Mendelian randomization - an introduction

Bjørn Olav Åsvold

K.G. Jebsen Center for Genetic Epidemiology, NTNU December 2020



This lecture will cover

- The purposes of Mendelian randomization studies
- Principles underlying Mendelian randomization
- Assumptions of Mendelian randomization
- Strengths and limitations

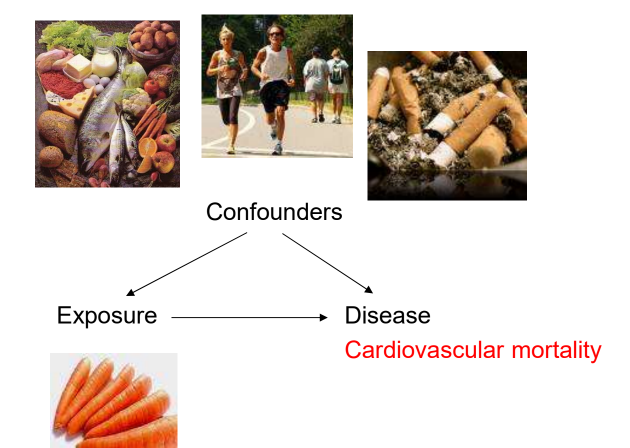


What's the purpose of Mendelian randomization?

- In epidemiology, we often aim to examine causes of disease
- Causal inference is often challenged by the possibility of confounding or reverse causality
- Mendelian randomization: a study approach where genetic variants are used to examine the causal effects of modifiable exposures

The shortcomings of observational studies – one out of many examples: ß-carotene & CVD mortality

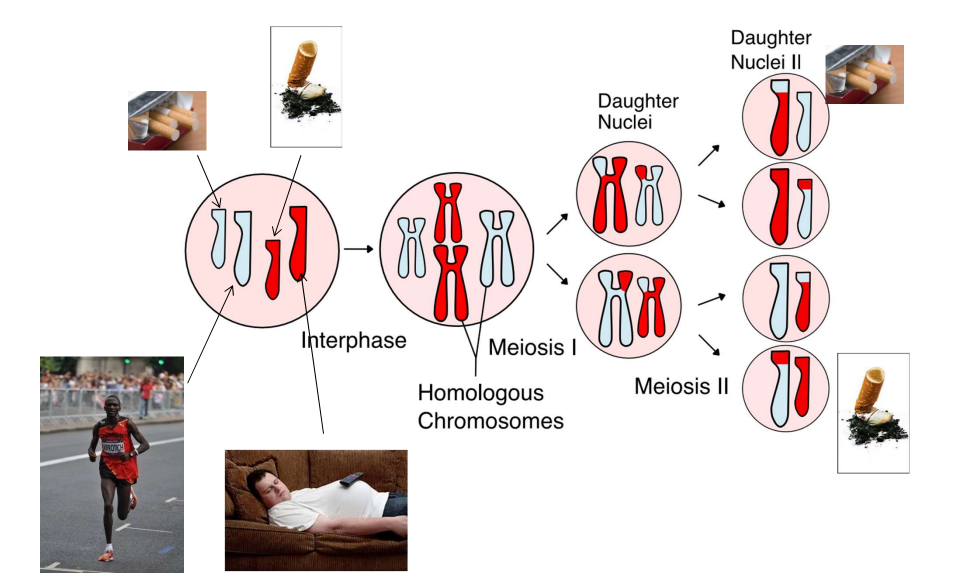
Cohorts Male health workers (United States) Male social insurance workers (Finland) Female social insurance workers (Finland) Male chemical workers (Switzerland) Hyperlipidaemic men (United States) Nursing home residents (United States) Cohorts combined 0.5 0.75 1.25 1.5 1.75 0.1 Relative risk (95% CI)



Example: CRP levels and coronary heart disease



- Mendelian randomization: Genetic variants are used as proxies instrumental variables – for modifiable risk factors in observational studies.
- Relies on Mendel's laws of inheritance:
- Generally, genes are randomly assorted from parents to offspring, and inheritance of one trait is independent of inheritance of other traits.

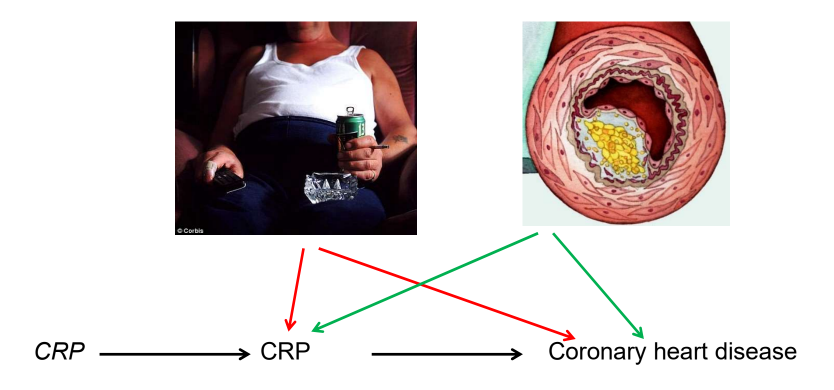


- Mendelian randomization: Genetic variants are used as proxies instrumental variables – for modifiable risk factors in observational studies
- Relies on Mendel's laws of inheritance:
- Generally, genes are randomly assorted from parents to offspring, and inheritance of one trait is independent of inheritance of other traits.

CRP level

Genetic variants associated with a specific trait will not be liable to confounding by behavioual, socioeconomic and physiological factors.
 obesity, smoking, preclinical CHD

Example: CRP levels and coronary heart disease

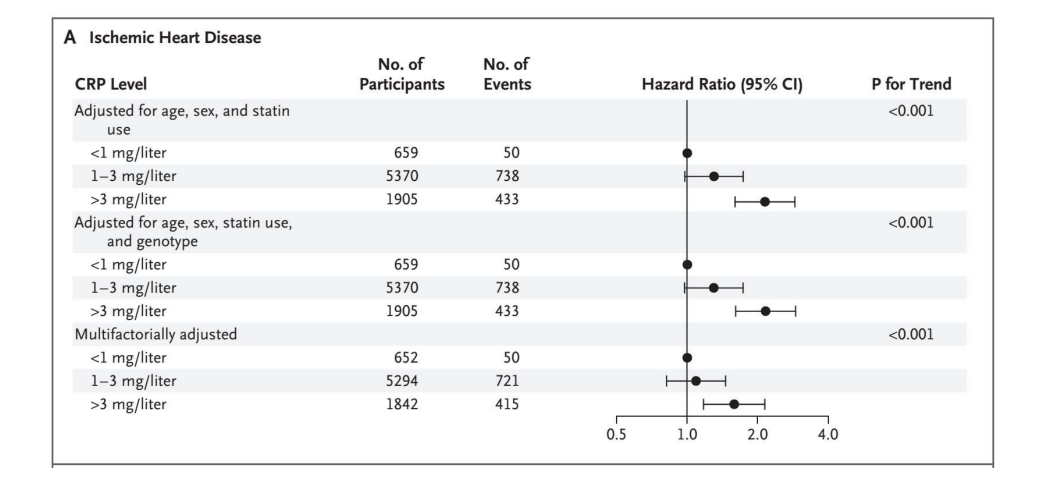


The NEW ENGLAND JOURNAL of MEDICINE

ORIGINAL ARTICLE

Genetically Elevated C-Reactive Protein and Ischemic Vascular Disease

Jeppe Zacho, M.D., Anne Tybjærg-Hansen, M.D., D.M.Sc., Jan Skov Jensen, M.D., D.M.Sc., Peer Grande, M.D., D.M.Sc., Henrik Sillesen, M.D., D.M.Sc., and Børge G. Nordestgaard, M.D., D.M.Sc.



e			Change in CRP Level %	No. of Partici- pants	Mean ±SE
type c	ombir	nation	ı		
CC	CC	TT	0	3,887	H
CC	CC	TT	7	6,876	1
CC	CC	TT	18	3,051	H
CT	CT	TT	18	7,006	H
CT	CT	TT	25	6,486	H
CC	CA	GT	35	1,125	H
TT	TT	TT	38	3,412	H
CC	CA	GT	36	998	H
CT	AT	GT	64	1,045	_
			22	17,668	H
			43	5,455	Н
				0	.0 1.0 2.0 3.0 4.0
					Plasma CRP (mg/liter)
	CC CC CT CT CC TT CC	otype combine CC CC CC CC CC CC CT CT CT CT CC CA TT TT CC CA	otype combination CC CC TT CC CC TT CT CT TT CC CA GT TT TT TT CC CA GT	in CRP Level % otype combination CC CC TT 0 CC CC TT 7 CC CC TT 18 CT CT TT 18 CT CT TT 25 CC CA GT 35 TT TT TT 38 CC CA GT 36 CT AT GT 64	in CRP Level pants otype combination CC CC TT 0 3,887 CC CC TT 7 6,876 CC CC TT 18 3,051 CT CT TT 18 7,006 CT CT TT 25 6,486 CC CA GT 35 1,125 TT TT TT 38 3,412 CC CA GT 36 998 CT AT GT 64 1,045

- Higher CRP levels were associated with increased risk of coronary heart disease,
- but genetically determined higher CRP levels were not associated with increased risk of CHD.
- This strongly suggests that elevated CRP levels do not cause coronary heart disease,
- and that the observed association between CRP levels and risk of coronary heart disease is explained by confounding or reverse causality.



Randomized study?

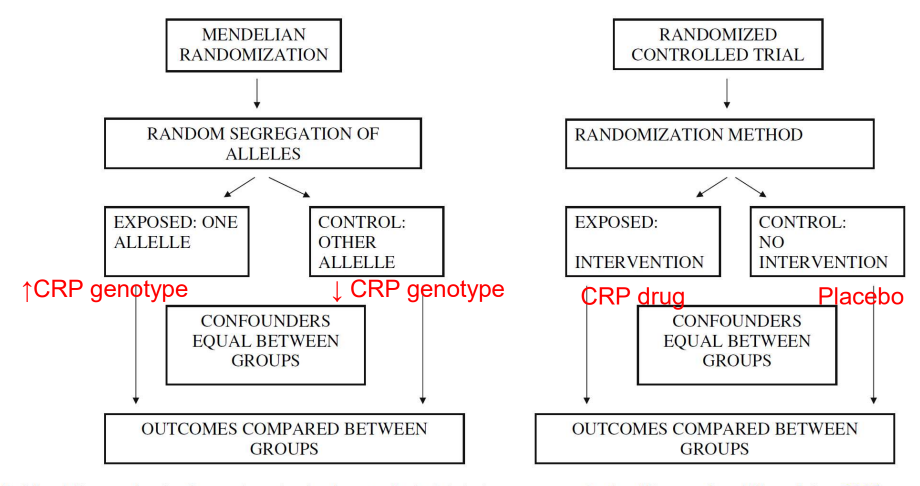
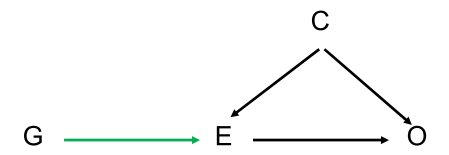


Fig. 7 Mendelian randomization and randomized controlled trial designs compared, after Hingorani and Humphries (2005)

3 key assumption of MR studies

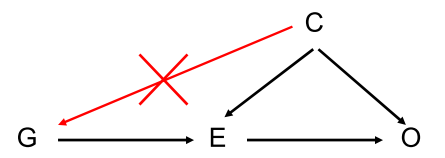
1. The genetic instrument is associated with the exposure (the relevance assumption).





3 key assumption of MR studies

2. There is no association between the genetic instrument and confounders of the exposure-outcome relation – in other words: the genetic instrument shares no common causes with the outcome (the independence assumption).



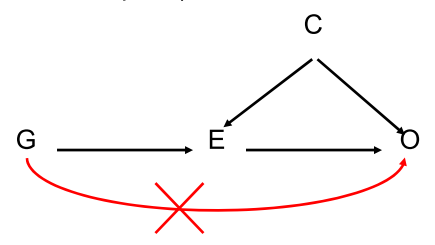
Such associations can occur due to

- Weak instrument bias
- Horizontal pleiotropy
- Linkage disequilibrium among gene loci
- Population stratification

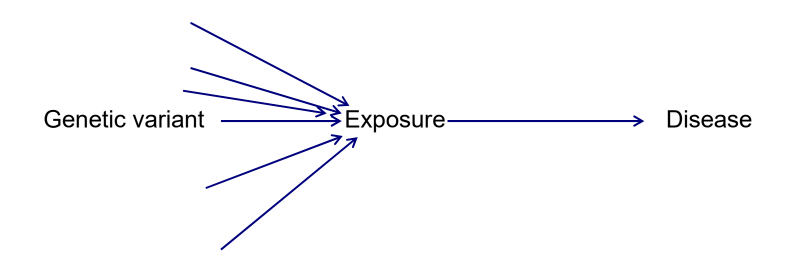
100

3 key assumption of MR studies

3. There is no association between the genetic instrument and the outcome except through the exposure of interest (the exclusion restriction assumption).



This assumption is violated if there is horizontal pleiotropy: a genetic variant affects multiple traits through separate pathways



MR study designs

- We need to estimate 2 associations:
- Gene-exposure association: G-X
- Gene-outcome association: G-Y
- We usually estimate the causal effect: G-Y / G-X
- One-sample MR: G-X and G-Y estimated in the same study sample
- Two-sample MR: G-X and G-Y estimated in different samples
- Bidirectional MR: Both causal directions examined
- Multivariable MR: Several correlated exposures examined
- Two-step MR: Mediation analysis
- Gene x environment interaction
-



How to select the genetic instruments?

- Two general approaches:
- 1. Select SNPs with proven or plausible biological effect on the target exposure
- 2. Select SNPs from GWAS

Some limitations of MR – and possible solutions

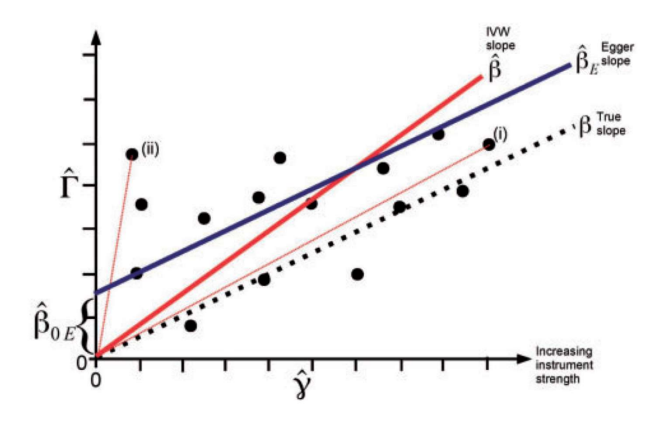
- Lack of suitable instruments
 - More GWAS
- Low power
 - □ Combine multiple SNPs (genetic risk score or meta-analysis), larger N
- Winner's curse (overestimation of SNP-exposure association)
 - consequences differ by study design
- Poor biological understanding possibilities for pleiotropy
 - Select variants with known biology
 - □ If multiple variants: numerous sensitivity analyses available
- Trait heterogeneity (SNPs associated with multiple dimensions of a single trait)
- Time-varying exposures
- Effects on disease risk vs. disease progression
- Dynastic effects, population stratification, assortative mating
 - Within-family MR

Some Mendelian randomization analyses

(sensitivity analyses to examine / correct for pleiotropy)

- Inverse-variance weighted MR: main analysis, meta-analysis
 (weighted average) of ratio estimates, assumes balanced pleiotropy
- MR-Egger regression: allows for directional pleiotropy, but has other assumptions and lower power
- Weighted median: valid if the majority of the information comes from valid instruments
- Weighted mode: assumes that the mode of instruments is valid
- MR-PRESSO, MR-Lasso, MR-Robust: assume that there is a small number of invalid instruments (outliers)
- MR-RAPS
- Contamination mixture
- MRMix

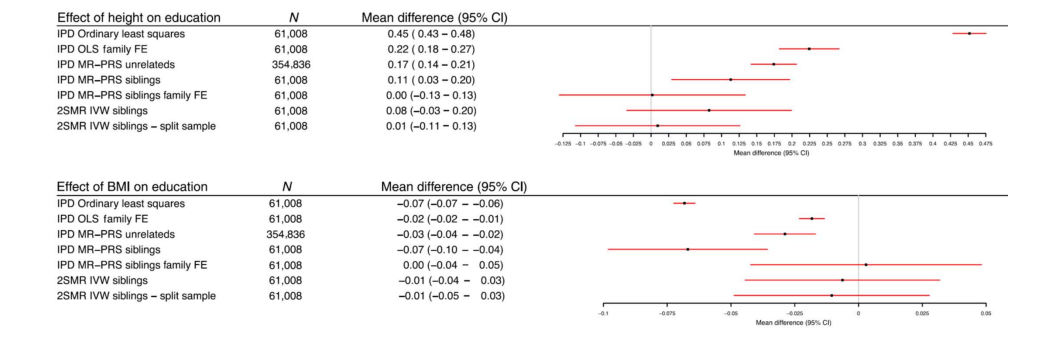
See for example: Slob & Burgess, Genetic Epidemiology 2020;44(4):313-29.



Bowden et al., IJE 2015

Some limitations of MR – and possible solutions

- Lack of suitable instruments
 - More GWAS
- Low power
 - □ Combine multiple SNPs (genetic risk score or meta-analysis), larger N
- Winner's curse (overestimation of SNP-exposure association)
 - consequences differ by study design
- Poor biological understanding possibilities for pleiotropy
 - Select variants with known biology
 - □ If multiple variants: numerous sensitivity analyses available
- Trait heterogeneity (SNPs associated with multiple dimensions of a single trait)
- Time-varying exposures
- Effects on disease risk vs. disease progression
- Dynastic effects, population stratification, assortative mating
 - Within-family MR





Another use of MR: Identify intended and unintended drug effects



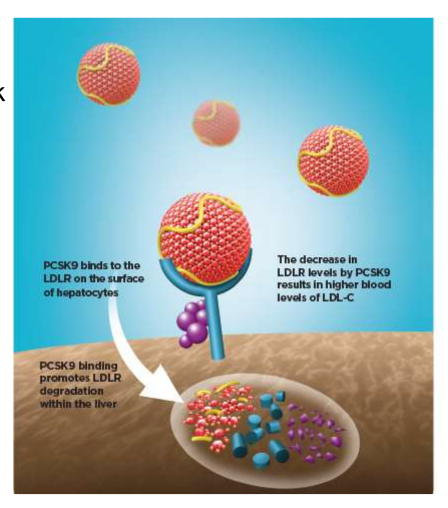


PCSK9 inhibitors: New, effective lipid-lowering drugs that reduce the risk of cardiovascular events

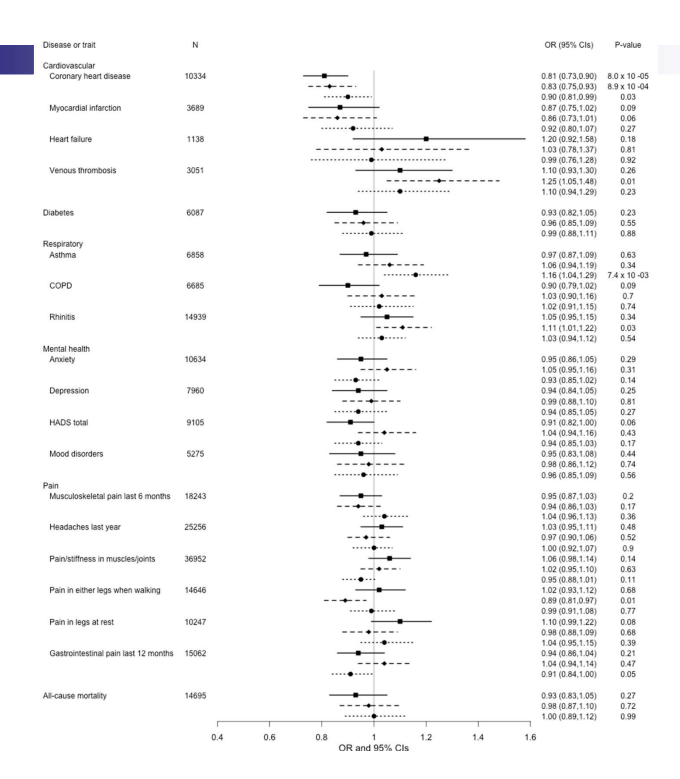
Is long-term PCSK9 inhibition hazardous?

We lack long-term RCT evidence

Ben Brumpton et al., Circ Genom Precis Med 2019



Trait	N		Beta (95% CI)	P-value
Cardiovascular				
LDL cholesterol (mmol/L)	68035	•	-0.26 (-0.29,-0.23)	8.3 x 10 -49
,		→ -	-0.25 (-0.28,-0.21)	2.8 x 10 -43
			-0.25 (-0.29,-0.22)	3.8 x 10 -53
HDL cholesterol (mmol/L)	69123	•	0.02 (0.01,0.03)	3.0 x 10 -03
		•	0.01 (-0.01, 0.02)	0.26
		•	0.02 (0.01,0.03)	2.5 x 10 -03
Total cholesterol (mmol/L)	69133	•	-0.27 (-0.30,-0.23)	9.4 x 10 -42
		+	-0.27 (-0.31,-0.23)	1.3 x 10 -40
		••	-0.26 (-0.29,-0.22)	3.9 x 10 -45
Triglycerides (mmol/L)	69379	-	-0.05 (-0.09,-0.01)	0.01
		•	-0.04 (-0.08,0.00)	0.04
		••	-0.05 (-0.09,-0.02)	2.9 x 10 -03
Systolic blood pressure (mmHg)	69076		0.00 (-0.61,0.61)	0.67
			0.16 (-0.46,0.78)	0.26
			-0.16 (-0.73,0.41)	0.14
Diastolic blood pressure (mmHg)	69076		-0.19 (-0.57,0.20)	0.77
			-0.03 (-0.42,0.37)	0.1
			-0.36 (-0.72,0.00)	0.02
Weight (kg)	68847		-0.08 (-0.52,0.37)	0.7
			-0.10 (-0.55,0.35)	0.07
DMI (Ice/ex2)	60036		-0.33 (-0.74,0.08)	0.12
BMI (kg/m2)	68836		-0.04 (-0.18,0.10)	0.57
			0.00 (-0.14,0.15)	0.96
Glucose non-fasting (mmol/L)	69123		-0.04 (-0.17,0.10) 0.04 (-0.01,0.09)	0.6 0.13
Glucose Hori-lasting (Hillion)	03123		-0.02 (-0.07,0.03)	0.13
		Ţ <u>.</u> .	0.04 (-0.01,0.09)	0.12
C-reactive protein (mg/L)	48434		0.04 (-0.01,0.09)	0.52
C-reactive protein (mg/L)	40404		0.11 (-0.13,0.35)	0.37
			-0.13 (-0.35,0.10)	0.27
Respiratory		•	0.10 (0.00,0.10)	0.21
FEV1 (L/s)	17136	-	0.04 (-0.01,0.08)	0.09
(2.5)			-0.03 (-0.08,0.01)	0.15
			0.01 (-0.03,0.05)	0.66
FVC (L)	16700	-	0.03 (-0.02,0.08)	0.22
		-+	-0.04 (-0.09,0.01)	0.12
		· • ·	0.02 (-0.03, 0.06)	0.49
FEV1/FVC ratio	16700	•	0.00 (0.00,0.01)	0.19
		•	0.00 (-0.01,0.00)	0.67
		•	0.00 (-0.01,0.01)	0.96
Mental health				
HADS-Anxiety	59071		-0.17 (-0.29,-0.05)	6.7 x 10 -03
			0.03 (-0.10,0.15)	0.67
		•	-0.08 (-0.18,0.05)	0.26
HADS-Depression	61863		-0.08 (-0.19,0.03)	0.14
			0.03 (-0.09,0.12)	0.77
		•.	-0.08 (-0.18,0.02)	0.1
HADS-Total	63365		-0.23 (-0.43,-0.03)	0.02
			0.05 (-0.16,0.24)	0.7
			-0.18 (-0.36,0.01)	0.07
	-	0.8 -0.4 0 0.4 0.8	3	
		Beta and 95% CI		





PCSK9 inhibition

- Genetically determined, life-long PCSK9 inhitibion leads to lower LDL cholesterol and lower risk of coronary heart disease.
- We did not find convincing evidence of side effects.
- Such a study does not replace the need for RCTs,
- but gives information about anticipated effects and side effects long before evidence from long-term RCTs is available.

	MR Study	RCT
Cost	Low cost, mostly use of existing data	High cost, generating data
Time	Quick and efficient	Very intensive and lengthy
Population	Very large sample size, usually representative of target population	Small sample size, usually not representative of target population
Exposure	Exposed to genetic variant at birth by random	Exposed to the drug by random
Outcomes	Multiple outcomes at once, hypothesis generating, can study life time diseases	Specific outcome, hypothesis testing, difficult to study diseases with long latent period
Drug Effects	Both common and uncommon Short-term and life-time effects	Mostly common Mostly short-term effects
Ethics	No ethical issues, no exposure to the drug	Ethical issue due to exposure to a new drug

Summary

- Mendelian randomization is a way of utilizing genetic variation to study causal effects of modifiable exposures
- Mendelian randomization is not generally prone to confounding, is immune to reverse causation, and can inform about effects of life-long exposure,

but has other limitations and relies on assumptions that are only partially testable