

Animal Shelter Outcomes for Labradors in Austin Animal Center

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Abstract

Although Labradors are widely viewed as one of the most popular breeds in the United States, they are also a popular breed found in animal shelters https://www.cleartheshelters.com. Taking a look at the outcomes of over 6,000 Labradors in the Austin Animal Center (the largest "no-kill" animal shelter in the U.S.), we build a Cox Proportional Hazards model to see how neuter status affects Labrador adoption rates.

Data Source

The data we used was from the Austin Animal Center, downloaded from kaggle at https://www.kaggle.com/aaronschlegel/austin-animal-center-shelter-outcomes-and. The raw data included cats, dogs, birds, livestock and other. Since we were only interested in labradors we created a subset of the data which included 2,128 Labrador and Labrador Mixed breeds for a total of 6,980 observations. We decided to delete one observation which had NULL values for outcome and age upon outcome since we thought our analysis without this observation would not change. We also excluded 48 data points where the gender and neuter status of the Labrador were not known. We split the column of sex upon outcome into two columns for analysis: sex with only Male/Female entries and neuter with Intact/Fixed entries where "Fixed" means the dog is either neutered or spayed and intact means it was not. Another covariate we changed was color. There were a ridiculous amount of "color" descriptions. Our favorite color that we saw was "peach." We decided to change this column to entries of either Mixed fur pattern or Solid fur patterns to denote dogs with more than one color versus dogs that are solid colored. Our final dataset included the following:

- color: The descriptive fur pattern of Mixed color fur pattern or solid color fur pattern.
- event: The event was 1 if the Labrador had been adopted, and censored was 0 if the Labrador had been transferred, missing (off-site, in foster care or in kennel), possible theft, euthanasia or death.
- date_diff: The amount of time in days the Labrador spent in the shelter.
- sex: The gender of the dog.
- neuter: If the dog is fixed or intact.

color date diff event neuter Mixed 63 days Fixed 1 Female Mixed 95 days Female Fixed 1 Mixed 4445 days Male Fixed Mixed 745 days Male Fixed Mixed 1 1764 days Male Fixed

1794 days

Male

Fixed

Table 1: Final Dataset

Research Question

On average how much time do Labradors spend in animal shelters before they go to a home?

0

We will plot a Kaplan-Meier curve and find the median time that Labradors spent in this animal shelter.

Does the color, gender or neuter status of a Labrador affect it's chance for adoption?

Mixed

We'll build a Cox Proportional Hazards (Cox PH) model with the covariates neuter status, sex, and color to answer this question. We will also explore an Accelerated Failure Time (AFT) model as our extension component, to look at a comparison of survival times.

Data Exploration

Before we start our analysis we must understand our data. We found that we had a total of 6,929 observations. Out of these, 3211 are Female (46%), 3713 have mixed fur pattern (53%), and 5726 were fixed (82%). Using the quantile function on the length of time spent in the shelter, we found that 50% of the dogs spent more than 395 days in the shelter, this is over one year!

Figure 1: As a preliminary assessment of our data, and using the quantile function we find that half of the Labradors spend more than a year in the animal shelter (395 days, which is shown as the blue line). However, this form includes both censored and uncensored labradors which underestimates our value so we should take this figure lightly.

Days

We plotted a Kaplan-Meier Curve without controlling for any variables. We noticed that the confidence interval curves were tight around the actual survival curve. This proves that the data is very detailed and the information is concrete because we have enough observations to do analysis on Labradors in the Austin Animal Center.

```
## Call: survfit(formula = labradors.surv ~ 1)
##
## n events median 0.95LCL 0.95UCL
## 6929 5285 732 730 734
```

Kaplan-Meier

To visually see the survival curves we must plot Kaplan-Meier Curves for each covariate color, sex and neuter on time to adoption for labradors. The Kaplan-Meier curves for sex in Figure 3 shows that male Labs have higher estimated survival rates since the blue male line is above the red female line. As time increases, male and female Labs have the same or similar survival rates because the lines look like they begin to overlap after the 3,000 days mark. The Kaplan-Meier curves for neuter status in Figure 4 shows that intact Labs (labs that have not been neutered) have higher estimated survival rates than fixed labs. This means that more fixed labs are being adopted or going to homes (having the event) than labs that have not been neutered. The Kaplan-Meier curves for fur patterns in Figure 5 show labs with single colored fur patterns have higher survival rates than those with mixed fur patterns. The gaps immediately widen after the 50% survival rate mark and begin to close again nearer to the bottom.

Kaplan-Meier Curve for All Austin Animal Center Labradors

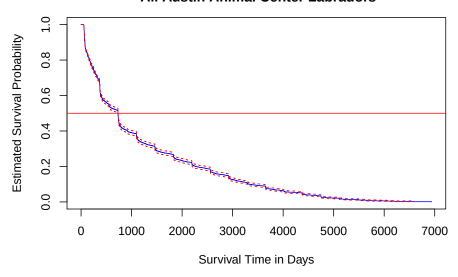


Figure 2: Kaplan-Meier curve

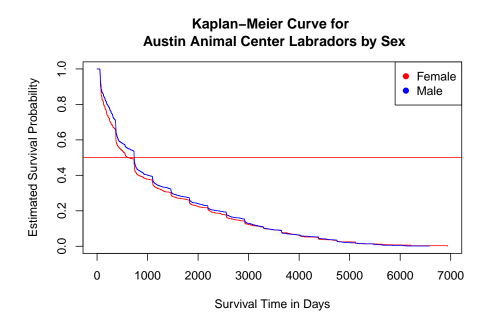


Figure 3: There is not a noticeable difference between the two curves so our pre-analysis hypothesis is that there is not going to be a significant difference.

Kaplan-Meier Curve for **Austin Animal Center Labradors by Fixed Status** Fixed **Estimated Survival Probability** Intact 0.8 9.0 0.4 0.2 0.0 0 1000 2000 4000 3000 5000 6000 7000 Survival Time in Days

Figure 4: We see a bigger difference in the two survival curves between fixed and intact Labradors, mainly that fixed has a lower survival probability over time. However, this might be because there are many more observations on fixed labs than on intact labs, (fixed has about 4,000 more observations) so we will proceed with caution in our analysis because of this fact.

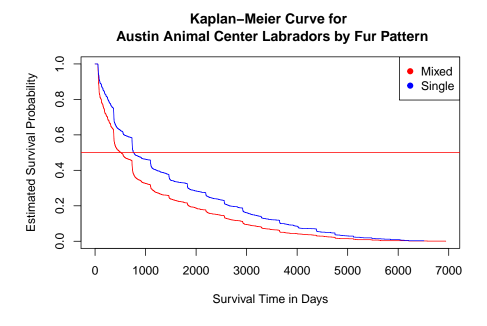


Figure 5: There appears to be an even amount of labs with mixed and solid fur patterns. There seems to be a slight difference in survival probability between the two fur patterns with mixed having a lower survival probability.

Log Rank Tests

The log-rank test allows us to compare 2 or more survival curves, more specifically, that there is a significant difference between the survival curves. The null hypothesis that we are testing is that there is no difference in survival curves. The low p-value means we reject our null hypothesis.

```
## Call:
  survdiff(formula = labradors.surv ~ labradors.df$sex)
##
##
##
                               N Observed Expected (O-E)^2/E (O-E)^2/V
  labradors.df$sex=Female 3211
                                     2442
                                               2347
                                                         3.87
  labradors.df$sex=Male
                                                         3.09
                                                                       7
                            3718
                                     2843
                                               2938
##
    Chisq= 7 on 1 degrees of freedom, p= 0.008
##
  survdiff(formula = labradors.surv ~ labradors.df$neuter)
##
##
                                  N Observed Expected (O-E)^2/E (O-E)^2/V
  labradors.df$neuter=Fixed
                                        4884
                                                  4585
                                                            19.5
                                                                        149
  labradors.df$neuter=Intact 1203
                                         401
                                                   700
                                                           127.9
                                                                        149
##
    Chisq= 149 on 1 degrees of freedom, p= <2e-16
##
## survdiff(formula = labradors.surv ~ labradors.df$color)
##
##
                                N Observed Expected (O-E)^2/E (O-E)^2/V
## labradors.df$color=Mixed 3713
                                      2853
                                                2437
                                                          70.9
                                                                      134
                                                2848
  labradors.df$color=Solid 3216
                                      2432
                                                          60.7
                                                                      134
##
##
    Chisq= 134 on 1 degrees of freedom, p= <2e-16
```

After performing log-rank tests for the 3 covariates (sex, fur pattern, fixed status) individually, we see that the p-value for each log-rank test is less than $\alpha = 0.05$. The null hypothesis for each test is that the survival curves for the covariate are not significantly different, but since the p-values for each test are less than $\alpha = 0.05$, we can conclude that each of the covariates' survival curves are statistically significantly different.

Model Building

To pick the right covariates for our model we use the anova function on the full model which includes all three covariates that we are examining. The anova function below indicates that each of the covariates are significant to include in our model since each one has a p-value of less than 0.05. Since the last covariate also has a significant p-value then we can conclude that each previous covariate in the model is also included and thus also significant. In anova, order of covariates matter and each new covariate is added to the previous model and evaluated.

```
## Analysis of Deviance Table
## Cox model: response is Surv(date_diff, event)
## Terms added sequentially (first to last)
##
```

```
Chisq Df Pr(>|Chi|)
##
          loglik
          -41263
## NULL
## neuter -41177 171.8988
                                  < 2e-16 ***
          -41120 113.6214
                                  < 2e-16 ***
                            1
## sex
          -41117
                    6.0749
                                 0.01371 *
##
  ---
                    0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

We also use the backward elimination method in the step function which also indicates that we should include all three of our covariates. The step function uses the Akaike Information Criterion (AIC) to measure each model quality and gives us the best model for the specificied stepwise search (i.e. backward, forward, or both).

```
## Start: AIC=82239.47
## Surv(date_diff, event) ~ neuter + color + sex
##
##
            Df
                 AIC
## <none>
               82239
## - sex
             1 82244
## - color
             1 82351
## - neuter
             1 82390
## Call:
  coxph(formula = Surv(date_diff, event) ~ neuter + color + sex,
##
       data = labradors.df)
##
##
                    coef exp(coef) se(coef)
                                                           р
## neuterIntact -0.59223
                            0.55309
                                     0.05215 -11.357 <2e-16
                -0.29497
                                     0.02778 -10.617 <2e-16
## colorSolid
                            0.74455
## sexMale
                -0.06811
                            0.93416
                                     0.02761
                                             -2.467 0.0136
##
## Likelihood ratio test=291.6 on 3 df, p=< 2.2e-16
## n= 6929, number of events= 5285
```

Model Checking

In order to build our Cox PH model we must check to see if the proportional hazard assumptions are met. We will do this by graphical evaluation of the log-log plots and by doing a residuals test using the cox.zph() function

When evaluating the log-log plots we want to make sure that the survival curves don't cross and that the curves look proportional to one another through time. These criteria imply that the proportional hazards assumption is met. Figure 6 shows the log-log plot for the sex covariate. We see that the curves cross and overlap both at the beginning and end of the curves. The suggests that the PH assumption is violated. Figure 7 shows the log-log plot for the neuter covariate. We see that the survival curves cross in the beginning, but then seem to be proportional until the very end. Because of this, we will also assume the PH assumption is violated. In Figure 8, the log-log plot for the color covariate shows an increase in the gap between the curves, and then a decline in the size of the gap through time. This suggests that the hazard proportion is not constant and that the PH assumtions do not hold.

Using the Goodness-of-Fit test (table below) to further evaluate the PH assumptions, we test the null hypothesis that there is no correlation between the residuals and survival times. To do this we use the cox.zph() function in R. In the table below we see that for color and sex we reject the null hypothesis. This

Log-log plot of Sex Covariate

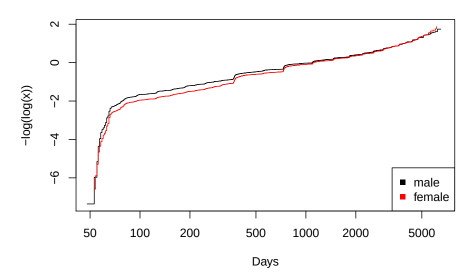


Figure 6: The log-log plot for the sex covariate violates the PH assumptions we are checking because the curves appear to be crossing in various places.

Log-log plot for Neuter Covariate

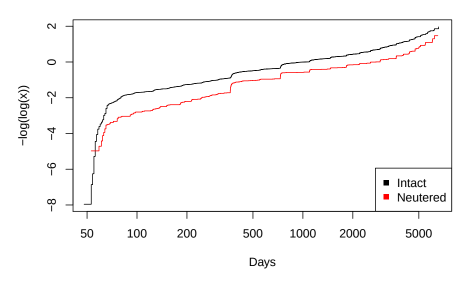


Figure 7: The log-log plot for the neuter covariate shows crossing lines in the very beginning but overall looks parallel after that. We may want to perform another test because there may be a PH violation.

Log-log plot for Color Covariate

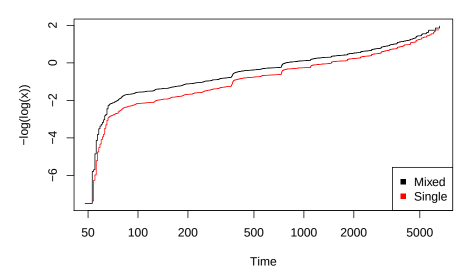


Figure 8: The survival curves appear to get closer to each other as time progresses. This means that the hazard ratio is no longer constant and it depends on time.

means that color and sex violate the PH assumptions. Notice that we do not reject the null hypothesis for the neuter variable which means that PH assumptions hold. Since this is a test and relied more on data than on visual analysis, we will follow the goodness-of-fit test conclusion that the PH assumptions hold for the neuter covariate. Our next step is to deal with the covariates that violate the PH assumption. We will do this by stratifying those covariates.

```
## rho chisq p
## neuterIntact 0.00847 0.379 5.38e-01
## colorSolid 0.07001 25.564 4.28e-07
## sexMale 0.06162 20.026 7.64e-06
```

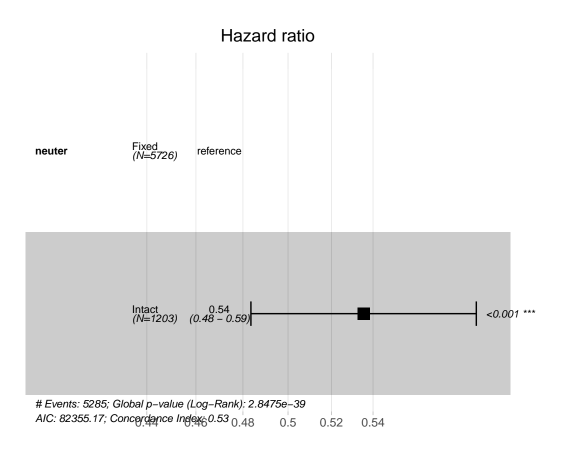
Stratified Cox PH Model

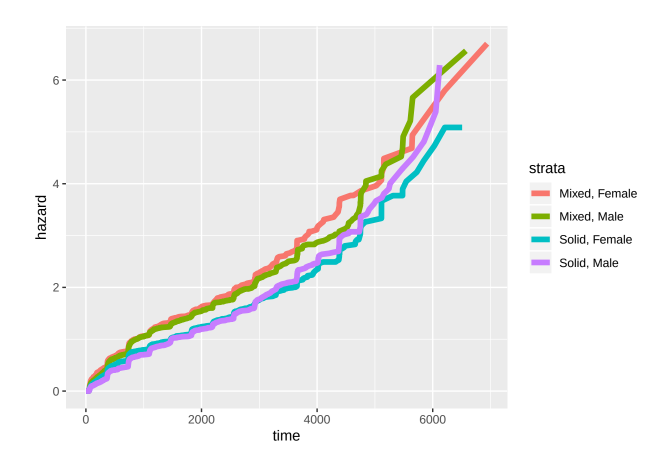
```
## Call:
  coxph(formula = Surv(date_diff, event) ~ neuter + strata(color) +
       strata(sex), data = labradors.df)
##
##
##
     n= 6929, number of events= 5285
##
                   coef exp(coef) se(coef)
##
                                                  z Pr(>|z|)
## neuterIntact -0.5964
                            0.5508
                                     0.0522 -11.43
                                                      <2e-16 ***
  Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
##
                exp(coef) exp(-coef) lower .95 upper .95
## neuterIntact
                   0.5508
                                1.816
                                         0.4972
                                                    0.6101
##
```

Interaction Terms

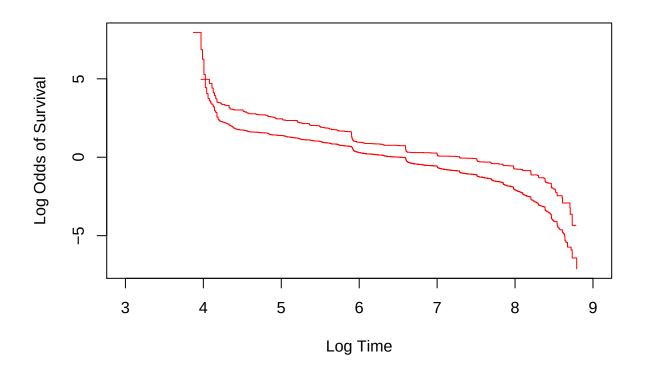
We use the anova function

```
## Analysis of Deviance Table
## Cox model: response is Surv(date_diff, event)
## Model 1: ~ neuter + strata(color) + strata(sex)
## Model 2: ~ neuter + strata(sex) * strata(color)
## loglik Chisq Df P(>|Chi|)
## 1 -33898
## 2 -33898
              0 0 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
## Cox model: response is Surv(date_diff, event)
## Model 1: ~ neuter + strata(color) + strata(sex)
## Model 2: ~ strata(color) + strata(sex) * neuter
##
   loglik Chisq Df P(>|Chi|)
## 1 -33898
## 2 -33894 7.5675 1 0.005943 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table
## Cox model: response is Surv(date_diff, event)
## Model 1: ~ neuter + strata(color) + strata(sex)
## Model 2: ~ strata(sex) + strata(color) * neuter
   loglik Chisq Df P(>|Chi|)
## 1 -33898
## 2 -33898 0.0124 1
## Analysis of Deviance Table
## Cox model: response is Surv(date_diff, event)
## Terms added sequentially (first to last)
##
##
                     loglik
                               Chisq Df Pr(>|Chi|)
## NULL
                     -33975
## neuter
                     -33898 154.2903 1 < 2.2e-16 ***
## neuter:strata(sex) -33894 7.5675 1 0.005943 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Warning in .get_data(model, data = data): The `data` argument is not
## provided. Data will be extracted from model fit.
## Warning: Removed 1 rows containing missing values (geom_errorbar).
```





Extension: Accelerated Failure Times



```
Df Deviance Resid. Df -2*LL
                                      Pr(>Chi)
## NULL NA NA 6927 85292.53
## neuter 1 169.3364
                     6926 85123.20 1.033014e-38
      Df Deviance Resid. Df -2*LL
                                   Pr(>Chi)
## NULL NA NA 6927 85292.53
                     6926 85284.95 0.005911505
## sex 1 7.577082
       Df Deviance Resid. Df
                            -2*LL
                                     Pr(>Chi)
## NULL NA NA 6927 85292.53
## color 1 145.9612
                    6926 85146.57 1.323773e-33
    Terms Resid. Df -2*LL Test Df Deviance
            6927 85335.44
                            NA NA
## 1 neuter
## 2 neuter
              6926 85123.20
                           = 1 212.243 4.451633e-48
    Terms Resid. Df
                    -2*LL Test Df Deviance
                                           Pr(>Chi)
## 1 sex 6927 85494.30
                          NA
                                 NA
             6926 85284.95
                           = 1 209.3457 1.908173e-47
   Terms Resid. Df
                    -2*LL Test Df Deviance
                                           Pr(>Chi)
## 1 color 6927 85324.52 NA NA
            6926 85146.57 = 1 177.9542 1.355545e-40
## 2 color
```

```
##
## Call:
## survreg(formula = Surv(date_diff, event) ~ neuter, data = labradors.df,
      dist = "loglogistic")
                 Value Std. Error
                                       z
                 6.3626
                           0.0195 326.7 <2e-16
## (Intercept)
## neuterIntact 0.7392
                            0.0620 11.9 <2e-16
## Log(scale)
              -0.2045
                           0.0111 -18.5 <2e-16
##
## Scale= 0.815
##
## Log logistic distribution
## Loglik(model) = -42574 Loglik(intercept only) = -42649.5
## Chisq= 151.06 on 1 degrees of freedom, p= 1e-34
## Number of Newton-Raphson Iterations: 4
## n= 6929
##
## Call:
## survreg(formula = Surv(date_diff, event) ~ sex, data = labradors.df,
       dist = "loglogistic")
##
                 Value Std. Error
                                       z
## (Intercept) 6.3480
                           0.0272 233.06 < 2e-16
## sexMale
               0.1708
                           0.0368 4.64 3.4e-06
## Log(scale) -0.2064
                           0.0110 - 18.69 < 2e - 16
##
## Scale= 0.814
##
## Log logistic distribution
## Loglik(model) = -42638.8
                            Loglik(intercept only) = -42649.5
## Chisq= 21.51 on 1 degrees of freedom, p= 3.5e-06
## Number of Newton-Raphson Iterations: 3
## n= 6929
##
## Call:
## survreg(formula = Surv(date_diff, event) ~ color, data = labradors.df,
       dist = "loglogistic")
##
                 Value Std. Error
                                      z
## (Intercept) 6.2143
                          0.0248 250.6 <2e-16
              0.4791
## colorSolid
                           0.0362 13.2 <2e-16
## Log(scale) -0.2203
                           0.0110 -20.0 <2e-16
## Scale= 0.802
## Log logistic distribution
                           Loglik(intercept only)= -42649.5
## Loglik(model) = -42563.1
## Chisq= 172.91 on 1 degrees of freedom, p= 1.7e-39
## Number of Newton-Raphson Iterations: 3
## n= 6929
```