, mydata.ped and mydata.map.

When PLINK starts it will attempt to contact the web, to check whether there is a more up-to-date version available or not. After checking, PLINK writes a file called .pversion to the working directory and use this cached information for the rest of the day. This option can be disabled with the --noweb option on the command line. When using PLINK on a machine with no, or a very slow, web connection, it may be desirable to turn this feature off. This feature is turned on by default so that users are aware of new versions that may contain important new features or bug fixes. If your current version of PLINK is out of date, then a warning message will be displayed, suggesting that you download and install the current version. (This is the only reason the web connection is made – no other data is transmitted to the server.) If the current version is up-to-date, you will see something like the following:

```
Web-based version check ( --noweb to skip )
Connecting to web... OK, v1.04 is current
whereas, if the current version is not up-to-date, you will see something like the following:
Web-based version check ( --noweb to skip )
Connecting to web...

*** UPDATE REQUIRED ***

This version : 1.03
Most recent version : 1.04
Please upgrade your version of PLINK as soon as possible!
```

```
(visit the above website for free download)
Old versions of PLINK (<1.04) contain bugs fixed in 1.04
```

The web-based version check will also produce warning if an command used was found to have some issue discovered since that version was released (the warning will contain a link to a web page describing the issue).

To re-run a previous job, use the --rerun option, which takes a PLINK LOG file as the parameter. This option will scan the LOG file, extract the previous PLINK commands and re-execute them. If new commands are added to the command line, they will also be included; if the command also appeared in the original file, any parameters will be taken from the newer version. For example, if the original command was

```
plink --file mydata --pheno pheno.raw --assoc --maf 0.05 --out run1
then the command
plink --rerun run1.log --maf 0.1
```

would repeat the analysis but with the new minor allele frequency threshold of 0.1, not 0.05. Note that commands in the old LOG file can be overwritten but not removed with the rerun command.

Note By default, the --out statement would also be copied, and so the new output would overwrite any old results (i.e. with the run1 fileroot). It is often a good idea to also add a new --out command, therefore:

```
plink --rerun run1.log --maf 0.1 --out run2
```

For very long a complex commands, --rerun can save typing and help reduce mistakes.

HINT MS-DOS only allows command lines to be 127 characters in length – sometimes, PLINK command lines can grow longer than this. In this case, use the --script option, where the remaining options will be read from a text file. For example,

```
plink --script myscript1.txt
where the file myscript1.txt is a plain text file containing
--ped ..\data\version1\50K\allsamples.ped
--map ..\data\allmapfiles\finalversion\autosomal.map
--out ..\results\working\sample-missingness-v1.22
--from rs66537222
--to rs8837323
--geno 0.25
--maf 0.02
--missing
```

would be the same as typing all these options in at the command line (note that the commands do not need to be all on the same line now). Another advantage of using script files is that it aids attempts at making one's research reproducible.

#### 2.2 PED files

As well as the **--file** command described above, PED and MAP files can be specified separately, if they have different names:

```
plink --ped mydata.ped --map autosomal.map
```

**Note** Loading a large file (100K+ SNPs) can take a while (which is why we suggest converting to binary format). PLINK will give an error message in most circumstances when something has gone wrong.

The PED file is a white-space (space or tab) delimited file: the first six columns are mandatory:

```
Family ID
Individual ID
Paternal ID
Maternal ID
Sex (1=male; 2=female; other=unknown)
Phenotype
```

The IDs are alphanumeric: the combination of family and individual ID should uniquely identify a person. A PED file must have 1 and only 1 phenotype in the sixth column. The phenotype can be either a quantitative trait or an affection status column: PLINK will automatically detect which type (i.e. based on whether a value other than 0, 1, 2 or the missing genotype code is observed).

If an individual's sex is unknown, then any character other than 1 or 2 can be used. When new files are created (PED, FAM, or other which contain sex) then the original coding will be preserved. However, these individuals will be dropped from any analyses (i.e. phenotype set to missing also) and an error message will arise if an analysis that uses family information is requested and an individual of 'unknown' sex is specified as a father or mother.

HINT To disable the automatic setting of the phenotype to missing if the individual has an ambiguous sex code, add the --allow-no-sex option. When using a data generation command (e.g. --make-bed, --recode, etc) as opposed to an analysis command, then by default the phenotype is not set to missing is sex is missing. This behaviour can be changed by adding the flag --must-have-sex.

**HINT** You can add a comment to a PED or MAP file by starting the line with a # character. The rest of that line will be ignored. Do not start any family IDs with this character therefore.

Affection status, by default, should be coded:

- -9 missing
- 0 missing
- 1 unaffected
- 2 affected

If your file is coded 0/1 to represent unaffected/affected, then use the --1 flag:

```
plink --file mydata --1
```

which will specify a disease phenotype coded:

- -9 missing
- 0 unaffected
- 1 affected

The missing phenotype value for quantitative traits is, by default, -9 (this can also be used for disease traits as well as 0). It can be reset by including the --missing-phenotype option:

```
plink --file mydata --missing-phenotype 99
```

Other phenotypes can be swapped in by using the --pheno (and possibly --mpheno) option, which specify an alternate phenotype is to be used, described below.

Genotypes (column 7 onwards) should also be white-space delimited; they can be any character (e.g. 1,2,3,4 or A,C,G,T or anything else) except 0 which is, by default, the missing genotype character. **All markers should be biallelic**. All SNPs (whether haploid or not) must have two alleles specified. Either Both alleles should be missing (i.e. 0) or neither. No header row should be given. For example, here are two individuals typed for 3 SNPs (one row = one person):

```
FAMO01 1 0 0 1 2 A A G G A C FAMO01 2 0 0 1 2 A A A G 0 0
```

• • •

The default missing genotype character can be changed with the --missing-genotype option, for example:

```
plink --file mydata --missing-genotype N
```

**NOTE** Different values to the missing phenotype or genotype code can be specified for output datasets created, with --missing-output-phenotype and --missing-output-genotype.

#### 2.2.1 Different PED file formats: missing fields

Sometimes data arrive in a number of different formats: for example, where the genotype information just has a single ID column followed by all the SNP data, with the other family and phenotype information residing in a separate file. Rather than have to recreate new files, it is sometimes possible to read in such files directly. The standard behavior of PLINK when reading a PED file with --file or --ped can be modified to allow for the fact that one or more of the normally obligatory 6 fields are missing:

```
--no-fid
```

indicates there is no Family ID column: here the first field is taken to be individual ID, and the family ID is automatically set to be the same as the individual ID (i.e. obviously, all individuals would be treated as unrelated). In other files that require family and individual ID (e.g. alternate phenotype file and cluster files, for which this flag has no effect), the individual ID would need to be entered also as the family ID therefore.

```
--no-parents
```

indicates that there are no paternal and maternal ID codes; all individuals would be assumed to be founders in this case

```
--no-sex
```

indicates that there is no sex field; all individuals set to have a missing sex code (which also sets that individual to missing unless the allow-no-sex option is also used)

```
--no-pheno
```

indicates that there is no phenotype filed; all individuals are set to missing unless an alternate phenotype file is specified.

It is possible to use these flags together, so using all of them would specify the most simple kind of file mentioned above: a single, unique ID code followed by all genotype data.

**IMPORTANT** These options only work for the basic PED file (i.e. specified by --file or --ped. They do not work for transposed files, when merging in a file with --merge, or with binary filesets or covariate, cluster or alternate phentype files.

If the genotype codes in a PED file are in the form AG rather than A G, for example, such that every genotype is exactly two characters long, then then flag

```
./plink --file mydata --compound-genotypes </tt>
```

can be added. Note that this only works for input for PED files (not TPED or LGEN files, and not for any output options, e.g. --recode, etc).

**Note** To load the PED file from the standard input stream instead of a file, use the - symbol as the file name, e.g.

```
perl retrieve_data.pl | ./plink --ped - --map mymap.map --make-bed
```

The MAP file still needs to be a normal file; this currently only works for --ped files.

#### 2.3 MAP files

By default, each line of the MAP file describes a single marker and must contain exactly 4 columns:

```
chromosome (1-22, X, Y or 0 if unplaced) rs# or snp identifier
Genetic distance (morgans)
Base-pair position (bp units)
```

Genetic distance can be specified in centimorgans with the --cm flag. Alternatively, you can use a MAP file with the genetic distance excluded by adding the flag --map3, i.e.

```
plink --file mydata --map3
In this case, the three columns are expected to be
    chromosome (1-22, X, Y or 0 if unplaced)
    rs# or snp identifier
    Base-pair position (bp units)
```

**Note** Most analyses do not require a genetic map to be specified in any case; specifying a genetic (cM) map is most crucial for a set of analyses that look for shared segments between individuals. For basic association testing, the genetic distance column can be set at 0.

SNP identifiers can contain any characters except spaces or tabs; also, you should avoid \* symbols in names also.

To exclude a SNP from analysis, set the 4th column (physical base-pair position) to any negative value (this will only work for MAP files, not for binary BIM files).

```
1 rs123456 0 1234555

1 rs234567 0 1237793

1 rs224534 0 -1237697 <-- exclude this SNP

1 rs233556 0 1337456
```

The MAP file must therefore contain as many markers as are in the PED file. The markers in the PED file do not need to be in genomic order: (i.e. the order MAP file should align with the order of the PED file markers).

#### 2.3.1 Chromosome codes

The autosomes should be coded 1 through 22. The following other codes can be used to specify other chromosome types:

The numbers on the right represent PLINK's internal numeric coding of these chromosomes: these will appear in all output rather than the original chromosome codes.

For haploid chromosomes, genotypes should be specified as homozygotes: for most analyses, PLINK will treat these appropriately. For example, consider the following example PED file, containing two males (1 and 2) and two females (3 and 4):

```
1 1 0 0 1
                 A A
                                        A A
                                                A A
2 1 0 0 1
                 A C
                         A C
                                 A C
                                        A C
                                                A C
             1
3 1 0 0 2
                 A A
                         A A
                                A A
                                        A A
                                                A A
4 1 0 0 2
                 A C
                         A C
                                 A C
                                        A C
                                                A C
```

and MAP file

```
snp1
                   1000
1
X
                   1000
      snp2
              0
Y
      snp3
              0
                   1000
XY
      snp4
              0
                   1000
                   1000
MT
      snp5
              0
```

Generating frequencies for these SNPs,

```
plink --file test --freq
```

we see plink.frq is

CHR	SNP	A1	A2	MAF	NM
1	snp1	C	Α	0.25	8
23	snp2	C	Α	0.2	5
24	snp3	C	Α	0	1
25	snp4	C	Α	0.25	8
26	snp5	C	Α	0	2

There are several things to note. First, the numeric chromosome codes are used in the output to represent X, Y, XY and MT. Second, haploid chromosomes are only counted once (i.e. male X and Y chromosome SNPs and all MT SNPs). Third, several genotypes have been set to missing if they are not valid (female Y genotype, heterozygous haploid chromosome). The NM field represents the number of non-missing alleles for each SNP – this is because invalid genotypes are automatically set to missing.

We can see which genotypes have been set to missing by running the --recode command; however, usually PLINK preserves all genotypes when generating a new file (i.e. if one is just reformatting a file, say from text to binary format, it is not necessarily desirable to change any of the content; as above, summary statistic and analysis commands do set these genotypes missing automatically still). However, if we also add the --set-hh-missing flag, any invalid genotypes will be set to missing in the new file:

```
plink --file test --set-hh-missing
```

which creates the new PED file plink.recode.ped

In other words, the actual alleles that PLINK pays attention to are shown in **bold**, all non-bold alleles are ignored.

```
1 1 0 0 1
                                             \mathbf{A} \mathbf{A}
                                                                 \mathbf{A} A
                                                                                       \mathbf{A} A
                                                                                                             \mathbf{A} \mathbf{A}
                                                                                                                                   \mathbf{A} A
2 1 0 0 1
                                             \mathbf{A} \mathbf{C}
                                                                   A C
                                                                                        A C
                                                                                                            \mathbf{A} \mathbf{C}
                                                                                                                                  A C
                                  1
3 1 0 0 2
                                  1
                                             \mathbf{A} \mathbf{A}
                                                                 \mathbf{A} \mathbf{A}
                                                                                        A A
                                                                                                            \mathbf{A} \mathbf{A}
                                                                                                                                  \mathbf{A} A
4 1 0 0 2
                                             \mathbf{A} \mathbf{C}
                                                                 \mathbf{A} \mathbf{C}
                                                                                       A C
                                                                                                            \mathbf{A} \mathbf{C}
                                                                                                                                  A C
```

#### 2.3.2 Allele codes

By default, the minor allele is coded A1 and the major allele is coded A2 (this is used in many output files, e.g. from --freq or --assoc). By default this is based on all founders (unless --nonfounders is added) with sex-codes specified (unless --allow-no-sex is added). This coding is applied after any other filters have been applied. It is sometimes desirable to present this automatic flipping of A1 and A2 alleles, by use of the --keep-allele-order option. For example, if one wishes to dump the genotype counts by use of the --model command, for two groups of individuals (using the --filter command), this ensures that the same minor allele will always be used in grp1.model as grp2.model (which can facilitate downstream processing of these files, for instance).

```
plink --bfile --filter pop.dat POP1 --model --keep-allele-order --out pop-1-genotypes plink --bfile --filter pop.dat POP2 --model --keep-allele-order --out pop-2-genotypes
```

That is, for any SNP that happens to have a different minor allele in POP1 versus POP2, the output in the two .model files will still line up in an easy manner.

## 2.4 Transposed filesets

Another possible file-format called a *transposed* fileset, containing two text files: one (TPED) containing SNP and genotype information where one row is a SNP; one (TFAM) containing individual and family information, where one row is an individual.

The first 4 columns of a TPED file are the same as a standard 4-column MAP file. Then all genotypes are listed for all individuals for each particular SNP on each line. The TFAM file is just the first six columns of a standard PED file. In otherwords, we have just taken the standard PED/MAP file format, but swapped all the genotype information between files, after rotating it 90 degrees. For each, the above example PED/MAP fileset

```
<---- normal.ped ---->
1 1 0 0 1 1 A A G T
2 1 0 0 1 1 A C T G
3 1 snp1 0 5000650
3 1 0 0 1 1 C C G G
4 1 0 0 1 2 A C T T
5 1 0 0 1 2 C C G T
6 1 0 0 1 2 C C T T
```

would be represented as TPED/TFAM files:

```
<----> trans.tped ---->
                                         <- trans.tfam ->
1 snp1 0 5000650 A A A C C C A C C C C
                                         1 1
                                              0
                                                 0 1
                                         2
1 snp2 0 5000830 G T G T G G T T G T T T
                                         3
                                           1
                                              0
                                                 0
                                                 0
                                         5
                                           1
                                                 Ω
                                                    1
                                                      2
                                              0
```

This kind of format can be convenient to work with when there are very many more SNPs than individuals (i.e. WGAS data). In this case, the TPED file will be very long (as opposed to the PED file being very wide).

To read a transposed fileset, use the command

```
plink --tfile mydata
```

which implies mydata.tped and mydata.tfam exists; alternatively, if the files are differently named, they can be individually, fully specified:

```
plink --tped mydata.tped --tfam pedinfo.txt
```

**HINT** You can generate transposed filesets with the --transpose option, described in the data management section

# 2.5 Long-format filesets

Another possible file-format called a *long-format* fileset, containing three text files:

• a LGEN file containing genotypes (5 columns, one row per genotype)

- a MAP file containing SNPs (4 columns, one row per SNP)
- a FAM file containing individuals (6 columns, one row per person)

The MAP and FAM/PED files are described elsewhere this page. Consider the following example: A MAP file test.map

```
1 snp2 0 2
     2 snp4 0 4
     1 snp1 0 1
     1 snp3 0 3
     5 snp5 0 1
as described above. A FAM file test.fam
     1 1 0 0 1 2
     2 1 0 0 2 2
     2 2 0 0 1 1
     9 1 1 2 0 0
as described below. Finally, an LGEN file, test.lgen
     1 1 snp1 A A
     1 1 snp2 A C
     1 1 snp3 0 0
     2 1 snp1 A A
     2 1 snp2 A C
     2 1 snp3 0 0
     2 1 snp4 A A
     2 2 snp1 A A
     2 2 snp2 A C
     2 2 snp3 0 0
     2 2 snp4 A A
The columns in the LGEN file are
     family ID
     individual ID
     snp ID
     allele 1 of this genotype
     allele 2 of this genotype
```

Not all entries need to be present in the LGEN file (e.g. snp5 or person 9/1) or snp4 for person 1/1. These genotypes will be set to missing internally. The order also need not be the same in the LGEN file as for the MAP or FAM files. If a genotype is listed more than once, the final version of it will be used.

LGEN file can be reformatted as a standard PED file using the following command:

```
plink --lfile test --recode
which creates these two files: a PED file, plink.recode.map
     1 1 0 0 1
                    A A A C
                              0 0
                2
                                   0 0
     2 1 0 0 2
                    A A
                         A C
                              0 0
                                    A A
     2 2 0 0 1 1
                    AAACOO AA OO
     9 1 1 2 0 0
                    0 0 0 0 0 0 0 0 0 0
and the MAP file, plink.recode.map (note: it has been put in genomic order)
     1
             snp1
                     0
                              1
     1
             snp2
                     0
                              2
     1
             snp3
                     0
                              3
     2
             snp4
                              4
                     0
```

```
5 snp5 0 1
```

**NOTE** All individuals must be uniquely identified by the combination of the family and individual IDs. To read a long-format fileset, use the command

```
plink --lfile mydata
```

which implies mydata.lgen, mydata.map and mydata.map exist.

**NOTE** Currently, you cannot output a fileset in this format in PLINK.

# 2.6 Binary PED files

To save space and time, you can make a binary ped file (\*.bed). This will store the pedigree/phenotype information in separate file (\*.fam) and create an extended MAP file (\*.bim) (which contains information about the allele names, which would otherwise be lost in the BED file). To create these files use the command:

The .fam and .bim files are still plain text files: these can be viewed with a standard text editor. Do not try to view the .bed file however: it is a compressed file and you'll only see lots of strange characters on the screen...

NOTE Do not make any changes any of these three files; e.g. setting the position to a negative value will not work to exclude a SNP for binary files

You can specify a different output root file name (i.e. different to "plink") by using the --out option:

```
plink --file mydata --out mydata --make-bed
which will create
   mydata.bed
   mydata.fam
   mydata.bim

To subsequently load a binary file, just use --bfile instead of --file
plink --bfile mydata
```

When creating a binary ped file, the MAF and missingness filters are set to include everybody and all SNPs. If you want to change these, use --maf, --geno, etc, to manually specify these options: for example,

```
plink --file mydata --make-bed --maf 0.02 --geno 0.1
```

More information... If you want to write your own software that uses the BED file format, please follow this link for more information of the specification.

## 2.7 Alternate phenotype files

To specify an alternate phenotype for analysis, i.e. other than the one in the \*.ped file (or, if using a binary fileset, the \*.fam file), use the --pheno option:

```
plink --file mydata --pheno pheno.txt
```

where pheno.txt is a file that contains 3 columns (one row per individual):

Family ID Individual ID Phenotype

The original PED file must still contain a phenotype in column 6 (even if this is a dummy phenotype, e.g. all missing), unless the --no-pheno flag is given.

If an individual is in the original file but not listed in the alternate phenotype file, that person's phenotype will be set to missing. If a person is in the alternate phenotype file but not in the original file, that entry will be ignored. The order of the alternate phenotype file need not be the same as for the original file. If the phenotype file contains more than one phenotype, then use the --mpheno N option to specify the Nth phenotype is the one to be used:

```
plink --file mydata --pheno pheno2.txt --mpheno 4
```

where pheno2.txt contains 5 different phenotypes (i.e. 7 columns in total), this command will use the 4th for analysis (phenotype D):

Family ID
Individual ID
Phenotype A
Phenotype B
Phenotype C
Phenotype D
Phenotype E

Alternatively, your alternate phenotype file can have a header row, in which case you can use variable names to specify which phenotype to use. If you have a header row, the first two variables **must** be labelled FID and IID. All subsequent variable names cannot have any whitespace in them. For example,

```
FID IID qt1 bmi site
F1 1110 2.3 22.22 2
F2 2202 34.12 18.23 1
...
```

then

```
plink --file mydata --pheno pheno2.txt --pheno-name bmi --assoc
```

will select the second phenotype labelled "bmi", for analysis

Finally, if there is more than one phenotype, then for basic association tests, it is possible to specify that all phenotypes be tested, sequentially, with the output sent to different files: e.g. if bigpheno.raw contains 10,000 phenotypes, then

```
plink --bfile mydata --assoc --pheno bigpheno.raw --all-pheno
```

will loop over all of these, one at a time testing for association with SNP, generating a lot of output. You might want to use the --pfilter command in this case, to only report results with a p-value less than a certain value, e.g. --pfilter 1e-3.

WARNING Currently, all phenotypes must be numerically coded, including missing values, in the alternate phenotype file. The default missing value is -9, change this with --missing-phenotype, but it must be a numeric value still (in contrast to the main phenotype in the PED/FAM file).

#### 2.7.1 Creating a new binary phenotype automatically

To automatically form a one-versus-others binary phenotype (note: binary meaning dichotomous here, rather than a BED/binary-PED file) from a categorical covariate/phenotype file, use the command

```
plink --bfile mydata --make-pheno site.cov SITE3 --assoc
which assumes the file
     site.cov
contains exactly three fields
     Family ID
     Individual ID
     Code from which phenotype is created
For example, if it were
     A1
        1 SITE1
     В1
         1 SITE1
     C1
         1 SITE2
     D1
         1 SITE3
     E1
            SITE3
     F1
         1
            STTE4
     G2
            SITE4
```

then the above command would make individuals D1 and E1 as *cases* and everybody else as *controls*. However, if individuals present in mydata were not specified in site.cov, then these people would be set to have a missing phenotype.

An alternate specification is to use the \* symbol instead of a value, e.g.

```
plink --bfile mydata --make-pheno p1.list * --assoc
which assumes the file
    p1.list
contains exactly two fields
    Family ID
    Individual ID
```

In this case, anybody in the file p1.list would be made a case; all other individuals in mydata but not in p1.list would be set as a control.

#### 2.8 Covariate files

Certain PLINK commands support the inclusion of one or more covariates. Note that for stratified analyses, namely using the CMH (--mh) options, the strata are specified using the --within option to define clusters, rather than --covar.

To load a covariate use the option:

```
plink --file mydata --covar c.txt
```

The covariate file should be formatted in a similar manner to the phenotype file. If an individual is not present in the covariate file, or if the individual has a missing phenotype value (i.e. -9 by default) for the covariate, then that individual is set to missing (i.e. will be excluded from association analysis).

To select a particular subset of covariates, use one of the following commands, which either use numbers or names (i.e. if a header row exists in the file),

```
plink --file mydata --covar c.txt --covar-number 2,4-6,8
```

```
plink --file mydata --covar c.txt --covar-name AGE, BMI-SMOKE, ALC
```

Note that ranges can be used in both cases, with the - hyphen symbol, e.g. if the first row were

```
FID IID SITE AGE DOB BMI ETH SMOKE STATUS ALC
```

then both the above commands would have the same effect, i.e. selecting AGE, BMI, ETH, SMOKE, ALC.

To output a new covariate file, possibly with categorical variables downcoded to binary dummy variables (v1.01 onwards) use the --write-covar option as described here

Exception If the --gxe command is used, that selects only a single covariate, then use the command --mcovar, that works similarly to --mpheno to select which single covariate to use: with the --gxe command, the --covar-name and --covar-number options will not work.

NOTE Not all commands accept covariates, and PLINK will not always give you an error or warning. The basic association (--assoc, --mh, --model, --tdt, --dfam, and --qfam) do not accept covariates, neither do the basic haplotype association methods (--hap-assoc, --hap-tdt). Among the commands that do are --linear, --logistic, --chap and --proxy-glm. Also --gxe accepts a single covariate only (the others listed here accept multiple covariates).

#### 2.9 Cluster files

To load a cluster solution, or indeed any categorical grouping of the sample, use the --within option:

```
plink --file mydata --within f.txt
```

If this option is used, then permutation procedures will permute within-cluster only, effectively controlling for any effect of cluster membership. Similarly, tests that perform stratified analyses, such as the Cochran-Mantel-Haenszel, this option is used to define the strata.

This file should have a similar structure to the alternate phenotype file. The clusters can be coded either numerically or as strings:

```
F1 I1 A
F2 I1 B
F3 I1 B
F4 I1 C1
F5 I1 A
F6 I1 C2
F7 I1 C2
```

Here, individuals would be grouped in four groups:

```
Cluster A: F1/I1 F5/I1
Cluster B: F2/I1 F3/I1
Cluster C1: F4/I1
Cluster C2: F6/I1 F7/I1
```

All individuals in the file should be assigned to a single cluster in the cluster file.

#### 2.10 Set files

Various options (e.g. set based tests, gene-based tests of epistasis) require sets of SNPs to be specified. This is performed by including the --set option on the command line, followed by a filename that defines the sets. The file mydata.set should be in the following format:

```
SET_A
rs10101
rs20234
rs29993
END
SET_B
rs2344
rs888833
END
```

That is, each set must start with a *set name* (e.g. SET\_A). This name can not have any spaces in it. The name is followed by a list of SNPs in that set. The keyword END specifies the end of that particular set. Do not name any SNPs to have the name END!

Sets can be overlapping. Any SNPs specified in the set that do not appear in the actual data, or that have been excluded due to filters used, will be ignored.

The format is flexible in terms of whether each item appears on one line: the set file only needs to be whitespace delimited. For example, the file above could be specified as:

```
SET_A rs10101 rs20234 rs29993 END
SET_B rs2344 rs888833 END
```

**HINT** It is possible to automatically create a set-file, given a list of genomic co-ordinates, using the --make-set command, described here.

# Chapter 3

# Data management tools

PLINK provides a simple interface for recoding, reordering, merging, flipping DNA-strand and extracting subsets of data.

## 3.1 Recode and reorder a sample

A basic, but often useful feature, is to output a dataset:

- with the PED file markers reordered for physical position,
- with excluded SNPs (negative values in the MAP file) excluded from the new PED file
- possibly excluding other SNPs based on filters such as genotyping rate
- possibly recoding the SNPs to a 1/2 coding
- possibly recoding the SNPs between letters and numbers (A,C,G,T / 1,2,3,4)
- possibly transposing the genotype file (SNPs as rows)
- possibly recoding the SNP to an additive and dominant pair of components
- possibly listing the data with each specific genotype as a distinct row

The basic option to generate a new dataset is the --recode option:

```
plink --file data --recode
```

which will output the allele labels as they appear in the original; also, the missing genotype code is preserved if this is different from 0. Also, if --missing-output-genotype is specified (which can be as well as --missing-genotype) then this value will be used instead (i.e. so that input and output files can have different missing codes; this also applies to the phenotype with --missing-output-phenotype and --missing-phenotype).

The --make-bed option does the same as --recode but creates binary files; these can also be filtered, etc, as described below.

In contrast,

```
plink --file data --recode12
```

will recode the alleles as 1 and 2 (and the missing genotype will always be 0). Both these commands will create two new files

```
plink.ped
```

```
plink.map
```

(where, as usual, "plink" would be replaced by any specified -out filename).

Unless manually specified, for all these options, the usual filters for missingness and allele frequency will be set so as not to exclude any SNPs or individuals. By explicitly including an option, e.g. --maf 0.05 on the command line, this behaviour is overriden (see this page).

By default, any --recode option, and also --make-bed will preserve all genotypes exactly as they are. To set to missing Mendel errors or heterozygous haploid calls, use the options --set-me-missing and --set-hh-missing respectively. For the former, you will also need to specify --me 1 1 (i.e. to invole an evaluation of Mendel errors, which does not occur by default, by not excluding any individuals or SNPs based on the results, i.e. if you only want to zero-out certain genotypes).

To recode SNP alleles from A,C,G,T to 1,2,3,4 or vice versa, use --allele1234 (to go from letters to numbers) and --alleleACGT (to go from numbers to letters). These flags should be used in conjunction with a data generation command (e.g. --make-bed), or any other analysis or summary statistic option. Alleles other than A,C,G,T or 1,2,3,4 will be left unchanged. It is sometimes useful to have a PED file that is tab-delimited, except that between alleles of the same genotype a space instead of a tab is used. A file formatted in this way can load into Excel, for example, as a tab-delimited file, but with one genotype per column instead of one allele per column. Use the option --tab as well as --recode or --recode12 to achieve this effect.

To make a new file in which non-founders without both parents also in the same fileset are recoded as founders (i.e. pat and mat codes set both to 0), add the --make-founders flag.

#### 3.1.1 Transposed genotype files

When using either --recode or --recode12, you can obtain a transposed text genotype file by adding the --transpose option. This generates two files:

```
plink.tped
plink.fam
```

The first contains the genotype data, with SNPs as rows and individuals as columns, for example: if the original file was

```
1 1 0 0 1 1 1 1 G G
1 2 0 0 2 1 0 0 A G
1 3 0 0 1 1 1 1 A G
1 4 0 0 2 1 2 1 A A
```

then this would generate

```
1 snp1 0 10001 1 1 0 0 1 1 2 1 1 snp2 0 20001 G G G A G A A A
```

The first four columns are from the MAP file (chromosome, SNP ID, genetic position, physical position), followed by the genotype data. The plink.fam gives the ID, sex and phenotype information for each individual. The order of individuals in this file is the same as the order across the columns of the TPED file. The FAM file is just the first six columns of the PED file (or literally the same FAM file if the input where a binary fileset).

#### 3.1.2 Additive and dominance components

The following format is often useful if one wants to use a standard, non-genetic statistical package to analyse the data, as here genotypes are coded as a single allele dosage number. To create a file with SNP genotypes recoded in terms of additive and dominant components, use the option:

```
plink --file data --recodeAD
```

which, assuming C is the minor allele, will recode genotypes as follows:

SNP		$\mathtt{SNP}_{-}\mathtt{A}$	,	SNP_HET		
A A	->	0	,	0		
A C	->	1	,	1		
C C	->	2	,	0		
0 0	->	NA	,	NA		

In otherwords, the default for the additive recoding is to count the number of minor alleles per person. The --recodeAD option produces both an additive and dominance coding: use --recodeA instead to skip the SNP\_HET coding.

The --recodeAD option saves the data to a single file

```
plink.raw
```

which has a header row indicating the SNP names (with \_A and \_HET appended to the SNP names to represent additive and dominant components, respectively).

For example, consider the following PED file, which has two SNPs:

```
1 1 0 0 1 1 1 1 G G
1 2 0 0 2 1 0 0 A G
1 3 0 0 1 1 1 1 A G
1 4 0 0 2 1 2 1 A A
```

Using the --recodeAD option generates the file plink-recode.raw:

```
FID IID PAT MAT SEX PHENOTYPE snp1_2 snp1_HET snp2_G snp2_HET
1 1 0 0 1 1 0 0 0 2 0
1 2 0 0 2 1 NA NA 1 1
1 3 0 0 1 1 0 0 0 1 1
1 4 0 0 2 1 1 1 0 0
```

The column labels reflect the snp name (e.g. snp1) with the name of the minor allele appended (i.e. snp1\_2 in the first instance, as 2 is the minor allele) for the additive component. The dominant component (a dummy variable reflecting heterozygote state) is coded with the \_HET suffix.

This file can be easily loaded into R: for example:

```
d <- read.table("plink.raw",header=T)</pre>
```

For example, for the first SNP, the individuals are coded 1/1, 0/0, 1/1 and 2/1. The additive count of the number of common (1) alleles is therefore: 2, NA, 2 and 1, which is reflected in the field snp1\_2. The field snp1\_HET is coded 1 for the fourth individual who is heterozygous – this field can be used to model dominance effect of the allele.

The behavior of the --recodeA and --recodeAD commands can be changed with the --recode-allele command. This allows for the 0, 1, 2 count to reflect the number of a pre-specified allele type per SNP, rather than the number of the minor allele. This command takes as a single argument the name of a file that lists SNP name and allele to report, e.g. if the file recode.txt contained

```
snp1 1
snp2 A

then

plink --file data --recodeAD --recode-allele recode.txt

would now report in the LOG file
   Reading allele coding list from [ recode.txt ]
   Read allele codes for 2 SNPs
and the plink.raw file would read
   FID IID PAT MAT SEX PHENOTYPE snp1_1 snp1_HET snp2_A snp2_HET
```

If the SNP is monomorphic, by default the allele code out will be 0 and all individuals will have a count of 0 (or NA). If an allele is specified in --recode-allele that is not seen in the data, similarly all individuals will receive a 0 count (i.e. rather than an error being given).

**NOTE** For alleles that have exactly 0.50 minor allele frequency, as for the second SNP in the example above, then which allele is labelled as minor will depend on which was first encountered in the PED file.

#### 3.1.3 Listing by genotype

Another format that might sometimes be useful is the --list option which genetes a file

```
plink.list
```

that is ordered one genotype per row, listing all family and individual IDs of people with that genotype. For example, if we have a file with two SNPs rs1001 and rs2002 (both on chromosome 1):

```
A 1 0 0 1 2 A A 1 1
    B 2 0 0 1 2 A C 0 0
    C 3 0 0 1 1 A C 1 2
    D 4 0 0 1 1 C C 1 2
then then option
 plink --file mydata --list
will generate the file plink.list
    1 rs1001 AA A 1
     1 rs1001 AC B 2 C 3
     1 rs1001 CC D 4
    1 rs1001 00
    1 rs2002 22
    1 rs2002 21 C 3 D 4
     1 rs2002 11 A 1
    1 rs2002 00 B 2
which has columns
    Chromosome
    SNP identifier
    Genotype
    Family ID, Individual ID for 1st person
    Family ID, Individual ID for 2nd person
    Family ID, Individual ID for final person
```

Obviously, different rows will have a different number of columns. Here, we see that individual A 1 has the A/A genotype for rs1001, etc. This option is often useful in conjunction with --snp, if you want an easy breakdown of which individuals have which genotypes.

#### 3.2 Write SNP list files

To output just the list of SNPs that remain after all filtering, etc, use the --write-snplist command, e.g. to get a list of all high frequency, high genotyping-rate SNPs:

```
plink --bfile mydata --maf 0.05 --geno 0.05 --write-snplist
which generates a file
    plink.snplist
```

This file is simply a list of included SNP names, i.e. the same SNPs that a --recode or --make-bed statement would have produced in the corresponding MAP or BIM files.

## 3.3 Update SNP positions

To automatically update either the genetic or physical positions for some or all SNPs in a dataset, use the --update-map command, which takes a single parameter of a filename, e.g.

```
plink --bfile mydata --update-map build36.txt --make-bed --out mydata2
```

where, for example, the file build36.txt contains new physical positions for SNPs, based on db-SNP126/build 36, in the simple format of SNP/position per line, e.g.

```
rs100001 1000202
rs100002 6252678
rs100003 7635353
```

To change genetic position (3rd column in map file) add the flag --update-cm as well as --update-map. There is no way to change chromosome codes using this command. Normally, one would want to save the new file with the changed positions, as in the example above, although one could combine other commands instead (e.g. association testing, etc) although the updated positions would then be lost (i.e. the changes are not automatically saved).

Not all SNPs need feature in the file supplied here – these SNPs will keep there old position. If a SNP is listed more than once in this file, an error will be reported. Importantly, if this command changes the implied ordering of SNPs, a message will be written to the command line. Note, the order of SNPs will not be changed in the existing dataset with this command, only the positions. If the order has changed, then any command which relies on relative SNP positions (e.g. --hap-window, --homozyg, etc) should not be used on that dataset. In this case, it is necessary to save the file; then when reloading it, the SNPs will be automatically re-ordered upon reloading. If the LOG file does not show a message that the order of SNPs has changed, one need not worry.

#### 3.4 Write covariate files

If a covariate file is specified along with any of the above --recode options or with --make-bed, then that covariate file will also be written, as plink.cov by default. This option is useful if the covariate file has a different number of individuals, or is ordered differently, to produce a set of covariate values that line up more easily with the newly-created genotype and phenotype files.

```
plink --file data --covar myfile.txt --recode
```

creates also plink.cov. If you want just to create a revised version of the covariate file, but without creating a new set of genotype files, then use the --write-covar option. This can be used in conjunction with filters, etc, to output, for example, only covariates for high-genotyping (99%) cases, as in this example:

```
plink --file data --write-covar myfile.txt --filter-cases --mind 0.01
```

will output just the relevant lines of myfile.txt to plink.cov, sorted to match the order of data.ped. To also include phenotype information in the plink.cov file add the flag --with-phenotype. This can be useful, for example, when used in conjunction with --recodeA to generate the files needed to replicate an analysis in R (e.g. extracting the appropriate genotype data, and applying filters, etc).

#### 3.5 Write cluster files

Similar to --write-covar, the --write-cluster will output the *single* selected cluster from the file specified by --within. Unlike covariate files, this allows string labels to be used.

```
plink --bfile mydata --within clst.dat --write-cluster --out mynewfile
```

## 3.6 Flip DNA strand for SNPs

This command will read the list of SNPs in the file list.txt and flip the strand for these SNPs, then save a new PED or BED fileset (i.e. by using either the --recode or --make-bed commands):

```
plink --file data --flip list.txt --recode
```

The list.txt should just be a simple list of SNP IDs, one SNP per line.

Flipping strand means changing alleles

A -> T

C -> G

G -> C

T -> A

so, for example, a A/C SNP will become a T/G; alternatively, a A/T SNP will become a T/A SNP (i.e. in this case, the labels remain the same, but whether the minor allele is A or T will still depend on strand).

To flip strand for just a subset of the sample (e.g. if two samples have already been merged, and subsequently a strand issue has been identified for one of those samples) use the option --flip-subset, for example

```
plink --file data --flip list.txt --flip-subset mylist.txt --recode
```

where mylist.txt is a text file containing the individuals (family ID, individual ID) to be flipped.

**HINT** When merging two datasets, it is clearly very important that the two sets of SNPs are concordant in terms of positive or negative strand. Whereas some mismatches will be easy to spot as more than two alleles will be observed in the merged dataset, other instances will not be so easy to spot, i.e. for A/T and C/G SNPs.

# 3.7 Using LD to identify incorrect strand assignment in a subset of the sample

If cases and controls have been genotyped separately and then the data merged, it is always possible that strand has been incorrectly or incompletely assigned to each SNP, meaning that the merged data may contain a number of SNPs for which the allele coding differs between cases and controls (or between any other grouping, such as collection site, etc).

If the two mis-matched groups correspond to cases and controls exactly, then rare SNPs will show a very strong association with disease (e.g. 5% MAF in cases, 95% in controls) and be easy to spot as potential problems. More common SNPs could show intermediate levels of association that might be easier to confuse with a real signal.

A simple approach to detect some proportion of such SNPs uses differential patterns of LD in cases versus controls: the command --flipscan will query each SNP, and calculate the signed correlation between it and a set of nearby SNPs in cases and controls separately (of course, with the --pheno command, case and control status can be set to represent any binary split of the sample).

For each index SNP, PLINK identifies other SNPs in which the absolute value of the genotypic correlation is above some threshold. For these SNP pairs, it counts the number of times the signed correlation is different in sign between cases and controls (a *negative* LD pair) versus the same (a *positive* LD pair). For example, the command

```
plink --bfile mydata --flipscan
```

produces the output file

plink.flipscan

with the fields

CHR	Chromosome
SNP	SNP identifier for index SNP
BP	Base-pair position
A1	Minor allele code
A2	Major allele code
POS	Number of positive LD matches
$R\_POS$	Average correlation of these
NEG	Number of negative LD matches
$R\_NEG$	Average correlation of these
NEGSNPS	The SNPs showing negative correlation

For example, the majority of this file should show SNPs have a NEG value of 0; the value of POS will be zero or greater, depending on the extent of LD. For example:

CHR	SNP	BP	A1	A2	F	PLS	$R_PLS$	NEG	$R_NEG$	NEGSNPS
1	rs9439462	1452629	T	C	0	0	NA	0	NA	(NONE)
1	rs1987191	1457348	C	T	0	0	NA	0	NA	(NONE)
1	rs3766180	1468016	C	T	0.285	2	0.893	0	NA	(NONE)

However, occasionally one might observe different patterns of results. Of particular interest is when one SNP shows a large number of NEG SNPs. For example, here we show rs2240344 and nearby SNPs, all of which have at least one NEG SNP (lines truncated)

CHR	SNP	BP	A1	A2	F	PLS	$R\_PLS$	NEG	$R\_NEG$	NEGSNPS
14	rs12434442	72158039	T	C	0.249	5	0.515	1	0.46	rs2240344
14	rs4899437	72190986	G	C	0.394	5	0.802	1	0.987	rs2240344
14	rs2803980	72196284	G	Α	0.41	5	0.808	1	0.95	rs2240344
14	rs2240344	72197893	C	G	0.489	0	NA	7	0.807	rs12434442 rs4899437 .
14	rs2286068	72198107	C	T	0.407	7	0.741	1	0.962	rs2240344
14	rs7160830	72209491	T	C	0.414	6	0.801	1	0.922	rs2240344
14	rs10129954	72220454	T	C	0.413	6	0.729	1	0.73	rs2240344
14	rs7140455	72240734	T	C	0.469	4	0.72	1	0.64	rs2240344

This pattern of results quite clearly points to rs2240344 as being the odd man out: for 7 other SNPs, there is strong LD (r above 0.5) in either cases or controls, but with a SNP-SNP correlation in the other phenotype class that has the opposite direction. In contrast, there is not a single SNP for which both cases and controls have a consistent pattern of LD. For the nearby SNPs, all of which have only 1 NEG SNP, it is with rs2240344. So, in this particular case, it would suggest that stand is flipped in either cases or controls.

To display the specific sets of correlations in cases and controls for each SNP, add the option

#### --flip-scan-verbose

which generates a file

#### plink.flipscan.verbose

which lists for any SNP with at least one NEG pair of LD values, the correlations between the index SNP and the other flanking SNPs, showing the correlation in cases (R\_A) and controls (R\_U):

CHR_INDX	$\mathtt{SNP}_{\mathtt{INDX}}$	$\mathtt{BP}_{-}\mathtt{INDX}$	$A1_{-}INDX$	SNP_PAIR	BP_PAIR A:	1_PAIR	$R_A$	$R_{-}U$
14	rs2240344	72197893	C	rs12434442	72158039	T	-0.504	0.416
14	rs2240344	72197893	C	rs4899437	72190986	G	-0.99	0.983
14	rs2240344	72197893	C	rs2803980	72196284	G	-0.969	0.931
14	rs2240344	72197893	C	rs2286068	72198107	C	-0.971	0.952

14	rs2240344	72197893	C	rs7160830	72209491	T	-0.935	0.91
14	rs2240344	72197893	C	rs10129954	72220454	T	-0.782	0.679
14	rs2240344	72197893	C	rs7140455	72240734	T	-0.671	0.609

Here we see a clear pattern in which the correlation is similar between cases and controls in magnitude but has the opposite direction, strongly suggestive of a strand flip problem for this C/G SNP. In this case, the allele frequency turns out to be quite different between cases and controls (60% versus 40%) but the LD approach would have clearly detected this particular SNP being flipped in either cases or controls even if the true allele frequency were exactly 50%. This latter class of SNP would not cause problems of spurious association in single SNP analysis, but it could cause severe problems in haplotype and imputation analysis.

Naturally, if a SNP does not show strong LD with nearby SNPs, then this approach will not be able to resolve strand issues. Also, if more than one SNP in a region shows strand flips, or if there is a higher level of mis-coding alleles in general, then this approach may indicate that there are problems (many NEG scores above 0) but it might be less clear how to remedy them.

To know which to resolve (cases or controls) one would need to look at the frequency in other panels, or even the correlations, e.g. in HapMap. Ideally, one would only need to do this for a small number of SNPs if any. The --flip and --flip-subset commands described above can then be used to flip the appropriate genotypes.

Finally, the default threshold for counting can be changed by the following command:

```
--flip-scan-threshold 0.8
```

The default is set at 0.5 (i.e. the pair needs to have a correlation of 0.5 or greater in either cases or controls). The number of flanking SNPs with are considered for each index SNP can be modified with the commands

```
--ld-window 10
```

to set the number of SNPs considered upstream and downstream; the maximum physical distance away from the index SNP (1Mb by default) is specified in kb with the command:

```
--ld-window-kb 500
```

# 3.8 Merge two filesets

To merge two PED/MAP files:

```
plink --file data1 --merge data2.ped data2.map --recode --out merge
```

The --merge option must be followed by 2 arguments: the name of the second PED file and the name of the second MAP file. A --recode (or --make-bed, etc) option is necessary to output the newly merged file; in this case, --out option will create the files merge-recode.ped and merge-recode.map.

The --merge option can also be used with binary PED files, either as input or output, but not as the second file: i.e.

```
plink --bfile data1 --merge data2.ped data2.map --make-bed --out merge
```

will create merge.bed, merge.fam and merge.bim, as the --make-bed option was used instead of the --recode option. Likewise, the data1.\* files point to a binary PED file set.

If the second fileset (data2.\*) were in binary format, then you must use --bmerge instead of --merge

```
plink --bfile data1 --bmerge data2.bed data2.bim data2.fam --make-bed --out merge
```

which takes 3 parameters (the names of the BED, BIM and FAM files, in that order).

The two filesets can either overlap completely, partially, or not at all both in terms of markers and individuals. Imputed genotypes will be set to missing (i.e. if SNP\_B is not measured in the first file, but it is in the second, then any individuals in the first file who are not also present in the second file will be set to missing for SNP\_B.

By default, any existing genotype data (i.e. in data1.ped) will not be over-written by data in the second file (data2.ped). By specifying a --merge-mode this default behavior can be changed. The modes are:

- 1 Consensus call (default)
- 2 Only overwrite calls which are missing in original PED file
- 3 Only overwrite calls which are not missing in new PED file
- 4 Never overwrite
- 5 Always overwrite mode
- 6 Report all mismatching calls (diff mode -- do not merge)
- 7 Report mismatching non-missing calls (diff mode -- do not merge)

The default (mode 1) behaviour is to call the merged genotype as missing if the original and new files contain different, non-missing calls; otherwise: i.e.

				Merge mode				
data1.pe	ed ,	data2.ped	->	1	2	3	4	5
0/0	,	0/0	->	0/0	0/0	0/0	0/0	0/0
0/0	,	A/A	->	A/A	A/A	A/A	0/0	A/A
A/A	,	0/0	->	A/A	A/A	A/A	A/A	0/0
A/A	,	A/T	->	0/0	A/A	A/T	A/A	A/T

Modes 6 and 7 effectively provide a means for comparing two PED files – no merging is performed in these cases; rather, a list of mismatching SNPs is written to the file

```
plink.diff
```

They should also report the concordance rate in the LOG file, based on all SNPs that feature in both sets.

A warning will be given if the chromosome and/or physical position differ between the two MAP files.

NOTE Alleles must be exactly coded to match: that is, PLINK will not assume that a 1,2,3,4 SNP coding maps onto a A,C,G,T coding. You can use the --allele1234 and --alleleACGT commands *prior* to merging to convert datasets and then merge these consistently coded files (you cannot convert and merge on the fly, i.e. simply do putting --allele1234 on the command line along with --merge will not work: you need to use --allele1234 and --make-bed first).

## 3.9 Merge multiple filesets

To merge more than two standard and/or binary filesets, it is often more convenient to specify a single file that contains a list of PED/MAP and/or BED/BIM/FAM files and use the --merge-list option. Consider, for an extreme example, the case where each fileset contains only a single SNP, and that there are thousands of these files – this option would help build a single fileset, in this case.

For example, consider we had 4 PED/MAP filesets (labelled fA.\* through fD.\*) and 4 binary filesets, labelled fE.\* through fH.\*). Then using the command

```
plink --file fA --merge-list allfiles.txt --make-bed --out mynewdata would create the binary fileset
```

mynewdata.bed
mynewdata.bim
mynewdata.fam

(alternatively, the --recode option could have been used instead of --make-bed to generate a standard ASCII PED/MAP fileset). In this case, the file allfiles.txt was a list of the to-be-merged files, one set per row:

```
fB.ped fB.map
```

```
fC.ped fC.map
fD.ped fD.map
fE.bed fE.bim fE.fam
fF.bed fF.bim fF.fam
fG.bed fG.bim fG.fam
fH.bed fH.bim fH.fam
```

Important Each fileset must be on a line by itself: lines with two files are interpreted as PED/MAP filesets; lines with three files are interpreted as binary BED/BIM/FAM filesets. The files on a line must always be in this order (PED then MAP; BED then BIM then FAM)

Note In this case the first of the 8 files must be the starting file, i.e. associated with --file on the command line; this file only contains the 8-1 remaining files therefore. The final mynewdata.\* files will contain information from all 8 files.

The --merge-mode option can also be used with the --merge-list option, as described above: however, it is not possible to specify the "diff" features (i.e. modes 6 and 7).

## 3.10 Extract a subset of SNPs: command line options

There are multiple ways to extract just specific SNPs for analysis; this section describes options that use the command-line directly; the next section describes other methods that read a file containing the information.

#### 3.10.1 Based on a single chromosome (--chr)

To analyse only a specific chromosome use

```
plink --file data --chr 6
```

#### 3.10.2 Based on a range of SNPs (--from and --to)

To select a specific range of markers (that must all fall on the same chromosome) use, for example:

```
plink --bfile mydata --from rs273744 --to rs89883
```

#### 3.10.3 Based on single SNP (and window) (--snp and --window)

Alternatively, you can specify a single SNP and, optionally, also ask for all SNPs in the surrounding region, with the --window option:

```
plink --bfile mydata --snp rs652423 --window 20 which extracts only SNPs within \pm 20kb of rs652423.
```

#### 3.10.4 Based on multiple SNPs and ranges (--snps)

Alternatively, the newer --snps command is more flexible but slower than the previously described --snp and --from/--to commands. The --snps command will accept a comma-delimited list of SNPs, including ranges based on physical position. For example,

```
plink --bfile mydata --snps rs273744-rs89883,rs12345-rs67890,rs999,rs222
```

selects the same range as above (rs273744 to rs89883) but also the separate range rs273744 to rs89883 as well as the two individual SNPs rs999 and rs222. Note that SNPs need not be on the same chromosome; also, a range can span multiple chromosomes (the range is defined based on chromosome code order in that

case, as well as physical position, i.e. a range from a SNP on chromosome 4 to one on chromosome 6 includes all SNPs on chromosome 5). No spaces are allowed between SNP names or ranges, i.e. it is

```
--snps rs1111-rs2222,rs3333,rs4444
and not
--snps rs1111 - rs2222, rs3333 ,rs4444
```

Hint As mentioned above, unlike other methods mentioned above, --snps will load in all the data before extracting what it needs, whereas --snp only loads in what it needs, as so is a much faster way to extract a region from a very large dataset: as a result, if you really do want only a single SNP or a single range, use --snp (with --window) or some variant of the from/--to commands.

### 3.10.5 Based on physical position (--from-kb, etc)

One can also select regions based on a window defined in terms of physical distance rather than SNP ID, using the command: e.g.

```
plink --bfile mydata --chr 2 --from-kb 5000 --to-kb 10000
```

to select all SNPs within this 5000kb region on chromosome 2 (when using --from-kb and --to-kb you always need to specify the chromosome with the --chr option).

**HINT** Two alternate forms of the --from-kb command are --from-bp and --from-mb that take a parameter in terms of base-pair position or megabase position, instead of kilobase (to be used with the corresponding --to-bp and --to-mb options).

#### 3.10.6 Based on a set file (--gene)

Finally, if a SET file is also specified, you can use the --gene option to extract all SNPs in that gene/region. For example, if the SET file genes.set contains two genes:

```
GENE1
rs123456
rs10912
rs66222
END
GENE2
rs929292
rs288222
rs110191
END
then
plink --file mydata --set genes.set --gene GENE2 --recode
```

would, for example, create a new dataset with only the 3 SNPs in GENE2.

These options can be used either with standard pedigree files (i.e. using --ped or --file) or with binary format pedigree (BED) files (i.e. using --bfile). One must combine this option with the desired analytic (e.g. --assoc), summary statistic (e.g. --freq) or data-generation (e.g. --make-bed) option.

## 3.11 Extract a subset of SNPs: file-list options

To extract only a subset of SNPs, it is possible to specify a list of required SNPs and make a new file, or perform an analysis on this subset, by using the command

```
plink --file data --extract mysnps.txt
```

where the file is just a list of SNPs, one per line, e.g.

snp005
snp008
snp101

Alternatively, you can use the command --range to modify the behavior of --extract and --exclude. If the --range flag is added, then instead of a list of SNPs, PLINK will expect a list of chromosomal ranges to be given instead, one per line.

```
plink --file data --extract myrange.txt --range
```

All SNPs within that range will then be excluded or extracted. The format of myrange.txt should be, one range per line, whitespace-separated:

```
CHR Chromosome code (1-22, X, Y, XY, MT, 0)
BP1 Start of range, physical position in base units
BP2 End of range, as above
```

For example,

- 2 30000000 35000000 2 60000000 62000000
- X 10000000 20000000

would extract/exclude all SNPs in these three regions (5Mb and 2Mb on chromosome 2 and 10Mb on chromosome X).

One must combine these options with the desired analytic (e.g. --assoc), summary statistic (e.g. --freq) or data-generation (e.g. --make-bed) option.

#### 3.12 Remove a subset of SNPs

To re-write the PED/MAP files, but with certain SNPs excluded, use the option

```
plink --file data --exclude mysnps.txt
```

where the file mysnps.txt is, as for the --extract command, just a list of SNPs, one per line. As described above, the --range command can modify the behaviour of --exclude in the same manner as for --extract.

One must combine this option with the desired analytic (e.g. --assoc), summary statistic (e.g. --freq) or data-generation (e.g. --make-bed) option.

**NOTE** Another way of removing SNPs is to make the physical position negative in the MAP file (this can not be done for binary filesets (e.g. the \*.bim file).

## 3.13 Make missing a specific set of genotypes

To blank out a specific set of genotypes, use the following commands, e.g.

```
--zero-cluster test.zero --within test.clst
```

in conjunction with other data analysis, file generation or summary statistic commands, where the file test.zero is a list of SNPs and clusters, and test.clust is a standard cluster file.

If the original PED file is

1 1 0 0 1 1 A A C C A A 2 1 0 0 1 1 C C A A C C 3 1 0 0 1 1 A C A A A C

```
4 1 0 0 1 1
                   A A C C A A
    5 1 0 0 1 1
                   C C A A C C
                   A C A A A C
    6 1 0 0 1 1
    1b 1 0 0 1 1
                   A A C C A A
    2b 1 0 0 1 1
                   C C A A C C
    3b 1 0 0 1 1
                   A C A A A C
    4b 1 0 0 1 1
                   AA CC AA
    5b 1 0 0 1 1
                   C C A A C C
    6b 1 0 0 1 1
                   A C A A A C
and the MAP file is
    1 snp1 0 1000
    1 snp2 0 2000
    1 snp3 0 3000
and the list of SNPs/clusters to zero out in test.zero is
    snp2
           C1
           C1
    snp3
    snp1
           C2
and the cluster file test.clst is
    1b 1 C1
    2b 1 C1
    3b 1 C1
    4b 1 C1
    5b 1 C1
    6b 1 C1
    2 1 C2
    3 1 C2
then the command
 plink --file test --zero-cluster test.zero --within test.clst --recode
results in a new PED file, plink.ped,
     1 1001 1 A A C C A A
    2 1 0 0 1 1 0 0 A A C C
    3 1 0 0 1 1 0 0 A A A C
    4 1001 1 A A C C A A
    5 1 0 0 1 1 C C A A C C
    6 1 0 0 1 1 A C A A A C
    1b 1 0 0 1 1 A A 0 0 0 0
    2b 1 0 0 1 1 C C 0 0 0 0
    3b 1 0 0 1 1 A C 0 0 0 0
    4b 1 0 0 1 1 A A 0 0 0 0
    5b 1 0 0 1 1 C C 0 0 0 0
    6b 1 0 0 1 1 A C 0 0 0 0
```

i.e. with the appropriate genotypes zeroed out.

**HINT** See the section on handling obligatory missing genotype data, which can often be useful in this context.

#### 3.14 Extract a subset of individuals

To keep only certain individuals in a file, use the option:

```
plink --file data --keep mylist.txt
```

where the file mylist.txt is, as for the --remove command, just a list of Family ID / Individual ID pairs, one set per line, i.e. one person per line. (fields can occur after the 2nd column but they will be ignored – i.e. you could use a FAM file as the parameter of the --keep command, or have comments in the file. For example

```
F101 1
F1001 2_B
F3033 1_A Drop this individual because of consent issues
F4442 22
would be fine.
```

One must combine this option with the desired analytic (e.g. --assoc), summary statistic (e.g. --freq) or data-generation (e.g. --make-bed) option.

### 3.15 Remove a subset of individuals

To remove certain individuals from a file

```
plink --file data --remove mylist.txt
```

where the file mylist.txt is, as for the --keep command, just a list of Family ID / Individual ID pairs, one set per line, i.e. one person per line (although, as for --keep, fields after the 2nd column are allowed but they will be ignored).

One must combine this option with the desired analytic (e.g. --assoc), summary statistic (e.g. --freq) or data-generation (e.g. --make-bed) option.

#### 3.16 Filter out a subset of individuals

Whereas the option to keep or remove individuals are based on files containing lists, it is also possible to specify a filter to include only certain individuals based on phenotype, sex or some other variable.

The basic form of the command is **--filter** which takes two arguments, a filename and a value to filter on, for example:

```
plink --file data --filter myfile.raw 1 --freq
```

implies a file myfile.raw exists which has a similar format to phenotype and cluster files: that is, the first two columns are family and individual IDs; the third column is expected to be a numeric value (although the file can have more than 3 columns), and only individuals who have a value of 1 for this would be included in any subsequent analysis or file generation procedure. e.g. if myfile.raw were

```
F1 I1 2
F2 I1 7
F3 I1 1
F3 I2 1
F3 I3 3
```

then only two individuals (F3 I1 and F3 I2) would be included based on this filter for the calculation of allele frequencies. The filter can be any integer numeric value.

As with --pheno and --within, you can specify an offset to read the filter from a column other than the first after the obligatory ID columns. Use the --mfilter option for this. For example, if you have a binary fileset, and so the FAM file contains phenotype as the sixth column, then you could specify

```
plink --bfile data --filter data.fam 2 --mfilter 4
```

--filter-cases

to select cases only; i.e. cases have the value 2, and this is the 4th variable in the file (i.e. the first two columns are ignored, as these are the ID columns).

Because filtering on cases or controls, or on sex, or on position within the family, will be common operations, there are some shortcut options that can be used instead of --filter. These are

```
--filter-controls
--filter-males
--filter-females
--filter-founders
--filter-nonfounders
These flags can be used in any circumstances, e.g. to make a file of control founders,
plink --bfile data --filter-controls --filter-founders --make-bed --out newfile
or to analyse only males
plink --bfile data --assoc --filter-males
```

**IMPORTANT** Take care when using these with options to merge filesets: the merging occurs **before** these filters.

## 3.17 Create a SET file based on a list of ranges

Given a list of ranges in the following format (4 columns per row; no header file)

```
Chromosome
Start base-pair position
End base-pair position
Set/range/gene name
then the command
plink --file mydata --make-set gene.list
will generate the file
plink.set
```

in the standard set file format. The command --make-set-border takes a single integer argument, allowing for a certain kb window before and after the gene to be included, e.g. for 20kb upstream and downstream:

```
plink --file mydata --make-set gene.list --make-set-border 20
```

HINT See the resources page for pre-compiled RefSeq gene-lists that can be used here.

## Chapter 4

# **Summary statistics**

PLINK will generate a number of standard summary statistics that are useful for quality control (e.g. missing genotype rate, minor allele frequency, Hardy-Weinberg equilibrium failures and non-Mendelian transmission rates). These can also be used as thresholds for subsequent analyses (described in the next section).

Important! All the summary statistics described below are conducted after removing individuals with high missing genotype rates. This option can be set using the --mind option, the default value of which is set at 0.1 (i.e. exclude individuals with more than 10 percent missing genotype data).

## 4.1 Missing genotypes

To generate a list genotyping/missingness rate statistics:

```
plink --file data --missing
```

This option creates two files:

plink.imiss
plink.lmiss

which detail missingness by individual and by SNP (locus), respectively. For individuals, the format is:

FID Family ID IID Individual ID

MISS\_PHENO Missing phenotype? (Y/N) N\_MISS Number of missing SNPs

N\_GENO Number of non-obligatory missing genotypes

F\_MISS Proportion of missing SNPs

For each SNP, the format is:

SNP SNP identifier
CHR Chromosome number

N\_MISS Number of individuals missing this SNP N\_GENO Number of non-obligatory missing genotypes F\_MISS Proportion of sample missing for this SNP

HINT To test for case/control differences in missingness, see the --test-missing option.

**HINT** To produce summary of missingness that is stratified by a categorical cluster variable, use the —within *filename* option as well as —missing. In this way, the missing rates will be given separately for each level of the categorical variable. For example, the categorical variable could be which plate that sample was on in the genotyping. Details on the format of a cluster file can be found here.

## 4.2 Obligatory missing genotypes

Often genotypes might be missing obligatorarily rather than because of genotyping failure. For example, some proportion of the sample might only have been genotyped on a subset of the SNPs. In these cases, one might not want to filter out SNPs and individuals based on this type of missing data. Alternatively, genotypes for specific plates (sets of SNPs/individuals) might have been blanked out with the --zero-cluster option, but you still might want to be able to sensibly set missing data thresholds.

HINT See the section on data management to see how to make missing certain sets of genotypes.

Two functions allow these 'obligatory missing' values to be identified and subsequently handled specially during the filtering steps:

```
plink --bfile mydata --oblig-missing myfile.zero --oblig-clusters myfile.clst --assoc
```

This command applies the default genotyping thresholds (90% per individual and per SNP) but accounting for the fact that certain SNPs are obligatory missing (with the 90% only refers to those SNPs actually attempted, for example). The file specified by --oblig-clusters has the same format as a cluster file (except only a single cluster field is allowed here, i.e. only 3 columns). For example,

```
1 0 0 1 1
               A A C C
                         A A
  1 0 0 1 1
               C
                    A A
                         CC
               A C
  1 0 0 1 1
                    A A
                         A C
  1 0 0 1 1
               A A
                    C C
                         Α Α
  1 0 0 1 1
               C C
                    A A
                         C C
6 1 0 0 1 1
               A C
                    A A
                         A C
1b 1 0 0 1 1
               A A
                    0 0
2b 1 0 0 1 1
               C
                    0 0
                         0 0
3b 1 0 0 1 1
               A C
4b 1 0 0 1 1
               A A
                    0 0
                         0 0
5b 1 0 0 1 1
               C C
                    0 0
6b 1 0 0 1 1
               A C 0 0 0 0
and MAP file test.map
     1 snp1 0 1000
     1 snp2 0 2000
     1 snp3 0 3000
If the obligatory missing file, test.oblig is
     snp2
            C1
     snp3
            C1
```

it implies that SNPs snp2 and snp3 are obligatory missing for all individuals belonging to cluster C1. The corresponding cluster file is test.clst

```
1b 1 C1
2b 1 C1
3b 1 C1
4b 1 C1
5b 1 C1
```

indicating that the last six individuals belong to cluster C1. (Not all individuals need be specified in this file.)

**NOTE** You can have more than one cluster category specified in these files (i.e. implying different patterns of obligatory missing data for different sets of individuals).

Running a --missing command on the basic fileset, ignoring the obligatory missing nature of some of the data, results in the following:

```
plink --file test --missing
```

which shows in the LOG file that 6 individuals were removed because of missing data

.. 40 :-- 1

6 of 12 individuals removed for low genotyping (  $\mbox{MIND} \,>\, 0.1$  )

. . .

and the corresponding output files (plink.imiss and plink.lmiss) indicate no missing data (purely because the six individuals with 2 of 3 genotypes missing were already filtered out and everybody else left happens to have complete genotyping).

FID	IID	MISS_PHENO	$N\_MISS$	$F\_MISS$
1	1	N	0	0
2	1	N	0	0
3	1	N	0	0
4	1	N	0	0
5	1	N	0	0
6	1	N	0	0

and

CHR	SNP	$N\_MISS$	$F\_{MISS}$
1	snp1	0	0
1	snp2	0	0
1	snp3	0	0

In contrast, if the obligatory missing data are specified as follows:

```
plink --file test --missing --oblig-missing test.oblig --oblig-clusters test.clst
```

we now see

. . .

0 of 12 individuals removed for low genotyping (MIND > 0.1)

. . .

and the corresponding output files now include an extra field, N\_GENO, which indicates the number of non-obligatory missing genotypes, which is the denominator for the genotyping rate calculations

FID	IID	MISS_PHENO	$N\_MISS$	$N_{\text{GENO}}$	$F\_MISS$
1	1	N	0	3	0
2	1	N	0	3	0
3	1	N	0	3	0
4	1	N	0	3	0
5	1	N	0	3	0
6	1	N	0	3	0
1b	1	N	0	1	0
2b	1	N	0	1	0
3b	1	N	0	1	0
4b	1	N	0	1	0
5b	1	N	0	1	0
6b	1	N	0	1	0

and

CHR	SNP	${ t N\_MISS}$	$N_{-}GENO$	$F\_MISS$
1	snp1	0	12	0
1	snp2	0	6	0
1	snp3	0	6	0

Seen another way, if one specified --mind 1 to include all individuals (i.e. not apply the default 90% genotyping rate threshold for each individual before this step), then the results would not change with the obligatory missing specification in place, as expected; in contrast, without the specification of obligatory missing data, we would see

FID	IID	MISS_PHENC	N_MIS	S	$F\_MISS$
1	1	1	J	0	0
2	1	1	1	0	0
3	1	1	1	0	0
4	1	ľ	1	0	0
5	1	1	1	0	0
6	1	1	1	0	0
1b	1	1	1	2	0.666667
2b	1	1	1	2	0.666667
3b	1	1	1	2	0.666667
4b	1	1	1	2	0.666667
5b	1	1	1	2	0.666667
6b	1	1	1	2	0.666667
CHR	SNP	$N\_MISS$	F_MISS		
1	snp1	0	0		
1	snp2	6	0.5		

In this not particularly exciting example, there are no missing genotypes that are non-obligatory missing (i.e. that not specified by the two files) – if there were, it would counted appropriately in the above files, and used to filter appropriately also.

**NOTE** All subsequent analyses do not distingush whether genotypes were missing due to failure or were obligatory missing – that is, this option only effects the behavior of the --mind and --geno filters.

**NOTE** If a genotype is set to be obligatory missing but actually in the genotype file it is not missing, then it will be set to missing and treated as if missing.

## 4.3 Cluster individuals based on missing genotypes

Systematic batch effects that induce missingness in parts of the sample will induce correlation between the patterns of missing data that different individuals display. One approach to detecting correlation in these patterns, that might possibly idenity such biases, is to cluster individuals based on their *identity-by-missingness* (IBM). This approach use exactly the same procedure as the IBS clustering for population stratification, except the distance between two individuals is based not on which (non-missing) allele they have at each site, but rather the proportion of sites for which two individuals are both missing the same genotype.

To use this option:

and

1 snp3

```
plink --file data --cluster-missing
which creates the files:
    plink.matrix.missing
    plink.cluster3.missing
```

which have similar formats to the corresponding IBS clustering files. Specifically, the plink.mdist.missing file can be subjected to a visualisation technique such as multidimensinoal scaling to reveal any strong systematic patterns of missingness.

**Note** The values in the .mdist file are distances rather than similarities, unlike for standard IBS clustering. That is, a value of 0 means that two individuals have the same profile of missing genotypes. The exact value represents the proportion of all SNPs that are discordantly missing (i.e. where one member of the pair is missing that SNP but the other individual is not).

The other constraints (significance test, phenotype, cluster size and external matching criteria) are not used during IBM clustering. Also, by default, all individuals and all SNPs are included in an IBM clustering analysis, unlike IBS clustering, i.e. even individuals or SNPs with very low genotyping, or monomorphic alleles. By explicitly specifying --mind or --geno or --maf certain individuals or SNPs can be excluded (although the default is probably what is usually required for quality control procedures).

## 4.4 Test of missingness by case/control status

To obtain a missing chi-sq test (i.e. does, for each SNP, missingness differ between cases and controls?), use the option:

The actual counts of missing genotypes are available in the plink.lmiss file, which is generated by the --missing option.

**Note** This test is only applicable to case/control data.

## 4.5 Haplotype-based test for non-random missing genotype data

The previous test asks whether genotypes are missing at random or not with respect to phenotype. This test asks whether or not genotypes are missing at random with respect to the true (unobserved) genotype, based on the observed genotypes of nearby SNPs.

**Note** This test assumes dense SNP genotyping such that flanking SNPs are typically in LD with each other. Also bear in mind that a negative result on this test may simply reflect the fact that there is little LD in the region.

This test works by taking a SNP at a time (the 'reference' SNP) and asking whether haplotype formed by the two flanking SNPs can predict whether or not the individual is missing at the reference SNP. The test is a simple haplotypic case/control test, where the phenotype is missing status at the reference SNP. If missingness at the reference is not random with respect to the true (unobserved) genotype, we may often expect to see an association between missingness and flanking haplotypes.

Note Again, just because we might not see such an association does not necessarily mean that genotypes are missing at random – this test has higher specificity than sensitivity. That is, this test will miss a lot; but, when used as a QC screening tool, one should pay attention to SNPs that show highly significant patterns of non-random missingness.

This option is run with the command:

```
plink --file data --test-mishap
```

which generates an output file called

plink.missing.hap

which has the fields

LOCUS	Reference SNP
HAPLOTYPE	Flanking haplotype, or heterozygosity
F_0	Frequency of HAPLOTYPE if missing reference SNP
F_1	Frequency of HAPLOTYPE if not missing reference ${\tt SNP}$
M_H1	N missing/not missing for HAPLOTYPE
M_H2	N missing/not missing for not-HAPLOTYPE
CHISQ	Chisquare test for non-random missingness
P	Asymptotic p-value
SNPS	Identifier for flanking SNPs

The HAPLOTYPE typically represents each two-SNP flanking haplotype (i.e. not including the reference SNP itself); each reference SNP will also have a row labelled HETERO in this column, which means we are testing whether or not being heterozygous for the flanking haplotypes (which would, under many sets of haplotype frequencies, increase the chance of being heterozygous for the reference SNP). SNPs with no or very little missing genotype data are skipped. Only haplotypes above the <code>--maf</code> threshold are used in analysis.

Here is an example from real data (rows split into two sets for clarity):

LOCUS	HAPLOTYPE	F_0	$F_{-}1$	M_H1	M_H2
rs17012390	CT	0.5238	0.01949	55/104	50/5233
rs17012390	TC	0.4762	0.9805	50/5233	55/104
rs17012390	HETERO	1	0.04252	56/114	0/2567
LOCUS	HAPLOTYPE	CHISQ	P	SNPS	
rs17012390	CT	923.4	0	rs17012387 rs1	7012393
rs17012390	TC	923.4	0	rs17012387 rs1	7012393
rs17012390	HETERO	863.3	0	rs17012387 rs1	7012393

This clearly shows a huge chi-square (the sample is large, N of over 2500 individuals). We see that of 56 missing genotypes for this reference SNP, all occur when the flanking haplotypic background is heterozygous (i.e. M.H1 shows 56/114, indicating that there are 114 other instances of a heterozygous haplotypic background when the reference SNP is not missing) whereas we see not a single missing call when the flanking SNP background is homozygous, of which we see 2567 observations. This is clearly indicative of non-random association between the unobserved genotype and missing status.

Looking at the same data a different way, F\_1 indicates that the majority of the sample (people not missing at the reference SNP) have haplotype frequencies of CT and TC haplotypes at approximately 0.02 and 0.98 respectively). In contrast, because all people missing this SNP are on heterozygous backgrounds, these frequencies become approximately 50:50 in this group (shown in F\_0).

In the particular dataset this example comes from, this SNP would have passed a standard quality control test. The --hardy command shows that this SNP does not failure the HWE test; also, it does not show excessive amounts of missing data (the --missing command indicates a missing rate of 0.021). The genotype counts (obtained by the --hardy option) are, for the whole sample, 0/104/2584.

In contrast, here are the same results for a different SNP that does not show any evidence of non-random missingness.

LOCUS	HAPLOTYPE	$F_0$	F_1	$M_H1$	$M_H2$
rs3912752	CC	0.07692	0.06507	2/354	24/5086
rs3912752	TT	0.1154	0.205	3/1115	23/4325
rs3912752	CT	0.8077	0.73	21/3971	5/1469
rs3912752	HETERO	0.2308	0.4279	3/1164	10/1556
LOCUS	HAPLOTYPE	CHISQ	Р	SNPS	
rs3912752	CC	0.05967	0.807	rs3912751 rs	351596

rs3912752	TT	1.276	0.2586	rs3912751 rs351596
rs3912752	CT	0.7938	0.3729	rs3912751 rs351596
re3912752	HETERO	2 056	0 1516	re3912751   re351596

Here we do not see any deviation between the flanking haplotype frequencies between people missing versus genotyped for the reference SNP. Of course, there is less missingness for this SNP (26 missing genotypes) so we might expect power is lower, even if there were non-random missingness. This only highlights the point made above that, in general, significant results are more interpretable than non-signficant results for this test. But more importantly, if there are only a handful of missing genotypes, we do not particular care whether or not they are missing at random, as they would not bias the association with disease in any case. Of course, whether there is non-random genotyping error is another question...

By default, we currently just select exactly two flanking SNPs. This can be changed with the option --mishap-window. For example,

```
plink --bfile mydata --test-mishap --mishap-window 4
```

Future releases will feature a more intelligent selection of flanking markers.

Note This routine currently skips the SNPs on the X and Y chromosomes.

## 4.6 Hardy-Weinberg Equilibrium

To generate a list of genotype counts and Hardy-Weinberg test statistics for each SNP, use the option:

```
plink --file data --hardy
```

which creates a file:

plink.hwe

This file has the following format

SNP	SNP identifier
TEST	Code indicating sample
A1	Minor allele code
A2	Major allele code
GENO	Genotype counts: 11/12/22
O(HET)	Observed heterozygosity
E(HET)	Expected heterozygosity
P	H-W p-value

For case/control samples, each SNP will have three entries (rows) in this file, with TEST being either ALL, AFF (cases only) or UNAFF (controls only). For quantitative traits, only a single row will appear for each SNP, labelled ALL(QT).

Only founders are considered for the Hardy-Weinberg calculations – ie. for family data, any offspring are ignored.

**WARNING** By default, this procedure only considers founders, so no HW results would be given for sibling-only datasets (i.e. if no parents exist). To perform a rough, somewhat biased test, use the **--nonfounders** option which means that all individuals will be included. Alternatively, manually extract one person per family for this calculation and recode these individuals as founders (see the **--keep** option to facilitate this).

The default test is an exact one, described and implemented by Wigginton *et al* (see reference below), which is more accurate for rare genotypes. You can still perform the standard asymptotic test with the --hardy2 option.

```
A Note on Exact Tests of Hardy-Weinberg Equilibrium. Wigginton JE, Cutler DJ and Abecasis GR Am J Hum Genet (2005) 76: 887-93
```

## 4.7 Allele frequency

To generate a list of minor allele frequencies (MAF) for each SNP, based on all founders in the sample:

```
plink --file data --freq
will create a file:
     plink.frq
with five columns:
     CHR
              Chromosome
     SNP
              SNP identifier
     Α1
              Allele 1 code (minor allele)
     A2
              Allele 2 code (major allele)
     MAF
              Minor allele frequency
     NM
              Non-missing allele count
```

**HINT** To produce summary of allele frequencies that is stratified by a categorical cluster variable, use the --within *filename* option as well as --missing. In this way, the frequencies will be given separately for each level of the categorical variable. Details on the format of a cluster file can be found here.

**NOTE** If a SNP fails the genotyping rate threshold (as set by the --geno value, which is by default 0.10) the frequency will appear as NA in the plink.frq output file. To obtain frequencies on all SNPs irrespective of genotyping rate, set --mind 1.

## 4.8 Linkage disequilibrium based SNP pruning

Sometimes it is useful to generate a pruned subset of SNPs that are in approximate linkage equilibrium with each other. This can be achieved via two commands: --indep which prunes based on the *variance inflation factor* (VIF), which recursively removes SNPs within a sliding window; second, --indep-pairwise which is similar, except it is based only on pairwise genotypic correlation.

**Hint** The output of either of these commands is two lists of SNPs: those that are pruned out and those that are not. A separate command using the --extract or --exclude option is necessary to actually perform the pruning.

The VIF pruning routine is performed:

```
plink --file data --indep 50 5 2
will create files
    plink.prune.in
    plink.prune.out
```

Each is a simlpe list of SNP IDs; both these files can subsequently be specified as the argument for a --extract or --exclude command.

The parameters for --indep are: window size in SNPs (e.g. 50), the number of SNPs to shift the window at each step (e.g. 5), the VIF threshold. The VIF is 1/(1-R2) where R2 is the multiple correlation coefficient for a SNP being regressed on all other SNPs simultaneously. That is, this considers the correlations between SNPs but also between linear combinations of SNPs. A VIF of 10 is often taken to represent near collinearity problems in standard multiple regression analyses (i.e. implies R2 of 0.9). A VIF of 1 would imply that the SNP is completely independent of all other SNPs. Practically, values between 1.5 and 2 should probably be used; particularly in small samples, if this threshold is too low and/or the window size is too large, too many SNPs may be removed.

The second procedure is performed:

```
plink --file data --indep-pairwise 50 5 0.5
```

This generates the same output files as the first version; the only difference is that a simple pairwise threshold is used. The first two parameters (50 and 5) are the same as above (window size and step); the third parameter represents the r2 threshold. Note: this represents the pairwise SNP-SNP metric now, not the multiple correlation coefficient; also note, this is based on the genotypic correlation, i.e. it does not involve phasing.

To give a concrete example: the command above that specifies 50 5 0.5 would a) consider a window of 50 SNPs, b) calculate LD between each pair of SNPs in the window, b) remove one of a pair of SNPs if the LD is greater than 0.5, c) shift the window 5 SNPs forward and repeat the procedure.

To make a new, pruned file, then use something like (in this example, we also convert the standard PED fileset to a binary one):

```
plink --file data --extract plink.prune.in --make-bed --out pruneddata
```

#### 4.9 Mendel errors

To generate a list of Mendel errors for SNPs and families, use the option:

```
plink --file data --mendel
which will create files:
    plink.mendel
    plink.imendel
    plink.fmendel
    plink.lmendel
```

The \*.mendel file contains all Mendel errors (i.e. one line per error); the \*.imendel file contains a summary of per-individual error rates; the \*.fmendel file contains a summary of per-family error rates; the \*.lmendel file contains a summary of per-SNP error rates.

The \*.mendel file has the following columns:

```
FID Family ID

KID Child individual ID

CHR Chromosome

SNP SNP ID

CODE A numerical code indicating the type of error (see below)

ERROR Description of the actual error
```

The error codes are as follows:

$\mathbf{Code}$	Pat	,	Mat	-;	Offspring	5
1	AA	,	AA	->	AB	
2	BB	,	BB	->	AB	
3	BB	,	**	->	AA	
4	**	,	BB	->	AA	
5	BB	,	BB	->	AA	
6	AA	,	**	->	BB	
7	**	,	AA	->	BB	
8	AA	,	AA	->	BB	
9	**	,	AA	->	BB	(X chromosome male offspring)
10	**	,	BB	->	AA	(X chromosome male offspring)

The \*.lmendel file has the following columns:

```
CHR Chromosome
SNP ID
N Number of Mendel errors for this SNP
```

The  $\star$ .imendel file has the following columns:

```
FID Family ID

IID Individual ID

N Number of errors this individual was implicated in
```

The following heurtistic is used to provide a rough estimate of Mendel error rare 'per individual': error types 1 and 2 count for all 3 individuals (child, father, mother); error types 5 and 8 count only for the child (i.e. otherwise requires two errors, one in each parent); error types 3 and 6 count for the child and the father; all other types (4, 7, 9 and 10) count for the offspring and the mother. This metric might indicate that, for example, in a nuclear family with two parents and two offspring, many more Mendel errors can be associated with the first sibling; the remaining trio might not show any increased rate.

Currently, PLINK only scans full trios for Mendel errors. Families with fewer than 2 parents in the dataset will not be tested.

Finally, the \*.fmendel file has the following columns:

```
FID Family ID

PAT Paternal individual ID

MAT Maternal individual ID

CHLD Number of offspring in this (nuclear) family

N Number of Mendel errors for this (nuclear) family
```

#### 4.10 Sex check

This option uses X chromosome data to determine sex (i.e. based on heterozygosity rates) and flags individuals for whom the reported sex in the PED file does not match the estimated sex (given genomic data). To run this analysis, use the flag:

```
plink --bfile data --check-sex

which generates a file
    plink.sexcheck

which contains the fields

FID Family ID

IID Individual ID

PEDSEX Sex as determined in pedigree file (1=male, 2=female)

SNPSEX Sex as determined by X chromosome

STATUS Displays "PROBLEM" or "OK" for each individual

F The actual X chromosome inbreeding (homozygosity) estimate
```

A PROBLEM arises if the two sexes do not match, or if the SNP data or pedigree data are ambiguous with regard to sex. A male call is made if F is more than 0.8; a femle call is made if F is less than 0.2.

The command

```
plink --bfile data --impute-sex --make-bed --out newfile
```

will impute the sex codes based on the SNP data, and create a new file with the revised assignments, in this case a new binary fileset.

## 4.11 Pedigree errors

PLINK can accept multigenerational family data for family-based tests and Mendel error checks. It will break multigenerational families down into nuclear family units where appropriate. Extended family information is not used in an optimal manner, however (e.g. to help find Mendel errors using grandparental genotypes if parental genotypes are missing).

Unless PLINK is explicitly told to perform a family-based analysis, it will ignore any pedigree structure in the sample and analyse the data as if all individuals are unrelated (i.e. the --assoc option, for example, will ignore family structure). It is therefore the responsibility of the user to ensure that the data are appropriate for the type of test (e.g. if performing a standard association test with --assoc, this implies that all individuals should be unrelated for asymptotic significance values to be correct). The exception to this general rule is that certain summary statistics are based only on founders.

PLINK will spot most pedigree errors (e.g. if an individual has two fathers, for example). For a more comprehensive evaluation of pedigree errors (invalid or incompletely specified pedigree structures) please use a different software package such as PEDSTATS or famtypes http://pngu.mgh.harvard.edu/purcell/famtypes/.

## Chapter 5

# Inclusion thresholds

This section describes options that can be used to filter out individuals or SNPs on the basis of the summary statistic measures described in the previous summary statistics page.

#### 5.0.1 Summary statistics versus inclusion criteria

The following table summarizes the relationship between the commands to generate summary statistics (as described on the previous page, versus the commands to exclude individuals and/or markers, which are described on this page.

 Feature
 As summary statistic
 As inclusion criteria

 Missingness per individual
 --missing
 --mind N

 Missingness per marker
 --missing
 --geno N

 Allele frequency
 --freq
 --maf N

 Hardy-Weinberg equilibrium
 --hardy
 --hw N

 Mendel error rates
 --mendel
 --me N M

#### 5.0.2 Default threshold values

By default, PLINK does not impose any filters on minor allele frequency or genotyping rate. (Note that versions prior to 1.04 use to have thresholds of 0.01 for frequency and 0.1 for individual and SNP missing rate – this is no longer the case, i.e. it is as if the --all keyword is always specified).

To perform an analysis, or generate a new dataset, with filters applied, add the --mind, --geno or --maf options are to the command line, for example, when the --remove command is given.

## 5.1 Missing rate per person

The initial step in all data analysis is to exclude individuals with too much missing genotype data. This option is set as follows:

```
plink --file mydata --mind 0.1
```

which means exclude with more than 10% missing genotypes (this is the defalt value). A line in the terminal output will appear, indicating how many individuals were removed due to low genotyping. If any individuals were removed, a file called

```
plink.irem
```

will be created, listing the Family and Individual IDs of these removed individuals. Any subsequent analysis also specified on the same command line will be performed without these individuals.

One might instead wish to create a new PED file with these individuals permanently removed, simply add an option to generate a new fileset: for example,

```
plink --file data --mind 0.1 --recode --out cleaned
```

```
will generate files
```

cleaned.ped
cleaned.map

with the high-missing-rate individuals removed; alternatively, to create a binary fileset with these individuals removed:

```
plink --file data --mind 0.1 --make-bed --out cleaned
which results in the files
    cleaned.bed
    cleaned.bim
    cleaned.fam
```

**HINT** You can specify that certain genotypes were never attempted, i.e. that they are obligatory missing, and these will be handled appropriately by these genotyping rate filters. See the summary statistics page for more details.

## 5.2 Allele frequency

Once individuals with too much missing genotype data have been excluded, subsequent analyses can be set to automatically exclude SNPs on the basis of MAF (minor allele frequency):

```
plink --file mydata --maf 0.05
```

means only include SNPs with MAF i = 0.05. The default value is 0.01. This quantity is based only on founders (i.e. individuals for whom the paternal and maternal individual codes and both 0).

This option is appropriately counts alleles for X and Y chromosome SNPs.

## 5.3 Missing rate per SNP

Subsequent analyses can be set to automatically exclude SNPs on the basis of missing genotype rate, with the --geno option: the default is to include all SNPS (i.e. --geno 1). To include only SNPs with a 90% genotyping rate (10% missing) use

```
plink --file mydata --geno 0.1
```

As with the <code>--maf</code> option, these counts are calculated after removing individuals with high missing genotype rates.

## 5.4 Hardy-Weinberg Equilibrium

To exclude markers that failure the Hardy-Weinberg test at a specified significance threshold, use the option:

```
plink --file mydata --hwe 0.001
```

By default this filter uses an exact test (see this section). The standard asymptotic (1 df genotypic chi-squared test) can be requested with the --hwe2 option instead of --hwe.

The following output will appear in the console window and in plink.log, detailing how many SNPs failed the Hardy-Weinberg test, for the sample as a whole, and (when PLINK has detected a disease phenotype) for cases and controls separately:

```
Writing Hardy-Weinberg tests (founders-only) to [ plink.hwe ] 30 markers failed HWE test ( p <= 0.05 ) and have been excluded 34 markers failed HWE test in cases
```

#### 30 markers failed HWE test in controls

This test will only be based on founders (if family-based data are being analysed) unless the --nonfounders option is also specified. In case/control samples, this test will be based on controls only, unless the --hwe-all option is specified, in which case the phenotype will be ignored. This can be important if parents are coded as missing in an affected offspring trio sample.

Please refer to the --hardy option for more details on producing summary statistics of all HWE rates.

#### 5.5 Mendel error rate

For family-based data only, to exclude individuals and/or markers on the basis on Mendel error rate, use the option:

```
plink --file mydata --me 0.05 0.1
```

where the two parameters are:

- the first parameter determines that families with more than 5% Mendel errors (considering all SNPs) will be discarded.
- the second parameter indicates that SNPs with more than 10% Mendel error rate will be excluded (i.e. based on the number of trios);

Please refer to the summary statistics page for more details on generating summary statistics for Mendel error rates.

Note Currently, PLINK calculates the per SNP Mendel error rates at the same time as the per family error rates. In future releases, this may change such that the per family error rate is calculated after SNPs failing this test have been removed. Also, using this command currently removes entire nuclear families on the basis of high Mendel error rates: it will often be more appropriate to remove particular individuals (e.g. if a second sibling shows no Mendel errors). For this more fine-grained procedure, use the --mendel option to generate a complete enumeration of error rates by family and individual and exclude individuals as desired. Finally, it is possible to zero out specific Mendelian inconsistencies with the option --set-me-missing. This should be used in conjunction with a data generation command and the --me option. Specifically, the --me parameters should be both to 1, in order not to exclude any particular SNP or individual/family, but instead to zero out only specific genotypes with Mendel errors and save the dataset as a new file. (Both parental and offspring genotypes will be set to missing.)

```
plink --bfile mydata --me 1 1 --set-me-missing --make-bed --out newdata
```

## Chapter 6

# Population stratification

PLINK offers a simple but potentially powerful approach to population stratification, that can use whole genome SNP data (the number of individuals is a greater determinant of how long it will take to run). We use complete linkage agglomerative clustering, based on pairwise identity-by-state (IBS) distance, but with some modifications to the clustering process: restrictions based on a significance test for whether two individuals belong to the same population (i.e. do not merge clusters that contain significantly different individuals), a phenotype criterion (i.e. all pairs must contain at least one case and one control) and cluster size restrictions (i.e. such that, with a cluster size of 2, for example, the subsequent association test would implicitly match every case with its nearest control, as long as the case and control do not show evidence of belonging to different populations). In addition, external matching criteria can be specified, to match on age and sex, for example, as well as genetic information. Any evidence of population substructure (from this or any other analysis) can be incorporated in subsequent association tests via the specification of clusters.

All these analyses require genome-wide coverage of autosomal SNPs!

## 6.1 IBS clustering

To perform complete linkage clustering of individuals on the basis of autosomal genome-wide SNP data, the basic command is:

```
plink --file mydata --cluster
which generates four output files:
    plink.cluster0
    plink.cluster1
    plink.cluster2
    plink.cluster3
```

that contain similar information but in different formats. The

The \*.cluster0 file contains some information on the clustering process. This file can be safely ignored by most users.

The \*.cluster1 file contains information on the final solution, listed by cluster: e.g. for 4 individuals with the following Family and Individual IDs

- A 1 B 1
- C 1 D 1

we see 3 clusters, one line of output per cluster:

 $0 A_{-1}$ 

```
1 B<sub>-</sub>1 C<sub>-</sub>1
```

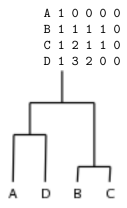
2 D\_1

(note how family and individuals IDs are concatenated with the underscore character in the output)

The \*.cluster2 file contains the same information but listed one line per individual: the three columns are family ID, individual ID and assigned cluster:

- A 1 0 B 1 1
- C 1 1
- D 1 2

The \*.cluster3 file is in the same format as cluster2 (one line per individual) but contains all solutions (i.e. every step of the clustering from moving from N clusters each of 1 individual (leftmost column after family and individual ID) to 1 cluster (labelled 0) containing all N individuals (the final, rightmost column): also shown is the dendrogram this represents: e.g.



**NOTE** If any constraints have been placed upon the clustering, then solutions represented in the \*.cluster3 file may not go as far as 1 cluster with all N individuals: in this case, the file \*.cluster2 will contain the final solution (i.e. as far as the clustering could go before running up against constraints, e.g. based on maximum cluster size, etc).

HINT! In large samples, cluster analyses can be very slow. Often the most time consuming step is calculating the pairwise IBS metrics: these only need to be calculated once however, even if you want to run the cluster analysis multiple times (e.g. with different constraints). This is achieved with the --read-genome option, assuming you have previously run the --genome command. It is a good idea to not impose a threshold of the --genome output in this case. For example:

```
plink --bfile mydata --genome --out mydata
```

followed by multiple clustering commands (see below for descriptions of the cluster constraint parameters used here)

```
plink --bfile mydata --read-genome mydata.genome --cluster --ppc 0.01
and
plink --bfile mydata --read-genome mydata.genome --cluster --mc 2 --ibm 0.01
etc.
```

ADVANCED HINT! In very large samples, cluster analyses can be very, very slow. When calculating the plink.genome file (as described above), if you have access to a cluster of computers for parallel computing, you can use the following approach to greatly reduce the time this step takes. In this case, we will assume you are familiar with and using a Linux operating system, and that the bsub prefix is used to send a job

to the cluster – obviously, change the script below as appropriate. This uses the --genome-lists option to calculate IBS statistics for only a subset of the sample at a time. If the binary fileset is data.\* then create multiple lists of, for example, 100 individuals per list

```
gawk 'print $1,$2' data.fam | split -d -a 3 -l 100 - tmp.list
```

If this creates, for example, 39 separate files (labelled 0 to 38), then run these in all unquie pairwise combinations in parallel with something like the following script: (i.e. edit the first line as appropriate)

Note the use of the --all command to ensure that the exact same list of SNPs is used for each subset of the sample in calculating the IBS/IBD scores; any pruning (e.g. for LD, or data quality) should therefore be performed prior to this step.

The finally, concatenate these individual files back into one, taking care to get only a single header line (assuming you have no individuals with FID1 in their ID...)

```
head -n1 data.sub.0.0.genome > header
cat data.sub*genome | fgrep -v FID1 | cat header - > data.genome
rm tmp.list*
rm data.sub.*
```

**NOTE** If you use this approach to calculate the IBD probabilities, then you should first perform --freq on the whole dataset, then add the line --read-freq plink.frq (obviously replacing the filename with your file) to make sure that everybody has the sample frequencies used in the IBD calculations.

## 6.2 Permutation test for between group IBS differences

Given that pairwise IBS distances between all individuals have been calculated, we can asked whether or not there are group differences in this metric, with respect to a binary phenotype. The command

```
./plink --bfile mydata --ibs-test
or, if an appropriate plink.genome file has already been created,
./plink --bfile mydata --read-genome plink.genome --ibs-test
```

will permute case/control label, and then recalculate several between-group metrics based on average IBS within that group. This command uses a fixed 10,000 permutations.

All results are written to the LOG file. First, the observed means and standard deviation of each of the 3 groups (case/control, case/case and control/control, in that order) will be displayed: e.g.

```
Between-group IBS (mean, SD) = 0.782377, 0.00203459
In-group (2) IBS (mean, SD) = 0.782101, 0.00232296
In-group (1) IBS (mean, SD) = 0.78273, 0.00170816
```

Then 12 separate tests are presented, which have self-explanatory names. If the label does not explicitly mention a comparison pair-type, it implies that the first pair type is being compared to the other two pair-types.

```
T1: Case/control less similar
                                                    p = 0.97674
 T2: Case/control more similar
                                                    p = 0.0232698
 T3: Case/case less similar than control/control
                                                    p = 0.00285997
 T4: Case/case more similar than control/control
                                                    p = 0.99715
                                                    p = 0.00430996
 T5: Case/case less similar
 T6: Case/case more similar
                                                    p = 0.9957
                                                    p = 0.99883
 T7: Control/control less similar
                                                    p = 0.00117999
 T8: Control/control more similar
                                                    p = 0.00726993
 T9: Case/case less similar than case/control
T10: Case/case more similar than case/control
                                                    p = 0.99274
T11: Control/control less similar than case/control p = 1
```

T12: Control/control more similar than case/control p = 9.9999e-06

For the purpose of stratification effects between cases and conrtols, the test T1 is probably most appropriate, as it directly asks whether or not, on average, an individual is less similar to another phenotypicallydiscordant individual than would be expected by chance (i.e. if we randomized phenotype labels). That is, to the extent that cases and controls are from two separate populations, you would expect pairs within a phenotype group to be more similar than pairs across the two groups, i.e. T1. Of course, the opposite could also be true (tested by T2), which would probably represent certain ascertainment procedures (i.e. taking this to an extreme, imagine a discordant sibling pair design: case/control pairs would on average be more similar than case/case and control/control pairs).

The other tests are provided for completeness and give a more general description of the variability between and within each group. The general pattern shown above would suggest that there is relatively more variability within the case sample than the control sample. Bear in mind when interpreting the empirical p-values that the relative sizes of case and control samples will have an impact on the exact p-value (i.e. these significance tests should not be taken to directly represent the magnitude of differences between groups).

Note This test assumes that individuals have a disease phenotype; obviously, one could swap in other labels (e.g. site of collection) via the --pheno command, as long as they are dichotomous.

#### 6.3 Constraints on clustering

This section describes the extra constraints that can be placed on the clustering procedure, specified via other options in addition to the --cluster option. As further described in the association analysis and permutation sections, these options can be used to set up various types of analyses that control for potential stratification in the sample.

#### 1) Based on pairwise population concordance (PPC) test:

This is a simple significance test for whether two individuals belong to the same random-mating population. To only merge clusters that do not contain individuals differing at a certain p-value:

```
--ppc 0.0001
```

**NOTE** This command has been changed from --pmerge in older versions of PLINK (pre 0.99n).

This test is based on the observed binomial proportion of IBS 0 loci pairs to IBS 2 het/het pairs: counts of these two types should be in the ratio of 1:2 if the two individuals come from the same population. The significant p-value indicates fewer IBS2 het/het loci than expected (based on normal approximation to binomial). These tests are also given by the --genome command.

WARNING! Unlike the basic IBS clustering, which places no restrictions on the SNPs that can be used in the analysis, this test assumes that the SNPs used are in linkage equilibrium. By default, this test will only count an 'informative' SNP pair (i.e. one that, for a particular pair of individuals, has two of each allele) as either an IBS 0 or IBS 2 count for this test (the HOMHOM and HETHET counts from the --genome option) if it is more than 500 kb more the previous informative pair of SNPs, for that particular pair of individuals. This gap parameter can be changed with the option

which would, in this case, reduce that gap to 100kb. (Note: all SNPs will still be used to calculate the main IBS distance metric, upon which the clustering is based).

HINT Also, this test is susceptible to non-random missingness in genotypes, particularly if heterozygotes are more likely to be dropped. It is therefore good practice to set the --geno very low for this analysis, i.e. so only SNPs with virtually complete genotyping are included.

#### 2) Based on phenotype:

To ensure that every cluster has at least one case and one control:

--cc

#### 3) Based on maximum cluster size:

To set the maximum cluster size to a certain value, e.g. 2:

--mc 2

Alternatively, to specify a maximum number of cases and a maximum number of controls per cluster, use the option:

--mcc 1 3

which, in this case, specifies that each cluster can have up to 1 case and 3 controls. Note the different syntax: -mcc as opposed to --mc. Using this in conjunction with the --cc constraint (that ensures at least 1 case and 1 control per cluster) this is an easy way to achieve a certain matching scheme, say 1 case to 3 controls, or 2 cases to 2 controls, etc.

#### 4) Based on fixed number of clusters:

To request that the clustering process stops at a certain fixed number of clusters, for example, a 2 cluster solution, use:

--K 2

**Note** If other clustering constraints are in place, it is possible that clustering may stop *before* reaching the specified number of clusters with the --K option; if other constraints are specified, you can think of this as stating the *minimum* number of clusters possible.

#### 5) Based on pattern of missing genotype data:

To only cluster individuals with sufficiently similar profiles of missing genotype data (genome-wide) use the option:

--ibm 0.02

which would only match people if they are discordantly missing (i.e. one person is missing a particular SNP but the other person is not) for 2 percent of the genome or less. Another way to incorporate missingness would be by defining overall call rate per individual as an external quantitative matching criteria (see below); this approach is preferrable however (as it does not match just on average rate, but also on whether it tends to be the same SNPs that are missing).

#### 6) Based on user-specified external matching criteria:

To use external matching criteria: for categorical matching criteria, use the option:

#### --match mydata.match

where the file mydata.match contains the following columns: family and individual ID and the one or more matching variables, one row per person:

```
Family ID
Individual ID
Matching criteria 1
Matching criteria 2
...
Matching criteria N
```

The default behavior is that only individuals with the same matching criteria across all the measures will be paired to make clusters. For example, if the file were:

```
F1 I1 1 1 1
F2 I2 1 2 1
F3 I3 2 2 2
F4 I4 1 2 1
F5 I5 1 1 1
```

then only F1/I1 and F5/I5 could be paired; also F2/I2 and F4/I4 could be paired. No other combinations of pairings would be possible. Therefore, no cluster would ever be formed that contained both individuals F1/I1 and F2/I2, for example.

One application of this option would be to ensure that individuals are sex-matched, or matched on some relevant environmental exposure, in addition to the genetic IBS matching.

It is possible to adjust the default behaviour to consider two individuals as potentially 'pairable' is they differ on a particular categorical criteria. This is achieved with the optional command:

```
--match-type mydata.bt
```

where mydata.bt is the name of a file that contains a series of 0s and 1s (or "-" and "+" characters), whitespace delimited, that indicate whether a criteria should be a "postive match" (i.e. two individuals are potentially pairable only if they have the *same* values for this variable) or a "negative match" (i.e. two individuals are potentially pairable only if they have *different values* for this variable). In the above example, if the file mydata.bt were

```
+ - +
```

then the following pairs are potentially pairable:

```
F1/I1 and F2/I2
F1/I1 and F4/I4
F5/I5 and F2/I2
F5/I5 and F4/I4
```

i.e. F1/I1 can no longer be paired with F5/I5 because they have the same value for the second matching variable, which is now a negative match criteria.

**Note** In this example, the matching variables only took two values: in practice, one can have any number of categories per matching variable.

Note Missing variables can be specified for matching variables – this means that the criteria will be ignored. As all pairs start out as potentially pairable, this means that missing matching criteria data will never be used to make a pair unpairable.

A second form a matching is based on quantitative traits – in this case, a maximum difference threshold is specified for each measure, such that individuals will not be matched if they differ beyond the threshold on the quantitative traits. This is achieved by the following options:

```
--qmatch mydata.match --qt mydata.qt
```

Note that a second --qt option is necessary as well as the --qmatch option. The --qt specifies a file that contains the thresholds, e.g. for 3 external quantitative criteria, this should contain 3 values:

```
5
0.333
120
```

The --qmatch should then contain the same number of quantitative matching criteria per person (again, one row per person):

```
F1 I1
         27
              -0.23
                       1003
F2 I2
               2.22
                       1038
         34
F3 I3
         45
               1.99
                        987
F4 I4
         19
             -9
                       2374
F5 I5
         18
             -0.45
                        996
```

In this case, for example, for the first measure only F4/I4 and F5/I5 are pairable, as —19-18— is not more than 5. This measure might represent age, for example. This pair is not matchable on the basis on the third metric, however, as —2374-996— ¿ 120. As such, no pairs could be formed between any of these five individuals, in this particular case. Note that individual is actually missing (default --missing-phenotype value is -9) for the second criterion: see below for a description of how missing data are handled in this context.

The .match and .qmatch files do not need to contain all individuals and do not need to be in the same order as the original PED files. Any individuals in these files who are not in the original files will be ignored.

Missing phenotypes are simply ignored (i.e. two individuals would not be called non-matching if either one or both had missing matching criteria). That is, the default for two individuals is that they are pairable – only non-missing, non-matching external criteria (as well as the p-value test based on genetic data, described above) will make a pair unpairable.

## 6.4 IBS similarity matrix

For the N individuals in a sample, to create a  $N \times N$  matrix of genome-wide average IBS pairwise identities:

```
plink --file mydata --cluster --matrix
creates the file
   plink.mibs
```

which contains a square, symmetric matrix of the IBS distances for all pairs of individuals. These values range, in theory, from 0 to 1. In practice, one would never expect to observe values near 0 – even completely unrelated individuals would be expected to share a very large proportion of the genome identical by state by chance alone (i.e. as opposed to identity by descent). A value of 1 would indicate a MZ twin pair, or a sample duplication. More details on pairwise relatedness can be obtained by using the --genome command.

The default behavior of --matrix to to output similarities (proportions of alleles IBS). To generate a distance matrix (1-IBS) then use the command

```
plink --file mydata --cluster --distance-matrix
instead. This will generate a file
    plink.mdist
```

**HINT** See the FAQ page for instructions on using using R to visualise these results; alternatively, use the --mds-plot option described below.

**NOTE** In versions prior to v1.00, there is no --distance-matrix command and --matrix outputs a file called plink.mdist rather than plink.mibs - these are still similarities, not distances.

## 6.5 Multidimensional scaling plots

To perform multidimensional scaling analysis on the  $N \times N$  matrix of genome-wide IBS pairwise distances, use the --mds-plot option in conjunction with --cluster. This command takes a single parameter, the number of dimensions to be extracted. For example, assuming we have already calculated the plink.genome file,

```
plink --file mydata --read-genome plink.genome --cluster --mds-plot 4
creates the file
     plink.mds
which contains one row per individual, with the fields
            Family ID
     IID
            Individual ID
     SOL
            Assigned solution code (from --cluster)
     C1
            Position on first dimension
     C2
            Position on second dimension
     C3
            Position on third dimension
     C4
            Position on fourth dimension
```

Plotting the C1 values against C2, for example, will give a scatter plot in which each point is an individual; the two axes correspond to a reduced representation of the data in two dimensions, which can be useful for identifying any clustering. Standard classical (metric) multidimensional scaling is used.

Instead of using each individual as the unit of analysis, you can make each point a cluster from the final solution (as determined by --cluster along with whatever constraints were imposed) and the distances between clusters are the average distances of all individuals in those clusters. Use the --mds-cluster flag (as well as --cluster --mds-plot K) for this.

## 6.6 Outlier detection diagnostics

Sometimes it can be useful to detect a handful of individuals who do not cluster with an otherwise fairly homogeneous sample. It is possible to generate some metrics describing much of an 'outlier' an individual is with respect to the other individuals in that sample, based on the genome-wide IBS information, as decribed above.

For any one individual, we can rank order all other individuals on the basis of how similar (in IBS terms) they are to this particular proband individual. We can then ask, is the proband's closest neighbour significantly more distant to the proband than all other individuals' nearest neighbour is to them. In otherwords, from the distribution of 'nearest neighbour' scores, one for each individual, we can calculate a sample mean and variance and transform this measure into a Z score. If an individual has an extreme low Z score, say less than 4 standard deviation units, this would indicate that this individual is an outlier with respect to the rest of the sample (i.e. this individual's nearest neighbour is a lot less near than the average nearest neighbour). As well as performing this test with the nearest neighbour, we can also perform it with the distribution of second-closest neighbours for each individual; then third closest neighbours, etc. It might sometimes be more informative to look at these 'second-closest' and 'third-closest' measures, to detect, for instance, a pair of individuals who are very similar to each other, but very distant from the rest of the sample – they would score normally on the 'first-closest' neighbour test, but not on the 'second-closest', 'third-closest' tests, etc. It might sometimes be informative to look at the whole distribution of these 'neighbour' metrics, going to 1st nearest to the Nth nearest.

Another metric which can be used to identify potential outliers is, for each individual, to calculate the proportion of binomial IBS tests (described in the constaints section above), for each individual, that showed a significant difference at the --ppc threshold.

The basic option is, for example:

```
plink --file data --cluster --neighbour 1 5
```

This command always takes two arguments, specifying, in this case, to consider from the 1st nearest neighbour to the 5th nearest neighbour; this option generates the output file:

#### plink.nearest

which contains the fields:

FID	Family ID
IID	Individual ID
NN	Nearest neighbour level (see below)
MIN_DST	IBS distance of nth nearest neighbour (see below)
Z	MIN_DST converted to a Z score (see below)
FID2	Family ID of the nth nearest neighbour
IID2	Individual ID of the nth nearest neighbour
PROP_DIFF	Proportion of significantly different others (see below)

Looking at some example output, in this case for two individuals from the Asian HapMap samples, measured on around 50K random SNPs, for nearest neighbours 1 to 5, we see:

FID	IID	NN	$MIN\_DST$	$oldsymbol{Z}$	FID2	IID2	$PROP\_DIFF$
JPT256	1	1	0.7422	0.8897	JPT265	1	0.01136
JPT256	1	2	0.742	1.223	JPT236	1	0.01136
JPT256	1	3	0.7408	0.6503	JPT261	1	0.01136
JPT256	1	4	0.7405	0.7285	JPT250	1	0.01136
JPT256	1	5	0.7402	0.6204	JPT269	1	0.01136
JPT257	1	1	0.7368	-3.701	JPT242	1	0.9318
JPT257	1	2	0.7364	-3.463	JPT238	1	0.9318
JPT257	1	3	0.7359	-3.832	JPT244	1	0.9318
JPT257	1	4	0.7356	-3.974	JPT245	1	0.9318
JPT257	1	5	0.7353	-4.046	JPT228	1	0.9318

Here we clearly see that the individual coded as JPT257 seems to be an outlier, with these first five measures being around 4 standard deviations below the group mean. In contrast, individual JPT256 does not appear to be an outlier, as the Z scores are above the mean (greater than 0). Plotting the Z scores for the entire sample makes it clear that JPT257 is indeed an outlier, as does the result for the IBS test – JPT257 is significant different from 93% of the rest of the sample (the threshold for the IBS test is set to be quite stringent here, 0.0005 – this is changed with the –-ppc option as described above). At this fairly strict level, the subtle differences between Japanese and Han Chinese individuals are not detected – using a threshold at 0.05, for example, one would see that many individuals show greater than the expected 0.05 in the PROP\_DIFF field, as it is now picking up this group difference.

## Chapter 7

# IBS/IBD estimation

As well as the standard summary statistics described above, PLINK offers some alternative measures such as estimated inbreeding coefficients for each individual and genome-wide identity-by-state and identity-by-descent estimates for all pairs of individuals. The latter can be used to detect sample contaminations, swaps and duplications as well as pedigree errors and unknown familial relationships (e.g. sibling pairs in a case/control population-based sample). PLINK also has functions to detect specific segments shared between distantly-related individuals.

All these analyses require a large number of SNPs!

#### 7.1 Pairwise IBD estimation

plink --file mydata --genome

The pairwise clustering based on IBS, as outlined in the previous section is useful for detecting pairs of individuals who look more different from each other than you'd expect in a random, homogeneous sample. In this section, we consider using the same genotype data to provide a complementary analysis: using estimates of pairwise IBD to find pairs of individuals who look too similar to each other, i.e. more than we would expect by chance in a random sample.

In a homogeneous sample, it is possible to calculate genome-wide IBD given IBS information, as long as a large number of SNPs are available (probably 1000 independent SNPs at a bare minimum; ideally 100K or more).

```
which creates the file
     plink.genome
which has the following fields:
     FID1
               Family ID for first individual
     IID1
               Individual ID for first individual
     FID2
               Family ID for second individual
     IID2
               Individual ID for second individual
     Z0
               P(IBD=0)
     Z1
               P(IBD=1)
     Z2
               P(IBD=2)
     PI_HAT
               P(IBD=2)+0.5*P(IBD=1) (proportion IBD)
     IBS0
               Number of IBS 0 nonmissing loci
               Number of IBS 1 nonmissing loci
     IBS1
               Number of IBS 2 nonmissing loci
     IBS2
     DST
               IBS distance (IBS2 + 0.5*IBS1) / ( N SNP pairs )
```

```
P IBS binomial test

HOMHOM Number of IBS 0 SNP pairs used in test

HETHET Number of IBS 2 het/het SNP pairs in test

RATIO R in ratio of 1:2 for IBS 0 : HETHET
```

This file will have as many rows as there are unique pairs of individuals in the sample – for large samples with thousands of individuals, this file can be very large (and take considerable time to generate).

HINT To produce a smaller version of this file use the command --genome-minimal instead; however, this is only useful if the purpose is to subsequently merge the data using --read-genome-minimal (i.e. when running --cluster or --segment. A disadvantage is that multiple plink.genome.min files cannot be concatenated in the same manner for normal plink.genome files; this will be remedied in future releases of PLINK (i.e. to allow parallel computation of the genome file.

As mentioned above, the IBD estimation part of this analysis relies on the sample being reasonably homogeneous – otherwise, the estimates will be biased (i.e. individuals within the same strata will show too much apparent IBD). It is therefore important to run the other population stratification measures provided by plink and other packages before estimating pairwise IBD. In addition, see the notes on the IBS test in the previous section where it is introduced as a constrain on clustering.

**HINT** To reduce the file size, use the  $--\min X$  option to only output to plink.genome pairs where PI\_HAT is greater than X. That is,

```
plink --file mydata --genome --min 0.05
```

will only display the pairs of individuals showing reasonably high levels of IBD sharing (i.e. typically it will be these pairs that are of interest, rather than the vast majority of pairs that show no excess sharing).

Hint Calculating the average pi-hat for each individual and looking for outliers is also useful (in particular, sample contamination will lead to too many heterozygote calls, which leads to fewer IBS 0 calls, which leads to over-estimated IBD with all other people in the sample). Be sure to set --min 0 and --max 1 in this case to obtain pairs for all individuals.

Advanced hint If you have access to a cluster, use the --genome-lists option to facilitate parallelization, as described in the IBS clustering section.

## 7.2 Inbreeding coefficients

Given a large number of SNPs, in a homogeneous sample, it is possible to calculate inbreeding coefficients (i.e. based on the observed versus expected number of homozygous genotypes).

```
plink --file mydata --het
which will create the output file:
    plink.het
```

which contains the fields, one row per person in the file:

```
FID Family ID

IID Individual ID

O(HOM) Observed number of homozygotes

E(HOM) Expected number of homozygotes

N(NM) Number of non-missing genotypes

F inbreeding coefficient estimate
```

This analysis will automatically skip haploid markers (male X and Y chromosome markers).

**Note** With whole genome data, it is probably best to apply this analysis to a subset that are pruned to be in approximate linkage equilibrium, say on the order of 50,000 autosomal SNPs. Use the **--indep-pairwise** and **--indep** commands to achieve this, described here.

**Note** The estimate of F can sometimes be negative. Often this will just reflect random sampling error, but a result that is strongly negative (i.e. an individual has *fewer* homozygotes than one would expect by chance at the genome-wide level) can reflect other factors, e.g. sample contamination events perhaps.

## 7.3 Runs of homozygosity

A simple screen for runs of homozygous genotypes within any one individual is provided by the commands --homozyg-snp and --homozyg-kb which define the run in terms of the required number of homozygous SNPs spanning a certain kb distance, e.g.

The algorithm is as follows: Take a window of X SNPs and slide this across the genome. At each window position determine whether this window looks 'homozygous' enough (yes/no) (i.e. allowing for some number of hets or missing calls). Then, for each SNP, calculate the proportion of 'homozygous' windows that overlap that position. Call segments based on this metric, e.g. based on a threshold for the average.

The exact window size and thresholds, relative to the SNP density and expected size of homozygous segments, etc, is obviously important: sensible default values are supplied for the context of dense SNP maps, scanning for large segments. In general, this approach will ensure that otherwise long runs of homozygosity are not broken by the occassional heterozygote. (For more accurate detection of smaller segments, one might consider approaches that also take population parameters such as allele frequency and recombination rate into account, in a HMM approach for example: but for now, PLINK only supports this basic detection of long, homozygous segments).

To run this option with default values, use the command

```
plink --bfile mydata --homozyg
which generates a file
   plink.hom
```

The plink.hom file has the following format, one row per identified homozygous region:

```
FID
         Family ID
IID
         Individual ID
CHR
         Chromosome
SNP1
         SNP at start of region
SNP2
         SNP at end of region
POS1
         Physical position (bp) of SNP1
POS2
         Physical position (bp) of SNP2
KΒ
         Length of region (kb)
NSNP
         Number of SNPs in run
DENSITY
         Average SNP density (1 SNP per kb)
PHOM
         Proportion of sites homozygous
PHET
         Proportion of sites heterozygous
```

The options to change the behavior of this function (along with the default values as parameters) are given below.

To change the definition of the sliding 'window', use the options

```
--homozyg-window-kb 5000
--homozyg-window-snp 50
```

To change the number of heterozygotes allowed in a window

```
--homozyg-window-het 1
```

To change the number of missing calls allowed in window, e.g.

```
--homozyg-window-missing 5
```

To change the proportion of overlapping windows that must be called homozygous to define any given SNP as 'in a homozygous segment', use

#### --homozyg-window-threshold 0.05

(i.e. this number is relatively low, so that SNPs at the edge of a true segment will be called, as long as the windows are sufficiently large, such that the probability of a window being homozygous by chance is sufficiently small).

The above options define the 'window' that slides across the genome; the options below relate to the final segments that are called as homozygous or not:

```
--homozyg-snp 100
```

--homozyg-kb 1000

You can also specify the required minimum density (in kb, i.e. 1 SNP per 50 kb)

--homozyg-density 50

Finally, if two SNPs within a segments are too far apart (measured in kb), that segment can be split in two:

```
--homozyg-gap 1000
```

**HINT** As is, this analysis should be performed on sets of SNPs that have been pruned for strong local LD (if the goal is to find long segments that are more likely to represent homozygosity by descent (i.e. autozygosity) rather than simply by chance).

To obtain pools of overlapping and potentially matching segments, we can use --homozyg-group in addition to the above, which generates the file

#### plink.hom.overlap

which contains the fields

```
FTD
        Family ID
IID
        Individual ID
PHE
        Phenotype of individual
CHR
        Chromosome
SNP1
        SNP at start of segment
        SNP at end of segment
SNP2
        Physical position of start of segment
BP1
BP2
        Physical position of end of segment
        Physical size of segment
KB
NS
        Number of segments in the pool that match this one
GRP
        Allelic-match grouping of each segment
```

For example, the command

```
plink --file test --homozyg --homozyg-group
```

might result in the file plink.hom.overlap containing:

FID	IID	PHE	CHR	SNP1	SNP2	BP1	BP2	KB	NS	GRP
1	1	2	1	snp1	snp7	1000000	7000000	6000	1	1
6	1	1	1	snp1	snp5	1000000	5000000	4000	1	1*
2	1	1	1	snp2	snp7	2000000	7000000	5000	0	2*
CON	3	1:2	1	snp2	snp5	2000000	5000000	3000		

This implies one pool (i.e. each pool is separated by a CON (consensus row) and a space. CON is the consensus region; the ratio is the case:control segment ratio; under IID we have the number of individuals.

When there is more than one pool, they are ordered by the number of segments in the pool, then physical position. To output only pools of a particular size, use the --pool-size N option (e.g. --pool-size 10 to only display pools with at least 10 segments).

A pool contains overlapping segments, which may or may not also allelically match. For allelic matching, segments are compared pairwise, and a match is declared if at least 0.95 of jointly non-missing, jointly homozygous sites are identical. This threshold can be changed with the option

```
--homozyg-match 0.99
```

The number of other segments in the pool that allelically match each segment is shown in the NS field. The GRP field shows how PLINK attempts to group allelically-matching segments within the pool of overlapping segments. It works as follows:

- For each segment, find the number of other segments that match (NS).
- Find segment with largest NS, denote as group 1, and put a \* to indicate this is the index for this group.
- Denote all other segments that match this index as being in GRP 1 (i.e. but without the \*)
- Continue to next ungrouped segment (2\*, etc)

By default, we compare all segments pairwise when asking if they match; if the --consensus-match flag is given, then for a pool of overlapping segments, matches are defined only on the basis of the consensus region (i.e. the overlapping region shared by all segments). This is probably not very sensible in many cases, as the consensus region can often be small (i.e. a single SNP).

The NS field can suggest any intransitivity in matching: e.g. if B matches A and C but A does not match C, then if B has already been grouped with A, C would not be added to this group as being an allelic match. In this case C would have NS ¿ 0 but belong to a GRP of its own.

Internally, all pools are formed but then pruned if, for instance, a smaller pool is included in a larger pool completely. That means that in certain circumstances you will see a segment in more than one pool. For example, imagine a grid with three people A, B and C along the columns, each row representing physical position, and the presence of a letter representing a homozygous run:

A . . A B .

A B C A B C

A . C

A . C

In this case, A,B and A,C and B,C pools will not be displayed, as they appear in the super-pool A,B,C. However, if we instead had:

. . .

Α..

АВ.

АВ.

A . . A . C

A . C

A . .

. . .

Then you will see A,B and A,C (i.e. with A shown twice) as we have two distinct consensus regions here. Finally, if the --homozyg-verbose option is added, the plink.hom.overlap file will then display the actual segments underneath each pool. Here each individual is listed across the page, so the 3 columns refers to the 3 segments in the pool. For example:

```
plink --file test --homozyg-snp 2 --homozyg-group --homozyg-verbose
```

now generates plink.hom.overlap as follows (with annotation added in *italics*):

FID	IID	Pl	HE CHR	SNP1	SNP2	BP1	BP2	KB	NS	GRP
1	1		2 1	snp1	snp7	1	7	0.006	1	1
6	1		1 1	snp1	snp5	1	5	0.004	1	1*
2	1		1 1	snp2	snp7	2	7	0.005	0	2*
CON	3	1	:2 1	snp2	snp5	2	5	0.003		
SNP	1	6	2		<	Family II	)			
	1	1	1		<	Individuo	al ID			
	1	1	2		<	GRP code				
snp1	[A/A]	[A/A]	C/A		<	now SNPs	are lis	ted down	the	page
snp2	[A/A]	[A/A]	[C/C]		<	start of	consens	us region	n	
snp3	[A/A]	[A/A]	[C/C]							
snp4	[A/A]	[A/A]	[C/C]							
snp5	[A/A]	[A/A]	[C/C]		<	end of co	onsensus	region		
snp6	[A/A]	A/C	[C/C]							
snp7	[A/A]	A/C	[C/C]							

A bracket indicates that that genotype is part of the homozygous segment: the consensus region is the intersection. The entire union of SNPs is displayed and the consensus region is indicated by spaces before and after. i.e. the consensus region is that where all genotypes are in [brackets].

Obviously, this file can get quite large (+wide) with real data and it is not very machine-readable.

# 7.4 Segmental sharing: detection of extended haplotypes shared IBD

**WARNING** This analysis is still in the *beta* development stage and is considerably more involved than many others provided by this package: currently, you should only perform these analyses if you consider yourself something of *an analytic expert* and are confident you will be able to interpret the output! Over time, we expect that the documentation and features supporting this analysis will improve.

There are five important steps to this analysis:

- Obtain a homogeneous sample
- Remove very closely related individuals
- Prune SNP set
- Detect segments
- Associate with disease

#### 7.4.1 Check for a homogenous sample

This analysis requires that all individuals belong to a single, homogeneous population. To ensure this assumption is reasonable: as described here, first run

```
plink --bfile mydata1 --genome
```

to generate a plink.genome file. This will be used subsequently in a number of steps.

Then, using the available tools, such as listed here and described more fully in the section on stratification, obtain a relatively homogeneous dataset. Some relevant options are listed here:

```
--cluster (cluster individuals)
--matrix (generate .mdist file, used to generate MDS plots)
```

```
--ppc (threshold for PPC test, not to cluster individuals)
--mds-plot (generate a multidimensional scaling plot)
--ibs-test (as case/control less similar on average?)
--neighbour (option to find individual outliers)
```

Also, remove individuals who appear to have higher levels of inbreeding than expected (see above). If you have a set of individuals you want to exclude from analysis based on these steps, for example, listed in the file outliers.txt (FID, IID) then use:

```
./plink --bfile mydata1 --remove outliers.txt --make-bed --out mydata2
```

#### 7.4.2 Remove very closely related individuals

The focus of this analysis is to look for extended haplotypes shared between distantly related individuals: having very closely related individuals (siblings, first cousins, etc) will likely swamp the results of the analysis. Scan the plink.genome file for any individuals with high PIHAT values (e.g. greater than 0.05). Optionally, remove one member of the pair if you find close relatives. (Alternatively, to keep them in but just exclude this pair from the segmental analysis, see below).

#### 7.4.3 Prune the set of SNPs

The segmental sharing analysis requires approximately independent SNPs (i.e. linkage equilibrium). Two options to prune are documented here.

A reasonable strategy might be as follows:

```
plink --bfile mydata2 --mind 1 --geno 0.01 --maf 0.05 --make-bed --out mydata3 followed by

plink --bfile mydata3 --indep-pairwise 100 25 0.2 followed by

plink --bfile mydata3 --extract plink.prune.in --make-bed --out mydata4
```

#### 7.4.4 Detecting shared segments (extended, shared haplotypes)

With a newly pruned fileset, ideally containing only independent, high quality SNPs in individuals who are not very closely related but are from the same population, run the command

```
plink --bfile mydata4 --read-genome plink.genome --segment
```

PLINK expects the 3rd column the MAP/BIM file to contain genetic distances in Morgan units. A reasonable approximation is to scale from physical position (i.e. column 4) at 1cM=1Mb. If the genetic distances are in cM instead of Morgans, add the --cm flag.

To set threshold on who to include/exclude based on genome wide IBD use

```
--min 0.01
```

For example, this would exclude pairs who share ¿10% of their genomes. Alternatively, to include all pairs, irrespective of whether we estimate any genome-wide sharing or not, add the option

```
--all-pairs
```

instead. This will use all pairs, allowing for a small prior probability of sharing for pairs that otherwise are at the boundary of IBD sharing (i.e. sharing 0% IBD). Naturally, for a large sample, it may become prohibitive to consider all possible pairs.

```
The --segment option generates a file
```

#### plink.segment

which has the fields:

FID1	Family ID of first individual
IID1	Individual ID of first individual
FID2	Family ID of second individual
IID2	Individual ID of second individual
PHE	Phenotype concordance: -1,0,1
CHR	Chromosome code
BP1	Start physical position of segment (bp)
BP2	End physical position of segment (bp)
SNP1	Start SNP of segment
SNP2	End SNP of segment
NSNP	Number of SNPs in this segment
KB	Physical length of segment (kb)

Here one row represents one segment. The PHE field is coded -1,0,1 for control/control, case/control, or case/case pairs respectively.

The option

```
--segment-length 2000
```

means to only select segments that are at least 2000 kb in length, for example. The option

```
--segment-snp 100
```

means only to select segments that contain at least 100 SNPs, for example.

For ease of interpretation, and to increase the probably that the segments are truly shared IBD and thus tags shared rare variation between two individuals, it makes sense to restrict ones focus to very extended segments (e.g. over 1Mb in size, for example).

Another option is the --segment-group option, which generates output similar to --homozyg-group, described above; similarly, --segment-verbose prints out the actual genotypes for the individuals that overlap. However, these can be large files that are not necessarily easy to interpret.

#### 7.4.5 Association with disease

Along with the --segment option, as above, if you also add:

```
--mperm N
```

then, for case/control data, this performs a test of whether segments stack up more in case/case pairs versus non-case/case pairs at any position, performing N permutations. Equivalently, you can use an already-created segment file:

```
./plink --bfile mydata4 --read-segment plink.segment --mperm 10000
```

This will generate two files:

```
plink.segment.summary
```

which contains one row corresponding to one SNP; there are five fields:

```
CHR Chromosome code

SNP SNP identifier

CONU Number of control/control segments over this SNP

DISC Case/control segments spanning this position

CONA Case/case segment count
```

The file

plink.segment.summary.mperm

contains empirical significance values for each position, asking whether there is a higher rate of case/case sharing than expected. It is important to note that the test statistic is still under developement: in this current release, it should merely be interpreted as a rough guide to the data. Naturally, the thresholds for declaring significance will be much lower than for genome-wide association analysis; precise guidelines will be put in place presently.

## Chapter 8

## Association analysis

The basic association test is for a disease trait and is based on comparing allele frequencies between cases and controls (asymptotic and empirical p-values are available). Also implemented are the Cochran-Armitage trend test, Fisher's exact test, different genetic models (dominant, recessive and general), tests for stratified samples (e.g. Cochran-Mantel-Haenszel, Breslow-Day tests), a test for a quantitative trait; a test for differences in missing genotype rate between cases and controls; multilocus tests, using either Hotelling's T(2) statistic or a sum-statistic approach (evaluated by permutation) as well as haplotype tests. The basic tests can be performed with permutation, described in the following section to provide empirical p-values, and allow for different designs (e.g. by use of structured, within-cluster permutation). Family-based tests are described in the next section

HINT The basic association commands (--assoc, --model, --fisher, --linear and --logistic) will test only a single phenotype. If your alternate phenotype file contains more than one phenotype, then adding the --all-pheno flag will make PLINK cycle over each phenotype, e.g. instead of a single plink.assoc output file, if there are 100 phenotypes, PLINK will now show

```
plink.P1.assoc
plink.P2.assoc
...
plink.P100.assoc
```

Naturally, it will take 100 times longer... If you are testing a very large number of phenotypes, it might be worth specifying --pfilter also, to reduce the amount of amount (e.g. only outputing tests significant at p=1e-4 if --pfilter 1e-4 is specified).

## 8.1 Basic case/control association test

To perform a standard case/control association analysis, use the option:

```
plink --file mydata --assoc
which generates a file
     plink.assoc
which contains the fields:
     CHR.
              Chromosome
     SNP
              SNP ID
     BP
              Physical position (base-pair)
              Minor allele name (based on whole sample)
     Α1
     F_A
             Frequency of this allele in cases
     F_{-}U
             Frequency of this allele in controls
```

```
A2 Major allele name
CHISQ Basic allelic test chi-square (1df)
P Asymptotic p-value for this test
OR Estimated odds ratio (for A1)
```

**Hint** In addition, if the optional command --ci X (where X is the desired coverage for a confidence interval, e.g. 0.95 or 0.99) is included, then two extra fields are appended to this output:

```
L95 Lower bound of 95% confidence interval for odds ratio
U95 Upper bound of 95% confidence interval for odds ratio
```

(where 95 would change if a different value was used with the --ci option, naturally).

See the next section on permutation to learn how to generate empirical p-values and use other aspects of permutation-based testing.

See the section on multimarker tests to learn how to perform haplotype-based tests of association.

This analysis should appropriately handle X/Y chromosome SNPs automatically.

### 8.2 Fisher's Exact test (allelic association)

To perform a standard case/control association analysis using Fisher's exact test to generate significance, use the option:

```
plink --file mydata --fisher
which generates a file
     plink.fisher
which contains the fields:
     CHR
             Chromosome
     SNP
             SNP ID
     BP
             Physical position (base-pair)
             Minor allele name (based on whole sample)
     Α1
     F_A
             Frequency of this allele in cases
     F_U
             Frequency of this allele in controls
     A2
             Major allele name
     Ρ
             Exact p-value for this test
     OR.
             Estimated odds ratio (for A1)
```

As described below, if --fisher is specified with --model as well, PLINK will perform genotypic tests using Fisher's exact test.

**Note** You can also use permutation to generate exact, empirical significance values that would also be valid in small samples, etc.

## 8.3 Alternate / full model association tests

It is possible to perform tests of association between a disease and a variant other than the basic allelic test (which compares frequencies of alleles in cases versus controls), by using the --model option. The tests offered here are (in addition to the basic allelic test):

- Cochran-Armitage trend test
- Genotypic (2 df) test
- Dominant gene action (1df) test

#### • Recessive gene action (1df) test

One advantage of the Cochran-Armitage test is that it does not assume Hardy-Weinberg equilibrium, as the individual, not the allele, is the unit of analysis (although the permutation-based empirical p-values from the basic allelic test also have this property). It is important to remember that SNPs showing severe deviations from Hardy-Weinberg are often likely to be bad SNPs, or reflect stratification in the sample, however, and so are probably best excluded in many cases.

The genotypic test provides a general test of association in the 2-by-3 table of disease-by-genotype. The dominant and recessive models are tests for the minor allele (which is the minor allele can be found in the output of either the --assoc or the --freq commands. That is, if D is the minor allele (and d is the major allele):

```
Allelic:
                             versus
Dominant:
                (DD, Dd)
                                           dd
                             versus
Recessive:
                   DD
                                       (Dd, dd)
                             versus
                   DD
Genotypic:
                                          Dd
                                                                  dd
                             versus
                                                       versus
```

As mentioned above, these tests are generated with option:

```
plink --file mydata --model
which generates a file
    plink.model
which contains the following fields:
```

CHR Chromosome number SNP SNP identifier

TEST Type of test

AFF Genotypes/alleles in cases
UNAFF Genotypes/alleles in controls

CHISQ Chi-squated statistic

DF Degrees of freedom for test

P Asymptotic p-value

Each SNP will feature on five rows of the output, corresponding to the five tests applied. The column TEST refers to either ALLELIC, TREND, GENO, DOM or REC, referring to the different types of test mentioned above. The genotypic or allelic counts are given for cases and controls separately. For recessive and dominant tests, the counts represent the genotypes, with two of the classes pooled.

The genotypic and dominant/recessive tests will only be conducted if there is a minimum number of observations per cell in the 2-by-3 table: by default, if at least one of the cells has a frequency less than 5, then we skip the alternate tests (NA is written in the results file). The Cochran-Armitage and allelic tests are performed in all cases. This threshold can be altered with the --cell option:

```
plink --file mydata --model --cell 20
```

If permutation (with the --mperm or --perm options) is specified, the -model option will by default perform a permutation test based on the most significant result: that is, for each SNP, the best original result will be compared against the best of the four tests for that SNP for every replicate (note that STAT in the permutation output files below is in this case 1 minus the best p-value); in max(T) permutation mode, this will also be compared against the best result from all SNPs for the EMP2 field. This procedure controls for the fact that we have selected the best out of four tests for each SNP. The output will be generated in the file

```
plink.model.best.perm
or
    plink.model.best.mperm
depending on whether adaptive or max(T) permutation was used.
```

The behavior of the --model command can be changed by adding the --model-gen, --model-trend, --model-dom or --model-rec flags to make the permutation use the genotypic, the Cochram-Armitage trend test, the dominant test or the recessive test as the basis for permutation instead. In this case, one of the the following files will be generated:

```
plink.model.gen.permplink.model.gen.mpermplink.model.trend.permplink.model.trend.mpermplink.model.dom.permplink.model.dom.mpermplink.model.rec.permplink.model.rec.mperm
```

It is also possible to add the --fisher flag to obtain exact p-values:

```
./plink --bfile mydata --model --fisher
```

in which case the CHISQ field does not appear. Note that the genotypic, allelic, dominant and recessive models use the Fisher's exact; the trend-test does not and will give the same p-value as without the --fisher flag. Also, by default, when --fisher is added, the --cell field is set to 0, i.e. to include all SNPs.

### 8.4 Stratified analyses

When a cluster variable has been specified, by pointing to a file that contains this information, with the --within command, it is possible to perform a number of tests of case/control association that take this clustering into account, or explicitly test for homogeneity of effect between clusters.

**Note** In many cases, permutation procedures can also be used to account for clusters in the data. See the next section for more details. The tests presented below are only applicable for case/control data, so permutation might be useful for quantitative trait outcomes, etc.

There are two basic classes of test:

- Testing for overall disease/gene association, controlling for clusters
- Testing for heterogeneity of the disease/gene assocation between different clusters

The type of cluster structure will vary in terms of how many clusters there are in the sample, and how many people belong to each cluster. At one extreme, we might have two only 2 clusters in the sample, each with a large number of cases and controls. At the other extreme, we might have a very large number of clusters, such that each cluster only has 2 individuals. These factors will influence the choice of stratified analysis.

The tests offered are:

- Cochran-Mantel-Haenszel test for 2x2xK stratified tables
- Cochran-Mantel-Haenszel test for IxJxK stratified tables
- Breslow-Day test of homogeneity of odds ratio
- Partitioning the total association chi-square to perform between and within cluster association, and a test of homogeneity of effect

The Cochran-Mantel-Haenszel (CMH) tests are valid with both a large number of small clusters and a small number of large clusters. These tests provide a test based on an "average" odds ratio that controls for the potential confounding due to the cluster variable.

The Breslow-Day test asks whether different clusters have different disease/gene odds ratios: this test assumes a moderate sample size within each cluster. The partitioning total association test, which is conceptually similar to the Breslow-Day test, also makes the same assumption.

As mentioned above, the CMH test comes in two flavours: 2x2xK and IxJxK. Currently, the 2x2xK test represents a disease x SNP | cluster test. The generalized form, the IxJxK, represents a test of cluster x SNP | disease, i.e. does the SNP vary between clusters, controlling for any possible true

SNP/disease association. This latter test might be useful in interpreting significant associations in stratified samples. Typically, the first form of the test will be of more interest, however. These two tests are run by using the options:

```
plink --file mydata --mh --within mycluster.dat
for the basic CMH test, or
  plink --file mydata --mh2 --within mycluster.dat
for the IxJxK test.
The --mh option generates the file
     plink.cmh
which contains the fields
     CHR.
                Chromosome number
     SNP
                SNP identifier
     Α1
                Minor allele code
     A2
                Major allele code
     BP
                Physical position (base-pair)
                Cochran-Mantel-Haenszel statistic (1df)
     CHISQ
     Ρ
                Asymptotic p-value for CMH test
     OR
                CMH odds ratio
     L95
                Lower bound on confidence interval for CMH odds ratio
     U95
                Upper bound on confidence interval for CMH odds ratio
The range of the confidence interval with the --mh option can be changed with the --ci option:
  plink --file mydata --mh --within mycluster.dat --ci 0.99
The --mh2 option generates the file
     plink.cmh2
which contains the fields:
     CHR
                  Chromosome
     SNP
                  SNP identifier
     CHISQ_CMH2
                  Cochran-Mantel-Haenszel test for IxJxK tables
```

Asymptotic p-value for this test It is not possible to obtain confidence intervals or odds ratios for --mh2 tests.

Hint A trick to analyse phenotypes with more two categories (but only with nominal, not ordinal outcomes) is to use the --mh2 option with the phenotype in the cluster file and the phenotype in the PED file set all to a single value.

#### 8.5 Testing for heterogeneous association

As mentioned in the previous section, two methods are provided to test for between-cluster differences in association when using a case/control design. The Breslow-Day test is specified with the option:

```
plink --file mydata --bd --within myclst.txt
```

which runs and generates the same files as the --mh option, described above, but with two extra fields appended:

```
Breslow-Day test
CHISQ_BD
P_BD
           Asymptotic p-value
```

P\_CMH2

where a significant value indicates between-cluster heterogeneous in the odds ratios for the disease/SNP association.

A similar test of the homogeneity of odds ratio tests based on partitioning the chi-square statistic is given by:

```
plink --file mydata --homog --within myclst.txt
which generates the file
     plink.homog
which contains the fields
     CHR
              Chromosome number
     SNP
              SNP identifier
     A1
              Minor allele code
     A2
              Major allele code
     F_A
              Case allele frequency
     F_U
              Control allele frequency
     N_A
              Case allele count
     N_U
              Control allele count
     TEST
              Type of test
     CHISQ
              Chi-squared association statistic
     DF
              Degrees of freedom
     Ρ
              Asymptotic p-value
     OR
              Odds ratio
The TEST type is either
     TOTAL
              Total SNP & strata association
     ASSOC
              SNP association controlling for strata
     HOMOG
              Between-strata heterogeneity test
     X_1
              Association in first stratum
     X_2
              Association in second stratum
```

## 8.6 Hotelling's T(2) multilocus association test

#### **IMPORTANT** This command has been temporarily disabled

For disease-traits, PLINK provides support for a multilocus, genotype-based test using Hotelling's T2 (T-squared) statistic. The --set option should be used to specify which SNPs are to be grouped, as follows:

```
plink --file data --set mydata.set --T2
```

where mydata.set defines which SNPs are in which set (see this section for more information on defining sets).

This command will generate a file

```
plink.T2
```

which contains the fields

SET	Set name
SIZE	Number of SNPs in this set
F	F-statistic from Hotelling's test
DF1	Degrees of freedom 1
DF2	Degrees of freedom 2
P_HOTEL	Asymptotic p-value

HINT Use the --genedrop permutation to perform a family-based application of the Hotelling's T2 test. This command can be used with all permutation methods (label-swapping or gene-dropping, adaptive or max(T)). In fact, the permutation test is based on 1-p in order to make the between set comparisons for the max(T) statistic more meaningful (as different sized sets would have F-statistics with different degrees of freedom otherwise). Using permutation will generate one of the following files:

```
plink.T2.perm
which contain the fields
     SET
               Set name
     SIZE
               Number of SNPs in this set
     EMP1
               Empirical p-value
     NR.
               Number of permutation replicates
or, if --mperm was used,
     plink.T2.mperm
which contain the fields
     SET
               Set name
     SIZE
               Number of SNPs in this set
     EMP1
               Empirical p-value
     EMP2
               max(T) empirical p-value
```

Note that this test uses a simple approach to missing data: rather than case-wise deletion (removing an individual if they have at least one missing observation) we impute the mean allelic value. Although this retains power under most scenarios, it can also cause some bias when there are lots of missing data points. Using permutation is a good way around this issue.

### 8.7 Quantitative trait association

Quantitative traits can be tested for association also, using either asymptotic (likelihood ratio test and Wald test) or empirical significance values. If the phenotype (column 6 of the PED file or the phenotype as specified with the --pheno option) is quantitative (i.e. contains values other than 1, 2, 0 or missing) then PLINK will automatically treat the analysis as a quantitative trait analysis. That is, the same command as for disease-trait association:

```
plink --file mydata --assoc
will generate the file
 plink.qassoc
with fields as follows:
     CHR
               Chromosome number
     SNP
               SNP identifier
     BP
               Physical position (base-pair)
     NMISS
               Number of non-missing genotypes
     BETA
               Regression coefficient
     SE
               Standard error
     R2
               Regression r-squared
     Т
               Wald test (based on t-distribtion)
               Wald test asymptotic p-value
If permutations were also requested, then an extra file, either
     plink.assoc.perm
or
     plink.assoc.mperm
```

will be generated, depending on whether adaptive or max(T) permutation was used (see the next section for more details). The empirical p-values are based on the Wald statistic.

### 8.8 Genotype means for quantitative traits

Adding the flag --qt-means along with the --assoc command, when run with a quantitative trait, will produce an additional file with a list of means and standard deviations stratified by genotype, called

```
plink.qassoc.means
and format
     CHR
              Chromosome code
     SNP
              SNP identifier
              Description of next three fields
     VALUE
     G11
              Value for first genotype
     G12
              Value for second genotype
     G22
              Value for third genotype
where VALUE is one of GENO, COUNTS, FREQ, MEAN or SD (standard deviation). For example:
      CHR
                     SNP
                           VALUE
                                       G11
                                                 G12
                                                           G22
                            GENO
                                                           1/1
        5
             hCV26311749
                                       2/2
                                                 2/1
        5
             hCV26311749 COUNTS
                                                           597
                                         1
                                                  60
        5
             hCV26311749
                            FREQ
                                  0.00152
                                            0.09119
                                                       0.9073
        5
             hCV26311749
                            MEAN
                                    0.9367
                                             0.4955
                                                       0.5074
        5
             hCV26311749
                              SD
                                         0
                                               0.273
                                                       0.2902
        5
               hCV918000
                            GENO
                                       2/2
                                                 2/1
                                                           1/1
               hCV918000 COUNTS
        5
                                        47
                                                 237
                                                           359
        5
               hCV918000
                            FREQ
                                   0.07309
                                              0.3686
                                                       0.5583
        5
               hCV918000
                                     0.505
                                              0.5091
                            MEAN
                                                       0.5074
        5
               hCV918000
                              SD
                                    0.2867
                                              0.3064
                                                       0.2797
```

i.e. each SNP takes up 5 rows.

## 8.9 Quantitative trait interaction (GxE)

PLINK provides the ability to test for a difference in association with a quantitative trait between two environments (or, more generally, two groups). This test is simply based on comparing the difference between two regression coefficients. To perform this test:

```
plink --file mydata --gxe --covar mycov.dat
where mycovar.txt is a file containing the following fields:
    Family ID
    Individual ID
    Covariate value
See the notes on covariate files for more details.
This option will generate the file
```

```
plink.qassoc.gxe
which contains the fields:
```

CHR	Chromosome number
SNP	SNP identifier
NMISS1	Number of non-missing genotypes in first group (1)
BETA1	Regression coefficient in first group

```
SE1 Standard error of coefficient in first group
NMISS2 As above, second group
BETA2 As above, second group
SE2 As above, second group
Z_GXE Z score, test for interaction
P_GXE Asymptotic p-value for this test
```

**IMPORTANT!** The covariate must be coded as an affection status variable, i.e. 1 or 2 representing the first or second group. Values of 0 or -9 can be used to indicate missing covariate values, in which case that individual will be excluded from analysis.

### 8.10 Linear and logistic models

These two features allow for multiple covariates when testing for both quantitative trait and disease trait SNP association, and for interactions with those covariates. The covariates can either be continuous or binary (i.e. for categorical covariates, you must first make a set of binary dummy variables).

**WARNING!** These commands are in some ways more flexible than the standard --assoc command, but this comes with a price: namely, these run more slowly...

In this section we consider:

- Basic uasge
- Covariate and interactions
- Flexibly specifying the precise model
- Flexibly specifying joint tests

#### 8.10.1 Basic usage

For quantitative traits, use

or

```
plink --bfile mydata --linear
```

For disease traits, specify logistic regression with

```
plink --bfile mydaya --logistic
```

instead. All other commands in this section apply equally to both these models.

These commands will either generate the output file

```
plink.assoc.linear
```

plink.assoc.logistic

depending on the phenotype/command used. The basic format is:

```
CHR Chromosome

SNP SNP identifier

TEST Code for the test (see below)

NMISS Number of non-missing individuals included in analysis

BETA Regression coefficient (--linear) or odds ratio (--logistic)

STAT Coefficient t-statistic

P Asymptotic p-value for t-statistic
```

For the additive effects of SNPs, the direction of the regression coefficient represents the effect of each extra **minor allele** (i.e. a positive regression coefficient means that the minor allele increases risk/phenotype mean). The **--freq** command will indicate which allele is the minor one.

HINT Adding the --ci 0.95, for example, option will given 95% confidence intervals for the estimated parameters, in additional L95 and U95 fields in the output files.

By itself, the --linear command will give identical results to the Wald test from the --assoc command when applied to quantitative traits. The --logistic command may give slightly different results to the --assoc command for disease traits, but this is because a different test/model is being applied (i.e. logistic regression rather than allele counting). The difference may be particularly large for very rare alleles (i.e. if the SNP is monomorphic in cases or controls, then the logistic regression model is not well-defined and asymptotic results might not hold for the basic test either).

The TEST column is by default ADD meaning the additive effects of allele dosage. Adding the option

```
--genotypic
```

will generate file which will have two extra tests per SNP, corresponding to two extra rows: DOMDEV and GENO\_2DF which represent a separate test of the dominance component or a 2 df joint test of both additive and dominance (i.e. corresponding the the general, genotypic model in the --model command). Unlike the dominance model is the --model, DOMDEV refers to a variable coded 0,1,0 for the three genotypes AA,Aa,aa, i.e. representing the dominance deviation from additivity, rather specifying that a particular allele is dominant or recessive. That is, the DOMDEV term is fitted jointly with the ADD term in a single model.

**NOTE!** The coding PLINK uses with the 2 df --genotypic model involves two variables representing an additive effect and a dominance deviation;

A D
AA 0 0
AB 1 1
BB 2 0

Although the 2df test will be identical, you would **not** expect to see similar p-values, etc for the two individual terms if instead you used a different version of "genotypic" coding, e.g. in another analysis package, such as using dummy variables to represent genotypes:

G1 G2
AA 0 0
AB 1 0
BB 0 1

That is, although fundamentally the same, in terms of the 2df test, the interpretation of the two individual terms is different in these two cases. To achieve this coding in PLINK (v1.02 onwards), add the --hethom flag as well as --genotypic.

In a related note, you would not always expect the ADD p-value to be the same when entering in the dominance term as it is without it; if in doubt, you are advised to stick to just interpreting the 2 df test if using the --genotypic option.

To specify a model assuming full dominance (or recessive) for the minor allele (i.e. rather than the 2 df model mentioned above), you can specify with either

```
--dominant or --recessive
```

#### 8.10.2 Covariates and interactions

If a covariate file is also specified, then **all** covariates in that file will be included in the regression model, labelled COV1, COV2, etc. Note: this is different to other commands which take only a single covariate

(possibly working in conjunction with the --mcovar option). For example, if the covariate file is made as described here and contains 2 covariates then the command

```
plink --bfile mydata --linear --covar mycov.txt
```

will add two extra tests per SNP, COV1 and COV2. The p-value for the SNP term or terms in the model will be adjusted for the covariates; that is, a single model is fit to the data

```
Y = b0 + b1.ADD + b2.DOMDEV + b3.COV1 + b4.COV2 + e
```

(Note, using this notation, the genotypic test is of b1=b2=0).

**HINT** To condition analysis on a specific SNP when using --linear or --logistic, use the --condition option, e.g.

```
plink --bfile mydata --linear --condition rs123456
```

will test all SNPs but adding the allelic dosage for rs123456 as a covariate. This command can be used in conjunction with --covar and the other options listed here. To condition on multiple SNPs, use, for example,

```
plink --bfile mydata --linear --condition-list snps.txt
```

where snps.txt is a plain text file contain a list of SNPs which are to be included as covariates. The output will now include terms that correspond to the SNPs listed in the file snps.txt.

The conditioning SNPs are entered into the model simply as covariates, using a simple 0, 1, 2 allele dosage coding. That is, for two conditioning SNPs, rs1001 and rs1002 say, and also a standard covariate, the model would be

```
Y = b0 + b1.ADD + b2.rs1001 + b3.rs1002 + b4.COV1 + e
```

If the b1 coefficient for the test SNP is still significant after entering these covariates, this would suggest that it does indeeed have an effect independent of rs1001, rs1002 and the other covariate. (The other coefficients may still be highly significant, but these reflect the effects of the conditioning SNPs and covariates, not the test SNP.)

If the --sex flag is added, then sex will be entered as a covariate in the model (coded 1 for male, 0 for female), e.g

```
plink --bfile mydata --logistic --sex
```

If the option --interaction is added, then terms will be entered which correspond to SNP x covariate interactions (with DOMDEV as well as ADD if --genotypic is specified). In the case of two covariates, without --genotypic, for example, the command

```
plink --bfile mydata --linear --covar tmp.cov --interaction
```

results in the model

```
Y = b0 + b1.ADD + b2.COV1 + b3.COV2 + b4.ADDxCOV1 + b5.ADDxCOV2 + e
```

**NOTE** Please remember that when interaction terms are included in the model, the significance of the main effects can not necessarily be interpreted straightforwardly (i.e. they will depend on the arbitrary coding of the variables). In otherwords, when including the --interaction flag, you should probably only interpret the interaction p-value. Please refer to any standard text of regression models if you are unclear on this.

Finally, a --test-all option drops all the terms in the model in a multiple degree of freedom test.

#### 8.10.3 Flexibly specifying the model

Use command such as --covar and --interaction will automatically enter all covariates and possible SNP x covariate interactions. If one does not want to test all of these, then use the --parameters flag to extract only the ones of interest.

For example, to take the example above:

```
Y = b0 + b1.ADD + b2.COV1 + b3.COV2 + b4.ADDxCOV1 + b5.ADDxCOV2 + e
```

If one only wanted ADD, the two covariates and the ADDxCOV2 but **not** the ADDxCOV1 interaction, then, from the above example, you could use

```
plink --bfile mydata --linear --covar tmp.cov --interaction --parameters 1,2,3,5
```

That is, --parameters takes a comman-separated list of integers, starting from 1, that represent the terms in the model (in the order in which they would appear if the command were run without the --parameters flag). In this case:

```
ADD [1]
COV1 [2]
COV2 [3]
ADD x COV1 [4] <-- excluded
ADD x COV2 [5]
```

#### 8.10.4 Flexibly specifying joint tests

To perform a user-defined joint test of more than one parameter, use the --tests option. This takes a comma-delimited set of parameter numbers, for example: if the model is

ADD	[1]
COV1	[2]
COV2	[3]
ADDxCOV1	[4]
ADDxCOV2	[5]

then

```
plink --bfile mydate --linear --covar file.cov --interaction --tests 1,4,5
```

represents a 3 degree of freedom test of ADD and the two interactions.

Note, if this is used in conjunction with the --parameters option, then the coding here refers to the reduced model – for example, the command

```
plink --bfile mydate --linear --covar file.cov --interaction --parameters 1,2,3,5
--tests 1,4
```

performs a joint test of ADD and ADDxCOV2 (2df test) whilst controlling for main effects of COV1 and COV2, i.e. we *do not* use --tests 1,5, as there are now only 4 terms in the model:

		parameters 1,2,3,5	tests 1,4
ADD	[1]	[1]	TEST
COV1	[2]	[2]	
COV2	[3]	[3]	
ADDxCOV1	[4]	n/a	
ADDxCOV2	[5]	[4]	TEST

In other words, we fit the model

```
Y = b0 + b1.ADD + b2.COV1 + b3.COV2 + b4.ADDxCOV2 + e
```

and jointly test the hypothesis

```
H0: b1 = b4 = 0
```

As mentioned above, use --test-all to drop all terms in the model in a single joint test.

#### 8.10.5 Multicollinearity

A common problem with multiple regression is that of multi-collinearity: when the predictor variables are too strongly correlated to each other, the parameter estimates will become unstable. PLINK tries to detect this, and will display NA for the test statistic and p-value for all terms in the model if there is evidence of multi-collinearity. One common instance where this would occur would be if one includes the --genotypic option but a SNP only has two of the three possible genotype classes: in this case, ADD and DOM will be perfectly correlated and PLINK will display NA for both tests; this is basically telling you that you should rerun without the --genotypic option for that particular SNP. Similar principles apply to including covariates and interactions terms: the more terms you include, the more likely you are to have problems.

The --vif option can be used to specify the variance inflation factor (VIF) used in the initial test for multicollinearity. The default value is 10 – smaller values represent more stringent tests.

HINT If you have a quantitative trait, only want an additive model and have only a single binary covariate, use the --gxe option (described above) instead of --linear: it will run much faster (being based on a more simple test of the difference of two regression slopes; it will not necessarily give numerically identical results to the multiple regression approach, but asymptotically both tests should be similar).

#### 8.11 Set-based tests

These set-based tests are particularly suited to large-scale candidate gene studies as opposed to whole genome association studies, as they use permutaiton.

**NOTE** The basis of the set-based test has been changed in version 1.04 onwards. This analysis works as follows:

- For each set, for each SNP determine which other SNPs are in LD, above a certain threshold R
- Perform standard single SNP analysis (which might be basic case/control association, family-based TDT or quantitative trait analysis).
- For each set, select up to N "independent" SNPs (as defined in step 1) with p-values below P. The best SNP is selected first; subsequent SNPs are selected in order of descreasing statistical significance, after removing SNPs in LD with previously selected SNPs.
- From these subsets of SNPs, the statistic for each set is calculated as the mean of these single SNP statistics
- Permute the dataset a large number of times, keeping LD between SNPs constant (i.e. permute phenotype labels)
- For each permuted dataset, repeat steps 2 to 4 above.
- Empirical p-value for set (EMP1) is the number of times the permuted set-statistic exceeds the original
  one for that set.

Note that the empirical p-values are corrected for the multiple SNPs within a set (taking account of the LD between these SNPs). They are not corrected for multiple testing if there is more than one set, however (i.e. there is no equivalent of EMP2 (see the page on permutation).

The critical parameters described above, R, N and P can all be altered by the user, as described below. To perform a set-based test the critical keywords are

```
--set-test
--set my.set
--mperm 10000
```

which state that we are performing a set-based test, which set-file to use and how many permutations to perform (this last command is necessary). As mentioned above, the **--assoc** command could be replaced by **--tdt**, or **--logistic**, etc.

```
The set file my.set is in form
```

```
SET1
rs1234
rs28384
rs29334
END
SET2
rs4774
rs662662
rs77262
END
```

For example,

```
plink --file mydata --set-test --set my.set --mperm 10000 --assoc
```

would display in the LOG file the following critical parameters with their default values

Performed LD-based set test, with parameters:

```
r-squared (--set-r2) = 0.5
p-value (--set-p) = 0.05
max # SNPs (--set-max) = 5
```

The output is written to a file with a .set.mperm extension, for example

```
plink.assoc.set.mperm
```

with the fields

SET	Set name
NSNP	Number of SNPs in set
NSIG	Total number of SNPs below p-value threshold
ISIG	Number of significant SNPs also passing LD-criterion
STAT	Average test statistic based on ISIG SNPs
EMP1	Empirical set-based p-value
SNPS	List of SNPs in the set

For example, here is output from a case/control dataset with SNPs for five related genes (lines truncated)

SET	NSNP	NSIG	ISIG	STAT	EMP1	SNPS
GABRB2	45	0	0	0	1	NA
GABRA6	6	4	3	5.199	0.09489	rs3811991 rs2197414
GABRA1	22	11	5	5.951	0.09459	rs4254937 rs4260711
GABRG2	24	0	0	0	1	NA
GABRP	17	2	1	7.64	0.0269	rs7736504

Here the first gene, GABRB2 has 45 SNPs, but none of these are significant at p=0.05, and so the empirival p-value is necessarily 1.00. The next gene has 6 SNPs, 4 of which are significant, but only 3 of which are independently significant based on an r-squared threshold of 0.5. The STAT of 5.199 is the average chi-squared statistic across these three SNPs. It should not be interpreted in itself – rather, you should consider the EMP1 significance value based on it. In this case, P=0.095. The final gene, GABRP is nominally significant here, P=0.027, but this does not correct for the 5 genes tested of course.

Naturally, different thresholds will produce different results. Depending on the unknown genetic architecture, these may vary substantially and meaningfully so. In general, if the set represents a very large pathway

(dozens of genes) you might want to increase --set-max. There are probably no hard and fast rules with regard to how to set --set-p and --set-r2, except to say that running under a large number of settings and selecting the most significant is not a good idea.

Running with a "stricter" set of values

```
--set-r2 0.1
--set-p 0.01
--set-max 2
```

we see a broadly similar pattern of results; naturally, the thresholding on p-value means that GABRA6 goes from showing some signal to as bolutely no signal.

SET	NSNP	NSIG	ISIG	STAT	EMP1	SNPS
GABRB2	45	0	0	0	1	NA
GABRA6	6	0	0	0	1	NA
GABRA1	22	2	2	7.464	0.05949	rs4254937 rs4260711
GABRG2	24	0	0	0	1	NA
GABRP	17	1	1	7.64	0.06309	rs7736504

Alternatively, a more inclusive setting might be something like

```
--set-r2 0.8
--set-p 1
--set-max 10
```

which, in this particular case, happens to yield slightly stronger signals for GABRA6 and GABRA1 but weaker for GABRp (lines truncated)

SET	NSNP	NSIG	ISIG	STAT	EMP1 SNPS
GABRB2	45	12	10	1.749	0.7162 hCV26311691
GABRA6	6	6	6	3.998	0.0184 rs3811991
GABRA1	22	13	10	5.277	0.0182 rs4254937
GABRG2	24	11	10	0.6976	0.9099 hCV3167705
GABRP	17	10	10	2.753	0.1225 rs7736504

HINT Two extremes are to perform a test based on a) the best single SNP result per set:

```
--set-max 1
--set-p 1
or to use all SNPs in a set:
--set-max 99999
--set-p 1
--set-r2 1
```

# 8.12 Adjustment for multiple testing: Bonferroni, Sidak, FDR, etc

To generate a file of adjusted significance values that correct for all tests performed, use the option:

UNADJ Unadjusted p-value

BONF Bonferroni single-step adjusted p-values

HOLM Holm (1979) step-down adjusted p-values

SIDAK\_SS Sidak single-step adjusted p-values

SIDAK\_SD Sidak step-down adjusted p-values

FDR\_BH Benjamini & Hochberg (1995) step-up FDR control

FDR\_BY Benjamini & Yekutieli (2001) step-up FDR control

This file is sorted by significance value rather than genomic location, the most significant results being at the top.

WARNING Currently, these procedures are only implemented for asymptotic significance values for the standard TDT and association (disease trait and quantitative trait, --assoc, --linear, --logistic) tests and the 2x2xK Cochran-Mantel-Haenszel test. Future versions will allow these results for empirical significance values and for other tests (e.g. epistasis, etc).

## Chapter 9

L95

**U95** 

## Family-based association analysis

The main focus of PLINK is for population-based samples. There is some support for family-based analyses however, described in this section, for disease traits and quantitative traits.

### 9.1 Family-based association (TDT)

PLINK supports basic family-based association testing for disease traits, using the TDT and a variant of this test that also incorporates parental phenotype information, the *parenTDT*.

To run a basic TDT analysis for family data:

```
plink --file mydata --tdt
which generates the file
     plink.tdt
If permutation has been requested, then either
     plink.tdt.perm
or
     plink.tdt.mperm
will be generated also. The main output file, plink.tdt, contains the following fields:
     CHR
                  Chromosome number
     SNP
                  SNP identifier
     Α1
                  Minor allele code
     A2
                  Major allele code
     Τ
                  Transmitted minor allele count
     U
                  Untransmitted allele count
     OR
                  TDT odds ratio
     CHISQ
                  TDT chi-square statistic
     Ρ
                  TDT asymptotic p-value
     A:U_PAR
                 Parental discordance counts
     CHISQ_PAR Parental discordance statistic
     P_PAR
                 Parental discordance asymptotic p-value
     CHISQ_COM
                 Combined test statistic
     P_COM
                 Combined test asymptotic p-value
```

If the --ci option has been requested, then two additional fields will appear after TDT\_OR:

Lower 95% confidence interval for TDT odds ratio

Upper 95% confidence interval for TDT odds ratio

(naturally, if a value other than 0.95 was used as the argument for the --ci option, it will appear here instead.)

The TDT statistic is calculated simply as

$$(b-c)\hat{2} / (b+c)$$

where b and c are the number of transmitted and untransmitted alleles as shown in plink.tdt; under the null, it is distributed as a 1df chi-squared.

The parental discordance test is based on counting the number of alleles in affected versus unaffected parents, treating each nuclear family parental pair as a matched pair. These counts can be combined with the T and U counts of the basic TDT to give a combined test statistic, also shown in the output. The parenTDT assumes homogeneity within families rather than between families, in terms of population stratification. If parents are measured on the phenotype, then this test can add considerable power to family-based association analysis, whilst providing a strong degree (but not complete) protection against population stratification. The increase in power will depend on the proportion of parents that are discordant for the disease. This approach is described in Purcell et al AJHG (2005). PLINK uses a more simple approach to calculate the PAR and COM statistics, however: if

```
Unaffeced parent A/A A/B B/B Affected A/A - p r parent A/B x - q B/B z y - i.e. such that the A:U_PAR fields represents p+q+2r: x+y+2z, then pAR = ((p+q+2r) - (x+y+2z))\hat{2}
```

/ ( p+q+x+y+4(r+z) )

and

or

```
COM = ((b+p+q+2r) - (c+x+y+2z))^2
/ (b+p+q+c+x+y+4(r+z))
```

Both statistics follow a 1 df chi-squared distribution under the null.

When running the --tdt option, PLINK will first perform a check for Mendel errors and make missing the offending genotypes.

Using the --tdt option, if permutation is requested (using either --perm or --mperm) a file entitled either plink.tdt.perm

```
plink.tdt.mperm
```

will be generated: the empirical p-value will be based on the standard TDT test. The permutation procedure will flip transmitted/untransmitted status constantly for all SNPs for a given family, thereby preserving the LD and linkage information between markers and siblings.

## 9.2 parenTDT

The parenTDT, described in the paragraph above, is automatically included when using the --tdt option. These alternate commands generate the same output as for the --tdt command, described above, except the permutation is based not on the standard TDT, but either the parenTDT if using the option

```
plink --file mydata --parentdt1
or, the combined test (TDT and parenTDT) if using the option
plink --file mydata --parentdt2
```

### 9.3 Parent of origin analysis

When performing family-based TDT analysis, it is possible to separately consider transmissions from heterozygous fathers versus heterozygous mothers to affected offspring. This is performed by adding the --poo to request parent-of-origin analysis:

```
plink --file mydata --tdt --poo
```

which generates the file plink.tdt.poo. If permutation is also requested, this also generates the file plink.tdt.poo.perm or plink.tdt.poo.mperm, depending which permutation procedure is used. The main output file has the following format:

```
CHR
           Chromosome number
SNP
           SNP identifier
A1:A2
           Allele 1 : allele 2 codes
T:U PAT
           Paternal transmitted: untransmitted counts
OR_PAT
           Paternal odds ratio
CHISQ_PAT Paternal chi-squared test
           Maternal, as above
T:U_MAT
           Matneral, as above
OR_MAT
CHISQ_MAT Maternal, as above
Z_P00
           Z score for difference in paternal versus maternal odds ratios
P_P00
           Asymptotic p-value for parent-of-origin test
```

If permutation is requested, the default test statistic is the absolute value of the Z score for the parent-of-origin test (i.e. making a two-sided test). The flags --pat and --mat indicate that the permutation statistic should be the paternal TDT chi-squared statistics, or the maternal statistic, instead.

**NOTE** When both parents are heterozygous, these ambiguous transmissions are counted as 0.5 for both mother and father – this is why the T:U counts will often not be whole numbers.

## 9.4 DFAM: family-based association for disease traits

The DFAM procedure in PLINK implements the sib-TDT and also allows for unrelated individuals to be included (via a clustered-analysis using the Cochran-Mantel-Haesnzel). To perform this test:

```
which generates the file
     plink.dfam
which contains the fields
     CHR
                 Chromosome code
     SNP
                 SNP identifier
     A1:A2
                 Minor and major allele codes
     OBS
                 Number of observed minor alleles
     EXP
                 Number of expected minor alleles
     CHISQ
                 Chi-squared test statistic
                 Asymptotic p-value
```

plink --bfile mydata --dfam

This test can therefore be used to combine discordant sibship data, parent-offspring trio data and unrelated case/control data in a single analysis.

**NOTE** If you are analysing a sibling-only sample (i.e. no parents) then also add the **--nonfounders** option; otherwise, all SNPs will be pruned out at the filtering stage, as PLINK will by default only consider founder alleles when calculating allele frequency, Hardy-Weinberg, etc.

### 9.5 QFAM: family-based association tests for quantitative traits

PLINK offers a somewhat ad-hoc procedure to perform family-based tests of association with quantitative phenotypes: the QFAM procedure, which uses permutation to account for the dependence between related individuals. It adopts the between/within model as used by Fulker et al (1999, AJHG) and Abecasis et al (2000, AJHG) as implemented in the QTDT package. However, rather than fitting a maximum likelihood variance components model, as QTDT does, PLINK performs a simple linear regression of phenotype on genotype, but then uses a special permutation procedure to correct for family structure.

There are several ways to run QFAM: a total association test (between and within components)

```
plink --bfile mydata --qfam-total --mperm 100000

or a within-family test

plink --bfile mydata --qfam --mperm 100000

or a test including parental phenotypes

plink --bfile mydata --qfam-parents --mperm 100000

(Also, --qfam-between will look only at the between-family component of association).
```

**NOTE** In all cases above, we have used **--mperm** to specify permutation; adaptive permutation can also be used with QFAM (**--perm**). Permutation is necessary for the QFAM test.

The columns in the QFAM permutation result files are:

```
CHR Chromosome code
SNP SNP identifier
STAT Test statistic (ignore)
EMP1 Pointwise empirical p-value
NP Number of permutations performed
```

The columns in the non-permutation file (e.g. plink.qfam.total, if plink.qfam.total.mperm contains the permuted results) are as follows:

```
CHR Chromosome code
SNP SNP identifier
A1 Minor allele (corresponds to beta given below; absent in earlier PLINK releases)
TEST Type of test, TOT, WITH and BET
NMISS Number of non-missing individuals in analysis
BETA Regression coefficient
STAT Test statistic (ignore; not corrected for family-structure)
P Asymptotic p-value (ignore; use empirical p-value)
```

These results are from a standard --linear type analysis, i.e. which ignores family structure. They are displayed so that the direction of effect may be determined (from the BETA) – but otherwise, only the empirical p-value from the permuted results file should be looked at.

The B and W components are calcalated using parental genotypes if they are available for both parents, otherwise siblings are used. Singletons can be included in this analysis (i.e. B=G and W=0 for them): for example, the scores are shown below for a few configurations, when parents are available:

```
Genotype G

AA 1

Aa 0

aa -1

B = ( Pat + Mat ) / 2

W = G - B

Pat Mat Offspring G B W

AA AA AA 1 1 1 0
```

```
AA
                    AA
                                             0.5
                                                    0.5
Aa
                                     1
                                     0
                                             0.5
                                                    -0.5
Аa
          AΑ
                    Aa
          AΑ
                                             0
                                                    0
ลล
                    Aа
```

The QFAM permutation procedure breaks down the genotypes into between (B) and within (W) components, permutes them independently (i.e. at the family level, either swapping the B component for one family with another family, or flipping the sign of all W's in a family with 50:50 chance) and then (for the total association test) reconstructs the individual level "genotypes" as the sum of the new B's and W's i.e:

```
1) G -> B + W (individual-level)
2a) Permute B (family-level) -> B'
2b) Permute W (family-level) -> W'
3) B' + W' -> G' (individual-level)
```

The logic is that we know how to permute both B and W separately whilst maintaining the familial structural component, and they are orthogonal components, so we should permute them separately, but then recombine them as a single individual-level genotypic score.

**NOTE** The total --qfam-total test is designed to extract all association information from a family-based sample, controlling for relatedness: it is not robust to stratification. Use the --qfam for a strictly within family test.

In many circumstances, the standard QTDT as implemented in Goncalo Abecasis' QTDT http://www.sph.umich.edu/csg/abecasis/QTDT/ program will perhaps be more appropriate. The disadvantages of the QFAM procedure are

- that it uses permutation and so is slower
- appears to be slightly less powerful when there is a higher residual correlation

On the plus side, the advantages of the QFAM procedure are

- that is uses permutation and so is appropriate for non-normal phenotypes; it could also be used for disease phenotypes, although it will not be appropriate for affected-only TDT style designs
- that it can be applied to genome-wide data easily (albeit not necessarily quickly)

**Technical note** As a technical point: when permuting genotype between families in this way, one has to be careful with missing genotype data, particularly in the instance in which a family is completely missing. Because a missing B component cannot be recombined with a non-missing W component, and vice versa, this process would tend to increase the amount of missingness in permutations versus the original data.

One could exclude individuals with missing genotypes first and permute separately for each SNP, but this would no longer maintain the correlation between SNPs (and require more computation). Instead, we use the following scheme. We permute once per replicate (e.g. a table of F (original family) and F' (permuted family), true and permuted families). e.g. but let's say that 2 is missing their B component (denoted 2\*) For example:

```
F
   F,
0
   5
1
   2*
        <- remove ?
2*
   4
        <- remove ?
3
   1
4
   0
5
   3
```

This would knock out families 1 and 2 from the permutation. We therefore permute once to create a single table for permutation for all SNPs, but then resursively edit the table on a SNP-by-SNP basis, to

regroup the missing families, by swapping missing F' families: in this case, swap  $2^*$  with 4 (the other partner of  $2^*$ ), e.g.

So now we have a permuted sample but the total level of missingness is the same. This procedure still generates valid, completely random permutations of the non-missing genotype data and trys to maintain as much of the correlation between SNPs as possible (i.e. as typically only a small % of genotypes are missing and so we do not need to edit the table much).

## Chapter 10

## Permutation procedures

Permutation procedures provide a computationally intensive approach to generating significance levels empirically. Such values have desirable properties: for example, relaxing assumptions about normality of continuous phenotypes and Hardy-Weinberg equilibrium, dealing with rare alleles and small sample sizes, providing a framework for correction for multiple testing, and controlling for identified substructure or familial relationships by permuting only within cluster.

#### 10.0.1 Conceptual overview of permutation procedures

Permutation procedures are available for a variety of tests, as described below. For some tests, however, these procedures are not available (e.g. SNP x SNP epistasis, Cochran-Mantel-Haenszel tests). For other tests, permutation is necessary to obtain any significance values at all (e.g. set-based tests).

The permutation tests described below come can be categorized in two ways:

- Label-swapping versus gene-dropping
- Adaptive versus max(T)

#### 10.0.2 Label-swapping and gene-dropping

In samples of unrelated individuals, one simply swaps labels (assuming that individuals are interchangeable under the null) to provide a new dataset sampled under the null hypothesis. Note that only the phenotype-genotype relationship is destroyed by permutation: the patterns of LD between SNPs will remain the same under the observed and permuted samples. For family data, it might be better (or in the case of affected-only designs such as the TDT, necessary) to perform gene-dropping permutation instead. In it's most simple form, this just involves flipping which allele is transmitted from parent to offspring with 50:50 probability. This approach can extend to general pedigrees also, dropping genes from founders down the generations.

For quantitative traits, or samples in which both affected and unaffected non-founders are present, one can then perform a basic test of association (with disease, or with a quantitative trait) treating the pedigree data as if they were all unrelated (i.e. just using the --assoc option) but creating permuted datasets by gene-dropping will both control for stratification and the non-independence of related individuals (i.e. as these will also be properties of every permuted dataset). It is possible to maintain LD between SNPs by applying the same series of 50:50 flip/no-flip decisions to all SNPs in the same permuted replicate for a given transmission. In addition, it is possible to control for linkage by applying the same series of flip/no-flip decisions to all siblings in the same nuclear family. Both these features are automatically applied in PLINK.

### 10.0.3 Adaptive and max(T) permutation

Using either label-swapping or gene-dropping, there are two basic approaches to performing the permutations. The default mode is to use an *adaptive* permutation approach, in which we give up permuting SNPs that

are clearly going to be non-significant more quickly than SNPs that look interesting. In otherwords, if after only 10 permutations we see that for 9 of these the permuted test statistic for a given SNP is larger than the observed test statistic, there is little point in carrying on, as this SNP is incredibly unlikely to ever achieve a highly significant result. This greatly speeds up the permutation procedure, as most SNPs (that are not highly significant) will drop out quite quickly, making it possible to properly evaluate significance for the handful of SNPs that require millions of permutations. Naturally, the precision with which one has estimated the significance p-value (i.e. relating from the number of permutations performed) will be correlated the significance value itself – but for most purposes, this is precisely what one wants, as it is of little interest whether a clearly un-associated SNP really has a p-value of 0.78 or 0.87.

In contrast, max(T) permutation does not drop SNPs along the way. If 1000 permutations are specified, then all 1000 will be performed, for all SNPs. The benefit of doing this is that two sets of empirical significance values can then be calculated – pointwise estimates of an individual SNPs significance, but also a value that controls for that fact that thousands of other SNPs were tested. This is achieved by comparing each observed test statistic against the maximum of all permuted statistics (i.e. over all SNPs) for each single replicate. In otherwords, the p-value now controls the familywise error rate, as the p-value reflects the chance of seeing a test statistic this large, given you've performed as many tests as you have. Because the permutation schemes preserve the correlational structure between SNPs, this provides a less stringent correction for multiple testing in comparison to the Bonferroni, which assumes all tests are independent. Because it is now the corrected p-value that is of interest, it is typically sufficient to perform a much smaller number of tests – i.e. it is probably not necessary to demonstrate that something is genome-wide significant beyond 0.05 or 0.01.

#### 10.0.4 Computational issues

PLINK performs the basic tests of association reasonably quickly – for small datasets both permutation procedures will be feasible. For example, for a dataset comprising 100,000 SNPs measured on 350 individuals, each permutation (for all 100K SNPs) takes approximately 2 seconds on a modern Linux workstation. At this speed, it will take just over 1 day to perform 50,000 permutations using the max(T) mode and label-swapping. With the same dataset, using adaptive mode, the entire analysis is finished much quicker (although the empirical p-values are, of course, not corrected for multiple testing). For larger datasets (e.g. 1000s of individuals measured on ¿500K SNPs) things will slow down, although this will be linear with the number of genotypes – if one has access to a cluster, however, the max(T) approach lends itself to easy parrallelization (i.e. if one can set many jobs running analysing the same data, it is easy to combine the empirical p-values afterwards).

By default, PLINK will select a random seed for the permutations, based on the system clock. To specify a fixed seed instead add the command

```
--seed 6377474
```

where the parameter a (large) integer seed.

## 10.1 Basic (adaptive) permutation procedure

The default method for permutation is the adaptive method. To obtain a max(T) permutation p-value, see the section below. For either either case/control or quantitative trait association analysis, use the option:

```
plink --file mydata --assoc --perm
```

to initiate adaptive permutation testing. As well as the plink.assoc or plink.qassoc output file, adding the --perm option will generate a file named:

```
plink.assoc.perm
```

which contains the fields:

```
CHR Chromosome
SNP SNP ID
STAT Test statistic
EMP1 Empirical p-value (adaptive)
NP Number of permutations performed for this SNP
```

An alternate scheme is also available, that may under some circumstances be useful. Specifically, this approach fixes the observed marginal counts for the 2-by-3 tables that is case/control status by the two alleles and the missing allele count. After permuting case/control label, only two cells in the table, e.g. missing and A2 alleles for controls, are counted, the rest of the table is filled in on the basis of the fixed marginal values. This speeds up the permutation procedure a little, and also implicitly downweights association results where there is a lot of missing genotype data that is non-random with respect to genotype and case/control status. Naturally, this approach can not provide total protection against the problem of non-random missing genotype data. Also, for SNPs with lots of missing data, this test will be conservative, whether the missingness is non-random or not. For these reasons, this is not the default option, although this approach might be one worth exploring further. To use this alternate permutation scheme, use the --p2 flag:

```
plink --file mydata --assoc --perm --p2
or
plink --file mydata --assoc --mperm 1000 --p2
```

### 10.2 Adaptive permutation parameters

Although the --perm option invokes adaptive permutation by default, there are various parameters that alter the behavior of the adaptive process that can be tweaked using the --aperm option, followed by six parameters: for example,

```
plink --file mydata --assoc --aperm 10 1000000 0.0001 0.01 5 0.001
```

The six arguments (along with the default values) are:

```
Minimum number of permutations per SNP 5

Maximum number of permutations per SNP 1000000

Alpha level threshold (alpha) 0

Confidence interval on empirical p-value (beta) 0.0001

Interval to prune test list (intercept) 1

Interval to prune test list (slope) 0.001
```

These are interpreted as follows: for every SNP, at least 5 permutations will be performed, but no more than 1000000. After 5 permutations, the p-values will be evaluated to see which SNPs we can prune. The first interval value means to perform this pruning every 5 replicates; the second pruning parameter (0.001) means that the rate of pruning slows down with increasing number of replicates (i.e. pruning is, in this case, performed every 5+0.001R replicates where R is the current number of replicates). At each pruning stage, a 100\*(1 - beta / 2T)% confidence interval is calculated for each empirical p-value, where beta is, in this case 0.01, and T is the number of SNPs. Using the normal approximation to the binomial, we prune any SNP for which the lower confidence bound is greater than alpha or the upper confidence bound is less than alpha.

## 10.3 max(T) permutation

To perform the max(T) permutation procedure, use the --mperm option, which takes a single paramter, the number of permutations to be performed: e.g. to use with the TDT test:

```
plink --file mydata --tdt --mperm 5000
which will generate (along with the plink.tdt file) an file named
    plink.tdt.mperm
which contains the fields:
    CHR     Chromosome
    SNP     SNP ID
    STAT     Test statistic
    EMP1     Empirical p-value (pointwise)
    EMP2     Corrected empirical p-valie (max(T) / familywise)
```

Hint If multiple runs of PLINK are performed on the same dataset in parallel, using a computer cluster to speed up the  $\max(T)$  permutations, then the resulting estimates of empirical significance can be combined across runs as follows. Empirical p-values are calculated as (R+1)/(N+1) where R is the number of times the permuted test is greater than the observed test; N is the number of permutations. Therefore, therefore, given  $p_i$ , the empirical p-value for the ith run, this implies that  $p_i*(N_i+1)-1$  replicates passed the observed value. The overall empirical p-value should then be:

```
(SUM_i p_i * (N_i + 1) - 1 + 1) / (SUM_i N_i + 1)
```

To produce output files that contain either the best statistic per replicate, or all statistics per replicate, use either option

```
--mperm-save
or
--mperm-save-all
along with the usual --mperm command. The first command generates a file
plink.mperm.dummp.best
```

which contains two columns. The first is the replicate number (0 represents the original data, the remaining rows 1 to R where R is the number of permutations specified). The second column is the maximum test statistic over all SNPs for that replicate. The second command, --mperm-save-all produces a file

```
plink.mperm.dump.all
```

that could be a very large file: the test statistic for all SNPs for all replicates. As before, the first row is the original data; the first column represents the replicate number; all other columns represent the test statistic values for each SNP (NA if this cannot be calculated). These two files might be of use if, for example, you wish to create your own wrapper around PLINK to perform higher-order corrections for multiple testing, e.g. if more than one phenotype is tested per SNP. In most cases, for this purpose, the first form should suffice.

## 10.4 Gene-dropping permutation

To perform gene-dropping permutation, use the --genedrop option, combined with the standard --assoc option. Either adaptive: e.g.

```
plink --file mydata --assoc --genedrop
or max(T) permutation: e.g.
plink --file mydata --assoc --genedrop --mperm 10000
can be specified.
```

This analysis option is equally applicable to disease and quantitative traits, although at least some non-founder individuals should be unaffected. Currently, an individual must have both parents genotyped

for genedropping. For founders and for individuals without two genotyped parents, their genotypes are unchanged throughout all genedropping permutations.

It is possible to combine label-swapping with gene-dropping, however, to handle different family/sample configurations. That is, the basic gene-dropping procedure will leave untouched all individuals without two parents, making them uninformative for the test of association. One can think of at least three classes of groups of people without two parents in the dataset: founders/parents, siblings and unrelated singletons. Label-swapping within these groups can provide additional sources information for association that control different levels of the between/within family components of association.

There are three options, which can be used together, are:

--swap-sibs within family
--swap-parents partial within-family
--swap-unrel between family

which label-swap between sibs without genotyped parents (swapping only within families), between parents only (swapping only within families), or between all singletons (unrelated individuals) (swapping between families).

#### 10.4.1 Basic within family QTDT

This test only considers information from individuals with two genotyped parents:

```
plink --file mydata --assoc --genedrop
```

#### 10.4.2 Discordant sibling test

Although gene-dropping only considers individuals with two parents to be informative, valid family-based tests can include information from full-siblings – by label-swapping only within each full sibship that does not otherwise have parents, it is possible to augment the power of the gene-dropping approach:

```
plink --file mydata --assoc --genedrop --swap-sibs
```

## 10.4.3 parenTDT/parenQTDT

This test additionally incorporates information from phenotypically discordant parents (for either quantitative or disease triats). This provides more information for association, but provides a weaker level of protection against stratification (i.e. it assumes that mother and father pairs are well matched in terms of subpopulation stratum).

```
plink --file mydata --assoc --genedrop --swap-parents
```

#### 10.4.4 Standard association for singleton, unrelated individuals

If a sample is a mixture of families and unrelated individuals (e.g. case/control and offspring/parent trios combined) then adding this option as well as the --gene-drop option will perform label-swapping permutation for all unrelated individuals.

```
plink --file mydata --assoc --genedrop --swap-unrel
```

One or more of these options can be included with the **--genedrop** option. These features allow between and within family components of association to be included in analysis. Below are the results of some simple, proof-of-principle simulations, to illustrate parental discordance test:

Here is a subset of the results: in all cases, we have an unselected quantitative trait measured in parent/offspring nuclear families. The four models:

• no stratification, no QTL

- no stratification, QTL
- stratification between families (i.e. mother and father from same subpopulation), no QTL
- stratification within families (i.e. mother and father might not be from same subpopulation), no QTL

The three analytic procedures:

- standard QTL test (i.e. ignore family structure, which we know is incorrect)
- gene-dropping permutation (i.e. within family QTDT)
- gene-dropping + parental label-swapping (i.e. parenQTDT)

From simulation, the empirically estimated power/type I error rates (for a nominal value of 0.05) are:

```
500 trios (QT)
  Ι
        ΙI
              TTT
A 0.121 0.045 0.053
B 0.841 0.239 0.563
C 0.461 0.056 0.056
D 0.505 0.055 0.501
500 tetrads (QT)
  Ι
        ΙI
              III
A 0.173 0.043 0.050
B 0.900 0.363 0.653
C 0.439 0.042 0.045
D 0.390 0.044 0.421
```

That is,

- method I is, as expected, liberal (e.g. for tetrads, we see type I error rate of 17.3% instead of 5%). Subsequent values for this test should therefore be ignored in the table
- the parenQTDT (III) (as implemented by gene-dropping) is considerably more powerful than the standard within-family test that ignores parental phenotypes (II) i.e. 65% versus 36% for tetrads, in this particular instance.
- the parenQTDT is robust to stratification so long as it is between-family (condition C) i.e. it only assumes that mum and dad are matched on strata, not the whole sample. When this does not hold (condition D), then we get spurious association, as expected.

**HINT** For disease traits, the parenTDT test is automatically performed by the --tdt option (as long as there are at least 10 phenotypically discordant parental pairs in the sample). See the section of standard association testing for more details.

## 10.5 Within-cluster permutation

To perform label-swapping permutaion only within clusters, you must supply either a cluster file with the --within option, or indicate that family ID is to be used as the cluster variable, with the --family option. Then any label-swapping permutation procedure will only swap phenotype labels between individuals within the same cluster. For example,

```
plink --file mydata --assoc --within mydata.clst --perm
```

if the file mydata.clst were (for a PED file containing only 6 individuals, the file format is family ID, individual ID, cluster):

this would imply that only sets 1,2,3 and 4,5 could be permuted. That is, 1 and 3 could swap phenotypes, but not 1 and 5, for example. In this way, any between-cluster effects are preserved in each permuted dataset, which thereby controls for them.

To permute with family ID as the cluster variable for label-swapping, use the

```
plink --file mydata --assoc --family --perm
```

Note that label-swapping within families is different from gene-dropping. This approach would be appropriate for sibship data, for example, when no parents are available. The assumption is that individuals within family unit are interchangeable under the null – as such, you should not include mixtures of full siblings and half siblings, or siblings and parents, for example, in the same cluster using this approach.

Note Other options for stratified analyses are described on the previous page

## Chapter 11

## Multimarker haplotype tests

All tests described above are based on single SNP tests. It is also possible to impute haplotypes based on multimarker predictors using the standard E-M algorithm and to perform simple tests based on the distribution of probabilistically-inferred set of haplotypes for each individual.

As well as the autosomes, X and haploid chromosomes should be appropriately handled. Phasing can either be based on a sample of unrelated individuals, or certain kinds of family data. First, all founders are phased using the E-M algorithm; then all descendents of these founders are phased given the set of possible parental phases and assuming random-mating. Currently it is not possible to phase sibships without parents. The current implementation of the phasing and haplotype testing algorithm is designed focus on relatively small regions of the genome, rather than to phase whole chromosomes at once.

**HINT!** Another approach to haplotype-testing can be found under the page describing proxy association. This set of methods essentially just provide a different interface to the exact same E-M phasing and haplotype-testing algorithms, one that is centered around a specific reference SNP.

## 11.1 Specification of haplotypes to be estimated

Haplotype testing in PLINK requires that the user supplies a file listing the haplotypes to be tested (Some precomputed lists are given below which might be useful in some circumstances.) The formats of these files are described below. An alternative is to specify a simple, sliding window of fixed haplotype size (also described below).

The command

```
plink --file mydata --hap myfile.hlist
```

will read the file myfile.hlist, each row of which is expected to have one of the three following formats:

1) Particular allele specified

The first format specifies a particular haplotype at a given locus. Two example rows of this format are:

```
rs1001 5 0 201 1 2 TC snp1 snp2
rs1002 5 0 202 A C TTA snp1 snp3 snp4
```

The columns represent:

Col 1 : Imputed SNP name
Col 2 : Imputed SNP chromosome

Col 3 : Imputed SNP genetic distance (default: Morgan coding)

Col 4 : Imputed SNP physical position (bp units)

Col 5 : Imputed SNP allele 1 name

Col 6 : Imputed SNP allele 2 name

Col 7 : Tag SNP allele/haplotype that equals imputed SNP allele 1

Col 8+ : Tag SNP(s) [in same order as haplotype in Col 7]

Here we have explicitly specified the TC and TTA haplotypes. For example, in the first case, SNPs snp1 and snp2 may have all four common haplotypes seen in the sample, TT, CT and CC as well as TC; this command would select only the TC haplotype to be imputed, or as the focus of haplotype analysis. The imputed SNP, rs1001 therefore has the following alleles:

```
TC/TC 1/1
TC/* 1/2
*/* 2/2
```

and will be positioned on chromosome 5, and base-positon 201. Haplotypes other than TC will be coded 2.

The imputed SNP details (alleles, etc) will only be used if the --hap-impute option has been requested. For --hap-assoc and --hap-tdt options (which consider all possible phases rather than just imputing the most likely) these are not considered (but they are still required in this input file).

#### 2) 'Wildcard' specification

Alternatively, all haplotypes at a given locus above the --maf threshold can be automatically estimated by entering a line in myfile.hlist as follows:

- \* snp1 snp2 snp3
- \* snp1 snp2

i.e. where the first character is an asterisk \*, which would, taking just the first line for example, create all 3-SNP haplotypes for the SNPs labelled in the MAP file as snp1, snp2 and snp3, above the minor allele frequency threshold. If the haplotypes were, for example, AAC, AGG and TGG, then the following names would be automatically assigned:

```
H1_AAC_
H1_AGG_
H1_TGG_
```

Haplotypes based on subsequent lines in the file would be labelled H2\_\*\_, H3\_\*\_, etc. In this case, all two-SNP haplotypes for snp1 and snp2 would start H2\_. The chromosome and position flags for the new haplotypes are set to equal the first SNP of the set.

#### 3) 'Named wildcard' specification

Finally, this format is identical to the previous wildcard specification, except a name can be given to the haplotype. This uses \*\* instead of \* to start a row; the second entry is then interpreted as the name of the haplotype locus rather than the first SNP. For example:

```
** BLOCK1 snp1 snp2 snp3
```

\*\* BLOCK2 snp6 snp7

The only difference is that BLOCK1 and BLOCK2 names will be used in the output instead of H1 and H2 being assigned automatically.

#### 4) Sliding window specification

Finally, instead of specifying a haplotype file with the --hap option, you can use the --hap-window option to specify all haplotypes in sliding windows of a fixed number of SNPs (shifting 1 SNP at a time).

```
plink --bfile mydata --hap-window 3 --hap-assoc
```

to form all 3-SNP haplotypes across the entire dataset (respecting chromosome boundaries, however). In this case the windows will be automatically named WIN1, WIN2, etc.

## 11.2 Precomputed lists of multimarker tests

Below are links to some PLINK-formatted lists of multimarker tests selected for Affymetrix 500K and Illumina whole genome products, based on consideration of the CEU Phase 2 HapMap (at r-squared=0.8 threshold). One should download the appropriate file and run with the --hap option (after ensuring that any strand issues have been resolved). These files were generated by Itsik Pe'er and others, as described in this manuscript:

```
Pe'er I, de Bakker PI, Maller J, Yelensky R, Altshuler D & Daly MJ (2006) Evaluating and improving power in whole-genome association studies using fixed marker sets. Nat Genet, 38(6): 605-6.
```

- Affymetrix.GeneChip.500k.both.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Affymetrix.GeneChip.500k.both.CEU.0.8.tests.zip
- Illumina.HumanHap.300k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.300k.CEU.0.8.tests.zip
- Illumina.HumanHap.550k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.550k.CEU.0.8.tests.zip
- Illumina.HumanHap.650k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.650k.CEU.0.8.tests.zip

These tables list all tags for every common HapMap SNP, at the given r-squared threshold. The same haplotype may therefore appear multiple times (i.e. if it tags more than 1 SNP). The haplotypes are specified in terms of the + (positive) strand relative to the HapMap. You might need to reformat your data prior to using these files (using the --flip command, for instance) before you can use them.

Note These tables obviously assume that all tags on present in the final, post-quality-control dataset: i.e. if certain SNPs have been removed, it will be better to reselect the predictors – that is, these lists should really only be used as a first pass, for convenience.

### 11.3 Estimating haplotype frequencies

To obtain the haplotype frequencies for all haplotypes in each window, use the option:

# 11.4 Testing for haplotype-based case/control and quantitative trait association

In a population-based sample of unrelated individuals, case/control and quantitative traits can be analysed for haplotype associations, using the option, for example,

```
plink --file mydata --hap myfile.hlist --hap-assoc
```

which will generate haplotype-specific tests (1df) for both disease and quantitative traits; for disease traits only, an omnibus association statistic will also be computed. This option generates the file

```
plink.assoc.hap
```

which contains the following fields:

LOCUS Haplotype locus / window name HAPLOTYPE Haplotype identifer / "OMNIBUS" F\_A Frequency in cases  $F_{-}U$ Frequency in controls CHISQ Test for association DF Degrees of freedom Ρ Asymptotic p-value SNPS SNPs forming the haplotype

or

plink.qassoc.hap

which contains the following fields:

LOCUS Haplotype locus / window name HAPLOTYPE Haplotype identifer NANAL Number of individuals in analysis **BETA** Regression coefficient RSQ Proportion variance explained STAT Test statistic (T) Ρ Asymptotic p-value SNPs forming the haplotype SNPS

In all cases, the tests are based on the expected number of haplotypes each individual has (which might be fractional). The case/control omnibus test is a H-1 degree of freedom test, if there are H haplotypes.

## 11.5 Haplotype-based TDT association test

If the case/control data are being analysed, use the option

```
plink --file mydata --hap myfile.hlist --hap-tdt
```

to test for TDT haplotype-specific association. This option generates the file

plink.tdt.hap

which contains the following fields:

LOCUS Haplotype locus / window name

HAPLOTYPE Haplotype identifer / "OMNIBUS"

T Number of transmitted haplotypes

U Number of untransmitted haplotypes

CHISQ Test for association
P Asymptotic p-value

## 11.6 Imputing multimarker haplotypes

If the --hap-impute option is also given, this will create two new files:

```
plink --file mydata --hap myfile.hlist --hap-impute
```

will generate the file:

```
plink.impute.ped
plink.impute.map
```

based on the most likely E-M phase reconstructed haplotypes. One could then simply treat the most likely haplotype assignments as SNPs and use all the standard analytic options of PLINK, e.g. --assoc.

Warning This represents a quick and dirty approach to haplotype testing. Depending on how accurately the haplotypes have been imputed (i.e. the range of maximum posterior probabilities per individual) some bias will be introduced into subsequent tests based on these 'SNPs'. Typically, as long as cases and controls are phased together, as they are here, this bias is likely to be quite small and so should not substantively impact results (unpublished simulation results, SMP). Furthermore, exact methods can be used to refine the association for the putative hits discovered by this approach.

**NOTE** Future versions will allow for a binary PED file to be created from the --hap-impute command. You do **not** need to specify --recode when using --hap-impute.

### 11.7 Tabulating individuals' haplotype phases

To obtain a summary of all possible haplotype phases and the corresponding posterior probabilities (i.e. given genotype data), use the command:

```
plink --file mydata --hap myfile.hlist --hap-phase
which will generate the file
    plink.phase-*
```

where \* is the name of the 'window' (i.e. the row of the haplotype list file). That is, if the haplotype list contains multiple rows, then multiple phase files will be generated. These files contain the fields, where each row is one possible haplotype phase for one individual:

```
FID Family ID

IID Individual ID

PH Phase number for that individual (0-based)

HAP1 First haplotype, H1

HAP2 Second haplotype, H2

POSTPROB P(H1,H2 | G )

BEST 1 if most likely phase for that individual
```

# Chapter 12

# Conditional haplotype-based association testing

This page describes PLINK functions that are aimed at dissecting a haplotypic association. These functions largely include and extend the functionality offered in the older WHAP http://pngu.mgh.harvard.edu/~purcell/whap/ software package, which is no longer supported.

For reference, the main ways of specifying conditional haplotype tests, that modify the behaviour of main --chap command, are given here; they are also described in more detail below. Each row here is mutually exclusive, e.g. you would not want to, or be able to, specify --control and --alt-snp at the same time:

- Test whether SNPs have independent haplotyic effects (--independent-effect SNP, SNP, SNP)
- Test whether a set of SNPs explain an omnibus association (--control SNP, SNP,...)
- Test whether a specific set of haplotypes explain an omnibus association (--control HAPLOTYPE, HAPLOTYPE,...)
- Test specific haplotypes for association (--specific-haplotype HAPLOTYPE)
- Specify alternative and null haplotypic models in terms of sets of SNPs (--alt-snp SNP,SNP-SNP,... and/or --null-snp SNP,SNP-SNP,...)
- Specify alternative and null haplotypic models in terms of sets of haplotypes (--alt-group HAPLOTYPE, HAPLOTYPE, ... and/or --null-group HAPLOTYPE, HAPLOTYPE,...)
- Test a one or more simple SNP effects, potentially controlling for haplotype effects (--test-snp SNP,SNP-SNP,...)

It is also possible to include one or more continuous or binary covariates, which can include other SNPs outside of the phased region.

This page contains the following sections:

- Basic usage
- Specifying the type of test
- General specification of haplotype groupings
- Including covariates and other SNPs

The value of using --chap over --hap-assoc is that covariates can be included, and more complex conditional tests can be specified. The value of using --hap-assoc over --chap is that it is designed to iterate over very many SNPs in a single go, whereas the --chap test is more designed to focus on one specific set of SNPs.

#### Basic usage for conditional haplotype-based testing 12.1

The --chap command is used in conjunction with the --hap-snps command to specify a set of SNPs to phase, form haplotypes and test for association (in samples of unreated individuals only):

```
plink --bfile mydata --hap-snps rs1001-rs1005 --chap
```

which generates a file

plink.chap

The --hap-snps command can take a comma-delimited list of SNPs, including ranges, e.g. if the MAP file specifies the following SNPs and physical positions:

```
1 rs1001 0 101200
```

1 rs1002 0 102030

1 rs1003 0 107394

1 rs1004 0 107499

1 rs1005 0 113990

HAPLO

then the command

```
--hap-snps rs1001-rs1003,rs1005
```

includes all SNPs except rs1004, for example. The hyphen/minus symbol specifies all SNPs within a range (based on sorted physical position).

**NOTE** No spaces are allowed in this kind of comma-delimited list. Also, note that currently this will not work if SNP names have hypen characters in them. In this case, to use a different delimter for any ranges specified on the command line, add the "-d" flag (which can be any non-whitespace character except a comma (although be cautious if using characters with special meanings on command lines)

```
--d + --hap-snps SNP-A10001+SNP-A10020
```

to obtain a range between SNP-A10001 and SNP-A10020.

The default test is an *omnibus* haplotype test: that is, if there are H haplotypes, then --chap performs an H-1 df test comparing the alternate (each haplotype having a unque effect) versus the null (no haplotypes having any different effect). In each case, one haplotype is arbitrarily chosen to be the reference haplotype. The coefficients must be interpreted with respect to that haplotype, but otherwise the coding makes no difference.

For binary disease traits, the test is based on a likelihood ratio test. For continuous traits, the test is based on an F-test comparing the alternate and null models. For continuous traits, the --chap command also displays the proportion of variance in the outcome explained by the regression model (R-squared) as well as an adjusted R-squared (that takes model complexity into account).

For example, here is a plink.chap output file representing a basic omnibus test:

```
+++ PLINK conditional haplotype test results +++
5 SNPs, and 6 common haplotypes (MHF >= 0.01) from 32 possible
 CHR
                BP
                            SNP
                                   A1
                                        A2
           101200
   1
                         rs1001
                                    C
                                                  0.45
                                         Α
   1
           102030
                         rs1002
                                         C
                                                0.2362
                                         C
   1
            107394
                         rs1003
                                    Α
                                                0.4325
                                    Т
                                         G
           107499
                         rs1004
                                                0.2362
   1
           113990
                         rs1005
                                    Α
                                         C
                                                0.4487
Haplogrouping: each set allowed a unique effect
Alternate model
    AAATA
             AACTA
                       CCCGA
                                 ACAGC
                                          CCCGC
                                                    ACCGC
Null model
    AAATA, AACTA, CCCGA, ACAGC, CCCGC, ACCGC
                    FREQ
                                 OR(A)
                                               OR(N)
```

AAATA	0.169	(-ref-)	(-ref-)
AACTA	0.06728	2.619	
CCCGA	0.2125	0.8942	
ACAGC	0.2635	0.6839	
CCCGC	0.2375	1.025	
ACCGC	0.05022	1.038	

Model comparison test statistics:

Alternate Null -2LL: 535.4 554.5

Likelihood ratio test: chi-square = 19.11

df = 5p = 0.001836

There are several points to note:

- At the top of the output, PLINK lists the SNPs (SNP) involved in the test, their chromosomal (CHR) and base-pair (BP) positions, their alleles (A1 and A2) and the minor allele frequency (F).
- It is reported that there are 5 common haplotypes: this filter (default value of 0.01) can be changed by adding, for example, the --mhf 0.05 command (minimum haplotype frequency).
- The next section presents the *haplogrouping* under the null and alternate models. If two haplotypes are in the same <code>set</code>, it means they are treated as identical in terms of their effect on phenotype (i.e. a single regression coefficient is used for that group). For the basic omnibus test the haplogrouping will always take this simple form: under the alternate all haplotypes in their own set, whilst under the null all haplotypes are in one set. This output is more useful in interpreting some of the other conditional haplotype tests that are introduced below.
- The next section contains the estimated regression coefficients for each haplotype under the alternate and null models, as well as the frequency (FREQ) of each haplotype. For continuous traits, the coefficients are labelled BETA; for disease traits they are labelled OR and are in fact transformed to be odds ratios, i.e. exp(beta). The (-ref-) indicates which haplotype has been selected to be the baseline, reference category. If a haplotype has instead a pipe (vertical bar) | symbol, it implies that this haplotype is grouped with the one above it (and so it will not have a regression coefficient of its own). In the case of this simple null model as shown here, this implies that all haplotypes are equated with AAATA, the reference haplotype (i.e. there is no effect of any haplotype).
- When the null model is not so straightforward (as in the examples below), the rows are separated into the null-model haplogroups for clarity. In this case, certain *sub-null* model comparisons are also presented, to the right of the table of coefficients: these are shown and described below.
- The final section presents the overall model statistics: for a linear trait these are the R-squared (sometimes called the coefficient of determination) and adjusted R-squared, as well as the F-test. For disease traits, as in this case, only the sample log-likelihood under each model (-2LL) and the likelihood ratio test are presented. In both cases, the degrees of freedom is the number of parameters in the alternate model minus the number in the null model.

The interpretation of this particular analysis would be that overall variation at this locus appears to influence the trait, with p = 0.001836. Using the commands introduced below, we can perform various conditional tests to explore this *omnibus* result.

### 12.2 Specifying the type of test

If no other commands are given, the --chap test will perform an omnibus haplotypic association test. Various other options can be used to refine the type of test. In this section we introduce three commonly used tests; in the section below we introduce a more general way in which any two (nested) models can be compared.

### 12.2.1 Testing a specific haplotype

It is possible to specify a particular haplotype to be tested against all others: for example, CCCGA

```
./plink --file mydata --hap-snps rs10001-rs10005 --chap --specific-haplotype CCCGA
```

This creates the following two haplogroupings:

```
Alternate model
AAATA, AACTA, ACAGC, CCCGC, ACCGC
Null model
AAATA, AACTA, CCCGA, ACAGC, CCCGC, ACCGC
```

which hopefully begins to indicate how these groupings should be interpreted in relation to the tests they imply.

The main body of the output is:

OR(N)	OR(A)	FREQ	HAPLO
(-ref-)	(-ref-)	0.169	AAATA
I		0.06728	AACTA
I	1	0.2635	ACAGC
1		0.2375	CCCGC
1		0.05022	ACCGC
1	0.9153	0.2125	CCCGA

which shows that now under the alternate all haplotypes are grouped together except for CCCGA; versus all other haplotypes, this has an estimated odds ratio of 0.9153.

NOTE Of course, the estimated odds ratio for CCCGA was different in the first example given above (when it was 0.8942) because the reference category was different (it was then only AAATA as opposed to all other SNPs). In other words, remember that the odds ratios are only interpretable in relation to some specific baseline, reference category.

Finally, we see the model compariston test is non-significant

```
Likelihood ratio test: chi-square = 0.2653 df = 1 p = 0.6065
```

The option --each-vs-others will add an extra column to the output, if there is more than one haplotype-grouping under the alternate model, which provides p-values for haplotype-specific tests of that haplotype (or haplotype group) versus all others. For example,

```
./plink --file mydata --hap-snps rs10001-rs10005 --chap --each-vs-others
```

which produces output with the new SPEC(A) field

OR(N)	SPEC(A)	OR(A)	FREQ	HAPLO
(-ref-)	0.537	(-ref-)	0.169	AAATA
1	0.0001791	2.619	0.06728	AACTA
1	0.6065	0.8942	0.2125	CCCGA
1	0.003466	0.6839	0.2635	ACAGC

ACCGC	0.05022	1.038	0.787	1
CCCGC	0.2375	1.025	0.5132	1

which contains p-values for all haplotype-specific tests (i.e. as above, the haplotype CCCGA has the p-value of 0.6065 as above, i.e. that haplotype versus all others). The benefit of the --specific-haplotype command versus --each-vs-others is that it also produces the odds ratio for that haplotype.

These haplotype specific tests are of course similar to the basic test given by the --hap-assoc command e.g.

```
./plink --file mydata --hap-snps rs10001-rs10005 --hap-assoc
which generates the output file
     plink.assoc.hap
which contains the line
   LOCUS
         HAPLOTYPE
                        F_A
                               F_{-}U
                                     CHISQ
                                             DF
                                                       Ρ
                                                         SNPS
    WIN1
              CCCGA
                      0.205
                              0.22
                                     0.2689
                                               1 0.6041
                                                          rs1001|rs1002|rs1003|rs1004|rs1005
```

This command frames the test in a slightly different way and presents different statistics (i.e. it does not use logistic regression, case and control frequencies are presented instead of odds ratios, etc) but the p-value is, as expected, very similar (p=0.6041 from --hap-assoc versus p=0.6065 from the --chap test). Note that they are not expected to be numerically identical however.

### 12.2.2 Testing whether SNPs have independent effects

It is possible to ask whether one or more SNPs have an effect that is independent of the other SNPs in the model, framing the question in terms of haplotypes. This conditional test essentially stratifies by the haplotyic background: for the SNP(s) under scruntiny, we only compare the alleles/haplotypes that have a similar haplotypic background.

Before proceeding to the conditional haplotype tests, let's first consider the simple, single SNP effects for the example dataset:

```
./plink --file mydata --assoc
```

which generates the file plink.assoc which is as follows:

CHR	SNP	BP	A1	$F_A$	$F_{-}U$	A2	CHISQ	P	OR
1	rs1001	101200	С	0.4525	0.4475	Α	0.0202	0.887	1.02
1	rs1002	102030	Α	0.2775	0.195	C	7.544	0.00602	1.586
1	rs1003	107394	Α	0.395	0.47	C	4.584	0.03228	0.7362
1	rs1004	107499	T	0.2775	0.195	G	7.544	0.00602	1.586
1	rs1005	113990	Α	0.4825	0.415	C	3.644	0.05495	1.314

Here we see that SNPs rs1002 and rs1004 have the strongest associations, although rs1003 and rs1005 show marginal trends.

Next, to obtain a quick view of the LD in this small region, we can generate the matrix of r-squared (LD) values (i.e. note: this is using r-squared as a measure of LD, which is distinct from the coefficient of determination which descibes the fitted regression models).

```
./plink --file mydata --r2 --ld-window-r2 0
```

This command, by default, only outputs values for SNPs that have an r-squared greater than 0.2, are within 1 Mb and 10 SNPs of each other; these can be changed with the options --ld-window-r2, ld-window-kb and --ld-window respectively; in this case, we requested all SNPs to be reported with --ld-window-r2. The file

```
plink.ld
```

contains the fields

CHR_A	$\mathtt{SNP}_\mathtt{A}$	CHR_B	$SNP_B$	R2
1	rs1001	1	rs1002	0.260769
1	rs1001	1	rs1003	0.628703
1	rs1001	1	rs1004	0.260769
1	rs1001	1	rs1005	0.000357147
1	rs1002	1	rs1003	0.0964906
1	rs1002	1	rs1004	1
1	rs1002	1	rs1005	0.398912
1	rs1003	1	rs1004	0.0964906
1	rs1003	1	rs1005	0.00919232
1	rs1004	1	rs1005	0.398912

Here we see that rs1002 and rs1004 are in complete LD, but that there is also moderate (r-squared above 0.2) LD between many other pairs of SNPs.

Moving then to the conditional tests: using the dataset above, to test for an independent effect of rs1003, for example (independent of the haplotypic effects formed by the remaining SNPs), one would issue the command:

```
./plink --file mydata --hap-snps rs1001-rs1005 --chap --independent-effect rs1003
```

The haplogroupings implied by this command are

Alternate model

AAATA AACTA CCCGA ACAGC CCCGC ACCGC
Null model
AAATA, AACTA CCCGA ACAGC, ACCGC CCCGC

The test SNP, rs1003, is the middle SNP in the 5-SNP haplotype (an A/C SNP). In comparison to the alternate model, we now see that the null is formed by grouping two pairs of haplotypes; each pair is identical except for rs1003: i.e.

AAATA, AACTA

and

ACAGC, ACCGC

In each case here, the comparison between alternate and null models is to equate the effects of these haplotypes (i.e. implicitly providing a test for whether rs1003 has any effect). A haplotype such as CCCGA is effectively left out of the analysis: although it contains a C allele for rs1003, we never see the corresponding CCAGA haplotype to perform a stratified analysis.

The main output for this test is shown below:

SUBNULL P	OR(N)	OR(A)	FREQ	HAPLO
0.008016	(-ref-)	(-ref-)	0.169	AAATA
		2.619	0.06728	AACTA
n/a	0.6907	0.8942	0.2125	CCCGA
0.2643	0.5628	0.6839	0.2635	ACAGC
	1	1.038	0.05022	ACCGC
n/a	0.7897	1.025	0.2375	CCCGC

Model comparison test statistics:

Alternate Null -2LL: 535.4 544.4

Likelihood ratio test: chi-square = 8.982

df = 2

$$p = 0.01121$$

There are two new features to note: first, the null model is no longer a simple unitary group; the rows are separated out into the groups defined by the null model. That is, *null* does **not** mean *no effect of any haplotype*; rather, it is used in the statistical sense of the default, more simple model compared to the alternate: the model which we want to try to *nullify*.

Under the null, haplotypes AAATA and AACTA have a single parameter (both are the reference category); haplotypes ACAGC and ACCGC have an estimated odds ratio of 0.5628 (versus the reference group).

The second new addition is of the *sub-null* test p-values in the right-most column. These will only appear when the null model contains more than one group for which there was more than one group in the alternate model (i.e. groups in which haplotype effects have been equated within group). Whereas the likelihood ratio test at the bottom is a joint 2df test (for whether the two sets of haplotypes can be equated; equivalently, for whether rs1003 has an independent effect), the sub-model p-values represent a test of just that part of the model, i.e. a 1 df likelihood ratio test for whether AAATA and AACTA do indeed have similar odds ratios has the p-value of 0.008016.

One way of interpreting these results would be that rs1003 has an effect on the AA-TA haplotype background, but not the AC-GC background. However, drawing such a conclusion in this simple manner is not advised – p-values should not be interpreted in this direct manner, and also the power of the test will vary by the frequency of the haplotype background. (A feature will be added that enables one to ask specifically whether or not the effect of rs1003 varies between these two haplotype backgrounds: this involves the specification of linear constraints between parameters.)

Note that it is not always possible to perform a test of independent effects: for example, consider rs1002: given the set of common haplotypes under study, we see it is perfectly correlated with rs1004 (i.e. we only ever see the AT and CG haplotypes for these two SNPs. We therefore never see both alleles of rs1002 on the same haplotypic background. As such, the null model is the same as the alternate: PLINK therefore reports

```
Likelihood ratio test: ( not a valid comparison: identical models, df = 0 )
```

It is also possible to see whether more than one SNP has an independent effect: this is still a haplotypic test (of haplotypes formed by the two or more SNPs), but the test is stratified by the haplotypic background formed by the remaining SNPs. For example:

./plink --file mydata --hap-snps rs1001-rs1005 --chap --independent-effect rs1003,rs1004 leads to the haplogrouping

Alternate m	odel				
AAATA	AACTA	CCCGA	ACAGC	CCCGC	ACCGC
Null model					
AA $\mathbf{A}\mathbf{T}$ A,	$\mathtt{AACTA}$	CCCGA	ACAGC,	ACCGC	CCCGC

	1	1.038	0.05022 0.2375	ACCGC CCCGC
n/a 0.2643	0.6907 0.5628	0.8942 0.6839	0.2125 0.2635	CCCGA ACAGC
/-	0 0007	2.619	0.06728	AACTA
0.008016	(-ref-)	(-ref-)	0.169	AAATA
SUBNULL P	OR(N)	OR(A)	FREQ	HAPLO

Model comparison test statistics:

and the main test statistics

Alternate Null -2LL: 535.4 544.4

Likelihood ratio test: chi-square = 8.982 df = 2

$$p = 0.01121$$

In this particular case, this test of independent effects of rs1003 and rs1004 happens to give exactly the same results as the test of rs1003 by itself, which will be made clear from examining the haplogroupings. Note that, in both cases, the test is a two degree of freedom test.

### 12.2.3 Omnibus test controlling for X

To perform an omnibus test but controlling for a particular haplotype of set of haplotypes, you can use the --control command. The haplotypes can either be directly specified, or implied through the list of SNPs specified. This test is a complement to the --independent-effect test.

Typically, one would use this test in the case of a significant omnibus assocation result. For example, we could ask whether we still see the association even if we control for haplotypes of SNPs rs1002 and rs1004 (the two most highly associated SNPs, that are in complete LD with each other):

which gives implied haplogroupings:

Alternate model

AAATA AACTA CCCGA ACAGC CCCGC ACCGC

Null model

AAATA, AACTA CCCGA, ACAGC, CCCGC, ACCGC

In this case, rather than make the null model a single set, the --control command separates the haplotypes out into distinct groups based on the sub-haplotypes at SNPs rs1002 and rs1004, i.e.

The regression coefficient table is:

HAPLO	FREQ	OR(A)	OR(N)	SUBNULL P
AAATA	0.169	(-ref-)	(-ref-)	0.008016
AACTA	0.06728	2.619	1	
CCCGA	0.2125	0.8942	0.6603	0.2087
ACAGC	0.2635	0.6839	1	
CCCGC	0.2375	1.025	1	
ACCGC	0.05022	1.038	1	

and model comparison statistics are:

Alternate Null 
$$-2LL$$
:  $535.4$   $547.7$  Likelihood ratio test: chi-square =  $12.32$  df =  $4$  p =  $0.01515$ 

This is a 4 df test because 4 haplotypes are grouped with another haplotype (i.e. the 4 | symbols in the output).

One would conclude from this analysis that there is still a significant effect at this locus even controlling from the haplotypic effects of rs1002 and rs1004. In otherwords, the command

```
--control rs1002,rs1004
```

is identical to

```
--indepedent-effect rs1001,rs1003,rs1005
```

in this instance. Unlike the --independent-effect, the --control command does allow for hapltoype(s) to be specified, instead of SNPs: for example, we might ask whether the omnibus test is significant controlling for ACAGC:

```
./plink --file mydata --hap-snps rs1001-rs1005 --chap --control ACAGC
```

which gives the following haplogrouping

Alternate model

AAATA AACTA CCCGA ACAGC CCCGC ACCGC
Null model

AAATA, AACTA, CCCGA, CCCGC, ACCGC ACAGC

i.e., effectively leaving ACAGC out of the test, and this table of coefficients

OR(N)	OR(A)	FREQ	HAPLO
(-ref-)	(-ref-)	0.169	AAATA
1	2.619	0.06728	AACTA
1	0.8942	0.2125	CCCGA
1	1.025	0.2375	CCCGC
1	1.038	0.05022	ACCGC
0.624	0.6839	0.2635	ACAGC

Model comparison test statistics:

Alternate Null -2LL: 535.4 546

Likelihood ratio test: chi-square = 10.56

df = 4p = 0.03194

In otherwords, there is still a marginal omnibus assocation (p=0.032) after controlling for ACAGC. Repeating this test for each haplotype:

HAPLOTYPE	(control)	P-VALUE	(omnibus	association)
AAATA		0.000889	5	
AACTA		0.2803		
CCCGA		0.000844	:1	
CCCGC		0.000908	4	
ACCGC		0.000773	8	
ACAGC		0.03194		

which would suggest that there is no significant signal after controlling for AACTA, at the p=0.05 level at least. This is consistent with the true model: these data are in fact simulated, and AACTA was in fact the disease haplotype.

Finally, it is possible to specify multiple, comma-delimited haplotypes for the --control command.

# 12.3 General specification of haplotype groupings

Rather than use any of the above *convenience* functions for specifying tests, one can directly specify the haplogrouping, in one of two ways: by manually specifying the haplotypes, or the SNPs, to include under both alternate and null models.

### 12.3.1 Manually specifying haplotypes

With the --alt-group and --null-group commands, it is possible to directly specify the haplogrouping. These commands take a commandelimited list of *sets*, where the equals symbol is used to specify equality of haplotypes. For example, the command

--independent-effect rs1003

which gives rise to the following haplogroups

Alternate model

AAATA AACTA CCCGA ACAGC CCCGC ACCGC

Null model

AAATA, AACTA CCCGA ACAGC, ACCGC CCCGC

which could instead have been directly specified

- --alt-group AAATA, AACTA, CCCGA, ACAGC, CCCGC, ACCGC
- --null-group AAATA=AACTA, CCCGA, ACAGC=ACCGC, CCCGC

Note how the = symbol is used to define sets. When using these commands, the default for the alternate is as specified above, so this command could have been excluded. Also, it is not necessary to specify all haplotypes: if a haplotype is not specified, it will revert to its default grouping (i.e. depending on whether this is for the alternate or null). In other words, the same effect could have been achieved just with the single command

--null-group AAATA=AACTA, ACAGC=ACCGC

Finally, there are two wild-cards, one of which can be used in these two commands:

- \* Group all haplotypes not otherwise explicitly mentioned
- % Separate all haplotypes not otherwise explicitly mentioned

In other words, implicitly there is always a base-line of

```
--alt-group %
--null-group *
```

To just equate two haplotypes, for instance, but keeping everything else the same, one might use

```
--null-group AAATA=AACTA,%
```

i.e. which means "under the null, allow each haplotype to have a unique effect (%), with the exception of AACTA and AACTA, which should be grouped with each other".

### 12.3.2 Manually specifying SNPs

With the --alt-snp and --null-snp commands, it is possible to specify which SNPs should be used to form haplotypes. By default, all SNPs are included in the alternate, no SNPs are included in the null: this leads to the default haplogrouping of the omnibus test.

To illustrate this command, by reference to the --independent-effect specification, for example: the command

```
\begin{tabular}{ll} --independent-effect $rs1003$\\ is equivalent to \end{tabular}
```

```
--alt-snp rs1001-rs1005 --null-snp rs1003
```

### 12.4 Covariates and additional SNPs

Covariates can be included with the --covar option, the same as for --linear and --logistic models. By default, all covariates in that file with be used. Covariates always feature under both the alternate and null models.

```
./plink --file mydata --hap-snps rs1001-rs1005 --chap --covar myfile.cov
```

which generates an additional set of entries in the plink.chap output file, representing the coefficients (no other statistical tests are performed for the covariates, i.e. no p-values, etc):

COVAR	OR(A)	OR(N)

COV1 0.7834 0.8499

In a similar manner, additional SNPs can be included, which can be SNPs other than those included in the --hap-snps command. These SNPs are not considered in any way during the phasing process: the alleles are simply entered in an allelic dosage manner. The command --condition and a list of SNPs, or --condition-list followed by a filename with a list of SNP names, includes these.

which adds the following lines in the output file

OR(N)	OR(A)	SNPS
2.899	1.038	rs1006

Unlike for standard covariates, it is also possible to request that a SNP effect be dropped under the null model, which allows, for example, for a test of a SNP controlling for a set of haplotypes at a different locus: here, one would want to include all haplotype effects under the null, and use the --test-snp command to drop one or more of the conditioning SNPs:

which would instead show

OR(N)	OR(A)	SNPS
(dropped)	1.038	rs1006

and an extra degree of freedom would be added to the model comparison test. As the --null-group % command was used to effectively control for all haplotypic effects whilst testing this particular SNP, rs1006, the test will be a 1 df test,

Likelihood ratio test: chi-square = 
$$0.0007377$$
 df =  $1$  p =  $0.9783$ 

It is also possible to specify more than one conditioning SNP (and to drop none, some or all of these under the null): for example,

```
./plink --file mydata --hap-snps rs1001-rs1005 --chap --null-group % --condition rs1006,rs1007 --test-snp rs1006
```

# 12.5 General setting of linear constraints

to be completed

# Chapter 13

# Proxy association

This page describes a convenience function designed to provide a quick representation of a single SNP association, in terms of the surrounding haplotypic background. Specifically, given a particular (reference) SNP this approach involves a) finding flanking markers and haplotypes (proxies) that are in strong linkage disequilibrium with the reference SNP and, b) testing these proxies for association with disease, within a haplotype-based framework.

There are three main applications of this utility, which are described in more detail and with examples in the main text below:

- technical validation of single SNP results ( by looking for flanking haplotypes involving different markers that also show the same result )
- refining a single SNP association signal ( is there a stronger association with a local haplotype? )
- more robust single SNP tests (by framing single SNP tests within a haplotypic framework, some degree of control against non-random genotyping failure can be achieved)

The proxy approach also forms the basis of the imputation methods in PLINK, described separately. The methods are identical in fact, the only difference in imputation mode is the presence of a reference set of individuals that is handled specially.

The proxy methods use the same basic EM algorithm used by the other haplotyping methods in PLINK. The only difference is that the proxy methods put a wrapper around the basic haplotyping procedure that a) provides some methods to automatically select proxies to phase given a designated reference SNP, and b) frames the subsequent tests and summaries in terms of groups of haplotypes that track the reference SNP.

# 13.1 Proxy association: basic usage

The basic proxy association method for a particular SNP is invoked with the --proxy-assoc option:

```
plink --file mydata --proxy-assoc rs6703905
```

which generates a file

plink.proxy.report

This file contains three main sections, describing the local flanking SNPs, haplotypes and "proxies" for the reference SNP, and will be described below in turn. The full output file is shown here:

*** Pr	oxy hapl	Lotype	association	report	for rs:	13232128 ***		
	SNP	MAF	GENO	KB	RSO	Q OR	CHISQ	P
rs138	9273	0.286	0.00173	-99.2	0.0932	0.916	2.61	0.106
rs1023	6783	0.253	0.0236	-66.9	0.214	0.875	5.7	0.017

rs17556689	0.328	0.00259	-66.7	0.282	1.1	3.09	0.079
rs17135491	0.153	0.00317	-2.59	0.153	0.934	0.955	0.328
rs13232128	0.494	0.0179	0	*	0.828	14.9	0.000112
rs1826529	0.487	0.00461	9.72	0.674	0.883	6.58	0.0103
*.	FREQ	OR	CHIS	Ç	P		
GTTGAG	0.0171	1.02	0.022	1 (	0.882		
AGTGAG	0.0166	0.876	0.712	2 (	0.399		
GGTGAG	0.103	0.91	1.56	3 (	0.212		
GGCAAG	0.0226	0.97	0.047	5 (	0.827		
ATTAAG	0.111	0.853	4.96	3 (	0.026		
GTTAAG	0.0615	0.877	2.09	9 (	0.149		
AGTAAG	0.0557	0.881	1.73	3 (	0.188		
GGTAAG	0.0513	0.949	0.306	3	0.58		
GGCAGG	0.0365	1.39	7.56	0.0	00596		
ATTAAT	0.0233	0.825	1.78	3 (	0.183		
GGCAGT	0.249	1.05	0.893	3 (	0.345		
ATTAGT	0.0201	1.31	3.26	3 0	.0711		
AGTAGT	0.049	1.08	0.74	1 (	0.389		
GGTAGT	0.13	1.16	5.1	7 (	0.023		

Haplotype frequency estimation based on 6938 of 6938 founder chromosomes Omnibus haplotype test statistic: 23.3, df = 13, p = 0.0377

Of	125	subhaplotypes	considered,	8	met	proxy	criteria
-		~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~		_		P = V	

HAP	FREQ	RSQ	OR	CHISQ	P
T. G	0.422	0.72	0.843	12.3	0.000449
.G T	0.453	0.705	1.15	8.62	0.00333
.G.A T	0.445	0.693	1.14	7.22	0.0072
GG T	0.399	0.561	1.14	7.15	0.0075
G	0.487	0.674	0.883	6.58	0.0103
A T	0.505	0.661	1.12	5.56	0.0184
${\tt G} \ {\tt T}$	0.415	0.542	1.11	4.99	0.0255
GA T	0.408	0.535	1.1	4.19	0.0408

The first section lists the reference SNP (rs13232128) and 5 flanking SNPs that have been automatically selected as proxies. For each SNP, the minor allele frequency (MAF), genotyping failure rate (GENO) and distance to the reference SNP (KB) is given. A measure of single SNP association is also given for each SNP: odds ratio (OR), chi-squared statistic (CHISQ) and asymptotic p-value (P).

Importantly, however, these single SNP tests are not quite the same as from the basic --assoc command, as they are formed within the haplotypic context of the flanking SNPs. That is, for example, a single SNP test of the 5th SNP is formed by grouping the haplotypes as shown below, and testing for a difference in the frequency of the first group (containing A at the 5th position) versus the second group (all containing G).

 $\mathsf{GTTG} ext{-}\mathbf{A} ext{-}\mathsf{G}$ 

AGTG-A-G

 $\mathtt{GGTG}\text{-}\mathbf{A}\text{-}\mathtt{G}$ 

GGCA-A-G

ATTA-A-G

 $\mathsf{GTTA} - \mathbf{A} - \mathsf{G}$ 

AGTA-A-G

 ${\tt GGTA-}{\bf A}{\tt -}{\tt G}$ 

ATTA-A-T

versus

GGCA-G-G

GGCA-G-T

ATTA-G-T AGTA-G-T GGTA-G-T

Because the test is conducted in the context of a haplotypic test, it has some slightly different properties to the standard association test, which can sometimes be used to advantage. In particular, when there is strong LD in the region, the haplotype information will often help to fill in missing genotype data for single SNPs. Therefore, rather than throwing away individuals with missing genotype data, it is possible to try to reconstruct it from the surrounding region: this can lessen the impact of non-random genotyping failure causing spurious associations, as described below.

In this example, note that no other surrounding SNPs appear to show strong association with disease, compared to the reference SNP: when looking at the pattern of LD (RSQ column) we see that there are no SNPs with very high LD (e.g. over 0.8) to the reference SNP, so this is not necessarily surprising. Other *haplotypes* might be, however: this is what the rest of the report considers. The second section lists the haplotypes formed in that region given all flanking proxy SNPs (including the reference SNP) and the frequency and association with disease of each of these haplotypes.

Finally, the third part of the report contains that information on single proxies SNPs or haplotypes of two of more proxies (subhaplotypes) but excluding the reference SNP, that are in LD with the reference SNP; this list is sorted by strength of association with disease and filtered by other criteria, described below. For example, the first line in the above example is:

HAP	FREQ	RSQ	OR	CHISQ	P
T. G	0.422	0.72	0.843	12.3	0.000449

This suggest although no single SNP shows a similar association to the reference SNP in this region, a haplotype does show association results of a similar magnitude and is correlated with the reference SNP (in this case, the TG haplotype formed by the third and last proxy SNPs).

So, in this particular example, this might be taken as additional support for the association: it is of course still possible that the association is just due to chance, or due to population stratification, etc, but this would suggest that it is unlikely to be due to some technical genotyping artefact that was specific to the reference SNP, as we are also seeing the same signal from other SNPs (or, as in this case, a haplotype formed from two other SNPs).

Naturally, if one considers enough proxy haplotypes, some are bound to show stronger association with disease than the reference SNP merely due to chance. One should therefore be careful in how these tests are interpreted, i.e. not to forget the multiple testing that is implicit here.

This kind of analysis represents the typical use case for proxy association: we may have a single SNP association result, but the SNP might be rare or have a higher genotyping failure rate than we would like. Rather than exclude that SNP altogether, one option is to include the SNP in analysis, assess evidence for association, and then also ask whether other SNPs show the same signal. The assumption is that although the true alleles at the proxy SNPs are (hopefully) not independent of the reference SNP (i.e. there is LD) any technical genotyping artefact that influenced the reference SNP is unlikely to also be impacting the proxy SNPs (i.e. the implicit model of genotyping failure is that most SNPs are okay, but a few SNPs might fail: as such, we can use the surrounding genotype data to fill-in failed genotypes, even if these SNPs failed in a very biased way, e.g. if only TT homozygotes tended to fail and only in cases).

### 13.1.1 Heuristic for selection of proxy SNPs

The main parameters for SNP selection are:

- LD thresholds between the index and proxies, and between the proxies themselves
- Maximum number of SNPs and kb range to search for proxies
- Maximum number of proxies to include

There are four main commands to influence the search strategy for proxies:

Proxies are chosen based on LD with the reference SNP as follows. Proxies are examined one at a time in order of strongest to weakest LD with the reference. A proxy must be above a certain minimum r-squared threshold with the reference (criterion A), although if we already have two proxies selected, a different threshold is used (criterion B). In both cases, for a proxy to be added, it must not have an r-squared greater than criterion C with any proxy already selected. For common SNPs, the default values for A, B and C are:

```
Is a proxy?
A) r-sq > 0.00 with reference if < 2 proxies selected
B) r-sq > 0.25 with reference if 2 or more proxies selected
C) r-sq < 0.50 with any other proxy</pre>
```

Setting A lower than B, and to 0 by default, ensures that we always allow a chance of finding a 2-SNP haplotype that might tag the reference SNP, even if no single SNP does.

By default, proxy association selects up to 5 (--proxy-maxsnp) SNPs flanking the reference SNP, from a search of 15 SNPs (--proxy-window) either side of the reference, at most 250 kb away (--proxy-kb).

The defaults vary depending on the frequency of the index SNP: for rarer SNPs (MAF less than 0.1), a slightly larger search space will be used. This threshold can be changed with the command

```
--proxy-b-threshold 0.05
```

In contrast to common SNPs, for which the defaults are:

```
--proxy-r2 0.00 0.25 0.50

--proxy-window 15

--proxy-kb 250

--proxy-maxsnp 5
```

these values for rarer SNPs (as defined by --proxy-b-threshold) and the commands that can be used to change them:

```
--proxy-b-r2 0.00 0.01 0.50

--proxy-b-window 30

--proxy-b-kb 500

--proxy-b-maxsnp 510
```

In other words, the search space is increased for rarer SNPs, to increase the chance that a good haplotypic proxy is found even if there is no other single SNP that well captures the variation at the index site.

In addition, proxies must by default be above 0.01 MAF and below 0.05 genotyping failure rate. To explicitly select only more common proxies with very high genotyping rate (e.g. to verify association at a reference SNP with lower genotyping rate and a very rare allele), then set values for

```
--proxy-maf
and
--proxy-geno
j/tt; appropriately (these mirror the basic --maf and --geno commands).
```

Finally, there are some parameters that determine the behavior of the haplotypic proxy search (the 3rd section of the verbose output). Haplotypes formed by proxies must have a frequency of at least 0.01; these haplotypes must show an r-squared of at least 0.5 with the reference; when considering all possible subhaplotypes, only permutations of up to 3 SNP-haplotypes are considered.

Overall, it is possible to change the behaviour of the basic proxy selection heuristic with the following commands:

• to select a different number of flanking SNPs (--proxy-window)

- to filter proxy SNPs on distance to reference (--proxy-kb)
- to specify the maximum number of proxies (--proxy-maxsnp)
- to filter proxy SNPs on LD with reference (--proxy-r2)
- to not filter proxy SNPs on LD (--proxy-no-r2-filter, i.e. same as --proxy-r2 0 0 1)
- to filter proxy SNPs on MAF (--proxy-maf)
- to filter proxy SNPs on genotyping rate (--proxy-geno)
- to select a specifc set of flanking SNPs (--proxy-flanking)
- to filter haplotypes based on frequency (--proxy-mhf)
- to filter haplotypes based on LD with reference (--proxy-sub-r2)
- to select different levels of subhaplotype search (--proxy-sub-maxsnp)

For example, to select up to 6 SNPs, that are above 0.10 MAF and 0.01 genotyping failure rate, that are within 100 kb and 10 SNPs of the reference SNP and that have an r-squared of at least 0.1 with the reference but no greater than 0.5 with an already-selected proxy SNP; and then to look at all haplotype proxies that are above 0.005 minor haplotype frequency and have an r-squared of at least 0.8 with the reference SNP, use the command (line breaks added for clarity):

```
plink --file mydata --proxy-assoc rs6703905
--proxy-maxsnp 6
--proxy-r2 0.1 0.1 0.5
--proxy-window 10
--proxy-kb 100
--proxy-maf 0.1
--proxy-geno 0.01
--proxy-sub-r2 0.8
--proxy-mhf 0.005
```

As mentioned, rather than use the heuristic above, you can specify a particular set of SNPs with the command

```
plink --file mydata --proxy-assoc rs6703905 --proxy-flanking my.proxy.list
```

where my.proxy.list is a file listing the SNPs you wish to use as proxies for rs6703905, for example.

Warning There will possibly be a very, very large number of possible combinations to consider if you make both --proxy-maxsnp and --proxy-sub-maxsnp too large, meaning that the analysis will take a very long time to run. You should probably keep --proxy-maxsnp less than 10 and --proxy-sub-maxsnp less than 6.

**HINT** To speed up the proxy report, you need only load in the relevant chromosomal region: that is, use the --snp and --window options:

```
plink --bfile mydata --proxy-assoc rs12345 --snp rs12345 --window 300
```

### 13.1.2 Specifying the type of association test

By default, the --proxy-assoc command only applies to population-based samples of unrelated individuals. It is suitable for either disease (case/control) or quantitative trait outcomes: the appropriate test will automatically be selected depending on the phenotype.

The basic command cannot include covariates: however, if the flag --proxy-glm is added, then the routines that correspond to --linear and --logistic are used instead to test the proxy association, meaning that covariates can be included (this is slightly slower than the default analysis), e.g.

```
plink --bfile mydata --proxy-assoc rs12345 --proxy-glm --covar mycov.txt
```

**BETA** There is preliminary support for the TDT in this context with the --proxy-tdt option; this has not vet been fully tested however, and we do not vet suggest you use it generally.

### 13.2 Refining a single SNP association

The proxy association report is primarily designed simply to provide a convenient way to automatically scan for evidence of the same association signal coming from different sets of markers (that are assumed to be independent in terms of technical artefact but not LD). Of course, it is entirely possible that a 'proxy' may show a markedly stronger association than the original reference SNP. In this way, one might think of using the --proxy-assoc method as a way to refine an association signal, or fine-map a region. In a whole genome context, there is clearly nothing special about the particular SNP genotyped that shows association: it may be representing just the tip of an iceberg in association space, and certain haplotypes might have a stronger association. One strategy and way of using haplotype inflormation in a whole genome context, therefore, might be to scan all single SNPs for modest levels of association, and then exhaustively search the haplotype space surrounding those SNPs, but constraining the search to only haplotypes that are in LD with the original SNP (in this way, keeping the multiple testing burden somewhat under control, as although many more tests are added, they will all be quite highly correlated).

As implied in the section above, remember that taking just the best proxy association result (i.e. the top listed in the 3rd section of the report) will capitalize on chance and so these best values will not follow asymptotic null test statistic distributions. These p-values are perhaps best interpreted either against a set genome-wide significance threshold, or corrected for the number of subhaplotypes tested for a given reference SNP.

# 13.3 Automating for multiple references SNPs

To faciliate looking at more than one reference SNP at a time, you can use the command

```
plink --bfile mydata --proxy-assoc all
or
plink --bfile mydata --proxy-assoc all --proxy-list hits.list
```

That is, instead of a SNP name after --proxy-assoc, put the keyword all. PLINK will then treat as the reference, one at a time, either all SNPs in the dataset (first usage) or in the subset listed in the file hits.list (second usage).

By default, only a restricted degree of output is given, and no "subhaplotype" tests are performed when more than one SNP is specified as the reference (i.e. these correspond to the third section in the above example output). To get the full report for every SNP (all listed in a single file) add the option

```
--proxy-verbose
```

In non-verbose mode, the output is as follows, in a file

```
plink.assoc.proxy
with fields
     CHR.
                Chromosome code
     SNP
                Reference SNP
     BP
                Physical position
     Α1
                Name of first allele
     A2
                Name of second allele
     GENO
                Genotyping for the reference SNP
     NPRX
                Number of proxy SNPs used to tag reference SNP
     INFO
                Information metric for each reference SNP
                Reference SNP allele frequency in cases (disease traits)
     F_A
     F_{-}U
                Reference SNP allele frequency in controls (disease traits)
     OR.
                Odds ratio (for disease traits)
     Ρ
                Asymptotic p-value for test of association
     PROXIES
                (Optional, given --proxy-show-proxies) Displays actual proxy SNPs used
```

For, example, here are some lines from such an output file, in this case with the

```
--proxy-show-proxies
```

flag added, which appends the final PROXIES field to the output (lines truncated)

```
F_{-}A
CHR
           SNP
                     BP A1 A2
                                 GENO NPRX
                                            INFO
                                                           F_U
                                                                OR.
                                                                       P PROXIES
                           A 0.00605
 17
      rs731971 29529017
                        Т
                                         3
                                            1.01 0.103 0.104 1.02 0.849 rs4794990|rs11652429|...
    rs4794990 29529302 T
                           C 0.00346
                                         3
                                           1.01 0.102 0.104 1.02 0.838 rs731971|rs11652429|...
 17 rs12938546 29530562 C A 0.00115
                                         3 0.996 0.0322 0.0381 1.19 0.186 rs7359592|rs887071|...
 17 rs11652429 29531710 G C 0.00115
                                         5
                                               1
                                                   0.32
                                                          0.32
                                                                  1 0.984 rs11080256|rs1024613|...
```

The p-values reported here take account of the fact that the SNP has been probabilistically reconstructed. For example, the first line indicates that for rs731971 three proxy SNPs were selected, rs4794990, rs11652429 and rs887071.

The GENO and INFO have more meaning in the context of *imputation*, as described here, which involves running proxy association/imputation with a reference panel, such as the HapMap.

# 13.4 Providing some degree of robustness to non-random genotyping failure

When performing tests in a haplotype-context, the E-M algorithm is used to estimate haplotpe frequencies and each individual's posterior haplotype phase probabilities. The association test is then based on these fractional counts (i.e. allowing for ambiguity in inferred haplotypes). As such, missing genotypes are quite naturally accommodated in this framework: if for example an individual has genotypes for these 3 SNPs, then two haplotype phases are considered:

```
Observed Possible genotypes --> haplotypes A/C A/C G/G --> AAG / CCG ACG / CAG
```

whereas if the third SNP has a missing genotype (and if the other allele is T, for example) then the standard approach is just to consider a larger, consistent set (which are of course weighted by the current estimate of the population haploytpe frequencies):

```
        Observed
        Possible
        Possible

        genotypes
        -->
        haplotypes

        A/C A/C O/O
        -->
        A/C A/C G/G
        -->
        AAG / CCG

        ACG / CAG
        ACG / CAG
        ACG / CAG
```

In this way, if there is strong LD between SNPs, we can use the genotypes at flanking SNPs to effectively 'fill-in' missing genotype data. One advantage of this is that, if the genotypes are not missing at random for any given SNP, then it can give a less biased test to fill in the true values using LD information, rather than just to treat those genotypes as missing. This motivates a reframing of the basic single SNP association statistic in terms of groups of haplotypes rather than just as single SNPs (as shown above in the first example). Consider this example, involving simulated data, where the following haplotypes were simulated with these frequencies (in both cases and controls, so we would not expect any association with disease; 500 cases and 500 controls were generated).

Haplotype	Population	frequency
AABAB	0.4	
AABBA	0.2	
ABBBA	0.2	
BBBBB	0.1	
AAABB	0.1	

We will label the five SNPs, snp1, snp2, etc. Some non-random genotyping failure was simulated: in cases only, the BB genotype of snp3 only has a genotyping rate of 0.5 (i.e. half were set to missing). Such as pattern of genotyping failure, which is non-random with respect to both phenotype and genotype, can tend to produce spurious association results. For example, here are the basic single SNP results:

```
plink --file sim1 --assoc
```

which gives the output

CHR	SNP	A1	$F\_A$	$F_{-}U$	A2	CHISQ	Р	OR
1	snp1	В	0.102	0.106	Α	0.08585	0.7695	0.958
1	snp2	В	0.297	0.31	Α	0.3997	0.5272	0.9403
1	snp3	Α	0.1812	0.118	В	12.02	0.0005271	1.654
1	snp4	Α	0.406	0.383	В	1.107	0.2927	1.101
1	snp5	Α	0.388	0.393	В	0.05252	0.8187	0.9792

Note how snp3 shows a strong association (this is solely due to the non-random drop-out of genotypes for this SNP). However, the proxy association will, in this case, correct this:

```
plink --file sim1 --proxy-assoc snp3 --mind 1 --geno 1
```

Note that we use --mind and --geno to ensure that PLINK does not discard any individuals, in this particular case (i.e. we will use the flanking SNPs to fill in the missing data). This analysis gives the following output

*** Proxy	haplotype	associa	ation repo	ort for sr	ıp3 ***	
MAF	GENO	KB	RSQ	OR	CHISQ	P
0.104	0	-0.002	0.0145	0.958	0.0859	0.77
0.303	0	-0.001	0.0544	0.94	0.4	0.527
0.141	0.213	0	*	0.868	0.993	0.319
0.394	0	0.001	0.0813	1.1	1.11	0.293
0.39	0	0.002	0.08	0.979	0.0525	0.819
*	FREC	Į	OR	CHISQ	P	
ABBBA	0.199	) (	.945	0.254	0.615	
AABBA	0.191	L	1.03	0.0518	0.82	
	MAF 0.104 0.303 0.141 0.394 0.39 *	MAF GENO 0.104 0 0.303 0 0.141 0.213 0.394 0 0.39 0* FREC ABBBA 0.199	MAF GENO KB 0.104 0 -0.002 0.303 0 -0.001 0.141 0.213 0 0.394 0 0.001 0.39 0 0.002* FREQ ABBBA 0.199 0	MAF GENO KB RSQ 0.104 0 -0.002 0.0145 0.303 0 -0.001 0.0544 0.141 0.213 0 * 0.394 0 0.001 0.0813 0.39 0 0.002 0.08* FREQ OR ABBBA 0.199 0.945	MAF         GENO         KB         RSQ         OR           0.104         0         -0.002         0.0145         0.958           0.303         0         -0.001         0.0544         0.94           0.141         0.213         0         *         0.868           0.394         0         0.001         0.0813         1.1           0.39         0         0.002         0.08         0.979           .*         FREQ         OR         CHISQ           ABBBA         0.199         0.945         0.254	0.104       0       -0.002       0.0145       0.958       0.0859         0.303       0       -0.001       0.0544       0.94       0.4         0.141       0.213       0       *       0.868       0.993         0.394       0       0.001       0.0813       1.1       1.11         0.39       0       0.002       0.08       0.979       0.0525         .*       FREQ       OR       CHISQ       P         ABBBA       0.199       0.945       0.254       0.615

AABAB	0.394	1.1	1.11	0.293	
AAABB	0.111	0.868	0.993	0.319	
BBBBB	0.104	0.958	0.0859	0.77	
Haplotype frequency	estimation	based on	2000 of	2000 founder	chromosomes
Omnibus haplotype to	est statist:	ic: 1.88,	df = 4,	p = 0.759	
HAP	FREQ	RSQ	OR	CHISQ	P
.A BB	0.111	1	0.868	0.993	0.319
A. BB	0.111	1	0.868	0.993	0.319

In otherwords, instead of removing individuals who are missing for snp3 (which is implicitly what a single SNP association statistic would do) we use the flanking data to fill in the unobserved genotypes. Even if these are missing not-at-random, if there is strong LD then we will often be able to do a good job at guessing the true genotype. Note that the other SNPs (that have no missing genotype data) have identical association p-values under basic association test as under this constrained haplotype test, as would be expected (i.e. under most normal conditions, there is no loss of power in using a proxy-association approach).

**IMPORTANT** It is very important to remember that this test is not a panacea for the problem of missing data: many times there will not be sufficient LD to accurately reconstruct the missing genotype within the E-M. Future versions of PLINK aim to add diagnostics to indicate when this is the case; also, one might select the SNPs that define the flanking region more intelligently (e.g. making use of known patterns of LD, etc).

As such, thie results of this test should most probably be interpreted as follows: if a highly significant basic single SNP association result is not significant by this method, one would worry about biased missingness for that SNP; if a highly significant basic single SNP result remains highly significant, this is only meaningful when there is strong LD.

Of course, it is possible that other biases that are specific to haplotype analysis (the ability to estimate rare haplotype frequencies, etc) will impact these proxy tests, the effects of stratification may be more pronounced, etc. As such, these tests should be interpreted only as complementary pieces of information along with the basic SNP result, rather than as water-tight proof of an unbiased association *per se*.

However, if one knew up front that non-random genotyping drop-out might be an issue (for example, cases and controls from from different labs, different genotyping procedures used, etc) then it might seem prudent to take this approach.

Note Normally individuals are removed from the haplotype analysis if they are missing more than 50% of their genotypes for a given haplotype: in this case, we try to not remove individuals, but rather let the E-M fill in the missing data, so the rate is changed to 0.9 by default; this can be altered with the --hap-miss option.

# Chapter 14

# SNP imputation and association testing

This page describes PLINK functions to impute SNPs that are not directly genotyped but are present on a reference panel such as the HapMap. As well as imputing genotypes (either making the most likely call, or outputting the posterior probabilities of each genotype, or the dosage) some simple association tests can be framed in this context. These methods do not necessarily need whole-genome data to work however: with dense SNP genotyping in a particular region, these methods could still straightforwardly be applied.

These methods utilise the proxy association set of commands. The approach is a simple one, essentially based around the concept of multi-marker tagging. This approach is designed to provide a straightforward approach to imputation for common variants: it is unlikely to be optimal, particularly for rarer alleles, when compared to other imputation methods available.

In the text below, an *observed* SNP refers to one that was genotyped in both the reference and the WGAS sample. An *imputed* SNP refers to one that only appears in the reference panel.

**IMPORTANT** These features are still in *beta* meaning that they are still actively being updated, optimised, changed and fixed. As such, you are advised only to use these routines in an exploratory manner, if at all. More details regarding the specific procedures, and interpretation of results, will be posted presently. That is, rather than e-mail for more details, please wait until they are posted here.

# 14.1 Basic steps for using PLINK imputation functions

The first step is to create a single fileset with the reference panel merged in with your dataset. We assume that the HapMap CEU founders will be used in this example.

**HINT** A PLINK binary fileset of the Phase 2 HapMap data can be downloaded from here. For studies of individuals of European ancestry, the CEU founder fileset will be the one to download from that link.

Given the HapMap data, hapmap-ceu.\* or hapmap-ceu-all.\*, for example, you merge in your WGAS data as follows,

./plink --bfile hapmap-ceu --bmerge mydata.bed mydata.bim mydata.fam --make-bed --out merged

In imputation mode, the reference panel is denoted by making those individuals have a *missing value* for the phenotype. You will therefore need to edit the .fam files to make the 6th column (phenotype) 0 for all HapMap individuals and 1 (control) or 2 (case) for the individuals in your sample. If you have trio data, make sure that no observed individuals have missing phenotypes (i.e. set parents to controls in a TDT context, rather than have a missing phenotype code).

### 14.1.1 Strand issues

The HapMap SNPs are all given on the +ve strand, and so it is your responsibility to ensure that your data are aligned also, for the merge to work. The --flip command can help changing strand. If there are strand problems, PLINK will report a list of SNPs that did not match in terms of strand. Naturally, if there are SNPs A/T or C/G SNPs in your dataset, these will potentially go unflagged. As such, it is always a good idea to check allele frequencies between the HapMap and the WGAS sample to identify grossly deviant SNPs and/or undetected strand issues (i.e. create an alternate phenotype file with the HapMap individuals coded as controls and the rest of WGAS data as cases, and run a basic association command). The --flip-scan command can also help to detect some incorrectly aligned variants.

**NOTE** This will create a **very large** dataset and take some time; particularly if you have a parallel computing environment available, you might want to split the files and the merge procedures up by chromosomes, e.g. first download the archive with the HapMap CEU founder fileset split by chromosome, then merge each chromosome separately:

```
./plink --bfile mydata --chr 1 --make-bed --out data-1
./plink --bfile mydata --chr 2 --make-bed --out data-2
etc, followed by
./plink --bfile hapmap-ceu-chr1 --bmerge data-1.bed data-1.bim data-1.fam --make-bed
--out merged-1
./plink --bfile hapmap-ceu-chr2 --bmerge data-2.bed data-2.bim data-2.fam --make-bed
--out merged-2
```

This will create 22 separate filesets (merged-1, merged-2, etc) and all the following routines can then be run separately on each.

# 14.2 Combined imputation and association analysis of case/control data

Given the merged fileset, containing both the reference panel and the (more sparse) WGAS samples, PLINK will attempt to perform case/control association for every SNP (both observed and imputed) with the following command:

```
./plink --bfile merged-1 --proxy-assoc all
which will generate an output file
     plink.assoc.proxy
with the fields
     CHR
             Chromosome code
     SNP
             SNP identifier
     BP
             Physical position (base-pairs)
     A 1
             First allele code (not necessarily minor allele)
     A2
             Second allele code (not necessarily major allele)
             Genotyping rate in entire sample and reference panel
     GENO
     NPRX
             Number of proxy SNPs selected
     INFO
             Information content metric
     F_A
             Allele 1 frequency in cases
     F_{-}U
             Allele 1 frequency in controls
```

- OR Odds ratio
- P Significance value of case/control association test

The fields INFO and NPRX refer to how well PLINK managed, if at all, to impute the SNP. If NPRX is zero, then it could not be even poorly imputed. If INFO ranges from between 0 and 1, although it can be greater than 1 occasionally. A higher value general means a better imputed SNP; roughly speaking, only looking at imputed SNPs with a INFO value greater than 0.8 or so is probably good practice. More specific details on these metrics will be posted soon.

### 14.3 Modifying options for basic imputation/association testing

One of the most important modofying options for the --proxy-assoc test is --proxy-drop, which means that the *observed* SNPs are dropped, one at a time, from the WGAS sample when they are tested as the reference SNP (i.e. they will be re-imputed given the surrounding SNPs). That is, the command,

```
./plink --bfile merged-1 --proxy-assoc all --proxy-drop
```

would mean that every single SNP test statistic in plink.assoc.proxy would not involve a single observed genotype for that particular SNP; as such, running this association test with the --proxy-drop command is a good idea as it will provide both a means to assess the performance of the imputation (by comparing the results against the results of the observed genotypes) but also of an extra level of QC (if you still see a significant result, it cannot be due to technical artifacts specific to that SNP, as no observed genotypes were used in the test for that SNP).

The value of not using --proxy-drop always with --proxy-assoc (given that the basic --assoc command more straightforwardly calculates association for observed SNPs) is if there is a reasonable amount of missing genotype data for an observed SNP and you want to use imputation to recover it. (Although, in this case, there is perhaps less need to use a separate reference panel in any case, and so the standard proxy association approach, without any reference panel, can be used.)

### 14.3.1 Parameters modifying selection of proxies

Imputation in this context works simply by selecting a set of proxy SNPs (using the reference panel information) and then phasing these SNPs in both reference panel and WGAS sample jointly. By grouping haplotypes, the corresponding single SNP tests of *imputed* SNPs can then be straightforwardly performed.

There are a number of parameters that impact the choice of proxy SNPs. Fine tuning of these parameters is still in progress. These parameters will be described in more detail shortly. For now, the default parameters should be sufficient in most cases. See the proxy association page for a description of the parameters, the defaults, and how they can be changed.

# 14.4 Imputing discrete genotype calls

The association test described above performs imputation on-the-fly and does not save the imputed genotype calls or probabilities. To do so, and to generate other metrics of imputation performance, use the --proxy-impute command.

To generate summary statistics for the imputation performance of each SNP, use the command

```
./plink --bfile merged-1 --proxy-impute all
which produces a file
    plink.proxy.impute
which has the fields
    CHR Chromosome
```

SNP	SNP ID
NPRX	Number of proxy SNPs
INFO	Information metric
${\tt TOTAL\_N}$	Total number of WGAS sample genotypes (exc. reference panel)
OBSERVD	Proportion of these w/ observerd genotypes
IMPUTED	Proportion of these imputed
OVERLAP	Proportion of SNPs with both an imputed and overlapping
CONCORD	Concordance rate in the overlapping set

Here are some example lines:

CHR	SNP	NPRX	INFO	$TOTAL_N$	OBSERVD	IMPUTED	OVERLAP	CONCORD
18	rs7233673	5	0.993	3469	0	0.991	0	NA
18	rs7233597	5	0.998	3469	0.999	0.993	0.992	0.986
18	rs7505507	4	0.632	3469	0.999	0.332	0.332	0.891

e.g. the first line represents an unobserved SNP, for which 99% of individuals were imputed; the second line was an observed SNP, but if we drop it and try to re-impute, we get 99.3%; the concordance rate between imputed and genotyped is 98.6% for this SNP. The final line represents a SNP that did not perform as well: we only impute a third of genotypes and these are less than 90% concordant (this was an observed SNP also). In this case, we see the INFO score is lower (below 0.8) for this third SNP than for the other two: at the standard 0.8 threshold this SNP would have been ignored in any case.

The required confidence threshold for making a call can be changed with, for example,

```
--proxy-impute-threshold 0.8
```

(it is set to 0.95 by default currently).

To give genotype-specific concordances, use the additional option:

```
--proxy-genotypic-concordance
```

then a set of extra fields are append to the plink.proxy.impute output

```
F_AA Frequency of true 'AA' genotype
I_AA Proportion imputed for true AA genotype
C_AA Concordance rate for true AA genotype
F_AB As above, for 'AB' genotype
```

That is, for a very rare SNP, overall concordance would be high just by chance, even if none of the rare genotypes were correctly called. This option is therefore useful to get a better picture of imputation performance (when the observed genotype is also available).

In addition, if

```
--proxy-show-proxies
```

is also specified, an extra PROXIES field will appear in plink.proxy.impute showing the specific SNPs selected.

To perform imputation and save the dosages (fractional count of 0 to 2 alleles for each genotype), add the --proxy-dosage option;

```
./plink --bfile merged-1 --proxy-impute all --proxy-dosage
```

which produces a file

```
plink.proxy.impute.dosage
```

in which each imputed SNP is represented as a row; the fields (which does not have any header row)

```
SNP Identifier
Allele 1 code
Allele 2 code
Information content score for SNP
```

```
Allele dosage for first individual in sample
Allele dosage for second individual in sample
Allele dosage for final individual in sample
```

This file can then be analysed outside of PLINK.

To perform imputation and save the called (most likely) genotypes in a new fileset, add the --make-bed option;

```
./plink --bfile merged-1 --proxy-impute all --make-bed --out imputed-1
```

By default, PLINK will only replace genotypes that were missing in the original WGAS sample; to make PLINK re-impute all genotypes (whether they were actually observed or not), add the --proxy-replace flag,

```
./plink --bfile merged-1 --proxy-impute all --proxy-replace --make-bed --out imputed-1
```

Note Future versions will do obvious things, like let you generate proxy-impute and proxy-assoc output files in the same run (you can't now).

Important Making discrete calls for the most likely genotype will necessarily introduce error and bias in the all but perfectly imputed SNPs. As such, one should take care in the analysis and interpretation of imputed datasets – they should not be treated as if they were directly observed with certainty. In particular, one should be particularly cautious when combining multiple imputed files, particularly if different platforms were used and/or if the files also differ by disease state. Indeed, such an analysis is currently not recommended.

#### Verbose output options 14.5

To get a verbose output for a single SNP in the association mode, use instead of the all keyword the specific SNP name:

```
--proxy-assoc rs123235
```

See the web-page on proxy association methods to interpret this output.

You can also specify verbose imputation for one or more SNPs, e.g.

```
--proxy-impute rs8096534 --proxy-verbose
```

which will add extra lines to the file plink.proxy.impute representing the actual calls per person:

```
rs8096534
                78-03C15376 TBI-78-03C15376-1
                                                 01 01 0 1 0
rs8096534
                78-03C15377 TBI-78-03C15377-1
                                                 00 00 1 0 0
rs8096534
                78-03C15378 TBI-78-03C15378-1
                                                 01 01 0 1 0
rs8096534
                78-03C15398 TBI-78-03C15398-1
                                                 00 00 1 0 0
                78-03C15448 TBI-78-03C15448-1
                                                 01 01 0 1 0
rs8096534
rs8096534
                78-03C20292 TBI-78-03C20292-1
                                                 11 11 0 0 1
                78-03C20300 TBI-78-03C20300-1
                                                 11 10 0 0.08199 0.918
rs8096534
rs8096534
                78-03C20317 TBI-78-03C20317-1
                                                 01 01 0 1 0
                78-03C20335 TBI-78-03C20335-1
                                                 01 01 0 1 0
rs8096534
```

where the fields are (note: currently there is no header for these fields)

```
SNP
        SNP identifier
FID
        Family ID
IID
        Individual ID
OBS
        Observed genotype (coded 00,01,11 = AA,AB,BB, 10 = missing)
        Imputed genotype (as above)
TMP
        Probability of 'AA' genotype
PAA
```

```
PAB Probability of 'AB' genotype
PBB Probability of 'BB' genotype (i.e. these last 3 numbers sum to 1.00)
```

In addition, after these lines you will see a table of counts which summarises the actual calls versus the true values (if known). Ideally, you would observe high numbers down the diagonal therefore (the columns are the same as the rows):

Imputati	ion matr	ix (rows	observed	d, columns imputed)
A/A	292	2	0	1
A/G	0	1389	8	55
G/G	0	5	1585	130
0/0	1	1	0	0

and this is then followed by the normal, single-line non-verbose report for that SNP

CHR	SNP	NPRX	INFO	${\tt TOTAL\_N}$	OBSERVD	IMPUTED	OVERLAP	CONCORD
18	rs8096534	5	0.961	3469	0.999	0.946	0.946	0.995

Although you are able to specify --proxy-impute all and --proxy-verbose together, be warned that this will typically result in a very large output file for real data. It is better used for single SNPs in its current format.

# Chapter 15

# LD-based result clumping procedure

This page describes PLINK's ability to group SNP-based results across one or more datasets or analyses, based on empirical estimates of linkage disequilibrium between SNPs. The basic procedure was inspired by a script written by Ben Voight.

There are probably two main applications for this method:

- To report the top X single SNP results from a genome-wide scan in terms of a smaller number of clumps of correlated SNPs (i.e. to assess how many independent loci are associated, for example)
- To provide a quick way to combine sets of results from two or more studies, when the studies might also be genotyped on different marker sets

### 15.1 Basic usage for LD-based clumping

The --clump command is used to specify one or more result files (i.e. precomputed analyses of some kind). By default, PLINK scans these files and extracts fields with the headers SNP and P. For example:

```
plink --file mydata --clump mytest1.assoc
which generates a file
    plink.clumped
```

The actual genotype dataset specified here (i.e. the mydata.\* fileset) may or may not be the same dataset that was used to generate the results in mytest1.assoc. The mydata fileset is only used to calculate linkage disequilibrium between the SNPs that feature in mytest1.assoc (i.e. the analyses are not re-run).

There are four main parameters that determine the level of clumping, listed here in terms of the command flag used to change them and their default values:

```
--clump-p1 0.0001 Significance threshold for index SNPs
--clump-p2 0.01 Secondary significance threshold for clumped SNPs
--clump-r2 0.50 LD threshold for clumping
--clump-kb 250 Physical distance threshold for clumping
```

The clumping procedure takes all SNPs that are significant at threshold p1 that have not already been clumped (denoting these as  $index\ SNPs$ ) and forms clumps of all other SNPs that are within a certain kb distance from the index SNP (default 250kb) and that are in linkage disequilibrium with the index SNP, based on an r-squared threshold (default 0.50). These SNPs are then subsetted based on the result for that SNP, as illustrated below. This is a greedy algorithm and so each SNP will only appear in a single clump, if at all.

In the default, non-verbose mode, the default output lists all index SNPs and a summary of the other SNPs that are clumped with this SNP: (note, SNP IDs and positions are made-up in the example below):

CHR	F	SNP	BP	P	TOTAL	NSIG	S05	S01	S001	S0001	SP2
8	1	rs1234564	15716326	5.01e-07	0	0	0	0	0	0	NONE
14	1	rs1205236	69831825	1.46e-06	0	0	0	0	0	0	NONE
2	1	rs16331058	114547107	2.33e-06	3	0	0	0	0	3	rs2366902(1)
2	1	rs759966	54902416	9.28e-06	4	0	0	0	3	1	rs12538389(1)
11	1	rs8031586	44633498	9.75e-06	1	0	0	0	0	1	rs802328(1)
12	1	rs12431413	30028246	9.89e-06	0	0	0	0	0	0	NONE
6	1	rs14966070	62091121	1.07e-05	0	0	0	0	0	0	NONE

where the fields are as follows

```
CHR
                Chromosome code
       F
                Results fileset code (1,2,...)
       SNP
                SNP identifier
       BP
                Physical position of SNP (base-pairs)
       TOTAL
                Total number of other SNPs in clump (i.e. passing --clump-kb and --clump-r2
thresholds)
                Number of clumped SNPs that are not significant (p > 0.05)
       NSIG
       S05
                Number of clumped SNPs 0.01 < p < 0.05
       S01
                Number of clumped SNPs 0.001 < p < 0.01
       S001
                Number of clumped SNPs 0.0001 < p < 0.001
                Number of clumped SNPs p < 0.0001
       S0001
       SP2
                List of SNPs names (and fileset code) clumped and significant at --clump-p2
```

That is, the TOTAL field lists all SNPs that are clumped with the index SNP, irrespective of the p-value for those SNPs. This number is then split into those clumped SNPs that are not significant ( $p_{\xi}0.05$ ) and various other groups defined by significance thresholds. For SNPs that are significant at the p2 threshold, they are listed explicitly. The (1) after each SNP name refers to the results file they came from (in this case, there is only a single result file specified, so all values are 1).

To specify more than a single result file, use a comma-delimited list after --clump (without any spaces between file names), for example:

```
plink --bfile mydata --clump mytest1.assoc,mytest2.assoc
```

To specify a field labelled other than P, use the command

```
{\tt plink --bfile \ mydata \, --clump \ mytest1.assoc \, --clump-field \ P\_CMH}
```

for example.

**NOTE** The same fields are extracted from all results files (e.g. SNP and P) - i.e. it is not possible to specify different fields from different files.

**NOTE** All results are interpreted as p-values – i.e. it is not possible to specify a Z-statistic, as significance is always defined as less than the threshold. Finally, by default a SNP is not allowed to appear in more than one clump, either as an index or non-index SNP. If you add the command, then a SNP that has appeared as a non-index SNP in one clump can appear as a non-index SNP in other clumps:

```
--clump-allow-overlap
```

# 15.2 Verbose report

For a more detailed report of the SNPs in each clump, add the flag --clump-verbose

```
plink --bfile mydata --clump mytest1.assoc --clump-verbose
```

which produces a report as follows:

I IUIAL	- NOIG	S05	S01	S001	S0001
-07 0	0	0	0	0	0
P TOTAL	NSIG	S05	S01	S001	S0001
-06 0 	)	0	0	0	0
P TOTAL	L NSIG	S05	S01	S001	S0001
-06 3	3 0	0	0	0	3
) ALLELES	F	P	)		
) A	1	2.34e-06	3		
AT/GC	1	4.42e-05	5		
AC/GT	1	1.28e-05	5		
AT/GC	1	2.68e-05	5		
1	P TOTAL  P TOTAL  P TOTAL  OF ALLELES  OF A  AT/GC  AC/GT	P TOTAL NSIG -06 0 0  P TOTAL NSIG -06 3 0  Q ALLELES F 0 A 1 1 AT/GC 1 5 AC/GT 1	P TOTAL NSIG S05 -06 0 0 0  P TOTAL NSIG S05 -06 3 0 0  Q ALLELES F F D A 1 2.34e-06 1 AT/GC 1 4.42e-05 5 AC/GT 1 1.28e-05	P TOTAL NSIG S05 S01 -06 0 0 0 0  P TOTAL NSIG S05 S01 -06 3 0 0 0  Q ALLELES F P 0 A 1 2.34e-06 1 AT/GC 1 4.42e-05 5 AC/GT 1 1.28e-05	P TOTAL NSIG S05 S01 S001 -06 0 0 0 0 0  P TOTAL NSIG S05 S01 S001 -06 3 0 0 0 0  Q ALLELES F P 0 A 1 2.34e-06 1 AT/GC 1 4.42e-05 5 AC/GT 1 1.28e-05

etc

For example, for the third SNP, rs16331058 we see there are 3 other SNPs that fulfil the specified criteria (kb distance less than 250kb, r-squared greater then 0.5 and p-value of less than p2 threshold of 0.01), and they are listed explicitly in verbose mode. As well as the kb and r-squared for each SNP (relative to rs16331058) we see listed the fileset which the result comes from (F – in this case, all are listed 1, as there was only one result file specified) and p-value. Also, the alleles column indicates for the index SNP what the minor allele is (A); for the other SNPs, the two haplotypes that are more common than expected are listed (e.g. for SNPs A/B and 1/2, then if P(A1) > P(A)P(1) it will list A1/B2, otherwise A2/B1.

### 15.2.1 Annotation by SNP details and genomic co-ordinates

Another useful verbose-mode option is --clump-anotate which takes as a parameter a comma-delimited list of header names, e.g.

### --clump-annotate A1,OR

and will then list these items in the verbose report mode (e.g. minor allele and odds ratio, in this case, if the results file were a plink.assoc file). The output would then appear as, for example,

CHR	F	SNP	BP	F	TOTAL	NSIG	S05	S01	S001	S0001
2	1	rs16331058	114547107	2.337e-06	3	0	0	0	0	3
			KB	RSQ	ALLELES	F	P		ANNOT	
(INDE	EX)	rs16331058	0.0	1.000	Α	1	2.34e-06	Α,	1.23	
		rs2366902	-75.4	0.611	AT/GC	1	4.42e-05	Τ,	1.17	
		rs1274528	-47.4	0.555	AC/GT	1	1.28e-05	С,	1.22	
		rs3200591	-22.3	0.964	AT/GC	1	2.68e-05	Τ,	1.19	

i.e. here we can see that for rs2366902 the minor allele T had an odds ratio of 1.17; this is consistent with the index SNP, as the haplotype AT is more common than expected (i.e. indicating the direction of the LD).

NOTE The allele coding in the ALLELES field is taken directly from the specified genotype data, i.e. mydata.\* in this case, whereas the allele coding in the ANNOT field is taken (if available and --clump-annotate selects an allele field) from the results file. It is up to the user to ensure that these match to be interpretable (i.e. in terms of number versus letter coding, but more importantly in terms of strand, etc, which might be an issue if the genotype data is a file different from that which the results were calculated on, e.g. see below for an example).

A further option is --clump-range, which takes a gene-list or region-list file as a parameter. For example, this might be a list of all RefSeq genes, as available here. The command

plink --bfile mydata --clump myresults.assoc --clump-range glist-hg18

would, for example, generate the additional file

plink.clumped.ranges

which has the fields

CHR Chromosome code
SNP Index SNP per clump
P p-value

N Number of clumped SNPs POS Genomic co-ordinates KB kb span of clumped SNPs

RANGES List of ranges/genes that intersect the clumped region

For example, the first four rows of a simulated, random study are:

CHR	SNP	Р	N	POS	KB	RANGES
17	rs9944528	1.927e-05	2	chr17:7789403977933018	38.979	[UTS2R,SKIP,FLJ35767
9	rs17534370	1.958e-05	1	chr9:7029717270297172	0	[PGM5]
11	rs12418173	1.965e-05	7	chr11:112102294112133479	31.185	[]

which indicates that rs9944528 has one other SNP that clumps with it (N=2), which is just under 40kb away, spanning three genes; the next SNP doesn't have any clumped partners and falls in the PGM5 gene; the third SNP has 6 other clumped SNPs, spanning just over 30kb, but no genes are in that interval.

If the --clump-range flag is added in --clump-verbose mode, the output looks slightly different. In this case, the special plink.clumped.ranges file is not produced: now all the output is in the plink.clumped file:

CHR	F		SNP	BP		P TOTA	AL	NSIG	S05	S01	S001	S0001
17	1	rs994	14528	77894039	1.93e-0	5	1	0	0	0	0	1
				KB	RSQ	ALLELES	F	7	Р			
(INI	DEX)	rs994	14528	0	1.000	С	1	L	1.93e-05			
		rs720	7095	39	0.648	CG/GA	1	L	2.83e-05			
				:77894039		,		-				
		SPAN:										
(	CENE	S w/SNPs		D								
`	GLIVE:	•		r 2R,SKIP,FLJ	125767							
		GENES	. 015	ZR,SKIF,FL	133707							
CHR	F		SNP	BP		P TOTA	AL	NSIG	S05	S01	S001	S0001
9	1	rs1753	34370	70297172	1.96e-0	5	0	0	0	0	0	0
		GENES	: PGM	5								
CHR	F		SNP	BP		P TOTA	AL	NSIG	S05	S01	S001	S0001
11	1	rs1241	8173	112133479	1.96e-0	5	6	0	0	0	2	4
				KB	RSQ	ALLELES	F	7	P			
(INI	DEX)	rs1241	8173	0	1.000	G	1	_	1.96e-05			
		rs1280	0322	-31.2	0.902	GG/AC	1	Ĺ	0.000133			
		rs187	70496	-30.7	0.853	GC/AT	1	_	0.000267			
		rs219	9197	-20.1	1	GG/AA	1	_	9.76e-05			
		rs793	31135	-16.7	1	GG/AA	1	_	1.96e-05			
		rs1241	8739	-10.8	1	GA/AC	1	_	3.5e-05			
			98311	-4.98	1	GT/AC	1		1.96e-05			
		RANGE:	chr11	:112102294.	.11213347	-						
		SPAN.										

SPAN: 31kb GENES w/SNPs: GENES: -----

Note, if there is more than 1 SNP in a clump, we distinguish here between whether or not one of the clumped SNPs is actually within the a specified region or gene (GENES w/SNPs) versus whether that gene or region is just within the general clumped range (GENES).

Naturally, any file can be used with --clump-range - the regions do not have to correspond to actual genes, but they could be regions of interest identified by other means.

Finally, the command

--clump-range-border 20

adds a 20kb border to the start and stop of each gene or region.

# 15.3 Combining multiple result files (potentially from different SNP panels)

When more than one output file is specified, e.g. as

```
plink --bfile mydata --clump mytest1.assoc,mytest2.assoc,mytest3.assoc
```

there are two other options that can modify the behaviour of --clump. First,

--clump-index-first

indicates that index SNPs should only taken from the first result file listed (mytest1.assoc in the example above). In other words, this allows for an asymmetric comparison, in which we ask only whether or not a result in a particular file has any other SNPs (in that same, or in different files) that could be clumped.

Second, the additional option

```
--clump-replicate
```

means that only clumps containing clumped SNPs with p2-significant results in *more than one* result file are shown. This could be used in the following context: imagine one had data for two different whole-genome scans, for the same phenotype but performed on different platforms, e.g. Affymetrix and Illumina. A quick way to compare these sets of results would be to use the HapMap as a common dataset (i.e. containing all SNPs on both platforms, or the majority of these in any case) as follows:

```
plink --bfile hapmap --clump affymetrix.assoc,illumina.assoc --clump-verbose --clump-replicate
```

This assumes that you have made the fileset hapmap.\* to contain all SNPs for one of the analysis panels, e.g. CEU. In this context, we are only interested in hits (e.g. p-values less than 1e-3) that are seen across the studies, by using the --clump-replicate flag (i.e. only clumps where F is seen to have values of both 1 and 2 for p2-significant SNPs). In this case, it also probably makes sense to equate the p1 and p2 thresholds, by adding, for example,

```
--clump-p1 1e-3 --clump-p2 1e-3
```

Finally, by also adding the

--clump-annotate A1,OR

flag, you can see whether or not there appears to be a consistent direction of effect also (by putting together the direction of odds ratios with the over-represented haplotype to tie together the two or three SNPs).

# 15.4 Selecting the single best proxy

The command

--clump-best

produces an additional file

### plink.clumped.best

which contains the fields

INDEX	Index SNP identifier
PSNP	Best proxy SNP
RSQ	LD (r-squared) between index and proxy
KB	Physical distance between index and proxy
P	p-value for proxy SNP
ALLELES	The associated haplotypes for the index and proxy SNP
F	Which file (fromclump) this result came from

For example, if we use the command

```
plink --bfile mydata --clump myresults-a.assoc, myresults-b.assoc --clump-best
```

based on dummy simulated data result files myresults-a.assoc and myresults-b.assoc, the first few lines of plink.clumped are as follows:

CHR	F	SNP	BP	P	TOTAL	NSIG	S05	S01	S001	S0001	SP2
11	1	rs2513514	75922141	2.27e-07	3	0	0	0	1	2	rs2508756(1),
20	1	rs6110115	13911728	8.24e-07	9	0	2	3	2	2	rs6079243(1),
11	1	rs2508756	75921549	1.07e-06	0	0	0	0	0	0	NONE
15	1	rs16976702	54120691	1.15e-06	1	0	0	0	1	0	rs16976702(2)

The corresponding plink.clumped.best file shows the single best proxy SNP for each index SNP. This information could have been extracted manually after using the --clump-verbose, but the --clump-best option simply makes this easier.

F	ALLELES	P	KB	RSQ	PSNP	INDEX
2	AA/GG	8.04e-05	0	1	rs2513514	rs2513514
2	CC/AA	0.00145	0	1	rs6110115	rs6110115
NA	NA	NA	NA	NA	NA	rs2508756
2	GG/CC	0.0009	0	1	rs16976702	rs16976702

For example, the best SNP, rs2513514 (which had the lowest p-value in this case for F1, i.e. myresults-a.assoc) has a single best proxy of rs2513514, the same SNP, but in F2, i.e. myresults-b.assoc. The third SNP here, rs2508756, does not have any proxy SNP that meets the criteria for clumping (--clump-r2, --clump-p2, etc).

One might often want to add the three options

- --clump-index-first
- --clump-replicate
- --clump-allow-overlap

along with --clump-best. This would pose the question: what is the best proxy in myresults-b.assoc (i.e. --clump-replicate forces a cross-file proxy) for the top results in myresults-a.assoc (e.g. --clump-index-first forces the first-listed file to contain index SNPs only). The --clump-allow-overlap will mean that a proxy SNP can be selected for more than one index SNP, if it is the best. These may sometimes be the same SNP, if it is present in both result sets, otherwise it will rely on all SNPs being present in the mydata fileset, and will use LD information to select the best proxy.

# Chapter 16

# Gene reporting tool

The functions listed here are designed to provide a quick and easy way to partition any PLINK results file that indexes SNPs based on chromosome and base-pair position in terms of genes.

### 16.1 Basic usage

The basic command to produce a gene-centric report of single SNP results, for example from run1.assoc, is

```
./plink --gene-report run1.assoc --gene-list glist-hg18
```

which assumes the file run1.assoc will have a standard header row containing the fields CHR and BP, which it will if it was created by the PLINK --assoc command previously. It is not necessary that the original genotype filesets be present when running this command.

The gene list, glist-hg18, should a standard text file in the following format: one row per gene, chromosome, start and stop positions (base-pair) and then gene name, e.g.

```
7 20140803 20223538 7A5
19 63549983 63556677 A1BG
10 52236330 52315441 A1CF
8 43266741 43337485 A26A1
15 19305252 19336667 A26B1
21 13904368 13935777 A26B3
```

These files are available for download from the resources section of this web-site.

This generates a file

```
plink.range.report
```

which simply takes the lines of the results file, and lists them by the genes specified in the gene-list file. The listing is alphabetical by gene name. For example,

	ACO2 chr22:4019507440254939 ( 59.865kb )									
	DIST	CHR	SNP	BP	A1	$F_{-}A$	$F_{-}U$	A2	CHISQ	Р
OR										
	13.22kb	22	rs2267435	40208294	3	0.3958	0.3537	1	0.3351	0.5627
1.197										
	24.84kb	22	rs2076196	40219909	1	0.3333	0.2683	3	0.8852	0.3468
1.364										
	57.13kb	22	rs1810460	40252200	4	0.04167	0.07317	2	0.8278	0.3629
0.5507										
	ADORA2A chr22:2315352923168325 ( 14.796kb )									

DIST CHR SNP BP  $F_A$  $F_{-}U$ A2 CHISQ Ρ A1 OR. 11.14kb 22 rs5760423 23164672 4 0.4592 0.4024 3 0.5854 0.4442 1.261

etc, which shows the lines of run1.assoc split by the genes the SNPs fall in. In this case, the first gene is ACO2; the location based on glist-hg18 is specified, along with the length. Then the SNPs within this gene are listed. If genes overlap, then the SNPs will be listed more than once. If a SNP does not fall within any gene or region specified, then it will not be listed here.

The first field, DIST is added, which represents the distance from the start position of the gene. (Note: if a border is added, with --gene-list-border, see below, then DIST can be negative, i.e. representing that the SNP is before the actual start of the gene.)

Naturally, the regions listed in the **--gene-list** file do not have to correspond to actual genes – for example, they might correspond to known linkage peaks, or regions with disease-related copy number variants, etc.

### 16.2 Other options

The following options modify this procedure:

```
--pfilter 0.01
```

will list only SNPs with p-values less than 0.01. This requires that the results file has a field labelled P in the header row.

The additional command

```
--gene-list-border 20
```

will add a 20kb border to the start and stop of each gene listed in the gene file.

The additional command

```
--gene-subset candidate.list
```

will make a report extracting only the genes listed in candidate.list from the file specified by --gene-list. For example, if the file candidate.list contained two schizophrenia candidate genes,

DISC1 COMT

then (assuming the genes listed here match a row in the gene-list file, glist-hg18)

will only report nominally significant (P=0.05) SNPs within or near (+/- 50kb) these two genes. This is designed to be a more convenient way to quickly query a focussed set of genes, so one can keep only a single, central gene-list file.  $jem_{\tilde{\ell}}$ 

This document last modified i/em;

# Chapter 17

# **Epistasis**

For disease-trait population-based samples, it is possible to test for epistasis. The epistasis test can either be case-only or case-control. All pairwise combinations of SNPs can be tested: although this may or may not be desirable in statistical terms, it is computationally feasible for moderate datasets using PLINK, e.g. the 4.5 billion two-locus tests generated from a 100K data set took just over 24 hours to run, for approximately 500 individuals (with the --fast-epistasis command). Alternatively, sets can be specified (e.g. to test only the most significant 100 SNPs against all other SNPs, or against themselves, etc). The output consists only pairwise epistatic results above a certain significance value; also, for each SNP, a summary of all the pairwise epistatic tests is given (e.g. maximum test, proportion of tests significant at a certain threshold, etc). To test for gene-by-environment interaction, see either the section on stratified analyses for disease traits, or the section on QTL GxE for quantitative traits.

**IMPORTANT!** These tests for epistasis are currently only applicable for population-based samples, not family-based.

# 17.1 SNP x SNP epistasis

To test SNP x SNP epistasis for case/control population-based samplee, use the command

```
plink --file mydaya --epistasis
which will send output to the files
    plink.epi.cc
    plink.epi.cc.summary
```

where cc = case-control; for quantitative traits, cc will be replaced by qt.

The default test uses either linear or logistic regression, depending on whether the phenoype is a quantitative or binary trait. PLINK makes a model based on allele dosage for each SNP, A and B, and fits the model

```
Y \sim b0 + b1.A + b2.B + b3.AB + e
```

The test for interaction is based on the coefficient b3.

Hint For disease traits only, an approximate but faster method can be used to screen for epistasis: use the --fast-epistasis command instead of --epistasis. This test is based on a Z-score for difference in SNP1-SNP2 assocation (odds ratio) between cases and controls (or in cases only, in a case-only analysis). If you use this to screen a large number of SNPs, you should probably report the more standard logistic regression test value also. In practice, both approaches usually give similar results, which justifies the use of --fast-epistasis as a screening tool for a computationally-demanding problem. Of course, given a specific (and often extreme) threshold, --epi1, the exact above-threshold list of SNPs will not always be the same;

if you choose to use this approach, it is probably wise to apply it to select a subset of pairs of SNPs below a reasonably liberal --epi1 threshold to be tested with the more standard --epistasis command.

Important The --epistasis command is set up for testing a potentially very large number of SNP by SNP comparisons, most of which would not be significant or of interest. Because the output may contains millions or billions of line, the default is to only output tests with p-values less than 1e-4, as specified by the --epi1 option (see below). If your dataset is much smaller and you definitely want to see all the output, add --epi1 1 . If you do not, odds are you'll see a blank output file except for the header (i.e. immediately telling you that none of the tests were significant at 1e-4).

#### Specifying which SNPs to test

There are different modes for specifying which SNPs are tested: ALL x ALL

```
plink --file mydata --epistasis

SET1 x SET1 where epi.set contains only 1 set

plink --file mydata --epistasis --set-test --set epi.set

SET1 x ALL where epi.set contains only 1 set

plink --file mydata --epistasis --set-test --set epi.set --set-by-all

SET1 x SET2 where epi.set contains 2 sets

plink --file mydata --epistasis --set-test --set epi.set
```

For the 'symmetrical' cases (ALLxALL and SET1xSET1) then only unique pairs are analysed.

For the other two cases (SET1xALL, SET1xSET2) then all pairs are analysed (e.g. will perform SNPA x SNPB as well as SNPB x SNPA, if A and B are in both SET1 and SET2). It will not try to analysis SNPA x SNPA however.

#### The output

The output can be controlled via

```
plink --file mydata --epistasis --epi1 0.0001
```

which means only record results that are significant p<sub>i</sub>=0.0001. (This prevents too much output from being generated). The output is in the form

```
CHR1 Chromosome of first SNP
SNP1 Identifier for first SNP
CHR2 Chromosome of second SNP
SNP2 Identifier for second SNP
OR_INT Odds ratio for interaction
STAT Chi-square statistic, 1df
P Asymptotic p-value
```

A second part of the output: for each SNP in SET1, or in ALL if no sets were specified, is information about the number of significant epistatic tests that SNP featured in (i.e. either with ALL other SNPs, with SET1, or with SET2). The threshold --epi2 determines this:

```
plink --file mydata --epistasis --epi1 0.0001 --epi2 0.05
```

The output in the plink.epi.cc.summary file containts the following fields:

```
CHR Chromosome SNP SNP identifier N\_SIG # significant epistatic tests (p <= "--epi2" threshold)
```

```
N_TOT # of valid tests (i.e. non-zero allele counts, etc)
PROP Proportion significant of valid tests
BEST_CHISQ Highest statistic for this SNP
BEST_CHR Chromosome of best SNP
BEST_SNP SNP identifier of best SNP
```

This file should be interpreted as giving only a very rough idea about the extent of epistasis and which SNPs seem to be interacting (although, of course, this is a naive statistic as we do not take LD into account – i.e. PROP does not represent the number of *independent* epistatic results).

#### 17.1.1 A faster epistasis option

## 17.2 Case-only epistasis

For case-only epistatic analysis,

```
plink --file mydata --fast-epistasis --case-only
sends output to (co = case-only)
    plink.epi.co
    plink.epi.co.summary
```

All other options are as described above.

Currently, in case-only analysis, only SNPs that are more than 1 Mb apart, or on different chromosomes, are included in case-only tests. This behavior can be changed with the --gap option, with the distance specified kb: for example, to specify a gap of 5 Mb,

```
plink --file mydata --fast-epistasis --case-only --gap 5000
```

This option is important, as the case-only test for epistasis assumes that the two SNPs are in linkage equilibrium in the general population.

## 17.3 Gene-based tests of epistasis

WARNING This test is still under heavy development and not ready for use.

## Chapter 18

## R plugin functions

This page describes PLINK's limited support for R-based 'plug-in' functions. In this manner, users can extend the basic functionality of PLINK to better meet their own needs.

R http://www.r-project.org/ is a powerful, freely-available package for statistical computing. PLINK uses the Rserve http://www.rforge.net/Rserve/ package to communicate with R. There are some notes on installing and running the Rserve package below.

The idea is that some analyses, such as survival analysis for example, are already implemented in R but not currently available in PLINK. Having a simple interface for accessing such R functionality, allows one to benefit from both the data-handling features of PLINK (i.e. it being designed specifically to handle large SNP datasets efficiently, in a way that the basic R package is not) as well as the ever-increasing library of statistical tools in R. Also, this should provide an easy way to prototype new methods, etc.

Currently there is only support for SNP-based analyses (i.e. there is a strict model that assumes we return one test statistic value for each and every SNP in the data set). Potentially (if there is interest/specific suggestions) this will be expanded such that other entities can be the unit of analysis, i.e. individuals, or groups of SNPs or individuals, or even the results of PLINK analyses, for graphing or post-processing.

That is, in this first release of R plug-in functionality only the basic, limited set of features is currently available, and there will likely still be some rough edges...

Note Currently, there is only support for R-plugins for Linux-based PLINK distributions.

**Note** Version 1.04 of PLINK has updated to client code to support the latest version of Rserve. You should re-install Rserve (see notes below) to make sure you have the latest version.

## 18.1 Basic usage for R plug-ins

Assuming Rserve has been installed and is running locally (see below) and that the file myscript.R contains the R code conforming to the standard for a PLINK plug-in (see here), then the command is simply

```
plink --file mydata --R myscript.R
which generates a file
    plink.auto.R
```

Currently this file just contains the raw output for each SNP, giving the value returned from R. Currently only a single value can be passed back: that is, there is no support for extracting more information per SNP. One approach it to explicitly write the desired information to a file from the R function you define (taking care not to overwrite it, as the function is recalled for every batch of SNPs sent to R).

The assumed use is more to screen a large number of SNPs, then extract ones of interest for more detailed follow-up analysis in R itself. The value you return need not be a p-value however – so you could always

set up functions to return other quantities (e.g. odds ratios). However, they must be single, floating point values (i.e. not strings or R objects).

**Note** Future developments will allow the plug-in function to be embedded within permutations, gene-based tests, etc, in the same way the basic -assoc command is.

## 18.2 Defining the R plug-in function

PLINK expects a function in the exact form

```
Rplink <- function(PHENO,GENO,CLUSTER,COVAR)</pre>
```

to be defined in the supplied file. This function is expected to return a numeric vector, with as many elements are there are SNPs. Internally, PLINK will call the Rplink function – it must be written exactly as shown here. The objects refer to:

```
PHENO vector of phenotypes (n)

GENO matrix of genotypes (n x 1)

CLUSTER vector of cluster membership codes (n)

COVAR matrix of covariates (n x c)
```

where n is the number of individuals (after pruing, filtering, etc) and c is the number of covariates (if any). PLINK generates these objects internally, so the user can assume these exist for when the Rplink() function is called. (In practice, the number of SNPs, 1 will probably be smaller than the total number of SNPs in the file, as PLINK passes the genotype data into R in batches rather than all in one go).

Genotypes are coded 0, 1, 2 and NA, as per the --recodeA option.

An example R plug-in is shown here – this is probably the most straightforward template for an R-plugin, in which the apply() function is used to iteratively call the nested function (f1()), once per SNP, in this case. For example, the file myscript.R might contain the following plug-in:

```
Rplink <- function(PHENO,GENO,CLUSTER,COVAR)
f1 <-function(x) mean(x, na.rm=T) / 2
as.numeric(apply(GENO, 2, f1))</pre>
```

If you are not familiar with the R language, there are a number of excellent resources available from the main R webpage http://www.r-project.org/.

The first line defines a function, f1() that calculates the allele frequency for each SNP (as the genotypes are coded as the count of the minor allele, 0,1,2). The second line applies this function to each column of the genotype data, using the apply( data, row/col, function) command.

Another, perhaps more useful, example is implementing survival analysis within PLINK: here we define a function, f1() to return the p-value for the first coefficient; we assume here that a censoring variable was loaded into PLINK as the first covariate (i.e. the R Surv function takes two parameters, the survival time and censoring status). (This is probably not the optimal way to implement this analysis, but is intended purely as an example of what can be done.)

```
library(survival)
Rplink <- function(PHENO,GENO,CLUSTER,COVAR)
f1 <- function(s)
  m <- coxph( Surv( PHENO , COVAR[,1] ) ~ s )
  summary(m)$coef[5]
apply( GENO , 2 , f1 )</pre>
```

## 18.3 Example of debugging an R plug-in

To generate a text file that contains the R commands PLINK would have run (rather than actually trying to run them – this is useful for debugging purposes), add the following flag

```
plink --file mydata --R myscript.R --R-debug
```

To illustrate how to use the debug function to figure out why things might not be working as expected, consider this example, in which we are trying to use this approach to implement a basic logistic regression. The file

```
mylog.R
```

which contains the function

Rplink <- function(PHENO,GENO,CLUSTER,COVAR)</pre>

```
f1 <- function(s)

m <- glm( PHENO ~ s , family="binomial" )
  summary(m)$coef[8]

apply( GENO , 2 , f1 )</pre>
```

and we have a dataset with three SNPs; the internal PLINK logistic regression command

```
plink --file mydata --logistic
```

yields

CHR	SNP	BP	A1	TEST	NMISS	ODDS	STAT	P
1	snp0	10000	Α	ADD	200	1.256	1.15	0.2501
1	snp1	10001	В	ADD	200	0.9028	-0.5112	0.6092
1	snp2	10002	В	ADD	200	0.6085	-2.242	0.02499

Trying to run the R implementation:

```
plink --file mydata --R mylog.R
```

we obtain a set of invalid p-values in plink.auto.R

```
snp0 -1
snp1 -1
snp2 -1
```

To find out what is happening, we will run the same command with the debug option

```
plink --file mydata --R mylog.R --R-debug
```

This writes to the file plink.auto.R the actual commands that would be passed to R, including the data and the function:

```
1 <- 3
    g \leftarrow c(1, 2, 1, 2, 2, 1, 1, 1, 1, 2, 2, 0, 0, 1, 1, 0, 1, 2, 1, 1, 1,
    2, 1, 1, 2, 1, 1, 0, 1, 1, 1, 0, 1, 2, 1, 1, 1, 0, 0, 0, 0, 1, 1, 1,
    1, 1, 1, 1, 0, 1, 2, 1, 1, 1, 1, 0, 1, 1, 0, 2, 2, 0, 1, 1, 2, 0, 1,
    1, 1, 2, 1, 1, 1, 1, 1, 0, 2, 2, 0, 0, 2, 1, 1, 1, 2, 1, 1, 0, 1, 1,
    1, 1, 2, 2, 2, 1, 0, 2, 0, 1, 1, 1, 0, 0, 1, 0, 1, 1, 0, 1, 1, 0, 1,
    0, 2, 2, 1, 0, 0, 0, 1, 0, 1, 2, 2, 2, 1, 0, 0, 0, 2, 1, 2, 2, 1, 1,
    1, 1, 0, 0, 1, 1, 1, 1, 1, 2, 1, 1, 0, 1, 2, 2, 1, 2, 2, 1, 2, 0, 1,
    1, 1, 1, 2, 1, 1, 0, 1, 0, 1, 1, 2, 1, 2, 1, 1, 1, 1, 1, 0, 1, 1, 1,
    2, 2, 1, 2, 1, 1, 2, 2, 0, 0, 1, 2, 1, 0, 0, 1, 1, 2, 1, 2, 2, 2, 0,
    1, 1, 0, 2, 1, 1, 2, 1, 0, 1, 1, 0, 1, 1, 1, 1, 2, 1, 1, 0, 1, 1, 0,
    0, 1, 1, 2, 1, 0, 1, 2, 0, 2, 1, 1, 1, 0, 0, 2, 1, 1, 1, 2, 0, 1, 1,
    1, 1, 1, 2, 1, 2, 0, 1, 1, 0, 1, 0, 2, 1, 0, 2, 1, 2, 2, 0, 0, 0, 1,
    1, 2, 1, 1, 1, 0, 2, 1, 0, 2, 2, 1, 1, 2, 1, 1, 1, 2, 0, 1, 1, 0, 1,
    2, 2, 2, 0, 1, 1, 1, 1, 1, 2, 1, 1, 2, 1, 2, 0, 0, 2, 0, 2, 1, 1, 1,
    1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 2, 1, 0, 0, 1, 1, 0, 2, 1, 1, 1, 2,
    2, 2, 1, 1, 0, 0, 2, 2, 1, 2, 2, 0, 2, 2, 2, 2, 0, 1, 2, 2, 2, 2, 0,
    0, 0, 1, 1, 1, 2, 1, 1, 2, 1, 1, 1, 2, 2, 2, 0, 1, 2, 0, 0, 1, 1, 1,
    0, 1, 1, 1, 0, 0, 1, 1, 2, 1, 0, 1, 0, 2, 2, 1, 2, 1, 1, 1, 0, 1, 1,
    1, 1, 1, 1, 1, 0, 1, 0, 1, 1, 2, 1, 1, 0, 0, 0, 0, 1, 0, 1, 0, 2, 2,
    2, 2, 1, 2, 1, 1, 1, 2, 1, 2, 0, 0, 1, 0, 1, 2, 1, 0, 2, 0, 1, 1, 0,
    1, 0, 1, 1, 0, 2, 0, 1, 2, 1, 1, 2, 2, 1, 2, 0, 2, 0, 2, 0, 0, 1, 1,
    1, 1, 2, 1, 0, 2, 0, 1, 1, 0, 1, 2, 2, 2, 1, 0, 1, 2, 1, 2, 1, 2, 0,
    0, 1, 0, 1, 1, 2, 0, 1, 1, 2, 1, 0, 1, 2, 1, 0, 2, 2, 2, 2, 2, 2, 1,
    0, 2, 1, 2, 1, 1, 1, 1, 2, 0, 1, 1, 1, 2, 2, 1, 0, 1, 1, 2, 1, 1, 0,
    1, 1, 2, 1, 1, 1, 2, 1, 1, 1, 1, 1, 2, 0, 2, 0, 2, 2, 1, 0, 1, 2, 1,
    0, 2, 0, 0, 1, 0, 2, 1, 0, 2, 1, 1, 1, 0, 1, 1, 1, 1, 1, 1, 1, 0, 2, 2,
    0, 1, 2, 1)
    GENO <- matrix( g , nrow = n ,byrow=T)</pre>
    GENO[GENO == -1] <- NA
    Rplink <- function(PHENO,GENO,CLUSTER,COVAR)</pre>
    f1 <- function(s)</pre>
      m <- glm( PHENO-1 ~ s , family="binomial" )</pre>
      summary(m)$coef[8]
    apply(GENO, 2, f1)
In R, load this function
 source("plink.auto.R")
and then try to run the Rplink function
 Rplink (PHENO, GENO, CLUSTER, COVAR)
```

and you will see the error message

```
Error in eval(expr, envir, enclos) : y values must be 0 \le y \le 1
```

which indicates that R is expecting a 0/1 coding for this particular function, not the default 1/2 coding used by PLINK for the phenotype/dependent variable. You might therefore want to change the relevant line of the function from

```
m <- glm( PHENO ~ s , family="binomial" )
to
    m <- glm( PHENO==2 ~ s , family="binomial" )
for example. Then, repeating the above debug procedure, you would see in R
    Rplink(PHENO,GENO,CLUSTER,COVAR)
gives
    [1] 0.25013412 0.60921037 0.02499268
which are the correct p-values. So, now the function is fixed running
    plink --file mydata --R mylog.R
would generate the same set of p-values as the PLINK logistic command, in plink.auto.R
    snp0 0.250134
    snp1 0.60921
    snp2 0.0249927</pre>
```

## 18.4 Setting up the Rserve package

First, you must ensure that you have Rserve installed on your system. Normally, this will involve just typing, at the R command prompt (not the system shell prompt)

```
install.packages("Rserve")
```

HINT For this to work, R must have been configured with --enable-R-shlib.

When using any R-based PLINK plug-in, Rserve must be running in the background before invoking the PLINK command. To start Rserve, just type at the shell prompt

```
R CMD Rserve
```

(note, you may need to change Rserve to the full path of where Rserve was installed), or, within R, type at the R prompt

```
library(Rserve)
Rserve()
```

Please see the Rserve documentation http://www.rforge.net/Rserve/doc.html for further support.

## Chapter 19

# SNP annotation database lookup

This page describes PLINK's ability to output basic annotation information on SNPs on common WGAS genotyping platforms, via a web-based lookup function.

The SNP annotation data were compiled by Patrick Sullivan's lab http://genetics.unc.edu/faculty/sullivan.htm; the original data files are available here https://slep.unc.edu/evidence/.

**NOTE** All gene names must be HUGO standard gene names . For example, the serotonin transporter is SLC6A4 (not HTT or SERT).

If you use these annotations in a publication, include the following sentence and corresponding references:

Using the PLINK retrieval interface, SNP annotations were created using the TAMAL database (1) based chiefly on UCSC genome browser files (2), HapMap (3), and dbSNP (4).

- Hemminger BM, Saelim B, Sullivan PF. TAMAL: An integrated approach to choosing SNPs for genetic studies of human complex traits. Bioinformatics 2006;22:626-7.
- Hinrichs AS, Karolchik D, Baertsch R, Barber GP, Bejerano G, Clawson H, Diekhans M, Furey TS, Harte RA, Hsu F, Hillman-Jackson J, Kuhn RM, Pedersen JS, Pohl A, Raney BJ, Rosenbloom KR, Siepel A, Smith KE, Sugnet CW, Sultan-Qurraie A, Thomas DJ, Trumbower H, Weber RJ, Weirauch M, Zweig AS, Haussler D, Kent WJ. The UCSC Genome Browser Database: update 2006. Nucleic Acids Res 2006;34:D590-8.
- Altshuler D, Brooks LD, Chakravarti A, Collins FS, Daly MJ, Donnelly P. A haplotype map of the human genome. Nature 2005;437:1299-320.
- Wheeler DL, Barrett T, Benson DA, Bryant SH, Canese K, Chetvernin V, Church DM, DiCuccio M, Edgar R, Federhen S, Geer LY, Helmberg W, Kapustin Y, Kenton DL, Khovayko O, Lipman DJ, Madden TL, Maglott DR, Ostell J, Pruitt KD, Schuler GD, Schriml LM, Sequeira E, Sherry ST, Sirotkin K, Souvorov A, Starchenko G, Suzek TO, Tatusov R, Tatusova TA, Wagner L, Yaschenko E. Database resources of the National Center for Biotechnology Information. Nucleic Acids Res 2006;34:D173-80.

## 19.1 Basic usage for SNP lookup function

The basic command is, for example,

```
plink --lookup rs1475515
```

which outputs to the LOG file the following information

PLINK-SNP (WGAS SNP annotation courtesy of Patrick Sullivan)

Connecting to web	
SNP ID	: rs1475515
Affy ID	:
Affy 5.0	: no
Affy 6.0	: no
Perlegen ID	:
Perlgen 600	: no
Illumina 650	: yes
Illumina 550	: no
Non-syn SNP	: no
SNP Error	: no
SNP Pos Duplication	: no
Chromosome	: 1
Strand	: -
HG17 Position (bp)	: 228459232
HG18 Position (bp)	: 230219120
Pseudo-autosomal region?	: N/A
NCBI reference allele	: T
UCSC reference allele	: A
Observed alleles	: C/T
Human alleles	: C/T
Predominant human allele	: A
Chimp allele	: T
Macaque allele	: T
dbSNP MAF	: 0.038
HapMap CEU MAF	: 0
HapMap ASI MAF	: 0
HapMap YRI MAF	: 0.15 : -
HapMap CEU Strand	: - : C
HapMap ASI Allala	: C
HapMap ASI Allele HapMap YRI Allele	. C
In gene transcript	:
In gene coding region	•
Nearby Genes(KB distance)	•
Segmental duplication?	: no
Copy Number Variant?	: no
Conservation >95% pctile?	: no
Conservation >99% pctile?	: no
Disease-causing region?	: no
miRNA target? (TargetScan)	: no
miRNA target? (PicTAR)	: no
Regulatory potential?	: yes
Promotor region? (Stanford)	: no
Promotor region? (firstEF)	: no
Transfactor binding site	: no
Enhancer?	: no
Exon?	: no
Consensus splice site?	: no
5' UTR?	: no
3' UTR?	: no

To perform a lookup query on a batch of SNPs rather than 1 at a time, use the command

```
plink --lookup-list hits.list
where hits.list is just a list of SNP IDs (RS numbers); this will generate a file
plink.snp.annot
```

containing multiple reports of the above kind. There is a limit to the number of SNPs that can be submitted at one time (currently 200).

## 19.2 Gene-based SNP lookup

It is possible to dump all SNPs in a gene with the command

```
plink --lookup-gene DISC1
```

which does two things: writes some gene-centric information to the LOG file, and lists all the SNPs that feature on common WGAS platforms to the file

```
plink.snp.list
```

By default, SNPs within 20kb upstream and downstream of the gene are recorded. To change this, add the command

```
--lookup-gene-kb 0
or
--lookup-gene-kb 100
```

for example.

In the information written to the LOG file, there is a strong bias towards neuropsychiatrically-relevant information, reflecting the research interests of the creator. For example, the output for DISC1 is: (note: there are a few relatively redundant or uninformative fields currently that will be removed in future releases)

```
Looking up gene information (and SNPs +/- 20 kb)
Connecting to web... Writing SNP details to [ plink.snp.list ]
Gene Name
Product
                                         : disrupted in schizophrenia 1 isoform Es
Entry
CCDS Name
                                         : CCDS31056.1
KG ID
                                         : uc001hux.1
SwissProt ID
                                         : Q9NRI5-4
                                         : 2888
Hugo ID
Hugo alias
Hugo old gene names
Has gene name?
                                         : no
HG18 strand
                                         : +
HG18 chrom
                                         : 1
HG18 TX Start
                                         : 229829236
HG18 TX End
                                         : 229924970
HG18 CDS Start
                                         : 229829236
HG18 CDS End
                                         : 229924970
                                         : 95734
HG18 TX Length
HG18 TX Length Percentile
                                         : 96
HG17 strand
HG17 chrom
                                         : 0
HG17 TX Start
                                         : 0
HG17 TX End
                                         : 0
```

```
HG17 CDS Start
                                               : 0
    HG17 CDS End
                                               : 0
    HG17 TX Length
                                               : 0
    Has HG17 pos
                                               : no
    mRNA accession numbers
                                               : NM_001012958.1 ENST00000317586 OTTHUMT00000092355
                                               : NP_001012976.1 ENSP00000320784 OTTHUMP00000035959
    Protein accession numbers
    Pseudoautosomal HG18
                                               : no
    Pseudoautosomal HG17
                                               : no
    Brain expressed 50th percentile
                                               : yes
    Brain expressed 75th percentile
                                               : yes
    Correlated cortex expression
                                               : NA
    Correlated lymphoblastoid expression
                                               : yes
    Number association studies from SZGene : 20
     Annotation from SLEP database
                                               : ? Schizophrenia [PMID=16033310]/Schizoaffective
                                               : disorder, susceptibility to, 181500 (3) [OMIM=605210]
                                               : /Schizophrenia, susceptibility to, 604906 (3)
[OMIM=605210]
    Association studies from GAD database
                                               : psych (16)
  It is possible to supply a list of genes to lookup, with the command
    plink --lookup-gene-list mygenes.txt
  that will dump the SNPs from multple genes in a SET file format, e.g. where the file
       mygenes.txt
  is something like
       COMT
       DISC1
       CACNA1C
  These could then be subsequently extracted with the command
        --extract plink.snp.list
  as the END comments and gene names will just be ignored if these are not SNP IDs in the MAP file.
```

## 19.3 Description of the annotation information

For a detailed description of the annotation fields and how they were compiled, please see Patrick Sullivan's PDF https://slep.unc.edu/evidence/files/README\_annotations.pdf

## Chapter 20

## SNP simulation routine

PLINK provides an interface to a very simplistic SNP simulation routine, designed to generate large SNP datasets for population-based, case/control studies. This function is largely intended as a convenience function for generating data to prototype new methods, comparing the power of different approaches, etc, rather than producing *realistic* whole genome data. Critically, all SNPs simulated are *unlinked and in linkage equilibrium*.

## 20.1 Basic usage

The basic command to simulate a SNP data file is the --simulate option,

```
./plink --simulate wgas.sim --make-bed --out sim1
```

which takes as a parameter the name of a file (here wgas.sim) that describes the to-be-simulated data. The simulation file wgas.sim is as follows:

```
100000 null 0.00 1.00 1.00
100 disease 0.00 1.00 2.00
```

These files can have 1 or more rows, where each row has exactly five fields, as follows

```
Number of SNPs in this set Label of this set of SNPs Lower allele frequency range Uppoer allele frequency range Odds ratio for disease
```

Given this file, PLINK would generate 100,000 SNPs with no association with disease. Each SNP would have its own population allele frequency, generated as a uniform number between, in this case, 0.00 and 1.00. In addition, 100 extra SNPs will be simulated that are associated with disease (population odds ratio of 2.00).

The names of each SNP would follow from the label (which must be unquie), with a number appended, e.g.

```
null_0
null_1
null_2
...
disease_99
```

An exception is that if a set only contains a single SNP, nothing is appended to the label. This is useful in generating multiple samples from the same population, as described below.

Obviously, a uniform allele frequency range is not realistic: one could instead specify a series of bins to enrich for rarer SNPs, if so desired, to build a more realistic spectrum of allele frequencies (not that the example below is meant to be more realistic).

```
      20000
      nullA
      0.00 0.05 1.00

      10000
      nullB
      0.05 0.10 1.00

      5000
      nullC
      0.10 0.20 1.00

      10000
      nullD
      0.20 0.99 1.00
```

As well as generating the actual data, the --simulate outputs to the LOG file the following:

```
Reading simulation parameters from [ wgas.sim ] Writing SNP population frequencies to [ plink.simfreq ] Read 2 sets of SNPs, specifying 100100 SNPs in total Simulating 100 cases and 100 controls Assuming a disease prevalence of 0.01
```

The plink.simfreq file is described below. By default, 100 cases and 100 controls are generated. This can be changed with the command-line options

```
--simulate-ncases 5000
and
--simulate-ncontrols 5000
for example. Likewise, the default disease prevalence is assumed to be 0.01. This can be changed with
--simulate-prevalence 0.05
```

for example.

In the example above, the simulated data were directly saved to a binary fileset: this need not be the case. For example, any other analysis command could instead have been applied, e.g. --simulate acts just like --file or --bfile:

```
./plink --simulate wgas.sim --assoc
```

although the actual simulated data would be subsequently lost of course.

Hint This tool only generates individuals drawn from a homogeneous population, but you can easily imagine using several --simulate runs then using PLINK commands to merge the resulting files to specify more complex scenarios, e.g. representing population stratification, allelic heterogeneity, etc.

## 20.2 Resimulating a sample from the same population

The --simulate command also generates the file plink.simfreq. This records, for each SNP of the two sets, null and disease from the wgas.sim example, the *actual* allele frequency chosen for that particular SNP when simulating the data. For example,

```
1 null_0 0.1885 0.1885 1
1 null_1 0.424675 0.424675 1
1 null_2 0.12797 0.12797 1
1 null_3 0.544394 0.544394 1
1 null_4 0.938641 0.938641 1
```

. . . .

Conveniently, this information is output in the same format as the original simulation file: note how the upper and lower allele frequency range is converged to specify a particular value, i.e. the first row shows a range of 0.1885 to 0.1885, i.e. effectively forcing the allele frequency for the first SNP to be 0.1885. This can be useful, as to generate a new independent dataset from the same population as the first, you would simply use the plink.simfreq output file, as input for a new --simulate command, see below.

Putting this together, one might imagine setting up a simple screen/replicate simulation design: first we generate the original WGAS screening data

```
./plink --simulate wgas.sim --make-bed --out screen
run our association test
./plink --bfile screen --assoc
```

gawk 'NR>1 && \$9 < 1e-3 print \$2 'plink.assoc > positives

and extract a list of significant SNPs (here using the Unix gawk command, to filter on the p-value column, 9)

```
and then generate and test these same SNPs in an independent sample

./plink --simulate screen.simfreq --extract positives --assoc --out replication
```

etc. By labeling true disease SNPs and null SNPs sensibly as above, you can tell how many true positives and false positives appear at the screening and the replication stages, e.g. using Unix bash shell scripting to summarise results:

```
t=1e-3
s0='fgrep null plink.assoc | gawk ' $9 < t ' t=$t | wc -1'
s1='fgrep disease plink.assoc | gawk ' $9 < t ' t=$t | wc -1'
echo "Detected $s1 true positives and $s0 false positives in screening"
t=1e-2
s0='fgrep null replication.assoc | gawk ' $9 < t ' t=$t | wc -1'
s1='fgrep disease replication.assoc | gawk ' $9 < t ' t=$t | wc -1'
echo "Of these, $s1 true positives and $s0 false positives replicate"</pre>
```

## Chapter 21

## SNP scoring routine

PLINK provides a simple means to generate *scores* or *profiles* for individuals based on a simple allelic scoring system involving one or more SNPs. One potential use of such would be to assign a single quantitative index of genetic load, perhaps to build simple multi-SNP prediction models.

**Note** This is an advanced function intended for exploratory analyses, that is still in a *beta* development phase. If the point of this routine isn't clear to you, you probably should just ignore this entire feature.

## 21.1 Basic usage

The basic command to generate a score is the --score option, e.g.

```
./plink --bfile mydata --score myprofile.raw
```

which takes as a parameter the name of a file (here myprofile.raw) that describes the scoring system. This file has the format of one or more lines, each with exactly three fields

```
SNP ID
Reference allele
Score (numeric)
for example
SNPA A 1.95
SNPB C 2.04
SNPC T -0.98
SNPD A -0.24
```

These scores can be based on whatever you want. One choice might be the log of the odds ratio for significantly associated SNPs, for example. Then, running the command above would generate a file

```
plink.profile
```

with one individual per row and the fields:

```
FID Family ID

IID Individual ID

PHENO Phenotype for that

CNT Number of non-missing SNPs used for scoring

SCORE Total score for that individual
```

The score is simply a sum across SNPs of the number of reference alleles (0,1 or 2) at that SNP multiplied by the score for that SNP. For, example,

Genotype	A/A	G/G	A/T	0/0
# ref alleles	2	0	1	n/a

Score 2\*1.95 + 0\*2.04 + 1\*-0.98 -> 2.92

The score 2.92/3 (the average score per non-missing SNP) could then be used, e.g. as a covariate, or a predictor of disease if it is scored in a sample that is independent from the one used to generate the original scoring weights. Obviously, a score profile based on some effect size measure from a large number of SNPs will necessarily be highly correlated with the phenotype in the original sample: i.e. this in no (straightforward) way provides additional statistical evidence for associations in that sample.

## Chapter 22

# Rare copy number variant (CNV) data

This page describes some basic file formats, convenience functions and analysis options for rare copy number variant (CNV) data. Support for common copy number polymorphisms (CNPs) is described here.

Copy number variants are represented as *segments*. These segments are essentially represented and analysed in a similar manner to how PLINK handles runs of homozygosity (defined by a start and stop site on a given chromosome). Allelic (i.e. basic SNP) information is not considered here: PLINK skips the usual procedure of reading in SNP genotype data.

Here we assume that some other software package such as the Birdsuite http://www.broad.mit.edu/mpg/birdsuite/ package has previously been used to make calls for either specific copy-number variable genotypes or to identify particular genomic regions in individuals that are deletions or duplications, based on the raw data. That is, PLINK only offers functions for downstream analysis of CNV data, not for identifying CNVs in the first place, i.e. similar to the distinction between SNP genotype calling versus the subsequent analysis of those calls.

In this section, we describe the basic format for rare CNV data; the steps involved in making a MAP file and loading the data. We consider ways to filter the CNV lists by type, genomic location or frequency. We describe options for relating CNVs to phenotype, either at the level of genome-wide burden or looking for specific associations. Finally, we detail the tools for producing reports of any genes intersected by CNVs and for displaying groups of overlapping CNVs.

## 22.1 Basic support for segmental CNV data

The basic command for reading a list of segmental CN variants is

```
plink --cnv-list mydata.cnv
--fam mydata.fam
--map mydata.cnv.map
which can be abbreviated
```

(note that the map file must have the .cnv.map map extension). The CNV list file mydata.cnv has the format

```
FID Family ID
IID Individual ID
```

plink --cfile mydata

```
CHR Chromosome
BP1 Start position (base-pair)
BP2 End position (base-pair)
TYPE Type of variant, e.g. 0,1 or 3,4 copies
SCORE Confidence score associated with variant
SITES Number of probes in the variant
```

Having a header row is optional; if the first line starts with FID it will be ignored.

Note The SCORE and SITES values are not used in any direct way, except potentially as variates to filter segments on, as described below. That is, the values of these do not fundamentally impact the way analysis is performed by PLINK itself (they might alter the meaning of the results of course, e.g. if including low-confidence calls into the analysis!). In other words, if whatever software was used to generate the CNV calls does not supply some conceptually similar values, it is okay to simply put dummy codes (e.g. all 0) in these two fields.

The first few lines of a small example file is shown here:

FID	IID	CHR	BP1	BP2	TYPE	SCORE	SITE
P1	P1	4	71338469	71459318	1	27	0
P1	P1	5	31250352	32213542	1	34.2	0
P1	P1	7	53205351	53481230	3	18.2	0
P2	P2	11	86736484	87074601	1	22	0
P2	P2	14	47817280	47930190	4	55.1	0

The FAM file format is the first 6 fields of a PED file, described here; this file lists the sex, phenotype and founder status of each individual. The MAP file format is described here, although the next section how this can be automatically created using the --cnv-make-map command.

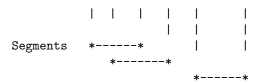
## 22.2 Creating MAP files for CNV data

Prior to any analysis, a dummy MAP first needs to be created (this step only needs to be performed once per CNV file). This PLINK-generated MAP file has dummy entries that correspond to the start and stop sites of all segments. This facilitates subsequent parsing and analysis of CNV data by PLINK. The --cnv-make-map command is used as follows:

```
plink --cnv-list mydata.cnv --cnv-make-map
which creates a file
     plink.cnv.map
which will look just like a standard MAP file but with dummy markers:
     1
              p1-51593
                           0
                                51593
     1
              p1-51598
                           0
                                51598
     1
              p1-51666
                           0
                                51666
     1
              p1-52282
                           0
                                52282
     1
              p1-69061
                           0
                                69061
```

where the marker names start with the p prefix and contain chromosome and base-position information. As an (unrealistic) example to illustrate how the mapping works, consider the following, with 3 segments, spanning "positions" 1 to 8, 4 to 12 and 16 to 23. In this case, 6 unque map positions would be created, the three start positions and the three stop positions.

```
Base 11111111111222222
Position 1234567890123456789012345
Marker # 1..2...3...4...5.....6..
```



The new MAP file would then be

```
1 p1-1 0 1
1 p1-4 0 4
1 p1-8 0 8
1 p1-12 0 12
1 p1-16 0 16
1 p1-23 0 23
```

Given such a MAP file, these three segments would then be perfectly mapped to the corresponding markers (p1-1 to p1-8, p1-4 to p1-12 and p1-16 to p1-23). The created MAP file is then specified in subsequent segmental CNV analyses (using --cnv-list) with the standard --map command (or --cfile command).

## 22.3 Loading CNV data files

Once a suitable MAP file has been created, i.e. with dummy markers that correspond to the position of every start and stop site of all segments, use the --cnv-list command again to load in the CNV segment data. As mentioned above, in addition to the basic CNV file, a MAP (previously generated) and FAM file (continuing ID and phenotype information) also need to be specified. For example.

```
plink --map plink.cnv.map --fam mydata.fam --cnv-list mydata.cnv
Alternatively, if the MAP, FAM and CNV list files all have the same root, the command
plink --cfile study1
is equivalent, i.e. it implies the following files exist
    study1.cnv
    study1.cnv.map
    study1.fam
```

By default either command will simply load in the CNV data and produce a report in the LOG file, enumerating the number of CN states in the total dataset and any filtering processes applied. For example,

```
Reading segment list (CNVs) from [ cnv1.list ]
714 of 2203 mapped as valid segments
1872 mapped to a person, of which 714 passed filters
CopyN Count
0 46
1 339
3 200
4 129
Writing segment summary to [ plink.cnv.indiv ]
```

This indicates that of 2203 total segments (i.e. should correspond to number of lines in the cnv1.list file, allowing for any header) 1872 are mapped to a person in the dataset. In other words, some of the segments in cnv1.list are for individuals not in cnv1.fam. These are simply ignored; for example, these individuals might have been filtered out of the study for other reasons, e.g. QC based on standard SNP genotypes. Of these, 714 passed the further set of filters, as described below. As described below, segments can be filtered based on genomic location, frequency, size, quality score/number of sites and type (duplication or deletion).

It will also be reported in the LOG file if some of the segments do not map to a marker in the MAP file: if this is because you've used --chr or similar commands to restrict the portion of the data examined, you can safely ignore this line; otherwise, it might mean that the appropriate MAP file wasn't created (e.g. using --cnv-make-map) for that CNV file.

By default, PLINK will create a file that summarises per individual events (after any filtering has been applied), in a file named

```
plink.cnv.indiv
```

which has the fields, one row per person, in the same order as the original FAM file:

```
FID Family ID

IID Individual ID

PHE Phenotype

NSEG Number of segments that individual has

KB Total kilobase distance spanned by segments

KBAVG Average segment size
```

PLINK will also create a file

```
plink.cnv.summary
```

that represents a count of CNVs, in cases (AFF) and controls (UNAFF) that overlap each map position.

**Note** PLINK does not check to see whether segments are overlapping for the same person or not (e.g. if a deletion and a duplication event had been specified for the same person in the same region, or if the same event is listed twice).

## 22.4 Filtering of CNV data based on CNV type

The segments read in can be filtered in a number of ways. First, one can specify to read in only either deletions (TYPE is less than 2) or duplications (TYPE is greater than 2), with the options,

```
--cnv-del
and
--cnv-dup
```

Segments can also be filtered based on a minimum size (kb), score or number of sites contributing with the following commands:

```
--cnv-kb 50
--cnv-score 3
--cnv-sites 5
```

The default minimum segment size is 20kb; none of the other filters have a default setting that would exclude anything. Also, corresponding maximum thresholds can be set:

```
--cnv-max-kb 2000
--cnv-max-score 10
--cnv-max-sites 10
```

As mentioned above, the SCORE and SITES fields are not used for any other purpose in analysis, and so if you do not have this information, can can safely enter dummy information (e.g. a value of 1 for every CNV).

The set of individuals for whom segment data are based on can be modified with the standard --keep and --remove options, to exclude people from the analysis.

## 22.5 Filtering of CNV data based on genomic location

It is possible to extract a specific set of segments that overlap with one or more regions as specified in a file, e.g. that might contain the genomic co-ordinates for genes or segmental duplications, etc. Use the command

```
--cnv-intersect regions.list
```

The file regions.list should be in the following format: one range per line, whitespace-separated:

```
CHR Chromosome code (1-22, X, Y, XY, MT, 0)
BP1 Start of range, physical position in base units
BP2 End of range, as above
MISC Any other fields after 3rd ignored
```

For example, if regions.list were

```
2 30000000 35000000 REGION1
```

2 60000000 62000000

X 10000000 20000000 Linkage hotspot

then

```
plink --cfile mydata --cnv-intersect regions.list
```

would extract all segments in mydata.cnv that at least partially span these three regions (5Mb and 2Mb on chromosome 2 and 10Mb on chromosome X), ignoring the comments or gene names. A typical type of file used with --cnv-intersect will often be a list of genes (such as available in the resources page).

Alternatively, you can use

```
--cnv-exclude regions.list
```

to filter out a specific set of segments, i.e. to remove any CNVs that overlap with one or more regions specified in the file regions.list.

Assuming the region file has consistent, unique names in the fourth field, the command

```
--cnv-subset mylist.txt
```

takes a list of region names and extracts just these from the main --cnv-intersect, --cnv-exclude (or --cnv-count, as described below) list. e.g. if mylist.txt contained

```
REGION1
REGION2
and region.list where
2 30000000 35000000 REGION1
2 60000000 62000000 GENE22
X 10000000 20000000 LinkageHotspot
```

then only the first region (chromosome 3, 30Mb to 35Mb, labelled REGION1) would be extracted, as REGION2 does not exist. The --cnv-subset command requires that the regions.list file has exactly four fields (i.e. always a unique region/gene name in the fourth field).

#### 22.5.1 Defining overlap for partially overlapping CNVs and regions

The basic intersection or exclusion commands will select all segments that are at least partially in the specified region. Alternatively, one can select only segments that have at least X percent of them in the specified region, for example

```
--cnv-overlap 0.50
```

would only include (--cnv-intersect), or exclude (--cnv-exclude), events that have at least 50% of their length spanned by the region.

There are two other variant forms of the overlap command, which change the denominator in calculating the proportion overlap:

```
--cnv-union-overlap 0.50
```

which defines overlap as the ratio of the intersection and the union, also

```
--cnv-region-overlap 0.50
```

which defines overlap as the ratio of the intersection and the length of the region (rather than the CNV). For example,

In this example, if we take each character to represent a standard length

```
Default overlap = 3 / 15
Union overlap = 3 / 19
Region overlap = 3 / 7
```

This next example illustrates how the overlap statistics can then subsequently be used to include or exclude specific CNVs: if overlap threshold were set to 0.5, then only the first of therse two CNVs would be selected by --cnv-intersect

The default setting is equivalent to setting --cnv-overlap 0 (i.e. more than 0% must overlap). Finally, the command

```
--cnv-disrupt
```

will select only CNVs that start or stop within a region specified in the region list (i.e. resulting in a partially deleted or duplicated gene or region). The normal overlap commands cannot be used in conjunction with the --cnv-disrupt defintion of whether or not a CNV overlaps a gene.

#### 22.5.2 Filtering by chromosomal co-ordinates

In addition, the standard commands for filtering chromosomal positions are still applicable, for example

```
--chr 5
or
--chr 2 --from-mb 20 --to-mb 25
```

Note that for a CNV to be included when using these filters, both the start and stop site must fall within the prespecified range (i.e. a CNV spanning from 19 to 24Mb on chromosome 2 would not be included in the above example).

## 22.6 Filtering of CNV data based on frequency

It is also possible to exclude based on the frequency of CNVs at a particular position. There are two main approaches to this: by assigning frequencies for *regions* and then applying the same routines as for the range-intersection command described above, or alternatively by assigning each CNV a single, specific count.

These commands, and the differences between them, are described more fully on this page. As well as the two basic approaches described above, one can specify different degrees of overlap when calculating frequencies, which can alter the result of frequency filtering.

The key commands and some examples are given here. To remove segments that map to regions with more than 10 segments

```
--cnv-freq-exclude-above 10
```

To remove any segments that only have at most 4 copies

```
--cnv-freq-exclude-below 5
```

To remove any segments not in regions with exactly 5 copies

```
--cnv-freq-exclude-exact 5
```

and correspondingly to include only segments in regions with exactly 5 copies

```
--cnv-freq-include-exact 5
```

As with the earlier range intersection commands, the definition of *intersection* can be *soft*, specified with the --cnv-overlap option. In most cases here, one would probably want to allow for a soft filtering, e.g. with --cnv-overlap 0.5 for example.

For example, given the following segments, and counts below



then --cnv-freq-exclude-above 1 would remove all three segments if --cnv-overlap 0 (the default) were set. This is because each CNV has at least some part of it that intersects with a region that contains more than 1 CNV. However, if --cnv-overlap were instead set to 0.5, for example, then only the top segment would be removed (as the other two segments have more than 50% of their length outside of a region with more than 1 segment). If the overlap were set higher still, then in this example no CNVs would be removed by the command --cnv-freq-exclude-above 1.

**NOTE** Because multiple CNVs at the same region will not all exactly overlap, and may be spanned by distinct larger events, or contain smaller events, in other individuals, then requesting that you include only CNVs with exactly five copies for example (--cnv-freq-include-exact 5) does **not** mean that at all positions in the genome you will always see either 0 or 5 copies. Rather, the selection process works exactly as specified above. Please see this page for further details.

#### 22.6.1 Alternative frequency filtering specification

The alternate approach is invoked with the command

```
--cnv-freq-method2 0.5
```

where the value following it represents an overlap parameter (there is no need to specify the --cnv-overlap command directly when using --cnv-freq-method2). Based on this overlap, PLINK will assign a specific count to each CNV that represents the number of CNVs that overlap it (including itself) based on a union intersection overlap definition with the specified proportion parameter, between that CNV and all CNVs.

This approach is illustrated in the page, that gives more details on the frequency filtering commands including a comparison to the region-based approach to filtering, described above.

If the --cnv-freq-method2 command is used, then the other frequency filtering commands will use the CNV-based counts to include of exclude CNVs, for example

```
If --cnv-write (see below) is specified with --cnv-freq-method2, then the additional command --cnv-write-freq
```

will add a field FREQ to the plink.cnv file generated that shows the frequency for each CNV. Also, the --cnv-seglist command (see below) can be modified with --cnv-write-freq (to report the frequency as a number at the start and stop of each CNV instead of the usual codes).

#### 22.6.2 Miscellaneous commands frequency filtering commands

To keep only segments that are unique to either cases or to controls

```
--cnv-unique
```

This can be used in conjunction with other frequency filter commands. To drop individuals from the file who do not have at least one segment after filtering, add the flag

```
--cnv-drop-no-segment
```

This can make the plink.cnv.indiv summary files easy to browse, for example.

## 22.7 Association analysis of segmental CNV data

To perform a set of global test of CNV burden in cases versus controls, add the

```
--cnv-indiv-perm
```

option as well as

```
--mperm 10000
```

for example (i.e. permutation is required). By default, this reports on four tests, which use these metrics to calculate burden in both cases and controls

```
RATE Number of segments
PROP Proportion of sample with one or more segment
TOTKB Total kb length spanned
AVGKB Average segment size
```

Tests are based (1-sided) on comparing these metrics in cases versus controls, evaluated by permutation. If a list of regions is supplied in a file, e.g. gene.list and the command

```
--cnv-count gene.list
```

then an extra test is added

```
GRATE Number of regions/genes spanned by CNVs
GPROP Number of CNVs with at least one gene
GRICH Number of regions/genes per total CNV kb
```

These tests respect all the normal filtering commands, with the exception that --cnv-intersect and --cnv-exclude cannot be used if --cnv-count is also being used.

The mean metrics in cases and controls are reported in the file

```
plink.cnv.grp.summary
```

when the **--cnv-indiv-perm** command is used. For example: this gives the number of events (N) in cases and controls, the rate per person, the proportion of cases/controls to have at least one event, the total distance spanned per person and the average event size per person.

TEST	GRP	AFF	UNAFF
N	ALL	528	362
RATE	ALL	0.1557	0.1138
PROP	ALL	0.1309	0.1041
TOTKB	ALL	290.8	265.4
AVGKB	ALL	249.8	243.3

As usual, if the --within command is added and a cluster file specified, then any permutations are performed within cluster. In this case, the statistics displayed in the plink.cnv.grp.summary file are also split out by the strata as well as presented in total (as indicated by the GRP field).

## 22.8 Association mapping with segmental CNV data

To perform a simple permutation-based test of association of segmental CNV data for case/control phenotypes, add the option

```
--mperm 50000
```

to perform, for example, 50,000 null permutations to generate empirical p-values. The results are saved in the file

```
plink.cnv.summary.mperm
```

This is a standard empirical p-value file: EMP1 and EMP2 represent pointwise and genome-wide corrected p-values, respectively. Both tests are 1-sided by default.

You can consult the corresponding

```
plink.cnv.summary
```

that is also generated for details of the association: this file has the fields

```
CHR Chromosome code
```

SNP SNP identifier (dummy SNP, see below)

BP Base-pair position

AFF Number of affected individuals with a segment at this position

UNAFF Number of unaffected individuals

To instead perform a 2-sided test (i.e. allowing that events might be more common in controls) add the flag

```
--cnv-2sided
```

To perform an analysis in which the total number of events within a sliding window is compared between cases and controls (rather than the number overlapping a single position) add the flag

```
--cnv-test-window 50
```

where the parameter is the kb window either side of the test position. As before, the association results are reported per marker, but now the counts indicate the total number of segments that overlap any of the 100 kb window surrounding the test position (+/- 50 kb), rather than just the test position itself. Significance is evaluated by permutation as before.

## 22.9 Writing new CNV lists

Given a set of filters applied, you can output as a new CNV file the filtered subset, with the command

```
--cnv-write
```

For example, to make a new file using only deletions over 200kb but not more than 1000kb, with a quality score of 10 or more, use the command

```
plink --cfile cnv1
--cnv-del
--cnv-kb 200
--cnv-max-kb 1000
--cnv-score 10
--cnv-write
--out hiqual-large-deletions
```

```
which will generate two new files
```

```
hiqual-large-deletions.cnv
hiqual-large-deletions.fam
```

To obtan a corresponding MAP file, so that you can subsequently use

```
\hbox{\it ---cfile hiqual-large-deletions}
```

give the command

plink --cnv-list hiqual-large-deletions.cnv --cnv-make-map --out hiqual-large-deletions (although note that this will overwrite the LOG file generated by the --cnv-write command).

#### 22.9.1 Creating UCSC browser CNV tracks

As opposed to listing CNVs in PLINK format with --cnv-write, the command --cnv-track will generate a UCSC-friendly BED file (note: this is distinct from a PLINK binary PED file) that can be uploaded to their browser for convenient viewing.

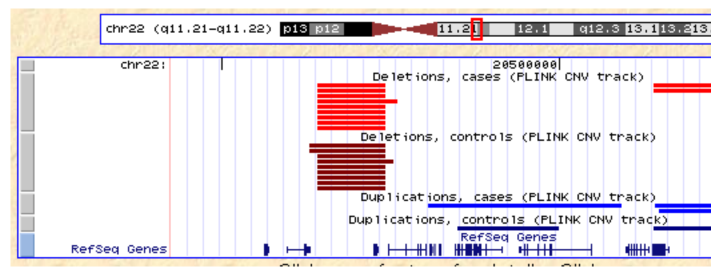
```
plink --cfile mydata --cnv-track --out mycnvs
which generates a file
    plink.cnv.bed
```

The filtering commands described above can be combined with this option.

By using the *Manage custom tracks* option on the UCSC genome browser http://genome.ucsc.edu/cgi-bin/hgGateway, one can easily visualise the CNV data, along side other genomic features. For example, the file (IID and SCORE, SITES information is omitted for clarity)

FID IID	CHR	BP1	BP2	TYPE	SCORE	SITES
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20129453	20241877	1		
	22	20140609	20241877	1		
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20639721	20793965	1		
	22	20639721	20765489	1		
	22	20305076	20591362	3		
	22	20646213	20756780	3		
	22	20140420	20259122	1		
	22	20639866	20787533	3		
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20348901	20498220	3		
	22	20140420	20241877	1		
	22	20140420	20241877	1		
	22	20639643	20793173	3		
	22	20140420	20241877	1		
	22	20141114	20241877	1		
	22	20140420	20254215	1		
	22	20140420	20241877	1		
	22	20129130	20241877	1		

is rendered



Note that the CNVs are split by deletion versus duplication (red versus blue) and case versus control (light versus dark).

Additionally, a poor-man's version of this plot can be obtained with the command

```
--cnv-seglist
which produces a file
plink.cnv.seglist
```

described below.

which, for the CNV list above, can be seen here. Deletions and duplications are represented by + and - symbols at the start of each CNV; case and control status is represented as A and U.

symbols at the start of each CNV; case and control status is represented as A and U.

Finally, it is also possible to report CNVs annotated by the regions or genes they span (see --cnv-verbose-report-region

## 22.10 Listing intersected genes and regions

```
With the --cnv-intersect (or --cnv-exclude) command, you can add the flag
--cnv-report-regions
which will create a file
plink.reg
```

listing only the regions that intersect (or do not intersect) with any of the CNVs (given the filtering and overlap commands that might also be specified). For example, to obtain a list of genes that are intersected by a rare case singleton deletions over 500kb (i.e. event seen only once)

#### Alternatively, the command

#### --cnv-verbose-report-regions

produces a verbose form of plink.reg, which does not just list the regions or genes intersected but lists the specific segmental CNVs also. This can be used in conjunction with, for example,

```
--cnv-subset sczgenes.txt
```

in order to produce reports on specific genes of interest. For example if sczgenes.txt contained

DISC1

COMT

then

plink --cfile mydata

- --cnv-verbose-report-regions
- --cnv-intersect glist-hg17
- --cnv-subset sczgenes.txt

would produce a file

plink.reg

that might contain something like the following report

O		0		0 1				
RANGE (+/- Ok	b )	[ 1 2	28069	295 228483751	DISC1 ]			
FID	IID	PHE	CHR	BP1	BP2	TYPE	SCORE	SITES
PT-2FET	1	1	1	228007236	228115017	DUP	31.36	59
PT-108L	1	2	1	228013674	228119443	DUP	13.15	24
PT-1036	1	2	1	228013674	228122852	DUP	16.06	25
PT-1PJD	1	2	1	228019276	228119443	DUP	22.21	23
PT-1S24	1	1	1	228019276	228119443	DUP	17.78	23
PT-2412	1	1	1	228019276	228122852	DUP	21	24
PT-28HG	1	1	1	228019304	228120053	DUP	34.43	56
PT-27R2	1	2	1	228019304	228120053	DUP	39.86	56
PT-2M7W	1	1	1	228019304	228120053	DUP	35.87	56
PT-2C1W	1	2	1	228019304	228120053	DUP	58.52	56
PT-27TB	1	2	1	228019304	228120053	DUP	35.47	56
PT-295W	1	1	1	228019304	228120053	DUP	34.32	56
PT-2M1L	1	2	1	228019304	228120053	DUP	17.69	56
PT-2DNH	1	2	1	228019304	228120053	DUP	42.2	56
PT-2D5K	1	2	1	228019304	228120053	DUP	28.67	56
PT-2CJS	1	1	1	228019304	228120053	DUP	51.11	56
PT-2C8H	1	2	1	228019304	228120053	DUP	29.98	56
RANGE (+/- Ok	b )	[ 22	18303	8862 18331082	COMT ]			
FID	IID	PHE	CHR	BP1	BP2	TYPE	SCORE	SITES
PT-285C	1	1	22	17107473	18686471	DEL	374.01	463
PT-1PGT	1	2	22	17237164	18683947	DEL	427.59	449
PT-22Q9	1	1	22	17237164	19787083	DEL	237.745	323
PT-229U	1	2	22	17237164	19921619	DEL	412.48	342
PT-2M6E	1	1	22	17250982	18686471	DEL	1051.29	919
PT-2BXV	1	2	22	17250982	18772410	DEL	1087.02	922
PT-2LZQ	1	1	22	17250982	19790389	DUP	591.505	810
PT-22LI	1	1	22	17259415	18684680	DEL	276.04	221
PT-232G	1	2	22	17259415	19787083	DEL	286.57	338
PT-1NQ3	1	1	22	17259415	19787083	DEL	168.843	218

PT-22RR	1	2	22	17259415	19787083	DEL	470.42	342
PT-101Z	1	1	22	17259415	19921619	DEL	434.3	341

That is, this is a list of any CNV that at least partially overlaps these two genes. The exact behavior can be modified with flags such as --cnv-del, --cnv-wb, --cnv-overlap, --cnv-disrupt, --filter-cases, etc.

## 22.11 Reporting sets of overlapping segmental CNVs

Finally, there are two option to group or report sets of segments that span a particular position. In the first case, use the option

#### --segment-group

which takes all segments in a given region (whole genome unless otherwise specified) and forms "pools" of overlapping segments. Several pools of overlapping segments will be created; these will be listed in order of decreasing size (number of segments); note that the same segment can appear in multiple pools (e.g. if A overlaps with C, and B overlaps with C, but A and B do not overlap). The pools give information as described below.

The more restricted form of this command forms a single pool of all segments that overlap a particular position, which takes a single parameter of a marker name; typically these will be the dummy pos\* markers created by the --cnv-make-map command.

#### --segment-spanning pos119

In this case, for some made-up data, we see from the plink.cnv.summary file that there are 8 cases and 6 controls with a segment spanning a particular position, pos586

UNAFF	AFF	BP	SNP	CHR
6	8	16631570	pos586	1

In this case, there is unsurprisingly no association between segmental CNVs and disease: for example, the corresponding position in the plink.cnv.summary.mperm file shows an empirical p-value of 0.35, but of p=1 if adjusted for multiple testing (EMP2)

EMP2	EMP1	STAT	SNP	CHR
• • •	• • •	• • •	• • •	• • •
1	0.351324	0.419408	pos586	1

Naturally, one would usually be more interested in following up significantly associated regions of course... Nonetheless, if so desired we can see which segments (given any of the filtering specified) are spanning this position, with --segment-spanning, which gives the following:

POOL	FID	IID	PHE	CHR	BP1	BP2	KB	TYPE	SCORE
S1	PT-2378	PT-2378	2	12	16631570	16751087	119.517	DEL	10.23
S1	PT-268D	PT-268D	2	12	16631494	16732162	100.668	DEL	9.3
S1	PT-2M80	PT-2M80	1	12	16631441	16751082	119.641	DEL	31.23
S1	PT-2FZ9	PT-2FZ9	2	12	16631436	16751045	119.609	DEL	15.2
S1	PT-287D	PT-287D	1	12	16616579	17183201	566.622	DUP	200.3
S1	PT-2C91	PT-2C91	2	12	16616579	16751045	134.466	DEL	14.3
S1	PT-28A8	PT-28A8	1	12	16616579	16751045	134.466	DEL	8.3
S1	PT-2FPB	PT-2FPB	1	12	16616579	16714372	97.793	DEL	11.1
S1	PT-28IG	PT-28IG	2	12	16616579	16708856	92.277	DEL	10.3
S1	PT-2E5N	PT-2E5N	2	12	16614664	16715703	101.039	DEL	9.87
S1	PT-2FVL	PT-2FVL	1	12	16614664	16751045	136.381	DEL	10.67
S1	PT-2DYE	PT-2DYE	2	12	16614664	16715489	100.825	DEL	11.82

S1	PT-264I	PT-264I	2	12	16614664	16751045	136.381	DEL	14.2
S1	PT-25WZ	PT-25WZ	1	12	16591338	16715767	124.429	DEL	14.7
S1	CON	14	8:6	12	16631570	16708856	77.286	NA	NA
S1	UNION	14	8:6	12	16591338	17183201	591.863	NA	NA

For CNV data (in contrast to shared segments based on homozygosity or IBD sharing) the extra fields of TYPE (deletion or duplication) and SCORE (some metric of quality/confidence of CNV call) are also presented.

Here we see the 14 segments listed, 8 cases and 6 controls. The CON and UNION lines at the end of the pool give the consensus region (i.e. shared by all segments) and the total distance spanned by all. The PHE field gives the phenotype for each individual.

Note that the way in which the dummy markers are selected will effectively mean that every possibly unique position, in terms of counts of segments, is evaluated. The actual base pair regions of any dummy marker is itself probably not of interest: given a significant (set of) SNPs, the strategy would be to select any one and generate the corresponding pool to see what and where the association maps to.

# 22.12 Illustration of the different CNV frequency filtering commands

To illustrate both the region-based and CNV-based methods of frequency filtering, consider this example CNV file, with 18 individuals and 18 CNVs, which contains a complex set of partially overlapping events:

```
FID IID CHR BP1 BP2 TYPE SCORE SITES
     1 10000
              20000 1 10 10
      1 10000
               20000 1 10 10
3
         9000
               21000 1 10 10
      1 10000
               32000 1 10 10
5
      1 20000
               31000 1 10 10
         5000
               50000 1 10 10
      1 40000
               51000 1 10 10
      1 44000
               48000 1 10 10
               46000 1 10 10
      1 42000
      1 41000
               49000 1 10 10
      1 39000
               48000 1 10 10
      1 38000
               52000 1 10 10
      1 80000
               85000 1 10 10
     1 90000
               99000 1 10 10
14 1
     1 91000
               99000 1 10 10
     1 89000
               98000 1 10 10
     1 90000
               99000 1 10 10
     1 90000
              99000 1 10 10
```

The files are available for you to download and play with: test1.cnv, test1.cnv.map and test1.fam. The command

```
./plink --cfile test1 --cnv-seglist
```

gives the following output in plink.cnv.seglist, but with the rightmost column being the AFF CNV count field from plink.cnv.summary (i.e. all 18 individuals are coded as cases; this number represents the number of CNVs spanning that particular MAP position):

		AFF
p1-5000	+	1
p1-9000	+	2
p1-10000	++  +	5
p1-20000	AA +	6

p1-20001 p1-21000 p1-21001 p1-31000 p1-31001 p1-32000 p1-32001 p1-38000 p1-39000 p1-40000	 A       A     A   A   +   +	4 4 3 3 2 2 2 1 2 3 4
p1-44000 p1-46000	A	7 7
p1-46001	111111	6
p1-48000	A   A	6
p1-48001		4
p1-49000	A	4
p1-49001	111	3
p1-50000		3
p1-50001	ii	2
p1-51000	A	2
p1-51001	İ	1
p1-52000	A	1
p1-52001	0	
p1-80000	+ 1	
p1-85000	A 1	
p1-85001	0	
p1-89000	+ 1	
p1-90000	+++	4
p1-91000	+	5
p1-98000	A	5
p1-98001	1 111	4
p1-99000	A AAA	4
p1-99001	0	

#### 22.12.1 Region-based, or locus-based, frequency filtering (default)

**NOTE** These commands are intended to illustrate how the filtering works, rather than provide useful examples of how to analyse data in practice.

For example, the command

```
plink --cfile test1 --cnv-seglist --cnv-freq-exclude-above 4 --cnv-overlap 1 will remove CNVs that completely span regions with 5 or more CNVs:
```

```
p1-5000 + | p1-9000 + | p1-10000 | + | p1-20000 | + | | p1-20001 | | | | | p1-21000 | A| | | p1-21001 | | | |
```

```
Al I
p1-31000
                  1 1
p1-31001
p1-32000
                  A |
                    p1-32001
p1-38000
                   +|
                + ||
p1-39000
                |+||
p1-40000
p1-41000
               +||||
p1-42000
               \Pi\Pi\Pi\Pi
p1-44000
               \Pi\Pi\Pi\Pi
p1-46000
                | | | | | |
p1-46001
               11111
p1-48000
               | A | | |
p1-48001
               | | | | |
p1-49000
               A |||
p1-49001
                  | | |
p1-50000
                  | | A
                  \prod
p1-50001
p1-51000
                  Αl
p1-51001
                   p1-52000
                   Α
p1-52001
p1-80000
               +
p1-85000
p1-85001
p1-89000
p1-90000
                 |+++
p1-91000
               +||||
p1-98000
                | A | | |
               1 111
p1-98001
               A AAA
p1-99000
p1-99001
```

The command

```
plink --cfile test1 --cnv-seglist --cnv-freq-exclude-above 6 --cnv-overlap 0
```

will remove CNVs that completely even partially overlap regions with 7 or more CNVs: ( this removes 7  $\,$  CNVs in total)

```
p1-5000
p1-9000
              ++| +
p1-10000
p1-20000
              AA|+|
p1-20001
                \Pi
                A | |
p1-21000
p1-21001
                 II
                 Αl
p1-31000
p1-31001
                  p1-32000
                  Α
p1-32001
p1-38000
p1-39000
p1-40000
```

p1-41000 p1-42000 p1-44000 p1-46000 p1-46001 p1-48000 p1-48001 p1-49000 p1-49001 p1-50000 p1-50001 p1-51000 p1-51001 p1-52000 p1-52001 p1-80000 p1-85000 Α p1-85001 p1-89000 p1-90000 |+++ +|||| p1-91000 p1-98000 | A | | | 1 111 p1-98001 p1-99000 A AAA p1-99001

#### 22.12.2 Alternative frequency filtering approach

The standard approach to frequency filtering considers the frequency of CNVs at each particular genomic location, defining *regions* with a particular number of CNVs spanning it; CNVs are subsequently filtered based on the extent to which each individual CNV overlaps or does not overlap with these regions.

An alternative approach (invoked with the --cnv-freq-method2 flag) is to define frequency as being a property of a particular CNV rather than of a region, which is perhaps more intuitive. Here we count for each CNV how many other CNVs overlap it. The overlap definition here is forced to be a union overlap that isn't allowed to be disruptive (--cnv-disrupt), in order to ensure symmetry (i.e. if A overlaps B, then B must overlap A). The frequency filtering is then based on these counts.

Below are the frequency counts for each CNV, given different values for the overlap parameter specified in the --cnv-freq-method2 command:

	cnv-freq-method2 0			-	cnv-freq-method2 0.5				0.5	cnv-freq-method2 1							
					 						- 						
p1-5000				12	i					1	i					1	
p1-9000		6		12	- 1			3		1	- 1		1	L		1	
p1-10000	6 6	6	6	12	-	3	3	3	2	1	1	2	2 1	L	1	1	
p1-20000	6 6	6 6	6	12	-	3	3	3	2 2	1	1	2	2 1	1 1	1	1	
p1-20001		6 6	6	12	-			3	2 2	1	1		1	1 1	1	1	
p1-21000		6 6	6	12	- 1			3	2 2	1	1		1	1 1	1	1	
p1-21001		6	6	12	-				2 2	1	1			1	1	1	
p1-31000		6	6	12	- 1				2 2	1	1			1	1	1	
p1-31001			6	12	- 1				2	1	1				1	1	
p1-32000			6	12	1				2	1	1				1	1	

p1-32001	12	1	1
p1-38000	7 12	4 1	1 1
p1-39000	7 7 12	4 4 1	1 1 1
p1-40000	7 7 7 12	4441	1 1 1 1
p1-41000	7 7 7 7 12	6 4 4 4 1	1 1 1 1 1
p1-42000	7 777712	2 64441	1 11111
p1-44000	7 7 7 7 7 7 12	2 2 6 4 4 4 1	111111
p1-46000	7 7 7 7 7 7 12	2 2 6 4 4 4 1	111111
p1-46001	7 7 7 7 7 12	264441	11111
p1-48000	7 7 7 7 7 12	2 6 4 4 4 1	1 1 1 1 1 1
p1-48001	7 7 7 12	6 441	1 111
p1-49000	7 7 7 12	6 441	1 111
p1-49001	7 7 12	4 4 1	1 1 1
p1-50000	7 7 12	4 4 1	1 1 1
p1-50001	7 7	1 4 4	1 1
p1-51000	7 7	1 4 4	1 1
p1-51001	7	1 4	1
p1-52000	7	1 4	1
p1-52001		I	I
p1-80000	1	1	1
p1-85000	1	1	1
p1-85001		I	I
p1-89000	5	l 5	1
p1-90000	5 5 5 5	5555	1 3 3 3
p1-91000	5 5 5 5 5	55555	1 1 3 3 3
p1-98000	5 5 5 5 5	55555	1 1 3 3 3
p1-98001	5 5 5 5	5 5 5 5	1 3 3 3
p1-99000	5 5 5 5	5 5 5 5	1 3 3 3
p1-99001		1	I

Any additional commands such as <code>--cnv-freq-exclude-above 5</code> would work in a straightforward manner based on these counts. For example

```
plink --cfile test1
        --cnv-freq-method2 0
        --cnv-freq-include-exact 5
        --cnv-write
        --cnv-write-freq
will include just the group of segments starting after position 89000
     Filtering segments based on frequencies
     Will remove 13 CNVs based on frequency (after other filters)
     18 mapped to a person, of which 18 passed filters
     5 of 18 mapped as valid segments \,
e.g. as shown in file
     plink.cnv
which contains
    FID IID CHR
                            BP1
                                          BP2
                                                TYPE
                                                             SCORE
                                                                       SITES
                                                                                  FREQ
```

14	1	1	90000	99000	1	10	10	5
15	1	1	91000	99000	1	10	10	5
16	1	1	89000	98000	1	10	10	5
17	1	1	90000	99000	1	10	10	5
18	1	1	90000	99000	1	10	10	5

#### 22.12.3 In summary

For a complex set of partially overlapping CNVs, any attempt to collapse CNVs into discrete groups or counts will inevitably be somewhat artificial. Nonetheless, the commands presented here provide a range of options, to either strictly or loosely filter as desired. This made-up example dataset is particularly complex – in most real cases, these frequency filters will yield sensible results.

To select CNVs below some overall frequency (e.g. 1%, which if there are 1000 individuals would mean 10 events) the option

```
--cnv-freq-exclude-above 10
```

would be a good default.

To select strictly defined singleton CNVs (those seen only once in a dataset), use

<sup>--</sup>cnv-overlap 0.5

<sup>--</sup>cnv-freq-exclude-above 1

## Chapter 23

# Common copy number polymorphism (CNP) data

This page describes some basic file formats, convenience functions and analysis options for common copy number polymorphism (CNP) data. Support for rare copy number variant (CNV) data is described here.

Common copy number variation is represented for specific SNP genotypes, for example, allowing A, AAB or AABB calls (being copy number 1,3 and 4 respectively) as well as the canonical AA, AB and BB genotypes. These formats are specified via the "generic variant" (--gfile) option.

Here we assume that some other software package such as the Birdsuite http://www.broad.mit.edu/mpg/birdsuite/ package has previously been used to make calls for either specific copy-number variable genotypes or to identify particular genomic regions in individuals that are deletions or duplications, based on the raw data. That is, PLINK only offers functions for downstream analysis of CNV data, not for identifying CNVs in the first place, i.e. similar to the distinction between SNP genotype calling versus the subsequent analysis of those calls.

## 23.1 Format for common CNVs (generic variant format)

For common CNVs, that might also have meaningful allelic/SNP variation, it can be desirable to represent and analyse these not as segments. The rest of the page considers non-segmental specification of CNVs: that is, copy-number variable specific genotype calls, such as A or AAB.

Such data are represented with the *generic variant* file format, and read into PLINK with the command:

```
plink --gfile mydata
```

where three files are assumed to exist

```
mydata.fam (describes individuals, as usual)
mydata.map (describes variants, as usual)
mydaya.gvar (new file format)
```

The .gvar file is in long-format: always with 7 fields, one row per genotype (note that the reference to the first and second parents above does not imply that paternal or maternal origin should be known or is used)

```
FID Family ID

IID Individual ID (i.e. person should appear in .fam file)

NAME Variant name (should appear in .map file)

ALLELE1 Code for allele from first parent

DOSAGE1 Copy number for first allele

ALLELE2 Code for allele from second parent
```

#### DOSAGE2 Copy number for second allele

Some example of using this format to represent different genotypes are shown here:

```
1 1 var1
           A 1
                C 1
                       -> normal het
1 1 var2
           A 2
                C 1
                       -> AAC genotype
           0 1
                0 1
                       -> missing individual
1 1 var3
1 1
          0 0 0 0
                       -> homozygous deletion
    var4
          4 1 7 1
                       \rightarrow e.g. 4/7 genotype
1 1 var5
2 1 var5
          4 1 8 1
                       -> e.g. 4/8 genotype
1 1 var6 A 0.95 C 1.05 -> expected allele dosage (e.g. from imputation)
```

As currently implemented, all the codings below would be equivalent, i.e. specifying an AA homozygote:

```
1 1 var7 A 1 A 1
2 1 var7 A 0 A 2
3 1 var7 A 2 A 0
4 1 var7 X 0 A 2
5 1 var7 0 0 A 2
```

That is, for a missing (null) genotype, ALLELE1 and ALLELE2 should both be set to 0, and by convention, DOSAGE1 and DOSAGE2 should be 1 (indicating a 0 0 genotype). But if a DOSAGE value is 0, then the value of the corresponding ALLELE column does not matter. Thus, genotypes can have DOSAGE >= 1 for one allele, and DOSAGE for the other allele: A 0 B 3 means 3 copies of allele B and no copies of A; X 0 B 3 means the same thing because the X is ignored when DOSAGE=0.

When loading this kind of file, PLINK will parse allelic and copy number variation; currently by default it looks for integer dosage calls in this part of the process. There are currently no functions implemented yet for fractional counts, but the datatype exists.

Alleles and CNVs are then appropriately counted. PLINK assesses and records for each variant whether there is allelic and/or copy number variation, and this influences downstream analysis. Currently variation is defined as at least one individual varying, but in the future thresholds will be added (e.g. to treat a site of a CNV only if, say, 1% of all individuals have a non-canonical copy number).

The basic summary output is also in "long format": in the future this will be expanded and reformated, e.g. to include specific allelic/CNV frequencies or counts; stratification by phenotype, etc. This summary file is called

plink.gvar.summary
and always contains three columns, as illustrated here

NAME	FIELD	VALUE
var1	CHR	1
var1	BP	1
var1	CNV	yes
var1	ALLELIC	yes
var1	GCOUNT	1000
var1	В	0.6031
var1	Α	0.3969
var1	[2]	0.56
var1	[3]	0.378
var1	[4]	0.062
var1	B/B	30:38
var1	BB/B	66:60
var1	BB/BB	42:20
var1	A/B	142:101
var1	A/BB	161:91
var1	A/A	162:87

The CN counts are always in [x] to distingush from allele codes, if they are also numeric. e.g. in this example, 37.5% of sample have the deletion for example. There can be more than 2 CN states for a given variant.

If the trait is binary, then the counts for copy-number specific genotypes (e.g. A/BB) will be given separately for cases and controls, separated by a colon.

## 23.2 Association models for combined SNP and common CNV data

PLINK has implemented the following regression models (logistic or linear) currently applicable to biallelic SNPs residing within CNPs:

$$Y \sim b0 + b1.(A+B) + b2.(A-B)$$

When an association test is performed, extra lines will be appended to the plink.gvar.summary file

var1	B(SNP)	-0.05955
var1	P(SNP)	0.09085
var1	B(CNP)	0.09314
var1	P(CNP)	0.3809
var1	B(CNP SNP)	0.5638
var1	P(CNP SNP)	0.0006768
var1	B(SNP CNP)	-0.2042
var1	P(SNP CNP)	0.0002242
var1	P(SNP&CNP)	0.0007413

Covariates can be added with --covar as with --linear or --logistic. The coefficients and p-values for the SNP and CNP will reflect this, although the specific coefficients and p-values for the covariates themselves are not shown in the output.

This section is not finished - more details will be added online presently.

## Chapter 24

## Resources available for download

This page contains links to several freely-available resources, mostly generated by other individuals. All these resources are provided "as is", without any guarantees regarding their correctness or utility.

### 24.1 The Phase 2 HapMap as a PLINK fileset

The HapMap http://www.hapmap.org genotype data (the latest is release 23) are available here as PLINK binary filesets. The SNPs are currently coded according NCBI build 36 coordinates on the forward strand. Several versions are available here: the entire dataset (a single, very large fileset: you will need a computer with at least 2Gb of RAM to load this file).

The *filtered* SNP set refers to a list of SNPs that have MAF greater than 0.01 and genotyping rate greater than 0.95 in the 60 CEU founders. This fileset is probably a good starting place for imputation in samples of European descent. Filtered versions of the other HapMap panels will be made available shortly.

Description	riie size	riie name
Entire HapMap (release 23, 270 individuals, 3.96 million SNPs)	110M	hapmap_r23.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap_r23.zip
CEU founders (release 23, 60 individuals, 3.96 million SNPs)	49M	hapmap-ceu-all-r23.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap-ceu-all-r23.zip
CEU founders (release 23, 60 individuals, filtered 2.2 million SNPs)	29M	hapmap-ceu-r23.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap-ceu-r23.zip
CEU founders (release 23, as above, files split by chromosome, 1-22 and X)	29M	$hapmap\text{-ceu-by-chr-r} 23.zip\ \text{http://pngu.mgh.harvard.edu/}^{\sim} purcell/\text{dist/hapmap-ceu-by-chr-r} 23.zip\ \text{http://pngu.mgh.harvard.edu/}^{\sim} purcell/\text{dist/hapmap-ceu-by-ceu-by-chr-r} 23.zip\ \text{http://pngu.mgh.harvard.edu/}^{\sim} purcell/\text{dist/hapmap-ceu-by-ceu-by-chr-r}^{\sim} purcell/\text{dist/hapmap-ceu-by-ceu-by-chr-r}^{\sim} purcell/\text{dist/hapmap-ceu-by-ceu-by-chr-r}^{\sim} purcell/dist/hapmap-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-ceu-by-c$
Entire HapMap (release 22, 270 individuals, 3.96 million SNPs)	110M	hapmap_r22.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap_r22.zip
CEU founders (release 22, 60 individuals, 3.96 million SNPs)	49M	hapmap-ceu-all.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap-ceu-all.zip
CEU founders (release 22, 60 individuals, filtered 2.2 million SNPs)	$_{29M}$	hapmap-ceu.zip http://pngu.mgh.harvard.edu/~purcell/dist/hapmap-ceu.zip
CEU founders (release 22, as above, files split by chromosome, 1-22 and X)	$_{29M}$	${\tt hapmap-ceu-by-chr.zip~http://pngu.mgh.harvard.edu/} {\sim} {\tt purcell/dist/hapmap-ceu-by-chr.zip}$

## 24.2 Teaching materials and example dataset

A tutorial can be downloaded from here; the material is similar to the online tutorial but slightly more involved. As it currently stands, it is designed to first use gPLINK to perform a set of basic tests and QC procedures and then move to standard PLINK for more in-depth analysis.

It is designed to work on a standard modern laptop computer or equivalent desktop. It was written for vesion 1.02 of PLINK, but should remain compatible with future releases.

Description	File size	File name	37 6 1 6
ZIP archive containing data	15M	example.zip http://pngu.mgh.harvard.edu/~purcell/dist/exam	You are feel free to
ZIP archive containing teaching materials	1.3M	teaching.zip http://pngu.mgh.harvard.edu/~purcell/dist/teac	hing.zip
use, modify or distribute these	files in	any way you wish, although giving m	e appropriate credit for the
materials would be appreciated.			

The example.zip archive contains

wgas1.ped	Whole-genome SNP data example PED file
wgas1.map	Corresponding MAP file
extra.ped	Follow-up genotyping for a particular region
extra.map	Corresponding MAP file
pop.cov	Population membership variable
command-list.txt	List of all commands for 2nd part of practical

The teaching.zip archive contains a PowerPoint and a Word file:

```
practical-1-slides.ppt
practical-2-notes.doc
```

These two files cover the first and second half of the tutorial respectively. The second document assumes the first half has already been completed (but also contains some introductory remarks concerning the data). I will probably update the Word document to also include the early commands covered in the PowerPoint/gPLINK part (i.e. so that the entire practical can be performed from the command line rather than using gPLINK). The list of commands (command-list.txt) is included so that people can cut-and-paste commands in, rather than type. If using DOS, it is a good idea to first increase the window width (right click on header on DOS window, Properties, Layout and increase buffer and window width to around 120 characters).

Everything should be fairly self-explantory after looking through the PowerPoint file and Word document.

#### 24.3 Multimarker test lists

These files, generated by Itsik Pe'er and others, facilitate the 'multi-marker predictor' approach to association testing, as described in the manuscript:

```
Pe'er I, de Bakker PI, Maller J, Yelensky R, Altshuler D & Daly MJ (2006) Evaluating and improving power in whole-genome association studies using fixed marker sets. Nat Genet, 38(6): 605-6.
```

They are PLINK-formatted lists of multimarker tests selected for Affymetrix 500K and Illumina whole genome products, based on consideration of the CEU Phase 2 HapMap (at r-squared=0.8 threshold). One should download the appropriate file and run with the --hap option (after ensuring that any strand issues have been resolved).

- Affymetrix.GeneChip.500k.both.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Affymetrix.GeneChip.500k.both.CEU.0.8.tests.zip
- Illumina.HumanHap.300k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.300k.CEU.0.8.tests.zip
- Illumina.HumanHap.550k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.550k.CEU.0.8.tests.zip
- Illumina.HumanHap.650k.CEU.0.8.tests.zip http://pngu.mgh.harvard.edu/~purcell/dist/mmtests/Illumina.HumanHap.650k.CEU.0.8.tests.zip

**Note** These haplotypes are specified in terms of the +ve (positive) strand relative to the HapMap. You might need to reformat your data prior to using these files (using the --flip command, for instance) before you can use them.

**Note** These tables list all tags for every common HapMap SNP, at the given r-squared threshold. The same haplotype may therefore appear multiple times (i.e. if it tags more than 1 SNP).

Note These tables obviously assume that all tags on present in the final, post-quality-control dataset: i.e. if certain SNPs have been removed, it will be better to reselect the predictors – that is, these lists should really only be used as a first pass, for convenience.

In general, however, quite possibily an easier and better strategy is instead to analyse the data within an imputation context, e.g. utilising the proxy association procedures rather than using these fixed lists.

### 24.4 Gene sets

**NOTE** The gene range lists below have replaced this old gene SET file: you are advised to use the lists below rather than this file.

Here is a PLINK-format SET file, containing a genome-wide set of genes (N=18272). The co-ordinates are based on NCBI B36 assembly, dbSNP 126; a gene is arbitrarily defined as including 50kb upstream and downstream.

Download (ZIP archive): gene-list.zip http://pngu.mgh.harvard.edu/~purcell/dist/gene-list.zip

### 24.5 Gene range lists

These are gene lists: files containing lists of genes, based on either hg17 or hg18 co-ordinates. The format is one gene per row,

```
Chromosome
Start position (bp)
Stop position (bp)
Gene name
```

These lists can be used with PLINK commands such as --make-set, --range, --gene-list, --cnv-intersect, --clump-range, etc. These gene lists were downloaded from UCSC table browser for all RefSeq genes on July 24th 2008. Overlapping isoforms of the same gene were combined to form a single full length version of the gene. Isoforms that didn't overlap were left as duplicates of that gene.

Rather than using the gene sets (described above), we suggest using these gene lists to make gene sets on the fly (using --make-set-border if so desired, to add a fixed kb border on the fly).

```
Gene list (hg18): glist-hg18 
 Gene list (hg17): glist-hg17 http://pngu.mgh.harvard.edu/\simpurcell/dist/glist-hg17
```

## Chapter 25

## Miscellaneous

This page details a collection of options and commands that did not get proper mention elsewhere.

### 25.1 Output modifiers

One convenient filter is

```
--pfilter 1e-3
```

which will, for example, only report statistics with p-values less than 1e-3.

**NOTE** This is operation for the basic association tests, but do not expect this to work for all methods that return a p-value.

To obtain -log10(p) values instead of p-values in the \*adjusted file, add the flag (this does not change the output of p-values in other files)

```
--log10
```

To fix the value of lambda used for the genomic control in the \*adjusted file, instead of estimating it from the data, use the option, for example

```
--lambda 1.2
```

To obtain an extra set of columns that facilitates making a Q-Q plot in the \*.adjusted file, add the option

```
--qq-plot
```

This will work with either basic p-values, or with --log10 p-values.

## 25.2 Analyses with different species

In this respect, PLINK differentiates between species only in terms of how many chromosomes there are, and which are sex-linked or haploid. Several non-human species are supported, by adding each analysis the extra flag

```
or --horse
or --cow
or --sheep
```

### 25.3 Matrix of pairwise LD (genotype correlation)

Correlations based on genotype allele counts (i.e. w/out phasing, and for founders only) can be obtained with the commands

```
plink --file mydata --r
or
  plink --file mydata --r2
These both create a file called
  plink.ld
with a list of R or R-squared values in it.
```

#### 25.3.1 Filtering the output

By default, several filters on imposed on which pairwise calculations are calculated and reported. To only analyse SNPs that are not more than 10 SNPs apart, for example, use the option (default is 10 SNPs)

```
--ld-window 10
to specify a kb window in addition (default 1Mb)
--ld-window-kb 1000
```

and to report only values above a particular value (this only applies when the --r2 and not the --r command is used) (default is 0.2)

```
--ld-window-r2 0.2
```

The default for --ld-window-r2 is set at 0.2 to reduce the size of output files when many comparisons are made: to get all pairs reported, set --ld-window-r2 to 0.

#### 25.3.2 Obtaining LD values for a specific SNP versus all others

To obtain all LD values for a set of SNPs versus one specific SNP, use the --ld-snp command in conjunction with --r2. For example, to get a list of all values for every SNP within 1Mb of rs12345, use the command

```
plink --file mydata
--r2
--ld-snp rs12345
--ld-window-kb 1000
--ld-window 99999
--ld-window-r2 0
```

The --ld-window and --ld-window-r2 commands effectively means that output will be shown for *all* other SNPs within 1Mb of rs12345.

#### 25.3.3 Obtaining a matrix of LD values

Alternatively, it is possible to add the --matrix option, which creates a matrix of LD values rather than a list: in this case, all SNP pairs are calculated and reported.

#### 25.3.4 Haplotype-based LD calculations

A different command, --1d, instead takes two SNP IDs as parameters and calculates the R-squared based on the four haplotype frequencies: i.e. unlike the basic --r2 command, which is based simply on the genotypic correlation, this involves phasing, just for one particular pair of SNPs. For example:

```
plink --file mydata --ld rs12345 rs67890
```

No output files are generated apart from the LOG file, which reports the estimated R-squared value:

```
LD information for SNP pair [ rs12345 rs67890 ] r-sq = 0.944388
```

Again, these calculations are based only on founders.

#### 25.4 Known issues

Development of PLINK is ongoing: as such, there is always likely to be a list of features, listed here, that are only partially implemented, or have known problems not yet fixed. A list of known issues can be found on the warnings page:

http://pngu.mgh.harvard.edu/purcell/plink/warnings.shtml

## Chapter 26

## FAQ and Hints

This section contains a small but expanding set of answers to questions and hints.

- Can I convert my binary PED fileset back into a standard PED/MAP fileset?
- Can I speed up loading large files?
- Why are no individuals included in my analysis?
- Why are my results different from an analysis using program X?
- How large a file can PLINK handle?
- Why does my linear/logistic regression output have all NA's?
- What kind of computer do I need to run PLINK?
- Can I analyse multiple phenotypes in a single run (e.g. for gene expression datasets)?
- How does PLINK handle the X chromosome in association tests?
- Can/why can't gPLINK perform a particular PLINK command?
- When I include covariates with --linear or --logistic, what do the p-values mean?

# 26.1 Can I convert my binary PED fileset back into a standard PED/MAP fileset?

Yes. Use the --recode option, for example:

```
plink --bfile mydata --recode --out mynewdata
```

You might also want to use the variant --recode12 and --recodeAD forms, described here.

## 26.2 To speed up input of a large fileset

As well as using the binary fileformat, which greatly increases speed of loading relative to the PED/MAP format, if you know that you have already excluded all the individuals you want (with the per-individual genotyping threshold option), then setting

```
--mind 1
```

will skip the step where per-individual genotyping rates are calculated, which can reduce the time taken to load the file. Note, the command --all is equivalent to specifying --mind 1 --geno 1 --maf 0 (i.e. do not apply any filters).

### 26.3 Why are no indidividuals included in the analysis?

A common cause for this is either that all individuals are non-founders (e.g. a sibling pair dataset) and PLINK, by default, only uses founders to calculate allele frequencies. The

```
--non-founders
```

option can force these individuals in.

An alternative is that none of the individuals have a valid sex code – in this case, they are all set to missing status, unless the

```
--allow-no-sex
```

option is given. You are strongly recommended to enter the correct sex codes for all individuals however, so they can be appropriate treated in any subsequent analyses involving the sex chromosomes.

## 26.4 Why are my results different from an analysis using program X?

This is obviously a difficult question to answer without specific details. Therefore, if you send me a question along these lines and want to get an answer, please make it as specific as possible, to put it bluntly! Ideally, include example data that replicates the problem / illustrates the difference.

There is always the possibility that the difference could be due to a bug in PLINK, which is obviously something I would want to track down and fix. Similarly, it could be due to a bug in the other software. Perhaps more likely, the difference might arise from one of two general sources

- The analytic routines themselves are slightly different. Are the results dramatically different? Do not expect exact numerically similarity between similar analyses (i.e. even for a simple case, --assoc, --fisher and --logistic will give slightly different p-values for a simple single SNP test, but this is to be expected). So, is the difference really meaningful? Perhaps more importantly, are you sure the other routine really is implementing a similar test, with similar assumptions, etc?
- A common reason for apparent differences between PLINK and other analysis packages is that PLINK implements some default filtering of the data, i.e. first removing individuals or SNPs with below threshold genotyping rate. Look at the LOG file to check that exactly the same set of individuals were actually included in both analyses. In other words: be sure to check how missing data were handled in each case.

## 26.5 How large a file can PLINK handle?

There are no fixed limits to the size of the data file; it uses currently 1 byte for 4 SNP genotypes and some overhead per SNP and per individual. This means that you should be able to get datasets of, say, 1 million SNPs and up to 5000 individuals, in a machine with 2GB RAM without causing too much stress/swapping, etc. That is,

```
5000 * 1e6 / 4 = 1.25e9 bytes = \tilde{1}GB.
```

Things scale more or less linearly after that. So for a very large file 4 times the size (20K individuals for example), an 8GB or 16GB machine would be required to load the data in a single run).

For datasets with very many SNPs, even the list of SNP names and storage information can take a reasonable amount of space, even if the number of individuals is small (i.e. for the Phase 2 HapMap data, most of the space is taken up with the SNP name and position information, rather than the genotypes themselves).

You can test the capacity of PLINK and your machine by entering the commands

```
plink --dummy 15000 500000 --make-bed --out test1
```

to generate a dummy file of, in this instance, 15,000 individuals genotyped on 600,000 SNPs. If you do not get an Out of memory error, then it has worked. Note that dealing with files this size will take a while. Of course, in many cases it would be easy to split up the data and do per-chromosome analyses if need be, which would help on smaller machines.

### 26.6 Why does my linear/logistic regression output have all NA's?

PLINK will set the output to be all NAs if it was unable to fit the regression model. Common causes for this are:

- There is no variation in the phenotype or one or more of the predictor variables: are you sure the right variables were selected, and that no filters were applied meaning that the individuals left are all cases, for example? Is the SNP monomorphic?
- The second reason is that the correlation between predictor variables is too strong. PLINK uses the variance inflation factor criterion (VIF) to check for multi-collinearity. If two or more variables perfectly predict each other, PLINK will (correctly) print all NAs to the output, indicating that the model can not be fit. Sometimes, PLINK may be overly-conservative in calling such problems however, which is particularly likely to occur if you add more covariates and allow for interactions between terms (as the interaction terms will correlate with the main effect variables). The default VIF is 10; try setting this value higher with the --vif option, to say 100. The VIF is 1/(1-R) where R is the multiple correlation coefficient between one predictor variable and all others. A value of 100 implies R=0.99. If one variable or more variables fail the VIF test, then the entire model is not run and NAs appear in the output.

### 26.7 What kind of computer do I need to run PLINK?

There are no special requirements: PLINK should be able to be compiled for any machine for which a recent C/C++ compiler is available. Pre-compiled binary versions are distributed from this website for Linux, MS-DOS and Mac machines.

In terms of speed, memory and diskspace, obviously more is usually better. The suggestions below are really minimum values to make life easy for a "normal" sized study (i.e. many analyses could easily be run on much smaller machines; some analyses will require more resources, etc).

The FAQ above about dataset limits gives some indication of the amount of RAM needed for large studies. Basically, for any whole genome scale studies you would want at least 2Gb of RAM; 4 or 8Gb would be desirable.

In terms of disk space: the main storage requirements will result from the raw data (e.g. CEL files, etc) rather than genotype files or most PLINK results files. However, certain PLINK files can be large: e.g. .genome files for large samples, dosage output for whole-genome imputation of all HapMap SNPs, etc. Therefore, a large hard drive is desirable: not including storage for CEL files, a drive of at least 200Gb would be good.

PLINK does not specifically take advantage of multi-core processors. For large datasets, a fast processor is desirable (e.g. at least 3GHz). The majority of analyses described in these pages can be performed on a single processor. For certain analyses (e.g. epistasis, using permutation procedures on very large datasets, IBS calculation on very large datasets, etc) then access to a parallel computing cluster, if possible, is very desirable and sometimes necessary.

In terms of operating systems, there should not be major differences in performance: using a Linux/Unix environment probably has some advantages in terms of the existing text file processing utilities typically available, and the more powerful shell scripting options, but probably personal preference and institutional support is a bigger consideration. There is a definite advantage to ensuring a C/C++ compiler exists on the system so that the source code version of PLINK can be compiled for your particular system however – this may give some performance advantages and allows access to the development source code (i.e. to receive a

patched version that fixes a particular problem or adds a new feature before the next release in generally available).

# 26.8 Can I analyse multiple phenotypes in a single run (e.g. for gene expression datasets)?

For most association commands, you can specify the --all-pheno option to automatically loop over all phenotypes in an alternate phenotype file:

```
plink --bfile mydata --pheno phenos.raw --all-pheno --linear --covar covar.dat
```

If there are N phenotypes, this will generate N separate output files. If a header row was supplied in the alternate phenotype file, then each file will have the phenotype name appended (it is up to the user therefore to ensure that the phenotype names are unique). If not, the output files are simply numbered, P1, P2, etc, (e.g. plink.P1.assoc, etc).

This works for most basic association commands that consider all SNPs (e.g. --assoc, --logistic, --fisher, --cmh, etc) but currently not for any haplotype analysis or epistasis options.

## 26.9 How does PLINK handle the X chromosome in association tests?

By default, in the linear and logistic (--linear, --logistic) models, for alleles A and B, males are coded

A -> 0 B -> 1

and females are coded

AA -> O

AB -> 1

BB -> :

and additionally sex (0=male,1=female) is also automatically included as a covariate. It is therefore important not to include sex as a separate covariate in a covariate file ever, but rather to use the special --sex command that tells PLINK to add sex as coded in the PED/FAM file as the covariate (in this way, it is not double entered for X chromosome markers). If the sample is all female or all male, PLINK will know not to add sex as an additional covariate for X chromosome markers.

The basic association tests that are allelic (--assoc, --mh, etc) do not need any special changes for X chromosome markers: the above only applies to the linear and logistic models where the individual, not the allele, is the unit of analysis. Similarly, the TDT remains unchanged. For the --model test and Hardy-Weinberg calculations, male X chromosome genotypes are excluded.

Not all analyses currently handle X chromosomes markers (for example, LD pruning, epistasis, IBS calculations) but support will be added in future.

## 26.10 Can/why can't gPLINK perform a particular PLINK command?

gPLINK is intended only as a lightweight interface to some of the basic PLINK commands. It is designed to provide an easy way to become familiar with PLINK and to perform certain very basic operations for users who are not yet familiar with command line interfaces. It is not the recommended mode for using PLINK for anything beyond the most basic analyses and there are no immediate plans to extend gPLINK any further to incorporate new commands that are added to PLINK.

# 26.11 When I include covariates with --linear or --logistic, what do the p-values mean?

If one or more covariates are included (by --covar) when using --linear or --logistic, PLINK performs a multiple regression analysis and reports the coefficients and p-values for each term (i.e. SNP, covariates, any interaction terms). The only term omitted from the report is the intercept.

The p-values for the covariates **do not** represent the test for the SNP-phenotype association after controlling for the covariate. That is the first row (ADD). Rather, the covariate term is the test associated with the covariate-phenotype association. These p-values might be extremely significant (e.g. if one covaries for smoking in an analysis of heart disease, etc) but this does not mean that the SNP has a highly significant effect necessarily. For example:

```
CHR.
            SNP
                      BP
                          A1
                                      NMISS
                                                  BETA
                                                                             Ρ
                                TEST
                                                            STAT
  1
     rs1234567
                 742429
                           G
                                 ADD
                                        1495
                                              -0.03335
                                                         -0.1732
                                                                        0.8625
  1
     rs1234567
                 742429
                           G
                               COV1
                                       1495
                                                0.1143
                                                           9.748
                                                                   8.321e-022
```

suggests that the covariate is highly correlated with the outcome (which will often be already known, presumably), but there is no evidence that the SNP is in any way correlated with phenotype. These correspond to the partial regression coefficient terms of a muliple regression

```
Y \sim m + b1.ADD + b2.COV1 + e
```

where p=0.8625 is the Wald test for b1, p=8e-22 is the Wald test for b2, the covariate-phenotype relationship. To repeat: it does not mean that the SNP-phenotype test has a p=8e-22 after controlling for COV1.

## Appendix A

## Reference Tables

## A.1 Options

Option	Parameter/default	Description
Basic input/output	,	•
file	plink	Specify .ped and .map files
ped	plink.ped	Specify .ped file
map	plink.map	Specify .map file
тар	ринк.шар	Specify map me
no-sex		PED file does not contain column 5 (sex)
no-parents		PED file does not contain columns 3,4 (parents)
no-fid		PED file does not contain column 1 (family ID)
no-pheno		PED file does not contain column 6 (phenotype)
liability		PED file does contain liability (column 7)
map3		Specify 3-column MAP file format
шарз		Specify 3-column WAT the format
tfile	plink	Specify .tped and .tfam files
tped	plink.tped	Specify .tped file
tfam	plink.tfam	Specify .tfam file
	-	
lfile	plink	Specify long-format: LGEN, FAM and MAP
bfile	plink	Specify .bed, .bim and .fam
bed	plink.bed	Specify .bed file
bim	plink.bim	Specify .bim file
fam	plink.fam	Specify .fam file
<u>-</u>	piiiik.iaiii	Specify train the
out	plink	Specify output root filename
silent		Suppress output to console
pheno	phenofile	Specify alternate phenotype
make-pheno	file value	Specify binary phenotype, with cases have value
make-pheno	file *	Specify binary phenotype, with cases are present
mpheno	var #	Specify which, if >1 phenotype column
pheno-name	var name	Instead ofmpheno, if a header row exists
all-pheno		Perform association for all phenotypes in file
loop-assoc	clusterfile	Perform association for each level of cluster versis all others
100P assoc		1 criorin association for each rever of classer versis an others
covar	covarfile	Specify covariate
mcovar	var #	Specify which, if >1 covariate column
within	filename	Specify clustering scheme
mwithin	var #	Specify which, if >1 cluster column
MWI CHIII	vai #	specify which, if >1 classer column
script	filename	Include command-line options from file
Selection of SNPs and individuals		
chr	N	Select a particular chromosome N
gene	name	Select a particular gene, given a SET file (set)
	~	
from	SNP	Select range from this SNP
to	SNP	to this SNP (must be on same chromosome)
snps	SNP list	Select comma-delimited list of SNPs, allowing for ranges, e.g. snp1,snp2,snp6-snp12
snp	SNP	Select this SNP
window	kb	and (optionally) all SNPs in the surrounding kb window
from-bp	bp	Select SNPs within this window
to-bp	bp	specified in base-pair position

Select SNPs within this window... --from-kb kb --to-kb kb specified in kilobases Select SNPs within this window...
... specified in megabases --from-mb --to-mb  $_{\mathrm{mb}}$ --extract snplist snplist Extract list of SNPs Exclude list of SNPs --keep --remove Keep only these individuals Remove these individuals indlist indlist Perform keep before remove (default opposite) --keep-before-remove --exclude-before-extract Perform exclude before extract (default opposite) Filter individuals matching value Specify filter value, if >1 filter column filename value --mfilter var # Include only cases
Include only controls
Include only males
Include only females
Include only founders
Include only nonfounders --filter-cases --filter-controls --filter-males --filter-females --filter-founders --filter-nonfounders Remove individuals with missing phenotypes --prune Other data management options Make .bed, .fam and .bim Output new .ped and .map files As above, with 1/2 allele coding --make-bed --recode --recode12 As above, with Haploview .info file Ouput fastphase format file Ouput bimbam format file Ouput structure format file --recodeHV --recode-fastphase --recode-bimbam --recode-structure Raw data file with additive coding
Raw data file with additive/dominance coding
Delimit --recode and --recode12 with tabs
Output one genotype per line, list of FIDs and IIDs
Pairwise listing of genotypes for two individuals --recodeA --recodeAD --list FID1 IID1 FID2 IID2 List only the (filtered) SNPs in the dataset --write-snplist --update-map Update physical positions in a map file Update genetic distances in a map file filename Output ordered, filtered covariate file --write-covar --with-phenotype --dummy-coding Include PED/phenotype information in new covariate file Downcode categorical covariates to binary dummy variables Merge in a PED/MAP fileset Merge in a binary fileset Merge multiple standard and/or binary filesets Specify merge mode (1-7) pedfile, mapfile bedfile, bimfile, famfile list file --merge --bmerge --merge-list --merge-mode --zero-cluster --oblig-missing --oblig-cluster filename filename filename Zero-out specific SNPs for specific clusters SNPs/clusters that are obligatory missing Individuals/clusters defining obligatory missingness --flip --flip-subset snplist individual-list Flip strand of SNPs in list Flip strand of SNPs only for these individuals in list LD-based heuristic to look for SNPs flipped between cases and controls --flip-scan 0/1 unaffected/affected coding Use AA, AG, 00 coding (no spaces between alleles in PED file) Missing phenotype code Missing genotype code Missing phenotype code for output Missing genotype code for output Convert (A,C,G,T) to (1,2,3,4) Convert (1,2,3,4) to (A,C,G,T) Do not flip A1 to be the minor allele --compound-genotypes --missing-phenotype -9 0 -9 0 --missing-genotype
--missing-output-phenotype
--missing-output-genotype --allele1234 --alleleACGT --keep-allele-order Do not set ambiguously-sexed individuals missing When making a new dataset, do set ambiguously-sexed individuals missing Making new fileset, set heterozygous haploids missing Making new fileset, set Mendel errors missing --allow-no-sex --must-have-sex --set-hh-missing --set-me-missing --make-founders Set non-founders without two parents to founders When performing TDT, dump parsed family structure Reporting summary statistics Allele frequencies
Modifies --freq to report actual allele counts
Include all individuals in MAF/HWE calculations --freq

--nonfounders

--missing --test-missing

Missing rates (per individual, per SNP) Test of missingness differing by case/control status

--test-mishap Haplotype-based test for non-random missingness --cluster-missing IBM clustering Report Hardy-Weinberg disequilibrium tests (exact) Report Hardy-Weinberg disequilibrium tests (asymptotic) --hardy --hardy2 --mendel Report Mendel error checks Use X chromosome data to check an individual's assigned sex Use X chromosome data to impute an individual's assigned sex --check-sex --impute-sex --within cluster file Stratify frequencies and missing rates by clusters Inclusion thresholds Minor allele frequency Maximum minor allele frequency Maximum per-SNP missing 0.01 --max-maf 0.1 --geno Maximum per-SNP missing
Maximum per-person missing
Hardy-Weinberg disequilibrium p-value (exact)
Hardy-Weinberg disequilibrium p-value (asymptotic)
Hardy-Weinberg disequilibrium p-value (asymptotic)
HW filtering based on all founder individuals for binary trait (instead of just unaffecteds)
Mendel error rate thresholds (per SNP, per family)
Minimum genotype cell count for --model
Minimum pi-hat for --genome output
Maximum pi-hat for --genome output --mind 0.1 --hwe  $0.001 \\ 0.001$ --hwe-all 0.1 0.1 --me --cell 0 --max IBS stratification / clustering Calculate IBS distances between all individuals --genome --cluster Perform clustering
Output IBS (similarity) matrix
Output 1-IBS (distance) matrix
Maximum cluster size --matrix --distance-matrix --mc 0 Maximum cluster size
Cluster by phenotype
Maximum number of cases/controls per cluster
Constrain IBS matching on IBM matching
IBS test p-value threshold (was --pærge)
Skip SNPs within this for PPC test --cc --mcc --ibm --ppc 0.0  $0.01 \\ 0.01$ --ppc-gap --match --match-type 500kb match-file match-type-file match-file Specify external categorical matching criteria Specify external categorical matching direction (+/- match) Specify external quantitative matching criteria --qmatch threshold-file N M --qt --neighbour Specify quantitative matching thresholds Outlier statistics (for nearest neighbours N to M) Whole genome summary statistics Output genome-wide IBS/IBD
Read previously-computed genome values
Adjusted estimated IBD values
Indicate 'impossible' estimated IBD values
Individual inbreeding F / heterozygosity
Identify runs of homozygosity (# SNPs)
Allow for N hets in run of homozygosity
Group pools of overlapping segments
Identify threshold for allelic matching overlapping segments
Display actual genotypes for each pool --genome
--read-genome
--nudge
--impossible
--het genome-file --homozyg-kb --homozyg-snp N SNPs N hets --homozyg-het --homozyg-group --homozyg-match 0.95 --homozyg-verbose Association analysis procedures Case/control or QTL association Fisher's exact (allelic) test Cochran-Armitage and full-model C/C association Exact full-model tests --assoc --fisher --model --model --fisher Hotelling's T(2) multilocus test --T2 Cochran-Mantel-Haenszel SNPxDISEASE—STRATA Cochran-Mantel-Haenszel SNPxSTRATA—DISEASE Breslow-Day homogeneity of odds ratios test --mh2 --bd --homog Partitioning chi-square homogeneity of odds ratios test QTL interaction test (dichotomous covariate only) --gxe Test for quantitative traits and multiple covariates Test for disease traits and multiple covariates Include dominance term in model, and 2df model --linear --logistic --genotypic Include dominance term in model, and 2df mode Fit dominant model for minor allele Fit recessive model for minor allele Include additive effects of these SNPs in model Include additive effects of these SNPs in model Include sex effect in model Include all covariates in a covariate file Include SNP x covariate interactions

Joint test of all terms in model --dominant --recessive SNP filename --condition-list --sex --covar --interaction filename --test-all --parameters 1,2,...1,2,...Fit only a subset of model terms Joint test of user-specified set of parameters Family-based TDT and parenTDT (permute TDT) --tdt As above, except permuted statistic is parental test As above, except permuted statistic is combined test Parent-of-origin analysis in TDT --parentdt1 --parentdt2

dfam		Disease family-test (families and unrelateds)
ci	0.95	Confidence interval for CMH odds ratios
set-test		Set-based association (requiresmperm)
set-p	p-value r2	p-value threshold for set-based test R-squared threshold for set-based test
set-r2 set-max	N SNPs	Maximum number of SNPs in set
	1, 51,15	Maximum number of pitt's in sec
Permutation procedure optionsperm		Run permutations (adaptive-mode)
mperm	1000	# of permutations in max-perm mode
aperm		Parameters (six) for adaptive permutation mode
rank		Modifiesmperm for rank-based permutation
model-trend		Use CA-trend test frommodel
model-gen model-dom		Use genotypic test frommodel Use dominant test frommodel
model-rec		Use recessive test frommodel
,		D (() 1 1 () (C 11 1 ()
genedrop swap-parents		Permutation by gene-dropping simulation (family-data)  Labal-swap permutation for parents when gene-dropping
swap-sibs		Labal-swap permutation for siblings when gene-dropping
swap-unrel		Labal-swap permutation for unrelateds when gene-dropping
family		Make Family ID the cluster
p2		Alternate permutation scheme (C/C only)
Epistasis analysis		
epistasis		Perform SNP x SNP epistatic analysis
fast-epistasis twolocus	SNP SNP	Quick SNP x SNP screening for C/C data Display contingency table for two SNPs
case-only		Case-only epistatic analysis
gap	1000	Gap (kb) for SNP x SNP case-only epistasis tests
epi1 epi2	0.0001 0.01	Output p-value threshold: pairs Output p-value threshold: summary
->set-by-all		Test set 1 SNPs paired with all others
nop		Do not calculate p-values (fast screening)
genepi		Gene-based test for epistasis
Haplotype inference and linkage disequilibriumhap-snps	snplist	Specify a list of SNPs to phase
hap-window	N	Specify haplotype sliding window
hap whap	tagfilename tagfilename	Multimarker predictor / haplotype list Weighted haplotype test list
wnap	tagmename	weighted haplotype test list
hap-assoc		Perform haplotype-based case/control association
hap-tdt hap-freq		Perform haplotype-based TDT Output haplotype frequencies for entire sample
hap-phase		Output individual haplotype phases
hap-phase-wide		Output individual haplotype phases, wide-format
hap-impute hap-pp	0.8	Create fileset with imputed haplotypes as SNPs Posterior probability threshold
hap-miss	0.5	Proportion of missing genotypes allowed
hap-min-phase-prob	0.01 N	Minimum reported phase probability
hap-max-phase mhf	0.01	Maximum number of phases considered per person Minor haplotype frequency threshold
Proxy association and imputation methods		
proxy association and imputation methods	SNP/all	Proxy association methods
proxy-glm	•	Use linear models in proxy association
proxy-drop proxy-tdt	SNP/all	Drop then re-impute observed genotypes Proxy TDT association methods
	•	·
proxy-imputeproxy-replace	SNP/all	Proxy imputation methods Replace observed genotypes
proxy-dosage		Also output dosage file
proxy-impute-threshold	0.95	Per-genotype threshold to impute for an individual
proxy-list	file	Specify SNPs to impute/test
proxy-flanking	file	Specify proxies for single reference SNP
proxy-r2	0 0.05 0.5	Proxy selection LD parameters
proxy-maxsnp proxy-window	5 15	Maximum number of proxies tto select Proxy SNP search space (SNPs)
proxy-window proxy-kb	250	Proxy SNP search space (kb)
proxy-b-threshold	0.1	MAF threshold for rare alleles (plan B)
proxy-b-r2	0.05 0.5	Alternate proxy selection LD parameters
proxy-b-maxsnp	0.1	Alternate maximum number of proxies to use
proxy-b-window proxy-b-kb	0.1 250	Alternate proxy SNP search space (SNPs) Alternate proxy SNP search space (kb)
From a me	200	in proxy one contain space (RD)

Proxy SNP MAF threshold Proxy SNP missingness threshold No LD-based proxy selection --proxy-maf 0.01 --proxy-geno --proxy-r2-no-filter 0.05 --proxy-mhf 0.05 Proxy haplotype frequency threshold --proxy-sub-maxsnp 0.8 Maximum number of SNPs per haplotypic proxies (verbose mode) --proxy-verbose Verbose mode --proxy-show-proxies
--proxy-genotypic-concordance List actual proxies in non-verbose mode In imputation, show genotypic-specific concordance Conditional haplotype association tests Main conditional-haplotype test command Test for specific haplogroup effect Test for independent effect --chap --specific-haplotype haplotype(s) --independent-effect snp(s)/haplotype(s) Control for certain effects Specify SNP groupings under alternate Specify SNP groupings under null --alt-snp --null-snp --alt-group Specify haplogroupings under alternate Specify haplogroupings under null --test-snp --each-versus-others --each-vs-others Drop 1 or more conditioning SNPs Each all haplogroup-specific p-values As above LD-based result clumping Comma-delimited result files p-value threshold for index SNPs p-value threshold for clumped SNPs r<sup>2</sup> (LD) threshold for clumping kb-threshold for clumping --clump --clump-p1 file(s) 1e-4 1e-2 --clump-p2 --clump-r2  $0.2 \\ 250$ Only report multi-file clumps
For each SNP in the first file, find the best proxy from the other files
Specifty verbose output
Add gene/region range information to clumped output
Use a kb border around each gene/region
Include these fields in verbose mode
Specifty p-value field other than P
Only index based on first results file --clump-replicate --clump-best --clump-verbose --clump-range --clump-range-border --clump-annotate --clump-field filename field(s) field --clump-index-first
--clump-allow-overlap Only index based on first results file Specify that a SNP can appear in more than one clump Gene annotation of SNP results Results file to perform gene-report on List of genes/regions for reporting Add a kb border aroud each gene/region Only report on a subset of genes, listed here filename --gene-list-border filename kb filename --gene-subset --gene-report-empty Report genes without any informative SNPs LD pruning and pairwise LD N M VIF VIF pruning (N-SNP window, shifted at M-SNP intervals) --indep --indep-pairwise r2 pruning (as above) Pairwise SNPxSNP LD (r) N M r2 Pairwise SNPxSNP LD  $(\hat{r2})$ Limit pairwise SNPxSNP to within a N SNP window --r2 Ν Definition of SETs setfilename SET definitions Copy number variants (CNV) analysis fileset --cfile fileset --cnv-list filename Filter only deletions --cnv-del Filter only duplications --cnv-dup --cnv-intersect filename Include segments intersecting with regions Exclude segments intersecting with regions Include/Exclude segments that start or stop within a gene/region Count number of regions intersected by CNVs Add a kb border around each region --cnv-exclude filename --cnv-disrupt
--cnv-count
--cnv-border filename kb Exclude CNVs overlapping regions with more than N CNVs Exclude CNVs overlapping regions with fewer than N CNVs Exclude CNVs overlapping regions with exactly N CNVs Include CNVs overlapping regions with exactly N CNVs Use alternative method for determining CNV frequency --cnv-freq-excldue-above --cnv-freq-excldue-below --cnv-freq-excldue-exact N N N --cnv-freq-incldue-exact --cnv-freq-method2 Define overlap of CNV and region by CNV length Define overlap of CNV and region by union Define overlap of CNV and region by region length --cnv-overlap N N N --cnv-union-overlap Create a new CNV and FAM file --cnv-write --cnv-write-freq Include frequency counts if --cnv-freq-method2 specified

190

cnv-make-map cnv-report-regions		Create a new MAP file from a CNV and FAM file List regions that are intersected by CNVs
cnv-report-regionscnv-verbose-report-regions		Verbose listing of regions that are intersected by CNVs
cnv-subset	filename	Define overlap of CNV and region by region length
CIV Subsec	mename	Define overlap of Civv and region by region length
cnv-track	kb	Create a UCSC-compatible BED track for viewing CNVs
cnv-blue	kb	Make this CNV track blue
cnv-red	kb	Make this CNV track red
cnv-green	kb	Make this CNV track green
cnv-brown	kb	Make this CNV track brown
cnv-kb	N	Exclude segments below N kb
cnv-max-kb	N	Exclude segments above N kb
cnv-score	N	Exclude segments below N score
cnv-max-score	N	Exclude segments above N score
cnv-drop-no-segment		Remove individuals with no segments
cnv-unique		Exclude CNVs seen in both cases and controls
cnv-seglist	kb	Create a printout of CNVs
cnv-indiv-perm		Permutation test for genome-wide CNV burden
cnv-test-2sided		Use 2-sided approach for empirical p-values
cnv-test-window	kb	Extend test to a region extending kb distance on either side of position
Data simulation options		
simulate	filename	Simulate SNP population-based data
simulate-ncases	100	Number of cases to simulate
simulate-ncontrols	100	Number of controls to simulate
simulate-prevalence	0.01	Disease prevalence in population
dummy	N M	Generate dataset of N individuals on M SNPs
Misc analysis output options		
adjust		Output adjusted p-values and calculate genomic control
lambda	X	Set lambda to X instead of estimating from data
qq-plot		Generate entries to faciliate a Q-Q plot in adjusted output
Misc.		
help		Display list of options
dog		Set chromosome codes for dog
mouse		Set chromosome codes for mouse
horse		Set chromosome codes for horse
cow		Set chromosome codes for cow
sheep		Set chromosome codes for sheep
lookup	SNP rs#	Lookup WGAS SNP annotation information
lookup-gene	gene name	List all SNPs in gene
lookup-list	snplist filename	SNP annotation for multiple SNPs
=	=	

## A.2 Output files (alphabetical listing)

Filename	Main associated command(s)	Description
plink.adjust	adjust	Adjusted significance values (multiple testing
plink.assoc	assoc	Association results
plink.assoc.hap	hap-assoc	Haplotype-based association results
plink.assoc.linear	linear	Linear regression model
plink.assoc.logistic	logistic	Logistic regression model
plink.assoc.mperm	assocmperm	maxT permutation empirical p-values
plink.assoc.perm	assocperm	Adaptive permutation empirical p-values
plink.assoc.proxy	proxy-assoc	Proxy association results
plink.assoc.set	assocset	Set-based association results
plink.bed	make-bed	Binary PED file
plink.bim	make-bed	Binary MAP file
plink.chap	chap	Conditional haplotype tests
plink.cov	write-covar	Ordered, filtered covariate file
plink.clumped	clump	LD-based results clumping
plink.clumped.best	clump-best	Single best LD-based clumping
plink.clumped.ranges	clump-range	Gene/region report for clumps
plink.cluster0	cluster	Progress of IBS clustering
plink.cluster1	cluster	IBS cluster solution, format 1
plink.cluster2	cluster	IBS cluster solution, format 2
plink.cluster3	cluster	IBS cluster solution, format 3
plink.cluster3.missing	cluster-missing	IBM cluster solution, format 3
plink.cmh	mh	Cochran-Mantel-Haenszel test 1
plink.cmh2	mh2	Cochran-Mantel-Haenszel test 2
plink.cnv.indiv	cnv-list	Copy number variant per individual summary
plink.cnv.overlap	cnv-list	Copy number variant overlap
plink.cnv.summary	cnv-list	Copy number variant summary
plink.cnv.summary.mperm	cnv-list	Copy number variant test
plink.diff	merge-mode 6/7	Difference file
plink.epi-cc1	epistasis	Epistasis: case/control pairwise results
plink.epi-cc2	epistasis	Epistasis: case/control summary results
plink.epi-co1	epistasiscase-only	Epistasis: case-only pairwise results
plink.epi-co2	epistasiscase-only	Epistasis: case-only summary results
plink.fam	make-bed	Binary FAM file
plink.fmendel	mendel	Mendel errors, per family
plink.frq	freq	Allele frequency table
plink.frq.count	freqcounts	Allele counts table
plink.frq.hap	hap-freq	Allele frequency table
plink.genepi.dat	genepi	Gene-based epistasis R dataset
plink.genepi.R	genepi	Gene-based epistasis R script
plink.genome	genome	Genome-wide IBD/IBS pairwise measures
-		• •

```
plink.het
                                                                                                                                                                               Individual inbreeding coefficients
                                                                                         --het
                                                                                                                                                                               Runs of homozygosity
Pools of overlapping runs of homozygosity
 plink.hh
plink.hom
plink.hom.overlap
                                                                                       --homozyg-snp --homozyg-kb
--homozyg-group
                                                                                                                                                                               Between strata homogeneity test
Hardy-Weinberg test statistics
Mendel errors, per individual
Missing rates, per individual
Info file for Haploview filesets
List of individuals removed for low genotyping
Imputed from multi-marker predictors
Imputed from multi-marker predictors
plink.homog
                                                                                         --homog
 plink.hwe
                                                                                        --hardy
plink.imendel
plink.imiss
                                                                                         --missing
                                                                                         --recodeHV
plink.info
plink.irem
plink.imputed.map
                                                                                        --mind
                                                                                         --hap-impute
                                                                                                                                                                               Imputed from multi-marker predictors
Imputed from multi-marker predictors
Recoded LIST file
Mendel errors, per locus
Missing rates, per locus
Log file (always generated)
Recoded MAP file
plink.impute.ped
                                                                                          --hap-impute
 plink.list
                                                                                         --list
 plink.lmendel
                                                                                         --mendel
plink.lmiss
plink.log
                                                                                         --missing
plink.map
plink.mdist
plink.mdist.missing
                                                                                         --recode
                                                                                        --cluster --matrix
--cluster-missing
                                                                                                                                                                               IBS distance matrix
IBM distance matrix
                                                                                                                                                                             IBM distance matrix
Mendel errors, per error
List of SNPs that show problem phasing (could not be found or on wrong chromosome)
Test of differences in C/C missing rates
Haplotype-based test of non-random genotyping failure
List of SNPs that show strand problems when merging files (more than 2 alleles)
Full-model association results
Best full-model association max(T) permutation results
Genotypic association max(T) permutation results
Genotypic association max(T) permutation results
Genotypic association max(T) permutation results
Dominant association adaptive permutation results
Trend test association max(T) permutation results
Trend test association max(T) permutation results
Recessive association adaptive permutation results
Recessive association adaptive permutation results
List of SNPs with no observed founders
List of individuals with ambiguous sex code
Nearest neighbour (IBS) statistics
Information on pedigree structure
Recoded PED file
Haplotype phases (one file per locus)
Pairwise list of two people's genotypes
Proxy imputation output
Proxy imputation dosage output
 plink.mendel
                                                                                          --mendel
                                                                                                                                                                                Mendel errors, per error
                                                                                        --hap
--test-missing
--test-mishap
 plink.mishap
plink.missing
plink.missing.hap
 plink.missnp
                                                                                         --merge
 plink.model
                                                                                         --model
plink.model.best.mperm
plink.model.best.perm
                                                                                        --model --mperm
                                                                                       --model --mperm
--model --mperm
--model --mperm --model-gen
--model --mperm --model-dom
--model --mperm --model-dom
--model --mperm --model-dom
--model --mperm --model-trend
--model --perm --model-trend
--model --perm --model-trend
--model --perm --model-rec
--model --perm --model-rec
 plink.model.gen.mperm
plink.model.gen.perm
plink.model.dom.mperm
plink.model.dom.perm
plink.model.trend.mperm
plink.model.trend.perm
plink.model.rec.mperm
plink.model.rec.perm
plink.nof
plink.nosex
plink.nearest
                                                                                            -cluster --neighbour
plink.pdump
                                                                                         --pedigree
plink.pdd
plink.ped
plink.phase-*
plink.plist
                                                                                        --recode
--hap --phase
--plist
                                                                                                                                                                               Proxy imputation output
Proxy imputation dosage output
Verbose proxy association output
List of remaining SNPs (i.e. not pruned)
List of pruned-out SNPs
plink.proxy.impute
plink.proxy.impute.dosage
plink.proxy.report
plink.prune.in
                                                                                         --proxy-impute
                                                                                        --proxy-impute --proxy-dosage
--proxy-assoc
--indep --indep-pairwise
--indep --indep-pairwise
 plink.prune.out
                                                                                                                                                                               List of pruned-out SNPs
Quantitative trait association results
Quantitative trait interaction results
Listing of CNVs by genes/regions
Recoded additive/dominance format file
plink.qassoc
plink.qassoc.gxe
                                                                                        --assoc
                                                                                        --gxe
--cnv-verbose-report-regions
 plink.range.report
 plink.raw
                                                                                         --recodeAD
                                                                                                                                                                               Recoded additive/dominance format file
List of SNPs in the dataset
Hotelling's T(2) test results
TDT/parenTDT asymptotic results
TDT/parenTDT permutaion results
TDT/parenTDT max(T) permutation results
TDT/parenTDT adaptive permutation results
plink.raw
plink.snplist
plink.T2
plink.tdt
                                                                                         --write-snplist
                                                                                        --T2
--tdt
 plink.tdt.hap
                                                                                        --tdt
plink.tdt.mperm
plink.tdt.perm
                                                                                       --tdt
--tdt
                                                                                                                                                                               TDT parent-of-origin results
TDT parent-of-origin max(T) permutation results
TDT parent-of-origin adaptive permutation results
TDT parent-of-origin set-based results
 plink.tdt.poo
                                                                                         --tdt
                                                                                                        --poo
                                                                                       -tdt --poo --mperm
-tdt --poo --mperm
-tdt --poo --perm
-tdt --poo --set --mperm
-tdt --set --mperm
-transpose / --tfile
--transpose / --tfile
plink.tdt.poo.mperm
plink.tdt.poo.perm
plink.tdt.poo.set
                                                                                                                                                                               TDT/parenTDT set-based results
FAM for for transposed fileset
PED file for transposed fileset
SNP x SNP contingency table
 plink.tdt.set
 plink.tfam
plink.tped
plink.twolocus
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192