

hlabud: HLA genotype analysis in R

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Summary

The human leukocyte antigen (HLA) genes have thousands of different alleles in the human population, and have more associations with human diseases than any other genes. Data for all known HLA genotypes are curated in the international ImMunoGeneTics (IMGT) database, and the Allele Frequency Net Database (AFND) provides allele frequencies for each HLA allele across human populations. Our opensource R package *hlabud* facilitates access to HLA data from IMGT/HLA and AFND, and provides functions for HLA divergence calculations, fine-mapping analysis of amino acid (or nucleotide) positions, and low-dimensional embedding.

Availability

Source code and documentation are available at github.com/slowkow/hlabud

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1. Introduction

Human leukocyte antigen (HLA) genes encode the proteins that enable cells to display antigens to other cells, so the immune system can recognize pathogens such as bacteria and viruses. Geneticists have identified thousands of variants (e.g. single nucleotide polymorphisms) in the human genome that are associated with hundreds of different diseases and phenotypes [1].

HLA nomenclature consists of allele names like *HLA*01:01* to indicate the genotype of each individual in a study. Each allele name corresponds to multiple mutations at different positions throughout the gene's sequence, so it is difficult to estimate the similarity of two alleles solely from the allele names. This ambiguity about specific amino acid positions means that allele names are not ideal for statistical analysis.

Researchers have developed software tools for calling HLA genotypes (Figure 1) with high accuracy from DNA-seq or RNA-seq next-generation sequencing reads [2], so there may be opportunities to use this type of data for HLA association studies. Most software tools report allele names, not genotypes at specific

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nucleotide positions. Providers of HLA typing services often report genotypes with the traditional HLA allele names (i.e. *HLA*01:01*) instead of reporting alleles at specific nucleotide positions (Figure 1).

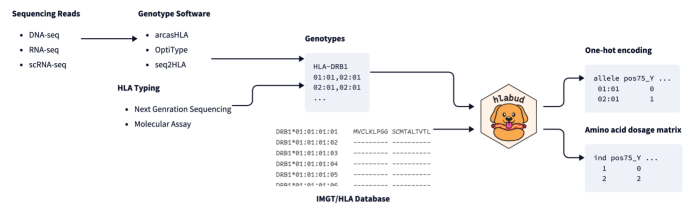


Figure 1: hlabud converts HLA genotypes to amino acid position matrices.

In contrast to allele-level analysis, fine-mapping analysis associates a phenotype with each amino acid at each position. Many amino acid residues at specific loci have been associated with human diseases and blood protein levels [3]. Published amino acid associations represent opportunities for experimental validation that could advance understanding of the disease-associated mechanisms related to HLA proteins.

Fine-mapping results can be interpreted in the context of the protein structures that are affected by the associated amino acid positions. We might have different hypotheses about the function of a mutation in the peptide binding groove than a mutation in the interior region of the protein.

To facilitate HLA fine-mapping, we developed *hlabud*, a free and open-source R package that downloads data from the IMGT/HLA database [4] and automatically creates amino acid (or nucleotide) position matrices that are ready for analysis (Figure 1). *hlabud* functions return simple lists, where each item in the list is a matrix or a data frame. This design makes it easy to integrate *hlabud* with any downstream R packages for data analysis or visualization.

2. EXAMPLES

2.1. Downloading data for a gene

Curated HLA genotype data is provided by the IMGT/HLA database at GitHub (github.com/ANHIG/IMGTHLA). In the example below, we use *hlabud* to download the sequence alignment data for *HLA-DRB1*, read it into R, and encode it as a one-hot matrix:

a <- hla alignments("DRB1")</pre>

With one line of code, hlabud will:

- Download data from the IMGT/HLA Github repository.
- Cache files in a local folder that supports multiple data releases.

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- Read the data into matrices and dataframes for downstream analysis.
- Create a one-hot encoding of the multiple sequence alignment data.

2.2. Computing a dosage matrix

Once we have obtained a list of genotypes for each individual (e.g. "DRB1*04:01, DRB1*05:01"), we can use *hlabud* to prepare data for fine-mapping regression analysis that will reveal which amino acid positions are associated with a phenotype in a sample of individuals. To calculate the number of copies of each amino acid at each position for each individual, we can run:

```
dosage(genotypes, a$onehot)
```

where genotypes is a vector of HLA-DRB1 genotypes and asonehot is a one-hot matrix representation of HLA-DRB1 alleles. The dosage matrix can then be used for omnibus regression [5] or fine-mapping (i.e. regression with each single position) (Figure 2A).

2.3. Visualizing alleles in two dimensions

Visualizing data in a two-dimensional embedding with algorithms like UMAP [6] can help to build intuition about the relationship between all objects in a dataset. UMAP accepts the one-hot matrix of HLA alleles as input, and the resulting embedding can be used to visualize the dataset for exploratory data analysis (Figure 2B).

2.4. Allele frequencies in human populations

hlabud provides direct access to the allele frequencies of HLA genes in the Allele Frequency Net Database (AFND) [7] (http://allelefrequences.net) (Figure 2C).

2.5. HLA divergence

Each HLA allele binds a specific set of peptides. So, an individual with two highly dissimilar alleles can bind a greater number of different peptides than a homozygous individual [8]. hlabud implements the Grantham divergence calculations [9] (based on the original Perl code) to estimate which individuals can bind a greater number of peptides (higher Grantham divergence):

```
my genos <- c("A*23:01:12,A*24:550", "A*25:12N,A*11:27", "A*24:381,A*33:85")
hla_divergence(my_genos, method = "grantham")
3.3333333
        0.4924242
                                    4.9015152
```

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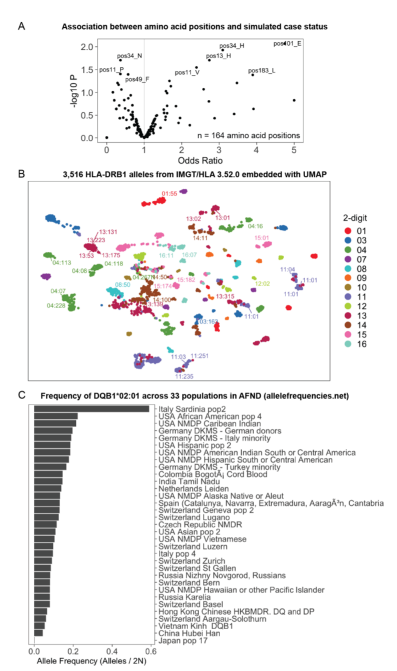


Figure 2: (**A**) Association between amino acid positions and simulated case-control status. The x-axis represents the odds ratio and the y-axis represents $-\log_{10}P$ from a logistic regression analysis in R. (**B**) 3,516 HLA-DRB1 alleles represented as dots in a two-dimensional embedding computed by UMAP from a one-hot encoding of amino acids. (**C**) Allele frequencies for HLA-DQB1*02:01 in the AFND.

3. Installation and documentation

The easiest way to install *hlabud* is to run this command in an R session:

remotes::install_github("slowkow/hlabud")

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The complete manual is available at https://slowkow.github.io/hlabud. hlabud has been tested on Linux/Unix. Mac OS (Darwin) and Windows.

4. Discussion

Our open-source R package hlabud gives users access to HLA data from two public databases, and implements HLA divergence calculation [9]. hlabud downloads HLA genotype data from the IMGT-HLA GitHub repository [10], caches it in a userconfigurable folder, and prepares the data for downstream analysis in R.

We provide tutorials for HLA divergence, fine-mapping association analysis with logistic regression, and embedding with UMAP. hlabud provides allele frequencies for all HLA genes, obtained from the Allele Frequency Net Database (AFND) [7].

5. ACKNOWLEDGMENTS

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6. COMPETING INTERESTS

No competing interest is declared.

7. AUTHOR CONTRIBUTIONS STATEMENT

K.S. wrote the software and the manuscript. A.C.V. reviewed the manuscript.

8. RELATED WORK

BIGDAWG is an R package that provides functions for chi-squared Hardy-Weinberg and case-control association tests of highly polymorphic genetic data like HLA genotypes [11]. HATK is set of Python scripts for processing and analyzing IMGT-HLA data [12].

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