

# Haploview Documentation

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# Chapter 1. Using Haploview

## Loading a Dataset

Data can be loaded in six formats. Ped and Haps files can also load an optional marker info file and PLINK files normally require an accompanying map or binary map file. Further options are presented on the load screen:

- Haploview saves time by only computing pairwise LD statistics for markers within a certain distance of each other. The default is 500KB. Enter a value of zero to force all pairwise computations.
- Haploview excludes individuals with less than 50% complete genotypes. This threshold can be adjusted in the load dialog. Additional details about excluded individuals are available from the marker check tab.
- When loading a file dumped from the HapMap project website, it is possible to automatically display SNP and gene tracks from the HapMap above the data by checking the "Download and show HapMap info track" box. More information is available with the LD Display help. [hapmap file only]
- If you wish to perform association tests, you must inform the program now and select either family trios or case/controls. For family datasets a standard TDT or parenTDT are available. More details are available under association. [pedfile only]
- If your data is from the X chromosome in the linkage formats, tick the box so that Haploview will correctly process your data. In other formats, select the X chromosome in the dropdown menu. X chromosome data is not supported by the phased haplotype format. All functionality now works with the X chromosome.
- Haploview will maximize the information available from a pedigree for both LD analyses and association tests. For the former it creates a maximal set of unrelated individuals, using trio data only for obligate parent/offspring phasing. For TDT association testing, all available transmissions from parent-offspring will be used. More detailed information about specific situations is available in the FAQ.
- Haploview can be configured to support proxy host settings using the "Proxy Settings" button on the load screen.

Haploview allocates 512MB of memory by default. This is usually sufficient to handle datasets with several thousand markers. If you are running the program on very large datasets (>20,000 markers) you may need to force more memory (presuming your computer has sufficient resources available). This can be accomplished using the following command:

```
java -jar Haploview.jar -memory 2000
```

Where "2000" in this case specifies 2000 megabytes of memory and can be adjusted as necessary. Previous versions of Haploview required a slightly different command to adjust available memory, which still works:

```
java -Xmx2000M -cp Haploview.jar edu/mit/wi/haploview/Haploview
```

---

# Data Quality Checks

## Marker Checks

After loading a file, Haploview shows some basic data quality checks for the markers. Markers are filtered out based on some default criteria which can be adjusted as necessary. Markers can be added or removed from analyses by hand via the checkboxes. The data in this table can be sorted by clicking on any of the column headers. Compound sorts can be done by clicking on the first column header then CTRL clicking on the next one.

- **#** is the marker number.
- **Name** is the marker ID specified (only if an info file is loaded).
- **Position** is the marker position specified (only if an info file is loaded).
- **ObsHET** is the marker's observed heterozygosity.
- **PredHET** is the marker's predicted heterozygosity (i.e.  $2 * \text{MAF} * (1 - \text{MAF})$ ).
- **HWpval** is the Hardy-Weinberg equilibrium p value, which is the probability that its deviation from H-W equilibrium could be explained by chance.
- **%Geno** is the percentage of non-missing genotypes for this marker.
- **FamTrio** is the number of fully genotyped family trios for this marker (0 for datasets with unrelated individuals).
- **MendErr** is the number of observed Mendelian inheritance errors (0 for datasets with unrelated individuals).
- **MAF** is the minor allele frequency (using founders only) for this marker.
- **Alleles** are the major and minor alleles for this marker.
- **Rating** is checked if the marker passes all the tests and unchecked if it fails one or more tests (highlighted in red).

You can adjust the filtering thresholds and click "Rescore" to refilter the markers using the new values. These thresholds can be reset to values by clicking "Reset Values". Markers can also be selected/unselected by hand by clicking the "Rating" checkbox or using the "Select All" and "Deselect All" buttons. Any marker which fails one of the quality tests will have the relevant field(s) highlighted in red.

## Duplicate Markers

If two markers in an input file have the same chromosomal position, Haploview will ignore the less completely genotyped marker by default and highlight both in yellow on the check markers panel. When running in nogui mode Haploview always ignores the less completely genotyped version of two markers with the same position. If you want to use both from the command line, you'll need to adjust one of the positions.

If two markers in an input file have the same name, Haploview renames the second one in the file by appending ".X" to the filename, where "X" is a running integer count starting with 1. So if you have marker1, marker1 and marker2, Haploview would adjust this to: marker1, marker1.1 and marker2. Note

that if the markers with the same name have different positions, Haploview won't deselect any of them; if they do have identical positions, it will filter all but one out as described above.

## Filtered Individuals

The top of the tab contains information about individuals filtered during the loading of the file. It will show overview information about the number of singletons and trios used and the number of independent families loaded. Further information can be shown by clicking the "Advanced Views" button. This will present a list of up to four buttons depending on the nature of the loaded dataset. The "Individual Summary" button will show genotyping percentage by family and individual. If individuals have been excluded, the "Excluded Individuals" button will present a list of excluded individuals as well as the reason for exclusion. If Mendel errors are present, view detailed Mendel error information by clicking the "Mendel Errors" button. If male heterozygotes are present in x chromosome data, information about them can be viewed by clicking "Male Heterozygotes". All of these advanced views can also be exported using the "Export to File" button. Details about individual filtering can be found in the FAQ.

## LD Display

### Perusing the LD Display

- The color scheme option (Display menu) allows you to choose among several LD color schemes. The following tables provide details on the color schemes, and a key to the meaning of the currently selected scheme can be dropped down from the "Key" menu in the upper right corner of the screen.

**Table 1.1. Standard Color Scheme**

	D' < 1	D' = 1
LOD < 2	white	blue
LOD # 2	shades of pink/red	bright red

**Table 1.2. Confidence Bounds Color Scheme**

Strong Evidence of LD	dark grey
Uninformative	light grey
Strong Evidence of Recombination	white

**Table 1.3.  $r^2$  Color Scheme**

$r^2 = 0$	white
$0 < r^2 < 1$	shades of grey
$r^2 = 1$	black

**Table 1.4. Alternate D'/LOD Color Scheme**

	Low D'	High D'
Low LOD	white	shades of pink
High LOD	white	black

( $r^2$  and Alt D'/LOD courtesy of Will Fitzhugh)

**Table 1.5. 4 Gamete Color Scheme**

4 distinct 2-marker haplotypes	white
< 4 distinct 2-marker haplotypes	black

- In order to help keep the display uncluttered, D prime values of 1.0 are never shown (the box is empty). These values can be switched on or off with the "Show LD values" option in the Display menu.
- The zoom option (Display menu) allows you to select one of three zoom modes. The two zoomed out versions can be useful for browsing large datasets.
- Large datasets also show a "map" in the lower left corner which gives an overview of the D prime display and allows you to navigate quickly. Clicking on an area of the map will cause the main display to jump to that area. This map also shows the currently defined blocks as small black lines across the top.
- Markers with additional notes (as loaded from the info file) are highlighted (the names are green in the zoomed-in view and the lines from the SNP position to the LD chart are green in the zoomed-out view. Details can be viewed by right clicking on the marker number (as mentioned below).
- Right clicking on the marker number (or the equivalent space in the zoomed out views) shows the marker name, minor allele frequency and any additional notes specified in the info file. This can be especially helpful in the zoomed out views which do not display marker names. The last such piece of popup information clicked will be shown at the top of the LD plot. This reminder can be dismissed by left clicking anywhere on the LD plot.
- Right clicking on any pairwise LD comparison will show a more detailed summary of the LD between the two markers in question. This information is also shown at the top of the screen as described above and can be dismissed by left clicking anywhere on the LD plot.

## Additional Data Tracks

### Analysis Track

A graph of any variable versus chromosomal location can be added above the LD plot with the "Load Analysis Track" option. Simply create a file with two columns: <position> <value> . Haploview will plot the values in a continuous line along the top of the screen, along with a scale bar on the Y-axis. You can load several analysis tracks which will all be plotted in the same box at the top of the LD plot.

### HapMap Gene/SNP Track

The "Download HapMap info track" option (with an internet connection) allows you to connect to the HapMap Project server and download and display a track with HapMap genotyped SNPs and gene names. If an info file is specified, the default boundaries are the positions of the first and last markers (which is only valid if the info file is in genomic coordinates). You must specify the proper chromosome and genomic build in the dialog box. If you are using a file downloaded from the HapMap website the program will specify the correct default chromosome, build and start/end positions. This track display can be configured with the "HapMap Info Track Options" item in the "Display" menu. Available tracks include HapMap SNPs, Entrez genes, recombination rate, contigs, and GC content.

# Blocks and Haplotypes

## Blocks

Haploview generates blocks whenever a file is opened, but these blocks can be edited and redefined in a number of ways. In the Analysis menu, you can clear all the blocks in order to start over, define blocks based on one of several automated methods or customize the parameters of those algorithms. Additionally, the blocks can be edited by hand.

## Confidence Intervals [DEFAULT]

The default algorithm is taken from Gabriel et al, Science, 2002. 95% confidence bounds on D prime are generated and each comparison is called "strong LD", "inconclusive" or "strong recombination". A block is created if 95% of informative (i.e. non-inconclusive) comparisons are "strong LD". This method by default ignores markers with  $MAF < 0.05$ . The MAF cutoff and the confidence bound cutoffs can be edited by choosing "Customize Block Definitions" (Analysis menu). This definition allows for many overlapping blocks to be valid. The default behavior is to sort the list of all possible blocks and start with the largest and keep adding blocks as long as they don't overlap with an already declared block.

## Four Gamete Rule

This is a variant on the algorithm described in Wang et al, Am. J. Hum. Genet., 2002. For each marker pair, the population frequencies of the 4 possible two-marker haplotypes are computed. If all 4 are observed with at least frequency 0.01, a recombination is deemed to have taken place. Blocks are formed by consecutive markers where only 3 gametes are observed. The 1% cutoff can be edited to make the definition more or less stringent.

## Solid Spine of LD

This internally developed method searches for a "spine" of strong LD running from one marker to another along the legs of the triangle in the LD chart (this would mean that the first and last markers in a block are in strong LD with all intermediate markers but that the intermediate markers are not necessarily in LD with each other).

Markers can be removed from blocks by clicking on the marker number (along the top of the D prime graph). Blocks can be defined by hand by clicking and dragging along the marker number row. Any block which overlaps with an existing block will take precedence and delete the existing block.

# Haplotypes

## Display

View haplotypes for selected blocks by clicking on the "Haplotypes" tab or selecting "Haplotypes" from the Display menu. Haplotypes are estimated using an accelerated EM algorithm similar to the partition/ligation method described in Qin et al, 2002, Am J Hum Genet. This creates highly accurate population frequency estimates of the phased haplotypes based on the maximum likelihood as determined from the unphased input.

The haplotype display shows each haplotype in a block with its population frequency and connections from one block to the next. In the crossing areas, a value of multiallelic  $D'$  is shown. This represents the level of recombination between the two blocks. Note that the value of multiallelic  $D'$  is computed for

only the haplotypes ("alleles") currently displayed. This usually does not have a strong effect, as the rare haplotypes contribute only slightly to the overall value. Above the haplotypes are marker numbers along with a tick beneath haplotype tag SNPs (htSNPs).

## Display Controls

The display can be edited using the controls at the bottom of the screen to display only more common haplotypes or to adjust the connecting lines. By default, alleles are displayed using A,C,G,T along with the special symbol 'X' which represents a fairly rare situation in which only one allele is unambiguously observed in phased data. The 'X' represents the allele of unknown identity. The display can also be changed to show the alleles numerically from 1-4 with 8 being the equivalent of 'X', or as blue and red boxes, with blue being the major allele and red the minor.

## Tag SNPs

Haplotype tag SNPs are no longer displayed by default in the Haplotypes tab. It is recommended that all tagging be done via the Tagger tab. The block-by-block tags can be displayed by ticking the "Show tags in blocks" option in the Display menu.

# Tagger

## Introduction

We have developed a tagging strategy that combines the simplicity of pairwise methods with the potential efficiency of multimarker approaches. We avoid overfitting and unbounded haplotype tests in the association phase by (a) using only those multiallelic combinations in which the alleles are themselves in strong LD, and (b) explicitly recording the allelic hypotheses that are to be tested in the subsequent association analysis. Attractive practical features include the ability to force in or exclude sets of tags.

Haploview is based on Paul de Bakker's *Tagger*. It and more information are available at the Tagger website. There are a number of differences between the implementations, although they are constructed around the same concept. Tagger currently searches a much broader space of available multi-marker tests (up to 6-mers) whereas Haploview allows only 2- or 3-marker tests in the interest of computational efficiency.

## Features

Haploview's Tagger operates in either pairwise or aggressive mode. In either case it begins by selecting a minimal set of markers such that all alleles to be captured are correlated at an  $r^2$  greater than a user-editable threshold with a marker in that set. Certain markers can be forced into the tag list or explicitly prohibited from being chosen as tags. You can also specify which markers in the dataset you want to be captured.

Aggressive tagging introduces two additional steps. The first is to try to capture SNPs which could not be captured in the pairwise step (N.B. these must have been "excluded" since otherwise they would simply be chosen to capture themselves) using multi-marker tests constructed from the set of markers chosen as pairwise tags. After this, it tries to "peel back" the tag list by replacing certain tags with multi-marker tests. Tagger avoids overfitting by only constructing multi-marker tests from SNPs which are in strong LD with each other, as measured by a pairwise LOD score. This LOD cutoff can be adjusted to loosen or tighten this requirement; in general, the default cutoff of 3.0 is appropriate for selecting tags from a HapMap-sized reference panel of 120 chromosomes.

Much more information about the development of this algorithm is available at the Tagger website.

## Tagger Configuration Panel

**N.B.** Haploview's Tagger requires either an info file or a hapmap style input file, because it references the marker names specified in those files. If you load a pedigree or phased haplotypes input file without an info file, the Tagger panels will not be available.

This panel shows all SNPs available for tag selection. SNPs which are deselected in the Check Markers tab will not be in this list. There are three checkboxes for each SNP:

Force Include	Checking this box will force this SNP to be chosen as a tag SNP.
Force Exclude	Checking this box will prohibit this SNP from being chosen as a tag SNP.
Capture this Allele?	If this box is checked, Haploview will include this SNP in the list of alleles to be captured by the chosen tag set.

**N.B.** The include and exclude checkboxes are mutually exclusive, and "Capture this Allele" must be checked in order to either include or exclude a marker.

Directly below the marker list are buttons to quickly manipulate the table above. Use "Include All" to check all of the "Force Include" boxes, and "Exclude All" to check all of the "Force Exclude" boxes. "Uncapture All" will uncheck the "Capture this Allele?" column for all markers, and "Reset Table" will return the table to its initial state. Beneath these buttons are several additional tagging options. You can choose from among pairwise and two aggressive tagging strategies discussed above. You can also set the  $r^2$  and LOD thresholds as previously mentioned. Additionally, you can specify the maximum number of tags to pick, as well as the minimum distance (in base pairs) between picked tags. You can load a set of SNPs to include or exclude using the "Load Includes" and "Load Excludes" buttons. These buttons take in a file with a single column of SNPs to include or exclude. The "Alleles to Capture" button also takes in a file with a single column of SNPs to be captured. Design scores can also be loaded in using the "Design Scores" button. Design score files should contain two columns containing the SNP and the design score to assign to that SNP. Clicking "Run Tagger" will run the tagging algorithm. When finished it will switch from the Configuration to the Results Panel.

## Tagger Results Panel

This panel is split into a "Tests" section on the left and a marker-by-marker report on the right. The marker report lists all SNPs, the test which best captures them, and their  $r^2$  with that test. SNPs which were unchecked from the "Capture this allele?" list on the Configuration panel are greyed out. SNPs which could not be successfully tagged are shown in red.

The first list in the "Tests" section shows all the tests (both single marker and multi-marker alleles) chosen by Haploview. Selecting tests in this list will show which alleles are captured by those tests in the second list in the panel. Beneath these lists is a summary of the tagging results.

Captured N alleles with mean $r^2$ of X.	This shows how many of the SNPs in the dataset have been successfully tagged by the set of chosen tests. The mean $r^2$ represents the mean for only those SNPs successfully captured.
Captured N percent of alleles with $r^2 >= 0.8$	This shows what fraction of the alleles captured by the tests have an $r^2 >= 0.8$ . Of course, if your tagging $r^2$ threshold is $>= 0.8$ this value will always be 100%.

Using N SNPs in M tests. This shows that N unique SNPs have been chosen to create M tests, which can either be one of the set of N SNPs or some combination of those SNPs.

The "Dump Tests File" button exports a file with the list of tests in the format used by Haploview's custom association test file and Tagger's export. This file contains the list of all tests (single SNPs and multi-marker tests) selected by Tagger for subsequent association analysis. In pairwise-only tagging this file will be identical to the "Tags" file, below.

The "Dump Tags File" button exports a file with the list of Tag SNPs in the format used by Haploview's custom association test file and Tagger's export. It is the concise list of SNPs selected by Tagger for genotyping. In pairwise-only tagging this file will be identical to the "Tests" file, above.

The "Export Tab to Text" option in the File menu will export a summary file showing the best tag for each marker and the list of tests along with the alleles tagged by each test.

## Association Tests

If selected when loading the data, Haploview computes single locus and multi-marker haplotype association tests. For case/control data, the chi square and p-value for the allele frequencies in cases vs. control are shown. For family trios, all probands (affected individual with genotyped parents) are used to compute TDT values. If the parenTDT option is selected, additional information is gained from parental phenotypes. More information about this method can be found in the Citations list in the About Haploview section.

The haplotype association test is performed on the set of blocks selected on the LD and haplotype tabs. Results are shown only for those haplotypes above the display threshold on the haplotype tab. Counts for both TDT and case control association tests are obtained by summing the fractional likelihoods of each individual for each haplotype. In other words, if a particular individual has been determined by the EM to have a 40% likelihood of haplotype A and 60% likelihood of haplotype B, 0.4 and 0.6 would be added to the counts for A and B respectively.

Additional information about the way in which pedigrees are filtered for TDT purposes can be found in the FAQ.

Haploview is not intended to be the only way of testing association results, but to provide a straightforward way to do simple association tests. It's always a good idea to try out multiple approaches to analyzing your data.

You can load a set of custom association tests in the format exported by Haploview and Tagger. This format is discussed below.

## Permutation Testing

Haploview provides a framework for permuting your association results in order to obtain a measure of significance corrected for multiple testing bias. You can choose to permute one of several test sets:

Single Markers Only	Permute just association tests to the individual SNPs in your dataset.
Single Markers and Haplotypes in Blocks	Permute the individual SNPs as above, along with all the haplotypes shown in the Haplotypes tab.
Haplotypes in blocks only	Permute only the haplotypes in the Haplotypes tab, ignoring the single marker results.

Custom Tests from File                      Permute the set of tests loaded from an external file. Note that this choice is only available if you provided a tests file when you loaded your dataset.

Specify how many permutations to do and press the "Do Permutations" button to start the permutations. While the permutations are running, Haploview shows the following:

- A progress bar which tracks the progress of the permutations.
- The highest permuted chi square so far.
- The fraction of permutations whose strongest association exceeds the best observed chi square.

You can stop the permutations at any time with the "Stop" button. Once the permutations are complete, Haploview displays:

- A table listing all tests (single SNP and haplotype) along with their association chi squares and permuted p-values.
- A histogram of the highest chi square from each of the permutations.

You can save the permutation summary by using the "Export Tab to Text" option in the File menu.

## PLINK

Haploview can now take in PLINK outputs. These files require a separate map file or binary map file corresponding to each marker in the output file in order to load. Any output file from PLINK can be loaded provided that it contains a SNP column corresponding to the map file. The map file can contain SNPs that are not present in the associated output file and the SNPs need not be in the same order in the two files. PLINK output is displayed in a single tab containing a sortable table of results and a variety of filtering options below the table. In SNP-based files, you can also load in additional columns using the "Load Additional Results" button.

Filtering options include chromosome and position. The "Filter1" and "Filter2" dropdown boxes can be used to further filter on any unrecognized columns in the table. All filters are jointly applied (logically equivalent to an "AND" operator as opposed to an "OR"). You can also go directly to a specified marker name in the table using the "Go" button on the second row, and remove columns from the table by selecting a column from the "Remove Columns:" dropdown menu and using the adjacent "Remove" button. The "Reset" button can be used to revert the table and filters back to their original state (though the sorting state is retained). Please note that non-SNP based files which can be loaded in without a map file do not have the chromosomal or marker filters.

You can use the Fisher method for combining p-values in your PLINK results using the "Combine P-Values" button under the filtering options. This will bring up a dialog that allows you to choose between 2 and 5 p-value columns for use in the Fisher-combined algorithm. Once you click "Go", a new column designated as "P\_COMBINED" will appear as the last column in the table.

You can create graphical plots from your results table using the "Plot" button under the filtering options. Use the Plot Options dialog to specify a title for your plot and various plotting options. At the top of the dialog, select an optional title for your plot. Then choose which columns to use on the X and Y axes of the plot as well as the scale for each using the appropriate drop-down boxes. If you have loaded a SNP based file with an accompanying map file, selecting "Chromosomes" as the X-Axis will plot your results across the chromosomes and will color code them separately. You can also select "Index" for either axis which will simply plot sequential numbers for each result shown in the table for that axis. Additionally, you can specify up to two thresholds for use in the plot, along with which axis to place them on and which direction

they should be. Threshold 1 or the "Suggestive" threshold for the  $-\log_{10}$  scale will create a blue line and Threshold 2 or the "Significant" threshold for the  $-\log_{10}$  scale will create a red line. Datapoints which pass the thresholds will be larger in size than the standard datapoints. Directly beneath the thresholds, you can choose the base datapoint size for results in your plot using the dropdown box. To the right of that, you can use the optional "Color Key" dropdown menu to select a column to be used as a coloring key in the plot. Please note that this functionality will only work when the chosen color key column has 50 or fewer unique values. On the next line, you can use the "Show Gridlines?" checkbox to select whether to show or hide the gridlines in the plot. To the right of that, you can specify the initial width and height of the plot (in pixels). Finally, on the next line you can use the "Export to SVG" checkbox and browse to a location to save your plot to a high quality SVG file. Please note that the SVG option generally takes a great deal of processing power and memory and should only be used when very high quality images are required. In most cases, you can save the plot images as PNG files using the right click context menu described below.

Once the plot has loaded, you can hover over individual data points to see information about that point in a tooltip popup. SNP based data will display the corresponding marker name, chromosome, position, and the value that is being plotted. Non SNP-Based data will either display the corresponding FID and IID values if they are available, or simply the X and Y values for that datapoint. You can also click a datapoint to be taken back to that result in the results table. For many more plotting options including export options, right click anywhere on the plot.

By highlighting a specific result in the table and clicking "Go to Selected Region", you can bring up a dialog to automatically fetch that region from the HapMap website and load it into Haploview. The dialog allows you to specify the size of the region and the HapMap analysis panel that you wish to download. You can also optionally choose to annotate the columns from the PLINK tab to annotate in the LD Plot. Once the region has been successfully loaded into Haploview, the initially selected marker will be highlighted in blue on the Check Marker and PLINK tabs. SNPs that appear in the PLINK tab are now marked in green on the LD Plot, and the specific result that you specified is further highlighted in white. You can view the annotated data from the PLINK table by right clicking on the marker number in the LD Plot. You can use the "Force in PLINK SNPs" button in the Tagger panel to force include all the SNPs contained in the PLINK results tab. Please note that using "Go to Selected Region" requires an active internet connection.

The "Export Tab to Text" option in the File menu will export a text file containing the current view of the results table. This file will preserve any sorting and filtering that you've enabled in the table.

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# Chapter 2. Files

## Input File Formats

Haploview currently accepts input data in five formats, standard linkage format, completely or partially phased haplotypes, HapMap Project data dumps, PHASE format, and PLINK outputs. The program can also automatically fetch phased HapMap data off the HapMap website. It also takes in a separate file with marker position information, as well as several auxiliary input files, described below. The four formats are explained in depth below.

### Linkage Format

Linkage data should be in the Linkage Pedigree (pre MAKEPED) format, with columns of family, individual, father, mother, gender, affected status and genotypes. The file should not have a header line (i.e. the first line should be for the first individual, not the names of the columns). Please note that Haploview can only interpret biallelic markers — markers with greater than two alleles (e.g. microsatellites) will not work correctly. A sample line from such a file might look something like:

```
3      12     8     9     1     2     1 2     3 3     0 0     4 2  
a      b     c     d     e     f     -----g-----
```

- |                      |  |
|----------------------|--|
| (a) pedigree name    | A unique alphanumeric identifier for this individual's family. Unrelated individuals should not share a pedigree name.                                       |
| (b) individual ID    | An alphanumeric identifier for this individual. Should be unique within his family (see above).  |
| (c) father's ID      | Identifier corresponding to father's individual ID or "0" if unknown father. Note that if a father ID is specified, the father must also appear in the file. |
| (d) mother's ID      | Identifier corresponding to mother's individual ID or "0" if unknown mother. Note that if a mother ID is specified, the mother must also appear in the file. |
| (e) sex              | Individual's gender (1=MALE, 2=FEMALE).  |
| (f) affection status | Affection status to be used for association tests (0=UNKNOWN, 1=UNAFFECTED, 2=AFFECTED).   |

## (g) marker genotypes

Each marker is represented by two columns (one for each allele, separated by a space) and coded either ACGT or 1-4 where: 1=A, 2=C, 3=G, T=4. A 0 in any of the marker genotype position (as in the the genotypes for the third marker above) indicates missing data.

It is also worth noting that this format can be used with non-family based data. Simply use a dummy value for the pedigree name (1, 2, 3...) and fill in zeroes for father and mother ID. It is important that the "dummy" value for the ped name be unique for each individual. Affection status can be used to designate cases vs. controls (2 and 1, respectively).

Files should also follow the following guidelines:

- Families should be listed consecutively within the file (i.e. all the lines with the same pedigree ID should be adjacent)
- If an individual has a nonzero parent, the parent should be included in the file on his own line.

## Phased Haplotypes

Haplotype data for Haploview's input must be formatted in columns of Family, Individual and Genotypes. There should be two lines (chromosomes) for each individual. This is the standard format of Genehunter's TDT output. See the sample below:

FAM1	FAM1M01	0	4	2	2
FAM1	FAM1M01	0	4	2	2
FAM1	FAM1F02	3	h	1	2
FAM1	FAM1F02	3	h	1	2

The data format uses the numerals 1-4 to represent genotypes, the number zero to represent missing data, and the letter "h" to represent a heterozygous allele. That is, if an individual is heterozygous at a locus, both alleles should be "h" if the phasing (which allele falls on which chromosome) is uncertain.

## HapMap Project Data Dumps

Data from the HapMap Project can be dumped by region using the GBrowse interface. The saved data file is in a marker-per-line format which can be loaded in Haploview.

GBrowse dumps only one file, which has one marker per line and which includes familial relationships among the HapMap samples as well as marker position information. The file format has several header lines (beginning with "#") which Haploview parses. Open the file by selecting "Browse HapMap Data" option and selecting the downloaded file.

If you wish to load data from another source in HapMap style format, you will need to specify pedigree information in the header of the file

you've created. This can be done by creating lines of the following format at the top of your file:

```
#@ FAM01 NA0001 0 0 1 1
```

This data is the same as the pedfile format discussed above. That is, the fields are family,individual,father,mother,gender,affected status. You would then replace the NAXXXX identifiers in the header row of the HapMap file with your identifiers, subject to two important constraints: they must be unique across the entire dataset, not just within a family and they must begin with the characters NA.

#### PHASE Format

Data in the HapMap *PHASE* format can be loaded into Haploview using three separate files. The first is the data file containing binary allele information. The second is a sample file containing a single column of the individual IDs used in the dataset. The third is a legend file containing four columns: marker, position, 0, and 1. Only the legend file requires a header and is used to decode the information in the data file. These files can be loaded in as GZIP compressed files using the "Files are GZIP compressed" checkbox on the initial loading screen.

#### Marker Information File

The marker info file is two columns, marker name and position. The positions can be either absolute chromosomal coordinates or relative positions. It might look something like this:

```
marker01 190299
marker02 190950
marker03 191287
```

An optional third column can be included in the info file to make additional notes for specific SNPs. SNPs with additional information are highlighted in green on the LD display. For instance, you could make note that the first SNP is a coding variant as follows:

```
marker01 190299 CODING_SNP
marker02 190950
marker03 191287
```

#### PLINK Format

Output files from *PLINK* can be loaded into Haploview using the *PLINK* tab on the initial loading screen. *PLINK* files must contain a header and at least one column header must be titled "SNP" and contain the marker IDs for the results in the file. *PLINK* loading also requires a standard *PLINK* map or binary map file corresponding to the markers in the output file. The map file can be either three or four headerless columns (the Morgan distance column is optional). The map file can also be embedded in the results file as the first few columns of the file using the "Integrated Map Info" checkbox. You can load in non-SNP based files as well by checking the "Non-SNP" box. These files do not require a map file. You can choose to only load in one chromosome from your results file using the "Only load results from Chromosome" checkbox and selecting a chromosome from the dropdown list. You can also select which columns to load from your results file by checking the "Select Columns" checkbox. For a great

deal more information on PLINK outputs, please see Shaun Purcell's PLINK website

#### Batch Load File

The "-batch" flag on the command line allows you to run Haploview automatically (in nogui mode) on several files. Batch input files should have one genotype file per line, along with an info file (if desired) separated by a space. Filenames must conform to the following rules:

- Pedfile names must end in ".ped"
- Phased haplotype file names must end in ".haps"
- HapMap file names must end in ".hmp"
- Info file names must end in ".info"

The following example shows 2 pedfiles (with info files) and a hapmap file:

```
sample1.ped   sample1.info
sample2.ped   sample2.info
sample3.hmp
```

## Output Files

For any given tab the information in the display can be saved. For the data check and association test tabs, a simple tab-delimited text file is generated from the tables. For the LD and Haplotype tabs, data can either be dumped to text files or the image can be saved to a PNG.

#### LD Text Output File

LD text output is a tab delimited set of columns containing the various measures of LD used by the program. Details for each column are shown below:

- L1 and L2 are the two loci in question, referenced by their number or name (if marker info file is provided)
- D' is the value of D prime between the two loci.
- LOD is the log of the likelihood odds ratio, a measure of confidence in the value of D'
- $r^2$  is the correlation coefficient between the two loci
- CI<sub>low</sub> is 95% confidence lower bound on D'
- CI<sub>hi</sub> is the 95% confidence upper bound on D'
- Dist is the distance (in bases) between the loci, and is only displayed if a marker info file has been loaded
- T-int is a statistic used by the HapMap Project to measure the completeness of information represented by a set of markers in a region

Details about additional options for this output type can be found below in the Export Options section.

#### LD PNG Output

When saving the LD table to a PNG, Haploview saves an image using the current display settings. This includes color scheme, zoom and proportional spacing. Thus, in order to save a less detailed image to a PNG, first zoom out, then export the tab. Note that Haploview cannot save large datasets at the higher zoom levels. For more information see the Export Options section below.

#### Haplotype Text Output File

Haplotype output shows a block, its markers, the haplotypes and their population frequencies, the crossover percentages to the next block and the multiallelic D prime. Crossover percentages are shown as a matrix with this block's haplotypes as the rows and the next block's haplotypes as the columns. An example might look like:

```
BLOCK 1.  MARKERS: 1 2 3 4
3312 (0.825)   | 0.800  0.025  0.000 |
1144 (0.163)   | 0.031  0.125  0.007 |
3342 (0.013)   | 0.006  0.000  0.006 |
Multiallelic Dprime: 0.802
BLOCK 2.  MARKERS: 10 11 12
441 (0.837)
222 (0.150)
242 (0.013)
```

In this example, the first block has 4 markers with 3 haplotypes displayed and the second block has 3 markers and 3 haplotypes. The tag SNPs for each block are (3,4) and (10,11) respectively. The crossover percentage matrix can be read as follows: 80% of all samples have the pattern 3312-441, 3.1% have the pattern 1144-441 and so forth.

#### Haplotype PNG Output

Saving the haplotype tab to a PNG produces an image using the current display settings (such as haplotype frequency cutoff).

#### Single Marker Association Text Output File

Single marker association results are saved in a tab-delimited text file with the following columns:

- # is the marker number.
- Name is the marker ID specified if an info file is loaded.
- Chi Square is the chi square value for the marker.
- p value is the significance level for the above chi square.

Trio (TDT) data only:

- Overtransmitted is the allele overtransmitted to affected offspring.
- T:U is the ratio of transmissions to non transmissions of the overtransmitted allele (see above).

Case-Control data only:

---

		<ul style="list-style-type: none"> <li>• <code>Major Alleles</code> are the major alleles in the case and control populations respectively.</li> <li>• <code>Case Control Ratios</code> are the ratios (shown as either counts or quotients, depending on selected options) for the case and control populations, respectively.</li> </ul>
Haplotype Association Text Output		<p>Haplotype association text output is a tab-delimited file, broken into sections by block. The columns are:</p> <ul style="list-style-type: none"> <li>• <code>Haplotype</code> is the sequence of alleles for this haplotype in this block.</li> <li>• <code>Frequency</code> is the population frequency for this haplotype.</li> <li>• <code>Chi Square</code> is the chi square value for the haplotype.</li> <li>• <code>p value</code> is the significance level for the above chi square.</li> </ul> <p>Trio (TDT) data only:</p> <ul style="list-style-type: none"> <li>• <code>T:U</code> is the ratio of transmissions to non transmissions of the haplotype to affected offspring.</li> </ul> <p>Case-Control data only:</p> <ul style="list-style-type: none"> <li>• <code>Case Control Ratios</code> are the ratios (shown as either counts or quotients, depending on selected options) for the case and control populations, respectively.</li> </ul>
Permutation Text Output File		<p>The output from the permutations tab shows the number of permutations performed and then a tab-delimited table with one row per permuted test and the following columns:</p> <ul style="list-style-type: none"> <li>• <code>Name</code> is the test name, which is either a marker name or a comma separated list of marker names then a tab then a comma separated set of alleles for those markers.</li> <li>• <code>Chi Square</code> is the observed association chi square for that test.</li> <li>• <code>Permutation p-value</code> shows the significance of the test among the permutation tests.</li> </ul>
Tagger Text Output File		<p>The Tagger text output begins with several pieces of summary information. More details on this can be found in the Tagger section. The rest of the output is divided into two sections. The first lists each marker, with the following rows:</p> <ul style="list-style-type: none"> <li>• <code>Marker</code> is the marker name.</li> <li>• <code>Best Test</code> is the test with the highest <math>r^2</math> to this marker.</li> <li>• <code>r<sup>2</sup> w/test</code> is the <math>r^2</math> between this marker and its test.</li> </ul> <p>The second part consists of a list of the tests and the alleles they capture best.</p>

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Tagger Tests Dump	This file is the same format used by Haploview for custom association tests and exported by Tagger. It is discussed below in the auxiliary files section.
Tagger Tags Dump	This file is the same format used by Haploview for custom association tests and exported by Tagger. It is discussed below in the auxiliary files section.
Marker Check Text Output File	<p>The marker check data is a tab-delimited file with the following columns:</p> <ul style="list-style-type: none"> <li>• # is the marker number.</li> <li>• Name is the marker ID specified (only if an info file is loaded).</li> <li>• Position is the marker position specified (only if an info file is loaded).</li> <li>• ObsHET is the marker's observed heterozygosity.</li> <li>• PredHET is the marker's predicted heterozygosity (i.e. <math>2 * \text{MAF} * (1 - \text{MAF})</math>).</li> <li>• HWpval is the Hardy-Weinberg equilibrium p value, which is the probability that its deviation from H-W equilibrium could be explained by chance.</li> <li>• %Geno is the percentage of non-missing genotypes for this marker.</li> <li>• FamTrio is the number of fully genotyped family trios for this marker (0 for datasets with unrelated individuals).</li> <li>• MendErr is the number of observed Mendelian inheritance errors (0 for datasets with unrelated individuals).</li> <li>• MAF is the minor allele frequency (using founders only) for this marker.</li> <li>• Alleles are the major and minor alleles for this marker.</li> <li>• Rating is "BAD" if the marker failed any of the above tests and blank otherwise.</li> </ul>
PLINK Table Text Output File	The PLINK text output is a tab-delimited file of the current view of the data in the PLINK tab. Please note that while the filtering state is preserved in this output, the sorting state is not.

## Export Options

The "Export Options" item in the File Menu allows adjustment of several parameters and allows the user to save any tab without having to switch to it. Specifically, the LD tab allow the markers to be filtered to output only some of the markers:

All	The default setting (and only one available for most tabs) is to use all the markers.
-----	---

Marker Range	Generates the LD text or PNG file for only a specific range of markers.
Adjacent Markers	Generates the LD text file for only adjacent markers. This can be useful to view the T-int stat, which measures LD information content in the gaps between markers.

There is also an option to generate a "compressed" LD PNG, which is useful for very large datasets. The image is shrunk to an arbitrary zoom level which allows Haploview to save the PNG with minimal memory usage.

## Auxiliary Input Files

**Blocks File** You can specify a set of blocks by loading a blocks file. Each line is a space separated list of markers with one block per line. For example:

```
1 2 3 4
9 10 11 12 13 14 15
```

Would create one block from markers 1-4 and another from 9-15. The first marker in the file is number 1 (not 0).

**Analysis Track File**

You can add an analysis track along the top of the LD display by loading a file with two columns, <position> <value>. Haploview will plot the values continuously with respect to the positions of the markers, so the positions should use the same coordinates as the marker info file. For example:

```
1000 0.3
2000 1.7
3000 11.0
4000 2.3
5000 4.6
```

Would plot a line from position 1000 to 5000. The values can be of any units or magnitude, as the Haploview scales the analysis track to the bounds of the values.

**Custom Association Tests File**

You can specify a set of custom association tests for Haploview to perform. The format takes both single marker tests and multi-marker tests (which require you to specify alleles for those markers). The format is one test per line with each line containing one of the following: a single marker name or several comma separated names, then a tab, then comma separated alleles for each marker. This format is exported by Haploview using the "Dump Tests" button in the Tagger Results panel and by Paul deBakker's Tagger webpage.

For instance, the following example would create 5 tests: markers 1, 2 and 3 individually, all the alleles (haplotypes) of the block 4,5,6 and the CAA haplotype of the block 12,13,14:

```
marker1
marker2
marker3
marker12,marker13,marker14    2,1,1
```

**N.B.** Using a Custom Association Tests File requires a marker info file, since the tests file reads the marker names as specified in the info file.

Tagger    Marker    Include/  
Exclude File

You can specify a list of markers for Tagger to include or exclude from those markers available for selection as tag SNPs. In either case the format is the same: one marker name per line. The following file could be used to either include or exclude markers 1,7 and 9:

```
marker1  
marker7  
marker9
```

**N.B.** Using a Tagger Include/Exclude File requires a marker info file, since it reads the marker names as specified in the info file.

---

# Chapter 3. Command Line Options

Haploview can be run from the command line without the display in order to do processing of multiple datasets or quick computation on very large datasets. In order to run Haploview without the display, add the "-nogui" flag. The "-help" flag shows a condensed explanation of all the command line options explained below. Haploview can be run from the command line using:

```
java -jar Haploview.jar
```

## General Options

-h, -help	Print help information.
-n, -nogui	Command line mode—does not launch display.
-q, -quiet	Quiet mode—minimizes output to command line.
-log <filename>	Outputs logfile information to specified filename (defaults to haploview.log if no name specified)
-memory <memsize>	Allocate <memsize> megabytes of memory to the Haploview process (default is 512MB).

## Input Options

-pedfile <filename>	Specify a genotype input file (or http:// location) in pedigree format. This option works in GUI mode.
-hapmap <filename>	Specify a HapMap format input file (or http:// location). This option works in GUI mode.
-phasedhmpdata <filename>	Specify a PHASE format data input file (or http:// location). This option works in GUI mode.
-phasedhmppsample <filename>	Specify a PHASE format sample input file (or http:// location). This option works in GUI mode.
-phasedhmplegend <filename>	Specify a PHASE format legend input file (or http:// location). This option works in GUI mode.
-hapmapDownload	Specify a phased HapMap download. This option works in GUI mode.
-haps <filename>	Specify a phased input file (or http:// location). This option works in GUI mode.
-info <filename>	Specify a marker information file (or http:// location). This option works in GUI mode.
-plink <filename>	Specify a PLINK or other results file (or http:// location). This option only works in GUI mode.
-map <filename>	Specify a map or binary map file (or http:// location). This option only works in GUI mode.

-batch <filename>	Specify a batch load file.
-blocks <filename>	Specify a block definition file (or http:// location). This will automatically use this block definition for haplotype output.
-track <filename>	Specify an analysis track file (or http:// location)
-chromosome <1-22,X,Y>	Specify which chromosome these data come from. This is especially critical when analyzing data from the X chromosome or direct HapMap downloads.
-panel <CEU,YRI,CHB+JPT>	Specify which analysis panel to use for phased HapMap downloads.
-startpos <integer>	Specify the start position in kb for phased HapMap downloads.
-endpos <integer>	Specify the end position in kb for phased HapMap downloads.
-release <16a,21>	Specify the HapMap release version for phased HapMap downloads (defaults to 21).
-gzip	Specify PHASE format inputs using GZIP compression
-nonSNP	Specify that the accompanying PLINK file is non-SNP based output. This option only works in GUI mode.
-selectCols	Activate the preloading column filter for PLINK loads. This option only works in GUI mode.

## Data Check Options

-skipcheck	Skip all the genotype data quality checks and uses all markers for all analyses.
-minMAF <threshold>	Exclude all markers with minor allele frequency below <threshold>, which must be between 0 and 0.5. Default of 0. This option works in GUI mode.
-maxMendel <integer>	Exclude markers with greater than <integer> Mendelian inheritance errors. Default of 1. This option works in GUI mode.
-minGeno <threshold>	Exclude markers with less than <threshold> fraction of nonzero genotypes. <threshold> must be between 0 and 1 with a default of 0.5. This option works in GUI mode.
-hwcutoff <threshold>	Exclude markers with a Hardy Weinberg p-value less than <threshold>, which ranges from 0 to 1 with a default of 0.001 This option works in GUI mode.
-maxDistance <distance>	Maximum intermarker distance for LD comparisons (in kilobases). Default is 500. This option works in GUI mode.
-missingCutoff <threshold>	Exclude individuals with more than <threshold> fraction missing data, where <threshold> is a value between 0 and 1 with a default of 0.5. This option works in GUI mode.

## Block Output Options

-blockoutput <type>	Generate haplotypes and population frequencies for blocks of <type>, which can be GAB (Gabriel et al), GAM (4 gamete blocks), SPI (solid spine blocks) or ALL (each of the previous 3). The default block type is Gabriel. More information can be found with the blocks help.
-blockCutHighCI <thresh>	Gabriel 'Strong LD' high confidence interval D' cutoff.
-blockCutLowCI <thresh>	Gabriel 'Strong LD' low confidence interval D' cutoff.
-blockMAFthresh <thresh>	Gabriel MAF threshold. Markers below this allele frequency will be skipped in building Gabriel blocks.
-blockRecHighCI <thresh>	Gabriel recombination high confidence interval D' cutoff.
-blockInformFrac <thresh>	Gabriel fraction of informative markers required to be in strong LD.
-block4GamCut <thresh>	4 Gamete block cutoff for frequency of 4th pairwise haplotype.
-blockSpineDP <thresh>	Solid Spine blocks D' cutoff for 'Strong LD'.

## Other Output Options

-check	Output marker quality checks to <inputfile>.CHECK
-mendel	Output Mendel error information to <inputfile>.MENDEL
-malehets	Output chromosome X male heterozygote information to <inputfile>.MALEHETS
-dprime	Output pairwise LD text table to <inputfile>.LD. Note that -dprime and -check default to no haplotype output unless the -blockoutput flag is also specified.
-png	Output PNG image file of LD display to <inputfile>.LD.PNG
-compressedpng	Output low-resolution (smaller file) PNG image of LD display to <inputfile>.LD.PNG
-infoTrack	Include HapMap info track in PNG image outputs
-spacing <threshold>	Use proportional spacing for dumped LD pngs. <threshold> ranges from 0 (no spacing) to 1 (max spacing) with a default of 0.
-ldcolorscheme <type>	Use a particular color scheme for dumped LD pngs. <type> can be DEFAULT, RSQ, DPALT, GAB or GAM. More information can be found with the LD display help
-hapthresh <threshold>	Only output haplotypes with frequency # <threshold>. Note that multiallelic D' and htSNPs are computed using only displayed haplotypes.
-excludeMarkers <markers>	Exclude markers in a comma separated list with ranges specified as start.end. So, to exclude markers 3, 5 and 10 through 15 you'd use "-excludeMarkers 3,5,10..15"

## Association Output Options

-assocCC	Output case/control association results. Saves single marker results to <inputfile>.ASSOC and haplotype results to <inputfile>.HAPASSOC. Haplotype association results are not generated if block type is set to ALL.
-assocTDT	Output TDT association results. Saves single marker results to <inputfile>.ASSOC and haplotype results to <inputfile>.HAPASSOC. Haplotype association results not generated if block type is set to ALL.
-customAssoc <file>	Loads a set of custom tests for association.
-permtests <numtests>	Performs <numtests> permutations on default association tests (or custom tests if a custom association file is specified) and writes to <inputfile>.PERMUT

## Tagging Output Options

-pairwiseTagging	Generates pairwise tagging information in <inputfile>.TAGS and .TESTS
-aggressiveTagging	As above but generates 2-marker haplotype tags unless specified otherwise by -aggressiveNumMarkers
-tagsqcounts	Generates conditional haplotype probabilities from tagger in <inputfile>.CHAPS
-aggressiveNumMarkers <2,3>	Specifies whether to use 2-marker haplotype tags or 2 and 3-marker haplotype tags.
-maxNumTags <n>	Only selects <n> best tags.
-includeTags <markers>	Forces in a comma separated list of marker names as tags.
-includeTagsFile <file>	Forces in a file (or http:// location) of one marker name per line as tags.
-excludeTags <markers>	Excludes a comma separated list of marker names from being used as tags.
-excludeTagsFile <file>	Excludes a file (or http:// location) of one marker name per line from being used as tags.
-captureAlleles <file>	Capture only the alleles contained in a file (or http:// location) of one marker name per line.
-designScores <file>	Specify design scores in a file (or http:// location) of one marker name and one score per line.
-mintagdistance <distance>	Specify a minimum distance in bases between picked tags.
-taglodcutoff <thresh>	Tagger LOD cutoff for creating multimarker tag haplotypes.
-tagsqcutoff <thresh>	Tagger r <sup>2</sup> cutoff.

---

# Chapter 4. About Haploview

Haploview was developed in and is maintained by Mark Daly's lab at the Broad Institute by Jeffrey Barrett, Julian Maller and David Bender. Questions and comments should be addressed to: haploview@broad.mit.edu

- The design of the LD and haplotype interfaces is the work of **Ben Fry** at the MIT Media Lab.
- Thanks to **Andrew Kirby and Hin-Tak Leung** for code contributions.
- Thanks to **Itsik Pe'er and Paul de Bakker** for their extensive contributions to methods development and testing.
- Hardy-Weinberg calculation code courtesy of **Goncalo Abecasis and Jan Wigginton** at the University of Michigan Center for Statistical Genetics
- The  $r^2$  and alternative  $D'$  color schemes are the work of **Will Fitzhugh**.
- The interface to the HapMap GBrowse track is courtesy of **Simon Twigger**.
- PLINK is the work of **Shaun Purcell** at the Center for Human Genetic Research of Massachusetts General Hospital

## Source Code

Haploview is an open source project hosted by SourceForge. The source can be downloaded at the SourceForge project site.

## Citations

Haploview can be cited with the following paper:

Barrett JC, Fry B, Maller J, Daly MJ. Haploview: analysis and visualization of LD and haplotype maps. *Bioinformatics*. 2005 Jan 15 [PubMed ID: 15297300]

Information about the exact test for HW can be found in the following paper:

Wigginton JE, Cutler DJ, Abecasis GR. A note on exact tests of Hardy-Weinberg equilibrium. *Am J Hum Genet*. 2005 May;76(5):887-93.

Information about parentTDT can be found in the following paper:

Purcell S, Sham P, Daly MJ. Parental phenotypes in family-based association analysis. *Am J Hum Genet*. 2005 Feb;76(2):249-59.

Questions, complaints and suggestion should be directed to haploview@broad.mit.edu.

# System Requirements

It is recommended that Haploview be run on a machine with at least 128M of memory. The Haploview jarfile should now automatically allocate extra memory when starting up, so the `-Xmx` flag is no longer required when running the program from the command line.

Haploview requires Java JRE 1.3 or later, but 1.4 is strongly recommended. It is worthwhile in any case to download the most recent Java release.

## Known Issues

We are working to correct a rare issue with Spineblock outputs for certain datasets.

Saving PNG files from the command line (the `-png` and `-compressedpng` flags) requires Java version 1.4 or later.

There appears to be an issue with the Windows JDK version 1.4.1\_03. Note that all other versions (both earlier and later versions of 1.4.1 included) seem to work fine. If you find this to be your current Java version (type `java -version` at the command prompt), please install the latest version.

## Updates

If you have an internet connection, Haploview will automatically check for an update upon startup. If a new version is available, it will show a message in the lower right corner of the screen for a few seconds. Details can be found by using the "Check for Updates" button in the File menu.

## Change Log

### Version 4.0, 21 August 2007

- added support for HapMap PHASE format data
- added support for phased HapMap downloads
- added support for PLINK output files including plotting functionality
- added a number of Tagger features to coincide with the Tagger website functionality
- added 2-marker aggressive tagging from the command line
- added log file functionality to command line mode
- added proxy support
- added table sorting to Tagger configuration
- added numerous minor display and interface tweaks
- added export options for Mendel errors and male heterozygotes on x chromosome
- added command line HapMap info track download
- added case control frequencies to command line association output
- added ld values selection to command line image export
- added citation information to the "About" dialog
- added "Uncapture All" button to Tagger
- added build 36 option to HapMap info track downloads
- added `http://` loading for files on the command line and initial load screen
- changed some labelling in Tagger and made pairwise the default option

- changed obsHet calculation to use founders only
- changed HW and het calculations to show as "NA" in X data with too few individuals
- enhanced GUI design with tabbed pane for initial loading dialog
- fixed bug with haplotype associations for extended trios and x chromosome
- fixed bug with case control counts for certain zeroed out markers
- fixed Mendel error reporting bug on x chromosome
- fixed Tagger bug affecting maximum number of tags input
- fixed bug when left-clicking on the LD plot
- fixed bug when changing tabs after multiple file loads
- fixed bug when changing tabs after running tagger and changing check marker thresholds
- fixed bug when loading custom association test file with multi-marker tests but no alleles
- fixed bug with long windows filenames in command line mode
- fixed bug with display when resizing the window to a very small size
- fixed bug with 'h' alleles in haps files not being properly calculated
- fixed bug with check panel export not accounting for user threshold adjustments
- fixed bug with LD display where some  $r^2$  values were invisible against the background
- fixed bug with integer overflow when loading files with tens of thousands of individuals
- fixed bug where HapMap info track would turn black if it was too wide
- fixed bug with null pointer exception on failed hapmap info track downloads
- fixed bug with null pointer exception when using Export Options Dialog
- fixed bug with LD Plot now showing the correct markers when changing marker ratings and using table sorting in the check panel
- fixed tagger results reporting of % captured when specifying alleles to capture
- fixed bug with marriage loops in pedigree files causing StackOverflow errors
- fixed bug with exception being thrown for certain input files with no markers
- fixed bug with multimarker tags showing in the tagger results even though they don't capture anything
- fixed bug with command line error handling
- fixed bug with loading improperly formatted info files from "Load marker data"

## **Version 3.32, 21 June 2006**

- fixed bugs in check markers display when using table sorting
- fixed line number output on error messages

- grammar corrections

## **Version 3.31, 2 June 2006**

- fixed bug where chromosomes were created in reverse

## **Version 3.3, 26 May 2006**

- support for X chromosome
- major memory usage improvements
- allow multiple analysis tracks and label Y axis
- added user defined hapmap sample option
- added haplotypes only permutation option
- added option to show different LD measures or hide them
- added -memory switch and changed default to 512M
- added minor allele to check panel
- added progress bar for data loading
- support for alphabetical (ACGT) input format
- added individual information dialog
- added check panel for phased haps files
- added dump tags button
- last popup dialog is shown at top of display
- added GOLD style color scheme
- can sort tables by clicking on headers
- changed case control allele order
- fixed grey screen with no text bug
- fixed tagger bugs for maxnumtags and forceinclude
- fixed batch mode for hapmap files
- fixed Tagger bug where some SNPs were tagged by more than one tag.
- fixed export options bug with association tab
- fixed generateHaplotype bug for matching alleles

## **Version 3.2, 13 April 2005**

- added Tagger interface

- added custom association test input
- added permutation testing of association results
- improved memory efficiency for EM haplotype reconstruction
- added Perlegen sample IDs
- fixed bug in PNG export cutting off final marker

## **Version 3.11, 04 February 2005**

- improved parsing of families with unrelated members
- fixed bug with singletons with large amounts of missing data
- fixed bug with java path differences from cmd line
- fixed export issues for LD display
- fixed bug with block output for files without blocks

## **Version 3.1, 27 January 2005**

- Additional HapMap info track display options.
- Improved parsing of complex pedigrees.
- Fixed bug with loading of haps style input files.
- Fixed bug with correctly parsing out-of-order info files.
- Fixed problem with association tests on datasets with no blocks.
- Fixed bug with exporting range of markers to LD PNG.

## **Version 3.0, 7 October 2004**

- bugfixes for block size display, checkdata tab, allele sorting, EM missing data.
- Added compressed PNG output mode.
- Lots of new command line options.
- Changed batch mode input format.
- Re-sort out of order info files.
- Substantial improvement in speed and memory usage in EM.
- New color schemes.
- Added additional sample information for 2nd and 3rd HapMap plate.
- Fixed handling of complex pedigrees.
- Added proportional spacing to LD display.

- Added HapMap GBrowse track.
- Filter individuals with lots of missing genotypes.
- Haplotype association tests.
- Update checker.

## **Version 2.05, 27 April 2004**

- Fixed problem with EM for long blocks
- Added "Export options" to allow exporting subsets of data and LD values for only adjacent marker pairs.
- Numerous minor bugfixes
- Added block labels to haplotype display
- Added block size to LD display
- Added saved block definition input file
- Added analysis track input option
- Enabled direct click from HapMap webpage to Haploview

## **Version 2.04, 21 January 2004**

- All color schemes now allowed with all block definitions, including hand-defined blocks.
- Enabled loading input file from command line while still opening GUI.
- Added command line HapMap Project input option.
- Added colored box haplotype display.
- Tweaked command line flags.
- Added command line checkdata only output.
- More accurate Hardy-Weinberg code implemented (courtesy of G. Abecasis & J. Wigginton).
- Correctly deals with genotype inputs with poorly genotyped markers.

## **Version 2.03, 18 December 2003**

- Correctly handles new HapMap Project dump format.
- Added minor allele frequency filter to data check tab.

## **Version 2.02, 05 December 2003**

- fixed confidence bounds coloring scheme bug
- fixed text output bug where marker numbers were incorrect

- added marker spacing map to top of HapMap datasets
- added T-int statistic to text dump of LD chart

## **Version 2.01, 31 October 2003**

- fixed bug involving European style decimal format (e.g. 0,45 vs. 0.45)
- when exporting data, default file name is now blank
- marker info file now works in either forward or reverse direction
- activated loading of HapMap Project dumped data