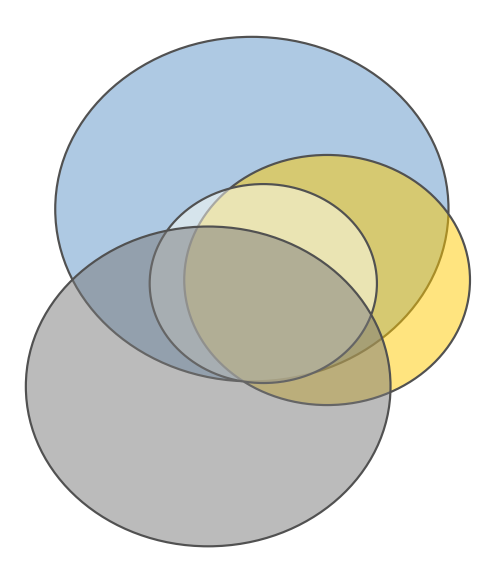
## BIOFILTER

Reference Manual rev. 1.0.1



biofilter - Biological Knowledge Integration Utility

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## Introduction

## Purpose of this manual

Contained within this manual are details for configuring and running the application, biofilter. If this is your first time to use the software, we highly recommend that you take a few minutes to download and work through one or more tutorials. Then, once familiar with the capabilities of the software, users can refer to this guide when making changes to the basic configuration settings.

## **Conventions Used**

There are two conventions used throughout this document. These text conventions are intended to help distinguish examples from configuration parameters.

## **Example commands**

```
biofilter sample.config --list-associations
```

The application itself will be named differently according to the platform it was compiled for. On Linux systems, it will be biofilter or biofilter64 depending on whether it runs on 32bit or 64bit systems. Windows and OS X will have be named slightly differently according to their platform as well.

## **Program Output**

```
Program output will be in a gray box. Some lines will often be removed when the output is lengthy.
```

## Configuration details are listed first in bold-left aligned with the rest of the text.

The first word(s) are the keywords which specify what is being changed. Each keyword (or phrase) has some number of parameters. These are listed in the order they should appear in the configuration line. In some cases, parameters can be repeated or are optional. Those are denoted inside []s.

Configuration details are generally followed immediately by an example line:

```
This is an example
```

Examples show how an actual entry would look and are followed by some descriptive information to help the user understand how the example would affect the biofilter application runtime.

## **Common Parameters**

There are a number of parameters which are used commonly across multiple configuration settings. In order to simplify the descriptions of the various properties of each command, we'll describe those properties here, and just refer to them as if they were a type.

#### Integer

Parameters specified in this way just simply refer to a whole number. In general, these values should be equal to or greater than 0, except when specified otherwise.

#### Float

Values specified as float are decimal values.

#### Index

If a parameter is listed as an index, it refers to the index, starting at 1 the user wishes to select.

#### max

This is generally an integer value representing the upper bound of some value. In some cases, such as minor allele frequency, it might represent a floating point value.

#### min

This is generally an integer value representing the lower bound of some value. In some cases, such as minor allele frequency, it is possible that it represents a floating point value.

#### On/Off

These parameters accept a boolean, Yes/No type setting. Users can use ON/OFF or YES/NO to set them.

## filename

When a configuration refers to a file for input or output, the filename is generally used. This can be either a fully qualified path (such as /home/torstees/wga) or it can specified as a path relative to the directory where the application was run (such as ../data/goodfilename). It can also be just a plain filename as long as the file itself is available from the directory in which the application was run.

#### label

A label refers to a parameter whose value can be any text string without whitespace. These labels are generally used for reporting but in many cases are used to determine filenames. As a result, users should avoid using unusual characters in the string that could possibly cause problems with filenames. Because spaces and tabs are used to separate each parameter on a given line, labels can not contain spaces.

#### description

A description is a chunk of text that can contain spaces. It will always be at the very end of a line and is generally optional.

# Using the Biofilter Application

The biofilter stand alone application can be used for more than one purpose. As a result, execution may take more than one set of parameters.

Many configuration parameters can be overridden on the command line. Those parameters are prepended with a specially designated keyword which is prefixed by a "-" sign. These parameters might take more than one value. Below is a list of the parameters currently supported by the biofilter application. All parameters except the configuration file are optional.

## **Command-Line Arguments**

biofilter config-file [--S (--sample-config)]

## Config-file

Specifies the filename to be used to control genomeSIMLAs overall specific behavior. If the configuration is available from within the current working directory, the filename alone is sufficient. If the filename exists in another directory, a fully qualified or relative path should be provided along with the filename itself.

## --sample-config (optional)

This parameter takes no arguments and causes the output of the biofilter to generate a basic configuration based on the default settings (and any that have been overridden by other parameters) No other execution is performed.

## -C (--coverage) filename

## filename

Adds a file to the coverage report list. Coverage files contain RS IDs for a platform used for a <u>coverage report</u>.

#### -D (--detailed-coverage)

Causes the coverage report to provide more detailed information.

## -d (--disease-dependent) filename

### filename

Adds a <u>disease dependent group</u> to the system. This is the same as adding a DISEASE\_DEPENDENT line to the configuration file.

## -G (--list-groups) [criteria]

## criteria (optional)

This string can be used to search the group's name and description. If a match is found, the Meta Group name, group ID, group name and group description are displayed for each match. If this command appears as the last flag on the command line, all groups will be listed.

This list does not refer to any setting other than the value at SETTINGS\_DB. As a result, all groups that are part of the SETTINGS\_DB are considered regardless of restrictive group limitations set within the configuration.

## -L (--list-models) filename

### filename

The name of the model file from a previous run.

This produces a basic model report.

## --model-report filename

#### filename

File containing pairwise snp models. Each line contains a single pairwise model listed as integer values separated by whitespace.

#### --marker-info

Produce <u>haploview</u> compatible marker-info files based on the <u>SNPS\_SOURCE</u> platform.

## --report-gene-coverage filename

#### filename

This file holds the list of genes for which coverage is to be reported in the coverage report.

## -s (--snps) filename

## filename

This overloads the setting, <u>SNPS SOURCE</u>, from the configuration file.

## --snp-report

Produces a report containing all genes associated with each of the SNPs found in **SNPS\_SOURCE** 

## -W (--write-models) filename

## filename

The filename to which the models will be written. The file generated will be a binary file, and should be usable with applications developed by the Ritchie Lab. Users wanting to peruse these resulting models should use the various reporting features upon the model file.

## -m (--show-models) filename

#### filename

Filename indicates the name of the model file (file produced during --write-models command).

This produces the **SNP-SNP Model report**.

### -P (--list-population-ids)

Lists the populations available in the database in use. Populations are used to adjust the gene boundaries to include additional SNPs that are observed to be within an LD threshold. By default, there should be 4 populations, with 3 to 6 different LD thresholds each.

## -d (--disease-dependent) filename

#### filename

This is a specially formatted file which contains user defined group information associated to disease specific knowledge. This is a command-line version of the configuration parameter, <u>Disease Dependent Groups</u>. A single run can have 0 or more disease specific files.

## -p (--print-count-estimates)

This flag forces the application to produce a Gene-Gene model report.

## --strip-optimization

This command causes the application to drop all of the Indexes. This can be done to speed up the insertions necessary during LD imports. Once the data has been properly imported, users should be sure to optimize the data once again.

## --optimize

This adds indexes to the underlying database, if they don't exist. This is generally only done once, but can be removed prior to performing LD import (see --strip-optimization above). Users generally shouldn't need to use this command.

## General Parameters

The following parameters control the basic behavior of the application through configuration options.

#### VARIATION\_FILENAME filename

```
VARIATION FILENAME variations.bn
```

This sets the path to the source file containing the binary variation data. This file should originally be downloaded with the application, however, as the data is updated, this file can be downloaded independently of the database. There is the possibility that the file is incompatible with the version of the biofilter in use. In such cases, the application will exit with an appropriate measure. To resolve this issue, the user should download the latest version of the application as well.

### SETTINGS\_DB filename

```
SETTINGS DB bio-settings.cn
```

This sets the path to the settings database. This file is required for all biofilter functionality and will periodically updated with new information. Users generally will be able to update their settings file without updating the application, however, in the event that major changes have been identified, the file require a newer version of the application. In this event, users will be notified and will have to update their software to use the newest data.

## MAX\_GENE\_COUNT integer

```
MAX GENE COUNT 30
```

This sets the upper limit for acceptable group size. Acceptable group size is the number of genes contained by a given group. If a group has as many or fewer genes in it, the biofilter uses it's contents to generate models. If a group exceeds this number, the biofilter queries each of it's children groups, performing the same check.

This setting is used to constrain the size of the resulting model counts to a number that is manageable according to modern computation resources.

## SNPS\_SOURCE filename

```
SNPS SOURCE /projects/ritchie/biofilter/Affy6.0-v27.txt
```

This setting allows the user to limit the SNPs considered to a specific platform (or some user-defined set of SNPs). By setting this value, the biofilter only loads SNP data for those found in the source.

## INCLUDE\_GROUPS group [group] [group] [...]

This allows the user to limit the search to those groups (and their children). This can be a specific group within a *meta-group* or it can be a meta-group ID (such as DIP). Each group is separated by spaces and is the group's unique ID. When including a group, all of that groups children are also included.

## INCLUDE\_GROUP\_FILE filename

```
INCLUDE_GROUP_FILE ../group_list.txt
```

This functions identically to INCLUDE\_GROUPS, however the source is a text file. This allows users to include a very large number of specially selected groups (and their children).

#### MODEL\_FILENAME filename

```
MDOEL FILENAME affy50-models.bin
```

This sets the output filename for the binary model file.

## DISEASE\_DEPENDENT string

```
DISEEASE DEPENDENT alz-bio-knowledge.txt
```

This instructs biofilter to load the knowledge associated with the file, alz-bio-knowledge.txt. A description of the format of these files can be found <u>here</u>.

## **POPULATION string**

```
POPULATION NO-LD
```

Setting the population for a given run allows the user to tap into expanded region boundaries associated with a particular LD cutoff and a given population. biofilter comes with a set of populations based on LD data found in hapmap, but users can contribute their own data. The string used as the parameter is one of a set of values known to the application. Users can query the application for a list of the valid settings by using the --list-populations flag.

## MODEL\_BUFFER\_INIT integer

```
MODEL BUFFER INIT 10000
```

The model buffer is an internal data structure which is used to maintain the massive amount of models generated without requiring huge amounts of RAM. This number is used to determine how large the buffer is at start. The buffer object will always have at least MODEL\_BUFFER\_INIT models in memory at one time, and can grow as large as MODEL\_BUFFER\_MAX + MODEL\_BUFFER\_INIT during the processing. For systems with 1 gigabyte of RAM or less should use the default values.

Increasing the init size will simply reduce the number of disk reads, and thus speed up the processing at the cost of increasing the actual memory foot print during processing. It is recommended that the init size be 1/4 the size of the max or smaller.

## MODEL\_BUFFER\_MAX integer

```
MODEL BUFFER MAX 100000
```

This number represents the size the buffer can get before triggering a disk write. Larger values will increase performance, and could reduce the number of disk reads by reducing the number of cycles required.

On 32bit Redhat intel systems, setting the MODEL\_BUFFER\_MAX to 20,000,000 allowed very fast processing of 500K sources with a memory footprint of 2.5 Gigabytes. The same setting for a million SNP source took 30 minutes and required 2.8 Gigabytes. Adding even more memory to the 64bit brought the run down to about 17 minutes

(500,000,000 MAX\_BUFFER). The same 1 million SNP source took over 2 hours at 4,000,000 but the footprint was around 1 gigabyte.

## MAX\_GENE\_COUNT integer

```
MAX_GENE_COUNT 30
```

During model production, the biofilter ignores groups that exceed this value. This is done to avoid generating too many models (which defeats the purpose, to some extent).

## PREFERRED\_ALIAS filename

```
PREFERRED ALIAS my favoriate genes.txt
```

The user specifies a list of gene aliases which they prefer. These aliases are used in place of the Ensembl ID in subsequent reports. See <u>formatting</u> details for help creating the file.

## LOAD\_ALL\_ALIASES YES/NO

```
LOAD ALL ALIASES NO
```

Loads all region aliases (gene names) and generates a report *report-prefix.aliases*. This makes it easier for the user to lookup the Ensembl IDs that are used by default in the reports.

## REPORT\_PREFIX string

```
REPORT PREFIX data/bio-alz
```

When a report is produced that is sent to file instead of std-out, it will use the value of REPORT\_PREFIX as the first part of the file name. Acceptable options can be anything that is acceptable for the filesystem except for whitespace (i.e. don't use spaces or tabs).

#### IMPLICATION\_IDX\_DUPLICATE\_WEIGHT float

```
IMPLICATION_IDX_DUPLICATE_WEIGHT 0.25
```

When computing implication index values, disease dependent groups grant only 1 point toward the implication index even if both genes are members of the one or of the disease dependent groups. The duplicate weight allows the user to nudge the relevance of a model where both genes are present in the same group(s) by multiplying the IMPLICATION\_IDX\_DUPLICATE\_WEIHT by the number of redundant disease dependent occurrences.

#### HTML REPORTS YES/NO

```
HTML REPORTS YES
```

When set to yes, most reports will be written in html format.

## DISEASE\_DEPENDENT\_LEVEL ALL\_MODELS/GROUP\_LEVEL/DD\_ONLY

```
DISEASE DEPENDENT LEVEL DD ONLY
```

Users can choose to filter models based on their association with disease dependent genes. When set to ALL\_MOD-ELS (default), all models will be produced. When set to GROUP\_LEVEL, only groups where gene-gene models are

produced that has one or more genes found in a disease dependent group will yield models. When set to DD\_ONLY, only the gene-gene models in which one of the genes is found in one of the disease dependent groups are produced.

## COLLAPSE\_ASSOCIATION\_REPORT YES/NO

COLLAPSE\_ASSOCIATION\_REPORT YES

When set to yes, the associations shown reflect only the groups which could generate models. The report properly respects the MAX\_GENE\_COUNT setting, but it does not obey the DISEASE\_DEPENDENT\_LEVEL setting (it shows all groupings, despite their relationship with disease dependent groups).

## BINARY\_MODEL\_ARCHIVE YES/NO

BINARY\_MODEL\_ARCHIVE YES

Setting this value to YES causes the gene-gene model archive and the snp-snp model archive to be written in binary format. In most cases, this won't be necessary, since both files will probably be reasonable in length. However, if either are expected to contain more than a few hundred thousand entries, it is recommended to use.

## ASSOCIATION\_REPORT YES/NO

ASSOCIATION REPORT YES

When Yes, this causes the biofilter to produce an association report.

# Input File Formats

With very few exceptions, files are space delimited ascii files.

## **Preferred Aliases**

This file provides one or more common gene names which can be substituted for Ensembl IDs in reports. Aliases must be alphanumeric (no spaces) and must match an alias known to the biofilter. Known aliases are those that were found in Ensembl's external synonyms associated with EntrezGene & Uniprot (TrEMBL and Swiss-Prot). Currently, only aliases which map back to a single Ensembl Gene ID are used.

This file is present only as a convenience for the user and is optional.

Example file:

NMT1 FURIN NRD1 S100B ATP2A2

# SNPS\_SOURCE

The SNP Source file contains all SNPs to be used in the analysis. Generally, this will match the SNPs from the platform to be used in the analysis. However, it is also possible to use a highly restricted set for other types of analysis (such as identifying which genes a set of interesting SNPs might be found in.)

The format is very simple. List all RS IDs in their integer format. Each ID should be separated by whitespace.

10000169 10000185 10000201 1000022 10000226 1000025 10000255 10000266

## **Disease Dependent Genes**

Users can tag genes as being disease dependent and add grouping information. Assigning genes to disease dependent groups causes two things to happen:

First, it allows the user to increment the implication index of disease dependent genes for every disease dependent meta-group they assign it to. Second, by providing groups of genes that have been known to be related to a given disease, it is possible to add models that represent interactions that a highly relevant to the disease that might not exist in more traditional grouping paradigms.

## **Disease Dependent Definition**

Users can create many disease dependent groupings. A high level grouping (or *meta-group*) will contain one or more *group*. Each of these *groups* will contain one or more regions. At present, the biofilter only looks at genes within a *group* for constituent models and there is no support for hierarchical groupings. However, users can simulate the effects of hierarchy by creating different disease dependent *meta-groups*, and correctly choosing the one appropriate for their needs. Models found inside multiple *groups* within a single *meta-group* will have their implication index incremented only by one. If a model occurs inside multiple *meta-groups*, it's implication index will reflect each *meta-group* it is found inside.

## Disease Dependent File Format

Disease dependent configurations are done outside of the application using a simple text file format. Each file represents a *super group* (*meta-group*) which can contain one or more *groups*.

## [Name] [Description]

The first line of the file must contain the disease dependent "name", a string with no spaces, followed by a short description (this can have spaces but must fit on a single line). The total length of the line should be 4096 characters or less. These are used by for reporting purposes and should be as meaningful as possible (i.e. if the user will have more than one meta-group for a single disease, they should properly name them so that they can be distinguished from one another.)

## GROUP [group name] [group description]

**GROUP** This keyword is required and should be correctly capitalized.

**Group Name** String with no spaces or tabs inside. This is used for reporting purposes and should be meaningfully distinguishable from any other groups.

**Group Description** Simple description for the group. This can have spaces, but must fit on the line with a total length of 4096 or less.

gene alias [gene alias]

Gene aliases are gene identifiers from one of the following sources: Uni-Prot IDs (TREMBL or SwissProt), Entrez Gene. These aliases must only identify a single entity, have no spaces in them, and must be available from within the Ensembl build upon which the biofilter data was based. Aliases can appear on separate lines or with spaces/tabs separating them or a mix of the two.

ALZHEIMERS GROUP alz-assoc Genes previously recognized through association studies AGT APH1A APOA1BP APOA2 CAMK1G CFH CHRNB2 CLCNKB
---

## **Model Production**

#### Overview

The Biofilter uses biological information about gene-gene relationships and gene-disease relationships to construct multi-SNP models before conducting any statistical analysis. Rather than annotating the independent effect of each SNP in a GWAS dataset, the Biofilter allows the explicit detection and modeling of interactions between a set of SNPs. In this manner, the Biofilter process provides a tool to discover significant multi-SNP models with non-significant main effects that have established biological plausibility. This approach has the added benefit of reducing both the computational and statistical burden of exhaustively evaluating all possible multi-SNP models.

Model production is gene centric, and thus requires that any SNPs to be considered be mapped to genes. The gene mapping takes place internally using local copies of current data sources such as Ensembl, HapMap and dbSNP. A structured mapping is made based on relationships from one of the knowledge sources and this information is used to identify candidates for snp-snp models.

The biological knowledge used by the Biofilter is derived from various sources which are identified as Meta Groups as well as optional user defined groupings. Currently, the data-sources represented include: Gene Ontology, KEGG, Net Path, pfam, Reactome and DIP. These sources are updated periodically, and made available as updates at the biofilter website.

There are two distinct types of data sources. Disease-dependent sources are user defined and reflect gene-disease associations. Disease independent sources represent key relationships between genes in important biological processes such as a metabolic pathway. (Bush 2009)

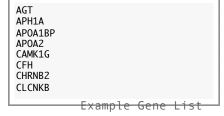
Users provide a set of SNPs that reflect the platform on which their analysis will be run. This can be a GWAS platform such as Illumina Human 1M-DUo BeadChip or one designed for the user's specific study. Biofilter requires only the rsNumber. It uses it's own copy of SNP data to insure that positional information matches it's internal representation for gene mapping. As a result, only those SNPs available in the Biofilter's local copy will be considered.

# Reports

# **Coverage Report**

The coverage report shows the amount of coverage a set of genes has on one or more platforms. The report requires 2

pieces of information: A single gene list, and 1 or more coverage lists. Depending on the settings chosen, the report might be plain text or in HTML format. As with any other HTML formatted report, users should be aware of the potential size of the report they are generating, since a huge report will be difficult to open in most browsers.



**Gene List:** The gene list is a series of gene aliases listed on individual lines. These gene aliases must be UniProt IDs or entrez genes and must not be defined such that they represent more than one ensembl ID.

**Coverage Files:** The user should provide one or more coverage files, which are just the individual RS IDs that can be found on a platform of interest. By default, the biofilter comes with 4 different platform files, but the user can use any list they wish, as long as the file contains only integer representation of the RS ids found on the platform. Any file that can be used as a <u>SNPS\_SOURCE</u> file can be used as a coverage file.

The baseline coverage is based on the SNPs found using the regular <u>SNPS SOURCE</u> value (or -s). This is treated as the total. Additional coverage sources are displayed as additional columns. Each entry shows the number of SNPs covered by that platform for a given gene.

A coverage report might look as follows:

```
biofilter\ sample.config\ -s\ Illumina-660Quad.txt\ --report-gene-coverage alz-genes.txt\ -C\ Illumina-660Quad.txt
```

Gene	Ensembl_id	Total	Illumina-660Quad.txt(593544)	
AGT	ENSG00000135744	78	78	
APH1A	ENSG00000117362	11	11	
APOA1BP	ENSG00000163382	5	5	
AP0A2	ENSG00000158874	12	12	
CAMK1G	ENSG00000008118	22	22	
CFH	ENSG00000000971	82	82	
CHRNB2	ENSG00000160716	1	1	
CLCNKB	ENSG00000184908	18	18	
CLCNND	EN2000000104900	10	18	

The gene alias is shown in the first column, followed by the ensembl\_id. The total represents the number of SNPs found in the <u>SNPS SOURCE</u> file and is followed by the SNP count for each of the coverage files. Using the option, --detailed-coverage, lists all RS IDs associated with the genes (and their position for the text report.)

Additional coverages can be added using additional -*C filename* options. Below is the same report in HTML format using --detailed-coverage. Ensembl IDs and RS Numbers provide links to ensembl's website for convenience.

Gene	Ensembl ID	Chromosome	Begin(kB)	End(kB)	Total	Illumina-660Quad.txt(593544)	SNP
AGT	ENSG00000135744	1	228777551	228945111	78	78	rs740490 rs740492 rs11122545 r rs11122547 rs853459 rs200584 r
APH1A	ENSG00000117362	1	148432473	148515725	11	11	rs3118124 rs11588837 rs112053 rs12086155 rs11205347 rs10888
APOA1BP	ENSG00000163382	1	154820731	154863290	5	5	<u>rs4661188</u> <u>rs11264505</u> <u>rs120234</u>
APOA2	ENSG00000158874	1	159458707	159489274	12	12	rs6413453 rs4233368 rs4073054 rs2307425 rs2501873 rs6686001
CAMK1G	ENSG00000008118	1	207793089	207874438	22	22	rs10746420 rs7525534 rs111193 rs11587591 rs7528698 rs669055
CFH	ENSG00000000971	1	194718611	195171294	82	82	rs12065463 rs6428342 rs133995 rs10754192 rs10494744 rs42943
CHRNB2	ENSG00000160716	1	152806881	152818975	1	1	rs3811450
CLCNKB	ENSG00000184908	1	16240720	16272971	18	18	rs10803410 rs6683445 rs868950 rs12015135 rs10803414 rs12047
*Boundaries	for genes are based o	n: CEU Popula	tion. DPrime	cutoff of 0.	80.		

# **Model Summary Report**

During model generation, a report is made containing all gene-gene pairs that were used to generate SNP-SNP models. This report contains SNP counts for each gene, the groups in common between the two genes and the number of models produced by the pairing. By default, Genes are reported using their Ensembl Stable ID. However, users can configure a <u>Preferred Gene Alias</u> file. Genes which appear in the preferred alias file will be reported according to the preferred alias.

	Gene-G	ene Models				
Gene	SNP	Gene	SNP	Impl.	Models	Groups
Name	Count	Name	Count	Index	Count	DI,DD
ABCG1	7	ACVR2B	7	2 2 2	49	1,485522
ABCG1	7	ABCB1	58		406	1,485522
ABCG1	7	CYP3A4	37	2	259	1,485522
ABCG1	7	SLC22A11	7	2	49	1,485522
ABCG1	7	SCUBE3	47	2 3	329	1,485522
ABCG1	7	ANXA9	6	3	42	112,485523
ABCG1	7	SULF1	3	2 2	21	1,485522
ABCG1	7	STX4	17	2	119	1,485522
ACVR2B	7	ABCB1	58	2	406	1,485522
ACVR2B	7	CYP3A4	37	1	259	1,
ACVR2B	7	SLC22A11	7	1	49	1,
ACVR2B	7	SCUBE3	47	1	329	1,
ACVR2B	7	ANXA9	6	2	42	1,485523
SULF1	3	ACVR2B	7	1	21	1,
STX4	17	ACVR2B	7	1 2	119	1,
ABCB1	58	CYP3A4	37	2	2146	1,485522
SLC22A11	7	ABCB1	58	2	406	1,485522
SCUBE3	47	ABCB1	58	2	2726	1,485522
		(truncated)				

## **Assocations List**

Suffixed by "associations.txt" or "associtions.html", association reports provide the user with a hierarchical representation of groups in their search. Only groups that have been loaded using INCLUDE\_GROUPS (or all, if no groups were provided) will appear.

biofilter sample.config --list-associations

```
Associations (Gene Ontonology):
     G0:0048154(1)
     $100B ( rs2839357 rs2839362 )
G0:0048155(1)
          ATP2A2 ( rs3026445 rs3026457 rs1860561 )
     G0:0048156(1)
          $100B ( rs2839357 rs2839362 )
     G0:0042982(2)
          G0:0042987(4)
               ABCG1 ( rs4148083 rs4148084 rs4148085 rs9975740 rs4148087 rs1117640 rs4148088 rs4148089
rs4148090 ..
               PSÉN1 ( rs214273 rs8006497 rs362350 rs214260 rs165933 rs362377 )
               ENSG00000162736 ( rs10494342 rs16831846 rs12239946 rs6664438 rs6677637 ) ENSG00000143801 ( rs2073489 rs1295640 )
          ENSG00000167755 ( rs1654537 )
                                                           (truncated)
Associations (KEGG):
     KEGG-05010(127)
          ENSG00000015475 ( rs181390 rs181396 rs181402 rs181405 rs9604787 rs181408 rs181417 rs5746474
rs5747351 rs9605401 rs738095 )
          ATP2A2 ( rs3026445 rs3026457 rs1860561
          PSEN1 ( rs214273 rs8006497 rs362350 rs214260 rs165933 rs362377 )
          NCSTN ( rs10494342 rs16831846 rs12239946 rs6664438 rs6677637 )
PSEN2 ( rs2073489 rs1295640 )
          BACE1 ( rs7083 rs522843 rs687740 rs473210 rs551662 rs676134 )
ENSG00000132906 ( rs6685648 rs2020902 rs4646018 rs2042370 rs1862710 rs4645989 rs1052571 )
NDUFV2 ( rs4148964 rs11081459 rs4148965 rs1039825 rs1472944 rs977581 rs874250 rs4148966 rs4148967
rs4148968 rs12966444 )
                                                            (truncated)
Associations (NetPath):
Associations (pfam):
Associations (Reactome):
```

Each relationship is nested with tabs on separate lines. In the example above, GO:0042987 has some child relationship to GO:0042982. The numbers beside a group ID indicate the number of genes associated with the group.

An optional setting, **COLLAPSE\_ASSOCIATION\_REPORT**, can be set to true to collapse groups where models would be generated. This allows the user to see more clearly which genes will be combined to produce models.

# **Disease Dependent Contents Report**

Suffixed with dd-contents, this report is produced when the user includes disease dependent information. This report is provided to allow the user to verify that the genes being used are the same as the genes were intended.

Alias	Ensembl ID	ID (	Chrom	Start Pos.	Stop Pos.	Snp Count
AGT APH1A APOA1BP APOA2 CAMKCF CFH CHRNB2 CLCNKB	ENSG00000135744 ENSG00000117362 ENSG00000163382 ENSG00000158874 ENSG00000008118 ENSG0000000971 ENSG00000160716 ENSG00000184908	98998 96653 97981 98229 98856 98642 97575 78893	1 1 1 1 1 1 1	228777551 148432473 154820731 159458707 207793089 194718611 152806881 16240720	228945111 148515725 154863290 159489274 207874438 195171294 152818975 16272971	78 11 5 12 22 82 1

# **SNP Report**

SNP reports provide information about the genes for which they are associated. If the user has selected LOAD\_ALL\_ALIASES YES in their configuration, a list of corresponding aliases will also be provided. This report is available as plain text and HTML format. In the HTML format, the RS Numbers and gene IDs will appear as links to the corresponding page at the ensembl website. As with any other HTML document, users should be aware of the potential size of their report before choosing HTML due to potential memory problems.

biofilter sample.config --snp-report

	ı: 🥢 İntro	duction to makin 🧳 PythonSpeed/Perform 🧳 http://drp.rubyforge.
RS ID	Chrom.	Region Name
10000010	4	ENSG00000185774 80333,KCNIP4
10000023	4	ENSG00000138696 658,BMPR1B
10000030	4	ENSG00000138821 64116,SLC39A8
10000092	4	ENSG00000185774 80333,KCNIP4
10000160	4	ENSG00000198092 389208,FLJ16046,TMPRSS11F
10000100	4	ENSG00000215127 401135,SYT14L
10000169	4	ENSG00000138771 57619,SHROOM3
10000185	4	ENSG00000184985 57537,SORCS2
1000022	13	ENSG00000125246 171425,CLYBL
10000226	4	ENSG00000163629 5783,PTPN13
10000220	4	ENSG00000145283 345274,SLC10A6
10000255	4	ENSG00000168843 56884,FSTL5

# **SNP-SNP Model Report**

The SNP-SNP Report lists the details for each model from a previously generated model file. Details include:

• Left/Right SNPs that make up the model.

• The Group(s) in which the pair of Loci were found

• The Genes from which each of the SNPs were found.

9 rs1805488	rs2072539 rs2072539 rs176590882
-------------	---------------------------------------

When possible, each of elements reported will reflect the highest form of the name provided by the user. At the very least, Ensembl IDs will be used. However, if the user specifies gene "Aliases", those will be used preferentially over the less familiar Ensembl ID.

SNP 1	Gene	Groups	SNP 2	Gene	Groups
2072539	GRIN2B	KEGG-05010	1805488	GRIN2B	KEGG-05010
2072539	GRIN2B	KEGG-05010	1990310	ENSG00000139180	KEGG-05010
176590882			17701871	ENSG00000139180	KEGG-05010

The excerpt to the left shows an example of what might be produced by such a report. Each snp is listed with it's corresponding genes provided with a link to ensembl.

The text report is similar except the SNPs are listed together as can be seen in the example below.

SNPS	Genes Groups	Genes	Groups
[ 2072539 1805488 ]	GRIN2B KEGG-05010	GRIN2B	KEGG-05010
[ 2072539 1990310 ]	GRIN2B KEGG-05010	ENSG00000139180	KEGG-05010
[ 176590882 1770187	1 ]	ENSG00000139180	KEGG-05010

## **Gene-Gene Models**

This isn't a report, but a data product which can be passed to applications which have been linked to the biofilter library and have support for gene-gene models. However, when the setting BINARY\_MODEL\_ARCHIVE is off, this is written in plain text, allowing the user to see exactly what models their run produced.

The first line contains the number of models. Each subsequent line contains the 4 columns: Gene ID 1, Gene ID 2, Implication Index, Disease Independent groups associated with the pairing. The Gene IDs are the numerical ID value (this is the second column from the genes file). This file requires the .genes output file in order to be used to fully expand gene-gene models into snp-snp models.

The file is sorted by implication index, such that those models with the highest score will be first.

```
11925
          55016
54991
                     4
                                1 | 2 | 4 | 9
75847
          91412
                     4
                                1 | 2 | 4 | 9
75837
          91412
                    4
                               1 | 2 | 4 | 9
75837
          75847
                               1 | 2 | 4 | 9
                    4
64546
          91412
                     4
                               1 | 2 | 4 | 9
64546
          75847
                     4
                               1 | 2 | 4 | 9
64546
          75837
                               1 | 2 | 4 | 9
. . . .
```

# **Genes Report**

This is actually part of the gene-gene model output and is a text file which contains the information required to associate genes with their contents (SNPs) and their potential contributions toward implication index.

Each column is separated by a tab. For compound fields such as SNP lists and group IDs, constituent members are separated by a "|" character. There are 5 Columns (not all will be filled): Alias, Gene ID, SNPs, Disease Independent groups, Disease Dependent Groups.

TYMS	44549	596909 3786362 11540152 11540153 2 4	9	360684 360685
ENOSF1	44559	596909 3786362 11540152 11540153 1 4		360685
YES1	44568	596909 3786362 11540152 11540153 1 2	4 9	
TWSG1	44857	3322 10898 12680 11559053 28552921 35867116		4
RALBP1	44861	3322 10898 12680 11559053 28552921 35867116		1 2 4 9
PPP4R1	44868	3322 10898 12680 11559053 28552921 35867116		1   4
CHST9	45509	417808 418546 1155514 17694469 28693844 1 4		
0XA1L	45871	1061040 1805059 1805061 2281677 8016634 801	8462	1   2   4
SLC7A7	45894	1061040 1805059 1805061 2281677 8016634 801	8462	1   4   9
MRPL52	45927	1805059 1805061 2281677 8016634 8018462 1		
MMP14	45945	1061040 1805059 1805061 2281677 8016634 801	8462	1   2   4

# **SNP Cleanup Report**

After loading the SNPs from the SNPS\_SOURCE file, the biofilter attempts to identify any RS IDs that have been merged into previous RS IDs by dbSNP. Any RS IDs that have been merged will be replaced with the proper RS ID. Those SNPs that have been identified as having been deleted by NCBI are removed. Each removal and renaming is noted in the file, *projectname.snp-cleanup*.

This file is a tab separated file listing first the SNPs that were removed due to having been deleted at NCBI followed by the mapping details for those that have been merged.

```
4 Expired SNPs Encountered:
       45469397
       rs45552437
       rs45616434
       rs45628831
504 rs IDs were updated
       Original ID
                      New ID
       rs41429248
                       rs2305130
       rs41480744
                      rs17145687
       rs12782608
                       rs9422653
       rs41396045
                      rs36130286
        rs41495349
                       rs16865746
        rs4365706
                       rs4026962
        rs11242845
                       rs9501985
        rs41503946
                       rs11077998
        rs4446752
                       rs2314691
                                    (truncated)
```

# Output Control (TBD)

# Example Run - Alzheimer's

The following tutorial will walk the user through performing many typical biofilter tasks including preparing a highly specialized model set to use with plato for performing an association study using knowledge specific to Alzheimer's disease. This search is intentionally highly selective and is intended for instructional purposes only. Under most circumstances, users will want to incorporate more knowledge than we will be including in this example.

The examples listed below use the following command to execute the biofilter, *biofilter*. This command may differ from machine to machine depending on hardware. This is done to allow different versions of the software to exist side by side. In general, 32bit linux distributions will simply be called biofilter. When compiled for 64bit systems, the name will be biofilter64. For OSX and windows, the name will be biofilter-OSX and biofilter-win32, with the possible 64 following the word biofilter when built for 64bit.

## **Listing Options**

If you are completely new to the biofilter, a good first step is to take a quick look at the various options available. As with many Unix application, simply running the program with no parameters will generate a basic list of options.

\$ biofilter

```
biofilter 0.5.0 (600) Debug Tue Sep 1 16:07:51 CDT 2009
Marylyn Ritchie, William Bush and Eric Torstenson
Please forward any comments or errors to <a href="mailto:biofilter@chgr.mc.vanderbilt.edu">biofilter@chgr.mc.vanderbilt.edu</a>
usage: biofilter <configuration file>
biofilter is a standalone application for use in investigating possible SNP associations
    in a set of data which, through biological knowledge, might be worth investigating
Optional Commands Include:
     -S [--sample-config]
                                                       -- Print sample configuration to std-out
     --report-gene-coverage gene-list-filename -- Reports the snp count for the genes in genelist
                                                       -- for the snps in snp-source
                                                      -- Reports each SNP and it's position/chromosome
     --marker-info
                                                      -- in a format acceptable by haploview
-- For each SNP in the SNP Source, lists the genes
     --snp-report
                                                      -- where that RS number is found
     --list-associations
                                                      -- Lists the associations for each group
Optional Parameters Include:
    -s [--snps] <snps filename>
-C [--coverage] <snps filename>
                                                      -- Override the snp source file ont he commandline
                                                      -- Add a file to coverage report list
    -D [--detailed-coverage]
-L [--list-models]
                                                      -- (used with -C) adds extra details to coverage report
-- Writes model list to std-out
        [--write-models] <model filename>
                                                      -- Writes model list to file (in binary format)
     -m [--show-models] <model filename>
                                                      -- Writes contents of model file to screen in human
                                                      -- readable form
    -p [--print-count-estimates
-l [--load-ld] <model filename>
                                                      -- Lists count estimates for gene - gene models
                                                      -- Loads LD information from the file, filename, and
                                                      -- adjusts the gene boundaries accordingly
                                                      -- Adds a meta group containing data from the file, filename
-- Adds group search criteria and produces a list of
    -d [--disease-dependent] <filename>
-G [--list-groups] [criteria]
                                                      -- group IDs that match the criteria
                                                      -- Lists all available Population based LD boundary options
     -P [--list-populations]
     --optimize
                                                      -- Updates internal structures to allow faster access. This
                                                      -- is usually done prior to release
     --strip-optimization
                                                      -- Strips the optimization out (this is helpful to allow data
                                                       -- imports to run more quickly)
```

## **Generating Sample Configuration**

Users can use biofilter to generate a skeleton sample configuration. Most command line options will be incorporated. Users should edit this configuration with appropriate settings. Filenames follow standard unix convention for location. This allows them to be expressed with full paths as well as paths relative to the directory in which the application is run (this includes just using the filename, if the file exists in the same directory as the application was run.) Fields that have no default value will be commented out in the example configuration. In some cases, such as SNPS\_SOURCE, a value is required before certain types of execution can be made.

```
$ biofilter --sample-config > sample.config
```

```
# Variations data
VARIATION_FILENAME
                       variations.bn
# BioFilter data
                       bio-settings.cn
SETTINGS_DB
# Max number of genes before we ignore the group
MAX_GENE_COUNT
                       30
# The source file for the RS numbers in your dataset
#SNPS_SOURCE
# List the various groups (by group name) separated by spaces
#INCLUDE_GROUPS
# Set the filename for the output model list (none writes to std-out)
MODEL_FILENAME
                       NONE
# Set the initial size of the model buffer.
MODEL_BUFFER_INIT
                      10000
# Set the upper limit to the buffer. Bigger -> faster, but must remain within
# the limits of the hardware or could cause the application
# to fail or become so slow that it will never complete.
MODEL_BUFFER_MAX
                       100000
# Set the population ID to match the population your data is drawn from so that
# LD patterns can be used to expand the gene boundaries.
# Add one or more files containing disease dependent genes
#DISEASE_DEPENDENT
\mbox{\#} User can specify aliases for genes (the alias must be present in the database \mbox{\#}\mbox{PREFERRED\_ALIAS}
# Prefix used for all reports
#REPORT_PREFIX
# Loads all aliases and generates a text report containing their associations
LOAD_ALL_ALIASES
                                                 sample.config
```

Many configuration options also have a command line override available. This is useful for performing tasks which are repeated from time to time.

## **General Reporting**

Except when generating a sample configuration, a summary report is produced containing the configuration details being used along with some details about the database in use.

```
Dependency Versions -----dbSNP: 36
Ensembl: 27
        Hap Map LD: 53
---- Configuration Parameters -----
  DISEASE_DEPENDENT : alz.txt
INCLUDE_GROUPS : 53887,5
                         53887,53892,53894,101057,101064,106734,160871
    LOAD_ALL_ALIASES
      MAX_GENE_COUNT
                         30
   MODEL_BUFFER_INIT
                         10000
                         100000
    MODEL_BUFFER_MAX
      MODEL_FILENAME
                         NONE
                         CEU-DP0.80
           POPULATION
     PREFERRED_ALIAS
                         gene_aliases.txt
       PROJECT : REPORT_PREFIX :
                         sample.config
                         tutorial
          SETTINGS_DB :
                         bio-settings.cn
 SNPS_SOURCE : VARIATION_FILENAME :
                         Illumina-660Quad.txt
                         variations.bn
```

The first three lines describe the source versions used in the local database. These refer to the versions from which the data was captured. The remaining lines represent the various configuration options in use.

For the purposes of our example, let's make a few changes. The following represent some changes that we might make:

## **Target Platform**

#### Whenever we generate models, we need to tell the biofilter what SNPs exist on that platform.

Users can specify a platform using either configuration changes or on the command line. For this example, we'll make a single change to the configuration file.

Open the file and edit the line that contains SNPS\_SOURCE. Remove the '#' sign and add an appropriate filename after the command. It should look something like the following:

```
# The source file for the RS numbers in your dataset
SNPS_SOURCE Illumina-660Quad.txt
```

The file Illumina-660Quad.txt is just a list of RS Numbers (without the letters 'RS') in a simple ascii file that represent each of the SNPs on our platform. In this case, the list was extracted from documentation on Illumina's 660 Quad platform. When specified as we have (along with no path information), the application assumes that the file is located in the same directory as the application is run.

## **Gene Aliases**

In general, the biofilter will supply the gene's Ensembl ID for all gene names. This ID is stable and has a single meaning. However, most users will prefer to see IDs listed in more familiar terms. Providing a list of familiar aliases allows the application to use a known synonym for the gene without it having to make any assumptions for the user. For more information on the format of this file, please see the <u>documentation</u> on gene aliases.

```
NMT1
FURIN
NRD1
S100B
ATP2A2
ABCG1
PSEN1
-gene aliases.txt —
```

For the purposes of example, we'll use a really small one with some genes we know will be in our report. This file is called, gene\_aliases.txt. To use these aliases, edit your configuration file and change the line containing PRE-FERRED\_ALIAS. Remove the '#' from the beginning of the line and add the filename to the end of the line. It should look something like what you see below.

```
# User can specify aliases for genes (the alias must be present in the database
PREFERRED_ALIAS gene_aliases.txt
```

#### Selective Search

The biofilter contains a large amount of information, much of which might be completely outside of the user's interest. Users can select as much or as little as they want to use. We'll be restricting the search to include only groups that have Alzheimer's listed in the comments. This is probably not a very good example, since it will only capture a tiny amount of knowledge, but for the purposes of our example, it will do just fine.

Biofilter gives the user the ability to query it's built-in group information. To get a list of groups that contain the letters 'alz' in the name or description field, type the following:

biofilter sample.config -G alz

```
Meta Group Group IDNameDescription
Gene Ontonology53887G0:0048154 "Interacting selectively with S100 beta protein. S100 is a small calcium and zinc binding protein produced in astrocytes that is implicated in Alzheimer's disease, Down Syndrome and ALS." [GOC:jic]
Gene Ontonology53892G0:0048155 "Interacting selectively with S100 alpha protein. S100 is a small calcium and zinc binding protein produced in astrocytes that is implicated in Alzheimer's disease, Down Syndrome and ALS." [GOC:jic]
Gene Ontonology53894G0:0048156 "Interacting selectively with tau protein. tau is a microtubule-associated protein, implicated in Alzheimer's disease, Down Syndrome and ALS." [GOC:jic]
Gene Ontonology101057G0:0042982 "The chemical reactions and pathways involving amyloid precursor protein (APP), the precursor of beta-amyloid, a glycoprotein associated with Alzheimer's disease." [GOC:go_curators]
Gene Ontonology101064G0:0042987 "The chemical reactions and pathways resulting in the breakdown of amyloid precursor protein (APP), the precursor of beta-amyloid, a glycoprotein associated with Alzheimer's disease."
[GOC:go_curators]
Gene Ontonology106734G0:0050435 "The chemical reactions and pathways involving beta-amyloid, a glycoprotein associated with Alzheimer's disease, and its precursor, amyloid precursor protein (APP)." [GOC:ai]
KEGG160871KEGG-05010Alzheimer's disease
```

Users can perform more selective searches by adding additional -*G keyword* phrases to the command-line. The more keywords you add, the more selective the search will be.

The second column contains the group IDs, which is what we need to update our configuration file so that it only includes the groups we are interested in. Open the configuration file and change the line containing "IN-CLUDE\_GROUPS" so that it looks similar to the text below:

```
# List the various groups (by group name) separated by spaces INCLUDE_GROUPS 53887 53892 53894 101057 101064 106734 160871
```

Notice that groups are separated by spaces, not commas.

When users specify no groups, all groups are used. However, if one or more groups are added to this line, only those groups and any that are hierarchically contained within them are used to generate models. Users can include an entire group by using the group ID associated with the top level group (such as the ID for GO.)

## **Report Prefix**

One last detail is the report prefix. By default, any text output that doesn't go to the command line will be named after the configuration file. However, users can override this behavior by providing a Report Prefix. For our purposes, we'll simply change the reports to start with the word tutorial.

# Prefix used for all reports
REPORT\_PREFIX tutorial

#### **Assocations List**

Users can visualize the layout of the grouping information by generating the Association Report. This is a hierarchical view of the data associated with the requested groups, genes and SNPs that will make up the requested model summary. The output of the associations list is written to a file, ending with "-associations.txt" or "-associations.html".

biofilter --list-associations

```
Associations (Gene Ontonology):
     G0:0048154(1)
         S100B ( rs2839357 rs2839362 )
     G0:0048155(1)
         ATP2A2 ( rs3026445 rs3026457 rs1860561 )
     GO:0048156(1)
         S100B ( rs2839357 rs2839362 )
     G0:0042982(2)
         G0:0042987(4)
              ABCG1 (rs4148083 rs4148084 rs4148085 rs9975740 rs4148087 rs1117640 rs4148088 rs4148089
rs4148090 ...
              PSÉN1 ( rs214273 rs8006497 rs362350 rs214260 rs165933 rs362377
              ENSG00000162736 ( rs10494342 rs16831846 rs12239946 rs6664438 rs6677637 ) ENSG00000143801 ( rs2073489 rs1295640 )
         ENSG00000167755 ( rs1654537 )
                                                       (truncated)
Associations (KEGG):
    KEGG-05010(127)
         ENSG000000015475 ( rs181390 rs181396 rs181402 rs181405 rs9604787 rs181408 rs181417 rs5746474
rs5747351 rs9605401 rs738095
         ATP2A2 ( rs3026445 rs3026457 rs1860561
         PSEN1 ( rs214273 rs8006497 rs362350 rs214260 rs165933 rs362377 )
         NCSTN ( rs10494342 rs16831846 rs12239946 rs6664438 rs6677637 )
         PSEN2 ( rs2073489 rs1295640 )

BACE1 ( rs7083 rs522843 rs687740 rs473210 rs551662 rs676134 )

ENSG00000132906 ( rs6685648 rs2020902 rs4646018 rs2042370 rs1862710 rs4645989 rs1052571 )
         NDUFV2 ( rs4148964 rs11081459 rs4148965 rs1039825 rs1472944 rs977581 rs874250 rs4148966 rs4148967
rs4148968 rs12966444 )
(truncated)
Associations (NetPath):
Associations (pfam):
Associations (Reactome):
Associations (DIP):
```

There is a lot there, even though we included only 7 groups. Many of the GO groups had one or more child groups, and the KEGG group we added had a large number of genes associated with it (127 to be exact).

If we were to use all 127 genes to generate models, the resulting model count could be tremendous, depending on the numbers of SNPs in each of those genes. If you look through the report as generated by the biofilter, you will see that some have quite a few SNPs.

This problem brings up the value in one of the configuration options. In the configuration file, you will find a line similar to the following:

```
# Max number of genes before we ignore the group
MAX_GENE_COUNT30
```

While the comment might suggest that we ignore the group, it's a bit misleading. The application won't actually create models with any group larger than 30. However, it will attempt to traverse any child groups and consider producing models with those (if they have 30 genes or less).

In the case of the KEGG group above, there are no child groups-so, that group will not yield any models with our current setting. Should you want to ensure that the group is used, the variable above can be set high enough to catch the group. Users should make these types of changes very carefully, though. Setting the threshold too high could result in massive delays. In order to maintain large lists, the bilfilter uses a file cache to keep up with the models as they are generated. As this cache becomes really large, it can alter the runtimes dramatically.

## Population Selection and LD

The database that holds group information also contains information about regions (genes). Users can select a population and an LD threshold to extend gene boundaries during model generation. This allows the system to potentially capture SNPs that might carry a signal but are missing from the platform. This LD information is drawn from the hapmap project.

To get a list of populations and their LD thresholds, use the --list-populations (-P).

biofilter sample.config -P

```
Label Comment
NO-LDNo LD
YRI-RS1.00 YRI Population. RSquared cutoff of 1.00
YRI-RS0.80 YRI Population. RSquared cutoff of 0.80
YRI-RS0.70 YRI Population. RSquared cutoff of 0.70
YRI-DP1.00 YRI Population. DPrime cutoff of 1.00
YRI-DP0.80 YRI Population. DPrime cutoff of 0.80
YRI-DP0.70YRI Population. DPrime cutoff of 0.70
CEU-RS1.00 CEU Population. RSquared cutoff of 1.00
CEU-RS0.80CEU Population. RSquared cutoff of 0.80
CEU-RS0.70CEU Population. RSquared cutoff of 0.70
CEU-DP1.00 CEU Population. DPrime cutoff of 1.00
CEU-DP0.80CEU Population. DPrime cutoff of 0.80
CEU-DP0.70CEU Population. DPrime cutoff of 0.70
CHB-RS1.00 CHB Population. RSquared cutoff of 1.00
CHB-RS0.80CHB Population. RSquared cutoff of 0.80
CHB-RS0.70CHB Population. RSquared cutoff of 0.70
CHB-DP1.00 CHB Population. DPrime cutoff of 1.00
CHB-DP0.80CHB Population. DPrime cutoff of 0.80
CHB-DP0.70CHB Population. DPrime cutoff of 0.70
JPT-RS1.00 JPT Population. RSquared cutoff of
JPT-RS0.80 JPT Population. RSquared cutoff of 0.80
JPT-RS0.70JPT Population. RSquared cutoff of 0.70
JPT-DP1.00JPT Population. DPrime cutoff of 1.00
JPT-DP0.80JPT Population. DPrime cutoff of 0.80
JPT-DP0.70JPT Population. DPrime cutoff of 0.70
```

By default, the system uses NO-LD, which interprets a gene explicitly by it's beginning and end base pair location.

The procedure used to extend the boundaries is called LD-Spline, a technique developed by the Ritchie Lab. The cutoff mentioned in the report above is the minimum LD statistic (R-Squared or D') a pair can have to extend the region's boundaries.

Our study most closely matches the CEU population, so we'll use one of those options. We'll let the system be quite liberal in the definition of a region's boundaries, so we'll choose the middle D' option, 0.80. To indicate to the biofilter that a particular population and LD threshold is to be used, the configuration file must be edited once again. Find the line that starts with POPULATION and change NO-LD to CEU-DP0.80.

```
# Set the population ID to match the population your data is drawn from so that
# LD patterns can be used to expand the gene boundaries.
POPULATIONCEU-DP0.80
```

It is important to realize that the populations listed are the only ones available in the database that is in use. Users can process their own LD using thresholds of their own choosing. However, that is beyond the scope of this tutorial.

## **Defining Disease Dependent Groups**

When building a model list, users have the option of adding in their own knowledge into the system. This is done through the use of Disease Dependent groups. A disease dependent group functions similarly to one of the larger disease independent ones (such as KEGG and GO).

To create a small example disease dependent group, create a text file named alz.txt containing the following lines:

```
ALZHEIMERS
GROUP alz-assoc Genes previously recognized through association studies
AGT
APH1A
APOA1BP
APOA2
CAMK1G
CFH
CHRNB2
CLCNKB
```

The first line is the name of the meta group name and is used to identify the various groups associated with this disease dependent set. The next line defines an actual group. Each group definition must start with the keyword, GROUP followed by it's name and some descriptive commentary. Each group should have a unique name, since it will be used in the reporting. As in all other cases, names must contain no spaces. However, the description can contain any character the user prefers, except for new line characters.

Following the group definition are the gene names. These should be common names, and must appear in the list of known aliases. In general, those names should be recognized by EntrezGene or Uniprot and must identify a single gene. Gene aliases should be separated by whitespace, however, they must start on the line after the group definition.

We could add more groups to this file using addition GROUP definitions, but this is sufficient for our needs.

Adding the disease dependent information to a run can be done using either a configuration file, or on the command line. We'll add it to the configuration file here. Open the configuration and add the new filename to the end of the line starting with #DISEASE\_DEPENDENT. Be sure to remove that '#' as well, or else the line will be ignored.

```
# Add one or more files containing disease dependent genes
DISEASE_DEPENDENT alz.txt
```

## **Model Generation**

We are now ready to produce our model list. The most important step is the production of a gene-gene model list. This is a complete listing of all gene-gene models that were defined based on the Disease Independent information in our database, and the disease specific information provided by the user. This model list can be passed directly to some programs (such as the application, *athena*) or it can be used to produce a selective list of snp-snp models. Both of these files can be written as text or in a binary format. The gene-gene models are sorted by Implication Index such that those with the highest score are at the beginning of the file.

To produce a gene-gene model file, simply pass the argument -W on the command line. This option can take 2 optional parameters: Minimum Implication Index and the Maximum SNP-SNP model count. These determine the number and quality of SNP-SNP models to be generated immediately upon completion of generating the Gene-Gene model file. The user must provide both values or neither.

For our purposes, let's produce up to One million models with the worst Implication Index of 2.0.

biofilter sample.config -W 2.0 1000000

Illumina-660Quad.txt : 59	2652 SNPs		(593548	matches i	n our da	tabase	)
Group G	roup ID	Group-Count	Gene-Count				
Gene Ontonology KEGG	1 2	6305 204	10673 4640				
NetPath pfam Reactome	3 4 9	21 3904 4024	161 16261 675				
DIP Disease Dependent (alz.txt ALZHEIMERS)	12	1310	1045				
alz-assoc	176647	1	8				

The first few lines just describe the state of our database with regard to the input we have provided. Out of our 593,548 SNPs, 592,652 were found in our local database. We also are provided a list of meta-groups and their various counts and IDs. These "Group IDs" are useful when selectively using one or more meta-groups. The group and gene counts represent the various children found.

The next portion describes the estimated number of SNP-SNP models for each Implication Index. This number is an overestimation, since it doesn't take into account actual overlap between gene-gene models..

To describe the output of our snp-snp models, we get a summary describing how many models were produced. This number is the exact number that was produced. If you do the math, you will see that there aren't exactly 1,000,000 models produced. This has to do with the fact that gene-gene models produce varying numbers of snp-snp models, including some models that might be produced by another gene-gene model. biofilter keeps the requested model count in mind and tries to get close to the requested models. These models will have 0 overlapping members, and will represent the highest Implication Index possible. In our case, we only represented 930,386 out of the 4,812,558 possible models with an implication index of 2.0. The ones that were generated were simply the ones produced by the first N gene-gene models.

```
Alias Report: tutorial-aliases.html
Gene-Gene Model Summary: tutorial-model-summary.txt
Gene-Gene Models: tutorial.gene-gene
Snp Models: tutorial.snpsnp
```

Finally, we get the list of reports that were generated. According to the list above, we should find an Alias report (this describes which genes were given an alias, a gene-gene model summary, the actual gene-gene model file and the snp-snp model file.

## tutorial-model-summary.txt

This file lists the contents of our disease dependent groups. In this case, there is only one group. Since we only recognize a subset of gene names, we produce this list to help the user recognize that

- A) We got the important genes,
- B) They correspond to the same gene that the user expected.

Adding new group Alias	(176647), alz-assoc Ensembl ID		Chrom	Start Pos.	Stop Pos.	Snp Count
AGT	ENSG00000135744	98998		228904897	228916564	8
APH1A	ENSG00000117362	96653		148502512	148508156	0
APOA1BP	ENSG00000163382	97981		154828178	154830715	0
APOA2	ENSG00000158874	98229		159458707	159460042	1
CAMK1G	ENSG00000008118	98856		207823668	207853906	12
CFH	ENSG00000000971	98642		194887764	194983255	16
CHRNB2	ENSG00000160716	97575		152806881	152818975	1
CLCNKB	ENSG00000184908	78893		16242834	16256390	7

The ID column (#3) is the internal ID that we use with the configuration parameter, INCLUDE\_GROUPS.

## tutorial.gene-gene

```
146491
786099375641|4|9|12
800488012641|2|4|9
889669431541|2|4|9
791098951541|2|4|12
791098353041|2|4|12
868319825841|2|4|9
800918240841|2|4|12
783738037741|2|9|12
834768362941|4|9|12
883629336641|2|9|12
796329344932|4|9
854548734934|9|12
867989792131|4|9

(truncated)
```

This file lists all gene-gene pairings and the number of models that the pairing yielded. This is the actual model file and might be unreadable (binary) depending on the value of the configuration property **BINARY\_MODEL\_AR-CHIVE** is set to **YES**. The first line indicates the number of gene-gene models contained int the file. Subsequent lines contain the gene-IDs, their Implication Index and the Disease Independent information associated with the pairing. The file is sorted by Implication Index, then gene 1 then gene 2. No duplicate gene pairings should be found.

A more understandable report is the tutorial-model-summary.txt file:

Gene	Gene SNP	ene Models Gene	SNP	Impl.	ModelsGroups
Name	Count	Name	Count	Index	CountDI,DD
PAFAH1B1	8	GLI2	31	1	2481,
PAFAH1B1	8	CHRNB2	1	2	81,176647
GLI2	31	CHRNB2	1	2	311,176647
GLI2	31	DRD2	18	1	5581,
NR0B1	1	DRD2	18	1	181,
NR0B1	1	GLI2	31	1	311,
TGFB3	6	TGFB2	22	1	1321,
ROCK1	10	EZR	10	1	1001,
ROCK1	10	ICAM1	5	1	501,
ROCK1	10	MSN	2	1	201,
ROCK1	10	VCAM1	22	1	2201,
ICAM1	5	EZR	10	1	501,
		(truncated)			,

Both show the gene pairings and implication index and the components, even though the model file simply shows the ID. This is used by the program performing the snp-snp model expansion. Users can estimate the count of snp-snp models to be produced by multiplying the two snp-counts for any given gene-gene model.

Finally, there is the file, tutorial.snpsnp. This contains the actual snp-snp models of interest and may be binary for the same reason as the gene-gene model file.

The first line is simply the number of models found in the file. Each subsequent line lists two RS IDs (without the letters, "R" and "S") and the Implication Index. This file is sorted by rs-ID 1 and rs-ID2 respectively and should contain no duplicate rs pairings.

Another file can be found, called tutorial.genes. This file can be used by programs reading the gene-gene models and describes the SNPs associated with a particular gene-gene model production.

# References

Bush WS, Dudek SM, Ritchie MD. Biofilter: A Knowledge-Integration System for The Multi-locus Analysis of Genome-wide Association Studies. *Pacific Symposium on Biocomputing*. 2009:368–379