Event time prediction with tidymodels

2023-04-13

The censored package was released in June 2022, enabling users to fit event time/survival time models using the tidymodels framework. As of this writing, there are now a total of 11 different engines that can be used with 6 different models.

This document is intended as a tutorial for using the broader tidymodels framework for event time analysis, including model tuning, evaluation, and selection.

To reproduce these results, you might need to update some package versions:

```
# Get CRAN versions of
pak::pak(c("parsnip", "censored"), ask = FALSE)

# Get GitHub versions of:
pak::pak(c("tidymodels/tune@ipcw"), ask = FALSE)
pak::pak(c("tidymodels/yardstick"), ask = FALSE)
```

An Example

We'll use the heart valve data set in the joineR package (also described in this publication). There are 256 patients in the study that experienced aortic valve replacement surgery. The data has time-dependent covariates, but we will skip those to simplify the analysis here. The outcome is the time to death after surgery:

```
library(joineR)
data(heart.valve, package = "joineR")
str(heart.valve)
```

```
## 'data.frame':
                  988 obs. of 25 variables:
##
  $ num
                : int 1112222333...
## $ sex
                : int 0000000000...
                      75.1 75.1 75.1 45.8 45.8 ...
##
   $ age
                : num
                : num 0.011 3.679 4.696 6.364 7.304 ...
##
   $ time
##
  $ fuyrs
                : num 4.96 4.96 4.96 9.66 9.66 ...
## $ status
                : int 0000000000...
##
   $ grad
                : num 10 10 10 14 9 12 NA NA 10 10 ...
## $ log.grad
                : num 2.3 2.3 2.3 2.64 2.2 ...
  $ lvmi
                : num 119 119 138 115 110 ...
                : num 4.78 4.78 4.92 4.74 4.7 ...
## $ log.lvmi
##
   $ ef
                      93 93 93 68 70 56 75 38 74 49 ...
                : int
                : num 1.77 1.77 1.77 1.92 1.92 1.92 1.92 1.65 1.65 1.65 ...
## $ bsa
## $ lvh
                : int 1 1 1 1 1 1 1 0 0 0 ...
## $ prenyha
                : int 3 3 3 1 1 1 1 3 3 3 ...
```

```
##
   $ redo
                 : int 0001111000...
                : int 27 27 27 22 22 22 22 25 25 25 ...
##
   $ size
##
   $ con.cabg
                : int
                      1 1 1 0 0 0 0 0 0 0 ...
                      103 103 103 76 76 76 76 130 130 130 ...
##
  $ creat
                 : int
##
   $ dm
                : int
                       0 0 0 0 0 0 0 0 0 0 ...
## $ acei
                : int 1 1 1 0 0 0 0 1 1 1 ...
                       1 1 1 2 2 2 2 1 1 1 ...
##
  $ lv
                : int
##
   $ emergenc
                : int
                       0 0 0 0 0 0 0 0 0 0 ...
##
   $ hc
                 : int 0000000000...
##
   $ sten.reg.mix: int 1 1 1 1 1 1 2 2 2 ...
                 : Factor w/ 2 levels "Homograft", "Stentless valve": 2 2 2 1 1 1 1 1 1 1 ...
```

Loading needed tidymodels packages:

```
library(tidymodels)
library(censored)

# -----

tidymodels_prefer()
theme_set(theme_bw())
options(pillar.advice = FALSE, pillar.min_title_chars = Inf)
```

We'll retrieve the appropriate event times for the outcome (since there are multiple time points where patients were measured). Then, we'll identify the predictors that have the same values across the multiple time points and merge them. Functions in the joineR package will help us out here:

```
## # A tibble: 256 x 9
##
     fuyrs status
                    age emergenc
                                    hc hs
                                                          lv
                                                               sex sten.reg.mix
##
      <dbl>
            <int> <dbl>
                           <int> <int> <fct>
                                                       <int> <int>
                                                                          <int>
##
   1 4.96
                0 75.1
                               0
                                     O Stentless valve
                                                           1
                                                                 0
                                                                              1
##
      9.66
                0 45.8
                               0
                                     0 Homograft
                                                           2
                                                                 0
                                                                              1
  3 7.92
                                                                              2
##
                0 63.3
                               0
                                     0 Homograft
                                                                 0
                                                           1
##
  4 4.04
                0 61.4
                               0
                                     0 Homograft
                                                           2
                                                                 0
                                                                              2
  5 8.82
                0 53.6
                               0
                                                                              2
##
                                     0 Homograft
                                                           1
                                                                 0
##
   6 6.25
                1 67.3
                               0
                                     1 Homograft
                                                           1
                                                                 0
                                                                              1
                               0
                                     O Stentless valve
                                                                              3
##
  7 7.98
                0 67.8
                                                           2
                                                                 0
  8 4.90
                0 73.0
                               0
                                     0 Homograft
                                                                              2
                                                           1
                               0
                                     0 Homograft
##
  9 9.20
                0 47.8
                                                           1
                                                                 0
                                                                              1
```

```
## 10 8.47 0 72.1 0 0 Homograft 2 0 3
```

We'll reformat some of the categorical predictors since they are currently encoded as integers.

Also, tidymodels expects that the event times and corresponding status data are pre-formatted using the Surv function in the survival package. We'll do that, then remove the original fuyrs and status columns.

```
heart_data <-
 heart_data %>%
  mutate(
    event_time = Surv(fuyrs, status),
    lv =
      case_when(
        lv == 1 ~ "good",
        lv == 2 ~ "moderate",
        lv == 3 ~ "poor"
      ),
    emergenc =
      case_when(
        emergenc == 0 ~ "elective",
        emergenc == 1 ~ "urgent",
        emergenc == 2 ~ "emergency"
      ),
    hc =
      case_when(
        hc == 0 ~ "absent",
        hc == 1 ~ "present_treated",
        hc == 2 ~ "present_untreated"
     ),
    sten.reg.mix =
      case_when(
        sten.reg.mix == 1 ~ "stenosis",
        sten.reg.mix == 2 ~ "regurgitation",
        sten.reg.mix == 3 ~ "mixed"
      ),
    hs =
      case_when(
        hs == "Homograft" ~ "homograft",
        TRUE ~ "stentless_porcine_tissue"
    across(where(is.character), factor)
  ) %>%
  select(-fuyrs, -status)
```

Since our focus is on prediction, the standard tidymodels methods for data splitting are used to create training and test sets. We'll also make cross-validation folds:

```
set.seed(6941)
valve_split <- initial_split(heart_data)
valve_tr <- training(valve_split)
valve_te <- testing(valve_split)</pre>
```

In the training set, the observed time values range from 0.047 years to 11 years and 19.79% of the patients died (i.e. were events).

New Prediction Types

There are different types of predictions for event time analysis. *Dynamic* predictions require a specific time to make the prediction (sometimes called a "landmark time"). For example, we might want to know the probability of survival up to some time t. A *static* prediction is one that is not dependent on time. For example, we might predict the event time from a model.

To demonstrate, let's fit a bagged tree to the training data:

```
bag_spec <-
bag_tree() %>%
set_mode("censored regression") %>%
set_engine("rpart", nbagg = 50)

set.seed(29872)
bag_fit <-
bag_spec %>%
fit(event_time ~ ., data = valve_tr)
```

Instead of using the training or testing sets, let's make two fake patients by randomly selecting rows from the training set:

```
set.seed(4853)
fake_examples <-
   slice(valve_tr, sample(nrow(valve_tr), 2))
fake_examples</pre>
```

```
## # A tibble: 2 x 8
## age emergenc hc hs lv sex sten.reg.mix event_time
## <dbl> <fct> <fct> <fct> <fct> <int> <fct> <surv>
## 1 83.4 elective absent stentless_porcine_t~ mode~ 0 mixed 5.413699+
## 2 77.4 urgent absent stentless_porcine_t~ mode~ 1 stenosis 4.594521+
```

The standard predict() machinery can be used to get static (type = "time") or dynamic predictions (type = "survival"). We'll create a grid of 101 time points for the latter:

```
time_points <- seq(0, 10, by = .1)
bag_pred <-
predict(bag_fit, fake_examples, type = "survival", eval_time = time_points) %>%
bind_cols(
   predict(bag_fit, fake_examples),
   fake_examples %>% select(event_time)
) %>%
add_rowindex()
bag_pred
```

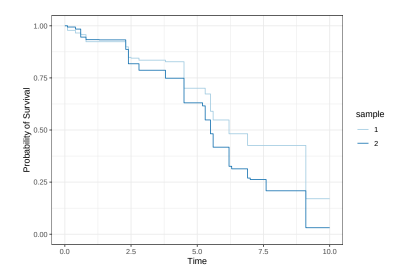
As usual, the prediction columns are prefaced with .pred_. What is unusual is that .pred is a list column, and each list element is a tibble with 2 columns and 101 rows. They contain the survival estimates for each patient:

bag_pred\$.pred[[1]] %>% slice(1:5)

```
## # A tibble: 5 x 2
     .eval_time .pred_survival
           <dbl>
##
                           <dbl>
## 1
             0
## 2
             0.1
                           0.978
                           0.978
## 3
             0.2
                           0.978
## 4
             0.3
## 5
             0.4
                           0.965
```

We can unnest these and plot the per-patient survival curves:

```
bag_pred %>%
  unnest(.pred) %>%
  mutate(sample = format(.row)) %>%
  ggplot(aes(.eval_time, .pred_survival, group = sample, col = sample)) +
  geom_step() +
  lims(y = 0:1) +
  labs(x = "Time", y = "Probability of Survival") +
  scale_color_brewer(palette = "Paired")
```



The static/dynamic prediction types make these models' tuning and evaluations a little more complex. In many tidymodels functions, there is a new argument called eval_time that is used to specify the time points for dynamic predictions (as we'll see in a minute).

Measures of Performance

Metrics to measure how well our model performs can also be split into dynamic and static metrics.

For static, a common choice is the concordance statistic, accessible via the concordance_survival() function. If we were looking at the test set results for the bagged tree model:

```
test_pred <-
   predict(bag_fit, valve_te, type = "survival", eval_time = time_points) %>%
bind_cols(
   predict(bag_fit, valve_te),
     valve_te %>% select(event_time)
)

test_pred %>%
   concordance_survival(truth = event_time, estimate = .pred_time)
```

Dynamic metrics usually are classification metrics re-purposed for survival analysis. For example, if we wanted to evaluate the model at t = 5, we could use the predicted survival probabilities and try to classify each data point as dead or alive. This ends up being a two class situation, and metrics like the Brier Score or the area under the ROC curve can be used to quantify how well the model works at time t.

The main difficulty is that, due to censoring, some data can't be cleanly classified. If we have a censored event at time 6, we definitely know that it should not be classified as an event. However, if the observed time were 2 and censored, we don't know if it is an event at t = 5 or not.

There are a lot of ways to deal with this issue. We've done an exhaustive reading of the literature and have come to a somewhat opinionated conclusion. Most of the survival metrics in the literature are developed to univariately score a collection of predictors, typically biomarkers, regarding how well they are associated with the event times. That's not what we are doing; we have model predictions.

Our choice for dynamic metrics is to use the inverse probability of censoring weights (IPCW), specifically the scheme used by Graf *et al.* (1999). They compute the probability that every data point might have been censored and uses the inverse of this value as a case weight. If the observed time is a censoring time, and that is before the evaluation time, the data point should make no contribution to the performance metric.

If you were to compute model performance manually (as above), these weights are computed using:

```
ipcw_data <-
  test_pred %>%
  .censoring_weights_graf(bag_fit, .) %>%
  select(-.pred_time)

ipcw_data
```

```
## # A tibble: 64 x 2
##
      .pred
                           event_time
##
      <list>
                               <Surv>
##
    1 <tibble [101 x 5]> 4.95616438+
##
   2 <tibble [101 x 5] > 8.81643836+
   3 <tibble [101 x 5]> 7.98082192+
##
    4 <tibble [101 x 5]> 9.20273973+
   5 <tibble [101 x 5] > 0.02191781+
##
   6 <tibble [101 x 5]> 4.37534247+
   7 <tibble [101 x 5] > 7.48493151
##
## 8 <tibble [101 x 5] > 3.20273973
```

```
## # i 54 more rows

# The adjusted data:
ipcw_data$.pred[[1]] %>% slice(1:5)
```

```
## # A tibble: 5 x 5
     .eval_time .pred_survival .weight_time .pred_censored .weight_censored
##
##
          <dbl>
                                                                          <dbl>
                          <dbl>
                                        <dbl>
                                                       <dbl>
## 1
            0
                          1
                                                       1
                                                                          1
## 2
            0.1
                          0.997
                                        0.100
                                                       1
                                                                          1
## 3
            0.2
                          0.997
                                        0.200
                                                       0.995
## 4
            0.3
                          0.997
                                       0.300
                                                                          1.01
## 5
            0.4
                          0.997
                                        0.400
                                                       0.995
                                                                          1.01
```

The last column is used as a case weight.

9 <tibble [101 x 5]> 6.31780800+ ## 10 <tibble [101 x 5]> 9.78630137+

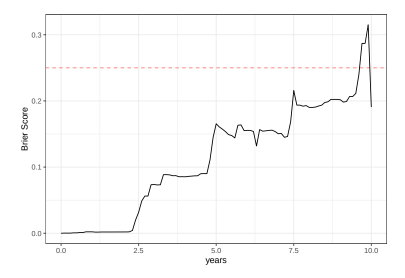
With the data in this format, we can use a yardstick function for dynamic metrics like brier_survival():

```
brier_scores <-
  ipcw_data %>%
  # No argument name is used for .pred
brier_survival(truth = event_time, .pred)
brier_scores %>% slice(1:5)
```

```
## # A tibble: 5 x 4
##
     .metric
                    .estimator .eval_time .estimate
##
     <chr>
                                              <dbl>
                   <chr>
                                    <dbl>
## 1 brier_survival standard
                                     0
                                           0
## 2 brier_survival standard
                                     0.1 0.000207
## 3 brier_survival standard
                                     0.2 0.000207
## 4 brier_survival standard
                                     0.3 0.000208
## 5 brier_survival standard
                                     0.4 0.000599
```

We compute a score for each landmark time:

```
brier_scores %>%
  ggplot(aes(.eval_time, .estimate)) +
  geom_hline(yintercept = 0.25, col = "red", alpha = 1 / 2, lty = 2) +
  geom_line() +
  labs(x = "years", y = "Brier Score")
```



The vertical line is the level of performance that you would get with a non-informative model.

The other dynamic metrics that are currently implemented are brier_survival_integrated() (for an AUC of the curve above) and roc_auc_survival().

Multiple static and dynamic metrics can be combined via a metric set.

Resampling the Model

tidymodels strongly focuses on empirical validation via resampling, which is also true of event time models.

We can use the fit_resamples() function with an rsample object to compute performance without using the test set. We need to tell the function what times to use for the dynamic metrics:

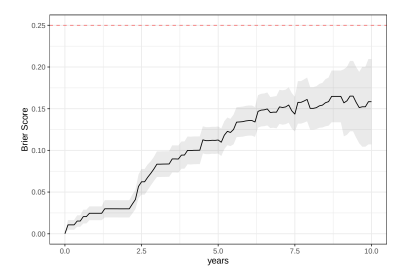
```
# Create resamples
set.seed(12)
valve_rs <- vfold_cv(valve_tr, repeats = 5)

bag_tree_res <-
bag_spec %>%
fit_resamples(event_time ~ ., resamples = valve_rs, eval_time = time_points)
```

By default, the Brier score is used:

```
collect_metrics(bag_tree_res) %>% slice(1:5)
```

```
## # A tibble: 5 x 7
##
     .eval_time .metric
                                .estimator
                                                      n std_err .config
                                             mean
          <dbl> <chr>
                                            <dbl> <int>
                                                          <dbl> <chr>
##
                                <chr>
## 1
                                                     50 0
               brier_survival standard
                                                                Preprocessor1_Model1
                                           0
                                                     50 0.00298 Preprocessor1_Model1
## 2
            0.1 brier_survival standard
                                           0.0107
            0.2 brier_survival standard
                                                     50 0.00298 Preprocessor1_Model1
## 3
                                           0.0107
## 4
            0.3 brier_survival standard
                                           0.0107
                                                     50 0.00298 Preprocessor1_Model1
            0.4 brier_survival standard
                                                     50 0.00350 Preprocessor1_Model1
## 5
                                           0.0152
```



Model Tuning

Suppose we try a regularized Cox model for these data. We'll add a recipe to the analysis and tune a lasso model. The code is pretty standard tidymodels syntax, with the added eval_time argument. We'll also use a metric set to include the integrated Brier score, which computes the AUC of the Brier/time curve:

```
lasso_spec <-
   proportional_hazards(penalty = tune(), mixture = 0) %>%
   set_engine("glmnet") %>%
   set_mode("censored regression")

lasso_rec <-
   recipe(event_time ~ ., data = valve_tr) %>%
   step_dummy(all_nominal_predictors()) %>%
   step_zv(all_predictors()) %>%
   step_normalize(all_numeric_predictors())

lasso_wflow <- workflow(lasso_rec, lasso_spec)

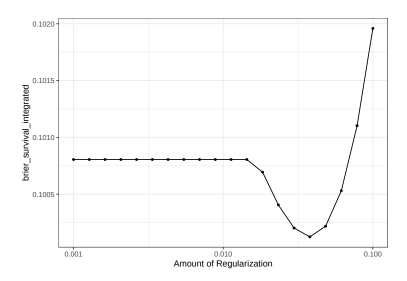
surv_metrics <- metric_set(brier_survival_integrated, brier_survival)</pre>
```

```
lasso_tune_res <-
lasso_wflow %>%
tune_grid(
   resamples = valve_rs,
   eval_time = time_points,
   grid = tibble(penalty = 10^seq(-3, -1, length.out = 20)),
   metrics = surv_metrics
)
```

We can plot the results for that specific metric:

3 Recipe Steps

```
autoplot(lasso_tune_res, metric = "brier_survival_integrated")
```



For these plot methods, eval_time can be passed in as shown. If a dynamic metric is used and eval_time is not set, the function will pick a time near the middle of the range.

We can also choose the best penalty. If we use an integrated method, no eval_time is needed:

```
best_penalty <- select_best(lasso_tune_res, metric = "brier_survival_integrated")</pre>
```

Now we can update the workflow and, assuming that this is the model that we want to keep, evaluate it on the test set:

For the test set, you can manually predict it or use last_fit() with the original split object:

```
test_res <-
last_fit(lasso_final_wflow, valve_split, eval_time = time_points)</pre>
```

As usual, you can get the test set statistics via:

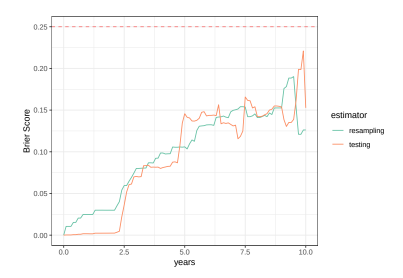
```
collect_metrics(test_res)
```

```
## # A tibble: 101 x 5
##
                   .estimator .eval_time .estimate .config
      .metric
                                  <dbl>
##
     <chr>
                   <chr>
                                             <dbl> <chr>
## 1 brier_survival standard
                                   0
                                                  Preprocessor1_Model1
                                    0.1 0.000267 Preprocessor1 Model1
## 2 brier survival standard
## 3 brier_survival standard
                                     0.2 0.000267 Preprocessor1_Model1
## 4 brier_survival standard
                                     0.3 0.000269 Preprocessor1_Model1
## 5 brier_survival standard
                                     0.4 0.000605 Preprocessor1_Model1
## 6 brier_survival standard
                                    0.5 0.000609 Preprocessor1_Model1
## 7 brier_survival standard
                                     0.6 0.00109 Preprocessor1 Model1
## 8 brier_survival standard
                                    0.7 0.00110 Preprocessor1_Model1
## 9 brier survival standard
                                     0.8 0.00173 Preprocessor1 Model1
                                     0.9 0.00173 Preprocessor1_Model1
## 10 brier_survival standard
## # i 91 more rows
```

How do the Brier Score estimates compare between the test set and resampling?

```
collect_metrics(test_res) %>%
  mutate(estimator = "testing") %>%
  select(.eval_time, estimator, Brier = .estimate) %>%
  bind_rows(
  lasso_tune_res %>%
    collect_metrics() %>%
    mutate(estimator = "resampling") %>%
    select(.eval_time, estimator, Brier = mean, penalty) %>%
    inner_join(best_penalty, by = "penalty")
) %>%
  ggplot(aes(.eval_time)) +
  geom_hline(yintercept = 0.25, col = "red", alpha = 1 / 2, lty = 2) +
```

```
geom_line(aes(y = Brier, col = estimator)) +
labs(x = "years", y = "Brier Score") +
scale_color_brewer(palette = "Set2")
```



Good!

Things still to do

- Update finetune to use eval_time
- Update Bayesian analysis methods in tidyposterior
- Update parsnip augment() to produce IPCW values.

Session Info

```
sessioninfo::session_info()
## - Session info
    setting value
##
##
    version R version 4.2.2 (2022-10-31)
             macOS Monterey 12.6
##
             aarch64, darwin20
##
   system
   ui
             X11
##
   language (EN)
##
##
    collate en_US.UTF-8
##
             {\tt en\_US.UTF-8}
   ctype
             America/New_York
##
##
    date
             2023-04-18
             2.19.2 @ /Applications/RStudio.app/Contents/Resources/app/quarto/bin/tools/ (via rmarkdown
##
    pandoc
##
## - Packages --
                               date (UTC) lib source
##
    package
                 * version
  backports
                   1.4.1
                               2021-12-13 [1] CRAN (R 4.2.0)
```

```
broom
                 * 1.0.4.9000 2023-03-23 [1] Github (tidymodels/broom@c0c2253)
##
    cachem
                               2023-02-24 [1] CRAN (R 4.2.2)
                   1.0.7
                               2023-04-13 [1] CRAN (R 4.2.0)
##
    censored
                 * 0.2.0
                   7.3-21
                               2023-01-23 [2] CRAN (R 4.2.0)
##
    class
##
                   3.6.1
                               2023-03-23 [1] CRAN (R 4.2.0)
##
    {\tt codetools}
                   0.2-19
                               2023-02-01 [2] CRAN (R 4.2.0)
                   2.1-0
                               2023-01-23 [1] CRAN (R 4.2.0)
    colorspace
                               2023-02-01 [1] CRAN (R 4.2.0)
##
    conflicted
                   1.2.0
##
    data.table
                   1.14.8
                               2023-02-17 [1] CRAN (R 4.2.0)
##
                 * 1.1.0.9000 2023-04-03 [1] Github (tidymodels/dials@6b59d71)
    dials
    DiceDesign
                   1.9
                               2021-02-13 [1] CRAN (R 4.2.0)
                               2022-12-11 [1] CRAN (R 4.2.0)
##
                   0.6.31
    digest
                               2022-02-05 [2] CRAN (R 4.2.0)
##
    doMC
                 * 1.3.8
##
                 * 1.1.1
                               2023-03-22 [1] CRAN (R 4.2.0)
    dplyr
##
                   0.3.2
                               2021-04-29 [1] CRAN (R 4.2.0)
    ellipsis
##
    evaluate
                   0.20
                               2023-01-17 [1] CRAN (R 4.2.0)
##
                   1.0.4
                               2023-01-22 [1] CRAN (R 4.2.0)
    fansi
##
    farver
                   2.1.1
                               2022-07-06 [1] CRAN (R 4.2.0)
                               2023-02-24 [1] CRAN (R 4.2.0)
##
    fastmap
                   1.1.1
    foreach
                 * 1.5.2
                               2022-02-02 [1] Github (cran/foreach@d4bac0e)
##
    furrr
                   0.3.1
                               2022-08-15 [1] CRAN (R 4.2.0)
##
    future
                   1.32.0
                               2023-03-07 [1] CRAN (R 4.2.0)
                               2022-11-05 [1] CRAN (R 4.2.0)
   future.apply
                   1.10.0
##
                   0.1.3.9000 2023-03-12 [1] Github (r-lib/generics@3873971)
##
    generics
                 * 3.4.2.9000 2023-04-11 [1] Github (tidyverse/ggplot2@2e649bb)
##
    ggplot2
                               2022-11-27 [1] CRAN (R 4.2.0)
    glmnet
                 * 4.1-6
##
                   0.16.2
                               2022-11-21 [1] CRAN (R 4.2.0)
    globals
    glue
                               2022-02-24 [1] CRAN (R 4.2.0)
##
                   1.6.2
##
                   1.0.1
                               2022-12-22 [1] CRAN (R 4.2.0)
    gower
##
    GPfit
                   1.0-8
                               2019-02-08 [1] CRAN (R 4.2.0)
##
    gtable
                   0.3.3
                               2023-03-21 [1] CRAN (R 4.2.0)
##
    hardhat
                   1.3.0.9000 2023-03-31 [1] Github (tidymodels/hardhat@ac2dfd0)
##
    htmltools
                   0.5.4
                               2022-12-07 [1] CRAN (R 4.2.0)
##
                 * 1.0.4.9000 2023-04-13 [1] local
    infer
##
    ipred
                 * 0.9-14
                               2023-03-09 [1] CRAN (R 4.2.0)
                 * 1.0.14
                               2022-02-05 [1] CRAN (R 4.2.0)
##
    iterators
##
    joineR
                 * 1.2.8
                               2023-01-22 [1] CRAN (R 4.2.0)
##
    knitr
                   1.42
                               2023-01-25 [1] CRAN (R 4.2.0)
##
                   0.4.2
                               2020-10-20 [1] CRAN (R 4.2.0)
    labeling
##
                   0.20-45
                               2021-09-22 [2] CRAN (R 4.2.2)
    lattice
                   1.7.2.1
                               2023-02-27 [1] CRAN (R 4.2.0)
##
    lava
                               2022-12-17 [1] CRAN (R 4.2.0)
##
                   1.1.6
   lhs
                               2022-10-07 [1] CRAN (R 4.2.0)
##
    lifecycle
                   1.0.3
##
   listenv
                   0.9.0
                               2022-12-16 [1] CRAN (R 4.2.0)
    lubridate
                   1.9.2
                               2023-02-10 [1] CRAN (R 4.2.0)
                               2022-03-30 [1] CRAN (R 4.2.0)
##
    magrittr
                   2.0.3
                               2023-01-23 [1] CRAN (R 4.2.0)
##
    MASS
                   7.3-58.2
##
                               2022-11-11 [1] CRAN (R 4.2.0)
    Matrix
                 * 1.5-3
   memoise
                   2.0.1
                               2021-11-26 [1] CRAN (R 4.2.0)
                               2023-01-25 [1] CRAN (R 4.2.0)
##
    modeldata
                 * 1.1.0
##
                   0.5.0
                               2018-06-12 [1] CRAN (R 4.2.0)
    munsell
## nlme
                   3.1-162
                               2023-01-31 [1] CRAN (R 4.2.0)
##
  nnet
                   7.3-18
                               2022-09-28 [2] CRAN (R 4.2.2)
    parallelly
                   1.35.0
                               2023-03-23 [1] CRAN (R 4.2.0)
```

```
parsnip
                 * 1.1.0.9000 2023-04-18 [1] Github (tidymodels/parsnip@51b0cd7)
                 1.9.0
##
                              2023-03-22 [1] CRAN (R 4.2.0)
    pillar
## pkgconfig
                 2.0.3
                              2019-09-22 [1] CRAN (R 4.2.0)
                   2023.03.31 2023-04-02 [1] CRAN (R 4.2.0)
## prodlim
##
    purrr
                 * 1.0.1.9000 2023-03-23 [1] Github (tidyverse/purrr@fd5a732)
## R6
                  2.5.1
                              2021-08-19 [1] CRAN (R 4.2.0)
## RColorBrewer 1.1-3
                              2022-04-03 [1] CRAN (R 4.2.0)
                  1.0.10
                              2023-01-22 [1] CRAN (R 4.2.0)
## Rcpp
## recipes
                 * 1.0.5.9000 2023-04-02 [1] local
## rlang
                 1.1.0.9000 2023-04-18 [1] Github (r-lib/rlang@9b50b7a)
                              2023-01-19 [1] CRAN (R 4.2.0)
## rmarkdown
                 2.20
## rpart
                  4.1.19
                              2022-10-21 [2] CRAN (R 4.2.2)
                * 1.1.1.9000 2023-03-31 [1] Github (tidymodels/rsample@690a1fb)
## rsample
## rstudioapi
                              2022-08-22 [1] CRAN (R 4.2.0)
                  0.14
## scales * 1.2.1
## sessioninfo 1.2.2
## shape 1.4.6
## statmod 1.5.0
## survival * 3.5-3
## tibble * 3.2.1
## tidymodels * 1.0.0
                              2022-08-20 [1] CRAN (R 4.2.0)
                              2021-12-06 [2] CRAN (R 4.2.0)
                              2021-05-19 [1] CRAN (R 4.2.0)
                              2023-01-06 [1] CRAN (R 4.2.2)
                              2023-02-12 [1] CRAN (R 4.2.0)
                              2023-03-20 [1] CRAN (R 4.2.0)
                              2022-07-13 [1] CRAN (R 4.2.0)
## tidyr
                 * 1.3.0.9000 2023-03-23 [1] Github (tidyverse/tidyr@0764e65)
                              2022-10-10 [1] CRAN (R 4.2.0)
## tidyselect
                  1.2.0
                  0.2.0
## timechange
                              2023-01-11 [1] CRAN (R 4.2.0)
## timeDate
                  4022.108 2023-01-07 [1] CRAN (R 4.2.2)
## tune
                 * 1.0.1.9003 2023-04-18 [1] Github (tidymodels/tune@c2ccb50)
## utf8
                  1.2.3
                              2023-01-31 [1] CRAN (R 4.2.0)
                  0.6.1.9000 2023-03-23 [1] Github (r-lib/vctrs@af29ad7)
## vctrs
                              2022-03-03 [1] CRAN (R 4.2.0)
## withr
                  2.5.0
## workflows * 1.1.3.9000 2023-03-31 [1] Github (tidymodels/workflows@f5bdfae)
## workflowsets * 1.0.0.9000 2022-12-19 [1] local
## xfun
                   0.37
                              2023-01-31 [1] CRAN (R 4.2.0)
                              2023-01-23 [1] CRAN (R 4.2.0)
## yaml
                   2.3.7
                 * 1.1.0.9001 2023-04-18 [1] Github (tidymodels/yardstick@d4a9d7a)
## yardstick
##
## [1] /Users/simoncouch/Library/R/arm64/4.2/library
## [2] /Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/library
##
```