Integrative multi-context Mendelian randomization method for identifying risk genes across human tissues

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Introduction

This vignette provides an introduction to the mintMR package. R package mintMR implements mintMR for integrative multi-context Mendelian randomization.

Install the development version of mintMR by use of the 'devtools' package. Note that mintMR depends on the 'Rcpp' and 'RcppArmadillo' package, which also requires appropriate setting of Rtools and Xcode for Windows and Mac OS/X, respectively. This package now depends on R (>= 3.5.0).

To install this package, run the following command in R

```
library(devtools)
install_github("ylustat/mintMR")
```

Load the package using the following command:

```
library("mintMR")
```

This vignette depends on R packages CCA and ggplot2, you can load these two packages using the following command:

```
suppressMessages(library("tidyverse"));
suppressMessages(library("CCA"));
library("ggplot2");
```

Fit mintMR for correlated SNPs using simulated data

In this section, we fit mintMR using simulated data (provided in the package).

```
load("/Users/yihaolu/Downloads/Research/mintMR/data/example_simulation_data.RData")
L <- 50
K1 <- K2 <- 5</pre>
```

The simulated data has 2 exposures, each with five contexts. It has 50 gene-CpG pairs. True effect status (0 indicates no effect, 1 indicates true non-zero effect). The mintMR requires the information for what contexts included for each exposure (the 'group' variable).

```
group <- list(exposure1 = 1:K1, exposure2 = 1:K2+K1)</pre>
```

In addition, mintMR takes IV-to-exposure effect (a list of $I_g \times \sum K_l$ matrices with length L), IV-to-outcome effect and corresponding standard errors. The LD matrices (corr_mat) and sample overlap (Lambda) are optional.

Default prior parameters used in the algorithm is

```
opts <- get_opts(L)
names(opts)</pre>
```

```
## [1] "a_gamma" "b_gamma"
## [ reached getOption("max.print") -- omitted 9 entries ]
```

It can be changed by specifying the parameters you want to use. For example:

```
opts <- get_opts(L, b_beta = rep(0.007,L))</pre>
```

MintMR can be run without specifying LD or sample overlap using following command.

When the LD information is available and formatted as a list of matrices, it can be accounted for:

When sample overlap information \widehat{C} is available as a $(1 + \sum_{l} K_{l}) \times (1 + \sum_{l} K_{l})$ matrix, it can be used in the function as

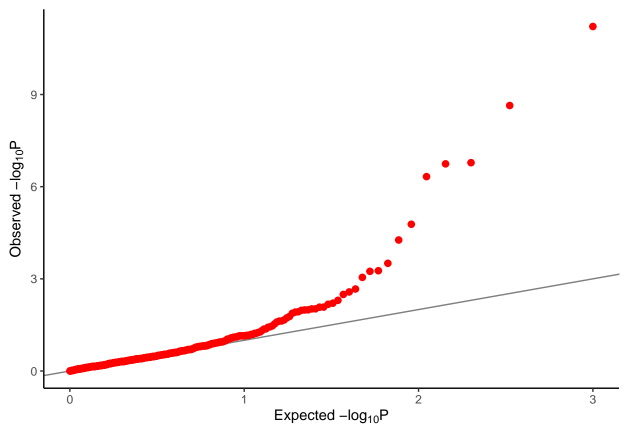
The output includes the effect estimates and p-values.

```
names(res)
```

```
## [1] "Estimate" "Pvalue"
```

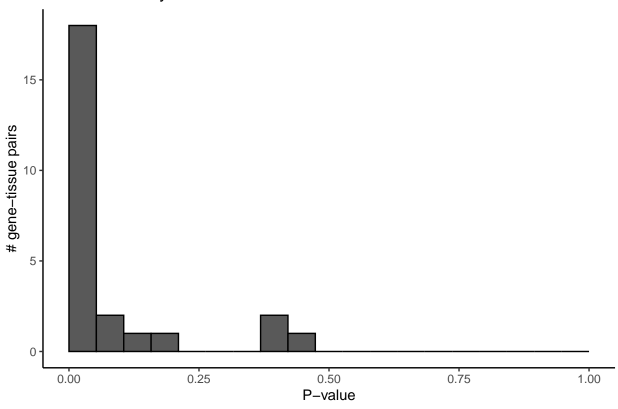
The p-value distribution for all gene-tissue pairs analyzed can be visualized as below.

```
p <- data.frame(mintMR = as.vector(res$Pvalue))
qqplot <- p %>%
    arrange(mintMR) %>%
    mutate(unif = ppoints(n())) %>%
    mutate(value = -log10(mintMR), unif = -log10(unif)) %>%
    ggplot(aes(y = value, x = unif))+
    geom_abline(intercept = 0, slope = 1, alpha = 0.5)+
    geom_point(size = 2, col = "red", alpha=1)+
    theme_classic()+
```



The p-value distribution for truly non-zero effects:

P-value for truly non-zero effect



The effect estimate for null effects:

```
estimate <- res$Estimate
estimate[dat$latent_status == 1] <- NA

estimate %>%
  reshape2::melt() %>%
  drop_na() %>%
  ggplot(aes(y = value))+
  theme_bw()+
  geom_boxplot(width=0.2)+
  geom_hline(yintercept = 0,col="red",linetype="dashed")
```

