Scenario 1: Absence of treatment effect heterogeneity (HTE)

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Generated: November 18, 2022

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1 Build Info

This report was generated using R version 4.2.2 (2022-10-31 ucrt). The following key packages were used:

- pandoc (version 2.19.2): Report output
- MatchIt (version 4.4.0): Nearest-neighbour matching
- sandwich (version 3.0.2): Estimation of (cluster-)robust standard errors
- dplyr (version 1.0.10): Facilitate data manipulation
- tibble (version 3.1.8): Facilitate data manipulation
- ggplot2 (version 3.3.6): Visualization of results
- mice (version 3.14.0): Multivariate Imputation by Chained Equations
- mvtnorm (version 1.1.3): Draw x1 and 1x from bivaraite normal distribution
- ggmice (version 0.0.1): Visualize predictor matrix of imputations

2 Methods

2.1 Data-generating mechanism

We generated data for a super-population of N=100,000 individuals treated with treatment A, B, or C. We generated two continuous variables x_1 and x_2 for each patient as baseline covariates, labeled as age (standardized) and disease severity (z-score). We generated a potential outcome, labeled as a test score for which higher values are better, under each of the three treatments. The test scores were generated from a linear model with age, disease severity, and the treatment indicators as main effects and an interaction term between the treatment indicators and age to introduce treatment effect heterogeneity. The outcome y is continuous, and generated as follows:

$$y_i \sim N(\mu_i, \sigma^2)$$

with

$$\mu_i = \beta_0 + \beta_1 x_{1i} + \beta_2 x_{2i} + \beta_3 I_{Bi} + \beta_4 I_{Ci} + (\beta_5 I_{Bi} + \beta_6 I_{Ci}) x_{1i}$$

Using the following regression coefficients:

- Baseline outcome risk (β_0) : **0**
- Confounder effect of x_1 (β_1): 1
- Confounder effect of x_2 (β_2): **0.25**
- Treatment effect of B versus A (β_3) : -0.5
- Treatment effect of C versus A (β_4) : -0.25
- Standard deviation of the residual error (σ): 1

Depending on the selected scenario, the values for β_5 and β_6 were modified.

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- Effect modification between B and x1 (β_5): **0**
- Effect modification between C and x1 (β_6): **0**

We assigned the treatment received as a function of the two baseline covariates x_1 and x_2 . Thus, the observable dataset consisted of the two baseline covariates, the treatment received and one outcome – the one generated under the treatment that the patient received.

2.2 Sampling distributions and parameter information

The treatment groups were sampled in equal sizes. The two continuous variables x_1 and x_2 were drawn from a bivariate normal distribution using the *mvtnorm* package, using the following mean values:

Table 1: Sampling means

Treatment	Mean age	Mean disease severity			
A	1	-1			
В	0	1			
C	-1	-1			

And the following covariance matrix:

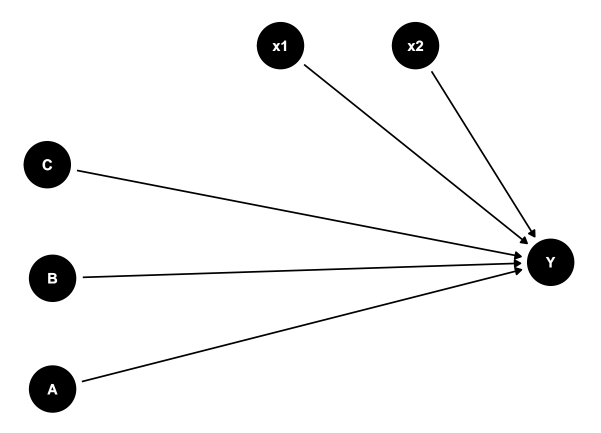
Table 2: Variance-covariance matrix of the baseline covariates

	x1	x2
	2	0
x2	0	2

2.3 Directed acyclic graph

To illustrate the relation between the treatment, the outcome and the treatment covariate interactions, a Directed acyclic graph is used. A, B, and C indicate the three treatment groups, Y is the outcome and x1 and x2 depict the covariates (age and disease severity). Both covariates influence the outcome y_i , but only in the presence of HTE do they also influence treatment B and C.

DAG in the absence of HTE



2.4 Implementation of pairwise matching

We applied 1:1 PS nearest-neighbor matching with a caliper and with replacement for each pairwise treatment comparison. We estimated the propensity score with a logistic regression model as a function of the two baseline covariates separately for each treatment comparison to mimic how pairwise matching would be repetitively applied in the context of three treatments being compared. We used a caliper of 0.05 standard deviation of the logit of the estimated PS. We assessed covariates balance pre- and post-matching with absolute standardized mean differences.

For each treatment comparison, we generated created two matched samples, separately targeting eachither of the ATTs. For example, for the comparison of treatment A vs B, we first match individuals treated with B to individuals treated with A, thus targeting the ATT-A. A first matched sample results from this matching. Second, starting from the original sample again, we now match individuals treated with A to individuals treated with B, targeting the ATT-B, which results in a second matched sample. For each treatment comparison and in each matched sample, we estimate the treatment effect (e.g., difference in means at a certain timepoint) using a linear regression model. We note that, in an application of pairwise matching to real data, researchers would typically generate only one matched sample, targeting one estimand or the other.

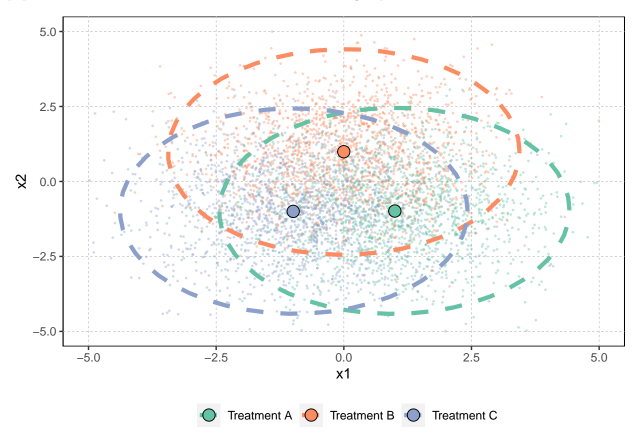
2.5 Evaluation and visualization

For the evaluation and visualization of covariate overlap, we compared the target populations before and after matching for all pairwise treatment comparisons with the bivariate ellipses. We also compared the estimated treatment effects in the matched sample.

3 Results

3.1 Scenario 1: Absence of HTE

We generate a superpopulation of 120000 patients, in which there are **40000** observations with drug A, **40000** observations with drug B and **40000** observations with drug C. Due to the presence of confounding, population characteristics differ across the three treatment groups.

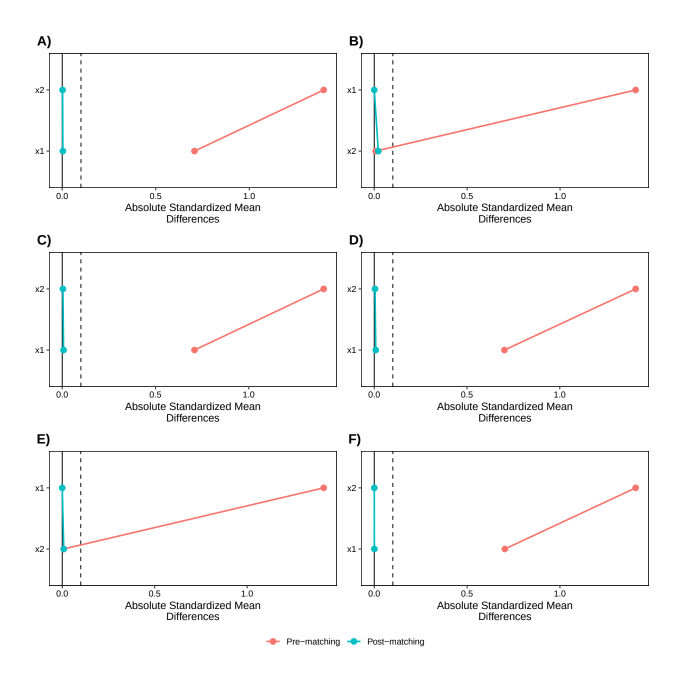


The continuous outcome y is generated from the linear predictor:

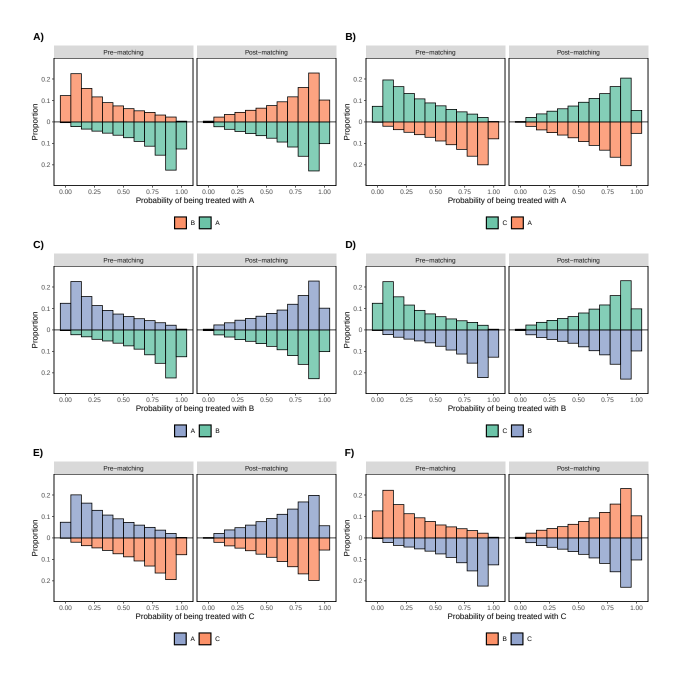
$$\mu_i = 0 + 1 x_{1i} + 0.25 x_{2i} + -0.5 I_{Bi} + -0.25 I_{Ci}$$

For all analyses, we apply nearest neighbor matching with a caliper width of 0.001 SD of the logit of the estimated PS. This results in a caliper of 0.002 for AB comparisons, a caliper of 0.002 for AC comparisons, and a caliper of 0.002 for BC comparisons. Afterwards, treatment effect estimates are derived using linear regression in the matched sample. A sandwich estimator is used to calculate cluster-robust standard errors.

3.2 Love plots

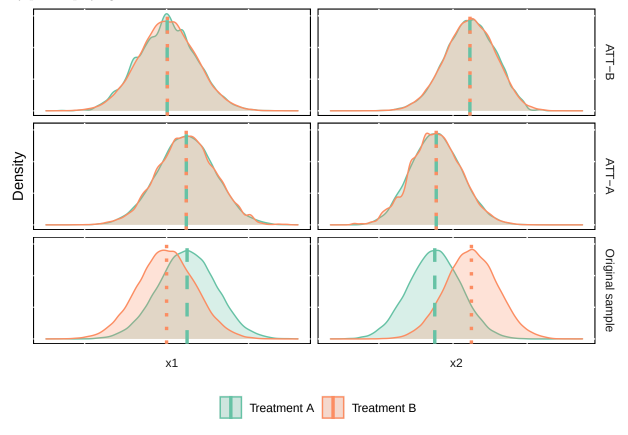


3.3 Balance plots



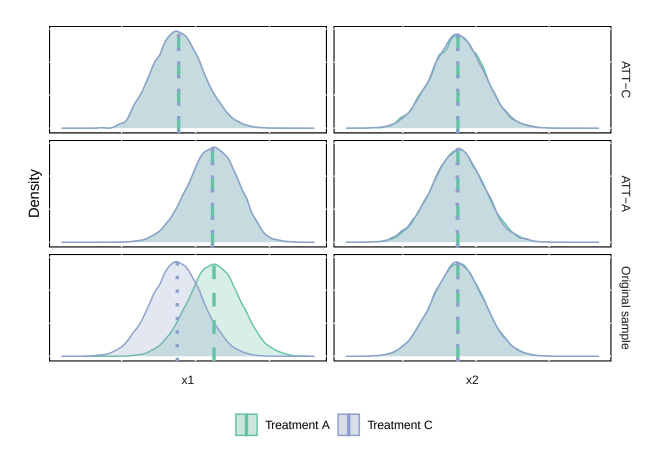
3.4 Difference between ATTA and ATTB

Joy plot displaying the differences between ATTA and ATTB



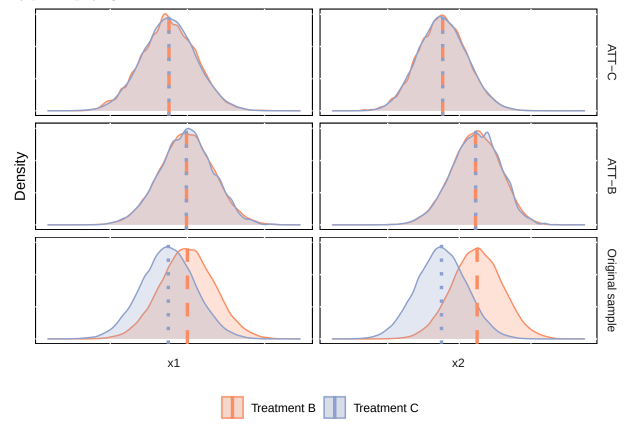
3.5 Difference between ATTA and ATTC

Joy plot displaying the differences between ATTA and ATTC



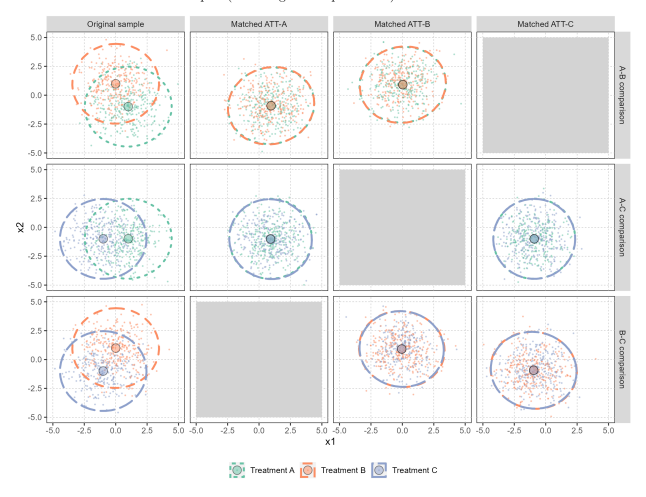
3.6 Difference between ATTB and ATTC

Joy plot displaying the differences between ATTB and ATTC



3.7 Overall distribution of covariates

Distribution of the matched samples (matching with replacement)



3.8 Treatment effect estimates

Table 3: Treatment effect estimates for Scenario 1 when adopting matching with replacement

	Comparison	Estimand	Replace	Reference	Estimate	SE	SS	ESS
x1	A versus B	ATTA	TRUE	0.50	0.5022952	0.0307740	51647	42111.28
x11	A versus B	ATTB	TRUE	0.50	0.4865111	0.0317254	51782	42363.85
x12	A versus C	ATTA	TRUE	0.25	0.2463661	0.0265947	53127	43732.07
x13	A versus C	ATTC	TRUE	0.25	0.2426073	0.0271994	53234	43911.66
x14	B versus C	ATTB	TRUE	-0.25	-0.2665614	0.0314961	51624	42238.68
x15	B versus C	ATTC	TRUE	-0.25	-0.2524470	0.0316183	51958	42334.52