R code and results for the lung function example

# BetaYG = vector of gene-outcome associations
# BetaXG = vector of gene-exposure associations
# seBetaYG = vector of gene-outcome association standard errors
# seBetaXG = vector of gene-exposure association standard errors
# '>' denotes the start of each block of R code

IVW approach (with MAF corrected weights)

> IVWfit = summary(lm(BetaYG ~ -1+BetaXG,weights=1/seBetaYG^2))
> IVWfit$coef  # Point estimate: note standard error may be incorrect

Coefficients:
      Estimate Std. Error t value Pr(>|t|)
BetaXG  0.58612   0.04387   13.36  <2e-16 ***

DF = length(BetaYG)-1
> IVWBeta = IVWfit$coef[1,1]
> SE = IVWfit$coef[1,2]/min(1,IVWfit$sigma)
> IVW_P = 2*(1-pt(abs(IVWBeta/SE),DF))
> IVW_CI = IVWBeta + c(-1,1)*qt(df=DF, 0.975)*SE

IVWResults = (point estimate, corrected standard error, 95% Confidence interval, t-statistic, p-value)

> IVWResults = c(IVWBeta,SE,IVW_CI,IVWBeta/SE,IVW_P)

MR-Egger regression (with MAF corrected weights)

> BYG = BetaYG*sign(BetaXG)  # Pre-processing steps to ensure all
> BXG = abs(BetaXG)  # gene--exposure estimates are positive
> MREggerFit = summary(lm(BYG ~ BXG,weights=1/seBetaYG^2))
> MREggerFit$coef # Point estimate: note standard errors may be incorrect

          Estimate Std. Error   t value Pr(>|t|)
(Intercept) -0.0008820533  0.002748935 -0.3208709  7.486846e-01
BXG          0.6042841068  0.071675342  8.4308507  1.137390e-14

# Inference with corrected standard errors

> MREggerBeta0  = MREggerFit$coef[1,1]
> MREggerBeta1  = MREggerFit$coef[2,1]
> SE0           = MREggerFit$coef[1,2]/min(1, MREggerFit$sigma)
> SE1           = MREggerFit$coef[2,2]/min(1, MREggerFit$sigma)
> DF            = length(BetaYG) - 2
> MRBeta0_p     = 2*(1-pt(abs(MREggerBeta0/SE0),DF))
> MRBeta1_p     = 2*(1-pt(abs(MREggerBeta1/SE1),DF))
> MRBeta0_CI    = MREggerBeta0 + c(-1,1)*qt(df=DF, 0.975)*SE0
> MRBeta1_CI    = MREggerBeta1 + c(-1,1)*qt(df=DF, 0.975)*SE1

# MREggerResults = (point estimate, corrected standard error, # 95% Confidence interval, t-statistic, p-value) for # intercept (row 1) and slope (row 2).

> MREggerResults     = matrix(nrow = 2,ncol = 6)
> MREggerResults[1,] = c(MREggerBeta0,SE0,MRBeta0_CI,MRBeta0_p)
> MREggerResults[2,] = c(MREggerBeta1,SE1,MRBeta1_CI,MRBeta1_p)

# Bootstrap method to obtain confidence interval # and standard error for slope parameter in # MR-Egger regression (with MAF corrected weights)

> boot = NULL; straps = 10000
> for (i in 1:straps) {
    BYG_boot = rnorm(length(BYG), mean=BYG, sd=seBetaYG)
    BXG_boot = rnorm(length(BXG), mean=BXG, sd=seBetaXG)
    BYG_boot = BYG_boot*sign(BXG_boot)
    BXG_boot = abs(BXG_boot)
    boot[i] = summary(lm(BYG_boot~BXG_boot,weights=seBetaYG^-2))$coef[2,1]
}
> boot_upper = sort(boot)[9751]
> boot_lower = sort(boot)[250]
> boot_se    = sd(boot)
# MREggerBoot = c(point estimate, standard error, 95% confidence interval)

> MREggerBoot = c(MREggerBeta1, boot_se, boot_lower, boot_upper)

Under a fixed-effect model, which is correct when all of the variants included in the analysis are valid IVs (i.e. no pleiotropic effects) and each variant identifies the same causal effect, the standard error automatically reported for the inverse-variance weighted method is incorrect. This is because the residual standard error in a weighted regression analysis should be unity [Thompson and Sharp, Explaining heterogeneity in meta-analysis: a comparison of methods. Stat Med 1999; 18:2693-2708]. This can be corrected by dividing the reported standard error of the coefficient by the reported residual standard error [Burgess, Dudbridge, and Thompson, Re: “Multivariable Mendelian randomization: the use of pleiotropic genetic variants to estimate causal effects” (letter). Am J Epidemiol 2015; 181(4):290-291]. However, in order to account for the heterogeneity introduced by pleiotropy under the framework outlined in this paper, we fit a model allowing for multiplicative random effects. No correction to the point estimate is required under this model, since it is mathematically equivalent to the fixed-effect estimate. Standard errors are calculated by only constraining the residual standard error to be 1 when it is less than 1 (which would imply under-dispersion), and otherwise allowing the residual standard error to take its estimated value.

The same rationale is applied to adjust the standard error reported by the MR-Egger method, and confidence intervals should be constructed using a t-distribution on the appropriate number of degrees of freedom (the number of genetic variants minus 2).

**Bootstrap standard errors**

Alternatively, a bootstrap approach to calculate standard errors, confidence intervals and p-values is also recommended. The above code performs a parametric bootstrap by drawing association estimates from their estimated sampling distributions, and performing the MR-Egger method for each randomly drawn dataset. We show this for the slope parameter only. The authors would like to thank Jon White (University College London) for helpful discussions on writing the code for implementing the methods.

Stata code for performing the same analysis is given below.

Disclaimer: The R and Stata code given here is, to the best of our knowledge and at the time of publication, correct. Please contact the lead author if you have concerns.

**Stata code**

To implement the IVW approach (with MAF corrected weights) in Stata, the equivalent code is

```
regress BetaYG BetaXG [aw=1/seBetaYG^2], nocons
```

To implement MR-Egger regression (with MAF corrected weights) in Stata, the equivalent code is

```
regress BYG BXG [aw=1/seBetaYG^2]
```
where BYG and BXG have been derived from BetaXG and BetaYG as above.

The corrected standard error for the IVW method can be calculated by:

```
regress BetaYG BetaXG [aw=1/seBetaYG^2], nocons
local wrongse = _se[BetaXG]
local degfree = e(df_r)
predict BetaYG_fit
gen BetaYG_sqres = (BetaYG_fit-BetaYG)^2*seBetaYG^2
summ BetaYG_sqres
local weighted_rss = r(sum)
di `wrongse'/min(1,sqrt(`weighted_rss'/`degfree'))
```

The correction for the Egger method is similar.

Bootstrap confidence intervals for the MR-Egger method (with MAF corrected weights) can be calculated by:

```
gen BYG_boot = .
gen BXG_boot = .
set obs 10000
generate float boot = .
forvalues i = 1/10000 {
    quietly replace BYG_boot = rnormal(BetaYG, seBetaYG)
    quietly replace BXG_boot = rnormal(BetaXG, seBetaXG)
    quietly replace BYG_boot = BYG_boot * sign(BXG_boot)
    quietly replace BXG_boot = abs(BXG_boot)
    quietly regress BYG_boot BXG_boot [aw=1/seBetaYG^2]
    quietly replace boot = _b[BXG_boot] in `i'
}
```

centile boot, centile (2.5 97.5)

**Further details of the simulation study setup**

In the simulation study, data were generated from the following model:

\[
U_i = \sum_{j=1}^J \phi_j G_{ij} + \epsilon_i^U
\]

\[
X_i = \sum_{j=1}^J \gamma_j G_{ij} + U_i + \epsilon_i^X
\]

\[
Y_i = \sum_{j=1}^J \alpha_j G_{ij} + \beta X_i + U_i + \epsilon_i^Y.
\]
This is a more general version of model (1) and (2), in that it additionally allows genetic variant \(G_j\) to be associated with confounding variable \(U_i\). This occurs when \(\phi_j\) is non-zero (a violation of IV1). If \(\phi_j = 0\) for all \(j\), then this model reduces to equations (1) and (2). In all simulations: the \(G_j\) were generated from a trinomial distribution, taking values (0, 1, 2) with probabilities (0.49, 0.42, 0.09) – this is equivalent to a single nucleotide polymorphism with minor allele frequency 0.3; error variables \(\delta^G_i, \delta^U_i\) and \(\delta^y_j\) were independently generated from a \(N(0, 2)\) distribution; and instrument strength parameters \(\gamma_j\) were generated from a Uniform(0.5, 4) distribution.

The performance of the standard IVW method and MR-Egger regression were investigated in a two-sample Mendelian randomization analysis context with \(J = 25\) variants, with a null (\(\beta = 0\)) and a positive (\(\beta = 0.05\)) causal effect. Two independent samples of \(N\) subjects were generated from the above model. For variant \(j\) out of 25, estimates for the gene-exposure associations (\(\hat{\gamma}_j\)) were obtained from the first sample and estimates for the gene-outcome associations (\(\hat{\gamma}_j\)) were obtained from the second sample, in order to calculate the ratio estimates \(\hat{\beta}_j = \hat{\gamma}_j / \hat{\gamma}_j\). Simulation scenarios (a)–(d) were implemented by additionally specifying values for \(\alpha_j\) and \(\phi_j\) as below:

- No pleiotropy, InSIDE satisfied: \(\alpha_j = 0, \phi_j = 0\);
- Balanced pleiotropy, InSIDE satisfied: \(\alpha_j \sim \text{Uniform}(-0.2, 0.2), \phi_j = 0\);
- Directional pleiotropy, InSIDE satisfied: \(\alpha_j \sim \text{Uniform}(0, 0.2), \phi_j = 0\);
- Directional pleiotropy, InSIDE not satisfied: \(\alpha_j \sim \text{Uniform}(0, 0.2), \phi_j \sim \text{Uniform}(0, 0.5)\).

In scenario (a), the ratio estimand based on the \(j\)th variant is equal to \(\beta\). In scenario’s (b) and (c), the ratio estimand based on the \(j\)th variant is equal to \(\beta + \frac{\alpha_j}{\gamma_j}\) but InSIDE holds. In scenario (d) the ratio estimand based on the \(j\)th variant is equal to

\[
\frac{\beta + \alpha_j + \phi_j}{\gamma_j + \phi_j}.
\]

The InSIDE assumption is not satisfied in this case because the numerator of the bias term (which represents the total ‘direct’ effect not via the exposure) and its denominator (which represents the instrument strength) contain the common term \(\phi_j\). Simulation results are shown in Table 1.

Data for the four funnel plots shown in Figure 5 were generated under the causal null hypothesis for scenarios (a)–(d), using the same two-sample approach. In order to accentuate the shapes of the funnel plots for illustrative purposes, we used \(J = 50\) genetic variants and doubled the range of the Uniform sampling densities for \(\alpha_j\) and \(\phi_j\). Web Figure A2 shows the equivalent scatter plots.
Results from the simulation study in a one-sample setting

The previous simulations were repeated in a one-sample Mendelian randomization setting. One sample of $N$ subjects was generated, and estimates for the gene-exposure associations ($\hat{\gamma}_j$) and estimates for the gene-outcome associations ($\hat{\Gamma}_j$) were obtained from the same sample.

Results are shown in Table A1 and A2. The pattern of results is generally similar to the two-sample case, but both methods perform slightly worse in terms of small sample bias and Type I error rate inflation, and markedly worse with weak instruments. Weak instrument bias is more problematic in the one sample context because it acts in the direction of the confounded observational association. It appears to be slightly worse for estimates from MR-Egger regression, while Type I error rate is worse for estimates from the IVW method.

These simulations were repeated using standard one-sample TSLS instead of IVW as the comparator method, and the results were very similar (data not shown). We conclude that IV analysis with weak instruments in a one-sample setting is troublesome, and that these difficulties are not resolved by the application of MR-Egger regression.

| $N$ | Inverse-variance weighted | | MR-Egger regression |
|-----|--------------------------|------------------|
|     | Mean F statistic | Mean estimate (mean SE) | Power to detect causal effect | Mean estimate (mean SE) | Power of MR-Egger test | Power to detect causal effect |
| No causal effect: $\beta = 0$ |
| Scenario (a) – no pleiotropy, InSIDE satisfied |
| 250  | 10.4 | 0.005 (0.021) | 0.077 | 0.016 (0.046) | 0.062 | 0.075 |
| 500  | 19.8 | 0.003 (0.015) | 0.067 | 0.011 (0.035) | 0.058 | 0.067 |
| 750  | 29.2 | 0.002 (0.013) | 0.060 | 0.009 (0.030) | 0.058 | 0.068 |
| 1000 | 38.6 | 0.002 (0.011) | 0.056 | 0.007 (0.026) | 0.056 | 0.060 |
| Scenario (b) – balanced pleiotropy, InSIDE satisfied |
| 250  | 10.4 | 0.005 (0.023) | 0.081 | 0.017 (0.051) | 0.058 | 0.075 |
| 500  | 19.9 | 0.003 (0.018) | 0.065 | 0.011 (0.041) | 0.056 | 0.068 |
| 750  | 29.2 | 0.002 (0.016) | 0.058 | 0.009 (0.037) | 0.053 | 0.063 |
| 1000 | 38.6 | 0.002 (0.014) | 0.063 | 0.007 (0.034) | 0.054 | 0.062 |
| Scenario (c) – directional pleiotropy, InSIDE satisfied |
| 250  | 10.4 | 0.043 (0.022) | 0.470 | 0.035 (0.047) | 0.052 | 0.124 |
| 500  | 19.8 | 0.040 (0.016) | 0.644 | 0.022 (0.037) | 0.081 | 0.100 |
| 750  | 29.2 | 0.039 (0.014) | 0.770 | 0.017 (0.032) | 0.113 | 0.086 |
| 1000 | 38.6 | 0.039 (0.012) | 0.853 | 0.013 (0.029) | 0.159 | 0.080 |
| Scenario (d) – directional pleiotropy, InSIDE violated |

Mean F statistic, Mean estimate (mean SE), Power to detect causal effect, Mean estimate (mean SE), Power of MR-Egger test, Power to detect causal effect.
<table>
<thead>
<tr>
<th></th>
<th>Scenario (a) – no pleiotropy, InSIDE satisfied</th>
<th>Scenario (b) – balanced pleiotropy, InSIDE satisfied</th>
<th>Scenario (c) – directional pleiotropy, InSIDE satisfied</th>
<th>Scenario (d) – directional pleiotropy, InSIDE violated</th>
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<td>250</td>
<td>10.4 0.055 (0.021) 0.678 0.067 (0.046) 0.063 0.306</td>
<td>10.4 0.056 (0.023) 0.609 0.068 (0.051) 0.059 0.277</td>
<td>10.4 0.093 (0.022) 0.968 0.085 (0.047) 0.051 0.426</td>
<td>10.6 0.179 (0.023) 1.000 0.138 (0.052) 0.127 0.710</td>
</tr>
<tr>
<td>500</td>
<td>19.8 0.053 (0.015) 0.886 0.061 (0.035) 0.058 0.397</td>
<td>19.9 0.053 (0.018) 0.786 0.062 (0.041) 0.057 0.317</td>
<td>19.8 0.090 (0.016) 0.999 0.073 (0.037) 0.080 0.483</td>
<td>20.2 0.176 (0.019) 1.000 0.112 (0.045) 0.322 0.661</td>
</tr>
<tr>
<td>750</td>
<td>29.2 0.052 (0.013) 0.962 0.059 (0.030) 0.059 0.479</td>
<td>29.2 0.052 (0.016) 0.868 0.059 (0.037) 0.053 0.345</td>
<td>29.2 0.089 (0.014) 1.000 0.067 (0.032) 0.112 0.523</td>
<td>29.7 0.176 (0.018) 1.000 0.101 (0.042) 0.463 0.642</td>
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<tr>
<td>1000</td>
<td>38.6 0.052 (0.011) 0.988 0.058 (0.026) 0.056 0.560</td>
<td>38.6 0.052 (0.014) 0.911 0.058 (0.034) 0.054 0.371</td>
<td>38.6 0.089 (0.012) 1.000 0.063 (0.029) 0.157 0.565</td>
<td>39.3 0.175 (0.017) 1.000 0.093 (0.039) 0.572 0.622</td>
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</tbody>
</table>

**Web Table A1:** Performance of inverse-variance weighted and MR-Egger regression estimates in simulation study for one-sample Mendelian randomization with a null (β = 0) and a positive (β = 0.05) causal effect. All tests are performed at 5% significance level. SE = standard error.
Scenario (c) – directional pleiotropy, InSIDE satisfied

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<td>0.008 (0.020)</td>
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<td>1.000</td>
<td>0.010 (0.015)</td>
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<td>100</td>
<td>20.7</td>
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<td>1.000</td>
<td>0.014 (0.010)</td>
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<td>150</td>
<td>14.2</td>
<td>0.038 (0.004)</td>
<td>1.000</td>
<td>0.018 (0.008)</td>
</tr>
</tbody>
</table>

**Web Table A2:** Performance of inverse-variance weighted and MR-Egger regression estimates in a simulation study for one-sample Mendelian randomization with a null causal effect ($\beta = 0$) and a fixed sample size, and varying the number of instruments ($J$)

### Web Figures

**Web Figure A1:** Genetic associations with blood pressure and coronary artery disease risk from 29 variants -- scatter plots of minor allele frequency corrected genetic associations with blood pressure ($\hat{C} \hat{C}$) against genetic associations with coronary artery disease ($\hat{\Gamma} \hat{\Gamma}$). Left: scatter plot for systolic blood pressure. Right: scatter plot for diastolic blood pressure. The inverse-variance weighted (IVW, red) and MR-Egger (blue) causal effect estimates are also shown as regression slopes.
Web Figure A2: Scatter plots of minor allele frequency corrected genetic associations with exposure ($\hat{\gamma}_j^C$) against genetic associations with the outcome ($\hat{\gamma}_j$) for 50 variants in four scenarios: (a) no pleiotropy; (b) balanced pleiotropy; (c) directional pleiotropy, InSIDE assumption satisfied; and (d) directional pleiotropy, InSIDE assumption not satisfied. The inverse-variance weighted (IVW, red) and MR-Egger (blue) causal effect estimates are also shown.
\( \hat{\Gamma}_j \)

\( \hat{C}_{\gamma_j} \)

IVW = 0.223

Egger = 0.086