

SOFTWARE TOOL ARTICLE

MendelianRandomization v0.9.0: updates to an R REVISED package for performing Mendelian randomization analyses using summarized data [version 2; peer review: 2 approved]

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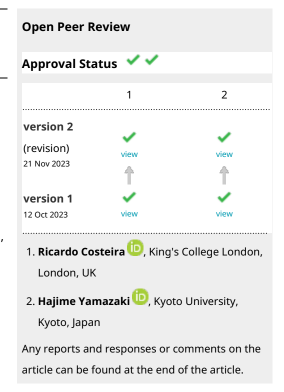
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Abstract

The MendelianRandomization package is a software package written for the R software environment that implements methods for Mendelian randomization based on summarized data. In this manuscript, we describe functions that have been added or edited in the package since version 0.5.0, when we last described the package and its contents. The main additions to the package since that time are: 1) new robust methods for performing Mendelian randomization, particularly in the cases of bias from weak instruments and/or winner's curse, and pleiotropic variants, 2) methods for performing Mendelian randomization with correlated variants using dimension reduction to summarize large numbers of highly correlated variants into a limited set of principal components, 3) functions for calculating first-stage F statistics, representing instrument strength, in both univariable and multivariable contexts, and with uncorrelated and correlated genetic variants. We also discuss some pragmatic issues relating to the use of correlated variants in Mendelian randomization.



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REVISED Amendments from Version 1

We have updated the manuscript in response to reviewer comments. In particular, we have expanded the introduction to provide more background information on the Mendelian randomization approach; we have clarified the source of the genetic association data; we have expanded and clarified the discussion about genetic correlation matrices and orientation; we have re-ordered the material, adding section headings on "Correlated variants" and "Variant correlation matrix and exposure correlation matrix"; we have expanded on how to obtain a genetic correlation matrix; and we have discussed specification of the exposure correlation matrix. All changes to the manuscript are minor in nature, and clarify the material in the initial submission rather than correcting or changing any interpretation of the original text.

Any further responses from the reviewers can be found at the end of the article

Introduction

Assessing causality in relationships between risk factors and outcomes is tricky and subject to many potential pitfalls. Mendelian randomization is an epidemiological technique that assesses such questions using genetic variants^{1,2}. Mendelian randomization uses genetic variants as instrumental variables to assess evidence for the causal effect of an exposure on an outcome.

An instrumental variable is a variable that behaves analogously to randomization in a randomized trial. It divides the population into groups with different distributions of the exposure, but the groups are otherwise indistinguishable (aside from any differences occurring due to downstream causal effects of the exposure)³. Formally, an instrumental variable should be associated with the exposure (relevance), not associated with the outcome via any confounding pathway (exchangeability), and does not affect the outcome directly, only potentially indirectly via an effect of the exposure (exclusion restriction)⁴. Generally speaking, genetic variants are plausible candidate instrumental variables, as they tend to have specific effects on particular biological mechanisms, they are uncorrelated with genetic variants that influence other biological mechanisms (due to Mendel's laws of inheritance), and they are fixed at conception (hence not subject to reverse causation or influence by environmental factors)⁵.

MendelianRandomization is a software package written for the R software environment⁶ that implements methods for Mendelian randomization based on summarized data⁷. By summarized data, we mean genetic associations with traits (beta-coefficients and standard errors) taken from regression analyses of a trait on a genetic variant⁸. Such associations are estimated in genome-wide association studies. They have been publicly reported for millions of variants by many large studies and consortia, and can be accessed through several different public portals^{9,10}.

This package has previously been introduced¹¹, and updates up to version 0.5.0 have been discussed¹². In this paper, we present updates to the package since version 0.5.0 up to the current version 0.9.0. A complete list of functions in the package is given in Table 1. The properties of the various methods are not discussed here in detail; we encourage users to read the relevant references for the specific methods or the guidelines paper for general advice on performing Mendelian randomization investigations¹³. We also encourage users to consult the documentation provided with the package, which describes all the options available for each method in greater detail. In this paper, we aim to provide an overview of recent updates to the package.

Table 1. Functions available in the MendelianRandomization package. Functions are divided into five categories: data entry functions, univariable estimation methods, multivariable estimation methods, data visualization functions, and functions that load data from PhenoScanner.

Function	Description	Status	Can include correlated variants?
Data entry function	ons		
mr_input	Data entry for univariable analysis		
mr_mvinput	Data entry for multivariable analysis		

Function	Description	Status	Can include correlated variants?
Univariable estim	ation methods		
mr_ivw	Inverse-variance weighted (IVW) method	†	✓
mr_median	Median method		
mr_egger	MR-Egger method		✓
mr_maxlik	Maximum likelihood method		✓
mr_mbe	Mode-based estimation method		
mr_hetpen	Heterogeneity penalized method		
mr_conmix	Contamination mixture method		
mr_lasso	Lasso method		
mr_cML	Constrained maximum likelihood method	*	
mr_divw	Debiased inverse-variance weighted method	*	
mr_pivw	Penalized inverse-variance weighted method	*	
mr_pcgmm	Principal component generalized method of moments method	*	✓
mr_allmethods	Runs several methods		
Multivariable esti	mation methods		
mr_mvivw	Multivariable IVW method	t	✓
mr_mvmedian	Multivariable median-based method		
mr_mvegger	Multivariable MR-Egger method		✓
mr_mvlasso	Multivariable lasso method		
mr_mvcML	Multivariable constrained maximum likelihood method	*	
mr_mvgmm	Multivariable generalized method of moments method	*	✓
mr_mvpcgmm	Multivariable principal component generalized method of moments method	*	✓
Data visualization	functions		
mr_plot	Scatter plot		
mr_forest	Forest plot		
mr_funnel	Funnel plot		
mr_loo	Leave-one-out plot		
Loading data from	n PhenoScanner		
extract.pheno.csv	Data entry from PhenoScanner .csv file (legacy)		
pheno_input			

^{* =} new since version 0.5.0, \dagger = updated since version 0.5.0, \checkmark = estimation method allows variants to be correlated (if not, then the method assumes variants are uncorrelated).

Methods

Implementation

Constrained maximum likelihood methods. The mr_cML and mr_mvcML functions perform the constrained maximum likelihood method, for the univariable Mendelian randomization¹⁴ (mr_cML) and multivariable Mendelian randomization¹⁵ (mr_mvcML) settings. These methods are robust to the violation of any of the three instrumental variable assumptions under mild assumptions. In a maximum likelihood framework, these methods constrain the number of invalid instruments with horizontal pleiotropy. The number of invalid instruments is asymptotically consistently selected by the Bayesian information criterion. To further account for the selection uncertainty with a finite sample, a data perturbation approach is employed.

The mr_cML function takes an *MRInput* object as input, created using the mr_input command. The syntax is:

```
mr_cML(mr_input(ldlc, ldlcse, chdlodds, chdloddsse),n=17723)
```

where ldlc and ldlcse are genetic associations with low-density lipoprotein (LDL) cholesterol and their standard errors for 28 uncorrelated genetic variants previously reported as associated with at least one of LDL-cholesterol, high-density lipoprotein (HDL) cholesterol, or triglycerides by Waterworth *et al.*¹⁶, and chdlodds and chdloddsse are genetic associations with coronary heart disease risk for the same variants. These data variables are provided with the package. n is the sample size for the genetic associations. In practice n could be the sample size either for the exposure or the outcome, the smaller value is recommended to get appropriately-sized confidence intervals; see reference for detailed discussions¹⁴.

The mr_mvcML function takes an *MRMVInput* object as input, created using the mr_mvinput command. The syntax is:

```
mr_mvcML(mr_mvinput(bx = cbind(ldlc, hdlc, trig),
    bxse = cbind(ldlcse, hdlcse, trigse),
    by = chdlodds, byse = chdloddsse), n = 17723)
```

where hdlc and hdlcse are genetic associations with HDL cholesterol and their standard errors, trig and trigse are genetic associations with triglycerides and their standard errors for the same 28 variants, and n is the sample size for the genetic associations with the exposures (or outcome if smaller). Again, these data variables are provided with the package.

The main options for these methods are DP = TRUE, which performs data perturbation and generally results in wider confidence intervals, but is recommended to provide appropriately-sized confidence intervals; MA = TRUE (for mr_cML), which performs model averaging, which again is recommended to provide appropriately-sized confidence intervals; and rho_mat (for mr_mvcML), which specifies the exposure correlation matrix between the summarized data for the exposures and outcome. If this is unspecified, then it is taken as the identity matrix, implying that the summarized data for the exposures and outcome are independent (typically because they were estimated in non-overlapping samples). Other options are provided to change the settings of the optimization functions used in the methods.

Debiased inverse-variance weighted method. The mr_divw function performs the debiased inverse-variance weighted method¹⁷. This method is an extension of the inverse-variance weighted (IVW) method that is more robust to weak instruments, with better bias and coverage properties.

The mr_divw function takes an MRInput object as input, created using the mr_input command. The syntax is:

```
mr_divw(mr_input(ldlc, ldlcse, chdlodds, chdloddsse))
```

The main options for this method are over.dispersion = TRUE, which allows for overdispersion in the variant-specific estimates (similar to a random-effects model for the IVW method), and diagnostics = FALSE, which provides a quantile—quantile plot of the variant-specific estimates, as a visual inspection for overdispersion and outliers.

Penalized inverse-variance weighted method. The mr_pivw function performs the penalized inverse-variance weighted method¹⁸. This method is an extension of the IVW method and the debiased inverse-variance weighted method, which handles weak instrument bias by a penalized log-likelihood function, and handles balanced horizontal pleiotropy by accounting for overdispersion in the variant-specific estimates.

The mr_pivw function takes an *MRInput* object as input, created using the mr_input command. The syntax is:

```
mr_pivw(mr_input(ldlc, ldlcse, chdlodds, chdloddsse))
```

The main options for this method are: lambda = 1, which is the penalty parameter. It plays a role in the bias-variance trade-off of the estimator. It is recommended to choose lambda = 1 to achieve the smallest bias and valid statistical inference; over.dispersion = TRUE, which allows for overdispersion in the variant-specific estimates; and delta = 0, which is a z-score threshold used for screening out weak instruments. delta should be greater than or equal to zero. When delta = 0, all variants provided will be used in the analysis. When delta > 0, the option sel.pval should be specified, which is the p-values of the genetic associations on the exposure. Then, the variants with sel.pval > 2*pnorm(delta,lower.tail = FALSE) will be removed from the analysis. The final option is Boot.Fieller = TRUE, which provides the p-value and the confidence interval of the causal effect calculated by the bootstrapping Fieller method.

Generalized method of moments method. The mr_mvgmm function performs the generalized method of moments (GMM) method¹⁹ for the multivariable Mendelian randomization setting. The key advantage of the GMM method is that estimates are more robust to weak instrument bias and/or measurement error in the exposures²⁰ compared with the standard IVW method, which is equivalent to a two-stage least squares approach with individual-level data⁷. Weak instrument bias is particularly important in the multivariable setting, as bias in the multivariable setting can be in any direction, particularly if instrument strength varies between the exposures.

Unlike the multivariable IVW method (mr_mvivw), the multivariable GMM method requires the sample sizes for genetic associations with the exposure, and the sample size for the genetic associations with the outcome. The method offers inferences that are robust to overdispersion in the variant-specific estimates (using the default option robust = TRUE).

If a genetic correlation matrix is not supplied in the mr_mvinput function, then the genetic variants will be assumed to be uncorrelated. There is also the option to use correlated variants if a genetic correlation matrix is supplied. If a genetic correlation matrix is supplied, the orientation of variants in the genetic correlation matrix (*i.e.* the assumed effect alleles) should be harmonized with the summary statistics used in the analysis as described in the mr_pcgmm section below.

The syntax for the mr_mvgmm function is:

```
mr_mvgmm(mr_mvinput(bx = cbind(ldlc, hdlc, trig), bxse = cbind(ldlcse, hdlcse, trigse), by = chdlodds, byse = chdloddsse), nx=rep(17723,3), ny=17723)
```

Principal component generalized method of moments methods. The mr_pcgmm and mr_mvpcgmm functions perform the principal component generalized method of moments (PC-GMM) method, for the univariable Mendelian randomization (mr_pcgmm) and multivariable Mendelian randomization (mr_mvpcgmm) settings²¹. This method is very similar to the GMM method, the key difference being that the GMM method uses individual variants as instruments rather than principal components.

These methods are designed for use when performing Mendelian randomization using genetic variants from a single gene region²². As an alternative to pruning and clumping approaches, which take a large number of variants from a gene region (potentially hundreds or thousands) and select a small number of uncorrelated (or weakly correlated) variants, the principal components approach performs dimension reduction on the full set of variant associations. The aim is to construct a small number of principal components which capture the information in the data, allowing the analysis to be performed using all available data, but avoiding numerical issues that would occur if highly-correlated genetic variants were included in the analysis. Previous investigations have shown that dimension reduction approaches can give additional precision compared with approaches using a small number of selected variants, and are less sensitive to variability arising from the variant selection process²³. Results from dimension reduction approaches are also less sensitive to misspecification of the variant correlation matrix, compared with other approaches using highly-correlated variants^{23,24}.

Compared with other dimension reduction approaches that have been proposed²⁵, the PC-GMM method has some important advantages: it uses the Continuously Updating Generalized Method of Moments method²⁶, which provides estimates that are more robust to weak instruments than some other instrumental variable methods²⁰; and it can allow for overdispersion in the variant-specific estimates (using the default option robust = TRUE), which is recommended to provide appropriately-sized confidence intervals and valid inferences.

The syntax for the univariable mr pcgmm function is:

```
mr_pcgmm(mr_input(bx = calcium, bxse = calciumse,
    by = fastgluc, byse = fastglucse, correlation = calc.rho),
    nx=6351, ny=133010)
```

where calcium, calciumse, fastgluc, fastglucse, and calcinho are genetic association data on six correlated variants from the *CASR* gene region provided with the package and their associations with serum calcium levels (calcium and associated standard errors calciumse) and fasting glucose levels (fastgluc and associated standard errors fastglucse), nx is the sample size for genetic associations with the exposure, and ny is the sample size for genetic associations with the outcome. These associations are:

```
> mr input(bx = calcium, bxse = calciumse,
> by = fastqluc, byse = fastqlucse, correlation = calc.rho)
   SNP exposure.beta exposure.se outcome.beta outcome.se
1 snp 1
        0.00625 0.00233 0.02805 0.0122
2 snp 2
            0.00590
                        0.00338
                                    0.00953
                                               0.0198
3 snp 3
            0.01822
                        0.00318
                                    0.03646
                                               0.0173
4 snp 4
            0.00598
                        0.00233
                                    0.01049
                                               0.0119
            0.00825
5 snp 5
                        0.00229
                                    0.02357
                                               0.0122
                        0.00352
            0.00651
                                    0.00204
                                               0.0179
6 snp 6
```

The syntax for the multivariable mr mvpcgmm function is:

```
mr_mvpcgmm(mr_mvinput(bx = cbind(ldlc, hdlc, trig),
    bxse = cbind(ldlcse, hdlcse, trigse),
    by = chdlodds, byse = chdloddsse,
    correlation = diag(length(ldlc))),
    nx=rep(17723,3), ny=17723)
```

We note that this example does not use genetic variants from a single gene region, and so does not represent a recommended use case for the method. It is provided to demonstrate the code syntax.

The default operation of the mr_pcgmm and $mr_mvpcgmm$ functions chooses the number of principal components to explain 99.9% of the variability in a weighted version of the variant correlation matrix. This can be varied, either by setting a different threshold (default thres = 0.999 corresponds to 99.9%) or by fixing the number of principal components using the r option (for example, r = 10 would select 10 principal components). As for the mr_mvcml function, the $mr_mvpcgmm$ function requires the exposure correlation matrix, set using the cor.x option (although for $mr_mvpcgmm$, correlations with the outcome are not needed). If not specified, this is set to the identity matrix, implying that the summarized data for the exposures are estimated in non-overlapping samples. If these associations were estimated in the same dataset and these correlations are not known, a sensitivity analysis may be worthwhile.

Correlated variants. Most methods in the MendelianRandomization package assume that all genetic variants are uncorrelated (see Table 1). The mr_ivw and mr_egger functions (and their multivariable counterparts) can account for correlations between variants, as can the mr_mvgmm and related methods (mr_pcgmm and mr_mvpcgmm) introduced above. While using partly correlated variants can provide additional precision compared with using uncorrelated (or minimally-correlated) variants, there are several cautions to the use of correlated variants in Mendelian randomization.

First, the estimate from the analysis with correlated variants should not be strikingly more precise than the result with the lead variant only (*i.e.* the variant having the lowest p-value in its association with the exposure) or with minimally-correlated variants. We would expect the correlated variants to explain slightly more variance in the exposure, and so we would expect the standard error to reduce and confidence intervals to narrow slightly when using correlated variants. However, if the standard error when using correlated variants is much

smaller than when using the lead variant only (say, it is two or three times smaller), then inputs should be checked carefully, as the increase in precision may represent a convergence issue. Reporting a sensitivity analysis using fewer variants or principal components is recommended.

Second, many software tools report the squared-correlation matrix (the r^2 matrix) between variants, not the (signed) correlation matrix. Mendelian randomization requires the correlation matrix, which provides correlations between variants.

Third, the signs of the correlations (positive or negative) must be correctly specified. If the correlation matrix is calculated using the same effect and non-effect alleles as the summarized data, then the correlations should have the correct signs²⁷. If not, then the correlation matrix should be harmonized to the same effect alleles by flipping signs in the relevant rows and columns. Failure to harmonize the genetic correlation matrix can result in erroneous estimates. This can be implemented in R by adapting the following example code:

```
flip = c(+1, +1, -1, +1, -1, +1)
calc.rho.signed = calc.rho*flip%o%flip
```

where flip is +1 for variants for which the alleles are correctly aligned, and -1 for variants for which the alleles are incorrectly aligned, and the elements of flip correspond to the genetic variants in order.

In this example, before orientation, the variant correlation matrix is:

```
> calc.rho

[,1] [,2] [,3] [,4] [,5] [,6]

[1,] 1.000 0.070 0.094 -0.172 0.079 -0.104

[2,] 0.070 1.000 -0.050 0.081 -0.080 -0.014

[3,] 0.094 -0.050 1.000 -0.306 0.349 -0.134

[4,] -0.172 0.081 -0.306 1.000 -0.129 0.446

[5,] 0.079 -0.080 0.349 -0.129 1.000 -0.289

[6,] -0.104 -0.014 -0.134 0.446 -0.289 1.000
```

After orientation, the variant correlation matrix is:

```
> calc.rho.signed

[,1] [,2] [,3] [,4] [,5] [,6]

[1,] 1.000 0.070 -0.094 -0.172 -0.079 -0.104

[2,] 0.070 1.000 0.050 0.081 0.080 -0.014

[3,] -0.094 0.050 1.000 0.306 0.349 0.134

[4,] -0.172 0.081 0.306 1.000 0.129 0.446

[5,] -0.079 0.080 0.349 0.129 1.000 0.289

[6,] -0.104 -0.014 0.134 0.446 0.289 1.000
```

As the third and fifth elements of flip are -1 (and all others are +1), all entries in the third row or column, and the fifth row or column, have flipped in sign (from positive to negative, or from negative to positive). For example, the third entry in the first row was +0.094, but is now -0.094. The element in the third row and fifth column (+0.349) is flipped in sign twice, and so its sign ends up unchanged; similarly for the element in the third row and third column, the fifth row and third column, and so on.

Fourth, the correlation matrix should be obtained from the same population (or same ancestry group) as the original data, to avoid mismatches between the correlation matrix and the summarized data. If the exposure and outcome data are from different ancestry groups, then correlated variant analyses should not be attempted. If the exposure and outcome data are from the same ancestry group, then correlations should be taken from the outcome data by preference, but either (or use of a suitable reference population) is acceptable.

Fifth, analysts should consider pruning out extremely highly correlated variants, as these are likely to contribute little to the analysis, even when using the mr pcgmm and mr mvpcgmm methods. A suggestion when using

these methods is to take pairs of variants that are correlated at $r^2 > 0.95$, and remove one of the pair at random until no such pairs remain. This can be implemented using the code below for a correlation matrix rho:

```
set.seed(496)
                       # for reproducibility
thres = sqrt(0.95) # threshold set to r2>0.95
omit = NULL
                       # set up list of variants to be omitted
rho.upper = rho
                       # correlation matrix
rho.upper[lower.tri(rho, diag=TRUE)] <- 0</pre>
                       # only consider upper triangle of correlations
j=1
                       # set counter to 1
while (max(abs(rho.upper), na.rm=TRUE)> thres) {
 omit[j] = ifelse(rbinom(1, 1, 0.5) == 1,
   which.max(apply(abs(rho.upper), 1, max, na.rm=TRUE)),
   which.max(apply(abs(rho.upper), 2, max, na.rm=TRUE)))
                       # find the highest correlation value
                       # select either the row or column at random
                       # add this to the list of omitted variants
 {\tt rho.upper[omit[j],]} <-0 ~\# {\tt set} {\tt the} {\tt correlations} {\tt in} {\tt this} {\tt row} {\tt to} {\tt zero}
 rho.upper[,omit[j]] <- 0 # set the correlations in this column to zero</pre>
                            # (to avoid selecting the same variant again)
 j = j + 1
                       # increment the counter
 } # stop when no more pairwise correlations exceed threshold
```

For other methods that allow for correlated variants, either pruning at a stricter threshold (say, $r^2 < 0.3$) or applying another approach for variant selection (such as fine-mapping or conditional modelling) is recommended to avoid high levels of variant multicollinearity.

Following these steps should ensure that the correlation matrix is relevant to the data under analysis, is correctly harmonized, and does not include pairs of variants with very highly correlations.

Variant correlation matrix and exposure correlation matrix. We note the distinction between the variant correlation matrix which represents correlations between variant association estimates occurring due to linkage disequilibrium, and the exposure correlation matrix which represents correlations between variant association estimates occurring due to correlations in exposure measurements, which arise if these estimates are obtained in the same individuals. If the analyst has access to individual-level data on allele counts, the variant correlation matrix can be calculated directly using the correlation (cor) function in R applied to the matrix of allele counts. The variant correlation matrix can be obtained from a reference population; for example, correlations for European ancestry individuals from UK Biobank are available here, and correlations for other ancestry groups (although for much smaller sample sizes) are available at https://ldlink.nci.nih.gov/?tab=ldmatrix (although note these are squared correlations, so not suitable for direct use in the MendelianRandomization package functions).

The exposure correlation matrix can only be estimated from individual-level data, and hence a sensitivity analysis for its value is suggested if it is unknown. However, investigations have indicated that estimates are generally insensitive to the correct specification of the exposure correlation matrix, and so the default value of the identity matrix is likely to be a reasonable choice. A sensitivity analysis varying this matrix is recommended, to assess whether findings change for different values of this matrix.

Operation. The R software environment (RRID:SCR_001905) runs on a wide variety of UNIX platforms, Windows, and MacOS, and requires minimal computer resources (256 kilobytes of RAM is recommended). The package requires R version 3.3.0 or higher. As the package now uses C++ code, an up-to-date version of Rtools should be installed to successfully install the package. This current work used R version 4.3.1.

Use case

Instrument strength

A further update to the MendelianRandomization package is the functionality to report F statistics in the mr_ivw and mr_ivw commands. The first-stage F statistic is a measure of the variance in the exposure

explained by the genetic variants in the exposure dataset²⁸. It is roughly equal to the R² statistic (the proportion of variance explained) multiplied by the sample size and divided by the number of instruments. Larger F statistics correspond to stronger instruments. The F statistic cannot be determined exactly based on summary statistics, but the simple formula implemented in the package provides a good approximation. While the mythical threshold of 10 for F statistics²⁹ is an oversimplification, this value does provide a reasonable yard-stick for judging the degree of potential bias due to weak instruments (although we would strongly caution against basing an analysis plan on measured values of the F statistic in the dataset under analysis²⁸). We note that although weak instrument bias is typically in the direction of the null for two-sample univariable Mendelian randomization³⁰, this is not necessarily the case for two-sample multivariable Mendelian randomization³¹.

The default implementation of the mr_ivw and mr_mvivw functions assumes that genetic variants are uncorrelated. Correlations can be specified in the mr input or mr mvinput command:

```
mr_ivw(mr_input(calcium, calciumse,
    fastgluc, fastglucse, corr=calc.rho))
```

If two uncorrelated variants both explain 3% of the variance in the exposure, then together they explain 6% of the variance in the exposure. If two correlated variants both explain 3% of the variance in the exposure, then the variance explained by both variants could be as low as 3% (if the variants are perfectly correlated) or as high as 6% (if the correlation is negligible), or a value between these two, depending on the magnitude of correlation. This correlation similarly affects estimates of the F statistic. When the variant correlation matrix is specified, the mr_ivw function accounts for these correlations in the calculation of the F statistic using a formula that involves the inverse of the Cholesky transform of the correlation matrix³².

For multivariable Mendelian randomization, the key measure of instrument strength is not the univariable F statistic for each exposure separately, but the conditional F statistic, representing the independent proportion of variance explained in each exposure, after accounting for associations with the other exposures^{33,34}. This is because if the genetic associations with one exposure were strong, but were near-perfectly correlated with the genetic associations with another exposure, then the multivariable model would not be able to differentiate between the effects of the two exposures.

The mr_mvivw function calculates estimates of the conditional F statistics based on summarized data. In order to identify and reliably estimate multivariable exposure effects through Mendelian randomization, we require that the genetic predictors of the exposures are not collinear. Hence, conditional F-tests assess whether the genetic predictors of any one exposure can be expressed as a linear combination of genetic predictors of other exposures in the model. Low values of conditional F-test statistics for a given exposure suggest that the exposure may not be identified, and hence estimates may suffer from weak instrument bias. This is generally a bigger concern for two-sample multivariable, rather than two-sample univariable, Mendelian randomization analyses since weak instrument biases in estimation are not necessarily toward the null of no causal effect, and hence inferences can suffer from inflated type I error rates (false positive findings). This calculation of conditional F-statistics involves the sample size for the genetic associations with the exposures, and conditional F statistics are only calculated if these sample sizes are provided. If a single value is provided for the sample size, then it is assumed that this sample size holds for all exposures. For example, the code:

```
mr_mvivw(mr_mvinput(bx = cbind(ldlc, hdlc, trig),
    bxse = cbind(ldlcse, hdlcse, trigse),
    by = chdlodds, byse = chdloddsse), nx = 17723)
```

gives output:

```
Number of Variants: 28

Exposure Estimate Std Error 95% CI p-value Cond F-stat exposure_1 1.925 0.439 1.064, 2.786 0.000 20.3 exposure_2 -0.590 0.555 -1.677, 0.498 0.288 12.9 exposure_3 0.723 0.230 0.272, 1.174 0.002 13.3
```

By comparison, the code:

```
mr_ivw(mr_input(ldlc, ldlcse, chdlodds, chdloddsse))
```

gives output including:

```
F statistic = 28.0.
```

We see that the univariable F statistic for LDL-cholesterol is 28.0, but the conditional F statistic is 20.3. Similarly, the univariable F statistic for HDL-cholesterol is 27.6, but the conditional F statistic is 12.9, and for triglycerides, the univariable F statistic is 41.6, but the conditional F statistic is 13.3. The instrument strength for LDL-cholesterol is similar in the multivariable analysis compared to the univariable analyses, but for HDL-cholesterol and triglycerides the instrument strength is weakened considerably.

F statistics and conditional F statistics are also provided by the mr_pcgmm and mr_mvpcgmm functions, as these may help the user to judge how varying the number of principal components selected affects instrument strength.

Conclusions

In summary, the MendelianRandomization package has added a number of features since the 0.5.0 version: to implement additional estimation methods, to implement methods for highly correlated variants, and to report F statistics.

We conclude by again repeating the warning that we stated at the end of the manuscript accompanying the initial package release¹¹: while this software simplifies the operational aspects of a Mendelian randomization, the truly difficult parts of an analysis are choosing sensible risk factors and outcomes, selecting genetic variants that are plausible instrumental variables, performing a reasonable range of analyses, and interpreting the results with care and caution¹³. These aspects of an analysis cannot be automated³⁵.

Data availability

Zenodo. MendelianRandomization package version 0.9.0. https://doi.org/10.5281/zenodo.8305056³⁶.

This project contains the following underlying data:

- Folder/Files containing input data used in the "use case section".

Software availability

Software available from: https://cran.r-project.org/web/packages/MendelianRandomization/index.html.

Archived source code at time of publication: http://doi.org/10.5281/zenodo.8305056³⁶

License: AGPL-3.0-only

Author contributions

Conceptualization: SB; Methodology: AP, TY, HX, ZL, SX, SB; Software: AP, TY, HX, ZL, SX, SB; Supervision: SB; Writing – Original Draft Preparation: SB; Writing – Review & Editing: all authors.

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Current Peer Review Status:





Version 2

Reviewer Report 30 November 2023

https://doi.org/10.21956/wellcomeopenres.22678.r70304

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Hajime Yamazaki 🗓



Section of Clinical Epidemiology, Department of Community Medicine, Graduate School of Medicine, Kyoto University, Kyoto, Japan

No further comments.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Clinical epidemiology, Gastroenterology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 23 November 2023

https://doi.org/10.21956/wellcomeopenres.22678.r70305

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Ricardo Costeira 🕛



Department of Twin Research and Genetic Epidemiology, King's College London, London, England, UK

No further comments.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Genomics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 23 Nov 2023

Stephen Burgess

Thank you Ricardo for the fast and positive response, and for your previous comments on this article, which have helped improve the work.

Competing Interests: No competing interests were disclosed.



Reviewer Report 30 October 2023

https://doi.org/10.21956/wellcomeopenres.22141.r68639

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Hajime Yamazaki 🗓



Section of Clinical Epidemiology, Department of Community Medicine, Graduate School of Medicine, Kyoto University, Kyoto, Japan

This article describes the updated functions of the MendelianRandomization package, which is widely used to conduct Mendelian randomization analyses. I believe that the updated functions and their detailed descriptions in this article will be immensely beneficial for researchers.

To enhance the readability and comprehensibility of the article, I propose the following suggestions:

- 1. At the beginning of the Methods section, the authors introduce 'three instrumental variable assumptions.' It would be prudent to both explicate these assumptions in the Introduction section and explain why genetic variants can serve as a proxy for random assignment.
- 2. This article utilizes data pertaining to LDL cholesterol, HDL cholesterol, triglycerides, and coronary heart disease risk as an illustrative example. It would be beneficial to clearly articulate the research question associated with this example, incorporate a graphical representation to delineate the relationship between genetic variants and these variables, and provide a brief overview of the selection process of these genetic variants within a multivariable Mendelian randomization context.
- 3. It would be beneficial to include a small portion of the example dataset, comprising the correlation matrix, within the article along with a reference to its availability in the

MendelianRandomization package.

- 4. The authors draw a distinction between the variant correlation matrix and the exposure correlation matrix. A more detailed explanation on how to estimate each of these, as well as guidance on which correlation matrix is pertinent for use with the MendelianRandomization package, would be beneficial.
- 5. The "Principal Component Generalized Method of Moments" section is extensive and could be made more readable by dividing it into two distinct subsections, each with its own subheading.

Is the rationale for developing the new software tool clearly explained?

Yes

Is the description of the software tool technically sound?

Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?

Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?

Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Clinical epidemiology, Gastroenterology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 12 Nov 2023

Stephen Burgess

Response to reviewers for "MendelianRandomization v0.9.0: updates to an R package for performing Mendelian randomization analyses using summarized data"

We would like to express thanks to the reviewers for their time and comments. Replies to points are indicated by angle brackets, and changes to the paper as a result of these comments are clearly indicated. We have numbered the comments for reference as A0, A1, A2, ... for the first reviewer; and B0, B1, B2, ... for the second reviewer.

Reviewer 2: Hajime Yamazaki, Section of Clinical Epidemiology, Department of Community Medicine, Graduate School of Medicine, Kyoto University, Kyoto, Japan

- B0. This article describes the updated functions of the MendelianRandomization package, which is widely used to conduct Mendelian randomization analyses. I believe that the updated functions and their detailed descriptions in this article will be immensely beneficial for researchers.
- > We thank the reviewer for their kind comments and positive view of their manuscript. <

To enhance the readability and comprehensibility of the article, I propose the following suggestions:

- B1. At the beginning of the Methods section, the authors introduce 'three instrumental variable assumptions.' It would be prudent to both explicate these assumptions in the Introduction section and explain why genetic variants can serve as a proxy for random assignment. > We have added a brief description of the three classical instrumental variable assumptions required to identify a causal effect, as well as a brief motivation as to why genetic variants are often suitable candidate instrumental variables. <
- B2. This article utilizes data pertaining to LDL cholesterol, HDL cholesterol, triglycerides, and coronary heart disease risk as an illustrative example. It would be beneficial to clearly articulate the research question associated with this example, incorporate a graphical representation to delineate the relationship between genetic variants and these variables, and provide a brief overview of the selection process of these genetic variants within a multivariable Mendelian randomization context.
- > We have added a brief description of how these genetic variants were chosen. While we appreciate the reviewer's point, we would not want to give the impression that the data included in the Mendelian randomization package are anything other than illustrative. They are not the ideal data to optimally address any research question. <
- B3. It would be beneficial to include a small portion of the example dataset, comprising the correlation matrix, within the article along with a reference to its availability in the MendelianRandomization package.
- > We have added the calcium and fasting glucose data to the manuscript, as well as their correlation matrix (see also point A2). <
- B4. The authors draw a distinction between the variant correlation matrix and the exposure correlation matrix. A more detailed explanation on how to estimate each of these, as well as guidance on which correlation matrix is pertinent for use with the MendelianRandomization package, would be beneficial. > There are various tools to estimate the variant correlation matrix. If you have access to individual-level genetic data (that is, allele counts for each SNP), you can calculate the variant correlation matrix using the correlation (*cor*) function in R. Alternatively, some pre-computed correlation matrices are available for download, for

example using the LDlink webtool (https://ldlink.nci.nih.gov/?tab=Idmatrix) or the Id extract function in the TwoSampleMR package. Variant correlation matrices for UK Biobank participants of European ancestries are available at

https://aws.amazon.com/marketplace/pp/prodview-4bhcvjnh4b4cs#resources. > We have listed some of these sources in the manuscript. > Exposure correlation estimates are required by some methods when genetic associations with any two exposures are measured from the same sample. Exposure correlation estimates may be more difficult to obtain than variant correlation estimates, but they could be estimated using individual-level data on the exposures from the same sample or an external sample used to measure genetic variant-exposure associations. In general, we have found estimation and inferences to be quite insensitive to mis-specified exposure correlations, and in practice we would recommend performing a sensitivity analysis that varies the inputted exposure correlations to assess sensitivity of findings to the specification of this matrix.

> We have expanded the discussion about exposure correlation matrices in the manuscript. <

B5. The "Principal Component Generalized Method of Moments" section is extensive and could be made more readable by dividing it into two distinct subsections, each with its own subheading.

> We have added separate sections marked "Correlated variants" and "Variant correlation matrix and exposure correlation matrix" that includes much of the text previously included in the "Principal Component Generalized Method of Moments" section, and hence breaking up this previously over-long section. <

Competing Interests: I am the corresponding author of the manuscript. I have no relevant conflicts to declare.

Reviewer Report 30 October 2023

https://doi.org/10.21956/wellcomeopenres.22141.r68641

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Ricardo Costeira 🗓



Department of Twin Research and Genetic Epidemiology, King's College London, London, England,

In this manuscript, the authors describe updates to the MendelianRandomization package implemented in R. In the package's latest version (v0.9.0), users can find functions to perform newer robust methods of MR (such as debiased and penalised IVW MR), consider variant correlation, and obtain F-statistics to estimate the strength of instrument variables.

The authors provide good rationale for when and how to implement the newer MR methods. Examples of code are abundant throughout the manuscript, and the package's functions are explained in detail. The authors set guidelines for implementing genetic correlation in MR and highlight potential caveats of the analysis. Altogether, the manuscript is useful to understand the newer functions available in MendelianRandomization v0.9.0 and has valuable insights into the practical considerations of applying more complex MR methods. Interpretation of newer output (the F-statistic) is included.

Minor comments:

Under the section of "Principal component generalized method of moments methods":

- "where calcium, calciumse, fastgluc, fastglucse, and calc.rho are data on six correlated variants from the CASR gene region provided with the package" – please reword to clarify the meaning of the calcium, calciumse, fastgluc and fastglucse variables.
- Is "flip = c(+1, +1, -1, +1, -1, +1)" length = 6 because "calc.rho" is a matrix of 6 variants? Please show "calc.rho" before and after applying the function "calc.rho*flip%o%flip". This can help understand the structure and signs of "calc.rho" going into the MR analysis.
- Regarding the correlation matrix coming from linkage disequilibrium data or exposure correlation: "However, our investigations have indicated that estimates are generally insensitive to the correct specification of this matrix." – Please expand on this comment.
 Under the section "Instrument Strength":
 - "We note that although weak instrument bias is typically in the direction of the null for two-sample univariable Mendelian randomization, this is not necessarily the case for two-sample multivariable Mendelian randomization." and "This is generally a bigger concern for two-sample multivariable, rather than two-sample univariable, Mendelian randomization analyses since weak instrument biases in estimation are not necessarily toward the null of no causal effect, and hence inferences can suffer from inflated type I error rates." Please explain why this is the case. Additionally, please comment on how weak instrument bias can affect other types of MR.

Is the rationale for developing the new software tool clearly explained?

Yes

Is the description of the software tool technically sound?

Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?

Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?

Yes

Are the conclusions about the tool and its performance adequately supported by the

findings presented in the article?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Genomics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 12 Nov 2023

Stephen Burgess

We would like to express thanks to the reviewers for their time and comments. Replies to points are indicated by angle brackets, and changes to the paper as a result of these comments are clearly indicated. We have numbered the comments for reference as A0, A1, A2, ... for the first reviewer; and B0, B1, B2, ... for the second reviewer.

Reviewer 1: Ricardo Costeira, Department of Twin Research and Genetic Epidemiology, King's College London, London, England, UK

A0. In this manuscript, the authors describe updates to the MendelianRandomization package implemented in R. In the package's latest version (v0.9.0), users can find functions to perform newer robust methods of MR (such as debiased and penalised IVW MR), consider variant correlation, and obtain F-statistics to estimate the strength of instrument variables. The authors provide good rationale for when and how to implement the newer MR methods. Examples of code are abundant throughout the manuscript, and the package's functions are explained in detail. The authors set guidelines for implementing genetic correlation in MR and highlight potential caveats of the analysis. Altogether, the manuscript is useful to understand the newer functions available in MendelianRandomization v0.9.0 and has valuable insights into the practical considerations of applying more complex MR methods. Interpretation of newer output (the F-statistic) is included.

> We thank the reviewer for their kind comments and positive view of this manuscript. <

Minor comments: Under the section of "Principal component generalized method of moments methods":

- A1. "where calcium, calciumse, fastgluc, fastglucse, and calc.rho are data on six correlated variants from the CASR gene region provided with the package" please reword to clarify the meaning of the calcium, calciumse, fastgluc and fastglucse variables.
- > We have clarified that *calcium* represents the genetic associations with serum calcium levels, and *fastgluc* the genetic associations with fasting glucose levels (*calciumse* and *fastglucse* are the standard errors respectively). <

- A2. Is "flip = c(+1, +1, -1, +1, -1, +1)" length = 6 because "calc.rho" is a matrix of 6 variants? Please show "calc.rho" before and after applying the function "calc.rho*flip%o%flip". This can help understand the structure and signs of "calc.rho" going into the MR analysis.
- > We have clarified that the entries in the *flip* vector correspond to the six genetic variants in turn. We have added to the manuscript a representation of the correlation matrix before and after orientation. <
- A3. Regarding the correlation matrix coming from linkage disequilibrium data or exposure correlation: "However, our investigations have indicated that estimates are generally insensitive to the correct specification of this matrix." Please expand on this comment.
- > We have clarified that this sentence refers to the exposure correlation matrix. <

Under the section "Instrument Strength":

- A4. "We note that although weak instrument bias is typically in the direction of the null for two-sample univariable Mendelian randomization, this is not necessarily the case for two-sample multivariable Mendelian randomization." and "This is generally a bigger concern for two-sample multivariable, rather than two-sample univariable, Mendelian randomization analyses since weak instrument biases in estimation are not necessarily toward the null of no causal effect, and hence inferences can suffer from inflated type I error rates." Please explain why this is the case. Additionally, please comment on how weak instrument bias can affect other types of MR.
- > Weak instrument bias is analogous to classical measurement error bias in a regression model. With a single regressor, error in the regressor leads to underestimation of the regression coefficient; this is known as regression dilution bias. With multiple regressors, bias due to measurement error can be in any direction. For example, suppose we have two regressors that are highly correlated, one of which is a causal risk factor for the outcome, and the other is not. Further, one is measured with substantial error, whereas the other is measured with no error. In a regression model, the coefficient for the regressor measured with error tends towards zero regardless of which is truly the causal risk factor. This could lead the coefficient for the precisely measured exposure being either over- or underestimated, depending on its true value and the true value for the coefficient of the imprecisely measured regressor. In a prediction model, this may be of little consequence, as the predicted values of the outcome may be insensitive to the exposure coefficients which is large, which is close to zero. However, if the model coefficients have a causal interpretation, then the coefficients matter, as the regressor with the non-zero coefficient is identified as the causal factor.
- > Generally speaking, bias towards the null in Mendelian randomization is of lesser consequence, as the primary goal of Mendelian randomization is not estimation. Bias towards the null may reduce power to detect a true causal effect, but it will not incorrectly suggest a false causal effect (which is regarded in the Neyman—Pearson hypothesis framework as a more serious error).

> As this explanation is fairly involved, we do not think it is appropriate to include in this manuscript. However, given that this journal performs open peer review, it can be found by the interested reader. <

Competing Interests: I am the corresponding author of the manuscript. I have no relevant conflicts to declare.