PSTAT-175-Final-Project

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```
library(readr)
library(tidyverse)
library(survival)
library(survminer)
library(ggplot2)
library(knitr)
```

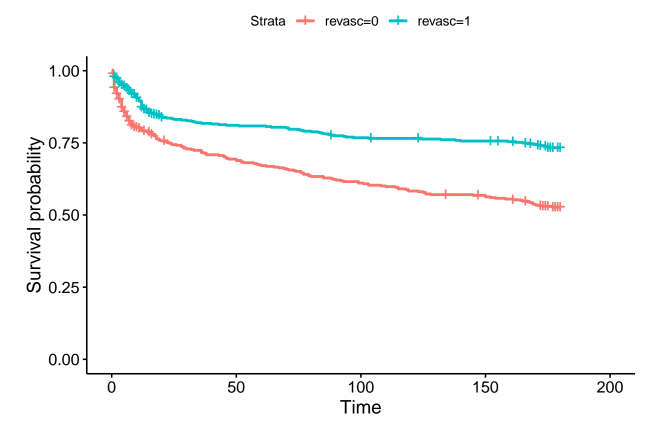
```
# Read in data
GRACE1000 <- read_table("GRACE1000.dat", col_names = FALSE)
#GRACE1000 <- read_table("~/Downloads/GRACE1000.dat", col_names = FALSE)
GRACE1000 <- GRACE1000 %>% select(-X10)
colnames(GRACE1000) <- c("id", "days", "death", "revasc", "revascdays", "los", "age", "sysbp", "stchange")</pre>
```

Final Project

1: Introduction

We will be analyzing the "GRACE1000" dataset from Hosmer and Lemenshow and May (2008), which contains data on 1000 patients who were part of a study on the effectiveness of revascularization by the Global Registry of Acute Coronary Events (GRACE). Our failure time of interest is the follow up time when the researchers checked-in with the patients, and our event of interest is whether the patient died during the follow-up period. Our main covariate of interest will be whether the patient had the revascularization procedure performed on them, which is coded by the "revasc" variable with value 1 if the patient had the procedure and 0 if they did not. The main research question of interest is whether revascularization is associated with higher survival rates after completing the procedure.

Here is a plot of the survival functions of patients who had the revascularization procedure in blue and patients who did not have the procedure done in red:



We can see that at first glance, patients who had revascularization done appear to have a higher survival probability compared to patients who did not have the procedure done. We will further explore this data with the tools we have learned:

2: Model Fitting

For this project, we plan to use AIC and BIC as model selection criteria to determine the most appropriate predictors for survival outcomes in the GRACE1000 dataset. Since this dataset contains multiple potential predictors, including age, systolic blood pressure, ST changes, and revascularization status, it's essential to balance model complexity with explanatory power. AIC and BIC provide objective measures to compare different Cox proportional hazards models, where AIC focuses more on prediction and BIC favors simpler, more interpretable models. This approach ensures that the final model explaining the time to death in relation to revascularization is both statistically robust and practically meaningful.

```
# Forward Stepwise Selection
# Possible covariates are revasc, sysbp, age, stchange

age.mod <- coxph(Surv(days,death) ~ age, data = GRACE1000)
sysbp.mod <- coxph(Surv(days,death) ~ sysbp, data = GRACE1000)
stchange.mod <- coxph(Surv(days,death) ~ stchange, data = GRACE1000)
revasc.mod <- coxph(Surv(days,death) ~ revasc, data = GRACE1000)

kable(AIC(age.mod, sysbp.mod, stchange.mod, revasc.mod))</pre>
```

	df	AIC
age.mod	1	4151.309
sysbp.mod	1	4245.999
stchange.mod	1	4252.286
revasc.mod	1	4231.999

kable(BIC(age.mod, sysbp.mod, stchange.mod, revasc.mod))

df	BIC
1	4155.090
1	4249.780
1	4256.067
1	4235.780
	1 1

Upon analyzing the Cox proportional hazards models, the results indicate that age alone provides the best balance between model fit and simplicity, as evidenced by the lowest AIC value of 4151.309 and BIC value of 4155.090. In comparison, models using systolic blood pressure (AIC = 4245.999), ST changes (AIC = 4252.286), and revascularization status (AIC = 4231.999) each demonstrate higher AIC values, suggesting that these individual predictors are less informative in explaining survival outcomes. The differences between AIC values are considerable, with the age model clearly outperforming others by a margin exceeding 4-10 AIC units, reinforcing the strength of age as a predictor.

```
age.sysbp.mod <- coxph(Surv(days,death) ~ age + sysbp, data = GRACE1000)
age.stchange.mod <- coxph(Surv(days,death) ~ age + stchange, data = GRACE1000)
age.revasc.mod <- coxph(Surv(days,death) ~ age + revasc, data = GRACE1000)
kable(AIC(age.sysbp.mod, age.stchange.mod, age.revasc.mod))</pre>
```

	df	AIC
age.sysbp.mod	2	4131.179
age.stchange.mod	2	4136.352
age.revasc.mod	2	4136.017

kable(BIC(age.sysbp.mod, age.stchange.mod, age.revasc.mod))

	df	BIC
age.sysbp.mod	2	4138.741
age.stchange.mod	2	4143.913
age.revasc.mod	2	4143.579

In this analysis, we compared three multivariable Cox models incorporating age with additional predictors: systolic blood pressure, ST change, and revascularization status. The model that included age and systolic blood pressure yielded the lowest AIC value (4131.179), indicating the best fit among the multivariable models tested. Models incorporating age with ST change (AIC = 4136.352) and age with revascularization (AIC = 4136.017) had slightly higher AIC values, suggesting marginally inferior fits. The consistently low AIC values across all models reinforce the importance of age as a key predictor while indicating that including systolic blood pressure may offer a slight improvement over the other two variables when combined with age.

```
sysbp.age.stchange.mod <- coxph(Surv(days,death) ~ sysbp + age + stchange, data = GRACE1000)
sysbp.age.revasc.mod <- coxph(Surv(days,death) ~ sysbp + age + revasc, data = GRACE1000)
kable(AIC(sysbp.age.stchange.mod, sysbp.age.revasc.mod))</pre>
```

	df	AIC
sysbp.age.stchange.mod	3	4117.788
sysbp.age.revasc.mod	3	4116.536

kable(BIC(sysbp.age.stchange.mod, sysbp.age.revasc.mod))

	df	BIC
sysbp.age.stchange.mod	3	4129.130
sysbp.age.revasc.mod	3	4127.879

```
# See if stchange lowers AIC and BIC
all.mod <- coxph(Surv(days,death) ~ stchange + sysbp + revasc + age, data = GRACE1000)
kable(data.frame(Metric = c("AIC", "BIC"), Value = c(AIC(all.mod), BIC(all.mod))))</pre>
```

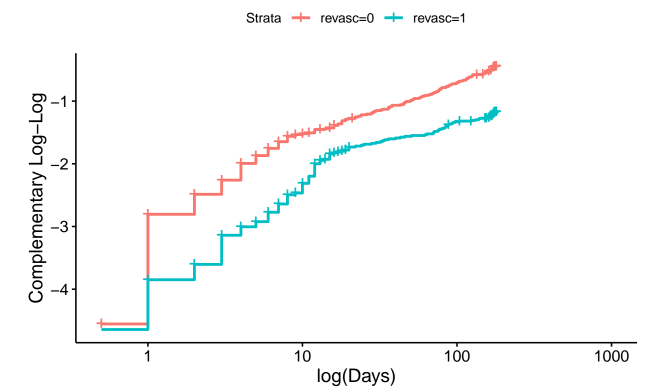
Metric	Value
AIC	4100.721
BIC	4115.844

After fitting a comprehensive Cox proportional hazards model incorporating all considered predictors: ST change, systolic blood pressure, revascularization, and age, the resulting AIC value was 4100.721 and BIC

was 4115.844, both lower than those from the previous models. This indicates that including all variables yields the best-fitting model based on forward stepwise selection criteria. These results suggest that while age alone and age in combination with other predictors offer substantial predictive power, the full model provides the most accurate explanation of survival probability in this dataset, balancing model complexity with fit.

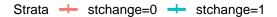
3: Check Proportional Hazards Assumptions

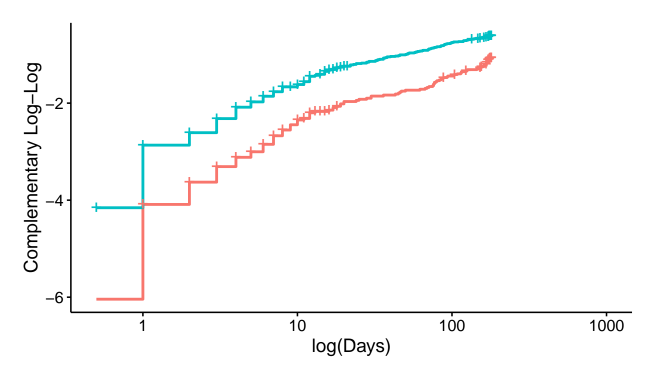
Log-Log Plot by Revasc



In this Complementary Log-Log plot of the treatment and control groups, the curves for patients who underwent revascularization (revasc=1) and those who did not (revasc=0) are presented. The relatively parallel nature of the curves suggests that the proportional hazards assumption holds reasonably well for the revascularization variable, as the log-log curves for the two groups do not cross and maintain a fairly consistent vertical separation throughout the range of time. Overall, the log-log plot supports the use of the Cox proportional hazards model with revascularization as a predictor

Log-Log Plot by Stchange





also very parallel lines, no assumptions violated

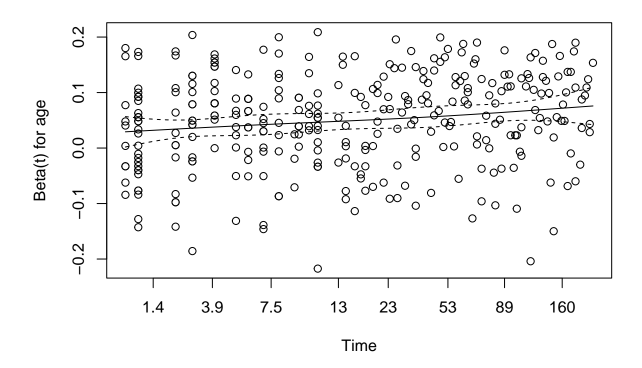
The Complementary Log-Log plot for Stchange also displays curves that do not cross into higher time values, meaning the proportional hazards assumption also holds here with the different Stchange values.

ZPH Plot for Age:

kable(as.data.frame(cox.zph(age.mod)\$table))

	chisq	df	p
age	7.758003	1	0.0053475
GLOBAL	7.758003	1	0.0053475

plot(cox.zph(age.mod))



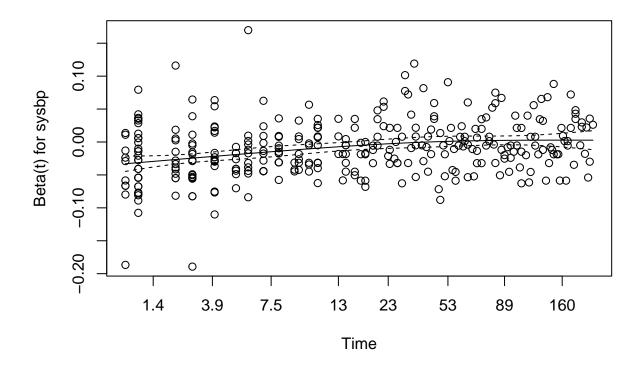
The statistical ouput shows a p-value of 0.0053 for both age and the global test, which is highly significant (p < 0.05). This indicates a violation of the proportional hazards assumption for age, meaning that the effect of age on the hazard is not constant over time. This zph plot visualizes Schoenfeld residuals for age over time in the Cox model. The residuals appear randomly scattered around the horizontal axis, and the smoothed curve does not display a clear trend. This pattern suggests that the proportional hazards assumption for age holds reasonably well. Although we can't see it in the graph, with the p-value being that low we will still assume the ph assumptions are being violated.

ZPH Plot for Systolic Blood Pressure:

kable(as.data.frame(cox.zph(sysbp.mod)\$table))

	chisq	df	р
sysbp	33.18016	1	0
GLOBAL	33.18016	1	0

plot(cox.zph(sysbp.mod))



Here we see Systolic Blood Pressure also violating the proportional hazards assumption with a p_value of 8.4e-09. Once again it doesn't appear to have any clear correlation in the graph but with a p_value that low we can not ignore that there is a clear time-varying effect in the model.

4: Conclusions

Our main scientific question of interest is whether the revascularization procedure significantly increases patients' survival probabilities, and while we have a general idea, we can take a look at the results of our models:

summary(revasc.mod)

```
## Call:
   coxph(formula = Surv(days, death) ~ revasc, data = GRACE1000)
##
##
     n= 1000, number of events= 324
##
##
##
             coef exp(coef) se(coef)
                                            z Pr(>|z|)
                               0.1144 -6.249 4.13e-10 ***
##
   revasc -0.7149
                      0.4892
                      '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
  Signif. codes:
##
##
          exp(coef) exp(-coef) lower .95 upper .95
##
             0.4892
                          2.044
                                    0.391
                                              0.6122
  revasc
##
```

```
## Concordance= 0.589 (se = 0.014)
## Likelihood ratio test= 40.46
                                  on 1 df,
                                             p=2e-10
## Wald test
                         = 39.05
                                  on 1 df,
                                             p = 4e - 10
## Score (logrank) test = 40.73
                                  on 1 df,
                                             p=2e-10
exp(coef(revasc.mod) + c(-1.96, 1.96) * sqrt(revasc.mod$var[1,1]))
## [1] 0.3909648 0.6122096
exp(confint(revasc.mod))
              2.5 %
                        97.5 %
##
## revasc 0.3909664 0.6122071
```

The summary of the revasc.mod Cox model indicates that revascularization is a highly significant predictor of survival (p < 0.0001), with a hazard ratio of 0.4892 and a 95% confidence interval of [0.391, 0.612]. This means that patients who underwent revascularization were approximately 51% less likely to die compared to those who did not undergo the procedure. The hazard ratio being less than 1 signifies a protective effect of revascularization.

```
summary(stchange.mod)
```

```
## Call:
   coxph(formula = Surv(days, death) ~ stchange, data = GRACE1000)
##
     n= 1000, number of events= 324
##
##
##
                coef exp(coef) se(coef)
                                                z Pr(>|z|)
                                    0.1185 4.38 1.19e-05 ***
## stchange 0.5189
                          1.6802
##
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
##
              exp(coef) exp(-coef) lower .95 upper .95
## stchange
                    1.68
                              0.5952
                                           1.332
                                                       2.119
##
## Concordance= 0.57 (se = 0.013)
## Likelihood ratio test= 20.17
                                       on 1 df,
                                                    p = 7e - 06
## Wald test
                            = 19.19
                                       on 1 df,
                                                    p=1e-05
## Score (logrank) test = 19.62 on 1 df,
                                                    p = 9e - 06
\exp(\operatorname{coef}(\operatorname{stchange.mod}) + \operatorname{c}(-1.96, 1.96) * \operatorname{sqrt}(\operatorname{stchange.mod}\operatorname{var}[1, 1]))
## [1] 1.332036 2.119339
```

```
## 2.5 % 97.5 %
```

exp(confint(stchange.mod))

stchange 1.332041 2.11933

Patients with an ST change on their initial ECG have a 68% higher hazard of mortality compared to those without such deviations, after controlling for time. This is a clinically meaningful result, as it highlights that ST changes detected early in an ECG are strongly associated with worse survival outcomes. The statistical significance (p < 0.00001) and a confidence interval for the hazard ratio of [1.332, 2.119] (an interval greater than 1) suggest that this association is reliable.

5: Advanced Methods

Binning Revascdays

In the GRACE1000 dataset, "revascdays" is coded as the censor time if the patient did not have the revascularization procedure done. We hypothesize that this could interfere with our models' performance because we have many observations that did not have revascularization done and were censored at the end of the study (180 days), so the range of values from 0 to 180 may not be the best way to analyze the covariate. We will instead experiment with binning "revascdays" into a categorical variable called "daysGroup" which will take on ranges of values from "revascdays" so we can analyze the covariate as a set of ranges rather than continuous. We will set nearly equally spaced intervals for the bins, but after binning them this way, we can analyze the results and look for other ways to group the values:

```
binnedGRACE1000 <- GRACE1000 %>% mutate(
   daysGroup = case_when(
        (revascdays >= 0 & revascdays <= 2) ~ "0-2 Days",
        (revascdays >= 3 & revascdays <= 6) ~ "3-6 Days",
        (revascdays >= 7 & revascdays <= 10) ~ "7-10 Days",
        (revascdays >= 11 & revascdays <= 14) ~ "11-14 Days",
        revascdays > 14 ~ "14+ Days"
    )
)
```

Now that we have our new data, lets see the results of our Cox PH model on the binned covariate:

```
summary(coxph(Surv(days,death) ~ daysGroup, data = binnedGRACE1000))
```

```
## coxph(formula = Surv(days, death) ~ daysGroup, data = binnedGRACE1000)
##
     n= 1000, number of events= 324
##
##
##
                          coef exp(coef) se(coef)
                                                        z Pr(>|z|)
## daysGroup11-14 Days -0.1825
                                   0.8332
                                            0.2277 - 0.801
                                                            0.4229
                                            0.1406 -2.266
## daysGroup14+ Days
                       -0.3186
                                   0.7272
                                                            0.0235 *
                                                            0.0522 .
## daysGroup3-6 Days
                        0.3400
                                   1.4050
                                            0.1752 1.941
## daysGroup7-10 Days
                      -0.0802
                                   0.9229
                                            0.1761 - 0.456
                                                            0.6487
##
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' 1
##
                       exp(coef) exp(-coef) lower .95 upper .95
## daysGroup11-14 Days
                          0.8332
                                      1.2002
                                                0.5332
                                                          1.3019
## daysGroup14+ Days
                          0.7272
                                      1.3752
                                                0.5520
                                                          0.9579
## daysGroup3-6 Days
                          1.4050
                                      0.7118
                                                0.9967
                                                          1.9805
## daysGroup7-10 Days
                          0.9229
                                      1.0835
                                                0.6536
                                                          1.3032
##
## Concordance= 0.592 (se = 0.014)
## Likelihood ratio test= 14.58 on 4 df,
                                             p=0.006
## Wald test
                        = 15.27
                                 on 4 df,
                                             p=0.004
## Score (logrank) test = 15.6 on 4 df,
                                            p=0.004
```

This model is using the "0-2 Days" group as a baseline which we know because it does not show up in the summary output. This means that all of the hazard ratios seen in this output are in relation to the "0-2

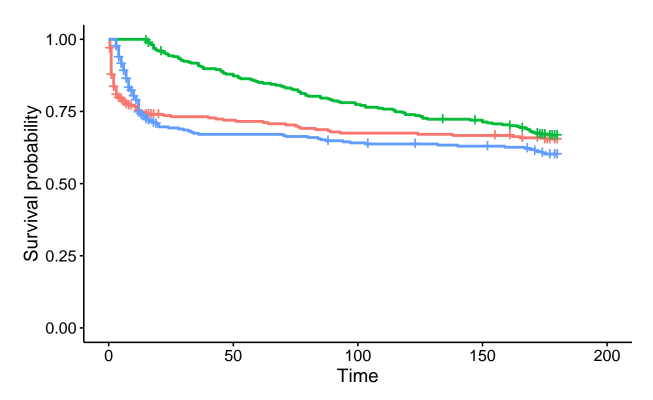
Days" till revascularization group. We see that the "14+ Days" group has a statistically significant difference in survival probabilities according to this model with a hazard ratio of 0.7272 compared to the "0-2 Days" group. We also see that the "3-6 Days" group is the only group to have a higher hazard rate than the "0-2 Days" group, so I would like to change the bins to explore this further:

```
binnedGRACE1000 <- GRACE1000 %>% mutate(
  daysGroup = case when(
    (revascdays >= 0 & revascdays <= 2) ~ "0-2 Days",
    (revascdays >= 3 & revascdays <= 14) ~ "3-14 Days",
    revascdays >= 15 ~ "15+ Days"
  )
)
summary(coxph(Surv(days,death) ~ daysGroup, data = binnedGRACE1000))
## Call:
## coxph(formula = Surv(days, death) ~ daysGroup, data = binnedGRACE1000)
##
##
    n= 1000, number of events= 324
##
##
                         coef exp(coef) se(coef)
                                                      z Pr(>|z|)
## daysGroup15+ Days -0.3176
                                 0.7279
                                          0.1406 - 2.259
                                                           0.0239 *
## daysGroup3-14 Days 0.0446
                                 1.0456
                                          0.1357 0.329
                                                           0.7425
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
                      exp(coef) exp(-coef) lower .95 upper .95
## daysGroup15+ Days
                         0.7279
                                    1.3738
                                                         0.9589
                                              0.5526
## daysGroup3-14 Days
                         1.0456
                                    0.9564
                                              0.8014
                                                         1.3643
##
## Concordance= 0.552 (se = 0.013)
## Likelihood ratio test= 8.61 on 2 df,
                                           p=0.01
## Wald test
                        = 8.32 on 2 df,
                                           p=0.02
## Score (logrank) test = 8.4 on 2 df,
                                          p=0.02
```

In this new binned data we see a similar outcome where the control group (revascdays = 180) has a statistically significant difference in hazard rates than the "0-2 Days" group. Lets take a look at a graph of this result:

```
ggsurvplot(survfit(Surv(days,death) ~ daysGroup, data = binnedGRACE1000))
```





I believe that the higher volume of censored observations in the 0-2 and 3-14 day groups along with the correlation of the patients needing the revascularization procedure could be affecting the model to calculate a higher hazard rate for patients with early revascularization, but this also is not the best way to deal with the revascularization times. Now that we have explored the "revascdays" covariate, let us focus on our covariate of interest, "revasc".

Treating Revascularization as a Time-Dependent Covariate

When trying to answer the question of whether the revascularization procedure is associated with higher survival rates, we run into a problem with our data in that not all of our observations had the revascularization procedure on day 0 or the first day they were a part of the study. This means that if a patient had the revascularization done on the 5th day (revascdays = 5), then our models are counting the first 5 days as this patient already having the procedure done, when in reality it has not happened until later. To fix this issue with the data, we will treat "revasc" as a binary time-dependent covariate that changes at some point in the study. To implement the time-varying covariate, we will split every observation that had the revascularization procedure done into 2 observations where the first observation has a start time of 0 and a stop time of the day they had the procedure done, and the second observation will have a start time of the day the procedure was done and a stop time of when the observation was censored. This way our models will properly understand that patients have not had the procedure done until the day given by "revascdays", and therefore are not a part of the treatment group until then.

We will manipulate the data to achieve this:

```
library(dplyr)
library(tidyr)
```

```
grace2 <- GRACE1000 %>%
  mutate(
   start_1 = 0,
   stop 1 = pmin(revascdays, days, na.rm = TRUE),
   status_1 = ifelse(days <= revascdays, death, 0),</pre>
   revasc_status_1 = 0, # Renamed for clarity
   start 2 = revascdays,
   stop_2 = ifelse(revasc == 1, days, NA),
   status_2 = ifelse(revasc == 1 & days > revascdays, death, NA),
   revasc_status_2 = 1 # Renamed for clarity
 ) %>%
 pivot_longer(
   cols = c(start_1, stop_1, status_1, revasc_status_1, start_2, stop_2, status_2, revasc_status_2),
   names_to = c("variable", "interval"),
   names_pattern = ([a-z]+)_([12])
  ) %>%
 pivot_wider(
   names_from = variable,
   values_from = value
 ) %>%
 filter(!is.na(stop)) %>%
 arrange(id, interval) %>%
 filter(!(start==0 & stop==0))
# Check output
kable(head(grace2))
```

id	days	death	revasc	revascdays	s los	age	sysbp	stchange	e interval	start	stop	status	revasc_status
1	180	0	0	180	9	28	107	1	1	0	180	0	0
2	5	0	1	0	5	32	121	1	2	0	5	0	1
3	2	0	0	2	2	33	150	0	1	0	2	0	0
4	5	0	1	2	5	35	172	0	1	0	2	0	0
4	5	0	1	2	5	35	172	0	2	2	5	0	1
5	180	0	1	9	10	35	106	0	1	0	9	0	0

Cox PH to show the effect of revascularization now that we have accounted for time

```
tvc <- coxph(Surv(start, stop, status) ~ revasc, data = grace2)</pre>
summary(tvc)
## Call:
## coxph(formula = Surv(start, stop, status) ~ revasc, data = grace2)
##
##
   n= 1353, number of events= 319
##
      (12 observations deleted due to missingness)
##
##
             coef exp(coef) se(coef)
                                          z Pr(>|z|)
## revasc -0.6920 0.5006
                             0.1150 -6.019 1.75e-09 ***
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
          exp(coef) exp(-coef) lower .95 upper .95
##
                         1.998
            0.5006
                                  0.3996
                                            0.6271
## revasc
##
## Concordance= 0.585 (se = 0.014)
## Likelihood ratio test= 37.41
                                on 1 df.
                                            p=1e-09
## Wald test
                        = 36.23
                                on 1 df,
                                            p = 2e - 09
## Score (logrank) test = 37.69 on 1 df,
                                            p=8e-10
```

In this Cox proportional hazards model, we analyzed the effect of revascularization on patient survival, accounting for the time-varying nature of the treatment by splitting the dataset at the point when revascularization was performed. The model revealed a significant association between revascularization and reduced hazard of death, with a hazard ratio of approximately 0.50 (95% CI: 0.40, 0.63), indicating that after revascularization, the risk of death was roughly half that of patients who had not yet undergone the procedure. This approach effectively modeled the time-dependent impact of revascularization by allowing the treatment's hazard to vary based on when it was performed during the follow-up period. Overall, these results suggest that revascularization is associated with a substantial and statistically significant improvement in survival, which becomes apparent when we account for the dynamic nature of treatment over time.

Test to see if we want to use the other covariates:

##

```
zph_test <- cox.zph(coxph(Surv(start, stop, status) ~ revasc + age + sysbp + stchange, , data = grace2)
kable(as.data.frame(zph_test$table))</pre>
```

	chisq	df	p
revasc	1.100055	1	0.2942539
age	7.441267	1	0.0063745
sysbp	24.201243	1	0.0000009
stchange	9.519219	1	0.0020333
GLOBAL	43.802813	4	0.0000000

We would not want to use any of the other covariates in this case as they all fail to meet the PH assumptions even while splitting on the day of revascularization

Following the textbook - "suppose the clinicians on the GRACE study wanted to determine whether the effect depended on the number of days from admission to revascularization."

```
tvc_interaction<- coxph(Surv(start, stop, status) ~ revasc_status * revascdays, data = grace2)

## Warning in Surv(start, stop, status): Stop time must be > start time, NA

## created

summary(tvc_interaction)

## Call:
## coxph(formula = Surv(start, stop, status) ~ revasc_status * revascdays,
## data = grace2)
```

```
##
     n= 1353, number of events= 319
##
      (12 observations deleted due to missingness)
##
##
                                   coef exp(coef)
                                                   se(coef)
                                                                   z Pr(>|z|)
## revasc_status
                             -3.119073
                                         0.044198
                                                   0.241760 -12.902
                                                                        <2e-16 ***
                             -0.023352
                                         0.976919
                                                   0.001717 -13.597
                                                                       <2e-16 ***
  revascdays
##
                             0.029756
                                        1.030203
  revasc status:revascdays
                                                   0.021072
                                                                        0.158
##
## Signif. codes:
                    0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
##
##
                             exp(coef) exp(-coef) lower .95 upper .95
                                                                0.07099
                                0.0442
                                           22.6254
##
  revasc_status
                                                     0.02752
                                0.9769
   revascdays
                                            1.0236
                                                     0.97364
                                                                0.98021
##
   revasc_status:revascdays
                                1.0302
                                            0.9707
##
                                                     0.98852
                                                                1.07364
##
## Concordance= 0.818 (se = 0.014)
  Likelihood ratio test= 308.3
                                  on 3 df,
                                              p=<2e-16
## Wald test
                         = 212.5
                                   on 3 df,
                                              p = < 2e - 16
                                  on 3 df,
## Score (logrank) test = 282.6
                                              p=<2e-16
```

In this interaction Cox proportional hazards model, we examined whether the effect of revascularization on survival depended on the number of days from admission to the revascularization procedure. The model included revasc_status, revascdays, and their interaction term. The analysis revealed that both revasc_status and revascdays were highly significant with p-values less than 2e-16, indicating that both receiving revascularization and the number of days from admission to the procedure independently affect survival. The interaction term (revasc_status:revascdays) was not statistically significant (p = 0.158), suggesting that the impact of revascularization on survival does not vary significantly based on how many days after admission it was performed. The hazard ratio for the interaction term was approximately 1.03 (95% CI: 0.97–1.07), meaning the modification effect is negligible. In summary, while both revascularization and time since admission to revascularization individually influence survival, their interaction does not significantly alter the risk of death in this dataset.

Conclusion

This analysis of the GRACE1000 dataset examined the relationship between revascularization and survival in patients with acute coronary events. Initial exploratory analysis, including Kaplan-Meier curves, suggested that patients undergoing revascularization had better survival probabilities compared to those who did not. Cox proportional hazards models, selected using AIC and BIC criteria, consistently highlighted age as a strong predictor of mortality, with lower AIC and BIC values compared to other covariates such as systolic blood pressure, ST changes, and revascularization status. However, a comprehensive model including all predictors provided the best overall fit, balancing complexity and explanatory power. The analysis of the proportional hazards assumption revealed that while age and systolic blood pressure violated the PH assumption, revascularization and ST changes appeared to satisfy it, supporting their use in Cox models. Advanced modeling approaches addressed the time-dependent nature of revascularization by splitting patient records at the time of treatment. This adjustment showed that revascularization significantly reduced the hazard of death (HR ~0.50, p < 0.0001), affirming its protective effect even after accounting for the timing of the procedure. Further exploration, including the binning of "revascdays" and testing for interaction effects, revealed that while both revascularization and time from admission to revascularization independently influenced survival, their interaction was not statistically significant. This suggests that the survival benefit of revascularization was consistent regardless of when it was administered during hospitalization. In summary, this project demonstrates that revascularization is associated with a substantial and statistically significant improvement in survival, supporting its role as an effective treatment in acute coronary events.

References

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