P8108 Group 2 Survival Analysis Project

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```

2022-12-1

```
library(survival)
library(tidyverse)
library(tidymodels)
library(glmnet)
library(ranger)
library(survminer)
library(survminer)
knitr::opts_chunk$set(message = FALSE, warning = FALSE)
```

Train Validation Test Split

Perform 10-fold Cross-Validation

The output contains 1 row for each fold/repeat. So, 10 folds * 5 repeats = 50 rows. The split_analysis column is a list column containing a data frame for each row with 9 folds combined, and the split_assessment column is a list column containing a data frame for each row with 1 fold.

Introduction

For our target population is hormone treatment an effective therapy in breast cancer survival? For our target population, is chemotherapy an effective therapy in breast cancer survival? How do predictions from non-parametric models like the random forest compare to semi-parametric in the Cox proportional hazard model?

Methods

The dataset of interest for this analysis comes from the Rotterdam tumor bank, including data from 2982 breast cancer patients. Follow up time for patients varied from just 1 month to as long as 231 months. Several prognostic variables are recorded including year of surgery, age at surgery, menopausal status (preor post-), tumor size (mm), differentiation grade, number of positive lymph nodes, progesterone receptors (fmol/l), estrogen receptors (fmol/l), and indicators for hormonal treatment and chemotherapy treatment. The outcome considered in this analysis was patient death.

(Placeholder for Cross-validation)

As part of this analysis, we consider the Cox Proportional Hazard (Cox PH) model, which allows us to model the hazard ratio based on covariates to understand their impact on the survival function. The Cox PH typically takes the form:

$$h(t|Z=z) = h_0(t)e^{\beta'z}.$$

In this application, we use the elastic net penalty, a mixture of the ℓ_1 and ℓ_2 norm regularization penalties. In the Cox PH framework, this penalty term takes the form of:

$$\lambda \left(\alpha \sum |\beta_i| + \frac{1}{2} (1 - \alpha) \sum \beta_i^2 \right)$$

where λ represents our penalty coefficient and α is the mixing parameter for the two regularization methods. This penalty helps to avoid over-fitting of our data. The algorithm used here in glmnet uses the Breslow approximation to handle ties. For more details on the derivation of this term and the algorithm used to fit the penalized Cox PH model, see Simon et al. (2011).

Exploratory Data Analysis

	0 (N=2643)	1 (N=339)	Total (N=2982)	p value
age				< 0.001
Mean (SD)	54.098 (12.984)	62.549 (9.921)	55.058 (12.953)	
Range	24.000 - 90.000	28.000 - 88.000	24.000 - 90.000	
meno				< 0.001
Mean (SD)	0.519 (0.500)	0.879 (0.327)	0.560 (0.496)	
Range	0.000 - 1.000	0.000 - 1.000	0.000 - 1.000	
size				< 0.001
<=20	1283 (48.5%)	104 (30.7%)	1387 (46.5%)	
20-50	$1119\ (42.3\%)$	172(50.7%)	$1291\ (43.3\%)$	
>50	241 (9.1%)	63 (18.6%)	304 (10.2%)	
grade	, ,	. ,		< 0.001

	0 (N=2643)	1 (N=339)	Total ($N=2982$)	p value
Mean (SD)	2.722 (0.448)	2.826 (0.380)	2.734 (0.442)	
Range	2.000 - 3.000	2.000 - 3.000	2.000 - 3.000	
nodes				< 0.001
Mean (SD)	2.327 (4.207)	5.720(4.576)	2.712(4.384)	
Range	0.000 - 34.000	1.000 - 24.000	0.000 - 34.000	
pgr				< 0.001
Mean (SD)	168.706 (300.337)	$108.233 \ (200.302)$	$161.831 \ (291.311)$	
Range	0.000 - 5004.000	0.000 - 1497.000	0.000 - 5004.000	
\mathbf{er}				0.069
Mean (SD)	$164.792\ (272.563)$	180.608 (271.693)	$166.590\ (272.465)$	
Range	0.000 - 3275.000	0.000 - 2444.000	0.000 - 3275.000	
chemo				< 0.001
Mean (SD)	0.209 (0.407)	$0.083 \ (0.276)$	0.195 (0.396)	
Range	0.000 - 1.000	0.000 - 1.000	0.000 - 1.000	
dtime				< 0.001
Mean (SD)	2679.067 (1309.178)	2030.534 (1043.971)	2605.340 (1298.078)	
Range	36.000 - 7043.000	45.000 - 6270.000	36.000 - 7043.000	
death				0.093
Mean (SD)	$0.421 \ (0.494)$	$0.469 \ (0.500)$	$0.427 \ (0.495)$	
Range	0.000 - 1.000	0.000 - 1.000	0.000 - 1.000	

	Overall (N=2982)
age	
Mean (SD)	$55.058\ (12.953)$
Range	24.000 - 90.000
meno	
Mean (SD)	$0.560 \ (0.496)$
Range	0.000 - 1.000
\mathbf{size}	
<=20	1387~(46.5%)
20-50	1291 (43.3%)
> 50	$304 \ (10.2\%)$
grade	
Mean (SD)	2.734(0.442)
Range	2.000 - 3.000
nodes	
Mean (SD)	2.712(4.384)
Range	0.000 - 34.000
pgr	
Mean (SD)	161.831 (291.311)
Range	0.000 - 5004.000
\mathbf{er}	
Mean (SD)	166.590 (272.465)
Range	0.000 - 3275.000
chemo	
Mean (SD)	0.195(0.396)
Range	0.000 - 1.000

Overall (N=2982)
0.114(0.317)
0.000 - 1.000
2605.340 (1298.078)
36.000 - 7043.000
0.427 (0.495)
0.000 - 1.000

Cross-Validation

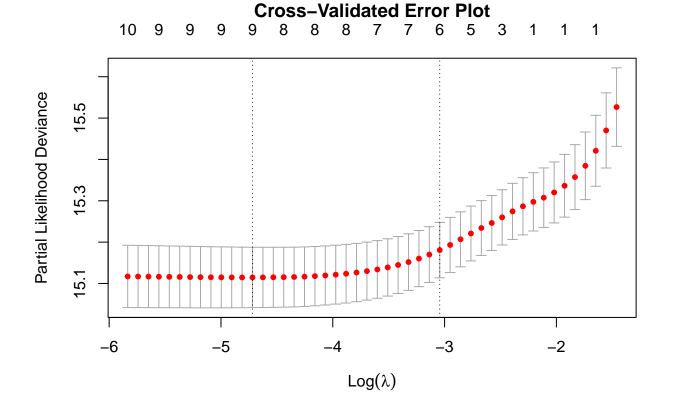
Cox/Cox with elastic net

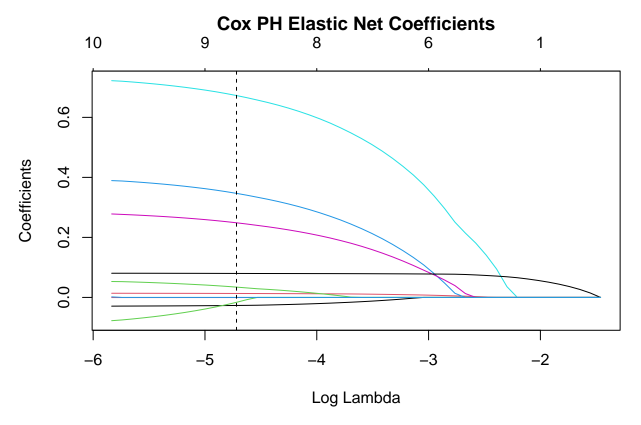
```
set.seed(2022)

cox_trn_x <- model.matrix(Surv(dtime, death) ~ ., rotterdam_training)[,-1]
cox_trn_y <- Surv(rotterdam_training$dtime, rotterdam_training$death)

cv_coxfit <- cv.glmnet(cox_trn_x, cox_trn_y, family = "cox", type.measure = "deviance")

par(mar = c(4,4,5,1))
plot(cv_coxfit, main = "Cross-Validated Error Plot")</pre>
```





```
coxnetfit_df <-
  data.frame(
    "coef" = as.vector(coef(coxnetfit, s = cv_coxfit$lambda.min)),
    "exp_coef" = as.vector(coef(coxnetfit, s = cv_coxfit$lambda.min)) %>% exp()
)

rownames(coxnetfit_df) <- labels(coef(coxnetfit, s = cv_coxfit$lambda.min))[[1]]

coxnetfit_df %>% round(digits = 4) %>%
    knitr::kable(caption = "Cox Proportion Hazard Elastic Net Coefficients")
```

Table 3: Cox Proportion Hazard Elastic Net Coefficients

	coef	exp_coef
year	-0.0268	0.9735
age	0.0132	1.0132
meno	0.0343	1.0349
size 20-50	0.3467	1.4144
size > 50	0.6727	1.9596

	coef	exp_coef
grade	0.2487	1.2823
nodes	0.0799	1.0832
pgr	-0.0004	0.9996
er	0.0000	1.0000
hormon	-0.0163	0.9838
chemo	0.0000	1.0000

In the table above, we can see that estrogen receptors and chemotherapy are selected out with a null value of 0 or $\exp(coef) = 1$. We can fit a cox proportional hazard model using only the selected covariates in the coxph function to find unbiased estimates of the coefficients along with standard errors and confidence intervals.

```
coxfit <- coxph(Surv(dtime, death) ~ year + age + meno + size + grade +</pre>
                  nodes + pgr + hormon,
                data = rotterdam_training, ties = "breslow")
coxfit %>%
  broom::tidy() %>%
 mutate(estimate = exp(estimate))
## # A tibble: 9 x 5
##
     term
               estimate std.error statistic p.value
##
     <chr>>
                  <dbl>
                            <dbl>
                                      <dbl>
                                                <dbl>
## 1 year
                  0.970 0.0129
                                      -2.34 1.92e- 2
                                      3.05 2.27e- 3
## 2 age
                  1.01
                         0.00464
```

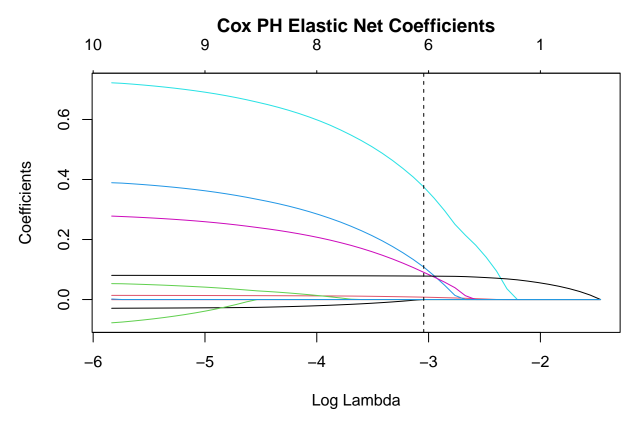
3 meno 0.517 6.05e- 1 1.07 0.122 ## 4 size20-50 1.51 0.0825 4.99 6.16e- 7 ## 5 size>50 6.51 7.56e-11 2.11 0.115 ## 6 grade 1.34 0.0903 3.24 1.19e- 3 ## 7 nodes 1.08 0.00633 12.8 2.11e-37 0.000147 -3.35 8.01e- 4 ## 8 pgr 1.00 ## 9 hormon 0.897 0.117 -0.928 3.53e- 1

<pre>confint(coxfit)</pre>	%>%	exp()	%>%	<pre>knitr::kable()</pre>

year 0.9462185 0.9951111 age 1.0050778 1.0235067 meno 0.8384713 1.3529343 size20-50 1.2837050 1.7740258 size>50 1.6860598 2.6446630 grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958			
age 1.0050778 1.0235067 meno 0.8384713 1.3529343 size20-50 1.2837050 1.7740258 size>50 1.6860598 2.6446630 grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958		2.5 %	97.5 %
meno 0.8384713 1.3529343 size20-50 1.2837050 1.7740258 size>50 1.6860598 2.6446630 grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958	year	0.9462185	0.9951111
size20-50 1.2837050 1.7740258 size>50 1.6860598 2.6446630 grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958	age	1.0050778	1.0235067
size>50 1.6860598 2.6446630 grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958	meno	0.8384713	1.3529343
grade 1.1227352 1.5993840 nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958	size 20-50	1.2837050	1.7740258
nodes 1.0709185 1.0978328 pgr 0.9992211 0.9997958	size > 50	1.6860598	2.6446630
pgr 0.9992211 0.9997958	grade	1.1227352	1.5993840
10	nodes	1.0709185	1.0978328
hormon 0.7133152 1.1282404	pgr	0.9992211	0.9997958
	hormon	0.7133152	1.1282404

In our (minimum error) model, we find significant effects for year of surgery, age at surgery, size of tumor, differentiation grade, number of positive lymph nodes, and progesterone receptors. The largest magnitude effects come from increasing size of tumor.

Below, we see the analogous results for a "1se" rule model.



```
coxnetfit_1se_df <-
  data.frame(
    "coef" = as.vector(coef(coxnetfit, s = cv_coxfit$lambda.1se)),
    "exp_coef" = as.vector(coef(coxnetfit, s = cv_coxfit$lambda.1se)) %>% exp()
)

rownames(coxnetfit_1se_df) <- labels(coef(coxnetfit, s = cv_coxfit$lambda.1se))[[1]]

coxnetfit_1se_df %>% round(digits = 4) %>%
  knitr::kable(caption = "Cox Proportion Hazard Elastic Net Coefficients (1se)")
```

Table 5: Cox Proportion Hazard Elastic Net Coefficients (1se)

	coef	exp_coef
year	0.0000	1.0000
age	0.0079	1.0079
meno	0.0000	1.0000
size 20-50	0.1081	1.1142
size > 50	0.3752	1.4553
grade	0.0907	1.0949
nodes	0.0781	1.0813

coef	exp_coef
-0.0001	0.9999
0.0000	1.0000
0.0000	1.0000
0.0000	1.0000
	-0.0001 0.0000 0.0000

Here, we remove meno, er, hormon and chemo and find the following results.

```
## # A tibble: 7 x 5
##
               estimate std.error statistic p.value
     term
##
     <chr>
                  <dbl>
                            <dbl>
                                       <dbl>
                                                <dbl>
## 1 year
                  0.967 0.0123
                                       -2.74 6.17e- 3
## 2 age
                         0.00281
                                       5.54 3.00e- 8
                  1.02
## 3 size20-50
                  1.50
                         0.0821
                                        4.91 9.00e- 7
## 4 size>50
                  2.09
                         0.114
                                        6.47 9.86e-11
## 5 grade
                  1.34
                         0.0899
                                        3.24 1.21e- 3
                         0.00627
## 6 nodes
                                       12.8 1.39e-37
                  1.08
## 7 pgr
                  1.00
                         0.000146
                                       -3.36 7.93e- 4
```

```
confint(coxfit_1se) %>% exp() %>% knitr::kable()
```

	2.5 %	97.5 %
year	0.9440155	0.9905000
age	1.0101039	1.0212800
size 20-50	1.2743768	1.7583626
size > 50	1.6722634	2.6147576
grade	1.1216338	1.5957779
nodes	1.0704169	1.0970519
pgr	0.9992240	0.9997963

Here we again find significant effects for year of surgery, age at surgery, size of tumor, differentiation grade, number of positive lymph nodes, and pgr.

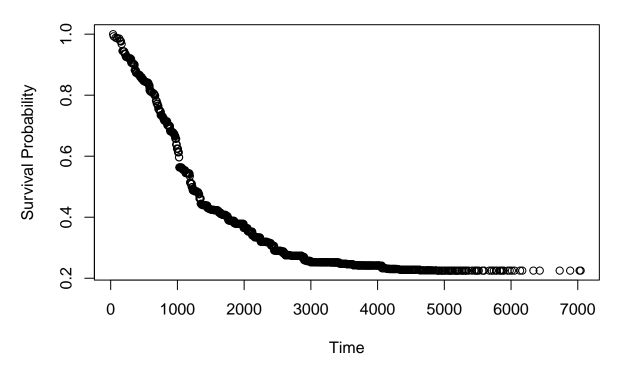
Random survival forest

The survival tree and the corresponding random survival forest (RSF) are highly favorable non-parametric methods when studying survival data. Generally, for a single survival tree, it will assign subjects to groups based on certain splitting rules regarding their covariates, and the subjects in each group will share a similar survival behavior.

```
set.seed(2023)
## Random Survival Forest
rsf <- ranger(Surv(time = dtime, event = death) ~ .,</pre>
              data = rotterdam_training,
              num.trees = 300,
              min.node.size = 15,
              importance = "permutation",
              scale.permutation.importance = TRUE)
## Remove variables not for prediction, and the outcome
rotterdam_test_d <-
  rotterdam_test %>%
  select(-death)
## Make prediction on all the test data points
pred_rsf <- predict(rsf, rotterdam_test_d, type = "response")</pre>
# Look at individual 7
pred_ref_7 <- data.frame(</pre>
 time = pred_rsf$unique.death.times,
  survival = pred_rsf$survival[7,])
head(pred_ref_7) %>% knitr::kable(align = "c")
```

time	survival
36	1.0000000
45	0.9929877
74	0.9881553
97	0.9877437
101	0.9857489
129	0.9857489

Survival Prediction for Patient 7

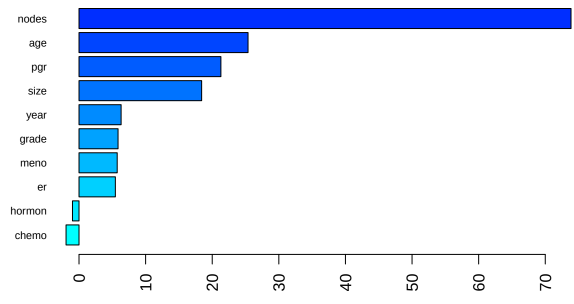


Find estimated median survival time for individual 7
head(pred_ref_7[pred_ref_7\$survival <= 0.5,]) %>% knitr::kable(align = "c") #1217

	$_{ m time}$	survival
306	1217	0.4984613
307	1218	0.4984613
308	1222	0.4984613
309	1226	0.4984613
310	1229	0.4927672
311	1231	0.4927672

```
# See the truth of individual 7
rotterdam_test[7,] %>% knitr::kable(align = "c")
```

	year	age	meno	size	grade	nodes	pgr	er	hormon	chemo	dtime	death
2463	1992	69	1	20-50	2	8	5	6	1	0	1869	0



With ranger package, we trained the random survival forest with training dataset used for survival prediction. As a non-parametric method, there is no parameters in RSF that could be interpreted. The ultimate goal of RSF is to predict the survival probability function of a given data point based on its covariate vector. Compared to semi-parametric Cox-PH model which forces the outcome and the covariates to have a special connection, the RSF makes prediction based on the survival time of training data points that shares similar propensity with the given input data point.

Since the "truth" of test data point (a single survival time) and the prediction we made here (a survival probability function) are not comparable, here we show the prediction result of the 7th test data point (pid = 58). The survival curve has been shown above, and the median survival time is 1217 days.

Comparison of Cox Proportional-Hazards Elastic Net with Random Survival Forest

We compared the Cox proportional-hazards elastic net model with the random survival forest by calculating the Brier score for each model on the validation set. The formula for the Brier score is as follows.

$$BS = \frac{1}{n} \sum_{i=1}^{n} (p_i - o_i)^2$$

The Brier score is used to evaluate the accuracy of probabilistic predictions from a model; its value ranges from 0 to 1 with 0 being perfect and 1 being the opposite. We calculated the Brier score using the validation set. For each observation in the validation set, predictions were made at the observed time of censoring or event. Our analysis proceeds as follows.

First, we calculated the Brier score for the Cox proportional-hazards elastic net model.

```
# Purpose: Calculates the Brier score for the Cox proportional-hazards elastic

# arguments: fit: The Cox proportional-hazards elastic net model.

# train: A dataframe, the training data used to fit the model.

# test: A dataframe, the data to use to calculate the Brier score.

# Returns: A double, the Brier score.

brier_coxnet <- function(fit, train, test) {
    train_x <- model.matrix(Surv(dtime, death) ~ ., train)[,-1]
    train_y <- Surv(pull(train, dtime), pull(train, death))</pre>
```

```
test_x <- model.matrix(Surv(dtime, death) ~ ., test)[,-1]</pre>
  test_y <- pull(test, death)</pre>
  num_obs <- nrow(test_x)</pre>
  p <- vector(mode = "double", length = num_obs)</pre>
  for(i in 1:num_obs) {
    surv_fit <- survival::survfit(fit, s = cv_coxfit$lambda.min,</pre>
                                    x = train_x,
                                    y = train_y,
                                    newx = test_x[i, ])
    time_index <- tail(which(surv_fittime <= test[i, "dtime"]), n = 1)
    p[i] <- 1 - surv_fit$surv[time_index]</pre>
  return(DescTools::BrierScore(resp = test_y, pred = p))
(brier_coxnet <- round(brier_coxnet(coxnetfit, rotterdam_training, rotterdam_validation), 3))
## [1] 0.329
```

The Brier score for the Cox elastic net model is 0.329.

Second, let's calculated the Brier score for the random survival forest model.

```
# Purpose: Calculates the Brier score for the random survival forest model.
# Arguments: fit: The random survival forest model.
             df: A dataframe, the data to use to calculate the Brier score.
# Returns: A double, the Brier score.
brier_ranger <- function(fit, df) {</pre>
  x <- df
  pred <- predict(fit, data = x)</pre>
  num obs <- nrow(df)</pre>
  p <- vector(mode = "double", length = num_obs)</pre>
  for(i in 1:num_obs) {
    time_index <- tail(which(pred$unique.death.times <= x[i, "dtime"]), n = 1)</pre>
    p[i] <- 1 - pred$survival[i, time_index]</pre>
  }
  return(DescTools::BrierScore(resp = df$death, pred = p))
(brier_ranger <- round(brier_ranger(rsf, rotterdam_validation), 3))</pre>
```

[1] 0.322

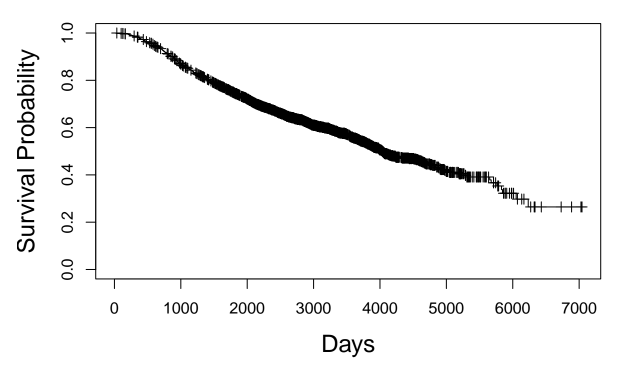
The Brier score for the random survival forest model is 0.322. The two models have very similar Brier scores.

Conformalized survival analysis

Supplemental analyses

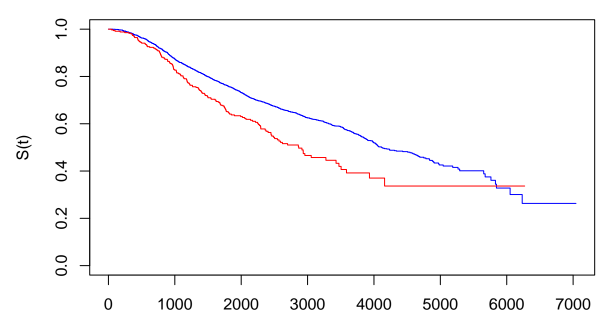
Kaplan-Meier Survival Estimate

Kaplan-Meier Survival Estimate

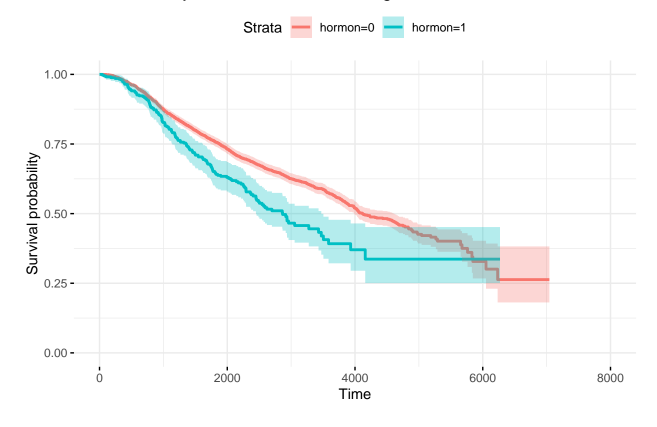


```
# make Kaplan-Meier estimates
kmfit <- survfit(Surv(dtime, death) ~ hormon, data = rotterdam, type=c("kaplan-meier"))
# print Kaplan-Meier table
#summary(kmfit)</pre>
```

stimates of Breast cancer survival by hormonal treatment assignments



days to death or last follow–up KM survival by hormonal treatment assignments



Log-rank Test

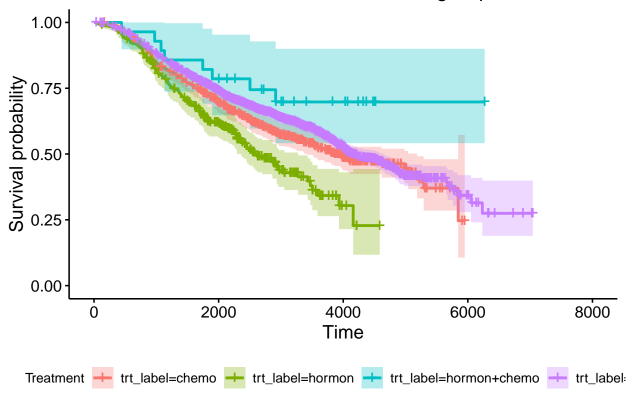
The null hypothesis of our log-rank test is: $H_0: S_1(t) = S_0(t)$, where $S_1(t)$ is the survival function of hormon treatment group, $S_0(t)$ is the survival function of control group.

Combined

```
# Add 1
rotterdam1 <-
  rotterdam %>%
 mutate(
    trt_label = case_when(
     hormon == 1 & chemo == 1 ~ "hormon+chemo",
     hormon == 1 & chemo == 0 ~ "hormon",
     hormon == 0 & chemo == 1 ~ "chemo",
      hormon == 0 & chemo == 0 ~ "none"
    )
  )
table(rotterdam1$trt_label)
##
##
          chemo
                      hormon hormon+chemo
                                                   none
##
            552
                         311
                                        28
                                                   2091
# Add 2
logrank1 <- survdiff(Surv(dtime, death) ~ trt_label, data = rotterdam1)</pre>
logrank1
## Call:
## survdiff(formula = Surv(dtime, death) ~ trt_label, data = rotterdam1)
##
##
                             N Observed Expected (0-E)^2/E (0-E)^2/V
## trt_label=chemo
                           552
                                     250
                                            233.7
                                                       1.13
                                                                  1.39
                                     151
                                             96.0
                                                                 34.43
## trt_label=hormon
                           311
                                                      31.45
## trt_label=hormon+chemo
                            28
                                       8
                                             14.3
                                                       2.79
                                                                  2.83
## trt_label=none
                           2091
                                            927.9
                                                       4.54
                                                                 16.84
                                     863
##
## Chisq= 40.4 on 3 degrees of freedom, p= 9e-09
logrank1$pvalue
## [1] 8.838693e-09
# Add 3
ggsurvplot(survfit(Surv(dtime,death) ~ trt_label, data = rotterdam1),
           conf.int = TRUE,
           legend = c("bottom"),
```

legend.title = c("Treatment")) +
ggtitle("Survival Curve of Hormon and Control group")

Survival Curve of Hormon and Control group



Hormon

```
logrank2 <- survdiff(Surv(dtime, death) ~ hormon, data = rotterdam)
logrank2</pre>
```

```
## Call:
## survdiff(formula = Surv(dtime, death) ~ hormon, data = rotterdam)
##
               N Observed Expected (O-E)^2/E (O-E)^2/V
##
                                         2.04
## hormon=0 2643
                     1113
                              1162
                                                   23.7
## hormon=1 339
                      159
                                110
                                        21.43
                                                   23.7
##
    Chisq= 23.7 on 1 degrees of freedom, p= 1e-06
```

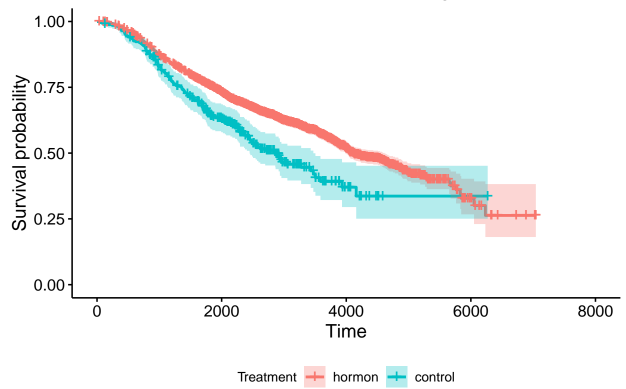
logrank2\$pvalue

[1] 1.133649e-06

The test statistic is 23.7, and the corresponding p-value is $1.133^{-6} \ll 0.05$, thus we reject the null and conclude that we are 95% confident that $S_1(t) \neq S_0(t)$. And since the test statistic is positive, we can conclude that the hormon treatment is significantly effective to breast cancer.

```
legend.labs = c("hormon", "control")) +
ggtitle("Survival Curve of Hormon and Control group")
```

Survival Curve of Hormon and Control group



Chemo

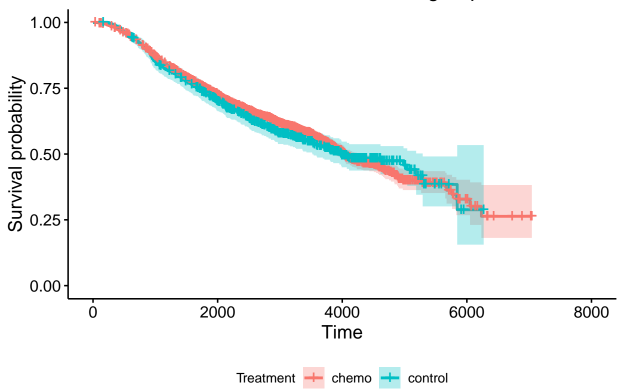
```
logrank3 <- survdiff(Surv(dtime, death) ~ chemo, data = rotterdam)
logrank3</pre>
```

```
## Call:
## survdiff(formula = Surv(dtime, death) ~ chemo, data = rotterdam)
##
## N Observed Expected (0-E)^2/E (0-E)^2/V
## chemo=0 2402    1014    1024    0.0963    0.495
## chemo=1 580    258    248    0.3977    0.495
##
## Chisq= 0.5 on 1 degrees of freedom, p= 0.5
```

logrank3\$pvalue

[1] 0.4818191

Survival Curve of Chemo and Control group



Results

Discussion

How our results compare with past research

Conclusion

References

—Note this reference is in MLA format—

Simon, Noah et al. "Regularization Paths for Cox's Proportional Hazards Model via Coordinate Descent." Journal of statistical software vol. 39,5 (2011): 1-13. doi:10.18637/jss.v039.i05