

# HaploGI – Haplotyping Given Inheritance

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## Introduction

HaploGI (Haplotyping Given Inheritance) is a C++ program for pedigree-based haplotyping of whole genome sequencing (WGS) data. It also identifies haplotype sharing among subjects in extended pedigrees.

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## Paper Citation

If you use **HaploGI** in your work, please cite:

Nafikov, R. A., Sohi, H., Nato Jr, A. Q., Horimoto, A. R., Bird, T. D., DeStefano, A., Blue, E. E., & Wijsman, E. M.

*Variant prioritization by pedigree-based haplotyping*. Submitted for publication to *Genetic Epidemiology*, 2025.

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## Software Citation

If you use **HaploGI** in your research, please also cite the following:

### [package] Software

DOI [xx.xxx/zenodo.xxx](https://doi.org/xx.xxx/zenodo.xxx)

```
@software{nafikov_2025_haplogi,  
  author      = {Rafael A. Nafikov},  
  title       = {HaploGI - Haplotyping Given Inheritance},  
  version     = {1.0.0},  
  year        = {2025},  
  publisher    = {Zenodo},  
  doi         = {xx.xxx/zenodo.xxx},  
  url         = {https://doi.org/xx.xxx/zenodo.xxx}  
}
```

This repository includes a CITATION.cff file.

On GitHub, click the “**Cite this repository**” button near the top to get citation details in various formats.

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## License

HaploGI is licensed under the GNU General Public License v3.0.  
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## Software URL

Repository: <https://github.com/RafPrograms/HaploGI>

Files available for download: - HaploGI.cpp (source code) - manual\_HaploGI\_v1.0.0.pdf (PDF) (user manual) - parameter\_file\_template.txt (parameter file template) - HaploGI\_utility\_scripts (Python utility scripts) - HaploGI\_test\_data.zip (example dataset)

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## Build and Install

You can build HaploGI from source using CMake:

Requirements CMake  $\geq 3.10$

C++17-compliant compiler (e.g., g++ ≥ 7, clang ≥ 5)

Unix-like environment (Linux/macOS recommended)

### Build Instructions

```
# Clone the repository (if not done yet)
git clone https://github.com/RafPrograms/HaploGI.git
cd HaploGI
```

```
# Create a separate build directory
mkdir build
cd build
```

```
# Generate Makefiles with CMake
cmake ..
```

```
# Compile the program
make
```

[rocket] Run from Build Directory After compiling, you can run HaploGI directly from the build directory:

```
./HaploGI -- [options] [parameter_file_path]
```

---

### Optional: Install System-Wide

To install the compiled binary to your system path (default: /usr/local/bin):

```
sudo make install
```

This will allow you to run HaploGI from anywhere in your terminal.

[idea] Note: The binary is installed to the bin/ directory under your system's CMAKE\_INSTALL\_PREFIX (default: /usr/local/bin).

**Custom Install Location** You can specify an install prefix:

```
cmake -DCMAKE_INSTALL_PREFIX=/your/custom/path ..
make
sudo make install
```

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### Getting Started

Launch HaploGI using the following syntax:

```
./HaploGI -- [options] [parameter_file_path]
```

## Run Options

Option	Description
--haplotyping	Pedigree-based haplotyping + core set of cases identification
--haplosharing	Evaluate haplotype sharing in predefined cases
--full	Combines both haplotyping and haplosharing

## General Options

Option	Description
--help	Display help
--version	Show current version

## Parameter File

See: `parameter_file_template.txt`

### Required for all run types:

Entry	Description
1#	Pedigree file path
2#	SNV genomic positions file path
3#	SNV genotypes file path
4#	Linkage region boundaries (cM)
5#	Max LOD marker position (cM)
6#	Output directory

### Required for --haplotyping and --full:

Entry	Description
7#	Linkage markers genomic positions file
8#	Meiosis indicators file
9#	Number of iterations in indicator file

**Required for --haplosharing:**

Entry	Description
10#	Haplotype sequences file
11#	Core set of cases file

**Optional:**

Entry	Description
12#	Seed number (default: 1234)

**Input File Formats**

All files must be **space-delimited**.

**Required per Run Option**

File	Required for
Pedigree file	All
SNV genomic positions	All
SNV genotypes	All
Linkage markers positions	haplotyping, full
Meiosis indicators	haplotyping, full
Haplotype sequences	haplosharing
Core cases	haplosharing

**Example Input Files**

**Pedigree File Format:**

```
subject father mother sex phenotype
*****
101 0 0 1 0
102 0 0 2 0
201 101 102 1 0
202 101 102 2 0
2010 0 0 2 0
301 201 2010 1 0
302 201 2010 2 2
```

The pedigree file contains **five space-delimited columns** with the following information:

1. **Subject ID**
2. **Father ID**
3. **Mother ID**
4. **Sex**
  - 1 = Male
  - 2 = Female
  - 0 = Unsexed / unknown
5. **Phenotype**
  - 1 = Control
  - 2 = Case
  - 0 = No phenotype data

#### **File Format Notes**

- **Header lines (above the main data) are ignored** by the program if they appear **before a line starting with \*.**
- You may include column headers before this marker.
- **IDs must not contain special characters** such as #, \*, or @.

**Note:** The **\*\*\*\*\*** line acts as a marker—any content above this line is ignored during processing.

---

#### **SNV Genomic Positions File Format:**

```
1052701 3.767099
1052874 3.767696
1053095 3.768460
1053154 3.768664
```

This file contains genomic position data for each single nucleotide variant (SNV), with one SNV per line.

#### **File Format**

- **No header row**
- Each line contains **two space-delimited columns:**

1. **Base pair (bp) position** – The physical location of the SNV on the chromosome
2. **Genetic position in centimorgans (cM)** – The corresponding genetic map position

**Note:** Ensure the order of positions in this file matches the order of SNVs used in related genotype and haplotype files.

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### SNV Genotype File Format:

```
variant_position 302 302 303 303 306 306 307 307 402 402 403 403 404 404 406 406 407 407 408
16:10414 1 1 1 1 1 1 2 1 1 1 1 1 1 1 1 1 1 1 2 1 1 1 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
16:10638 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
```

The SNV genotype file contains variant genotype data for subjects with whole genome sequencing (WGS) data. Genotypes are encoded as:

- 0 = Missing
- 1 = Reference allele (REF)
- 2 = Alternative allele (ALT)

### File Structure

- **First column:**  
Contains the SNV's genomic position in the format `chromosome:position` (e.g., `16:10414`).
  - **Remaining columns:**  
Each subject is represented by **two consecutive columns**, one for each of their diploid genotype alleles.
  - **Header row:**  
Lists subject IDs. Each subject ID appears **twice**, corresponding to their two genotype alleles.
- Note:** Ensure that subject IDs are consistent across files and that each subject has exactly two columns representing diploid genotype data.

[tools] Use `prepare_genotype_file.py` to generate an SNV genotype file from a VCF file for use in HaploGI runs.

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### Linkage Markers Genomic Positions File Format:

0.219846  
1.134855  
1.793034

This file contains the genetic positions (in centimorgans, cM) of linkage markers used to compute inheritance vectors with the **Morgan package**.

#### File Format

- **No header row**
- Each line contains a **single centimorgan (cM) position** for one linkage marker
- Markers are listed in the order expected by downstream analysis tools

**Note:** Ensure that the number and order of cM positions match the corresponding linkage marker set used in the analysis pipeline.

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**Meiosis Indicators File Format:** The **meiosis indicators file** is generated by the `gl_auto` program from the Morgan package.

It follows the same format as described in the official Morgan package manual.

**Note:** This file encodes inheritance information and is used in downstream linkage and haplotype analyses.

[tools] Use `decrease_number_of_MI_iterations.py` to generate a meiosis indicators file with a reduced number of iterations (recommended: 1000) to ease the computational burden on HaploGI.

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#### Haplotype Sequences File Format:

```
16:10414-23730 302_0 11111111112111111211
16:10414-23730 302_1 11112111222111111112
16:10414-23730 303_0 11111111222111111112
16:10414-23730 303_1 11111111222111111112
16:10414-23730 306_0 11112111222111111112
16:10414-23730 306_1 11111112112112111121
```

The haplotype sequences file is generated by **HaploGI** using either the `--haplotyping` or `--full` run options.

This file contains **three columns with no header**:

##### 1. Genomic Range

A string representing the chromosome and variant range in the format:

`chr:start-end`

- `chr`: Chromosome number



- **start and end:** Positions of the first and last genomic variants in the haplotype
2. **Subject ID and Chromosome**  
The subject identifier followed by an underscore and a digit:
    - **\_0:** Maternal chromosome
    - **\_1:** Paternal chromosome
  3. **Haplotype Sequence**  
A string of digits representing the sequence of genomic variants for the given region.
- Note:** This file does not include a header row. Be sure to account for that when parsing the file programmatically.

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### Core Set of Cases File Format:

302 306 403 408 411 501 504 506 511 512 513 516

This file contains a list of **case subject IDs**, separated by spaces, all on a **single line**.

- **No header row**
- IDs must match those used in other input files (e.g., pedigree, genotype, haplotype files)

HaploGI uses this set of cases to **check for the existence of haplotype sharing** among them.

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## Output File Formats

File	Generated by
Log	All
Haplotype sequences	haplotyping, full
Core cases	haplotyping, full
Allele inconsistencies	haplotyping, full
Shared haplotypes	haplosharing, full
Haplotype sharing patterns	haplosharing, full

### Example Output Files

**Haplotype Sequences File** – see **Haplotype Sequences File Format** [tools] Use create\_phased\_vcf.py to convert phased whole-genome sequencing (WGS) data generated by HaploGI into VCF format, enabling easier downstream analysis.

## Core Set of Cases – see Core Set of Cases File Format

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### Inconsistencies of Allele to FGL Assignments File Format:

```
bp_position
1445745
1455891
1458974
```

This file lists variants for which inconsistencies between alleles and Founder Genome Labels (FGL) were detected.

### File Format

- The file **includes a header row**.
- Each subsequent line contains the **base pair position** of a variant with detected inconsistency.
- One base pair position per line.

### Notes

- This file is intended for **exploratory purposes only**.
  - It is **not required** for haplotyping or determining haplotype sharing.
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### Shared Haplotype Sequence File Format:

```
16:1052701-1055604 22222212211222211212
16:1127696-1132994 11111222221112211212
16:1506499-1511322 11111121111111111111
16:1511338-1513919 11111111111111111111
16:1514349-1520077 11111111111111111111
```

This file contains haplotype sequences shared within genomic windows among cases listed in the core set of cases file.

### File Format

- The file has **two columns** and **no header**.
- The **first column** specifies the genomic window location, formatted as: chromosome\_number:first\_variant\_bp-last\_variant\_bp (e.g., 16:1052701-1055604)
- The **second column** contains the haplotype sequence corresponding to that genomic window.

[tools] Use risk\_haplotype\_sequence\_vcf.py to convert risk haplotype sequences generated by HaploGI into VCF format for convenient downstream analysis.

This file contains information about haplotype sharing across all genomic windows evaluated.

## File Format

- The file **includes a header row**.
- The **first column** contains genomic window numbers.
- Each subject with WGS data is represented by **two consecutive columns**:
  - One for the maternal chromosome
  - One for the paternal chromosome
- Entries indicate the presence of a shared haplotype:
  - 1 = Shared haplotype present in a **case**
  - 2 = Shared haplotype present in a **control**
  - 0 = No shared haplotype present.

## Usage

You can use the provided Python utility script `plot_haplo_type_sharing.py` to generate these plots.

[tools] Use `plot_haplotype_sharing.py` to visualize haplotype sharing data generated by HaploGI. This script helps identify the presence and boundaries of risk haplotypes. The resulting plots also provide a broader overview of haplotype sharing, supporting more informed decision-making during data analysis.

A number of Python utility scripts are available to assist with preparing HaploGI input files, processing output data, and visualizing results, at: [HaploGI utility scripts](#).

Other Python utility scripts that have not been introduced yet in this manual are:

[tools] risk\_alleles\_variants.py – Identifies and outputs variants whose alleles are uniquely present on the risk haplotype. The resulting file also includes associated metadata extracted from the input VCF file.

[tools] create\_genomic\_windows.py – Generates a genomic windows file from an SNV genomic positions file used in HaploGI runs. This facilitates easier cross-referencing of data to specific genomic regions.

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## Data Example

A full example with three run configurations is available in `data_example`.

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## Support

For questions, bug reports, or suggestions, please contact:  
nrafscience@gmail.com  
GitHub Issues

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## References

1. Nafikov et al., (2025). Variant prioritization by pedigree-based haplotyping. *Genetic Epidemiology*.
  2. Nafikov et al., (2018). Dealing with Admixture in Caribbean Hispanic Families. *Genetic Epidemiology*. DOI:10.1002/gepi.22133
  3. Tong & Thompson (2007). Multilocus lod scores. *Human Heredity*. DOI:10.1159/000109731
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## Web Resources

- **HaploGI**: <https://github.com/RafPrograms>
- **Morgan Package**: Morgan site
- **1000 Genomes Project**: <https://www.internationalgenome.org>