Multivariate Analysis for the Behavioral Sciences, Second Edition (Chapman and Hall/CRC, 2019)

Solutions to Exercises of Chapter 7: Survival Analysis

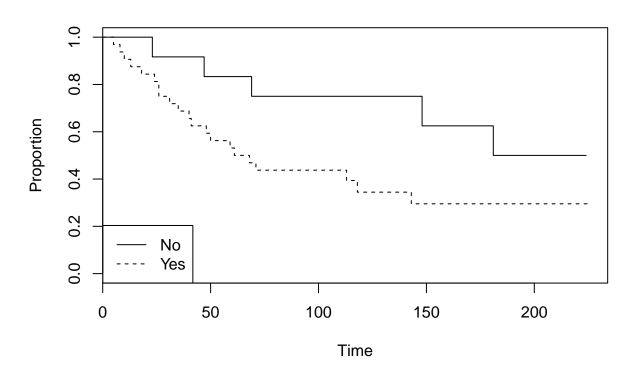
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Solutions

Exercise 7.2

```
brcancer <- structure(list(</pre>
 time = c(23, 47, 69, 70, 100, 101, 148, 181, 198, 208, 212, 224, 5, 8, 10, 13, 18,
        24, 26, 26, 31, 35, 40, 41, 48, 50, 59, 61, 68, 71, 76, 105, 107, 109,
        113, 116, 118, 143, 145, 162, 188, 212, 217, 225),
 1, 1, 1, 1, 1, 1, 0, 0, 0, 0, 1, 0, 1, 1, 0, 0, 0, 0, 0, 0),
 2L, 2L, 2L, 2L, 2L),
            .Label = c("No", "Yes"), class = "factor")),
 .Names = c("time", "event", "metastized"),
row.names = c(NA, -44L), class = "data.frame")
str(brcancer)
## 'data.frame':
               44 obs. of 3 variables:
            : num 23 47 69 70 100 101 148 181 198 208 ...
            : num 1 1 1 0 0 0 1 1 0 0 ...
## $ metastized: Factor w/ 2 levels "No", "Yes": 1 1 1 1 1 1 1 1 1 1 ...
head(brcancer)
   time event metastized
## 1
## 2
     47
                   No
## 3
     69
                   No
     70
           0
## 4
                   No
## 5 100
           0
                   No
## 6 101
                   No
attach(brcancer)
library(survival)
#plot of survival function
plot(survfit(Surv(time, event) ~ metastized), xlab = "Time", ylab = "Proportion", lty = 1:2)
legend("bottomleft", c("No", "Yes"), lty=1:2)
```



```
#logrank test
survdiff(Surv(time,event) ~ metastized)
## survdiff(formula = Surv(time, event) ~ metastized)
##
                   N Observed Expected (O-E)^2/E (O-E)^2/V
##
                                   9.2
## metastized=No 12
                            5
                                            1.91
                                                      3.04
                           21
## metastized=Yes 32
                                  16.8
                                            1.05
                                                      3.04
##
   Chisq= 3 on 1 degrees of freedom, p=0.0813
detach(brcancer)
```

Exercise 7.3

```
glioma <- structure(list(</pre>
 no. = c(1L, 2L, 3L, 4L, 5L, 6L, 7L, 8L, 9L, 10L, 11L, 12L, 13L, 14L, 15L, 16L,
         17L, 18L, 19L, 1L, 2L, 3L, 4L, 5L, 6L, 7L, 8L, 9L, 10L, 11L, 12L, 13L,
         14L, 15L, 16L, 17L, 18L),
 age = c(41L, 45L, 48L, 54L, 40L, 31L, 53L, 49L, 36L, 52L, 57L, 55L, 70L, 39L,
         40L, 47L, 58L, 40L, 36L, 27L, 32L, 53L, 46L, 33L, 19L, 32L, 70L, 72L,
         46L, 44L, 83L, 57L, 71L, 61L, 65L, 50L, 42L),
 sex = structure(c(1L, 1L, 2L, 2L, 1L, 2L, 2L, 2L, 2L, 2L, 2L, 1L, 2L, 1L, 1L,
                  1L, 2L, 1L, 2L, 2L, 2L, 1L, 2L, 1L, 1L, 1L, 2L, 2L, 2L, 2L,
                  1L, 1L, 1L, 2L, 2L, 2L, 1L),
       .Label = c("Female", "Male"), class = "factor"),
 1L, 1L, 1L, 1L, 1L, 1L, 2L, 2L, 2L, 2L, 2L, 2L, 1L,
                        .Label = c("GBM", "Grade3"), class = "factor"),
 2L, 2L, 2L, 2L, 2L, 1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L,
                    1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L, 1L),
         .Label = c("Control", "RIT"), class = "factor");
 event = c(TRUE, FALSE, FALSE, FALSE, FALSE, TRUE, FALSE, FALSE, FALSE, FALSE,
          FALSE, FALSE, TRUE, TRUE, FALSE, FALSE, TRUE, TRUE, TRUE, TRUE, TRUE,
           TRUE, TRUE, FALSE, FALSE, TRUE, TRUE, TRUE, TRUE, TRUE, TRUE, TRUE,
           TRUE, TRUE, TRUE, TRUE, TRUE),
 time = c(53L, 28L, 69L, 58L, 54L, 25L, 51L, 61L, 57L, 57L, 50L, 43L, 20L, 14L,
          36L, 59L, 31L, 14L, 36L, 34L, 32L, 9L, 19L, 50L, 48L, 8L, 8L, 11L, 12L,
          15L, 5L, 8L, 8L, 6L, 14L, 13L, 25L)),
 .Names = c("no.", "age", "sex", "histology", "group", "event", "time"),
 row.names = c("1", "2", "3", "4", "5", "6", "7", "8", "9", "10", "11", "12",
               "13", "14", "15", "16", "17", "18", "19", "20", "21", "22", "23",
                   "25", "26", "27", "28", "29", "30", "31", "32", "33", "34",
               "35", "36", "37"), class = "data.frame")
str(glioma)
## 'data.frame':
                  37 obs. of 7 variables:
             : int 1 2 3 4 5 6 7 8 9 10 ...
## $ no.
## $ age
              : int 41 45 48 54 40 31 53 49 36 52 ...
             : Factor w/ 2 levels "Female", "Male": 1 1 2 2 1 2 2 2 2 2 ...
## $ sex
## $ histology: Factor w/ 2 levels "GBM", "Grade3": 2 2 2 2 2 2 2 2 2 ...
## $ group
             : Factor w/ 2 levels "Control", "RIT": 2 2 2 2 2 2 2 2 2 2 ...
##
   $ event
              : logi TRUE FALSE FALSE FALSE TRUE ...
## $ time
              : int 53 28 69 58 54 25 51 61 57 57 ...
attach(glioma)
library(survival)
glioma_cox <- coxph(Surv(time, event) ~ age + sex + histology + group)</pre>
summary(glioma_cox)
## Call:
## coxph(formula = Surv(time, event) ~ age + sex + histology + group)
```

```
##
##
    n= 37, number of events= 23
##
                      coef exp(coef) se(coef)
##
                                                  z Pr(>|z|)
## age
                   0.04344
                             1.04440 0.02152 2.019
                                                      0.0435 *
                   0.06411
                             1.06621 0.49755 0.129
                                                      0.8975
## sexMale
## histologyGrade3 -1.48480
                             0.22655 0.61826 -2.402
                                                      0.0163 *
                             ## groupRIT
                  -2.44441
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
                  exp(coef) exp(-coef) lower .95 upper .95
## age
                    1.04440
                               0.9575
                                        1.00127
                                                   1.0894
                    1.06621
                                0.9379
                                        0.40210
                                                   2.8272
## sexMale
## histologyGrade3
                    0.22655
                                4.4141
                                        0.06744
                                                   0.7611
## groupRIT
                    0.08678
                               11.5238
                                        0.02776
                                                   0.2713
##
## Concordance= 0.855 (se = 0.068)
## Rsquare= 0.636
                   (max possible= 0.981 )
## Likelihood ratio test= 37.39 on 4 df,
                                          p=1.497e-07
## Wald test
                       = 27.32 on 4 df,
                                          p=1.714e-05
## Score (logrank) test = 40.28 on 4 df,
                                          p=3.788e-08
detach(glioma)
```

Required CI given in summary above - hazard on radioimmunotherapy (RIT) is between about 3% and 27% of hazard on standard therapy.

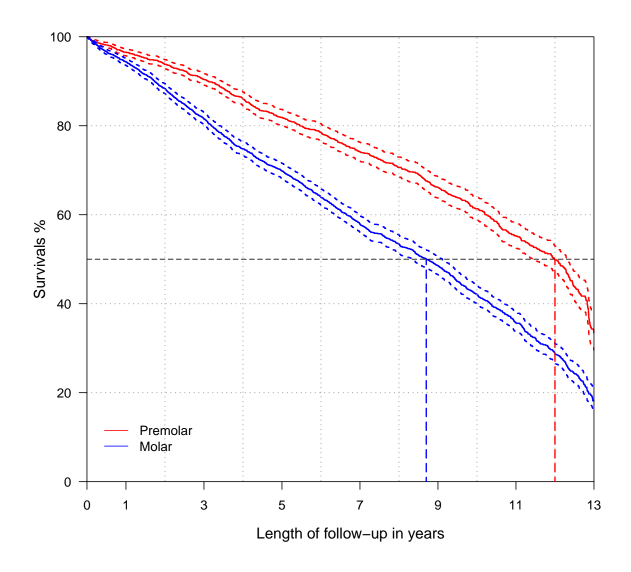
You should plot the survival curves of RIT and standard treatments.

Exercise 7.4

```
teeth <- read.table("data/teeth.txt", sep = '\t', header = TRUE)
head(teeth)
     ID Gender Age ISOcode Reint Survyear Type4 Material Type2
## 1 1
             1 30
                        24
                               0
                                     2.510
                                               1
## 2 2
             1 30
                        26
                               0
                                    2.510
                                               3
                                                        2
                                                              2
## 3 3
             1 30
                        16
                                    0.301
                                               3
                                                        2
                                                              2
                               1
             2 30
                                    0.545
                                                        2
## 4 4
                        14
                                               1
                                                              1
                               1
## 5 5
             2 30
                        15
                                    9.033
                                               1
                                                        2
                                                              1
                               1
## 6 6
             1 28
                        15
                               1
                                   12.526
                                               1
                                                        2
                                                              1
#tail(teeth)
# Set maximum longevity to 13 years and censor all at the end of the last year:
teeth$Survyear[teeth$Survyear > 13] <- 13</pre>
teeth$Reint[teeth$Survyear == 13] <- 0</pre>
```

As an example, Figure 2 of the original article (Palotie et al., 2017), "Survival (%) of restorations placed in posterior teeth (n = 5542) of 25- to 30-year-olds illustrated as medians and Kaplan-Meier curves with 95% confidence intervals by type of tooth", was created by *Anna Eronen* as follows:

```
library(survival)
#install.packages("plotrix")
library(plotrix)
plot(survfit(Surv(Survyear, Reint) ~ strata(Type2), data = teeth), conf.int = "both",
     col = c("red", "blue"), lwd = 1.7, axes = FALSE, yscale = 100,
     main = "", ylab = "", xlab = "", cex.axis = 1.2)
title(xlab = "Length of follow-up in years", line = 1.7, cex.lab = 1.2)
title(ylab = "Survivals %", line = 2.5, cex.lab = 1.2)
ablineclip(v=2, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(v=4, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(v=6, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(v=8, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(v=10, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(v=12, y1=0, y2=1.0, col="darkgrey", lty=3, lwd=1.4)
ablineclip(h=0.2, x1=0, x2=13, col="darkgrey", lty=3, lwd=1.4)
ablineclip(h=0.4, x1=0, x2=13, col="darkgrey", lty=3, lwd=1.4)
ablineclip(h=0.6, x1=0, x2=13, col="darkgrey", lty=3, lwd=1.4)
ablineclip(h=0.8, x1=0, x2=13, col="darkgrey", lty=3, lwd=1.4)
ablineclip(h=1.0, x1=0, x2=13, col="black",lty=1,lwd=1.0)
ablineclip(v=13, y1=0,y2=1,col="black",lty=1,lwd=1.0)
axis(1, at=c(0,1,3,5,7,9,11,13), labels=c(0,1,3,5,7,9,11,13), pos=0, las=1)
axis(2, at=c(0.0,0.2,0.4,0.6,0.8,1.0), labels=c(0,20,40,60,80,100), pos=0, las=2)
legend(x=0.15, y=0.15, c("Premolar", "Molar"), col=c("red", "blue"), lty=c(1,1), bty="n")
ablineclip(h=0.5, x1=0, x2=13, col="black", lty=5)
ablineclip(v=12.0, y1=0, y2=0.5, col="red", lty=5, lwd=1.7)
ablineclip(v=8.7, y1=0, y2=0.5, col="blue", lty=5, lwd=1.7)
```



```
# example of a log-rank test:
kmtest.teeth <- survdiff(Surv(Survyear, Reint) ~ Type2, data = teeth)</pre>
kmtest.teeth
## Call:
## survdiff(formula = Surv(Survyear, Reint) ~ Type2, data = teeth)
##
              N Observed Expected (0-E)^2/E (0-E)^2/V
##
                              1060
                                        92.8
## Type2=1 2216
                     746
                                                   167
                                        73.3
## Type2=2 3326
                    1654
                              1340
                                                   167
    Chisq= 167 on 1 degrees of freedom, p= 0
```