# Package 'MRlap'

October 8, 2021

<b>Title</b> MRlap is an R-package to perform two-sample Mendelian Randomisation (MR) analyses using (potentially) overlapping samples
<b>Version</b> 0.0.1.0000
Description MR estimates can be subject to different types of biases due to the overlap between the exposure and outcome samples, the use of weak instruments and Winner's curse. Our approach simultaneously accounts and corrects for all these biases, using crosstrait LD-score regression (LDSC) to approximate the overlap. It requires only GWAS summary statistics. Estimating the corrected effect using our approach can be performed as a sensitivity analysis: if the corrected effect do not significantly differ from the observed effect, then IVW MR estimate can be safely used. However, when there is a significant difference, corrected effects should be preferred as they should be less biased, independently of the sample overlap.
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Encoding UTF-8
LazyData true
Imports magrittr,  tibble, dplyr, data.table, rlang, stringr, TwoSampleMR, GenomicSEM
Remotes MRCIEU/TwoSampleMR, GenomicSEM/GenomicSEM
RoxygenNote 6.1.1
R topics documented:
MRlap
Index

2 MRIap

MRlap	MRlap - main function	

## Description

Performs cross-trait LD score regression, IVW-MR analysis and provide a correction that simultaneously accounts for biases due to the overlap between the exposure and outcome samples, the use of weak instruments and Winner's curse.

## Usage

```
MRlap(exposure, exposure_name = NULL, K_exposure = NA,
   P_exposure = NA, outcome, outcome_name = NULL, K_outcome = NA,
   P_outcome = NA, ld, hm3, MR_threshold = 5e-08,
   MR_pruning_dist = 500, MR_pruning_LD = 0, MR_reverse = 0.001,
   save_logfiles = FALSE, verbose = TRUE)
```

### Arguments

exposure	The path to the file containing the GWAS summary statistics for the exposure, or a data.frame (character, or data.frame)	
exposure_name	The name of the exposure trait, default="exposure" (character)	
K_exposure	If case-control exposure, prevalence in the population, default=NA for continuous traits (numeric) $$	
P_exposure	If case-control exposure, prevalence in the sample, default=NA for continuous traits (numeric)	
outcome	The path to the file containing the GWAS summary statistics for the exposure, or a data.frame (character, or data.frame)	
outcome_name	The name of the outcome trait, default="outcome" (character)	
K_outcome	If case-control outcome, prevalence in the population, default=NA for continuous traits (numeric) $$	
P_outcome	If case-control outcome, prevalence in the sample, default=NA for continuous traits (numeric)	
ld	The path to the folder in which the LD scores used in the analysis are located. Expects LD scores formated as required by the original LD score regression software. (character)	
hm3	The path to a file of SNPs with alt, ref alleles and rsid used to allign alleles across traits (character)	
MR_threshold	The threshold used to select strong instruments for MR, should be lower than 1e-5, default=5e-8 (numeric)	
MR_pruning_dist		
	The distance used for pruning MR instruments (in Kb), should be between 10 and 50000, default=500 (numeric)	
MR_pruning_LD	The LD threshold (r2) used for pruning MR instruments, should be between 0 and 1 (if 0, distance-based pruning is used), default=0 (numeric)	
MR_reverse	The p-value used to exclude MR instruments that are more strongly associated with the outcome than with the exposure,default=1e-3 (numeric)	
save_logfiles	A logical indicating if log files from LDSC should be saved, default=FALSE	
verbose	A logical indicating if information on progress should be reported, default=TRUE	

SmallExposure\_Data 3

#### **Details**

exposure and outcome are required arguments. The input file / data.frame should contain the following columns (lower or upper case) :

SNPID (rs numbers) should be: rs, rsid, snp, snpid, rnpid

CHR (chromosome) should be : chr POS (position) should be : pos

ALT (effect allele) should be: a1, alt, alts REF (reference allele) should be: a2, a0, ref

Z (z-score) should be: Z, zscore N (sample size) should be: N

If Z is not present, it can be calculated from BETA and SE.

BETA should be: b, beta, beta1, or

SE should be: se, std

If (at least) one of the datasets is coming from a case-control GWAS:\* The Sample size column should correspond to the total sample size. The number of cases (NCASES) and the number of controls (NCONTROLS) can also be provided (instead of the total sample size). NCASES should

be: n\_cases, ncases, n\_case, ncase

NCONTROLS should be: n\_controls, ncontrols, n\_control, ncontrol

SmallExposure\_Data

Exposure

#### **Description**

Subset of the original dataset containing the estimated effect of SNPs on the exposure

### Usage

SmallExposure\_Data

#### **Format**

A data frame with 750,000 rows and 13 variables:

chr chromosome

rsid rsid of the SNP

pos position

ref reference allele for the SNP

alt effect allele for the SNP

af allele frequency

info imputation quality

beta estimated effect size for the SNP

se standard error of the estimated effect size for the SNP

z z-score for the SNP

minuslog10p -log10(p) for the SNP

**p** p-value for the SNP

N sample size

4 SmallOutcome\_Data

SmallOutcome\_Data

Outcome

### Description

Subset of the original dataset containing the estimated effect of SNPs on the outcome

### Usage

SmallOutcome\_Data

#### **Format**

A data frame with 750,000 rows and 13 variables:

chr chromosome

rsid rsid of the SNP

pos position

ref reference allele for the SNP

alt effect allele for the SNP

af allele frequency

info imputation quality

beta estimated effect size for the SNP

se standard error of the estimated effect size for the SNP

z z-score for the SNP

 $\label{eq:minuslog10p} \textbf{minuslog10p} \ \ \text{-log10(p)} \ for \ the \ SNP$ 

**p** p-value for the SNP

N sample size

# Index

```
*Topic datasets
SmallExposure_Data, 3
SmallOutcome_Data, 4

MRlap, 2

SmallExposure_Data, 3
SmallOutcome_Data, 4
```