Survival Analysis with R: Exercises

Exercise set 1

Take a look at the built in colon dataset. If you type ?colon it'll ask you if you wanted help on the colon dataset from the survival package, or the colon operator. Click "Chemotherapy for Stage B/C colon cancer", or be specific with ?survival::colon. This dataset has survival and recurrence information on 929 people from a clinical trial on colon cancer chemotherapy. There are two rows per person, indidicated by the event type (etype) variable – etype==1 indicates that row corresponds to recurrence; etype==2 indicates death.

First, let's filter the data to only include the survival data, not the recurrence data. Let's call this new object colondeath. The filter() function is in the dplyr library, which you can get by running library(dplyr). If you don't have dplyr you can use the base subset() function instead.

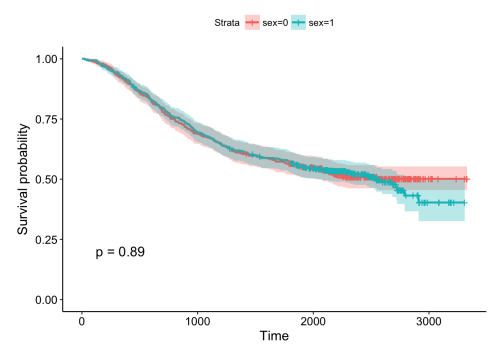
```
# using dplyr::filter
library(dplyr)
colondeath <- filter(colon, etype==2)

# Or, using base subset()
# colondeath <- subset(colon, etype==2)
head(colondeath)</pre>
```

- 1. Look at the help for ?colon again. How are sex and status coded? How is this different from the lung data?
- 2. Using survfit(Surv(..., ...,)~..., data=colondeath), create a survival curve separately for males versus females. Call the resulting object sfit. Run a summary() on this object, showing time points 0, 500, 1000, 1500, and 2000. Do males or females appear to fair better over this time period?

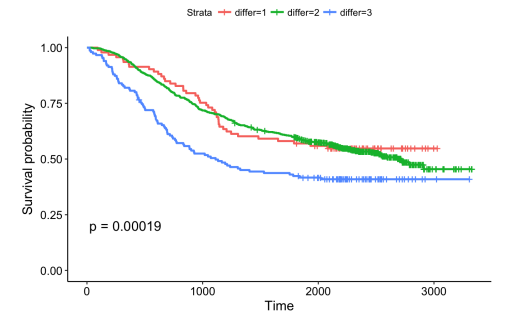
```
##
                    sex=0
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
       0
            445
                       0
                            1.000 0.0000
                                                   1.000
                                                                 1.000
            381
                            0.856 0.0166
                                                                 0.889
##
     500
                      64
                                                   0.824
##
    1000
            306
                      75
                            0.688 0.0220
                                                   0.646
                                                                 0.732
    1500
            265
##
                      40
                            0.598
                                    0.0232
                                                   0.554
                                                                 0.645
##
    2000
            218
                      22
                            0.547 0.0236
                                                   0.503
                                                                 0.596
##
##
                    sex=1
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
            484
                       0
                            1.000 0.0000
                                                   1.000
                                                                 1.000
       0
##
     500
            418
                      65
                            0.866
                                   0.0155
                                                   0.836
                                                                 0.897
            335
##
    1000
                      83
                            0.694
                                   0.0210
                                                   0.654
                                                                 0.736
##
    1500
            287
                      46
                            0.598
                                   0.0223
                                                   0.556
                                                                 0.644
##
    2000
            238
                      25
                            0.545 0.0227
                                                   0.503
                                                                 0.592
```

3. Using the survminer package, plot a Kaplan-Meier curve for this analysis with confidence intervals and showing the p-value. See ?ggsurvplot for help. Is there a significant difference between males and females?

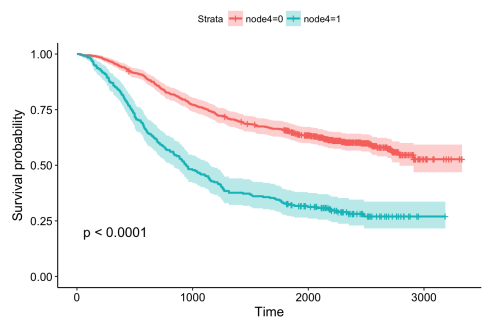


- 4. Create Kaplan-Meier plot stratifying by:
 - a. The extent of differentiation (well, moderate, poor), showing the p-value.
 - b. Whether or not there was detectable cancer in >=4 lymph nodes, showing the p-value and confidence bands.

Survival by tumor differentiation



Survival by involvement in >=4 lymph nodes



Exercise set 2

Let's go back to the colon cancer dataset. Remember, you created a colondeath object in the first exercise that only includes survival (etype==2), not recurrence data points. See ?colon for more information about this dataset.

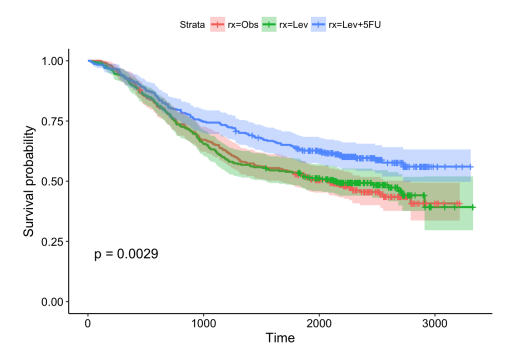
1. Take a look at levels(colondeath\$rx). This tells you that the rx variable is the type of treatment the patient was on, which is either nothing (coded Obs, short for Observation), Levamisole (coded Lev), or Levamisole + 5-fluorouracil (coded Lev+5FU). This is a factor variable coded with these levels, in that order. This means that Obs is treated as the baseline group, and other groups are dummy-coded to represent the respective group.

rx	Lev	Lev+5FU
Obs	0	0
Lev	1	0
Lev+5FU	0	1

2. Run a Cox proportional hazards regression model against this rx variable. How do you interpret the result? Which treatment seems to be significantly different from the control (Observation)?

```
## coef exp(coef) se(coef) z p
## rxLev -0.0266  0.9737  0.1103 -0.24 0.8092
## rxLev+5FU -0.3717  0.6896  0.1188 -3.13 0.0017
##
## Likelihood ratio test=12.2 on 2 df, p=0.0023
## n= 929, number of events= 452
```

3. Show the results using a Kaplan-Meier plot, with confidence intervals and the p-value.



4. Fit another Cox regression model accounting for age, sex, and the number of nodes with detectable cancer. Notice the test statistic on the likelihood ratio test becomes much larger, and the overall model becomes more significant. What do you think accounted for this increase in our ability to model survival?

```
coef exp(coef) se(coef)
##
## rxLev
             -0.08007
                         0.92305
                                  0.11161 -0.72 0.47312
   rxLev+5FU -0.40253
                         0.66863
                                  0.12054 -3.34 0.00084
              0.00533
                         1.00535
                                  0.00405
                                          1.32 0.18739
##
   age
## sex
             -0.02826
                         0.97214
                                  0.09573 -0.30 0.76786
              0.09275
                         1.09719
                                  0.00887 10.46 < 2e-16
## nodes
##
## Likelihood ratio test=87.8 on 5 df, p=0
   n= 911, number of events= 441
##
      (18 observations deleted due to missingness)
```

Exercise set 3

The "KIPAN" cohort (in KIPAN.clinical) is the pan-kidney cohort, consisting of KICH (chromaphobe renal cell carcinoma), KIRC (renal clear cell carcinoma), and KIPR (papillary cell carcinoma). The KIPAN.clinical has KICH.clinical, KIRC.clinical, and KIPR.clinical all combined.

1. Using survivalTCGA(), create a new object called clinkid using the KIPAN.clinical cohort. For the columns to extract, get both the disease code and the patient's gender (extract.cols=c("admin.disease_code", "patient.gender")). The first few rows will look like this.

```
times bcr_patient_barcode patient.vital_status admin.disease_code
## 1
      1158
                   TCGA-KL-8323
                                                                      kich
## 2
      4311
                   TCGA-KL-8324
                                                     0
                                                                      kich
## 3
       725
                   TCGA-KL-8325
                                                     1
                                                                      kich
## 4
      3322
                   TCGA-KL-8326
                                                                      kich
```

```
## 5
      3553
                   TCGA-KL-8327
                                                                        kich
## 6
     3127
                   TCGA-KL-8328
                                                      0
                                                                        kich
     patient.gender
##
## 1
              female
## 2
              female
## 3
              female
## 4
                male
## 5
              female
## 6
                male
```

2. The xtabs() command will produce tables of counts for categorical variables. Here's an example for how to use xtabs() for the built-in colon cancer dataset, which will tell you the number of samples split by sex and by treatment.

xtabs(~rx+sex, data=colon)

```
## rx 0 1
## Obs 298 332
## Lev 266 354
## Lev+5FU 326 282
```

Use the same command to examine how many samples you have for each kidney sample type, separately by sex.

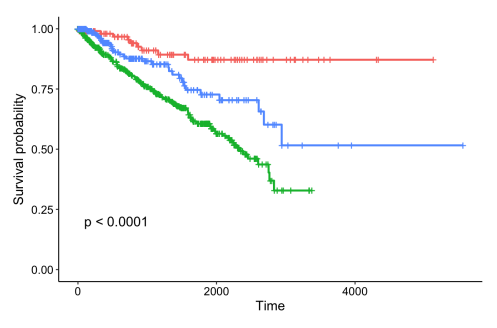
```
## patient.gender
## admin.disease_code female male
## kich 51 61
## kirc 191 346
## kirp 76 212
```

3. Run a Cox PH regression on the cancer type and gender. What's the effect of gender? Is it significant? How does survival differ by each type? Which has the worst prognosis?

```
##
                             coef exp(coef) se(coef)
                                                          z
                                                       4.62 3.9e-06
## admin.disease_codekirc 1.5929
                                     4.9179
                                               0.3450
## admin.disease_codekirp
                          0.9962
                                     2.7080
                                               0.3807
                                                       2.62
                                                             0.0089
## patient.gendermale
                          -0.0628
                                     0.9391
                                               0.1484 -0.42 0.6721
##
## Likelihood ratio test=39.4 on 3 df, p=1.4e-08
## n= 937, number of events= 203
```

- 4. Create survival curves for each different subtype.
 - a. Produce a Kaplan-Meier plot.
 - b. Show survival tables each year for the first 5 years.





Call: survfit(formula = Surv(times, patient.vital_status) ~ admin.disease_code, ## data = clinkid) ## ## admin.disease_code=kich ## time n.risk n.event survival std.err lower 95% CI upper 95% CI ## 0 111 0 1.000 0.0000 1.000 1.000 ## 365 86 2 0.980 0.0144 0.952 1.000 ## 730 72 2 0.954 0.0226 0.911 0.999 3 ## 1095 54 0.910 0.0329 0.848 0.977 ## 1460 44 1 0.893 0.0366 0.824 0.967 ## 1825 38 1 0.871 0.0415 0.794 0.957 ## ## admin.disease_code=kirc ## time n.risk n.event survival std.err lower 95% CI upper 95% CI 536 0 ## 0 1.000 0.0000 1.000 1.000 365 385 49 0.895 0.0142 0.924 ## 0.868 ## 730 313 32 0.816 0.0186 0.781 0.853 ## 1095 250 26 0.744 0.0217 0.703 0.788 ## 1460 181 20 0.678 0.0243 0.633 0.728 ## 1825 112 16 0.606 0.0277 0.554 0.663 ## ## admin.disease_code=kirp ## time n.risk n.event survival std.err lower 95% CI upper 95% CI ## 0 288 0 1.000 0.0000 1.000 1.000 ## 365 145 10 0.941 0.0182 0.906 0.977 730 100 ## 8 0.877 0.0278 0.824 0.933 ## 1095 67 2 0.853 0.0316 0.793 0.917 1460 3 ## 54 0.810 0.0388 0.737 0.889 ## 1825 36 5 0.727 0.0495 0.636 0.831