Causal inference in the social, health and business sciences: A very brief practical primer in R

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1 Introduction

Causal inference is about what works, when, for whom and under what circumstances. When an event occurs – a medical treatment, social intervention, marketing move, etc. – we want to know what happened as a result, if anything. But – and here's the crucial insight – we also need to know what would have happened *in the absence of* the event, all else being equal. That is, the causal effect of an event is the difference between what actually happened and the potential but actually-not-occurring counter-factual.

This working paper is a very brief practical primer on causal inference in the health and social sciences. We'll use minimal formal notation and instead emphasize practical computational implementation using the R scripting language. It assumes a basic understanding of Bayesian regression and generalized linear modeling in R. Most importantly, however, it assumes an interest in the question of how we can ever know whether something caused something else and to what extent.

Without further ado, let's dive in.

2 The fundamental issue of causal inference: Observed, potential and missing outcomes

We said just above that the causal effect of an event is the difference between what actually happened and the potential but actually-not-occurring counter-factual. But here immediately the fundamental issue of causal inference comes crashing down at our feet: We can never observe the counter-factual!

We can never re-play history or anything of the sort. For instance, we can never give a medical treatment to a patient and also *not* give the treatment to said patient. If we could, we could simply take the difference between the treatment and no treatment states to obtain an **individual treatment effect**. That'd be ideal. But it's not feasible, for obvious reasons. The counter-factual outcome is missing. Causal inference is, then, fundamentally a missing data problem.

So, the best we can do is to *infer* the counter-factual by estimation, approximation or informed (hopefully) guesswork and contrast that with the observed state. More precisely, the best we can do is to aim at an **average treatment effect** or variants thereof. An average treatment effect of, say, a medical treatment is the difference in outcomes under receiving the treatment vs. not receiving the treatment not within an individual, like the individual treatment effect, but instead *between* individuals. Further, these individuals have to be similar enough in relevant characteristics such that, whatever the difference in outcome between the treatment and no treatment groups is, in fact, explained by the treatment alone, and not by some other events or developments.

This, in a nutshell, is the fundamental issue of causal inference. It's simple. Deceptively so, because the real world is a mess. So we have assumptions to make. But in return we get something quite extraordinary: A rigorous, tried-and-tested framework for thinking about cause-and-effect-relationships. The fundamental issue of causal inference, then, is also a *promise*: A promise that, if we're careful, honest, and transparent, we

might actually get causal answers to the questions that matter most in life. Or, alternatively, we'll be told that there exist no valid answers to the questions we posed. Either is a very valuable yield.

2.1 Some notation

To make the discussion more succinct, we'll use some simple formal notation and terminology. It's useful to learn these for another reason, too: perhaps you'll want to navigate the causal inference literature on your own, when you're done here.

Throughout, Y is our outcome on which our cause has or hasn't an effect, and X is the cause, our predictor of interest – say, receiving a particular medical treatment X = 1 or not X = 0. That is, in its simplest form, an average treatment effect is simply the difference in expected outcomes between receiving the treatment $Y^{X=1}$ or not $Y^{X=0}$. This quantity is often written as $E[Y^{X=1}] - E[Y^{X=0}]$, where E(.) is the expectation operator, i.e. it computes the expected value.

2.2 DAGs and experimental randomization

One useful tool for thinking through a causal modeling problem is **DAG**s, directed acyclic graphs. A DAG graphs the assumed¹ causal relationships among variables and can thereby guide subsequent analytic strategies for recovering the causal effects of interest (or can reveal that no such effects can be recovered under anu circumstances, given the DAG). Directed means that the relationships, denoted by arrows, are causal. Acyclic means that variables can't cause themselves dynamically. And graph means... well, that should be obvious.

Figure 1 is one example of a DAG. It shows a simple but extremely common scenario in which both our cause X and outcome Y are influenced by a third variable Z. In other words, Z is a **confounder**. We'll discuss confounding in more depth in sections below. For now, suffice it to say that our estimate of the cause of X on Y will be biased if we do not take into account Z. How do we know that Z confounds the relationship between X and Y and thereby induces bias? Simple, we can read it off the DAG.



Figure 1: Directed acyclic graph (DAG) of confounding.

To see this, note that Z has a causal effect on both X and Y. This is technically called a **backdoor path**. A backdoor path is any set of arrows that both points into the outcome Y and predictor of interest X. In a basic sense, causal inference is about identifying such backdoor paths and then "blocking' them, using study design or statistical adjustment. This simple insight makes analyzing DAGs a sort of party game or puzzle. It can actually be fun to draw out more or less complicated DAGs and then identify how and whether a causal effect can be recovered under this particular graph, through backdoor paths and the likes. There's more to graph analysis than looking for and blocking backdoor paths, as we'll see below. But it's a good start.

So how do we block backdoor paths? I said just above that we can, among other approaches, use particular study designs. Randomization is the best known example of such a study design. What randomization does, from the perspective of DAGs, is that it deletes any arrows that go into X, meaning that there can longer be any backdoor paths, by definition. Figure 2 illustrates this. It's similar to figure 1, except that there's no arrow from Z to X. Under this model, an estimate of the effect of X on Y is no longer biased and we don't need to measure Z^2 .

Randomization allows us to delete arrows going into X from all other variables because randomization is, well, random. So, whether an individual receives a randomized medical treatment will not be dependent on anything else than the randomization mechanism. In the next section, we'll show in code and simple simulated data how this works.

¹Based on prior studies, theory, commonsense, or other sources of inference.

²Although, on a technical aside, knowing Z potentially increases the precision of our estimate of X.



Figure 2: DAG of randomized X.

In sum, randomization is an efficient strategy for causal estimation, because it makes X independent of all other variables in the graph. This is the reason why randomized controlled studies are often referred to as the "gold-standard" for causal inference. But, unfortunately, our work does not end here. Randomization is often imperfect (e.g., participant might not perfectly adhere to the randomized treatment) or not feasible for practical or ethical reasons (e.g., could we assign people to smoke in a randomized fashion such that we could estimate the causal effect of smoking on, say, cancer?).

This does not mean that all hope is lost. Causal inference in observational or pseudo-randomized contexts is possible. But it does force us to make some assumptions and add some steps to our workflow. For instance, through relatively simple statistical techniques, we can block backdoor paths that couldn't be deleted by randomization. Next section introduces the promises and pitfalls of this approach, using regression modeling.

3 Blocking backdoors

This section riffs on – and in a few ways goes slightly beyond – Cinelli's Crash Course (Cinelli, Forney, and Pearl 2020) and McElreath 2023 Rethinking lectures 5 and 6. Many other pedagogical pieces exist, see e.g. Lübke et al. (2020), (rohrer2018thinking?), Wysocki, Lawson, and Rhemtulla (2022).

As such, it will discuss:

Multiple regression as an alternative to randomization for blocking back-door paths; Table 2 fallacy (Westreich and Greenland 2013) and the danger of garbage can regression (e.g., blocking a mediator, collider bias, conditioning on post-treatment; Achen (2005)); conditional vs. marginal effects; some promises (decomposing effects, mediation, front-door criterion).

One critical take-home is that causes are not in the data (for instance, you cannot distinguish between a pipe and a fork from the data alone, because they have similar implications). A DAG cannot be built on the basis of a set of regressions. Regressions can be used to test the implications of a DAG or a set of them (for a discussion of this point in the wild, Purzycki, Bendixen, and Lightner 2022). All this means that variable selection – which variables to include in your adjustment set – cannot be automated; it strictly requires prior knowledge (see also books: Hernan and Robins (2020); Westreich (2019); etc.).

The average person (conditional) vs. people on average (marginal). Y \sim X is the marginal estimate. But adjustment is critical for de-confounding. But, adjustment returns only conditional estimates. Enter g-methods.

4 G-methods and marginal effects

As we saw above, multiple regression is a powerful tool for blocking backdoor paths. This allows for causal interpretation of the focal parameter(s) (but, importantly, *not* the control variables). However, regression has drawbacks that makes it insufficient for many common causal estimands, in particular when there are many covariates with complex relationships. Often, we're interested in marginal, not conditional, average treatment effects – that is, average treatment effects in a population as a whole, not just subsets thereof. In more complicated data structures, such as longitudinal studies with dynamical relationships between treatment and outcome, regression alone will also fall short. In all these cases, we need a few additional tools in the trunk.

One such family of tools is known as g-methods – g for g-eneral or g-eneralized (Robins 1986). Here, we'll focus on one g-method in particular, often referred to as g-computation (e.g. Snowden, Rose, and Mortimer 2011; Ahern, Hubbard, and Galea 2009) or s-tandardization (e.g. Vansteelandt and Keiding 2011; Hernan and

Robins 2020, ch. 13), while we'll also very briefly introduce another method, known as **inverse probability** of treatment weighting (IPTW). Standardization and IPTW rely on the same assumptions and will yield similar if not identical results (indeed, in many simpler cases, they are mathematically identical), but they arrive at their results at quite different analytic routes.

We focus primarily on standardization for a few reasons. First, since we want to perform our data analysis in a Bayesian framework, standardization is an obvious choice since the IPT weights are not obviously compatible with Bayes theorem (for some discussion see Robins, Hernán, and Wasserman 2015; Saarela et al. 2015).

Second, compared to IPTW, standardization and its practical implementation seem to be often overlooked in popular text books and primers on causal inference and econometrics in the social and health sciences (e.g., Morgan and Winship 2015; Pearl, Glymour, and Jewell 2016; Westreich 2019; Angrist and Pischke 2009; McElreath 2020), although the latter stresses contrasts. It is discussed in (Hernan and Robins 2020, ch. 13) but without explicit, reproducible practical/programming application, though there's a g-methods package). However, given its popularity, it's useful to at least be aware of the nuts and bolts of IPTW, too. So let's briefly present the two methods in turn.

4.1 Inverse Probability Weighting

Recall that the main aim of a covariate-adjusted analysis is to obtain conditional exchangeability: When we account for imbalances in covariates between treatment and control group, we say the two groups are exchangeable conditionally on the covariates. This means that the two groups are comparable such that if we detect a difference between the groups after the intervention, we can interpret that difference as a causal effect of the intervention.

The IPTW method obtains conditional exchangeability by, in effect, creating a "pseudo-population" in which covariates are balanced between treatment and control. As the name hints at, this pseudo-population is created by estimating the probability of receiving the treatment conditional on covariates. This probability is then inverted and used as weights in a regression predicting the actual outcome of interest. In code, the procedure looks like this. First, we simulate a simple, confounded data structure, with a binary treatment X with a coefficient of βx , binary confounder Z and a continuous outcome Y.

```
set.seed(123)

n <- 1e4

bX <- 0.5

Z <- rbinom(n, 1, 0.3)
X <- rbinom(n, 1, 0.4+Z*0.5)
Y <- rnorm(n, 10 + X*0.5 + Z*0.5)

gdat <- data.frame(Y=Y, X=X, Z=Z)</pre>
```

Next, we calculate IPT weights by, first, fitting a model that regresses X on Z. Then, for each row in the dataset, we get predictions from this model for the probability of receiving treatment. Then we compute the IPT weights and includes those weights in a regression that predicts the outcome by X. This latter step, the outcome regression, is known as a **marginal structural model** (MSM). It's a marginal model, because the coefficient of X is an average (or marginal, since the estimated weights marginalizes over the distribution of the covariate) treatment effect in the population; and structural, because X has a valid causal interpretation (structural is another word for causal).

```
# inverse logit function
inv_logistic <- function(x) exp(x)/(1+exp(x))

# receiving treatment conditional on Z
treat.mod <- glm(X ~ Z, data = gdat, family = "binomial")</pre>
```

```
# probability of treatment
gdat$pd <- predict(treat.mod) |> inv_logistic()

# compute inverse weights
gdat$w <- with(gdat, ifelse(X==1, 1/pd, 1/(1-pd)))

# MSM of outcome
glm(Y ~ X, data = gdat, weights = w)</pre>
```

The MSM recovers the simulated coefficient for X ($\beta x = 0.5$), even though the model does not adjust for Z in a traditional sense. This is because we adjusted for Z using weights. Note that this simple example could also, of course, be obtained using simple multiple regression that adjusted for X and Z. This, however, will not be so, when we tackle more complex longitudinal data in the next section.

There are at least two ways to improve on our simple IPTW workflow above. First, above we calculated unstabilized weights, but in many cases (and always with continuous exposures), so-called **stabilized weights** are preferred (and in all cases, it's good to know about both). In brief, the stabilized weights are different from the unstabilized in that the numerator is the unconditional inverse probability of treatment and they guard against extreme weights and, in turn, variance-inflation (for more detail, see Chesnaye et al. 2022). The stabilized weights can be calculated with an intercept-only regression on treatment.

```
# intercept-only model on treatment
treat.mod.sw <- glm(X ~ 1, data = gdat, family = "binomial")

# probability of treatment
gdat$pn <- predict(treat.mod.sw) |> inv_logistic()

# compute stabilized weights
gdat$sw <- with(gdat, ifelse(X==1, pn/pd, pn/(1-pd)))

# MSM of outcome with stabilized weight
glm(Y ~ X, data = gdat, weights = sw)</pre>
```

The other thing we can do to improve the workflow is to obtain a measure of uncertainty around our marginal estimate. However, since [placeholder], we can't rely on the standard error from our MSM. Instead, we must resort to bootstrapping, which involves running the same algorithmic routine R times. This will result in a distribution of estimates with length R. Here's one way to implement bootstrapping with the stabilized weights.

That's it for IPTW for now. They are popular and reasonably straightforward to implement. However, as mentioned, since there's no very clear way of obtaining uncertainty in the weights and then propagate that uncertainty to the MSM in a formal Bayesian framework, we leave IPTW here.

4.2 Standardization

Standardization is our main g-method here. It's arguably even more straightforward to implement than IPTW, as it requires us to only fit a single regression of the outcome and then do some post-fitting simulation.

The conceptual steps in standardization are as follows:

- Fit a theoretically informed model of the outcome including the treatment, covariates and possibly non-linear relationships and functional forms (e.g., interaction terms, quadratic terms, etc.)
- Duplicate the original data, set X = 1 for all individuals, and then obtain predictions of the outcome using the fitted model holding covariate(s) Z as observed z, $E[Y^{Z=z,X=1}]$.
- Duplicate the original data, set X = 0 for all individuals, and then obtain predictions of the outcome using the fitted model holding covariate(s) Z as observed z, $E[Y^{Z=z,X=0}]$.
- Compute contrast corresponding to the estimand of interest, $E[Y^{X=1}] E[Y^{X=0}]$.

In code, using Bayesian estimation with default priors via Stan and brms, the steps are these.

```
# E[Y{X=1} - Y{X=0}]
mean_hdci(Y.x1[,1] - Y.x0[,1])
```

The distribution of contrasts calculated in that final step are marginal estimates in that the two expectations $E[Y^{Z=z,X=1}]$ and $E[Y^{Z=z,X=0}]$ marginalize or average over the covariate(s) Z. We summarize the contrast by its posterior mean and HPDI, using the tidybayes package.

When performing standardization in a frequentist setting, we'd have to resort to bootstrapping in order to obtain valid uncertainty around the contrast (see e.g. Snowden, Rose, and Mortimer 2011), as with IPTW. However, in a Bayesian setting, we get uncertainty in one go, since the predicted values are valid posterior distributions of expectations.

Okay, now that we have introduced g-methods in a very simple data context, we're ready to tackle more complex data structures where regression modeling alone falls short.

4.3 Longitudinal analysis, time-varying treatments, doubly robust estimators

For illustrating a longitudinal data context with time-varying treatment, we'll simulate from the DAG in figure 3. Suppose that our treatment X is randomized but administered according to some time-varying covariate Z. We're interested in the *joint effects* of the treatment X_0 and X_1 on our outcome Y (that is, the effects of each treatment on the outcome not through the subsequent treatment), denoted by the dashed edges.

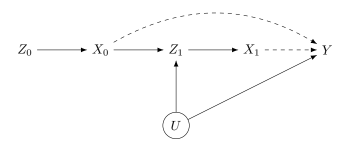


Figure 3: DAG of longitudinal treatment-confounder feedback.

However, some unobserved variable U affects both Z_1 and Y. This confounds the relationship between the treatment and the outcome via the backdoor path $Y \leftarrow U \rightarrow Z_1 \rightarrow X_1$. This in turn means that in order to estimate the hypothesized causal path $X_1 \rightarrow Y$, we'd want to adjust for Z_1 .

But here's the issue: Z_1 is also a collider on the path $Y \leftarrow U \rightarrow Z_1 \leftarrow X_0$, implying that if we condition on Z_1 , we open a non-causal path between X_0 and Y. All in all, under this DAG, we're not able to estimate the joint effects of X_0 and X_1 on Y using a single regression model, since the two treatment indicators imply different adjustment sets. G-methods to the rescue.

First, we simulate some data, under the model presented in figure 3. To keep things simple, we assume that the treatment, in fact, does not have any direct impact on Y (i.e., we could delete the dashed edges), such that the true coefficients of X_0 and X_1 should be (roughly) zero.

```
set.seed(9)

n <- 1e4

U <- rnorm(n, 0, 1)

Z_0 <- rbinom(n, 1, 1/(1+exp(0.5)))

X_0 <- rbinom(n, 1, 1/(1+exp(0.5 + Z_0*0.5)))

Z_1 <- rbinom(n, 1, 1/(1+exp(0.5 + X_0*0.5+U*0.5)))

X_1 <- rbinom(n, 1, 1/(1+exp(0.5 + Z_1*0.5)))</pre>
```

```
Y <- rnorm(n, 10 + U*2)

lgdat <- data.frame(Y=Y, X_0=X_0, X_1=X_1, Z_0=Z_0, Z_1=Z_1, U=U)
```

Then, we verify the chaos that Z_1 can havoc, if we haphazardly estimated the effect of X_0 on while adjusting for Z_1 . The coefficient is solidly non-zero.

```
mx0 <- glm(Y ~ X_0 + Z_1, data = lgdat)
summary(mx0)
confint(mx0)</pre>
```

And similarly, we check that *not* adjusting for Z_1 can bias our estimate of X_1 .

```
mx1 <- glm(Y ~ X_1, data = lgdat)
summary(mx1)
confint(mx1)</pre>
```

Then, we apply standardization to the data. The logic is the same as discussed above, only now we have two time points.

So, we first fit a model for the treatment at time point 0 and then compute the contrast in expectations between receiving and not receiving the treatment, $E[Y^{X_0=1,Z_0=z_0}] - E[Y^{X_0=0,Z_0=z_0}]$. Next, we fit a model for receiving treatment at time point 1 and similarly compute the contrast in expectations between receiving and not receiving the treatment, $E[Y^{X_1=1,X_0=x_0,Z_1=z_1}] - E[Y^{X_1=0,X_0=x_0,Z_1=z_1}]$.

```
# Outcome model for X_O
yXOmodel \leftarrow brm(Y \sim X_0 + Z_0,
                 data = lgdat, cores = 4)
\# E[Y\{X \ O=1, \ Z \ O=z \ O\}]
yX0_1 <- posterior_epred(yX0model, newdata = transform(lgdat, X_0=1))
\# E[Y\{X_0=0, Z_0=z_0\}]
yX0_0 <- posterior_epred(yX0model, newdata = transform(lgdat, X_0=0))
# Contrast: E[Y{X O=1}] - E[Y{X O=0}]
yX0contrast <- mean_hdci((yX0_1[,1] - yX0_0[,1]))</pre>
# Cutcome model for X_1
yX1model \leftarrow brm(Y \sim X_0 + X_1 + Z_1,
                 data = lgdat, cores = 4)
\# E[Y{X_1 = 1, X_0=x_0, Z_1=z_1}]
yX1_1 <- posterior_epred(yX1model, newdata = transform(lgdat, X_0=1))</pre>
\# E[Y{X_1=0, X_0=x_0, Z_1=z_1}]
yX1_0 <- posterior_epred(yX1model, newdata = transform(lgdat, X_0=0))
# Contrast: E[Y{X 1=1}] - E[Y{X 1=0}]
yX1contrast <- mean_hdci((yX1_1[,1] - yX1_0[,1]))</pre>
```

By breaking the procedure down into separate models – one for each time point and treatment – we by-pass the problem that a single regression runs into, namely that Z_1 is both a collider and a confounder. The resulting contrasts, stored in yX0contrast and yX1contrast, are valid causal estimates of the joint effects of each treatment on the outcome, to the extent assumptions hold (we discuss these assumptions in the next section).

Now for the finale. The above example comes from Hernan and Robins (2020), and it's useful in that it illustrates how regression modeling alone breaks if our estimands require different adjustment sets (e.g., if a variable is both a confounder that we want to adjust for and a collider that we want to keep unadjusted).

But let's take standardization for a spin in a real-world data analysis example. VanderWeele, Jackson, and Li (2016) sets the stage for our example: religion and (mental) health.

The DAG could look something like figure 4.

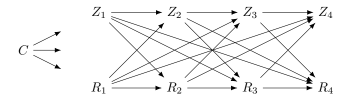


Figure 4: DAG of longitudinal exposure-outcome feedback.

C is a set of baseline covariates comprising standard demographics, including age, gender, income and education; the indefinite arrows indicate confounding on all observed variables). Our exposure R_{1-4} represent level of religiosity ("How important is religion to you?", measured on a 1 ("not at all important") to 4 ("very important") scale.) measured at four time points, and Z_{1-4} are subjective general health ("How would you describe your current health?", measured on a 1 ("very bad") to 5 ("very good") scale.), a set of time-varying covariates. Note the complex causal relationships across time between R and Z.

To provide some background, this dataset comes from a panel study on a pseudo-representative sample of Danes during the COVID-19 pandemic (for more detail, see Mauritsen et al., *in review*). As investigators, we are interested in assessing whether religiosity was a protective factor in terms of subjective general well-being in the context of the stress and uncertainty of a pandemic. If that was so, we'd expect a positive effect of religiosity on subjective health.

Suppose, then, that our estimand of interest is the contrast in subjective well-being at the final measurement time (that is, $Y = Z_4$) between being "maximally religious" $Y^{R_1 = R_2 = R_3 = 4}$ and being "minimally religious" $Y^{R_1 = R_2 = R_3 = 4}$. That is, we're interested in the *joint effects* of being religious on health at all time points up to the final measurement, meaning partitioning the respective effects of religiosity level at each time point on the outcome, except those running through the subsequent measurements of religiosity.

Note that we could've made the causal diagram even more involved by hypothesizing that our two time-varying variables also caused future versions of themselves beyond the immediately subsequent time point (e.g., adding paths $R_1 \to R_3$, $R_1 \to R_4$, $Z_1 \to Z_3$, $Z_1 \to Z_4$, etc.).

However, even as it stands, this analysis is not straightforward. Consider for instance that some of the effect of R_1 on the outcome works through Z_2 (and Z_3) and that if we adjust for Z_2 (or Z_3), we'll block some of the effect of R_1 . But note too that Z_2 (through Z_3) confounds the relationship between R_3 and the outcome, such that if we do not adjust for Z_2 (or Z_3), the estimate for the effect of R_3 will be biased.

All this implies that Z_2 is *both* a mediator that we'd want to keep unadjusted *and* a confounder that we'd want to adjust for. A single regression cannot handle this situation. But, as we've already seen, g-methods can.

First, we load the data, packages, and set the path to cmdstan, which helps speed up the sampling procedure. We also set the number of iterations to use in sampling. For now, we use only the complete cases (in all variables but for R_4 , which we're not actually using at this moment); below, we'll discuss in more detail what a complete cases analysis implies and ways of handling missing covariate values when using standardisation.

```
veluxdat <- read.csv("velux_data.csv")

d <- veluxdat[complete.cases(veluxdat[, !names(veluxdat) %in% c("religion_4")]),]</pre>
```

```
library(cmdstanr)
library(brms)
library(tidybayes)
set_cmdstan_path("C:\\cmdstan\\cmdstan-2.29.0")
iter <- 2000</pre>
```

Then, we fit the model for R_1 on the outcome Z_4 . We're interested in estimating all paths from $R_1 \to Z_4$ except those through subsequent religiosity measurements, while controlling for the demographic confounders. This means that, in addition to the baseline covariates and our main exposure R_1 , we only need to include R_2 in the model. Adjusting for R_2 blocks the paths from R_1 to the outcome through subsequent measures of religion. Had we assumed the path $R_1 \to R_3$, we'd have to include R_3 , too, to block the path $R_1 \to R_3 \to Z_4$.

Next, we fit a model for R_2 . In addition to the main exposure and the demographic covariates, we need to adjust for both R_1 (because it's a confounder on the path $R_2 \leftarrow R_1 \rightarrow Z_4$), R_3 (to block the path $R_2 \rightarrow R_3 \rightarrow Z_4$) and Z_1 (because it's a confounder on the path $R_2 \leftarrow Z_1 \rightarrow Z_2 \rightarrow Z_3 \rightarrow Z_4$).

Finally, the model for R_3 adjusts for the demographic covariates as well as R_2 (because it's a confounder on

the path $R_3 \leftarrow R_2 \rightarrow Z_4$) and Z_2 (because it's a confounder on the path $R_3 \leftarrow Z_2 \rightarrow Z_4$ through Z_3). Notice that, had we assumed a causal relationship $R_4 \rightarrow Z_4$, R_4 would have to be included, too.

The objects yr1contrast, yr2contrast and yr3contrast store the posterior means and HPDIs of the joint effects of religiosity on subjective health. You can inspect the contrasts to see that they huddle mostly around zero, meaning that we find little evidence for the notion that religion on average works as a protective factor during the pandemic, at least in these data.

One last comment, before we move on: In our estimation above, we silently cut some corners, for the sake of illustrating the basic principles of standardisation in a context with time-varying confounders. There are some additional modeling complexities that we could (and perhaps should) incorporate: Monotonicity in predictors; ordinal (not gaussian) outcome; random intercepts for individuals; Bayesian imputation.

In the next section, we discuss assumptions that are required for a causal interpretation of estimates obtained via standardization and IPTW.

4.4 It's assumptions all the way down

Causal inference in an observational setting generally requires several key conditions for a causal interpretation of the main exposure (e.g. Hernan and Robins 2020, ch. 13; Naimi, Cole, and Kennedy 2017). One way to think of this exercise is that when assumptions are (assumed) satisfied, an observational study will have emulated a randomized study.

First, we assume that the potential outcomes under varying levels of exposure are independent from the observed outcomes (conditional exchangeability). In a perfectly randomized trial, this is the case since randomization ensures that the probability of treatment is independent of the outcome — we say that the treatment and control groups are "exchangeable". However, in an observational setting, there can be countless factors that both influence exposure levels and the outcome. It's a main goal of a statistical model to adjust for these confounding factors, in order to obtain conditional exchangeability. For instance, as we've seen, a DAG is useful for guiding statistical adjustment. In essence, we aim to statistically block all backdoor paths that both influence our exposure M and outcome Y of interest.

Second, and relatedly, we assume *no model misspecification*, which entails that our model is specified correctly (e.g., in terms of functional relationships, no omitted confounding variables, etc.). However, whether any given statistical model and adjustment sets are sufficient to ensure these conditions hold is generally not empirically testable.

Third, and similarly, we assume that our variables are *measured without error*, another difficult-to-verify assumption.

Fourth, we assume that an individual's observed outcome under a given exposure is equivalent to the potential outcome that would've been observed under that exposure (counterfactual consistency). In other words, in the context of our g-computation procedure, when we obtain expected values for participants setting X = x, we assume that we obtain the values that we in fact would've observed, if those participants had been observed under X = x.

Fifth, we assume *positivity*, which implies that individuals have similar exposure levels within all confounder levels. While this is empirically unlikely to hold in particular (for instance, when we have several covariates, including continuous ones, several groups, etc.), lack of positivity can be ignored to the extent that we're willing to assume that estimates for the strata with zero observations can be extrapolated from the model fitted on the observed strata (Hernan and Robins 2020, 162).

Sixth, and more trivially, we assume temporal ordering of relevant variables, for instance such that exposures and mediating variables occur before an outcome.

5 To do:

- double check notation throughout within-document consistency and external consistency with e.g.,
 What if? book
- add references
- a little bit more formal detail on the IPTW weights and the difference btw. unstabilized and stabilized weights and why we can't rely on the standard errors (and therefore must resort to bootstrapping)
- Missing data MCAR, MAR, MNAR. See Morris et al. (2022, appendix) for discussion of missingness in X and/or Y in the context of g-comp.

Above, we conducted a complete cases analysis, under the assumption that missingness is unsystematic. One way to investigate this assumption a little bit further is to ask, does health predict dropout at the final measurement? This is not a bulletproof "test" by any means, but it's a start.

See end of Standardization chapter in What if?

• Econometric techniques

Regression discontinuity, instrumental variables, synthetic controls

• Threats to identification and some solutions

Attrition (simple MNAR analysis), selection bias, spill-over, non-compliance (IV of randomization), etc.

- Transportability of treatment effects (see What if? chapter); also extends naturally from standardisation/g-computation.
- Red herrings Testing for covariate imbalances
- Dictionary for common terminology in the causal inference literature

D-separation, Do-calculus,

• Principles for randomized trials/experiments

6 Appendix: Velux IPTW

I said above that we'd leave IPTW behind., Well, not quite. Here's how to use IPTWs and a MSM to analyze the panel data on health and religiosity.

Note that we use only the complete cases. Also, this illustration differs from the IPTW above, because now we have a continuous, not a binary, exposure (well, it's categorical, but we assume gaussian for the sake of simplicity). The IPTWs are calculated slightly differently, when the exposure is continuous. All this said, the results are qualitatively (although not numerically, likely due to difference in estimation and also different sample subsets) similar to our standardization approach, in that we find no noteworthy effect of religiosity on health.

However, try and run the bootstrap and subsequent steps without the weights (i.e., delete the weights argument from $iptw_fun()$); you'll see that the model then picks up a "near-statistically significant" negative association between R_3 and the outcome, which we can only guess is spurious.

```
diptw <- d[complete.cases(d),] # complete cases</pre>
## Computing IPTW with a continuous exposure: https://www.andrewheiss.com/blog/2020/12/01/ipw-binary-co.
## the stabilized weights for religion_t1 is 1 (since the numerator and the denominator is the same)
## cf., VanderWeel et al. (online appendix)
## numerator model for religion_t2
r2exppn <- glm(religion_2 ~ gender + age.c + education + household_income + religion_1, data = diptw)
diptw$pn_r2 <- dnorm(diptw$religion_2,</pre>
             predict(r2exppn),
             sd(r2exppn$residuals))
## denominator model for religion t2
r2exppd <- glm(religion_2 ~ gender + age.c + education + household_income + religion_1 + health_1, data
diptw$pd_r2 <- dnorm(diptw$religion_2,</pre>
             predict(r2exppd),
             sd(r2exppd$residuals))
## numerator model for religion_t3
r3exppn <- glm(religion_3 ~ gender + age.c + education + household_income + religion_2, data = diptw)
diptw$pn_r3 <- dnorm(diptw$religion_3,</pre>
             predict(r3exppn),
             sd(r3exppn$residuals))
## denominator model for religion_t3
r3exppd <- glm(religion_3 ~ gender + age.c + education + household_income + religion_2 + health_1 + hea
diptw$pd_r3 <- dnorm(diptw$religion_3,</pre>
             predict(r3exppd),
             sd(r3exppd$residuals))
## calculate weights
diptw$sw2 <- with(diptw, pn_r2/pd_r2)</pre>
diptw$sw3 <- with(diptw, pn_r3/pd_r3)</pre>
diptw$sw <- with(diptw, sw2*sw3)</pre>
# MSM without and with stabilized weights
```

```
velux_uw <- glm(health_4 ~ gender + age.c + education + household_income + religion_1 + religion_2 + re</pre>
summary(velux_uw)
velux_msm <- glm(health_4 ~ gender + age.c + education + household_income + religion_1 + religion_2 + r</pre>
summary(velux_msm)
### bootstrapping, when weights are calculated
library(boot)
# same function as above
iptw_fun <- function(formula, data, indices) {</pre>
 d <- data[indices,]</pre>
 fit <- glm(formula, family="gaussian", weights = sw, data=d)</pre>
 return(coef(fit))
iptw.velux.result <- boot(data = diptw,</pre>
                 statistic = iptw_fun,
                  R = 1e4,
                  formula = health_4 ~ gender + age.c + education + household_income + religion_1 + reli
# bootstrapped point estimate for religion_t1
iptw.velux.r1point <- iptw.velux.result$t0[6]</pre>
# bootstrapped point estimate for religion_t2
iptw.velux.r2point <- iptw.velux.result$t0[7]</pre>
# bootstrapped point estimate for religion_t3
iptw.velux.r3point <- iptw.velux.result$t0[8]</pre>
# bootstrapped interval for religion_t1
iptw.velux.r1interval <- boot.ci(iptw.velux.result,</pre>
                          type = "norm",
                          index = 6)$normal
# bootstrapped interval for religion_t2
iptw.velux.r2interval <- boot.ci(iptw.velux.result,</pre>
                          type = "norm",
                          index = 7)$normal
# bootstrapped interval for religion_t3
iptw.velux.r3interval <- boot.ci(iptw.velux.result,</pre>
                          type = "norm",
                          index = 8)$normal
```

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