# Survival Analysis, Algorithmic Fairness, and COMPAS Recidivism Algorithm Case Study

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# **Consulting Skills Focus**

- In real life consulting, the client will frequently understand the data and surrounding research better than the data scientist/statistician.
- Analysis should center around a well-defined research question that drive the analysis and the data should be able to provide insight on the question of interest.
- Human bias and data analysis: We all have bias. This can influence data analysis. A data analyst, we should do our best to objectively present the data. When necessary to make assumptions, state them explicitly.
- Publication Bias example: Researchers frequently want "positive results." Usually this means significant p-values. Variable selection is a simple way to change p-values, p-hacking. It's common to need to change variables in a model be it should be done a principled way.
- Question: Have you heard of field of algorithmic fairness (https://en.wikipedia.org/wiki/Fairness\_(machine\_learning))?
  - a. yes
  - o b. no

# Case Study Background

• US has more inmates, proportional to population size, than any other country. While Black Americans make up 13% of the total US population, they account for 40% of incarcerated population in the US.

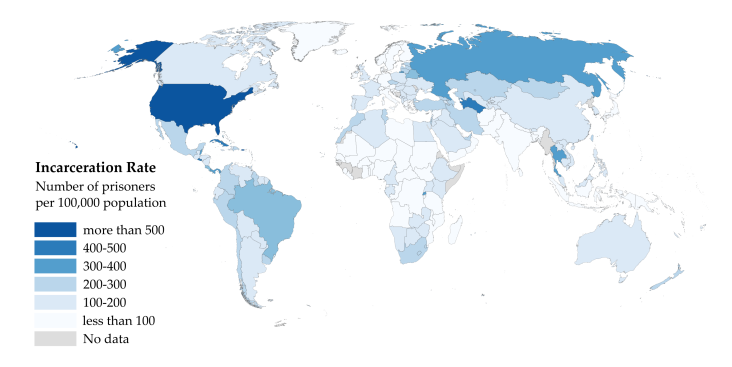


Image from Wikipedia (https://en.wikipedia.org/wiki/Incarceration in the United States#/media/File:Prisoners world map png2.png)

- In the US justice system, machine learning algorithms are sometimes used to assess a criminal defendant's risk of recidivism (arrest due to committing a future crime) are being used.
- Correctional Offenders Management Profiling for Alternative Sanctions (COMPAS) is the most widespread of these algorithms.
- Its goal according to COMPAS creators: assess "not just risk but also nearly two dozen so-called "criminogenic needs" that relate to the major theories of criminality, including "criminal personality," "social isolation," "substance abuse" and "residence/stability." Defendants are ranked low, medium or high risk in each category."
- In 2014, then U.S. Attorney General Eric Holder warned that the risk scores might be injecting bias into the courts. He called for the U.S. Sentencing Commission to study their use. "Although these measures were crafted with the best of intentions, I am concerned that they inadvertently undermine our efforts to ensure individualized and equal justice," he said, adding, "they may exacerbate unwarranted and unjust disparities that are already far too common in our criminal justice system and in our society."
- The questionnaire (https://www.documentcloud.org/documents/2702103-Sample-Risk-Assessment-COMPAS-CORE.html) for determining COMPAS does not directly ask for race, but some people question inherent racial bias in the algorithm.
- The COMPAS algorithm is proprietary and not available.

• More information in a 2016 ProPublica article (https://www.propublica.org/article/machine-bias-risk-assessments-in-criminal-sentencing).

# **Data**

- ProPublica requested two years of COMPAS scores from Broward County Sheriff's Office in Florida
- Discarded all but pre-trial COMPAS score assessments
- · ProPublica matched COMPAS scores with criminal records from Broward County Clerk's Office website
- COMPAS score screening date and (original) arrest date frequently differed. If they are too far apart, that may indicate an error. The days b screening arrest variable gives this difference in days.
- is\_recid is rearrest at any time. two\_year\_recid is rearrest within two years. Here, -1 indicates a
   COMPAS record could not be found and should probably be discarded
- COMPAS generates a general score, decile\_score, (1, 2,...,10) where 1 indicates a low risk and 10 indicates a high risk of recidivism. There is also a violence score as well, v\_decile\_score.

```
dat<-read.csv("./compas-scores.csv")
dim(dat)</pre>
```

## [1] 11757 47

names(dat)

```
[1] "id"
                                    "name"
##
##
    [3] "first"
                                    "last"
                                    "sex"
##
    [5] "compas screening date"
                                    "age"
##
    [7] "dob"
##
    [9] "age cat"
                                    "race"
## [11] "juv fel count"
                                    "decile score"
  [13] "juv misd count"
                                    "juv other count"
  [15] "priors count"
                                    "days b screening arrest"
## [17] "c_jail_in"
                                    "c jail out"
## [19] "c case number"
                                    "c offense date"
## [21] "c arrest date"
                                    "c days from compas"
## [23] "c_charge_degree"
                                    "c charge desc"
## [25] "is recid"
                                    "num r cases"
## [27] "r case number"
                                    "r charge degree"
                                    "r_offense_date"
## [29] "r_days_from_arrest"
## [31] "r_charge desc"
                                    "r jail in"
## [33] "r_jail_out"
                                    "is violent recid"
## [35] "num vr cases"
                                    "vr case number"
## [37] "vr_charge_degree"
                                    "vr_offense_date"
## [39] "vr charge desc"
                                    "v type of assessment"
## [41] "v_decile_score"
                                    "v_score_text"
## [43] "v screening date"
                                    "type of assessment"
## [45] "decile_score.1"
                                    "score_text"
## [47] "screening date"
```

### head(dat[,1:12])

```
##
     id
                               first
                       name
                                             last compas screening date sex
## 1
          miguel hernandez
                             miguel
                                       hernandez
                                                              2013-08-14 Male
      1
##
              michael ryan michael
                                                              2014-12-31 Male
                                            ryan
##
  3
      3
               kevon dixon
                               kevon
                                           dixon
                                                              2013-01-27 Male
                                           philo
                                                              2013-04-14 Male
##
  4
      4
                   ed philo
                                  ed
  5
               marcu brown
                                                              2013-01-13 Male
##
                               marcu
                                           brown
      6 bouthy pierrelouis
                             bouthy pierrelouis
                                                              2013-03-26 Male
##
            dob age
                              age_cat
                                                   race juv_fel_count decile_score
##
## 1 1947-04-18
                  69 Greater than 45
                                                                     0
                                                  Other
                                                                                   1
## 2 1985-02-06
                              25 - 45
                                              Caucasian
                                                                     0
                                                                                   5
                  31
## 3 1982-01-22
                              25 - 45 African-American
                                                                                   3
## 4 1991-05-14
                 24
                        Less than 25 African-American
                                                                     0
                                                                                   4
                        Less than 25 African-American
## 5 1993-01-21
                  23
                                                                     0
                                                                                   8
## 6 1973-01-22
                              25 - 45
                 43
                                                  Other
                                                                     0
                                                                                   1
```

```
summary(dat)
```

```
## id name first last
```

```
##
                     Length: 11757
                                                              Length: 11757
    Min.
           :
                 1
                                         Length:11757
    1st Qu.: 2940
                     Class :character
                                         Class :character
                                                              Class :character
##
##
    Median: 5879
                     Mode
                          :character
                                         Mode :character
                                                             Mode :character
##
    Mean
            : 5879
##
    3rd Qu.: 8818
##
    Max.
            :11757
##
##
    compas screening date
                                                    dob
                                sex
                                                                         age
##
    Length:11757
                           Length: 11757
                                                Length:11757
                                                                    Min.
                                                                           :18.00
##
    Class :character
                           Class :character
                                                Class :character
                                                                    1st Qu.:25.00
##
    Mode :character
                           Mode :character
                                                Mode :character
                                                                    Median :32.00
##
                                                                    Mean
                                                                            :35.14
##
                                                                    3rd Ou.:43.00
##
                                                                    Max.
                                                                            :96.00
##
##
      age cat
                            race
                                             juv fel count
                                                                  decile score
##
    Length: 11757
                        Length: 11757
                                            Min.
                                                    : 0.00000
                                                                 Min.
                                                                        :-1.000
                                                                 1st Qu.: 2.000
##
    Class :character
                        Class :character
                                            1st Qu.: 0.00000
##
    Mode :character
                        Mode :character
                                            Median : 0.00000
                                                                 Median : 4.000
##
                                            Mean
                                                    : 0.06158
                                                                 Mean
                                                                        : 4.371
##
                                             3rd Ou.: 0.00000
                                                                 3rd Ou.: 7.000
##
                                            Max.
                                                    :20.00000
                                                                 Max.
                                                                        :10.000
##
    juv misd count
                        juv other count
                                              priors count
##
                                                               days b screening arrest
            : 0.00000
                                : 0.00000
                                                    : 0.000
                                                              Min.
##
    Min.
                        Min.
                                            Min.
                                                                      :-597.000
##
    1st Ou.: 0.00000
                        1st Ou.: 0.00000
                                            1st Ou.: 0.000
                                                               1st Ou.:
                                                                        -1.000
##
    Median : 0.00000
                        Median : 0.00000
                                            Median : 1.000
                                                              Median: -1.000
##
            : 0.07604
                                                    : 3.082
                                                                      -0.878
    Mean
                        Mean
                                : 0.09356
                                            Mean
                                                              Mean
                        3rd Qu.: 0.00000
##
    3rd Ou.: 0.00000
                                            3rd Ou.: 4.000
                                                               3rd Ou.: -1.000
##
    Max.
           :13.00000
                                :17.00000
                                                    :43.000
                                                                      :1057.000
                        Max.
                                            Max.
                                                               Max.
##
                                                              NA's
                                                                      :1180
##
     c jail in
                         c jail out
                                            c case number
                                                                 c offense date
##
    Length:11757
                        Length:11757
                                            Length: 11757
                                                                 Length: 11757
    Class :character
                        Class :character
                                            Class :character
                                                                 Class :character
##
##
    Mode :character
                        Mode :character
                                            Mode :character
                                                                 Mode :character
##
##
##
##
##
                        c days from compas c charge degree
    c arrest date
                                                                 c charge desc
##
    Length:11757
                        Min.
                                    0.00
                                            Length: 11757
                                                                 Length:11757
##
    Class :character
                        1st Ou.:
                                    1.00
                                            Class :character
                                                                 Class :character
##
    Mode :character
                        Median:
                                    1.00
                                            Mode :character
                                                                 Mode :character
##
                        Mean
                                   63.59
##
                        3rd Qu.:
                                    2.00
                                :9485.00
##
                        Max.
                        NA's
                                :742
##
##
       is recid
                       num r cases
                                       r case number
                                                           r charge degree
##
    Min.
            :-1.0000
                       Mode:logical
                                       Length:11757
                                                           Length: 11757
```

```
1st Qu.: 0.0000
##
                       NA's:11757
                                      Class :character
                                                          Class :character
##
    Median : 0.0000
                                      Mode
                                             :character
                                                          Mode
                                                                :character
##
    Mean
           : 0.2538
##
    3rd Qu.: 1.0000
           : 1.0000
##
    Max.
##
##
    r days from arrest r offense date
                                           r charge desc
                                                                r jail in
                        Length: 11757
##
   Min.
           : -1.00
                                           Length:11757
                                                               Length:11757
##
    1st Qu.: 0.00
                        Class :character
                                           Class :character
                                                               Class :character
##
    Median:
             0.00
                        Mode :character
                                           Mode :character
                                                               Mode :character
##
    Mean
           : 20.41
##
    3rd Qu.: 1.00
           :993.00
##
    Max.
    NA's
           :9297
##
##
     r jail out
                        is violent recid
                                          num vr cases
                                                          vr case number
   Length:11757
                        Min.
##
                               :0.00000
                                          Mode:logical
                                                          Length: 11757
    Class :character
                                          NA's:11757
##
                        1st Ou.:0.00000
                                                          Class :character
    Mode :character
                        Median :0.00000
                                                          Mode :character
##
##
                        Mean
                               :0.07502
##
                        3rd Qu.:0.00000
##
                        Max.
                               :1.00000
##
                                                               v_type_of_assessment
##
    vr charge degree
                        vr offense date
                                           vr charge desc
    Length:11757
                        Length: 11757
                                           Length: 11757
##
                                                               Length: 11757
    Class :character
                        Class :character
                                           Class :character
                                                               Class :character
##
##
    Mode :character
                        Mode :character
                                           Mode :character
                                                               Mode :character
##
##
##
##
##
                     v score text
                                         v screening date
    v decile score
                                                             type of assessment
##
    Min.
           :-1.000
                     Length:11757
                                         Length:11757
                                                             Length: 11757
##
    1st Qu.: 1.000
                     Class :character
                                         Class :character
                                                             Class :character
    Median : 3.000
                     Mode :character
                                         Mode :character
                                                             Mode :character
##
##
   Mean
           : 3.571
##
    3rd Ou.: 5.000
##
    Max.
           :10.000
##
##
    decile score.1
                                         screening date
                      score text
##
   Min.
           :-1.000
                                         Length: 11757
                     Length:11757
    1st Qu.: 2.000
                                         Class :character
##
                     Class :character
##
    Median : 4.000
                     Mode :character
                                         Mode :character
##
           : 4.371
    Mean
    3rd Ou.: 7.000
##
##
    Max.
           :10.000
##
```

Are there multiple rows per person?

```
length(unique(dat$id))
 ## [1] 11757
 length(dat$id)
 ## [1] 11757
 length(unique(dat$name))
 ## [1] 11584
 sort(table(dat$name), decreasing=TRUE)[1:8]
 ##
 ##
         carlos vasquez
                                  john brown michael cunningham
                                                                         robert taylor
 ##
 ##
        anthony jackson
                               anthony smith
                                                gregory williams
                                                                           james brown
 ##
                                                                                      3
   • To me, these seem like common names, so it could be a coincidence, but I would check with the client as
     due diligence
   • What else could we do to check?
What about other demographics?
 table(dat$sex)
```

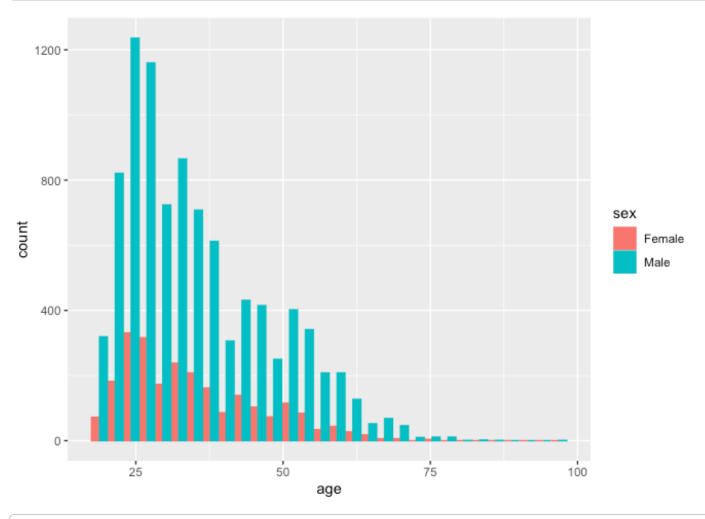
```
##
## Female Male
## 2421 9336
```

```
table(dat$sex)/sum(!is.na(dat$sex))*100
```

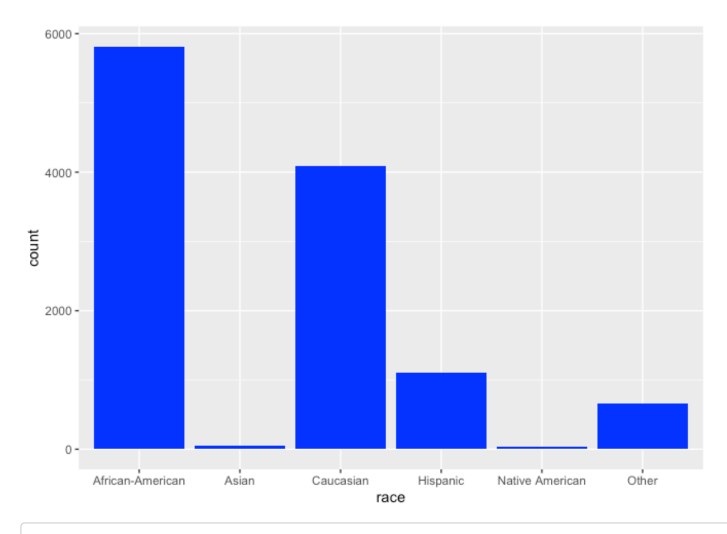
```
##
## Female Male
## 20.59199 79.40801
```

```
library(ggplot2)
ggplot(dat, aes(x=age, color=sex, fill=sex)) +
  geom_histogram(position="dodge")
```

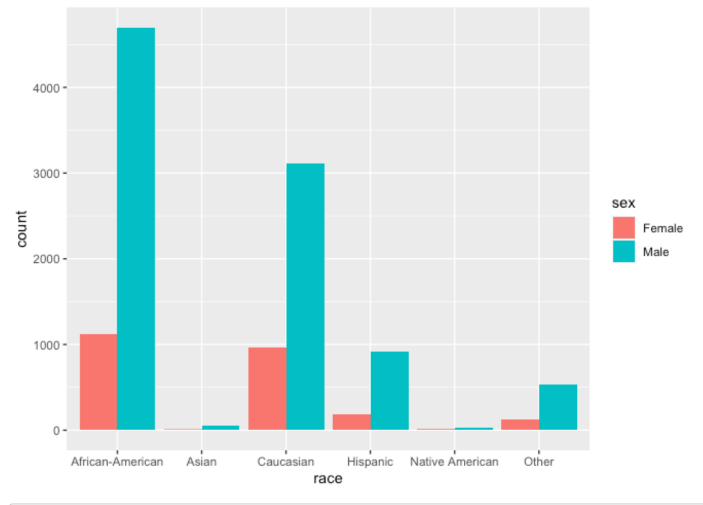
```
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
```



```
ggplot(dat, aes(race)) +
  geom_bar(fill='blue')
```

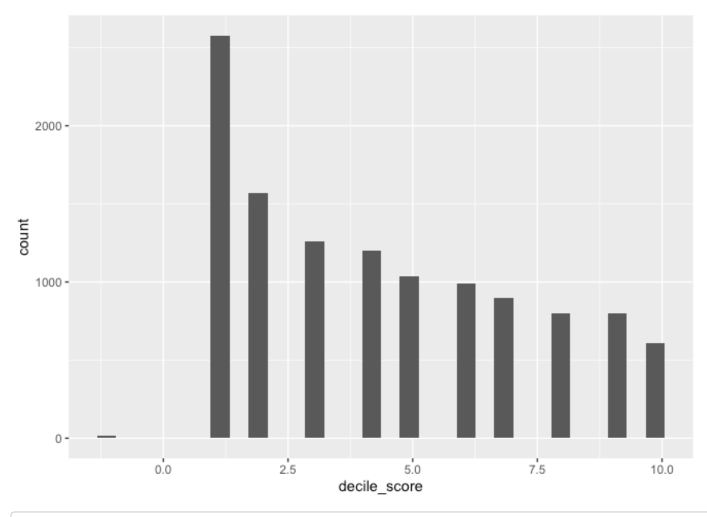


ggplot(dat, aes(x=race, fill=sex)) +
 geom\_bar(position='dodge')



```
ggplot(dat, aes(decile_score)) +
  geom_histogram()
```

## `stat\_bin()` using `bins = 30`. Pick better value with `binwidth`.



```
table(!is.na(dat$decile_score))
```

```
##
## TRUE
## 11757
```

### General recommendations:

- Look at the raw data and different plots of the data before doing any modeling.
- Look for missing data and for values that might not make sense.
- Make sure you understand what observations (rows) are included in the data and which of those observations serve your data analysis goals
- Try to understand what the variables (columns) represent and which ones will serve your data analysis goals

# Quantifying racial bias

• Before doing any analysis, let's look at recidivism, COMPAS, and race

```
df <- dat[dat$is_recid != -1,]
sum(is.na(df$race))

## [1] 0

sum(is.na(df$is_recid))

## [1] 0

table(df$race, df$is recid)[,2]/t(table(df$race))*100</pre>
```

```
##
## African-American Asian Caucasian Hispanic Native American Other
## [1,] 39.53827 20.75472 28.52279 25.86720 36.11111 24.79871
```

Above is the recidivism rate by race

COMPAS also gave Black Americans greater scores on average:

```
tapply(df$decile_score, df$race, mean)
## African-American
                                Asian
                                              Caucasian
                                                                 Hispanic
##
           5.326850
                             2.735849
                                               3.647459
                                                                  3.313181
##
    Native American
                                 Other
##
           4.805556
                             2.813205
```

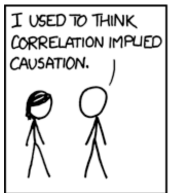
Is this the best way to present this information?

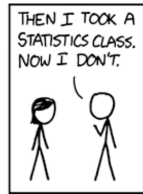
# How to model algorithmic bias?

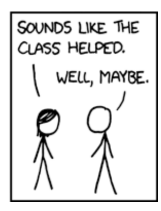
- What does bias mean here?
- Would COMPAS give someone a greater score solely due to being Black or some other demographic, without changing anything else?
- Stated differently, if two people have the same risk of recidivism, with race being their only difference, will the algorithm score them differently
- Remember COMPAS doesn't ask for race directly
- What does race affecting recidivism mean?
  - Incorrect: Someones race affects their behavior
  - Correct: The effect of race living in a racially biased society

- How could we quantify bias in this case? Are race and COMPAS still associated after taking recidivism into account?
- It is tempting to use decile\_score ~ is\_recid + race
  - This regression could answer the following: Is race helpful for predicting COMPAS score while controlling for recidivism?
  - If race were significant, that would indicate that race contributes to COMPAS beyond recidivism itself,
     so COMPAS would be racially biased
  - But, this is not a valid model because decile\_score is collected before is\_recid

# Causation and Collider Bias

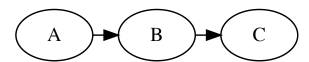






XKCD causal comic

# **Bayesian Network 1:**



Mental Model: Think of a dataset where *A*, *B*, *C* are collected

- A = Alcohol
- B = Hangover
- C = Miss Class

Question: What would a regression model of C ~ A + B yield?

- a. Both A and B should be statistically significant
- b. Only A should be statistically significant
- c. Only B should be statistically significant
- d. Neither A nor B should be statistically significant

```
set.seed(1234)
size <- 1000
A <- 6*rnorm(size)+50
B <- -2*A - 25 + rnorm(size)
C <- 5*B + 3 +rnorm(size)
summary(lm(C~A+B))</pre>
```

```
##
## Call:
## lm(formula = C \sim A + B)
##
## Residuals:
##
       Min
                  10
                      Median
                                    3Q
                                            Max
## -3.13161 -0.71957 0.03478 0.70215 3.05316
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.96001
                          0.87456
                                     2.241
                                             0.0252 *
## A
               -0.07084
                          0.06532 - 1.085
                                             0.2784
## B
                4.96310
                        0.03270 151.761
                                            <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.013 on 997 degrees of freedom
## Multiple R-squared: 0.9997, Adjusted R-squared:
## F-statistic: 1.739e+06 on 2 and 997 DF, p-value: < 2.2e-16
```

Question: What about this regression model: c ~ A?

- a. A should be statistically significant
- b. A should not be statistically significant

```
summary(lm(C~A))
```

```
##
## Call:
## lm(formula = C \sim A)
##
## Residuals:
##
      Min
                  Median
               1Q
                               3Q
                                      Max
  -15.9753 -3.4048 -0.0059
##
                            3.2714 16.5278
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
  ##
              -9.95096
                        0.02627 -378.80 <2e-16 ***
## A
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4.969 on 998 degrees of freedom
## Multiple R-squared: 0.9931, Adjusted R-squared: 0.9931
## F-statistic: 1.435e+05 on 1 and 998 DF, p-value: < 2.2e-16
```

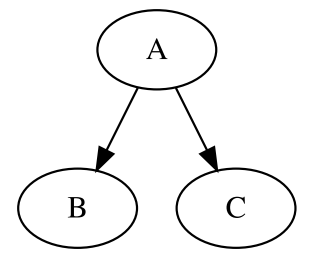
· Coefficient estimates:

$$C = 5B + 3 + \epsilon_B$$
  
= 5(-2A - 25 + \epsilon\_A) + 3 + \epsilon\_B  
= -10A - 122 + 5\epsilon\_A + \epsilon\_B

Question: Does this coefficient and intercept estimate make sense?

- a. yes
- b. nope

# **Bayesian Network 2:**



Mental Model:

- A = Smoker
- B = Yellow Teeth
- C = Cancer

Question: What would a regression model of C ~ A + B yield?

- a. Both A and B should be statistically significant
- b. Only A should be statistically significant
- c. Only B should be statistically significant
- d. Neither A nor B should be statistically significant

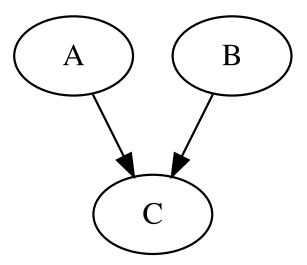
```
set.seed(1234)
size <- 1000
A <- 6*rnorm(size)+50
B <- -2*A - 25 + rnorm(size)
C <- 2*A +5 +rnorm(size)
summary(lm(C~A+B))</pre>
```

```
##
## Call:
## lm(formula = C \sim A + B)
##
##
  Residuals:
       Min
                  10
                      Median
##
                                    3Q
                                            Max
  -3.13161 -0.71957 0.03478 0.70215 3.05316
##
##
##
  Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
  (Intercept) 3.96001
                        0.87456 4.528 6.67e-06 ***
##
                           0.06532 29.533 < 2e-16 ***
## A
                1.92916
## B
               -0.03690
                           0.03270 - 1.128
                                              0.259
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 1.013 on 997 degrees of freedom
## Multiple R-squared: 0.9929, Adjusted R-squared: 0.9929
## F-statistic: 6.996e+04 on 2 and 997 DF, p-value: < 2.2e-16
```

Question: What about this regression model: c ~ A?

- a. A should be statistically significant
- b. A should not be statistically significant

# **Bayesian Network 3:**



### Mental Model:

- A =Allergies
- B = Flu
- C = Sinus

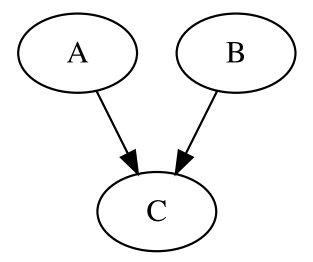
Question: What would a regression model of  $\, C \sim A + B \,$  yield?

- a. Both A and B should be statistically significant
- b. Only A should be statistically significant
- c. Only *B* should be statistically significant
- d. Neither A nor B should be statistically significant

```
set.seed(1234)
size <- 1000
A <- 6*rnorm(size)+50
B <- -2*rnorm(size) - 25 + rnorm(size)
C <- -4*A + 5*B + 3 +rnorm(size)
summary(lm(C~A+B))</pre>
```

```
##
## Call:
## lm(formula = C \sim A + B)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
##
  -3.03321 -0.68565 0.01655 0.66794 3.13811
##
##
  Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                                       6.888
   (Intercept) 2.967859 0.430869
                                                1e-11 ***
##
## A
               -4.000487
                           0.005264 - 759.946
                                               <2e-16 ***
## B
                4.998128 0.014068 355.283
                                               <2e-16 ***
##
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
  Signif. codes:
##
## Residual standard error: 0.9947 on 997 degrees of freedom
## Multiple R-squared: 0.9986, Adjusted R-squared: 0.9986
## F-statistic: 3.641e+05 on 2 and 997 DF, p-value: < 2.2e-16
```

# Bayesian Network 3 (again) with A as the outcome:



Question: What would a regression model of A ~ B + C yield?

- a. Both B and C should be statistically significant
- b. Only *B* should be statistically significant
- c. Only C should be statistically significant
- d. Neither B nor C should be statistically significant

```
summary(lm(A~B+C))
```

```
##
## Call:
## lm(formula = A \sim B + C)
##
##
  Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
##
  -0.75638 -0.17022 0.00544 0.16841
                                        0.80335
##
##
  Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
  (Intercept) 0.8215779 0.1070244
                                        7.677 3.89e-14 ***
##
## B
                1.2470301
                          0.0039408
                                      316.439 < 2e-16 ***
## C
               -0.2495388
                          0.0003284 - 759.946 < 2e-16 ***
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 0.2484 on 997 degrees of freedom
## Multiple R-squared: 0.9983, Adjusted R-squared: 0.9983
## F-statistic: 2.893e+05 on 2 and 997 DF, p-value: < 2.2e-16
```

Question: What would a regression model of A ~ B yield?

- a. *B* should be statistically significant
- b. B should not be statistically significant

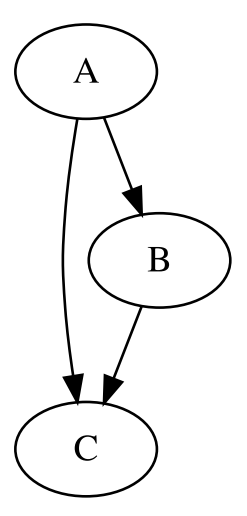
```
summary(lm(A~B))
```

```
##
## Call:
  lm(formula = A \sim B)
##
##
## Residuals:
##
       Min
                  10
                      Median
                                    30
                                            Max
##
   -19.9644 -3.8309 -0.0804
                                3.8547
                                       19.3418
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
  (Intercept) 46.99023
                           2.12137 22.151
                                             <2e-16 ***
##
## B
               -0.11401
                           0.08452 - 1.349
                                              0.178
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 5.982 on 998 degrees of freedom
## Multiple R-squared: 0.00182,
                                    Adjusted R-squared:
## F-statistic: 1.82 on 1 and 998 DF, p-value: 0.1777
```

• Even though A and B are independent, they are conditionally dependent if controlling for c.

- Why did this happen? Let's take a simple example
- Assume  $A \sim \text{Bernoulli}(0.4)$ , and  $B \sim \text{Bernoulli}(0.7)$
- Question: What is P(B = 1 | A = 1)?
- Define  $C = \begin{cases} 1 \text{ when } A = B \\ 0 \text{ when } A \neq B \end{cases}$
- Question: What is P(B=1|A=1,C=0)?
- *A* and *B* are independent; that is, knowledge of *B* give no information on the value of *A*. But, additional knowledge of *C* does give information about the value of *A*.

### **Bayesian Network 4**



### Mental Model:

- A =Study into the night
- B = Go to bed late
- C = Fail Test

Question: What would a regression model of C ~ A + B yield?

- a. Both A and B should be statistically significant
- b. Only A should be statistically significant
- c. Only B should be statistically significant
- d. Neither A nor B should be statistically significant

```
set.seed(1234)
size <- 1000
A <- 6*rnorm(size)+50
B <- A - 25 - 2*rnorm(size)
C <- -4*A + 5*B + 3 +rnorm(size)
summary(lm(C~A+B))</pre>
```

```
##
## Call:
## lm(formula = C \sim A + B)
##
## Residuals:
                      Median
##
       Min
                  10
                                    30
                                            Max
##
  -3.13161 -0.71957 0.03478 0.70215 3.05316
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
                           0.47704
                                      7.009 4.41e-12 ***
## (Intercept) 3.34366
                           0.01692 -237.358 < 2e-16 ***
## A
               -4.01550
## B
                5.01845
                           0.01635 306.907 < 2e-16 ***
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Residual standard error: 1.013 on 997 degrees of freedom
## Multiple R-squared: 0.992, Adjusted R-squared:
## F-statistic: 6.153e+04 on 2 and 997 DF, p-value: < 2.2e-16
```

Question: What about this regression model: c ~ A?

- a. A should be statistically significant
- b. A should not be statistically significant

```
summary(lm(C~A))
```

```
##
## Call:
## lm(formula = C \sim A)
##
##
  Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
  -30.7632 -6.7266 -0.2299
##
                                6.3962
                                        31.5156
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
  (Intercept) -117.61814
                             2.62464 -44.81
                                               <2e-16 ***
## A
                  0.90975
                             0.05229
                                       17.40
                                               <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 9.889 on 998 degrees of freedom
## Multiple R-squared: 0.2327, Adjusted R-squared: 0.232
## F-statistic: 302.7 on 1 and 998 DF, p-value: < 2.2e-16
```

Question: What about this regression model: B ~ A + C?

- a. Both A and C should be statistically significant
- b. Only A should be statistically significant
- c. Only C should be statistically significant
- d. Neither A nor C should be statistically significant

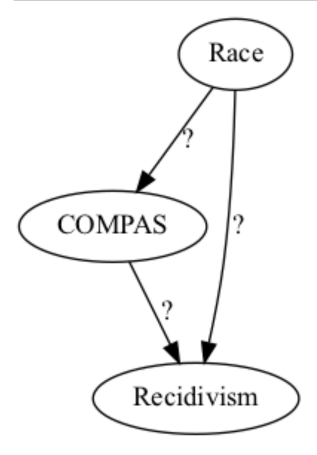
```
summary(lm(B~A+C))
```

```
##
## Call:
## lm(formula = B \sim A + C)
##
## Residuals:
##
        Min
                  10
                       Median
                                    30
                                             Max
  -0.61703 -0.13791 -0.00305 0.14136
##
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
  (Intercept) -0.9117518 0.0924550 -9.862
                                               <2e-16 ***
##
## A
                0.8020461
                          0.0012115 662.016
                                               <2e-16 ***
## C
                0.1971777 0.0006425 306.907
                                               <2e-16 ***
##
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.2007 on 997 degrees of freedom
## Multiple R-squared: 0.999, Adjusted R-squared: 0.9989
## F-statistic: 4.747e+05 on 2 and 997 DF, p-value: < 2.2e-16
```

# COMPAS and possible collider bias

- COMPAS uses questionnaire (https://www.documentcloud.org/documents/2702103-Sample-Risk-Assessment-COMPAS-CORE.html) responses to predict recidivism.
- Because COMPAS is used in sentencing, it may actually impact recidivism as well.
- One way to quantify racial bias in COMPAS would be to isolate the link between race and COMPAS that is not associated with recidivism. But, it is not clear how to untangle this from potential collider bias.

```
digraph {
  Race -> COMPAS [ label = "?"];
  COMPAS -> Recidivism [ label = "?"];
  Race -> Recidivism [ label = "?"]
}
```



- If we used decile\_score ~ is\_recid + race as a model to quantify bias, it seems very likely that there will be collider bias
- IMPORTANT: The model below is NOT informative because of the possible causal structure of the data
  - Included for teaching purposes only

```
summary(lm(decile_score ~ is_recid + race, data=df))
```

```
##
## Call:
## lm(formula = decile score ~ is recid + race, data = df)
##
## Residuals:
##
                           3Q
     Min
              1Q Median
                                  Max
  -7.225 -2.224 -0.225 1.776 7.555
##
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
                        4.73952
                                  0.04127 114.848 < 2e-16 ***
## (Intercept)
## is recid
                        1.48548
                                  0.05345 27.794 < 2e-16 ***
## raceAsian
                      -2.31198
                                  0.36300 -6.369 1.98e-10 ***
## raceCaucasian
                      -1.51576
                                0.05569 -27.217 < 2e-16 ***
                       -1.81059
                                  0.09033 -20.043 < 2e-16 ***
## raceHispanic
## raceNative American -0.47038
                                   0.43961 - 1.070
                                                     0.285
## raceOther
                       -2.29469
                                   0.11157 -20.566 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 2.629 on 11031 degrees of freedom
## Multiple R-squared: 0.1656, Adjusted R-squared:
## F-statistic: 364.9 on 6 and 11031 DF, p-value: < 2.2e-16
```

In the regression above, several race indicator variables are significant. But, because collider bias is possible here, we *cannot* conclude that COMPAS is racially biased.

# Survival Analysis

- Survival analysis is a set of statistical methods for modeling the time until an event occurs, especially when follow up is not complete for each observation.
- Example: Testing a new terminal cancer treatment, participants are either given the standard or test treatment. The goal is to prolong the patient's life. Each patient is followed until death from cancer. During follow up some participants die from cancer but some drop out while others might die from something else. Survival analysis allows us to use this data even though we do not have events for each participant.

### Set up

- Assume that T is the time until an event randomly occurs.
- For example, T might be the duration from cancer treatment until remission or death.
- $T \sim f$ 
  - That is, f(t) is the probability density function (pdf) of T where t is time
- $F(t) = P(T < t) = \int_0^t f(x) dx$  is cumulative distribution function (cdf) of T
- · Survival function:

$$S(t) = P(T > t) = 1 - F(t)$$

- The survival function gives the probability of not having an event before time t (survive until t)
- Hazard function:

$$\lambda(t) = \lim_{h \to 0} \frac{P(T \le t + h | T > t)}{h} = \lim_{h \to 0} \frac{P(t < T \le t + h)}{h P(T > t)} = \frac{f(t)}{S(t)} = -\frac{d \log S(t)}{dt}.$$

- Hazard function gives the instantaneous probability of an event at time t given survival until time t
- Notice that  $f(t) = \lambda(t)S(t)$
- Cumulative hazard function:

$$\Lambda(t) = \int_0^t \lambda(x) dx = -\int_0^t d\log S(x) = -\log S(t).$$

How to get the survival function from the hazard function:

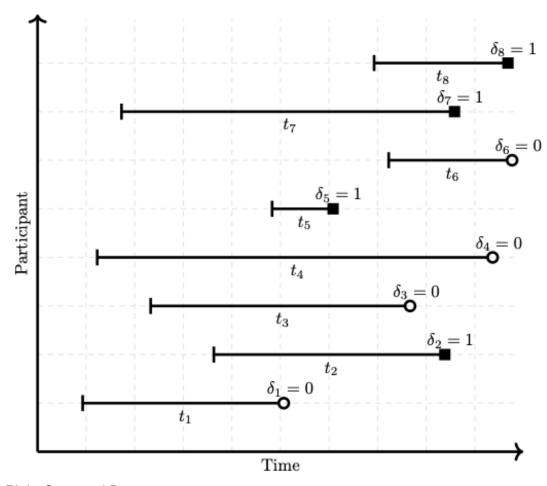
$$S(t) = \exp[-\Lambda(t)].$$

• Side note: If  $\lambda(t) = \lambda$  (constant function), then f is the exponential distribution:

$$\lambda(t) = \lambda \Leftrightarrow \Lambda(t) = \lambda t$$
  
 
$$\Leftrightarrow S(t) = \exp(-\lambda t)$$
  
 
$$\Leftrightarrow f(t) = \lambda(t)S(t) = \lambda \exp(-\lambda t)$$

### **Censoring at Random**

- Not always possible to wait for an event to occur for each participant before doing the analysis
- Cancer study example: participants may drop out of the study before an event is observed or the study may close before each participant experiences an event
- This is call right censored data: have start time but end times can either be at event or drop out time
- Question: For censored observations, how to make use of time duration without event?



### Right Censored Data

- Model:  $f(t|x;\theta)$  with corresponding hazard,  $\lambda(t|x;\theta)$ , and survival,  $S(t|x;\theta)$
- ullet Want heta to quantify difference in risk (until event) among observations
  - $\circ$  Note:  $\theta$  quantifies how fast an event will likely occur for an observation but communicated in terms of risk of event
- Assumption: censoring occurs at random (in independently from f)
- Censoring cumulative probability distribution model:

$$G(t; \phi)$$

• Corresponding censoring pdf model:

$$g(t;\phi)$$

· Data:

$$(t_1, \delta_1), \ldots, (t_n, \delta_n)$$

- $t_i$  for  $i=1,\ldots,n$  is duration of follow-up until either event or censor time
- $\delta_i$  is event indicator:

- $\delta_i = 1$  means that observation *i* had an event and  $t_i \sim f(t; \theta)$
- $\circ \ \delta_i = 0 \ \mathrm{means} \ \mathrm{that} \ \mathrm{observation} \ i \ \mathrm{was} \ \mathrm{censored} \ \mathrm{and} \ t_i \sim g(t;\phi)$
- Because f and g are independent (and because observations are independent), the likelihood is

$$L(\theta, \phi) = \prod_{i=1}^{n} [f(t_i; \theta)[1 - G(t_i; \phi)]]^{\delta_i} [g(t_i; \phi)S(t_i; \theta)]^{1 - \delta_i}$$

$$= \prod_{i=1}^{n} [f(t_i; \theta)]^{\delta_i} [S(t_i; \theta)]^{1 - \delta_i} \prod_{i=1}^{n} [g(t_i; \phi)]^{1 - \delta_1} [1 - G(t_i; \phi)]^{\delta_i}$$

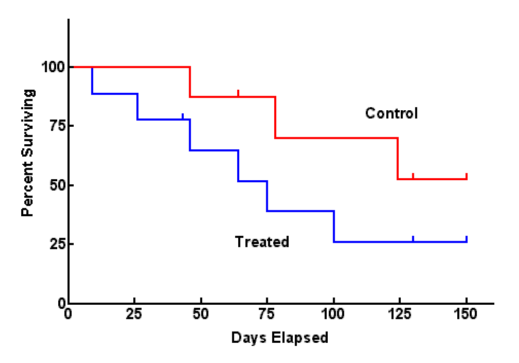
$$= L(\theta)L(\phi) \propto L(\theta).$$

- Observe an event for i ( $\delta_i=1$ ), then  $t_i\sim f$  and censoring did not occur prior  $[f(t_i;\theta)[1-G(t_i;\phi)]]^{\delta_i}$
- Observe censoring for i ( $\delta_i=1$ ), then  $t_i\sim g$  and an event did not occur prior  $[g(t_i;\phi)S(t_i;\theta)]^{1-\delta_i}$
- But, we do not care about the censoring distribution, only the time to event distribution.
- Partial likelihood

$$L(\theta) = \prod_{i=1}^{n} [f(t_i; \theta)]^{\delta_i} [S(t_i; \theta)]^{1-\delta_i} = \prod_{i=1}^{n} \lambda(t_i)^{\delta_i} S(t_i)$$

# Kaplan-Meier Estimator

- Question: Have you heard of/seen Kaplan-Meier Curves before this?
  - A: Yes
  - B: No
- Visualize the percent of population surviving until time *t* as *t* increases



### KM Curve Example

- Consider estimating survival: S(t) = P(T > t) from sample  $(t_1, \delta_1), \dots, (t_n, \delta_n)$
- Approximate S(t) as a non-parametric decreasing step function
  - S(t) is proportion of sample that has not experienced an event at time t
  - $\circ$  Problem: If i censored prior to t, we cannot know if their event occurred before or after t
- Order sample by event times  $t_i$  where  $\delta_i = 1$ :

$$t_{(1)}, t_{(2)}, \ldots, t_{(J)}$$

- ullet There are only J sample points in time where events occur
- Recall conditional probability rule  $P(A|B) = \frac{P(A,B)}{P(B)}$
- Because  $t_{(j)} > t_{(j-1)}$ ,

$$S\left(t_{(j)}\right) = P\left(T > t_{(j)}\right) = P\left(T > t_{(j)}, T > t_{(j-1)}\right) = P\left(T > t_{(j)} | T > t_{(j-1)}\right) \times P\left(T > t_{(j-1)}\right)$$

Repeating

$$S(t_{(j)}) = P(T > t_{(j)}|T > t_{(j-1)}) \times P(T > t_{(j-1)}|T > t_{(j-2)}) \times P(T > t_{(j-2)})$$

$$= P(T > t_{(j)}|T > t_{(j-1)}) \times P(T > t_{(j-1)}|T > t_{(j-2)}) \times \dots \times P(T > t_{(2)}|T > t_{(1)}) \times P(T > t_{(1)})$$

- This seems tautological (https://www.merriam-webster.com/dictionary/tautology), but it's helpful here because it allows us to include the censored observations in the denominator appropriately as *t* increases
- For j = 1, ..., J, the "instantaneous" probability of an event occurring at time  $t_{(i)}$ :

$$\pi_i = P\left(T \le t_{(i)}|T > t_{(i-1)}\right) = 1 - P\left(T > t_{(i)}|T > t_{(i-1)}\right)$$

Then

$$S(t_{(j)}) = (1 - \pi_j)(1 - \pi_{j-1}) \dots (1 - \pi_2)(1 - \pi_1).$$

- Calculate π<sub>i</sub>:
  - Let  $n_j = \#\{t_i \ge t_{(j)}\}$  be the number of participants who are still at risk (who haven't had an event or been censored) at time  $t_{(j)}$
  - Note: that  $n_i$  decreases as events occur or as they are censored.
  - Let  $d_j = \#\{t_i = t_{(j)}, \delta_i = 1\}$  be the number of events that occur at time  $t_{(j)}$ .
  - Maximizes the non-parametric likelihood

$$\pi_j = \frac{d_j}{n_i}$$

So, we can approximate the survival function as

$$\hat{S}(t) = \prod_{j=1}^{J} \left( 1 - \frac{d_j}{n_j} \right)^{I(t_{(j)} \le t)}.$$

• Using the delta-method, we can approxmiate the variance of the estimated survival function as

$$\hat{V}[\hat{S}(t)] = \hat{S}(t)^2 \sum_{j: t_{(i)} \le t} \frac{d_j}{n_j (n_j - d_j)}$$

With the variance, we can run statistical tests

This video (https://www.youtube.com/watch?v=NDgn72ynHcM) clearly illustrates how to calculate the KM survival function.

```
library(survival)

dat <- read.csv(url('https://raw.githubusercontent.com/propublica/compas-analysis/master/
cox-parsed.csv'))
names(dat)</pre>
```

```
"name"
    [1] "id"
##
##
   [3] "first"
                                    "last"
                                    "sex"
##
   [5] "compas screening date"
## [7] "dob"
                                    "age"
  [9] "age cat"
                                    "race"
##
## [11] "juv_fel_count"
                                    "decile score"
## [13] "juv misd count"
                                    "juv other count"
## [15] "priors count"
                                    "days b screening arrest"
## [17] "c_jail_in"
                                    "c jail out"
## [19] "c case number"
                                    "c offense date"
## [21] "c_arrest_date"
                                    "c days from compas"
## [23] "c_charge_degree"
                                    "c charge desc"
## [25] "is recid"
                                    "r case number"
                                    "r days from arrest"
## [27] "r_charge_degree"
## [29] "r_offense_date"
                                    "r charge desc"
## [31] "r_jail in"
                                    "r jail out"
## [33] "violent_recid"
                                    "is violent recid"
                                    "vr charge degree"
## [35] "vr case number"
## [37] "vr_offense_date"
                                    "vr charge desc"
## [39] "type_of_assessment"
                                    "decile score.1"
## [41] "score text"
                                    "screening date"
## [43] "v_type_of_assessment"
                                    "v decile score"
## [45] "v_score_text"
                                    "v_screening_date"
                                    "out custody"
## [47] "in custody"
## [49] "priors_count.1"
                                    "start"
## [51] "end"
                                    "event"
```

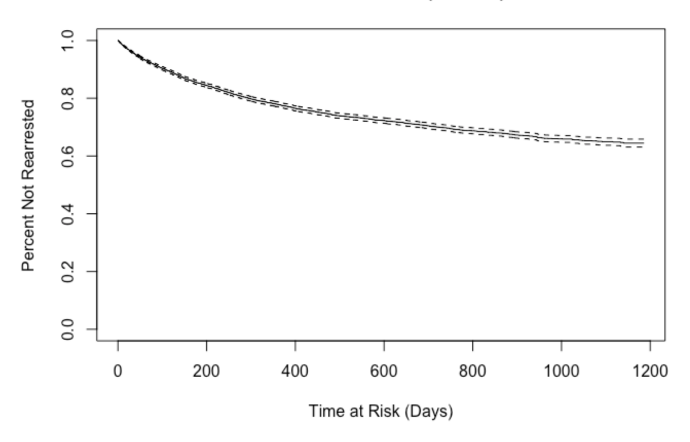
```
dim(dat)
## [1] 13419
                 52
dat2 <- dat[dat$end > dat$start,]
dim(dat2)
## [1] 13356
                 52
dat3 <- dat2[!duplicated(dat2$id),]</pre>
dim(dat3)
## [1] 10325
                 52
ph <- dat3[dat3$decile_score>0,]
dim(ph)
## [1] 10314
                 52
ph$t_atrisk <- ph$end - ph$start</pre>
survobj <- with(ph, Surv(t atrisk, event))</pre>
fit0 <- survfit(survobj~1, data=ph)</pre>
```

# summary(fit0)

plot(fit0, xlab="Time at Risk (Days)",
 ylab="Percent Not Rearrested",

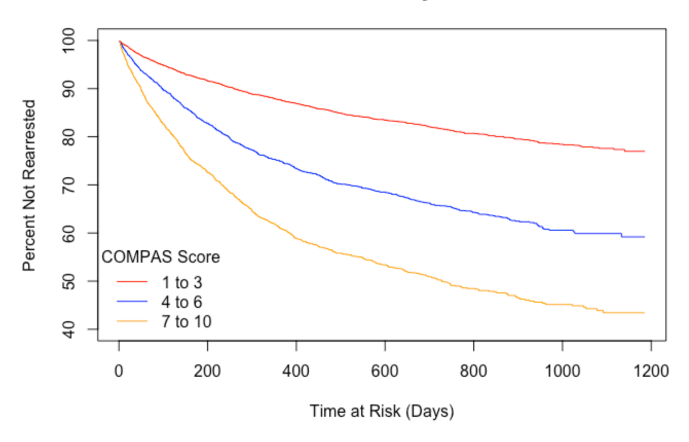
main ="Survival Function (Overall)")

### Survival Function (Overall)



```
ph$compas <- cut(ph$decile_score, breaks=c(0,3,6,10))
fitc <- survfit(survobj~compas, data=ph)
plot(fitc, xlab="Time at Risk (Days)",
   ylab="Percent Not Rearrested", yscale=100, ylim=c(1, 0.4),
   main="Survival Function by COMPAS",
   col = c('red', 'blue', 'orange', 'yellow', 'green', 'purple'))
legend_text = c('1 to 3', '4 to 6', '7 to 10')
legend('bottomleft', legend=legend_text, title='COMPAS Score', bty='n',
        col=c('red', 'blue', 'orange'), lty=1)</pre>
```

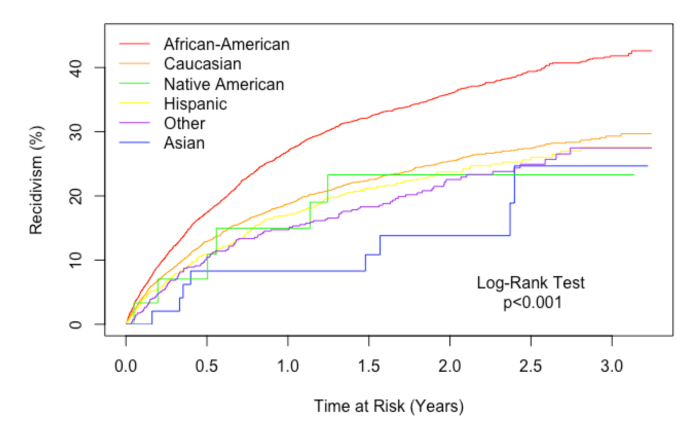
### Survival Function by COMPAS



```
ph$t_atrisk <- ph$end - ph$start

survobj_yr <- with(ph, Surv(t_atrisk/365.25, event))
fitr <- survfit(survobj_yr~race, data=ph)
# `fun` parameter only works here because plot is actually calling plot.survfit
plot(fitr, xlab="Time at Risk (Years)",
    ylab="Recidivism (%)", yscale=100, ylim=c(0, 0.45), fun = function(x) {1 - x},
    main="Recidivism by Race over Time",
    col = c('red', 'blue', 'orange', 'yellow', 'green', 'purple'))
recid_order = c(1,3,5,4,6,2)
legend('topleft', legend=levels(as.factor(ph$race))[recid_order], bty='n',
        col = c('red', 'blue', 'orange', 'yellow', 'green', 'purple')[recid_order], lty=1)
text(2.5, 0.05, 'Log-Rank Test\n p<0.001')</pre>
```

### Recidivism by Race over Time



- · Remember: For publication, make plots as easy as possible to understand
- In this case, the plot is easier to understand looking at recidivism rather than not recidivism
- Here S(t) give the probability that rearrest will occur after time t; that is, someone will not be arrested before time t
- F(t) = 1 S(t) is the cumulative probability of arrest before time t
- Each curve gives the proportion of recidivism in a racial group at time t
- Years seems easier to understand than day for this
- We can use the log-rank test (https://en.wikipedia.org/wiki/Logrank\_test) to see if at least one curve is different from others
- ullet  $H_0$ : There is no different between the curves;  $H_1$ : at least one curve is different from others

survdiff(survobj~race, data=ph)

```
## Call:
## survdiff(formula = survobj ~ race, data = ph)
##
##
                             N Observed Expected (O-E)^2/E (O-E)^2/V
## race=African-American 5147
                                   1607
                                         1293.97
                                                     75.725
                                                              142.951
                                           16.22
                                                      4.167
                                                                4.195
## race=Asian
                            51
## race=Caucasian
                          3569
                                    814
                                          994.40
                                                     32.727
                                                               51.230
## race=Hispanic
                           944
                                    206
                                          275.38
                                                     17.480
                                                              19.438
## race=Native American
                            32
                                            8.26
                                                      0.618
                                                                0.621
## race=Other
                           571
                                    118
                                          170.77
                                                     16.305
                                                               17.397
##
##
    Chisq= 147 on 5 degrees of freedom, p= <2e-16
```

- Statistical interpretation: This significant p-value indicates that at least one survival curve is different from the other
- · Research area interpretation: Recidivism rates differ by race
- Don't forget: documentation (https://cran.r-project.org/web/packages/survival/survival.pdf) is your best friend
- As a consultant, you will probably need to read the documentation a lot.

# Cox proportional hazards model

- Difficult to work with censored data using generalized linear models
- Question: We can use use a GLM here?
  - A: Yes
  - B: No
  - · C: Not Sure
- Assuming that each individual hazard function is proportional to some common baseline hazard function makes the problem workable:

$$\lambda(t|x_i) = \lambda_0(t) \exp(\beta x_i)$$

where  $x_i$  is the covariate vector for participant i and  $\beta$  is the parameter vector to be estimated

- $\lambda_0(t)$  is the hazard function for  $x_i = (0, ..., 0)$
- $\exp(\beta x_i)$  explains proportional differences in hazards as  $x_i$  changes as in parametric regression
- ullet Then the probability that individual j experiences an event at  $t_{(j)}$  given survival until  $t_{(j)}$  is

$$\lambda(t_{(j)}|x_{(j)}) = \lambda_0(t_{(j)}) \exp(x_{(j)}\beta)$$

• The total probability within the sample of an event occurring at time  $t_{(j)}$  given those who have survived until  $t_{(j)}$  is

$$\sum_{k:t_k \ge t_{(j)}} \lambda(t_{(j)}|x_k) = \sum_{k:t_k \ge t_{(j)}} \lambda_0(t_{(j)}) \exp(x_k \beta)$$

• Then probability of an event occurring at  $t_{(j)}$  conditioning on covariates  $x_{(j)}$  (partial likelihood) is

$$\tilde{L}_{j}(\beta) = P[(j) \text{ fails} | 1 \text{ failure from those at risk at } t_{(j)}] = \frac{P[(j) \text{ fails} | \text{still at risk}]}{\sum_{k:t_{k} \geq t_{(j)}} P(k \text{ fails} | \text{still at risk})}$$

$$= \frac{\lambda(t_{(j)}|x_{(j)})}{\sum_{k:t_{k} \geq t_{(j)}} \lambda(t_{(j)}|x_{k})} = \frac{\lambda_{0}(t_{(j)}) \exp(x_{(j)}\beta)}{\sum_{k:t_{k} \geq t_{(j)}} \lambda_{0}(t_{(j)}) \exp(x_{k}\beta)} = \frac{\exp(x_{(j)}\beta)}{\sum_{k:t_{k} \geq t_{(j)}} \exp(x_{k}\beta)}$$

- Notice that the baseline hazard function,  $\lambda_0(t)$ , cancels. So, now we can use use an optimization technique to maximize this function
- The joint likelihood for the sample is

$$\tilde{L}(\beta) = \prod_{j=1}^{J} L_j(\beta) = \prod_{j=1}^{J} \frac{\exp(x_{(j)}\beta)}{\sum_{k:t_k \ge t_{(j)}} \exp(x_k\beta)} = \prod_{i=1}^{n} \left[ \frac{\exp(x_i\beta)}{\sum_{\ell \in R(t_i)} \exp(x_{\ell}\beta)} \right]^{\delta_i}$$

log- partial likelihood:

$$\tilde{\ell}(\beta) = \sum_{j=1}^{J} \left[ x_{(j)}\beta - \log \left( \sum_{k: t_k \ge t_{(j)}} \exp(x_k \beta) \right) \right]$$

where  $R(t) = \{\ell : t_{\ell} \ge t\}$ 

· Maximize the likelihood with Newton-Raphson method

# Is COMPAS racially biased?

- To determine this, we must use interactions
- Let A and B be binary, input variables and Y a continuous outcome
  - assume linear regression
- How do we read the output?

Variable	Coef	p-value
A	1.5	0.01
В	0.1	0.35
A*B	0.5	0.02

• On average, Y increases by 1.5 when A=1 compared to A=0 while controlling for B and this change is statistically significant at an  $\alpha=0.05$  significance level

- There is no evidence to suggest that B is associated with Y (while controlling for A)
- Additionally, Y increases by 0.5 when both A and B are 1
- · Changing the baseline race
  - R uses alphabetical order so African-American (AA) would be the reference group without the relevel command
  - · Now, Caucasian (white) is the reference group
  - In most medical literature, white is the reference racial group, but this has come under some criticism
  - · Here, because AA is of particular interest, we probably don't want AA to be the reference group
- We divide age by 10 so that we can interpret change in risk per 10 years of age

```
ph$race = relevel(as.factor(ph$race), ref="Caucasian")
ph$age10 = ph$age/10
summary(coxph(survobj~decile_score*race+sex+age10, data=ph))
```

```
##
  coxph(formula = survobj ~ decile_score * race + sex + age10,
##
      data = ph)
##
    n=10314, number of events= 2759
##
##
##
                                        coef exp(coef) se(coef)
                                                                     z Pr(>|z|)
                                               1.20346 0.01302 14.226 < 2e-16
## decile score
                                     0.18520
## raceAfrican-American
                                     0.27475
                                               1.31620 0.09127 3.010 0.00261
## raceAsian
                                    -1.84443
                                               0.15812 0.82830 -2.227 0.02596
## raceHispanic
                                     0.07333
                                               1.07608 0.14037 0.522 0.60142
## raceNative American
                                    -2.60760
                                               0.07371 1.55362 -1.678 0.09327
## raceOther
                                    -0.22491
                                               0.79859 0.17648 - 1.274 0.20252
## sexMale
                                     0.39744
                                               1.48801 0.05268 7.545 4.54e-14
                                               0.90209 0.01875 -5.497 3.87e-08
## age10
                                    -0.10304
                                               0.96592 0.01558 -2.226 0.02604
## decile score:raceAfrican-American -0.03467
                                               1.36054 0.12496 2.464 0.01375
## decile score:raceAsian
                                     0.30788
## decile score:raceHispanic
                                    -0.03940
                                               0.96137 0.02765 -1.425 0.15418
## decile score:raceNative American
                                     0.30262
                                               1.35341 0.18380 1.646 0.09967
## decile score:raceOther
                                     0.04402
                                               1.04500 0.03573 1.232 0.21798
##
## decile score
## raceAfrican-American
## raceAsian
## raceHispanic
## raceNative American
## raceOther
## sexMale
## age10
## decile score:raceAfrican-American *
## decile_score:raceAsian
```

```
## decile_score:raceHispanic
## decile score:raceNative American
## decile_score:raceOther
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
                                     exp(coef) exp(-coef) lower .95 upper .95
## decile score
                                       1.20346
                                                   0.8309 1.173140
                                                                        1.2346
## raceAfrican-American
                                       1.31620
                                                    0.7598 1.100602
                                                                        1.5740
## raceAsian
                                       0.15812
                                                   6.3245 0.031183
                                                                        0.8017
## raceHispanic
                                                   0.9293 0.817258
                                                                        1.4169
                                       1.07608
## raceNative American
                                       0.07371
                                                  13.5665
                                                           0.003508
                                                                        1.5487
## raceOther
                                       0.79859
                                                   1.2522
                                                           0.565069
                                                                        1.1286
## sexMale
                                       1.48801
                                                   0.6720 1.342037
                                                                       1.6499
                                       0.90209
                                                   1.1085 0.869549
                                                                        0.9358
## age10
## decile score:raceAfrican-American
                                       0.96592
                                                   1.0353
                                                           0.936875
                                                                        0.9959
## decile score:raceAsian
                                       1.36054
                                                   0.7350 1.064977
                                                                        1.7381
## decile score:raceHispanic
                                       0.96137
                                                   1.0402
                                                           0.910651
                                                                        1.0149
## decile score:raceNative American
                                       1.35341
                                                   0.7389
                                                           0.944004
                                                                        1.9404
## decile score:raceOther
                                       1.04500
                                                   0.9569
                                                           0.974321
                                                                        1.1208
##
## Concordance= 0.667 (se = 0.005)
## Likelihood ratio test= 928.2 on 13 df,
                                             p = < 2e - 16
## Wald test
                        = 898.1 on 13 df,
                                             p = < 2e - 16
## Score (logrank) test = 965 on 13 df,
```

Factor	Hazard Rate Ratio (95% CI)	p-value
COMPAS		
Decile Score (per point)	1.20 (1.17, 1.23)	<0.001
Race (compared to White)		
Black	1.32 (1.10, 1.57)	0.003
Asian	0.16 (0.03, 0.80)	0.026
Hispanic	1.08 (0.82, 1.42)	0.601
Native American	0.07 (0.00, 1.55)	0.093
Other	0.80 (0.57, 1.13)	0.203
Sex (compared to Female)		
Male	1.49, (1.34, 1.65)	<0.001
Age		
(per 10 years)	0.90 (0.87, 0.94)	<0.001

### COMPAS Decile Score by Race (compared to White)

Black (per one point)	0.97 (0.94, 0.99)	0.026
Asian (per one point)	1.36 (1.06, 1.74)	0.014
Hispanic (per one point)	0.96 (0.91, 1.01)	0.154
Native American (per one point)	1.35 (0.94, 1.94)	0.100
Other (per one point)	1.05 (0.97, 1.12)	0.218

- Question: Does this model indicate that COMPAS is racially biased?
  - · A: Yes
  - B: No
  - · C: Not Sure
- Interpretations:
- For each unit increase in COMPAS decile score, risk of recidivism increases, on average, by a factor 1.2 (p<0.001).</li>
- Risk of recidivism is greater for Blacks (1.32, p=0.003) and smaller for Asians (0.16, p=0.026) compared to Whites; all other racial groups had similar risk to Whites.
- Risk is greater for men with 1.5 (p<0.001) times the risk of women.
- Risk decreases with age; for every 10 year increase, risk drops by a factor of 0.9 (10%) on average.
- Compared to Whites, a unit increase in the COMPAS decile score for African-Americans corresponds to a decrease in risk of recidivism by a factor of 0.97 (p=0.026).
- Said differently, among African-Americans and Caucasians with similar COMPAS scores, African-Americans, on average, have a 3% lower risk of recidivism compared to Caucasians. This indicates that COMPAS may be assigning higher scores to African-American than to Caucasians with a similar risk of recidivism.
- Asians, on the other hand, were assigned lower scores than Caucasians with a similar risk of recidivism (p=0.0128). There were no differences between other racial groups and Caucasians.
- Testing proportional hazards assumption
- Null Hypothesis: Proportional hazards
- Should consider transformation (next lecture)

```
test.ph <- cox.zph(coxph(survobj~race+age+decile_score, data=ph))
test.ph</pre>
```

```
## chisq df p
## race 6.79 5 0.2367
## age 4.66 1 0.0308
## decile_score 2.98 1 0.0841
## GLOBAL 18.89 7 0.0085
```

• Using our knowledge of regression with causation (Bayesian Networks above), how can we determine if the COMPAS algorithm is racially biased?

### **Time-Dependent Covariates**

- In cases, covariates can change over time
  - Here, zip code, or age can change over time
  - · This change may have an effect on the hazard function
- Recall that  $\lambda(t)$  is the instantaneous probability of an event at time t given survival up to t
- If one or more covariates change over time, x(t), we can model hazard as

$$\lambda(t|x(t)) = \lambda_0(t) \exp(\beta x(t))$$

· The partial likelihood become

$$\tilde{L}(\beta) = \prod_{i=1}^{n} \left[ \frac{\exp(x_i(t_i)\beta)}{\sum_{\ell \in R(t_i)} \exp(x_{\ell}(t_i)\beta)} \right]^{\delta_i}$$

### **Stratified Models**

If a sample of n observations are thought to have S mutually exclusive baseline hazards, we can choose to
use a stratified model

$$\lambda_h(t|x) = \lambda_{0h}(t) \exp(x\beta)$$
 for  $h = 1, \dots, S$ 

- Example: Want to assess effect of age and weight only on risk of death, we may want to stratify by gender
- If covariates are assumed to be different in different strata, we can estimate strata-specific parameters,  $\beta_h$ , for each strata

$$\lambda_h(t|x) = \lambda_{0h}(t) \exp(x\beta_h)$$
 for  $h = 1, \dots, S$ 

Partial likelihood:

$$\tilde{L}(\beta) = \prod_{h=1}^{S} \prod_{i=1}^{n_h} \left[ \frac{\exp(x_{i(h)}\beta_h)}{\sum_{\ell \in R_h(t_{i(h)})} \exp(x_{\ell(h)\beta_h}))} \right]$$

where  $n_h$  is the number in each strata, i(h) is the ith observation in the hth stratam,  $R_h$  is the stratam specific risk set

### Frailty model

- Some data will have associations among the observations themselves
- Example: COMPAS data could have multiple arrests, their associated COMPAS score, and their own follow up
- It is reasonable to assume that past scores, and arrests may provide information (association) on future data
- If there are associations among the observations in the data, the parameter point estimates will be accurate
- But, standard error will not be correct, so any inference (p-values, confidence intervals) will not be valid
- Solution: modify the information sandwich for GLMs with associated observations to Cox PH
- This provides a consistent estimator for the covariance matrix
- Note: so far we have not discussed sandwich estimator

These notes are based on chapter 9 of Lachin, John M. Biostatistical methods: the assessment of relative risks. Vol. 509. John Wiley & Sons, 2009.

# Consulting Case Study: Treating Syphilis in People living with HIV

- The typically, the first line treatment for syphilis is penicillin
- But, people living with HIV are sometimes thought to be immunocompromised
- Because of this, it was common for physicians to administer two or more standard doses to treat syphilis for someone living with HIV
- US treatment guidelines in the US recommended one standard dose regardless of HIV status
- But, there was disagreement in the medical community on this guideline
- This type of disagreement (equipoise) frequently leads to research

**Background:** Treatment guidelines recommend the use of a single dose of benzathine penicillin G (BPG) for treating early syphilis in human immunodeficiency virus (HIV)-infected persons. However, data supporting this recommendation are limited. We examined the efficacy of single-dose BPG in the US Military HIV Natural History Study.

**Methods:** Subjects were included if they met serologic criteria for syphilis (ie, a positive nontreponemal test [NTr] confirmed by treponemal testing). Response to treatment was assessed at 13 months and was defined by a  $\geq$ 4-fold decline in NTr titer. Multivariate Cox proportional hazard regression models were utilized to examine factors associated with treatment response.

**Results:** Three hundred fifty subjects (99% male) contributed 478 cases. Three hundred ninety-three cases were treated exclusively with BPG (141 with 1 dose of BPG). Treatment response was the same among those receiving 1 or >1 dose of BPG (92%). In a multivariate analysis, older age (hazard ratio [HR], 0.82 per 10-year increase; 95% con- fidence interval [CI], .73–.93) was associated with delayed response to treatment. Higher pretreatment titers

(refer- ence NTr titer <1:64; HR, 1.94 [95% CI, 1.58–2.39]) and CD4 counts (HR, 1.07 for every 100-cell increase [95% CI, 1.01–1.12]) were associated with a faster response to treatment. Response was not affected by the number of BPG doses received (reference, 1 dose of BPG; HR, 1.11 [95% CI, .89–1.4]).

**Conclusion:** In this cohort, additional BPG doses did not affect treatment response. Our data support the current recommendations for the use of a single dose of BPG to treat HIV-infected persons with early syphilis.

Ganesan, Anuradha, Octavio Mesner, Jason F. Okulicz, Thomas O'Bryan, Robert G. Deiss, Tahaniyat Lalani, Timothy J. Whitman, Amy C. Weintrob, Grace Macalino, and Brian K. Agan. "A single dose of benzathine penicillin G is as effective as multiple doses of benzathine penicillin G for the treatment of HIV-infected persons with early syphilis." Clinical Infectious Diseases 60, no. 4 (2015): 653-660. (https://academic.oup.com/cid/article/60/4/653/2895840)

### Look for in paper:

- Data inclusion criteria
- · Baseline table for individuals
- · Syphilis episodes table
- KM curves
- Cox PH Model

Response Letter (http://citeseerx.ist.psu.edu/viewdoc/download?doi=10.1.1.864.7969&rep=rep1&type=pdf)

- Because of the disagreement, some clinicians wrote a negative response to our study
- The response focused mainly on the methods

Our Response (https://academic.oup.com/cid/article/60/9/1444/404706)

# **High Level Summary**

- Always explore the data before running regressions and other statistical tests. Look at the raw data itself, try to understand variable names, variable distributions, missing data, etc
- Survival analysis tools, such as Kaplan-Meier curves and Cox PH regression, are helpful when follow times leading up to an event vary by observation, especially when censoring occurs.
- It is important to be aware of possible causal pathways. But, most of the time, it is not possible to use statistical models alone to attribute a causal relationships.