Target Trial Emulation in R

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This tutorial outlines the steps for implementing a Target Trial Emulation in R using the TrialEmulation package.

The process begins by defining the estimand, such as **intention-to-treat (ITT)** or **per-protocol (PP)**, and preparing observational data with specified columns for treatment, outcomes, and eligibility.

Censoring mechanisms, including **treatment switching** and **informative censoring**, are addressed using inverse probability of censoring weights (IPCW). Separate models are fitted to calculate these weights and adjust for biases.

The workflow proceeds to **expand the observational dataset** into a sequence of trials using pre-defined expansion options, creating trial-specific datasets. These datasets are then loaded and sampled for analysis.

A marginal structural model (MSM) is fit to estimate the causal effects. Predictions, such as survival probabilities and cumulative incidences, are generated for different treatment scenarios. The results are visualized to show survival differences over time, enabling meaningful inference on the treatment effects.

1. Setup

A sequence of target trials analysis starts by specifying which estimand will be used:

```
trial_pp <- trial_sequence(estimand = "PP") # Per-protocol
trial_itt <- trial_sequence(estimand = "ITT") # Intention-to-treat</pre>
```

Additionally it is useful to create a directory to save files for later inspection.

```
trial_pp_dir <- file.path(tempdir(), "trial_pp")
dir.create(trial_pp_dir)

trial_itt_dir <- file.path(tempdir(), "trial_itt")
dir.create(trial_itt_dir)</pre>
```

2. Data preparation

Next the user must specify the observational input data that will be used for the target trial emulation. Here we need to specify which columns contain which values and how they should be used.

```
data("data_censored") # dummy data in the package
head(data_censored)
```

```
##
    id period treatment x1
                                    x2 x3
                                                x4 age
                                                            age_s outcome
## 1 1
            0
                      1 1 1.146148362 0 0.734203 36 0.08333333
## 2 1
            1
                           0.002200337 0 0.734203 37 0.16666667
                                                                       0
            2
                      1 0 -0.481762418 0 0.734203 38 0.25000000
## 3 1
                                                                       0
            3
                      1 0 0.007872396 0 0.734203 39 0.33333333
## 4 1
                                                                       0
## 5 1
            4
                      1 1 0.216053715 0 0.734203 40 0.41666667
                                                                       0
            5
                      1 0 -0.057481504 0 0.734203 41 0.50000000
## 6 1
                                                                       0
##
    censored eligible
## 1
           0
## 2
           0
                    0
## 3
                    0
## 4
           0
                    0
## 5
           0
                    0
## 6
           1
                    0
```

```
# Per-protocol
trial_pp <- trial_pp |>
  set_data(
    data
             = data_censored,
   id
             = "id",
   period
           = "period",
   treatment = "treatment",
    outcome
            = "outcome",
    eligible = "eligible"
  )
# ITT
# Function style without pipes
trial_itt <- set_data(</pre>
 trial_itt,
 data
            = data_censored,
 id
            = "id",
         = "period",
 period
 treatment = "treatment",
 outcome = "outcome",
 eligible = "eligible"
)
```

```
trial_itt
```

```
## Trial Sequence Object
## Estimand: Intention-to-treat
##
## Data:
##
    - N: 725 observations from 89 patients
##
            id period treatment
                                     x1
                                                   x2
                                                         х3
                                                                    x4
                                                                          age
                                                                                    age_s
##
         <int>
                <int>
                           <num> <num>
                                                <num> <int>
                                                                 <num> <num>
                                                                                    <num>
                               1
                                                                           36 0.08333333
##
     1:
                                         1.146148362
                                                           0 0.7342030
##
     2:
             1
                    1
                               1
                                         0.002200337
                                                           0 0.7342030
                                                                           37 0.16666667
##
   724:
            99
                    6
                               1
                                      1 -0.033762356
                                                           1 0.5752681
                                                                           71 3.00000000
##
##
   725:
            99
                    7
                               0
                                      0 -1.340496520
                                                           1 0.5752681
                                                                           72 3.08333333
##
        outcome censored eligible time_on_regime
##
           <num>
                    <int>
                              <num>
               0
                         0
                                                   0
##
     1:
##
     2:
               0
                         0
                                  0
                                                   1
##
##
   724:
               0
                         0
                                  0
                                                   1
##
   725:
               1
                         0
                                  0
                                                   2
##
   IPW for informative censoring:
##
    - No weight model specified
##
## Sequence of Trials Data:
   - Use set_expansion_options() and expand_trials() to construct the sequence of trials datase
t.
##
## Outcome model:

    Outcome model not specified. Use set_outcome_model()
```

3. Weight models and censoring

To adjust for the effects of informative censoring, inverse probability of censoring weights (IPCW) can be applied. To estimate these weights, we construct time-to-(censoring) event models. Two sets of models are fit for the two censoring mechanisms which may apply: censoring due to deviation from assigned treatment and other informative censoring.

3.1 Censoring due to treatment switching

We specify model formulas to be used for calculating the probability of receiving treatment in the current period. Separate models are fitted for patients who had treatment = 1 and those who had treatment = 0 in the previous period. Stabilized weights are used by fitting numerator and denominator models.

There are optional arguments to specify columns which can include/exclude observations from the treatment models. These are used in case it is not possible for a patient to deviate from a certain treatment assignment in that period.

```
trial_pp <- trial_pp |>
  set_switch_weight_model(
   numerator = ~ age,
   denominator = ~ age + x1 + x3,
   model_fitter = stats_glm_logit(save_path = file.path(trial_pp_dir, "switch_models"))
  )
  trial_pp@switch_weights
```

```
## - Numerator formula: treatment ~ age
## - Denominator formula: treatment ~ age + x1 + x3
## - Model fitter type: te_stats_glm_logit
## - Weight models not fitted. Use calculate_weights()
```

This type of censoring is not used with an ITT estimand, so we cannot use <code>set_switch_weight_model()</code> with trial ITT objects.

3.2 Other informative censoring

In case there is other informative censoring occurring in the data, we can create similar models to estimate the IPCW. These can be used with all types of estimand. We need to specify censor_event which is the column containing the censoring indicator.

```
trial_pp <- trial_pp |>
  set_censor_weight_model(
    censor_event = "censored",
    numerator = ~ x2,
    denominator = ~ x2 + x1,
    pool_models = "none",
    model_fitter = stats_glm_logit(save_path = file.path(trial_pp_dir, "switch_models"))
    )
    trial_pp@censor_weights
```

```
## - Numerator formula: 1 - censored ~ x2
## - Denominator formula: 1 - censored ~ x2 + x1
## - Model fitter type: te_stats_glm_logit
## - Weight models not fitted. Use calculate_weights()
```

```
trial_itt <- set_censor_weight_model(
    trial_itt,
    censor_event = "censored",
    numerator = ~x2,
    denominator = ~ x2 + x1,
    pool_models = "numerator",
    model_fitter = stats_glm_logit(save_path = file.path(trial_itt_dir, "switch_models"))
)
trial_itt@censor_weights</pre>
```

```
## - Numerator formula: 1 - censored ~ x2
## - Denominator formula: 1 - censored ~ x2 + x1
## - Numerator model is pooled across treatment arms. Denominator model is not pooled.
## - Model fitter type: te_stats_glm_logit
## - Weight models not fitted. Use calculate_weights()
```

4. Calculate Weights

Next we need to fit the individual models and combine them into weights. This is done with calculate_weights().

```
trial_pp <- trial_pp |> calculate_weights()
trial_itt <- calculate_weights(trial_itt)</pre>
```

The full model objects are saved to disk in the directories we created above. The summaries are stored in the trial sequence object and can be printed:

```
show_weight_models(trial_itt)
```

```
## Weight Models for Informative Censoring
##
## [[n]]
## Model: P(censor_event = 0 | X) for numerator
##
##
                estimate
                           std.error statistic p.value
   (Intercept) 2.4480907 0.1405726 17.415128 6.334656e-68
##
                -0.4486482 0.1368765 -3.277759 1.046346e-03
##
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
##
   404.2156
                  724
                          -196.7002 397.4004 406.5727 393.4004 723
                                                                           725
##
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial itt/switch models\\model 29e8558426
4c.rds
##
## [[d0]]
## Model: P(censor_event = 0 | X, previous treatment = 0) for denominator
##
##
   term
                estimate
                           std.error statistic p.value
##
   (Intercept) 1.8941961 0.2071122 9.145746 5.921948e-20
                -0.5898292 0.1693402 -3.483101 4.956409e-04
##
   x1
                 0.8552603 0.3452930 2.476912 1.325247e-02
##
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
##
   283.0723
                  425
                          -132.1655 270.3309 282.4943 264.3309 423
                                                                           426
##
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_itt/switch_models\\model_29e84d26c1
3.rds
##
## [[d1]]
## Model: P(censor_event = 0 | X, previous treatment = 1) for denominator
##
##
   term
                estimate
                            std.error statistic p.value
##
   (Intercept) 2.81443372 0.3122688 9.0128570 2.007570e-19
##
                -0.03713196 0.2699579 -0.1375472 8.905983e-01
##
   x1
                 0.89351418 0.7771954 1.1496648 2.502819e-01
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
   113.0528
                  298
                                                                           299
##
                          -55.72938 117.4588 128.5601 111.4588 296
##
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_itt/switch_models\\model_29e8d84448
7.rds
##
```

```
show_weight_models(trial_pp)
```

```
## Weight Models for Informative Censoring
##
## [[n0]]
## Model: P(censor_event = 0 | X, previous treatment = 0) for numerator
##
##
                estimate
                           std.error statistic p.value
   (Intercept) 1.4026538 0.1993673 7.035525 1.985118e-12
##
                -0.5436594 0.2075654 -2.619220 8.813109e-03
##
##
   null.deviance df.null logLik
                                   AIC
                                            BIC
                                                     deviance df.residual nobs
##
                          -82.81349 169.627 175.8986 165.627 168
##
   172.8729
                  169
                                                                          170
##
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e87432689
d.rds
##
## [[n1]]
## Model: P(censor_event = 0 | X, previous treatment = 1) for numerator
##
##
   term
                estimate
                           std.error statistic p.value
##
   (Intercept) 2.7365126 0.3465786 7.8957919 2.884776e-15
                -0.1259467 0.3527451 -0.3570472 7.210564e-01
##
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
                          -34.04272 72.08543 78.11999 68.08543 149
##
   68.21358
                  150
                                                                           151
##
##
   path
   C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e8484d51d
9.rds
##
## [[d0]]
## Model: P(censor_event = 0 | X, previous treatment = 0) for denominator
##
##
   term
                estimate
                           std.error statistic p.value
##
   (Intercept) 1.0337903 0.2449150 4.221017 2.432028e-05
##
   x2
                -0.6189556 0.2153087 -2.874735 4.043662e-03
##
   x1
                0.9453986 0.4223668 2.238335 2.519919e-02
##
##
   null.deviance df.null logLik
                                   AIC
                                             BIC
                                                     deviance df.residual nobs
##
   172.8729
                  169
                          -80.12832 166.2566 175.664 160.2566 167
                                                                          170
##
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e85d0f1ec
e.rds
##
## [[d1]]
## Model: P(censor_event = 0 | X, previous treatment = 1) for denominator
##
##
   term
                estimate
                           std.error statistic p.value
    (Intercept) 2.5496142 0.3705515 6.8805941 5.960344e-12
##
##
                -0.1571042 0.3463174 -0.4536422 6.500864e-01
```

```
##
                1.0350346 1.0806645 0.9577761 3.381757e-01
   х1
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
   68.21358
                  150
                        -33.46038 72.92075 81.97259 66.92075 148
##
                                                                           151
##
##
   path
   C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e85e9c428
4.rds
##
## Weight Models for Treatment Switching
##
## [[n1]]
## Model: P(treatment = 1 | previous treatment = 1) for numerator
##
   term
                estimate
                           std.error statistic p.value
   (Intercept) 1.80162178 0.77463133 2.325780 0.02003031
##
                -0.02351116 0.01691961 -1.389581 0.16465623
##
   age
##
                                             BIC
##
   null.deviance df.null logLik
                                   AIC
                                                      deviance df.residual nobs
                  150
   188.829
                         -93.43779 190.8756 196.9101 186.8756 149
                                                                           151
##
##
## path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e8179e34b
7.rds
##
## [[d1]]
## Model: P(treatment = 1 | previous treatment = 1) for denominator
##
##
   term
                estimate
                            std.error statistic p.value
   (Intercept) 1.55485166 0.81706997 1.9029602 0.05704573
##
               -0.02312027 0.01696843 -1.3625460 0.17302562
##
   age
                0.52915871 0.43594855 1.2138100 0.22482028
##
   х1
##
   х3
                0.21319587 0.35744378 0.5964459 0.55087740
##
##
   null.deviance df.null logLik
                                   AIC
                                             BIC
                                                      deviance df.residual nobs
   188.829
                  150
                          -92.54787 193.0957 205.1649 185.0957 147
                                                                           151
##
##
##
   path
   C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e8365f2ff
h.rds
##
## [[n0]]
## Model: P(treatment = 1 | previous treatment = 0) for numerator
##
##
   term
                estimate
                            std.error statistic p.value
   (Intercept) 1.09212298 0.60582678 1.802698 0.07143559
##
##
   age
                -0.03360404 0.01439482 -2.334453 0.01957201
##
##
   null.deviance df.null logLik
                                    AIC
                                             BIC
                                                      deviance df.residual nobs
##
   232.2705
                  169
                      -113.2746 230.5492 236.8208 226.5492 168
                                                                           170
##
```

```
##
   path
## C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e86b1816e
##
## [[d0]]
  Model: P(treatment = 1 | previous treatment = 0) for denominator
##
##
   term
                estimate
                            std.error statistic p.value
##
    (Intercept) 1.03084683 0.63052714 1.6348969 0.10207067
                -0.03633255 0.01472576 -2.4672779 0.01361446
##
   x1
                 0.64473751 0.32346837 1.9932011 0.04623943
##
   х3
                -0.23411026 0.32147930 -0.7282281 0.46647397
##
   null.deviance df.null logLik AIC
                                         BIC
                                                  deviance df.residual nobs
##
   232,2705
                  169
                          -111.03 230.06 242.6032 222.06
                                                                        170
##
##
##
   path
   C:\\Users\\14379\\AppData\\Local\\Temp\\RtmpoDhrya/trial_pp/switch_models\\model_29e83d5c48d
##
d.rds
##
```

5. Specify Outcome Model

Now we can specify the outcome model. Here we can include adjustment terms for any variables in the dataset. The numerator terms from the stabilised weight models are automatically included in the outcome model formula.

```
trial_pp <- set_outcome_model(trial_pp)
trial_itt <- set_outcome_model(trial_itt, adjustment_terms = ~x2)</pre>
```

6. Expand Trials

Now we are ready to create the data set with all of the sequence of target trials.

```
trial_pp <- set_expansion_options(
    trial_pp,
    output = save_to_datatable(),
    chunk_size = 500 # the number of patients to include in each expansion iteration
)
trial_itt <- set_expansion_options(
    trial_itt,
    output = save_to_datatable(),
    chunk_size = 500
)</pre>
```

6.1 Create Sequence of Trials Data

```
trial_pp <- expand_trials(trial_pp)
trial_itt <- expand_trials(trial_itt)</pre>
```

```
trial_pp@expansion
```

```
## Sequence of Trials Data:
## - Chunk size: 500
## - Censor at switch: TRUE
## - First period: 0 | Last period: Inf
##
## A TE Datastore Datatable object
## N: 500 observations
##
           id trial period followup time outcome
                                                      weight treatment
                                                                                 x2
##
        <int>
                      <int>
                                     <int>
                                             <num>
                                                        <num>
                                                                              <num>
                                                                   <num>
##
                                         0
                                                 0 1.0000000
                                                                       1 1.1461484
     1:
     2:
##
                                                 0 0.8951447
                                                                       1 1.1461484
##
## 499:
                                                                       1 -0.3463778
           99
                          0
                                                 0 1.0000000
   500:
           99
                                                 0 1.0122336
                                                                       1 -0.3463778
##
          age assigned_treatment
##
##
        <num>
##
           36
                                1
     1:
##
     2:
           36
                                1
##
## 499:
                                1
## 500:
           65
                                1
```

7. Load or Sample from Expanded Data

Now that the expanded data has been created, we can prepare the data to fit the outcome model. For data that can fit comfortably in memory, this is a trivial step using load expanded data.

For large datasets, it may be necessary to sample from the expanded by setting the $p_control$ argument. This sets the probability that an observation with outcome == 0 will be included in the loaded data. A seed can be set for reproducibility. Additionally, a vector of periods to include can be specified, e.g., period = 1:60, and/or a subsetting condition, $subset_condition = "age > 65"$.

```
trial_itt <- load_expanded_data(trial_itt, seed = 1234, p_control = 0.5)</pre>
```

8. Fit Marginal Structural Model

To fit the outcome model we use fit_msm().

Model summary:

```
trial itt@outcome model
```

```
## - Formula: outcome ~ assigned treatment + x2 + followup time + I(followup time^2) + trial per
iod + I(trial_period^2)
## - Treatment variable: assigned_treatment
## - Adjustment variables: x2
## - Model fitter type: te_stats_glm_logit
##
## Model Summary:
##
##
                       estimate std.error statistic p.value conf.low conf.high
   term
                       -6.02
                                0.780
                                           -7.72
                                                    1.2e-14 -7.550
                                                                      -4.4916
##
   (Intercept)
   assigned_treatment 1.63
                                0.496
                                            3.28
                                                    1.0e-03 0.654
                                                                       2.5977
##
                                            0.74
##
                        0.31
                                0.418
                                                    4.6e-01 -0.511
                                                                       1.1282
##
   followup_time
                        0.34
                                0.244
                                            1.38
                                                    1.7e-01 -0.141
                                                                       0.8148
   I(followup_time^2) -0.02
                                0.014
                                           -1.42
                                                    1.5e-01 -0.049
##
                                                                       0.0077
   trial_period
                                            7.45
                                                    9.1e-14 5.371
##
                        7.29
                                0.978
                                                                       9.2040
##
   I(trial_period^2) -7.68
                                0.537
                                           -14.31
                                                    1.8e-46 -8.737
                                                                      -6.6325
##
##
   null.deviance df.null logLik AIC BIC deviance df.residual nobs
##
   158
                  800
                          -69.1 152 185 135
                                                  794
                                                               801
```

Depending on the model fitter used, we can also access the model object. For the default stats::glm logistic model, we have the glm object as well as the sandwich variance-covariance matrix.

```
trial_itt@outcome_model@fitted@model$model
```

```
##
## Call: glm(formula = formula, family = binomial("logit"), data = data,
       weights = weights, x = FALSE, y = FALSE)
##
##
## Coefficients:
##
          (Intercept) assigned_treatment
                                                                      followup time
                                                             x2
             -6.02067
                                                                            0.33673
##
                                   1.62585
                                                       0.30837
## I(followup time^2)
                             trial period
                                             I(trial period^2)
##
             -0.02049
                                   7.28762
                                                      -7.68478
##
## Degrees of Freedom: 800 Total (i.e. Null); 794 Residual
## Null Deviance:
                        157.8
## Residual Deviance: 134.7
                                 AIC: 152.2
```

trial_itt@outcome_model@fitted@model\$vcov

```
##
                       (Intercept) assigned_treatment
                                                                x2 followup_time
## (Intercept)
                       0.608651263
                                         -0.007606479
                                                       0.042942422 -0.143451214
## assigned_treatment -0.007606479
                                          0.245882729
                                                      0.087953406 -0.052364376
## x2
                       0.042942422
                                          0.087953406 0.174977954 -0.045052691
## followup time
                                         -0.052364376 -0.045052691
                      -0.143451214
                                                                     0.059487800
## I(followup time^2) 0.007130666
                                          0.002815736 0.002843807 -0.003362158
## trial_period
                      -0.105885453
                                         -0.341609248 -0.097440741
                                                                   0.104454026
## I(trial_period^2)
                                          0.165009684 0.046219048 -0.054969078
                       0.049055893
                      I(followup_time^2) trial_period I(trial_period^2)
##
## (Intercept)
                            0.0071306658 -0.10588545
                                                             0.04905589
## assigned_treatment
                            0.0028157357 -0.34160925
                                                             0.16500968
## x2
                            0.0028438066 -0.09744074
                                                             0.04621905
## followup_time
                           -0.0033621580
                                           0.10445403
                                                            -0.05496908
## I(followup_time^2)
                            0.0002067028 -0.00514379
                                                             0.00265172
## trial_period
                                                            -0.51334414
                           -0.0051437905
                                           0.95604134
## I(trial_period^2)
                            0.0026517200 -0.51328532
                                                             0.28822666
```

The complete object shows all the specifications:

```
trial_itt
```

```
## Trial Sequence Object
## Estimand: Intention-to-treat
##
## Data:
   - N: 725 observations from 89 patients
##
##
           id period treatment
                                   x1
                                                 x2
                                                       х3
                                                                 x4
                                                                       age
                                                                                age_s
        <int> <int>
##
                          <num> <num>
                                              <num> <int>
                                                               <num> <num>
                                                                                 <num>
##
                              1
                                    1 1.146148362
                                                        0 0.7342030
                                                                        36 0.08333333
##
     2:
            1
                   1
                              1
                                       0.002200337
                                                        0 0.7342030
                                                                        37 0.16666667
   ---
##
## 724:
           99
                   6
                              1
                                    1 -0.033762356
                                                        1 0.5752681
                                                                        71 3.00000000
## 725:
           99
                   7
                              0
                                    0 -1.340496520
                                                        1 0.5752681
                                                                        72 3.08333333
##
        outcome censored eligible time_on_regime
                                                                    wtC
                                                          wt
##
          <num>
                   <int>
                             <num>
                                                       <num>
                                                                  <num>
                        0
##
     1:
              0
                                 1
                                                 0 0.9835463 0.9835463
                                                 1 0.9429254 0.9429254
##
     2:
              0
                        0
                                 0
##
## 724:
              0
                        0
                                 0
                                                 1 0.9440988 0.9440988
## 725:
              1
                        0
                                 0
                                                 2 1.0092093 1.0092093
##
## IPW for informative censoring:
   - Numerator formula: 1 - censored ~ x2
   - Denominator formula: 1 - censored ~ x2 + x1
   - Numerator model is pooled across treatment arms. Denominator model is not pooled.
##
   Model fitter type: te_stats_glm_logit
   - View weight model summaries with show_weight_models()
##
##
## Sequence of Trials Data:
## - Chunk size: 500
## - Censor at switch: FALSE
## - First period: 0 | Last period: Inf
##
## A TE Datastore Datatable object
## N: 1558 observations
            id trial period followup_time outcome
##
                                                       weight treatment
                                                                                 x2
         <int>
                       <int>
                                     <int>
##
                                              <num>
                                                        <num>
                                                                   <num>
                                                                              <num>
             1
                                          0
                                                  0 1.0000000
##
      1:
                                                                       1 1.1461484
##
      2:
             1
                           0
                                         1
                                                  0 0.9429254
                                                                       1 1.1461484
##
## 1557:
            99
                           0
                                                  0 0.8917236
                                                                       1 -0.3463778
                                         6
## 1558:
            99
                                          7
                                                  1 0.8999358
                                                                       0 -0.3463778
##
         assigned_treatment
##
                       <num>
##
      1:
                           1
##
      2:
                           1
     ---
##
## 1557:
                           1
## 1558:
                           1
##
## Outcome model:
## - Formula: outcome ~ assigned_treatment + x2 + followup_time + I(followup_time^2) + trial_per
iod + I(trial_period^2)
```

```
## - Treatment variable: assigned_treatment
## - Adjustment variables: x2
   - Model fitter type: te_stats_glm_logit
##
## Model Summary:
##
##
                        estimate std.error statistic p.value conf.low conf.high
    term
    (Intercept)
                        -6.02
                                 0.780
                                             -7.72
                                                      1.2e-14 -7.550
                                                                        -4.4916
##
                                 0.496
                                              3.28
                                                      1.0e-03 0.654
##
    assigned_treatment 1.63
                                                                         2.5977
                                              0.74
                                                      4.6e-01 -0.511
##
                         0.31
                                 0.418
                                                                         1.1282
                                              1.38
##
    followup_time
                         0.34
                                 0.244
                                                      1.7e-01 -0.141
                                                                         0.8148
    I(followup_time^2) -0.02
                                             -1.42
                                                      1.5e-01 -0.049
                                                                         0.0077
##
                                 0.014
##
    trial period
                         7.29
                                 0.978
                                              7.45
                                                      9.1e-14 5.371
                                                                         9.2040
    I(trial period^2) -7.68
                                            -14.31
                                                      1.8e-46 -8.737
                                 0.537
                                                                        -6.6325
##
##
##
    null.deviance df.null logLik AIC BIC deviance df.residual nobs
##
    158
                  800
                           -69.1 152 185 135
                                                    794
                                                                 801
##
## Outcome data
## N: 801 observations from 76 patients in 18 trial periods
  Periods: 0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17
   Sampling control observations with probability: 0.5
##
           id trial_period followup_time outcome
                                                     weight treatment
                                                                               x2
        <int>
                      <int>
                                    <int>
##
                                             <num>
                                                      <num>
                                                                 <num>
                                                                             <num>
                          0
                                         0
                                                 1 1.000000
                                                                     1 -0.7365256
##
     1:
           15
##
     2:
           32
                          0
                                         0
                                                 1 1.000000
                                                                       1.9861380
##
## 800:
                          0
           39
                                        19
                                                 0 1.351756
                                                                     1 0.2189413
                          0
                                                                       1.2924128
##
   801:
           54
                                        19
                                                 0 1.359294
##
        assigned_treatment sample_weight
##
                      <num>
                                              <num>
##
     1:
                          1
                                         1 1.000000
##
     2:
                          1
                                         1 1.000000
##
                                         2 2.703512
## 800:
                          1
                          0
                                         2 2.718587
## 801:
```

9. Inference

We use the <code>predict()</code> method to estimate survival probabilities or cumulative incidences for different values of <code>assigned_treatment</code>.

```
preds <- predict(
    trial_itt,
    newdata = outcome_data(trial_itt)[trial_period == 1, ],
    predict_times = 0:10,
    type = "survival",
)

plot(preds$difference$followup_time, preds$difference$survival_diff,
    type = "l", xlab = "Follow up", ylab = "Survival difference")
lines(preds$difference$followup_time, preds$difference$`2.5%`, type = "l", col = "red", lty = 2)
lines(preds$difference$followup_time, preds$difference$`97.5%`, type = "l", col = "red", lty = 2)</pre>
```

