# vignette

## library(hlacoloc)

## Introduction

HLA-coloc performs colocalization in two steps, as described in the manuscript. These are:

- 1. HLA gene causal signature: this uses SuSiE to assign a posterior inclusion probability (PIP) to each HLA allele.
- 2. HLA PIP Bayesian regression: this uses stanR to perform Bayesian regression of the PIPs in each genes separately.

# Required inputs

The following dataframes are necessary to run it (for each of the two phenotypes):

- 1. The HLA association summary statistics. This is a dataframe with the following columns:
- Name: the name of the allele. These must be given in IMGT-HLA format, but can be of any field resolution though (as long as it's the same for both phenotypes).
- z: the z-scores of the associations.
- beta: the beta of the associations (if z is not provided).
- se: the standard errors (if z is not provided).
- N: the sample sizes.
- 2. The LD matrix (can be either in dataframe or matrix format). This is a square matrix of R coefficient for the HLA alleles above. Importantly, the order of the alleles needs to be the same in the summary statistics dataframe and in the LD matrix here.

## Example

#### First, load the data:

The data is included in the package.

```
data("ebna", "ms", "r_ebna", "r_ms")
```

# Data snippet

Here's a snippet of the EBNA data:

```
head(ebna)
          Name
                        beta
                                   se
#> 1 G*01:01:01 0.014137500 0.015906 7247
#> 2 G*01:04:04  0.076847400  0.070867 7247
#> 3 G*01:01:02 -0.017568100 0.018972 7247
#> 4 G*01:01:03  0.000931893  0.033494 7247
      G*01:05N -0.293138000 0.078049 7247
#> 6 G*01:06:01 0.002110380 0.035066 7247
r ebna[1:5,1:5]
              G*01:01:01 G*01:04:04 G*01:01:02 G*01:01:03
                                                           G*01:05N
#> G*01:01:01
             1.000000 -0.1573650 -0.5404320 -0.2478420 -0.0925080
#> G*01:04:04 -0.157365 1.0000000 -0.0359779 -0.0317737 -0.0171784
#> G*01:01:02 -0.540432 -0.0359779 1.0000000 -0.1419840 -0.0672568
#> G*01:01:03 -0.247842 -0.0317737 -0.1419840 1.0000000 -0.0257557
#> G*01:05N
              -0.092508 -0.0171784 -0.0672568 -0.0257557 1.0000000
```

And here's a snippet of the multiple sclerosis (MS) data:

```
head(ms)
         Name
                   beta
#> 1 G*01:01:01 -0.082764 0.032331 429822
#> 2 G*01:04:04  0.059263  0.144366  429822
#> 3 G*01:01:02 -0.050757 0.043057 429822
#> 4 G*01:01:03  0.120691  0.061987  429822
      G*01:05N -0.112573 0.158904 429822
#> 6 G*01:06:01 0.155703 0.059848 429822
r_ms[1:5,1:5]
            G*01:01:01 G*01:04:04 G*01:01:02 G*01:01:03
                                                        G*01:05N
#> G*01:01:01 1.0000000 -0.15260200 -0.5331010 -0.2476440 -0.09820330
#> G*01:01:02 -0.5331010 -0.03782300 1.0000000 -0.1369080 -0.05690370
#> G*01:01:03 -0.2476440 -0.01675320 -0.1369080 1.0000000 -0.02644930
#> G*01:05N -0.0982033 -0.00766048 -0.0569037 -0.0264493 1.00000000
```

### Now we run HLA-coloc on EBNA and MS:

This performs the 2 steps described above in one command.

#### Results:

Results include the colocalization table and a plot (optional).

The colocalization table includes 5 columns - gene: the HLA gene. - susie\_coloc\_prob: the probability that SuSiE selects at least one shared HLA allele between the two traits at the given gene. - bayes\_pd: the probability that the SuSiE posterior inclusion probabilities (PIP) correlate for each

genes. - direction\_of\_correlation: a check to make sure that the correlation between the PIPs is positive. - hla\_colocalization\_probability: the HLA colocalization probability (only valid if direction\_of\_correlation is correct). This is the product of susie\_coloc\_prob and bayes\_pd, and is the final probability of HLA colocalization.

```
coloc_res[["hla_colocalization"]]
#> # A tibble: 9 x 5
     gene susie_coloc_prob bayes_pd direction_of_correlat~1 hla_colocalization_p~2
#>
     <chr>
                      <dbl>
                              <dbl> <chr>
                                                                              <dbl>
#> 1 DQB1
                 1.00
                               1
                                     Correct
                                                                         1.00
#> 2 DRB1
                 0.990
                              0.927 Correct
                                                                         0.917
#> 3 DPB1
                 0.0856
                              1.00 Correct
                                                                         0.0856
                 0.00349
                              0.996 Correct
                                                                         0.00348
#> 4 H
#> 5 DQA1
                 0.00129
                              0.733 Correct
                                                                         0.000945
#> 6 B
                 0.000803
                              0.764 Incorrect
                                                                         0.000614
#> 7 C
                 0.000192
                              0.672 Incorrect
                                                                         0.000129
#> 8 A
                 0.000152
                               0.733 Correct
                                                                         0.000111
#> 9 G
                 0.00000352
                               0.549 Correct
                                                                         0.00000193
#> # i abbreviated names: 1: direction_of_correlation,
#> # 2: hla_colocalization_probability
```

The plot has two columns. Left (a): the betas of the HLA allele summary statistics for the two traits. Right (b): the PIPs obtained from SuSiE, and the final probabilities of HLA colocalization.

```
coloc_res[["plot"]]
```

