Technical Summary of Facial Attractiveness Project

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PCA:

I ran PCA on these features to see if we really need all these features. The expectation was that we would get a small set of components that upon rotation I could compare with a component for a particular type of facial fetaure. However, it turned out that almost 99% of the variation could be explained by just 1 component. Doing some more reading and spending time thinking about this result made us realize that all these points could be embedded on to a single manifold that upon rotation would collapse to a single linear representation. At this point, I decided to invalidate the use of PCA for our analysis since cutting down to just one feature to explain our variance in the data (which obviously isn't high at all) was not our intention to begin with.

Our plan was to try to run dimensionality reduction and test if it makes sense to use these reduction techniques before we process the data into a regression model. Since it might not make explanatory sense to reduce the entire data set to just one variable, I envision that we will probably keep the data and feed it into a regression model.

Then, I have done a preliminary linear regression, ridge and lasso regression model as well however, Ryan will go more into details for the ridge and lasso models so I'll not include that here. A preliminary linear regression helps us explain ~50% of the variability in the data as seen below. This is consistent with the results in the publication related to this dataset.

```
from sklearn.model_selection import train_test_split
X_train, X_test, y_train, y_test = train_test_split(combined, Y, test_size=0.33, random_state=42)
X_train.shape
(335, 136)
from sklearn import datasets, linear_model
from sklearn.metrics import mean squared error, r2 score
regr = linear model.LinearRegression()
regr.fit(X train, y train)
LinearRegression(copy_X=True, fit_intercept=True, n_jobs=1, normalize=False)
v pred = regr.predict(X train)
print('Coefficients: \n', regr.coef_)
 # The mean squared error
print("Mean squared error: %.2f"
        % mean_squared_error(y_train, y_pred))
# Explained variance score: I is perfect prediction
print('Variance score: % .2f' % r2_score(y_train, y_pred))
plt.scatter(X_test, y_test, color='black')
plt.plot(X_test, y_pred, color='blue', linewidth=3)
plt.xticks(())
plt.yticks(())
plt.show()
Coefficients:
 [[-0.03284617 0.02901486 -0.00622812 0.05476053 -0.0657278
                                                                                       0.12513919
  0.15072231 0.11357349 -0.1258242 -0.08112233 -0.09466987 0.0832176
    0.01377636 0.06988641 -0.11081145 0.01407732 0.00242719 -0.0118359
   0.01127598 -0.19906899 0.08834484 -0.06407751 0.01930778 0.11877733 0.06141545 -0.00673611 -0.11933928 -0.17228017 0.00354433 0.114841
  0.00313741 -0.13280379 0.11653157 -0.00159977 0.10630297 0.03267212 -0.06819599 -0.12826035 0.21393832 0.13062429 0.00376716 -0.23784209
    0.07326854 0.02226835 0.03963797 -0.06537726 -0.04037247 -0.00221746
    0.13002809 -0.05866689 -0.05287474 0.08921138 -0.02677864 -0.02598202
  -0.03856166 0.12217324 -0.08442543 -0.02639694 0.05381557 -0.01355733

        0.00415977
        0.0127093
        -0.01102282
        -0.02290586
        0.00059334
        0.02587144

        -0.01428466
        -0.02770865
        0.057688
        0.0505818
        -0.07432076
        0.00437421

        0.13848209
        -0.224472
        0.08953067
        0.02380393
        0.13310847
        0.0317529

        -0.07766298
        -0.09751175
        0.00786785
        0.14028074
        -0.04785564
        0.02610719

    0.01499493 -0.12629069 -0.0555311 -0.19292885 0.07931195 -0.0506167 
0.02838894 0.15966056 -0.04934273 0.05499867 -0.0956298 -0.02994016
   0.1265898 -0.04924849 0.04114478 0.03002002 -0.12041566 -0.05142445
    0.22281485 -0.06103927 -0.04632633 0.08721767 -0.10411409 0.01499757
    0.03416845 0.07447229 -0.07914449 -0.00583482]]
Mean squared error: 0.22
Variance score: 0.49
```

The other team members were researching the above mentioned so decided to test K- Nearest Neighbor Regression. The main idea is that the target is predicted by local interpolation of the targets associated of the nearest neighbors in the training set.

```
from sklearn.neighbors import KNeighborsRegressor

neigh = KNeighborsRegressor(n_neighbors=2)

neigh.fit(X_train, y_train)

KNeighborsRegressor(algorithm='auto', leaf_size=30, metric='minkowski', metric_params=None, n_jobs=1, n_neighbors=2, p=2, weights='uniform')

pred = (neigh.predict(X_train))
```

upon running a preliminary run in Python, we can see that we can make predictions using the KNN Regressor. The MSE is slightly reduced and the Variance is up by about 5 basis points. We will still go with our results for ridge and lasso regression for better explanation and accuracy.

```
Coefficients:
 [[-0.03284617 0.02901486 -0.00622812 0.05476053 -0.0657278
  -0.1647892 0.08400213 -0.00056747 -0.02802688 -0.0202565 0.02102697
 -0.01424751 0.03276084 -0.05077906 0.06901713 -0.04642176 0.02842491
 -0.06682542 0.07909213 0.00645239 -0.01457146 -0.01454085 -0.01203359
 -0.02614972 0.14420736 -0.04958517 -0.15893938 0.32534808 -0.23791386
  0.15072231 0.11357349 -0.1258242 -0.08112233 -0.09466987 0.0832176
  0.01377636  0.06988641 -0.11081145  0.01407732  0.00242719 -0.0118359
  0.01127598 -0.19906899 0.08834484 -0.06407751 0.01930778 0.11877733
  0.06141545 -0.00673611 -0.11933928 -0.17228017 0.00354433 0.114841
  0.00313741 -0.13280379 0.11653157 -0.00159977 0.10630297
                                                     0.03267212
  -0.06819599 -0.12826035 0.21393832 0.13062429 0.00376716 -0.23784209
  0.13002809 -0.05866689 -0.05287474 0.08921138 -0.02677864 -0.02598202
 -0.03856166 0.12217324 -0.08442543 -0.02639694 0.05381557 -0.01355733
  -0.01428466 -0.02770865 0.057688 0.0505818 -0.07432076 0.00437421
                      0.08953067 0.02380393 0.13310847 0.0317529
  0.13848209 -0.224472
  -0.07766298 -0.09751175 0.00786785 0.14028074 -0.04785564 0.02610719
  0.02838894 0.15966056 -0.04934273 0.05499867 -0.0956298 -0.02994016
  0.1265898 -0.04924849 0.04114478 0.03002002 -0.12041566 -0.05142445
  0.22281485 -0.06103927 -0.04632633 0.08721767 -0.10411409 0.01499757
  0.03416845 0.07447229 -0.07914449 -0.00583482]]
Mean squared error: 0.20
Variance score: 0.53
```

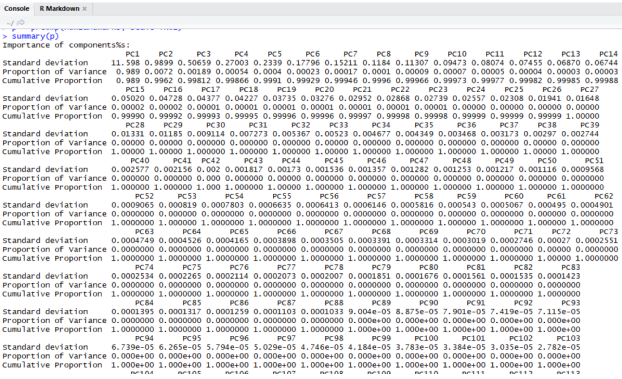
CFA:

Before performing model selection and multiple regression analysis, Principal Component Analysis, Common Factor Analysis and Cluster Analysis were performed on the facial landmarks data. The original data that was used had no definitions of columns. This led to attempting exploratory analyses without understanding the dataset, hence the x1, x2, x3 column names in picture 1. The column names were assigned manually since it represented nothing specific that was given with the data. It was guessed that the physical data were coordinates of some kind of facial structures, since all data were numerical.

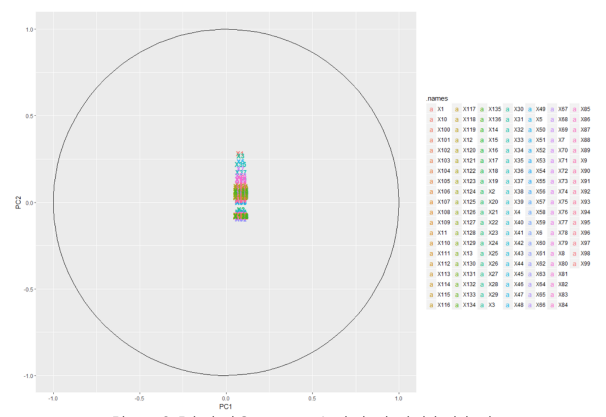


Picture 1: A view of the original dataset

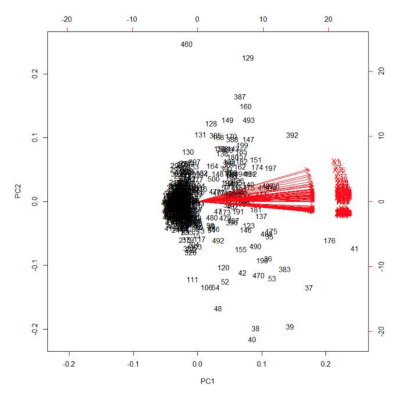
The data ranged anywhere from a low twenties up to more than five hundred. Due to the diverse range of the numerical variables, it was necessary to apply scaling in the Principal Component Analysis. All Principal Component Analyses and rotations used on the facial landmarks data were performed with correlations and not covariances. The output from the Principal Component Analysis demonstrated that it did not have anything significant. The first principal component explained 98.9% of the cumulative proportion in the dataset in output 1. It was logical to conclude that the Principal Component Analysis suggested using only one principal component. In the Principal Component Analysis plot and the bi plot, picture 2 and 3, it was obvious that the entire data showed an extremely linear relationship with each other. The screeplot also recommended to use only one principal component, since the first value was extremely high, followed by low values after in picture 4.



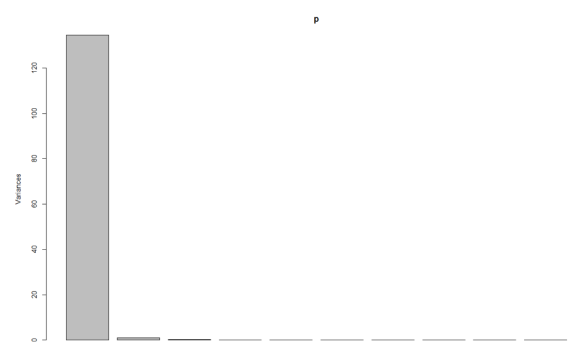
Output 1: Output from the Principal Component Analysis (original data)



Picture 2: Principal Component Analysis plot (original data)



Picture 3: Bi plot (original data)



Picture 4: Scree plot (original data)

After checking the insignificant results from the scaled Principal Component Analysis, rotation with "Varimax" was performed. In output 2, the rotation loadings were very close to 1. All loading values were positive as well. The first principal component explained 98.9 percent of the proportion variance in the data. This supported that only one principal component was recommended.

```
Loadings:
[1] 0.947 0.995 0.953 0.997 0.963 0.997 0.972 0.997 0.998 0.997 0.999 0.997 0.994 0.997 0.996 0.996 0.997 0.996 0.997 0.997 0.997 0.997 [22] 0.997 0.997 0.997 0.997 0.997 0.998 0.999 0.998 0.997 0.998 0.999 0.998 0.999 0.998 0.999 0.998 0.999 0.998 0.999 0.998 0.999 0.998 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999
```

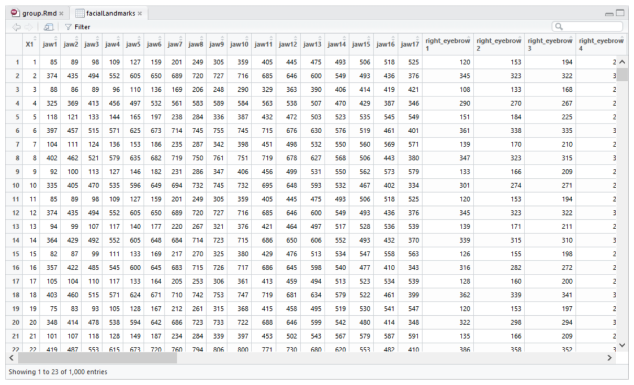
Output 2: Loadings output from the scaled rotation w/ "Varimax" (original data)

Common Factor Analysis was utilized, following the unsatisfying Principal Component Analysis and rotation. In output 3, nfactor of 1 and 2 were tried, but R gave an error stating "unable to optimize from this starting value." Singularity issues might have caused this error, since the relationship between the variables were extremely linear. The lowest nfactor was selected, which was 3. The output of the Common Factor Analysis illustrated that 2 factors were ideal for the facial data.

```
X15
      0.691
               0.713
      0.700
               0.703
X17
X35
      0.576
               0.813
X37
               0.785
      0.613
X39
      0.642
               0.760
               0.741
      0.661
X43
      0.675
               0.726
X55
      0.698
               0.706
X57
      0.697
               0.707
X59
      0.696
               0.707
X61
      0.695
               0.708
x63
      0.683
               0.722
               0.714
X65
      0.690
X67
      0.696
               0.707
               0.774
      0.628
X73
X75
               0.758
      0.647
x77
      0.667
               0.739
x79
      0.678
               0.727
               0.741
      0.665
      0.644
               0.760
X83
      0.668
               0.736
X97
x99
      0.682
               0.722
X101
      0.692
               0.712
X103
      0.698
               0.706
X115
      0.698
               0.705
X117
      0.691
               0.713
x119
      0.681
               0.723
      0.674
               0.731
X121
X123
      0.693
               0.712
X125
      0.698
               0.705
               0.705
X133
      0.698
      0.692
                Factor1 Factor2 Factor3
SS loadings
                 74.258 60.138
                                   1.325
Proportion Var
                  0.546
                          0.442
                                    0.010
Cumulative Var
                  0.546
                           0.988
                                   0.998
```

Output 3: Output from Common Factor Analysis (original data)

Since the original data did not have any given columns, further analysis was impossible. One of the team member, Ryan, processed the original data in python using a code from the developers' resource page in order to gather column names and details of the data. After the data processing, the newly processed data finally had specified column names in picture 5.



Picture 5: A view of the new processed dataset

Unfortunately, the odd rows had the coordinates for x-values, and the even rows had the coordinates for y-values. Therefore manual division into two datasets(x and y) in R was necessary. Although the physical numerical values changed during the python process, the two datasets are essentially the same data. Due to the ranges of the data values, scaling was used for the Principal Component Analysis. In output 4, it portrayed that the first principal component explained 99.2 percent of the cumulative proportion of the data for the x's. In output 5, it portrayed that the first principal component explained 99.7 percent of the cumulative proportion of the data for the y's. Both the x and y Principal Component Analysis outputs recommended only 1 principal components. Similar to the original dataset, the Principal Component Analysis did not show any great results.

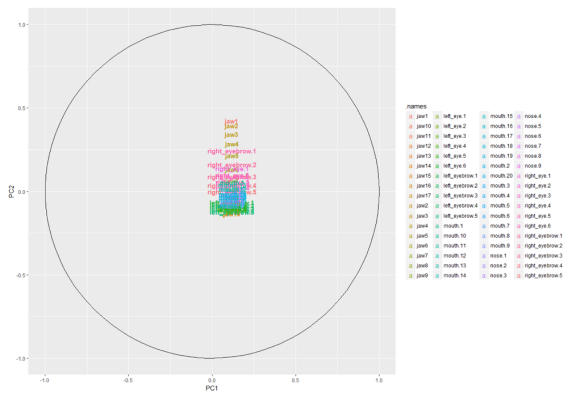
Importance of components%s:
PC1 PC2 PC3 PC4 PC5 PC6 PC7 PC8 PC9 PC10 PC11 PC12 PC13
Standard deviation 8.2151 0.60722 0.27500 0.15905 0.12210 0.09907 0.07232 0.05923 0.04597 0.04117 0.03315 0.02527 0.02440
Proportion of Variance 0.9925 0.00542 0.00111 0.00037 0.00022 0.00014 0.00008 0.00005 0.00003 0.00002 0.00002 0.00001 0.00001
Cumulative Proportion 0.9925 0.99788 0.99900 0.99937 0.99959 0.99973 0.99981 0.99986 0.99989 0.99992 0.99993 0.99994 0.99995
PC14 PC15 PC16 PC17 PC18 PC19 PC20 PC21 PC22 PC23 PC24 PC25 PC26
Standard deviation 0.02255 0.02109 0.01978 0.01796 0.0155 0.01432 0.01335 0.01148 0.01131 0.009883 0.009047 0.008272 0.007834
Proportion of Variance 0.00001 0.00001 0.00001 0.00000 0.00000 0.00000 0.00000 0.00000 0.000000
Cumulative Proportion 0.99996 0.99996 0.99997 0.99998 1.0000 0.99998 0.99999 0.99999 0.999990 0.999990 0.999990 0.999990
PC27 PC28 PC29 PC30 PC31 PC32 PC33 PC34 PC35 PC36 PC37 PC38
standard deviation 0.007224 0.007066 0.006155 0.005783 0.005587 0.005227 0.004991 0.004795 0.004543 0.00442 0.004025 0.003768
Proportion of Variance 0.000000 0.000000 0.000000 0.000000 0.000000
Cumulative Proportion 0.999990 0.999990 0.999990 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000
PC39 PC40 PC41 PC42 PC43 PC44 PC45 PC46 PC47 PC48 PC49 PC50
Standard deviation 0.003435 0.003323 0.003171 0.002925 0.002842 0.002641 0.002588 0.002345 0.002263 0.002177 0.002114 0.002036
Proportion of Variance 0.000000 0.000000 0.000000 0.000000 0.000000
Cumulative Proportion 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.00000000
PC51 PC52 PC53 PC54 PC55 PC56 PC57 PC58 PC59 PC60 PC61 PC62
Standard deviation 0.001967 0.001941 0.001883 0.00178 0.001688 0.00161 0.00157 0.001559 0.001529 0.001479 0.001443 0.001389
Proportion of Variance 0.000000 0.000000 0.000000 0.000000 0.000000
Cumulative Proportion 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.0000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.000000 1.00000000
PC63 PC64 PC65 PC66 PC67 PC68
Standard deviation 0.001368 0.001311 0.001291 0.001276 0.001248 0.001165
Proportion of Variance 0.000000 0.000000 0.000000 0.000000 0.000000
Cumulative Proportion 1.000000 1.000000 1.000000 1.000000 1.000000

Output 4: Output from the Principal Component Analysis for x (processed data)

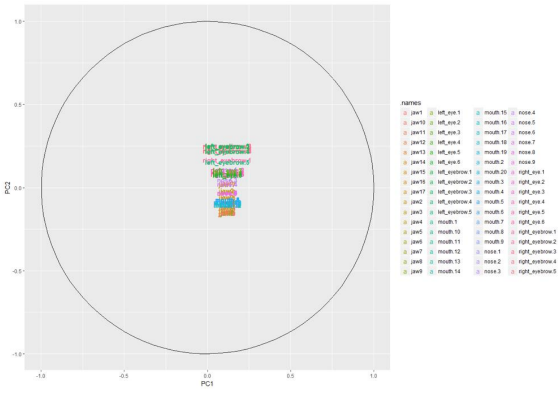
```
Importance of components%s:
                   PC1 PC2 PC3 PC4 PC5 PC6 PC7 PC8 PC9 PC10 PC11 PC12 PC13 8.2365 0.29800 0.17956 0.14775 0.06541 0.05570 0.04975 0.04427 0.03787 0.03245 0.02524 0.02169 0.01934
Standard deviation
                   0.9976 0.00131 0.00047 0.00032 0.00006 0.00005 0.00004 0.00003 0.00002 0.00002 0.00001 0.00001 0.00001 0.9976 0.99894 0.99942 0.99974 0.99980 0.99985 0.99988 0.99991 0.99993 0.99995 0.99996 0.99997 0.99997
Proportion of Variance
Cumulative Proportion
                   PC14 PC15 PC16 PC17 PC18 PC19 PC20 0.01814 0.01684 0.0139 0.01308 0.01186 0.01043 0.0104
                                                                  PC21 PC22
0.009601 0.007767
                                                                                 PC23 PC24 PC25
0.007318 0.007137 0.006707
                                                                                                         0.006262
Standard deviation
PC27 PC28 PC29 PC30 PC31 PC32 PC33 PC34 PC35 0.005524 0.004789 0.004066 0.003715 0.003568 0.003381 0.003351 0.003149 0.00297
Standard deviation
                                                                                        0.002832 0.002563 0.00243
1.000000 1.000000 1.00000
                                                                                        1.000000 1.000000 1.00000
                   PC39 PC40 PC41 PC42 PC43 PC44 PC45 PC46 PC47 0.002344 0.002268 0.00217 0.002065 0.001899 0.001862 0.001781 0.001633 0.001604
                                                                                        PC48 PC49 PC50
0.00155 0.001528 0.001483
Standard deviation
Proportion of Variance
                   Cumulative Proportion
                                     PC53
                                                                                            PC60
                    0.00138 0.001363 0.001317 0.001246 0.001208 0.001183 0.001127 0.001111 0.001084 0.001045 0.0009981 0.0009681
Standard deviation
PC63 PC64 PC65 PC66 PC67 PC68 0.0009478 0.0008826 0.0008522 0.0008373 0.000781 0.0007489
Standard deviation
Cumulative Proportion 1.0000000 1.0000000 1.0000000 1.0000000 1.0000000
```

Output 5: Output from the Principal Component Analysis for y (processed data)

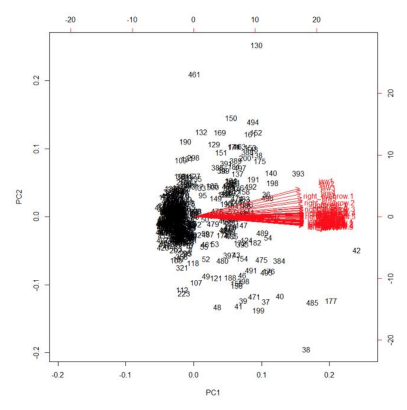
In the Principal Component Analysis plot and the bi plot for both x and y, picture 6, 7, 8 and 9, it was clear that the variables in the facial data had very strongly linear relationships. The screeplots for both x and y also suggested to use only one principal component. The first value had the highest value, followed by low values for both x and y in picture 10 and 11. This was expected in a way, since the original data had similar analyses outputs.



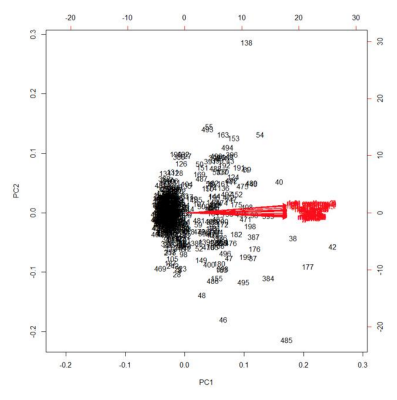
Picture 6: Principal Component Analysis plot for x (processed data)



Picture 7: Principal Component Analysis plot for y (processed data)

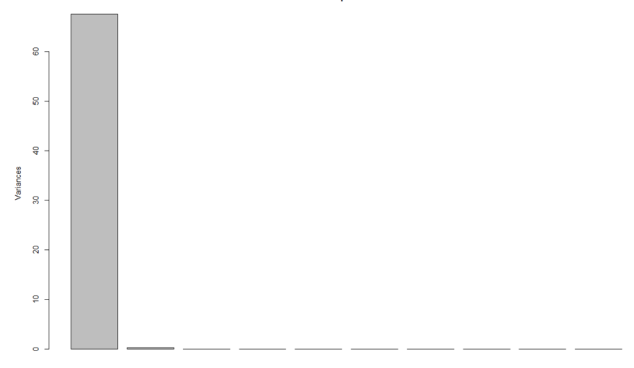


Picture 8: Bi plot for x (processed data)

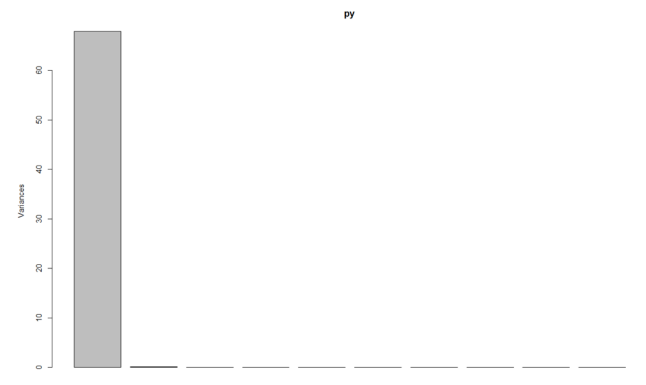


Picture 9: Bi plot for y (processed data)





Picture 10: Scree plot for x (processed data)



Picture 11: Scree plot for y (processed data)

Rotation with "Varimax" was utilized in order to hopefully find a different perspective of the data. However, all the loading values were extremely close to 1 for both x and y. All the loadings were positive as well for both x and y in output 6 and 7. The first principal component explained 99.2 percent and 99.8 percent of the proportion variance for x and y respectively. The root mean square of the residuals were also almost 0 for both x and y. This suggests strong linear relationships between the variables.

```
Loadings:
[1] 0.963 0.970 0.977 0.982 0.987 0.993 0.997 0.998 0.998 0.998 0.998 0.997 0.997 0.996 0.996 0.996 0.996 0.996 0.996 0.998 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.999 0.99
```

Output 7: Loadings output from the scaled rotation w/ "Varimax" for y (processed data)

Test of the hypothesis that 1 factor is sufficient. The degrees of freedom for the model is 2210 and the objective function was 300.51 The number of observations was 500 with Chi Square = 142694.4 with prob < 0

The root mean square of the residuals (RMSA) is 0

Then, Common Factor Analysis was performed to demonstrate any latent factors in the data. In output 8 and 9, nfactor of 1 was tried, but R gave an error stating "unable to optimize from this starting value." Therefore the next lowest value, 2, was selected as the nfactor. For both x and y, 2 factors were chosen, since the Common Factor Analysis stated that 2 factors explained close to 100% of the cumulative variances in the data. Most factor values were in the 0.6 to 0.8 range, which did not really show any significant results.

```
left_eye.4
                 0.799
                         0.600
                 0.796
                         0.604
left_eye.5
                 0.791
                         0.611
left_eye.6
                 0.734
                         0.676
mouth.1
                         0.658
                 0.752
mouth.2
mouth. 3
                         0.645
                         0.636
mouth.4
                 0.771
mouth.5
                         0.629
mouth.6
                 0.784
                         0.620
mouth.7
                 0.791
                         0.610
mouth.8
                 0.785
                         0.619
                 0.777
mouth.9
                         0.628
                 0.771
0.764
mouth.10
                         0.636
mouth.11
                         0.645
                 0.751
0.741
                         0.658
mouth.12
mouth.13
                         0.669
                 0.764
mouth.14
                         0.645
                 0.771
mouth.15
                         0.637
mouth.16
                 0.777
                         0.629
mouth.17
                 0.789
                         0.613
                 0.777
mouth.18
                         0.629
mouth.19
                 0.771
                         0.636
mouth.20
                 0.764
                 0.564
                         0.821
jaw1
jaw2
                 0.581
                         0.812
jaw3
                 0.607
                         0.792
jaw4
                 0.633
                         0.769
jaw5
                 0.664
                         0.741
jaw6
                 0.701
                         0.707
right_eyebrow.1 0.650
                         0.755
right_eyebrow.2 0.688
                         0.722
right_eye.1
                 0.699
                         0.713
                Factor1 Factor2
SS loadings
                 38.944
                        28.900
Proportion Var
                  0.573
                          0.425
Cumulative Var
                  0.573
                          0.998
```

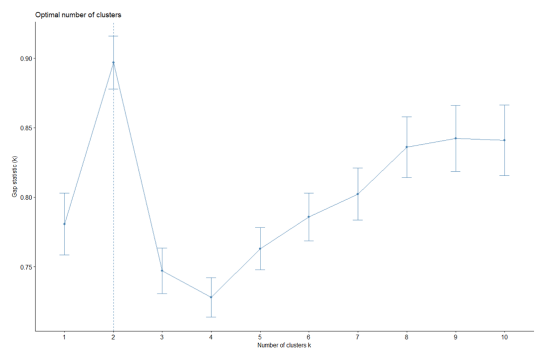
Output 8: Output from Common Factor Analysis for x (processed data)

```
right_eye.1
                 0.745
                         0.666
right_eye.2
right_eye.3
                 0.742
                         0.670
                 0.742
                         0.670
                         0.666
                 0.746
right_eye.4
                         0.665
right_eye.5
                 0.747
                 0.747
                         0.664
right_eye.6
left_eye.1
                 0.747
                         0.665
                 0.742
                         0.669
left_eye.2
left_eye.3
                 0.742
                         0.669
left_eye.4
                 0.746
                         0.665
left_eye.5
                 0.748
                         0.663
left_eye.6
                 0.748
                         0.664
mouth.1
                 0.781
                         0.624
mouth.2
                 0.778
                         0.628
mouth.3
                 0.776
                         0.631
mouth.4
                 0.776
                         0.630
mouth.5
                 0.776
                         0.631
mouth.6
                 0.778
                         0.628
                 0.782
mouth.7
                         0.623
                 0.782
mouth.8
                         0.623
mouth.9
                 0.781
                         0.624
                 0.781
mouth.10
                         0.624
                 0.781
mouth.11
                         0.624
                 0.782
mouth.12
                         0.623
                 0.781
mouth.13
                         0.624
                 0.780
                         0.626
mouth.14
                 0.780
                         0.626
mouth, 15
mouth.16
                 0.780
                         0.626
mouth.17
                 0.781
                         0.624
mouth.18
                 0.778
                         0.627
                 0.778
mouth.19
                         0.627
mouth.20
                 0.778
                         0.627
                Factor1 Factor2
ss loadings
                 39.494
                        28.425
Proportion Var
                  0.581
                          0.418
Cumulative Var
                  0.581
```

Output 9: Output from Common Factor Analysis for y (processed data)

Clustering:

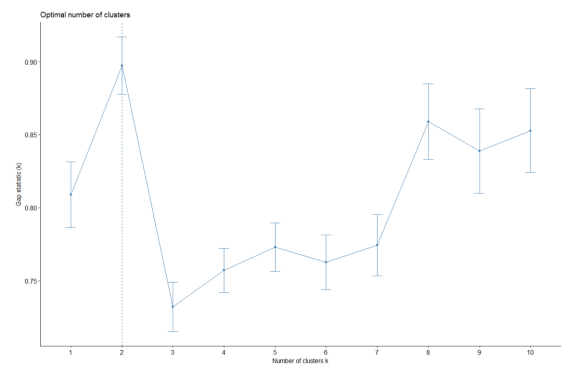
Since Principal Component Analysis and Common Factor Analysis did not show great results, Cluster Analysis was performed additionally. The picture 12 and 14 showed the optimal number of clusters for x and y. 2 clusters were chosen and the plot included how spread apart the data is in picture 13 and 15.



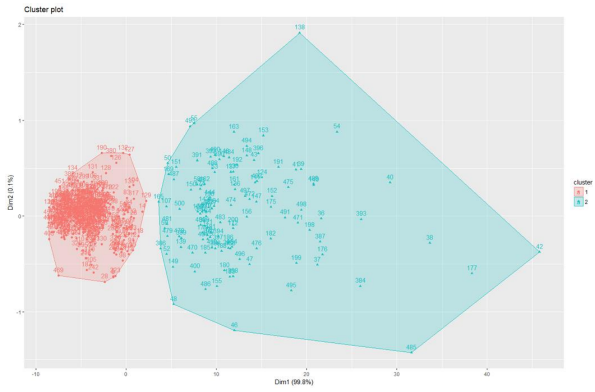
Picture 12: Plot of optimal number of clusters for x (processed data)



Picture 13: Cluster Analysis plot for x (processed data)



Picture 14: Plot of optimal number of clusters for y (processed data)



Picture 15: Cluster Analysis plot for y (processed data)

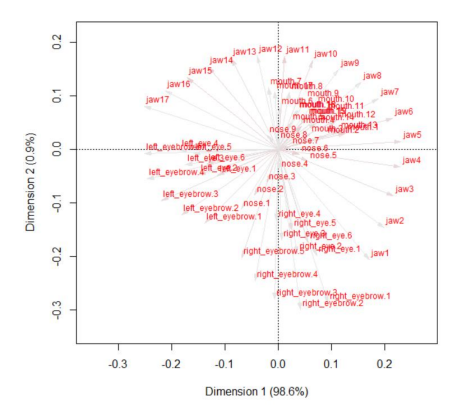
Due to the strong relationships/linearity issues in the data, Principal Component Analysis, rotation, Common Factor Analysis and Cluster Analysis did not bring any great conclusions in terms of model selection. It also does make sense that these analysis methods did not do much, because facial structure can change so much. It illustrated that y values had more linear relationships than x values, since there are more room to change vertically than horizontally when talking about facial landmarks. Additional exploratory analysis methods will be used to determine a suitable model.

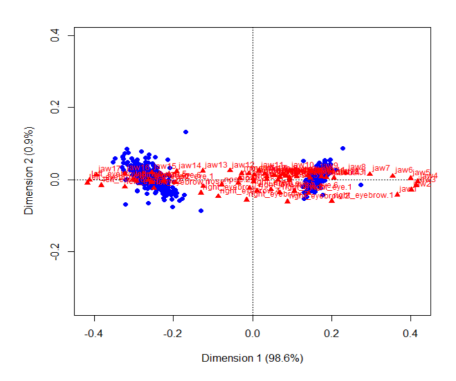
Correspondence Analysis:

I select correspondence analysis to explore the dataset. Firstly, I store all 68 variables and run CA on them. Using the plot, I demonstrate the facial landmarks on the graph. As a result, it shows which variables are best representative. In order to interpret the plot, you need to draw two arrows between the variables. The acute the angle, the more these 2 variables correspond. On the other hand, an obtuse angle between variables means that these 2 variables do not correspond a lot. For instance, as illustrated on the plot below, the angle between 'jaw16' and 'jaw17' is very acute. This means that these 2 variables correspond to each other very well. Also, to the contrary, 'jaw17' and 'righteyebrow' 5 depicts an obtuse angle, which means these 2 variables do not correspond well.

Another visualization of the CA is drawing a line of origin to create a scale. I decided not to use this plot as the visualization was not clearly illustrated due to many variables in our dataset. However, the process goes along like this. Firstly, draw a line of origin through a variable to create a scale. Then, with each variable draw a perpendicular line to the scale. The closer the perpendicular strikes the line, the more these two variables correspond and viceversa. To illustrate to some extent, drawing a line of origin through jaw7 means we created a 'jaw7' scale. By visualizing the perpendicular lines amongst the variables, we can depict that amongst the many variables which will correspond well with 'jaw7' some to mention are 'jaw6','righteyebrow1' and 'jaw8'. On the other side, 'jaw17','jaw14' and 'jaw13' would be some of the many variables which would correspond less.

Generally, our dataset is very correlated as we are dealing with a dataset of the same type. Correspondence analysis is an appropriate technique to explore relationships amongst variable response categories and can play a role in analyzing facial landmark data. Though of its effectiveness, further analysis should be explored to get a better understanding of the dataset.





Correspondence analysis was interesting and valuable to our understanding of the dataset. I used R to create these plots. I stored and ran CA on all 68 variables. After they were stored, I created two different types of plots. The figure below is a screenshot of my code in R which illustrates the process and acquiring of the plots.

Regression and Regularized Regression:

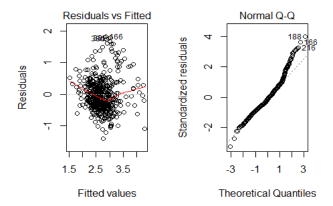
I analysed the face feature data by regression and classification. The goal is to predict exact attractiveness score for each person based on face features. I used multiple linear regression (MLR) as well as principal component regression (PCR).

Firstly, I used a multiple linear regression to explore the relations among the response attractiveness label. In multiple regression model, each part of faces such as left eye has two coordinates x and y. I flatted them into 1 row and gave appropriate names of the features with 'x' and 'y, so i have 136 predictors and 1 response. Then, I applied 10-folds cross validation to check the performance of model. For 10-folds cross validation, I randomized the raw data and divided the randomized data into equally 10 parts. For each part, I used the other 9 parts as the training data to train model and predict on the selected part. Overall, I used MSE as a measurement of performace. My result shows that the MSE is around 0.68, which appears to be pretty good.

```
mse = mean((dataVld$ratings - dataVld$pred)^2)
mse←

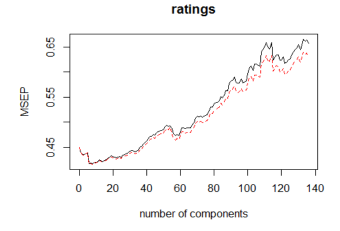
[1] 0.6757903←
```

The result shows that MSE is 0.6757903.



Residuals plot shows there is 'double bow' pattern. The normal qq plot shows there are lots of points far away from the straight line at the tail, indicating model transformation is needed to improve the model.

```
pcr.MSE = mean((predict(fit, dataVld, ncomp = 5)- dataVld$ratings)^2)
pcr.MSE ←
[1] 0.4271767←
```



```
pcr.MSE = mean((predict(fit, dataVld, ncomp = 5)- dataVld$ratings)^2)
pcr.MSE←
[1] 0.4271767←
```

Validation plot shows that best number of components for PCR is 5 as it has lowest MSE 0.4271767.

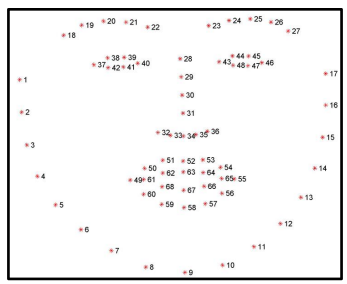
The goal of classification is to predict whether each person is attractive of not based on his/her face features. Here, I used linear discriminant analysis (LDA) and misclassification error rate as measurement to explore the face data.

Firstly, I divided the continuous ratings of faces into two groups. One group is 'Yes' and the other group is 'No' decided by whether the attractiveness ratings is above the mean of ratings. Then I applied 10-fold cross validation and computed the overall accuracy in classification. The score ranges from 1 to 5, with an average of 3. Above average score, I used label 'Yes', which means attractive. Below avarage score, I used label 'No', which means not attractive. My result shows that the overall accuracy is about 0.608. It appears that our LDA model is a little better than randomly selection of 'Yes' or 'No'. One reason might be collinearity in the features of faces. The other reason, I think, might be the cut-off of the attractiveness is not suitable. From the LDA result, we can see there are 108 mistakes when predicting people looking good as bad. Overall, LDA model is very efficient and useful. In order to improve our performance, we can try other techniques such as Lasso Regression.

There were some issues with the original data downloaded from the website. First, none of the data was labelled. There was no columns to go off of so we could not understand any of the results we would have gotten. Second, we did not have access to the python landmark tool to test on our own images. While this would not have been the end of the world it was something I wanted to do.

The process of getting the landmarks was involved and had many issues. I had very little experience with Python and the libraries needed only worked within Linux. The Python code will be listed in the Appendix. So after many hours of troubleshooting I was able to run our 500 image data set through the script using a for loop to code all at once. Below is an image of the facial landmarks that are numbered. Each one of those landmarks had an x and y coordinate and this is the data we would use for our modeling.

The way that the python script was grabbing facial features in of itself was a machine learning technique as I imported a pre-trained model to grab the features from our images. This model had been given hundreds of different faces with pre-mapped features and then was able to find these features on new images. I used for loops and array manipulation to efficiently process all of our images so once the script was running it was fairly easy to extract the data to a usable file.



68 landmarks making up 7 major portions of the face.

The regression process first started with splitting the data which now has the shape of 500x136. I used the floor function to split both the facial data (Dependent data) and the attractiveness ratings (Independent data). I started with the full model using the training data:

```
Residual standard error: 0.607 on 238 degrees of freedom Multiple R-squared: 0.5013, Adjusted R-squared: 0.2164 F-statistic: 1.759 on 136 and 238 DF, p-value: 7.305e-05

> mape <- mean(abs((fullPredict - testy))/testy)
> mape
[1] 0.2341761
```

As you can see this model is fairly average and most noteable you can see the difference between the R^2 and adj. R^2 because we are working with 136 variables. After this I then proceeded to perform Backwards stepwise selection to automatically select the variables. This gave pretty good improvements but still this model is no where near perfect. Adj R^2 of ~.35, MAPE of .224, RMSE of 1.22 and MSE of .69. This model improves in all aspects over the full model but the error values are still very high.

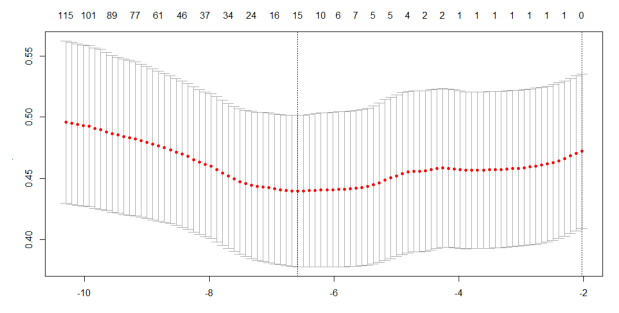
```
Residual standard error: 0.5529 on 308 degrees of freedom
Multiple R-squared: 0.4645, Adjusted R-squared: 0.3498
F-statistic: 4.049 on 66 and 308 DF, p-value: < 2.2e-16

> mape <- mean(abs((predictions - testy))/testy) [1] 1.224385
> mape
[1] 0.2239169

| mean(abs() | mean(abs() | mean() |
```

At this point I know that we have incredibly correlated data and that adding a penalty constraint on our regression could make a much more reliable model. I first performed ridge regression but halfway through I realized that it would be nice if we could automatically eliminate some of our 136 variables. So I decided to proceed with Lasso Regression. I did this using Cross validation and glmnet library to test for a wide range of lambda's to find the best one for our model. Below

(Graph 1) is the plot of the lambda's attempting to find the minimize the MSE (y axis). The final lambda we get is .001387 so that is the lambda we will use for our regression.



So after fitting our model we get 17 remaining variables. There seems to be a huge emphasis on jaw points in terms of attractiveness. Below is the full breakdown of the variables. The biggest representation amonst the variables is the Jaw group of x and y variables. 9 of our 17 variables were related to jaw structure. However if you look at the impact of each variable jaw7, mouth 5 y, jaw9 y and nose 7 y had the biggest individual impacts.

```
"2.51768583564636"
 (Intercept)"
"jaw1"
                   "-0.000846857956314382"
"jaw5"
                   "0.0083918385021834"
                   "-0.000346279554129708"
 iaw8"
"jaw12"
                   "-0.00148983913777072"
"jaw13"
                   "-2.26526868919727e-08"
"right_eyebrow 1" "-0.00635888417389142"
                   "0.00017826613898461"
"right_eyebrow 5"
                   "0.000833280498734204"
"left_eyebrow 4"
                   "-0.00458380337084622"
"jawl y
"jaw7 y"
                   "4.5306074801391e-05"
"jaw8 y"
                   "0.000129577379228305"
"jaw9 y"
                   "3.03443105097005e-09"
"nose 4 y"
                   "0.000864785092852011"
"nose 7 y"
                   "2.4683011315565e-05"
                   "0.0035838566350521"
"left_eye 5
                   "4.21562440709026e-05"
"mouth 5 y"
```

We can then run the same prediction analysis on this model as we did the other two. We got the output below. This is a much better model in terms of prediction on our testing data.

```
> rsePredict
[1] 0.6517649
> mse
[1] 0.3670251
> mapeLasso <- mean(abs((yPredict -
> mapeLasso
[1] 0.1710937
```

LASSO regression is a means of working with correlated data and minimizing the model to get the least amount of error. LASSO stands for least absolute shrinkage and selection. The name itself describes what it is doing. LASSO creates a lambda in which is a penalty parameter in order to reduce over fitting. The benefit of using LASSO over ridge in this example is that we wanted to eliminate some of our variables and LASSO sets weak impact features to 0 whereas Ridge keeps all variables and just makes coefficients smaller. Since our model had 136 variables and we were unable to do PCA

Lasso regression seemed like the perfect fit for our data. This showed as it gave us the best outcomes based on the 5 models we created.

When this model was used for prediction on the face of a team member and the face of a celebrity to test it on outside data it responded with unexpected results. The celebrity's face was at an angle and not completely centered. There are some issues when testing computer vision models like this however. The model is incredibly affected by the position, angle, and expression on the face. There are many ways this project could be improved. For example we could have created a much more massive data set that took the distances between all points on the face and use that instead of our data. This is much more complicated and harder to manage than our data but it would have ironed out the inconsistencies between faces in the original data set and the outside test images.

The practical uses of this analysis are pretty self explanatory as they could be used to rate attractiveness. However this is only for one ethnicity and attractiveness is incredibly subjective so building a perfect model would never be possible. One person who one group of raters thinks is attractive could be the opposite for another group. I do not believe the real world uses of this would extend far beyond personal use for recreation as it is interesting to see if what machine learning predicts your attractiveness as. I see no real applications even in professions such as modeling since there is much more than just base attractiveness in people.