# User Manual for GIGI v1.02

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## Introduction

GIGI is a C++ program to impute genotypes in pedigree data. It imputes missing genotypes of dense markers conditional on genotypes available on a subset of relatives in the pedigree and framework genotypes used to infer Inheritance Vectors (IVs). Our pedigree-based genotype imputation approach consists of two steps. The first step is to infer IVs at the positions of framework markers using gl\_auto, which is a MCMC-based program that is part of the MORGAN package. The second step is to impute dense genotypes by GIGI using the IVs output by gl\_auto and pedigree structure file from MORGAN.

GIGI is developed under the linux environment.

In this documentation, we use the following terminology. Framework markers are a relatively sparse set of markers that are used to infer IVs on a chromosome of interest. Dense markers are markers with missing genotypes on some subjects that we want to impute. For example, these dense markers may be genotypes obtained from sequence data or from a dense SNP panel, and may be typed on fewer and even different subjects in the pedigree. See the publication describing GIGI, below, for more information.

## **Citing GIGI**

Cheung, C.Y.K., Thompson, E.A., Wijsman, E.M. (2013). GIGI: An approach to effective imputation of dense genotypes on large pedigrees. *American Journal of Human Genetics*, [in press – to appear in the April, 2013 issue]

## Software URL

http://faculty.washington.edu/wijsman/software.shtml

## Files in GIGI software distribution

GIGI software code and its dependency files - the Mersenne random number generator

• example folder

## Inferring IVs using gl\_auto

The first step to impute genotypes is to use **framework markers** to infer IVs. For this purpose, we use gl\_auto, a program in the MORGAN package that is freely available at <u>http://www.stat.washington.edu/thompson/Genepi/MORGAN/Morgan.shtml</u>. To infer IVs in gl\_auto,

- we need to supply the following required files in MORGAN format:
  - (1) Pedigree file
  - (2) Marker file: this is a composite file that contains the map positions of framework markers (in centiMorgans assuming the Haldane map function), allele frequencies of framework markers , and genotype data of framework markers
  - (3) Parameter file used to run gl\_auto

Refer to the documentation of MORGAN for guidance on setting up these files and on running gl\_auto.

Example files used to infer IVs using gl\_auto are included under the "example/gl\_auto\_example" directory.

#### Overview

From running gl\_auto, we need to obtain 2 files:

- (a) Framework IVs file: GIGI uses this file, which contains IVs at framework positions
- (b) Pedigree-meiosis file: this file contains the information about the structure of the pedigree on which we want to impute genotypes. This file is different from the pedigree file used in gl\_auto. In addition to the pedigree structure, this file also contains information that GIGI needs to determine how the Inheritance Vectors (Meiosis indicators) are organized (i.e. the i<sup>th</sup> line of the meisosis indicator belongs to which subject in the pedigree, and whether this meiosis indicator is this person's maternal or paternal chromosome). We need to create this file from the console output of gl\_auto.

#### **Obtaining the files**

#### (a) Framework IVs file

In gl\_auto's parameter file, we must instruct gl\_auto to display the output as Meiosis Indicators instead of Founder Genome Labels. Please make sure we use the option:

#### output meiosis indicators

instead of "output founder genome labels" in the control/parameter file of gl\_auto.

#### (b) Pedigree-Meiosis file

We need to create the pedigree-meiosis file (b) from the console output of gl\_auto. It is very easy to make this file.

When we run gl\_auto, the program prints a huge amount of output to the console. This console output actually contains the content of the pedigree-meiosis file that we need to extract.

1. In order to extract this content, we first need to direct the console output to a file by using the ">" directive so we can subsequently extract the content from this file. i.e., type:

#### ./gl\_auto gl\_auto\_parameter\_file > glauto\_console\_output.txt

- 2. Then, we extract the pedigree-meiosis content from the console output to a new file. To simplify the creation of this file, use the Perl script "extractPedMeiosis.pl"
  - Usage: perl extractPedMeiosis.pl glauto\_console\_output.txt FILENAME\_PED\_MEIO
    - We need to have Perl installed under linux.
    - We assume that glauto\_console\_output.txt is in the same directory as extractPedMeiosis.pl
  - Alternatively, this file can also be easily created by the user. See Appendix B for the creation of this file manually and also for an example of what this file looks like.

## **Installing GIGI**

Simply unzip the files, navigate to the code directory, and type: make

If make does not work, go to the GIGI.cpp's directory and install the program by typing: g++ GIGI.cpp -o GIGI

GIGI was compiled successfully under Ubuntu 10.04, Mandriva Linux release 2009.0, and RedHat release 5.8, with g++ version-4.1.2, 4.3.2, and 4.5.2.

## **Running GIGI**

GIGI accepts a parameter file. To run GIGI, type :

./GIGI <parameter file> <options>

To run the example file, go to the main GIGI program directory, and type: ./GIGI example/param.txt

# **Options**

[The flags are case-sensitive.]

Flag	Purpose			
-verbose	to run GIGI with extended output to the console			
-seed=NUM	to change the default seed. NUM should be a number between 1 and 999999			
-prog=NUM	to print the progress of imputation at every NUM markers			
-longParam	to run GIGI using the old (GIGI v1.00's) format of the parameter file [General users			
-	should ignore]			
-outD=	to specify the absolute directory path of where the output files will be created. If this			
	flag is missing, the output files will be saved to the user's current directory.			
	e.g. /home/charles/GIGI_output [no character '/' at the end of the directory name]			

An example of running GIGI with additional options:

./GIGI example/param.txt -verbose -outD=/home/charles/GIGI\_output

## Making the Parameter File

The parameter file tells GIGI where to look for the required files and where to save the output files. GIGI needs the pedigree meiosis file (that user prepares from the output of gl\_auto), dense marker file, IVs file (from gl\_auto), map of the sparse marker file, map of the dense marker file, and allele frequencies of the dense marker file. It produces the called genotype file, genotype probability file, and another file that prints the number of IVs that are consistent with the observed dense genotypes [for sanity check].

An **example** of the parameter file is found in the example directory under: **example/param.txt** 

In GIGI\_v1.01, the parameter file is organized as follows:

- line 1 filename of the pedigree meiosis file
- line 2 framework IVs file
- line 3 number of sampled realizations in the IV file

line 4 - filename of the **map positions** of the **framework** markers file (cM based on Haldane map function)

- line 5 filename of the map positions of the dense markers file (cM based on Haldane map function)
- line 6 filename of the **dense marker genotype** file
- line 7 filename of the **allele frequencies** of the **dense** markers file
- line 8 Call\_method  $t_1 t_2$  (3 numeric values separated by space: e.g. 2 0.6 0.8)

On line 8, we specify up to 3 values.

1. Call\_method: use the value 1 to call the most likely genotype and 2 to use threshold-based calling

2 and 3. If threshold-based calling is used, then the threshold of  $t_1=0.8$  and  $t_2=0.9$  are set as defaults. This pair of thresholds can be changed by the user by specifying the 2<sup>nd</sup> and 3<sup>rd</sup> numerical values. The thresholds should be between 0.5 and <1. If only  $t_1$  is specified,  $t_2$  will be set to  $t_1+(1-t_1)/2$ .

The format of the parameter file used here is different than that of an older format used in GIGI\_v1.00. See Appendix A for the older format and how one can continue to use the older format of the parameter file. [not recommended for newer users]

Notes:

I suggest using the absolute paths of the filenames instead of relative paths. A relative path is relative to the directory containing the executable program. (The parameter file in the example folder is created using a relative path.)

\*line 1: user creates this file from the gl\_auto's console output. In version 1.01, GIGI can only process 1 pedigree at a time.

\*line 2: when you run gl\_auto, you should instruct gl\_auto to print Meiosis Indicators instead of Founder Genome Labels

\*line 3: – this corresponds to the number of samples that the user actually prints to the meisois indicator file.

## **File Formats**

# Examples of these files are provided in the example directory [refer to the param.txt for the filename of these files]

a. pedigree meiosis file [line 1]. The creation of this file was discussed in the **Inferring IVs using** gl\_auto section.

The pedigree meiosis specifies the pedigree structure and the index of meioses that GIGI needs to use to read in the corresponding Inheritance Vectors. GIGI imputes genotypes on subjects specified in this

pedigree. The content of this file is generated by gl\_auto.

b. framework IVs file [line 2]

The Inheritance Vectors file describes the descent pattern of chromosomes at the positions of the framework markers. It is the output file that **gl\_auto** generates.

c. map positions of framework markers [line 4]

The marker map positions of the framework markers file is a text-file which contains the map distance in centi-Morgans (cM) based on the Haldane map function. Markers must be ordered in ascending order and consistent with the order used in gl\_auto. Each line contains the position of a marker.

position of Marker1 position of Marker2 position of Marker3 ... position of MarkerN

e.g. 1.0 2.0 3.0 4.0

e. map positions of dense markers [line 5]

Similar to the marker map for framework panel of markers, the marker map of dense markers is a textfile which contains the map distance in centi-Morgans (cM) based on the Haldane map function. Markers must be ordered in ascending order. Each line contains the position of a marker.

e.g. 0.5 0.7 0.9 1.1 1.15

•••

b. dense marker genotype file [line 6]

The dense markers are the markers that we want to impute on some individuals in the pedigree. The dense marker file contains the genotypes of observed individuals.

- (a) This file is space-delimited
- (b) The markers should be sorted by ascending map positions.
- (c) Alleles are labeled numerically starting from 1,2,... in ascending order. We use 0 to indicate a missing allele. *If the original genotypes are in alpha-numeric coding, users first need to convert the genotypes to an indexed numerical format.* (GIGI supports imputation of multi-allelic markers.)

Each row of the file contains the genotypes for an individual. This file does not include the header line. The first column specifies the ID of an individual. The ID can be an alphanumeric string.

ID allele1\_marker1 allele2\_marker1 allele1\_marker2 allele2\_marker2 allele1\_marker3 allele2\_marker3 ...

An example of this file looks like:

101 1 1 1 2 2 2 102 1 2 0 0 2 2 103 1 1 1 2 1 2

6. allele frequencies of the dense marker file [line 7]

Each row contains the allele frequencies of the dense markers. The first column is the allele frequency of allele 1, the second column is the allele frequency of allele 2, etc... Each row must sum to 1. The allele frequencies file is space-delimited.

```
allele frequencies of marker 1
allele frequencies of marker 2
allele frequencies of marker 3
...
e.g.
```

0.4 0.6 0.2 0.3 0.5 0.8 0.2

## **Output files**

Three output files are created in the directory where GIGI is run:

1. impute.geno

The imputed genotype file has the same format as the input dense genotype file.

2. impute.prob

The imputed genotype probabilities file displays the genotype probabilities for each marker. For each bi-allelic marker, three columns are used to store the probability of genotype configuration 1/1, 2/1, and 2/2. For each tri-allelic marker, the probabilities are displayed and space-delimited with 1/1, 2/1, 2/2, 3/1, 3/2, 3/3. For multi-allelic markers in general, the appropriate number of columns are used to store all the combinations.

Markers are separated by tabs.

e.g.

ID genotype\_11\_marker1 genotype\_21\_marker1 genotype\_22\_marker1 genotype\_11\_marker2 genotype\_21\_marker2 genotype\_22\_marker2

•••

101 0.9 0.1 0	0.8 0 0.2
102 0.9 0.1 0	100

3. impute.consistentIV

The consistency file counts the number of IVs consistent with the observed genotypes of each dense marker. This file is used for a sanity check. A number that is low compared to the number of IVs used may indicate incompatibility between observed genotypes and the sampled IVs.

## Frequently Asked Questions (FAQs)

- 1. How do I choose framework markers? Because the framework markers are assumed to be sparse, we want to choose framework markers that are informative about which chromosomes are being transmitted at the framework loci. First, framework markers typed on a large number of subjects tend to be most informative. Second, framework markers that are multi-allelic tend to be more informative than di-allelic markers if they are available. Third, if framework markers are SNPs, markers with high minor allele frequencies in the sample tend to be more informative. Fourth, framework markers should be moderately but not too dense (e.g. not denser than 1 marker per 0.3 cM because of concern about MCMC mixing and violation of the assumption of Linkage Equilibrium). If the framework markers are multiallelic, this spacing should be greater, e.g., not denser than 1 marker per 2 cM for good MCMC mixing.
- 2. What is necessary about the genetic map that I am using? The map positions are in map distances based on the Haldane map function, instead of Kosambi map function or sequence positions. If this map is based on the Kosambi map function, the user will need to convert the map positions to Haldane map function using an appropriate conversion method.

Also, since recombination fractions are relative to each other, we strongly encourage the user to generate both the framework map positions and the dense marker positions at the same time.

- **3.** Can GIGI process multiple pedigrees? No. In version 1.01, GIGI can only process 1 pedigree at a time. For this reason, gl\_auto should also be run individually for each pedigree.
- 4. Can I use GIGI to impute genotypes on unrelated individuals? No.
- 5. What should I know when I use GIGI to impute genotypes on trios? Since there is no way to infer recombination on trios using unphased genotypes, gl\_auto produces an empty inheritance vectors file. From the pedigree-meiosis file, GIGI recognizes that this inheritance vectors file is empty but will proceed to attempt impute genotypes. For trios, imputing genotypes using GIGI is equivalent to imputing genotypes conditional on the pedigree structure and minor allele frequencies.

Information about linkage-disequilibrium is potentially useful for genotype imputation on trios. Since the current version of GIGI does not leverage information from linkage-disequilibrium, you may want to use existing linkage-disequilibrium-based software that supports imputation on trios (e.g. BEAGLE).

# Appendix A

GIGI v1.00 used an old format for the parameter file. It is not recommended for new users. For users of v1.00, if you want to continue to use this old format, use the flag "-longParam" when you run GIGI. This file format allows you to specify the exact filename and location of the output files. e.g.: ./GIGI example/oldparamFormat.txt -longParam

The old file format:

- line 1 filename of the pedigree meiosis file
- line 2 number of pedigrees, current max is 1. Just use the value  $\mathbf{1}$
- line 3 filename of the dense marker file
- line 4 IVs file
- line 5 number of saved realizations in the IVs file
- line 6 filename of the map positions of the framework markers file (cM Haldane map distance)
- line 7 filename of the map positions of the dense markers file (cM Haldane map distance)
- line 8 filename of the allele frequencies of dense markers file
- line 9 random seed number (e.g. 1234)
- line 10- filename of the output 1: called genotypes

line 11- Use the value **2**. This is the calling method to use. 1: most likely genotype calling, 2: threshold-based calling

line 12- the first threshold of threshold-based calling: t1- to call both alleles (this number should be between 0.5 and <1). e.g. 0.9

line 13- the second threshold of threshold-based calling: t2- to call 1 of 2 alleles, if GIGI can't call both alleles under the threshold specified in line 12. (a number that is between the first threshold <1). e.g. 0.9

line 14- filename of output 2: estimated genotype probabilities

line 15- filename of output 3: number of IVs consistent with each observed dense markers

# **Appendix B**

Creating the Pedigree-Meiosis file manually from the console output of gl\_auto.

Using a text editor, we open the console\_output.txt and fetch the line that begins with "name name.pa name.ma Compnt pat.meio mat.meio" We copy this line and table below. We paste this table to another file and save it

We copy this line and table below. We paste this table to another file and save it. The file includes the header line and looks like this:

name	name.p	a na	me.ma C	ompnt	pat.meio	mat.meio
2100_	6	0	0	1	0	0
2100_	21	0	0	1	0	0
2100_	25	0	0	1	0	0
2100_	29	0	0	1	0	0

2100_31	0	0	1	0	0			
2100_39	0	0	1	0	0			
2100_907	0	0	1	0	0			
2100_908	2100_901	21	00_9	02	1	2	1	
2100_909	0	0	1	0	0			
2100_910	2100_901	21	00_9	02	1	4	3	
2100_911	2100_901	21	00_9	02	1	6	5	
2100_915	2100_907	21	00_9	08	1	8	7	
(until the end of table)								

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## **Future Directions**

- 1. Enable GIGI to read marker genotype data in the long file format, similar to the format used in BEAGLE: e,g, each row contains a marker.
- 2. Convert the code from C++ to C