# **STATS 3DA3**

## Homework Assignment 6

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- 1. A classification problem based on the dataset by Rubini, Soundarapandian, and Eswaran (2015) is to classify an individual into has or doesn't have chronic kidney disease based on the following predictor variables: age, blood pressure (mm/Hg), specific gravity, albumin, sugar, red blood cells, pus cell, pus cell clumps, bacteria, blood glucose random (mgs/dl), blood urea (mgs/dl), serum creatinine (mgs/dl), sodium (mEq/L), potassium (mEq/L), hemoglobin (gms), packed cell volume, white blood cell count (cells/cmm), red blood cell count (millions/cmm), hypertension, diabetes mellitus, coronary artery disease, appetite, pedal edema, and anemia.
- 2. We import the data and transform it as necessary.

```
import pandas as pd
import numpy as np
from ucimlrepo import fetch_ucirepo
import certifi
chronic_kidney_disease = fetch_ucirepo(id=336)
X = chronic_kidney_disease.data.features
y = chronic_kidney_disease.data.targets
data = pd.concat([X, y], axis = 1)
```

To check if any variable transformations are necessary, we check that the data types of the variables in the dataframe match the description in the data dictionary.

### print(data.dtypes)

```
float64
age
          float64
bp
          float64
sg
al
          float64
          float64
su
rbc
           object
           object
рс
           object
рсс
```

```
object
ba
         float64
bgr
         float64
bu
         float64
sc
sod
         float64
         float64
pot
hemo
         float64
          float64
pcv
         float64
wbcc
rbcc
         float64
           object
htn
dm
           object
           object
cad
appet
           object
           object
ре
           object
ane
class
           object
dtype: object
```

We can see that age, blood pressure (bp), blood glucose random (bgr), blood urea (bu), serum creatinine (sc), sodium (sod), potassium (pot), hemoglobin (hemo), packed cell volume (pcv), white blood cell count (wbcc), and red blood cell count (rbcc) are all numerical values as they should be according to the data dictionary (note the data dictionary differentiates between integer and continuous values but changing numeric data types will not affect results). However, specific gravity (sg), albumin (al), and sugar (su) should be nominal but are float values. Red blood cells (rbc), pus cell (pc), pus cell clumps (pcc), bacteria (ba), hypertension (htn), diabetes mellitus (dm), coronary artery disease (cad), appette (appet), pedal edema (pe), anemia (ane), and class should all be nominal instead of object as they appear now. We correct the data type errors by converting the noted variables to categorical/nominal.

```
columns_to_convert = ['sg','al','su','rbc','pc','pcc','ba','htn','dm','cad','appet','pe','ane'
for col in columns_to_convert:
    data[col] = pd.Categorical(data[col])
```

In order to make the variables match the abbreviations found in the data dictionary, we rename whose and rbcc to we and rc, respectively.

```
data = data.rename(columns={'wbcc':'wc'})
data = data.rename(columns={'rbcc':'rc'})
```

3. We explore the data and give a detailed description of the dataset.

```
print(data.shape)
print(data.iloc[0:2,0:12])
print(data.iloc[0:2,12:26])
(400, 25)
    age
            bp
                   sg
                        al
                              su
                                  rbc
                                             рс
                                                         рсс
                                                                        ba
                                                                               bgr
   48.0
          80.0
                1.02
                       1.0
                             0.0
                                  {\tt NaN}
                                        normal
                                                 notpresent
                                                              notpresent
                                                                            121.0
          50.0
                1.02
                       4.0
                             0.0
                                  {\tt NaN}
                                        normal
                                                 notpresent notpresent
                                                                               NaN
     bu
           sc
   36.0
         1.2
   18.0
         0.8
   sod
        pot
              hemo
                      pcv
                                          htn
                                                 dm cad appet
                                                                 pe ane class
   NaN
        NaN
              15.4
                     44.0
                            7800.0
                                     5.2
                                                                           ckd
                                          yes
                                                yes
                                                      no
                                                          good
                                                                      no
   NaN
         NaN
              11.3
                     38.0
                            6000.0
                                     NaN
                                           no
                                                 no
                                                      no
                                                          good
                                                                 no
                                                                      no
                                                                           ckd
```

There are 25 variables and 400 observations. The variables include an individual's age in years, their blood pressue in mm/Hg, their specific gravity, their albumin level, their sugar level, the normality of their red blood cells, the normality of their pus cells, their presence of pus cell clumps, their presence of bacteria, their blood glucose random level in mgs/dl, their blood urea level in mgs/dl, their serum creatinine level in mgs/dl, their sodium level in mEq/L, their potassium level in mEq/L, their hemoglobinl level in gms, their packed cell volume, their white blood cell count in cells/cumm, their red blood cell count in millions/cmm, if they have hypertension, if they have

diabetes mellitus, if they have coronary artery disease, if they have a good or poor appetite, if they have pedal edema, if they have anemia, and if they have chronic kidney diease.

Next we create data summaries, including observation proportions.

```
for col in columns_to_convert:
    print(data[col].value_counts(normalize=True))
data.describe(include = 'all')
sg
1.020
         0.300283
1.010
         0.237960
1.025
         0.229462
1.015
         0.212465
1.005
         0.019830
Name: proportion, dtype: float64
al
0.0
       0.562147
1.0
       0.124294
2.0
       0.121469
3.0
       0.121469
4.0
       0.067797
5.0
       0.002825
Name: proportion, dtype: float64
su
0.0
       0.826211
2.0
       0.051282
3.0
       0.039886
1.0
       0.037037
4.0
       0.037037
5.0
       0.008547
Name: proportion, dtype: float64
rbc
```

normal 0.810484

abnormal 0.189516

Name: proportion, dtype: float64

рс

normal 0.773134

abnormal 0.226866

Name: proportion, dtype: float64

рсс

notpresent 0.893939

present 0.106061

Name: proportion, dtype: float64

ba

notpresent 0.944444

present 0.055556

Name: proportion, dtype: float64

htn

no 0.630653

yes 0.369347

Name: proportion, dtype: float64

 $\mathtt{dm}$ 

no 0.653266

yes 0.344221

\tno 0.002513

Name: proportion, dtype: float64

cad

no 0.914573

yes 0.085427

Name: proportion, dtype: float64

appet

good 0.794486

poor 0.205514

Name: proportion, dtype: float64

ре

no 0.809524

yes 0.190476

Name: proportion, dtype: float64

ane

no 0.849624

yes 0.150376

Name: proportion, dtype: float64

class

ckd 0.620

notckd 0.375

ckd\t 0.005

Name: proportion, dtype: float64

age	bp sg al	su rbc pc	pcc	ba bgr	]	ocv wc	rc htn	dm cad	appet pe	ane cla
count	391.000000	388.000000	353.00	354.0	351.0	248	335	396	396	356.000
unique	NaN	NaN	5.00	6.0	6.0	2	2	2	2	NaN
top	NaN	NaN	1.02	0.0	0.0	normal	normal	notpresent	notpresent	NaN
freq	NaN	NaN	106.00	199.0	290.0	201	259	354	374	NaN
mean	51.483376	76.469072	NaN	NaN	NaN	NaN	NaN	NaN	NaN	148.036
std	17.169714	13.683637	NaN	NaN	NaN	NaN	NaN	NaN	NaN	79.2817
min	2.000000	50.000000	NaN	NaN	NaN	NaN	NaN	NaN	NaN	22.0000
25%	42.000000	70.000000	NaN	NaN	NaN	NaN	NaN	NaN	NaN	99.0000
50%	55.000000	80.000000	NaN	NaN	NaN	NaN	NaN	NaN	NaN	121.000
75%	64.500000	80.000000	NaN	NaN	NaN	NaN	NaN	NaN	NaN	163.000
max	90.000000	180.000000	NaN	NaN	NaN	NaN	NaN	NaN	NaN	490.000

We can see that all variables, except chronic kidney disease, have missing values since no counts equal 400. We note that dm and class have 3 levels instead of their expected two. From the proportion counts of categorical variables, we can see that all categories are represented however some are severely under-represented. After correcting dm and ckd, we explore these distributions more through histograms and bar plots.

```
data = data.replace('\tno', 'no')
data = data.replace('ckd\t', 'ckd')
```

```
C:\Users\seeba\AppData\Local\Temp\ipykernel_23940\2882268402.py:1: FutureWarning: The behavior
data = data.replace('\tno', 'no')
C:\Users\seeba\AppData\Local\Temp\ipykernel_23940\2882268402.py:2: FutureWarning: The behavior
data = data.replace('ckd\t', 'ckd')
```

Next we re-examine the data types.

#### data.dtypes

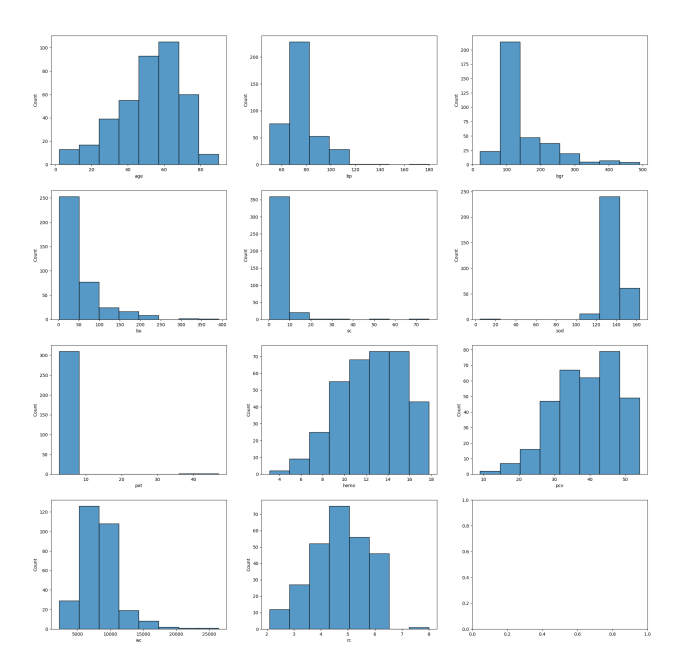
float64 age float64 bp sg category al category category su rbc category category рс category рсс category ba float64 bgr float64 bu float64 sc float64 sod float64 pot float64 hemo pcv float64 WC float64 float64 rccategory htn category dm

```
cad category
appet category
pe category
ane category
class category
dtype: object
```

The data types are only category (nominal) or numerical, as we previously transformed them to be such.

Finally, we look at variables' distributions, starting with numeric variables.

```
import seaborn as sns
import matplotlib.pyplot as plt
figure, axes = plt.subplots(4,3, sharex= False, figsize=(24,24))
k = 0
numeric_vars = data.select_dtypes(include='float64').columns
for i in range(3):
    for j in range(3):
        sns.histplot(ax = axes[i,j], data=data, x = str(numeric_vars[k]), binwidth=(data[str(numeric_vars.histplot(ax = axes[3,0], data=data, x = str(numeric_vars[9]), binwidth=(data[str(numeric_vars.histplot(ax = axes[3,1], data=data, x = str(numeric_vars[10]), binwidth=(data[str(numeric_vars.histplot(ax = axes[3,1], data=data, x = str(numeric_vars.histplot(ax = axe
```



Some variables are heavily skewed such as white blood cell count, potassium, serum creatinine, blood urea, blood pressure, and blood glucose random. Conversely, red blood cell count appears relatively normally distributed.

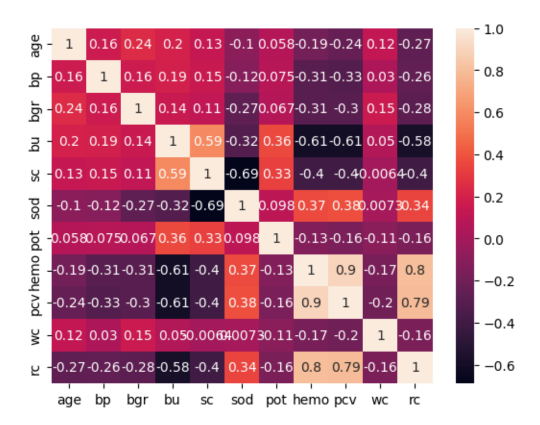
```
import seaborn as sns
import matplotlib.pyplot as plt
figure, axes = plt.subplots(7,2, sharex= False, figsize=(26,30))
k = 0
cat_vars = data.select_dtypes(include='category').columns
```

```
for i in range(7):
          for j in range(2):
                    sns.countplot(ax = axes[i,j], data=data, x = str(cat_vars[k]))
                   k = k+1
plt.show()
                                                                                                            200 - 175 - 150 - 125 - 100 - 75 - 25 - 0
                                                                                                           200 -
175 -
150 -
125 -
100 -
75 -
50 -
25 -
                                                                                                            300
250
250
200
150
100
300 -
250 -
50 200 -
150 -
                                                                                                            # 150
100
                                                                                                            300 -
250 -
50 -
150 -
100 -
50 -
200 -
150 -
                                                                                                            200
150
                                                                                                              100
300 -
250 -
200 -
150 -
                                                                                                            割 150
100
```

We notice most categorical variables are heavily skewed. For example, hypertension, diabetes mellitus, coronary artery disease, appetite, pedal edema, and anemia all have many more instances of "no" than "yes". Red blood cells and pus cell display similar trends where by "normal" is more frequent than "abnormal". Pus cell clumps and bacteria also display this trend where "present" is far less frequent than "not present". It is important to note that class (ckd, notckd) does not display this heavy skew. Although "ckd" is more frequent than "notckd", the two are relatively even.

4. We now analyze variable relationships. We use a heatmap to check for correlation between numerical variables.

```
num_col = data.select_dtypes(include = 'float64').columns
sns.heatmap(data[num_col].corr(), annot=True)
```



From this heatmap we can see hemoglobin and packed cell volume, hemoglobin and red blood cell count, and packed cell volume and red blood cell count have a high positive correlation (>0.75). Conversely, sodium and serum creatinine, hemoglobin and blood urea, and packed cell volume and blood urea have a high negative correlation (<-0.6). The least correlated variables are white blood

cell count and serum creatinine (0.0064). These correlations will likely cause feature selection to drop one of the variables in highly correlated variables.

5. We remove observations with missing values in order to conduct subgroup analysis and for future use.

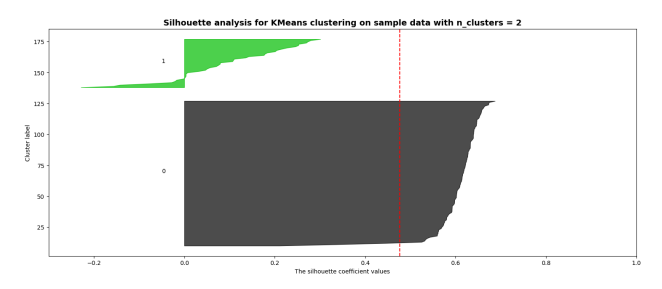
```
data = data.dropna(ignore_index=True) # drop observations with missing values
```

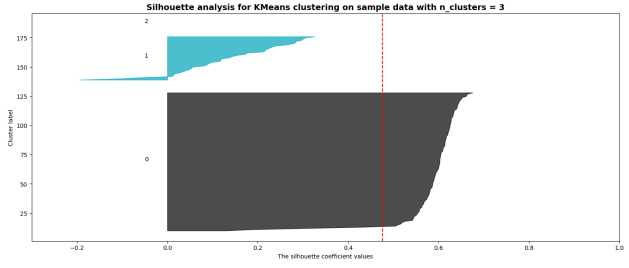
While this greatly reduces the data's sample size, it avoids adding bias by entering other values for missing values (eg. mean, median, or mode), or filling them in another way.

- 6. We do not conduct outlier analysis. This is because outliers can be very important in diagnosing diseases in health data. By removing outliers, we would miss the opportunity to see how extreme values can help in finding disease risks.
- 7. We now conduct using K-means clustering on the data, after scaling and creating dummy variables, to conduct sub-group analysis.

```
from sklearn.cluster import KMeans
from sklearn.metrics import silhouette_samples, silhouette_score
import matplotlib.cm as cm
from sklearn.preprocessing import scale
scaled_data = data.copy()
scaled_data[num_col] = pd.DataFrame(scale(data[num_col]))
dummy = pd.get_dummies(data = scaled_data, columns = cat_vars)
scaled_data = scaled_data.drop(cat_vars, axis = 1)
scaled_data = pd.concat([scaled_data, dummy], axis = 1)
scaled_data = scaled_data.drop(['class_ckd','class_notckd'], axis = 1)
range_n_clusters = [2, 3]
for n_clusters in range_n_clusters:
    km = KMeans(n_clusters = n_clusters, n_init = 20, random_state=0)
   cluster_labels_km = km.fit_predict(scaled_data)
   # average silhouette score
   silhouette_avg_km = silhouette_score(scaled_data, cluster_labels_km)
```

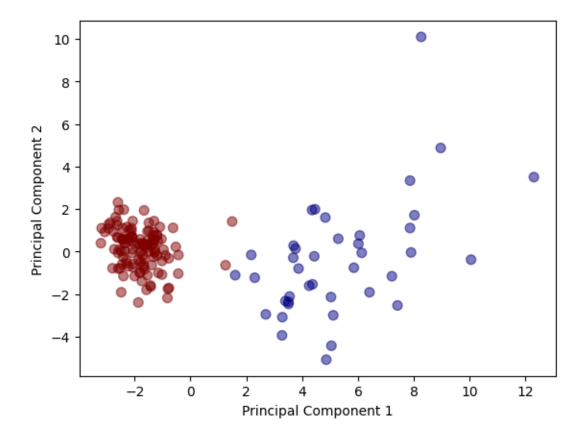
```
# compute the silhouette scores for each sample
sample_silhouette_values = silhouette_samples(scaled_data, cluster_labels_km)
fig, ax1 = plt.subplots(1, 1)
fig.set_size_inches(18, 7)
ax1.set_xlim([-0.3, 1])# change this based on the silhouette range
y_{lower} = 10
for i in range(n_clusters):
    # Aggregate the silhouette scores for samples belonging to
    # cluster i, and sort them
    ith_cluster_silhouette_values = sample_silhouette_values[cluster_labels_km == i]
    ith_cluster_silhouette_values.sort()
    size_cluster_i = ith_cluster_silhouette_values.shape[0]
   y_upper = y_lower + size_cluster_i
    color = cm.nipy_spectral(float(i) / n_clusters)
    ax1.fill_betweenx(
       y=np.arange(y_lower, y_upper),
       x1=0,
       x2=ith_cluster_silhouette_values,
       facecolor=color,
        edgecolor=color,
        alpha=0.7,
   )
    # label the silhouette plots with their cluster numbers at the middle
    ax1.text(-0.05, y_lower + 0.5 * size_cluster_i, str(i))
    # Compute the new y_lower for next cluster silhouette scores
   y_lower = y_upper + 10
ax1.set_title("The silhouette plot for various cluster")
ax1.set_xlabel("The silhouette coefficient values")
ax1.set_ylabel("Cluster label")
# vertical line for average silhouette score of all the values
ax1.axvline(x=silhouette_avg_km, color="red", linestyle="--")
plt.title(
```





There appears to be two subgroups, we now visualize them using PCA with two PCs.

```
km1 = KMeans(n_clusters=2, n_init=20, random_state=0)
km1.fit(scaled_data)
cluster_labels_km1 = km1.fit_predict(scaled_data)
```



From this analysis, we can see there is a clearly defined sub-group in the bottom left corner based on the data.

8. We now split the unscaled, original data into a 70/30 training/test split for testing, using a random seed of 1 and stratified sampling.

```
from sklearn.model_selection import train_test_split

y = data['class']

x = data.drop('class', axis = 1)

X_train, X_test, y_train, y_test = train_test_split(

x, y, test_size=0.3, random_state=1, stratify=y)
```

- 9. We choose two classifiers to address the classification problem. First we use k-nearest neighbours (KNN) then we use a decision tree. We choose these classifiers as they are supervised learning methods and decision trees are interpretable. Also, classic k-nearest neighbours is used for numerical variables only while decision trees can handle mixed variable types, allowing us to see how including categorical variables can affect performance.
- 10. We will use accuracy and sensitivity to compare the performance of the classifiers. Accuracy compares the overall performance while sensitivity will compare how good the classifier is at correctly identifying an individual with chronic kidney disease.
- 11. We use lasso regression to determine important features for KNN while decision trees automatically perform feature selection. We use the previous data which was scaled and uses dummy variables and select the corresponding testing and training values.

```
from sklearn.linear_model import Ridge, Lasso, ElasticNet, RidgeCV, LassoCV, ElasticNetCV
x_train_num = scaled_data.iloc[X_train.index]
y_train_num = y_train.cat.codes
x_test_num = scaled_data.iloc[X_test.index]
lasso_cv = LassoCV(alphas=np.logspace(-4, 4, 100), cv=5, max_iter=1500)
lasso_cv.fit(x_train_num, y_train_num)
m_lasso = Lasso(alpha=lasso_cv.alpha_)
m_lasso.fit(x_train_num, y_train_num)
m_lasso_pre = m_lasso.predict(x_test_num)
pd.DataFrame({'Feature': x_train_num.columns, 'Coefficient': m_lasso.coef_.reshape(len(x_train_num.columns), 'Coefficient': m_lasso.coef_.reshape(len(
```

	Feature	Coefficient
1	bp	-0.00000
2	bgr	-0.00000
3	bu	-0.00000
4	sc	-0.00000
5	$\operatorname{sod}$	0.00000
6	pot	-0.00000
7	hemo	0.00000
8	pcv	0.000000
9	wc	-0.00000
10	$\operatorname{rc}$	0.00000
11	age	-0.00000
12	bp	-0.00000
13	bgr	-0.00009
14	bu	-0.00000
15	sc	-0.00000
16	$\operatorname{sod}$	0.00010
17	pot	-0.00000
18	hemo	0.00013
19	pcv	0.00026
20	wc	-0.00006
21	rc	0.00000
22	$sg\_1.005$	0.00000
23	$sg\_1.01$	-0.00000
24	$sg\_1.015$	-0.00000
25	$sg\_1.02$	0.00000
26	$sg\_1.025$	0.00000
27	al_0.0	0.998354
28	al_1.0	-0.00000
29	al_2.0	-0.00000
30	al 3.0	-0.00000

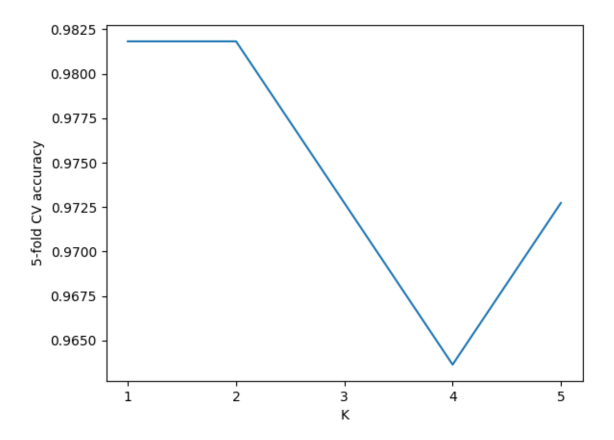
	Feature Coe	fficient
31	al_4.0	-0.000000
32	al_5.0	0.000000
33	su_0.0	0.000000
34	su_1.0	-0.000000
35	su_2.0	-0.000000
36	$su_3.0$	-0.000000
37	su_4.0	-0.000000
38	su_5.0	-0.000000
39	$rbc\_abnormal$	-0.000000
40	$rbc\_normal$	0.000000
41	pc_abnormal	-0.000000
42	$pc\_normal$	0.000000
43	pcc_notpresent	0.000000
44	pcc_present	-0.000000
45	ba_notpresent	0.000000
46	ba_present	-0.000000
47	htn_no	0.000000
48	htn_yes	-0.000000
49	dm_no	0.000000
50	$dm\_yes$	-0.000000
51	cad_no	0.000000
52	$cad\_yes$	-0.000000
53	$appet\_good$	0.000000
54	appet_poor	-0.000000
55	pe_no	0.000000
56	pe_yes	-0.000000
57	ane_no	0.000000
58	ane_yes	-0.000000

Using lasso regression, we see that many variables have been given a coefficient of zero. Specifically

all coefficients have been shrunk to zero except blood glucose random, sodium, hemoglobin, packed cell volume, white blood cell count, and albumin. Note that albumin received by far the largest coefficient.

12. We now implement 5-fold cross validation for KNN with the scaled numerical variables not given a coefficient of 0 in the lasso regression process, and decision tree with all the variables. First we perform cross-validation using the training set to select the best value of k.

```
## KNN Implementation
from sklearn.model_selection import cross_val_score
X_train_red = X_train.drop(['age', 'bp', 'sg', 'al', 'su', 'rbc', 'pc', 'pcc', 'ba', 'bu', 'sc', 'pot',
X_test_red = X_test.drop(['age', 'bp', 'sg', 'al', 'su', 'rbc', 'pc', 'pcc', 'ba', 'bu', 'sc', 'pot', 's
X_test_red = scale(X_test_red)
X_train_red = scale(X_train_red)
from sklearn import neighbors
from sklearn import metrics
k_range = range(1, 6)
cv_scores = []
for k in k_range:
    knn_cv = neighbors.KNeighborsClassifier(n_neighbors=k)
    cv_scores_k = cross_val_score(knn_cv, X_train_red, y_train, cv=5)
    cv_scores.append(np.mean(cv_scores_k))
plt.plot(k_range, cv_scores)
plt.xlabel('K')
plt.ylabel('5-fold CV accuracy')
plt.xticks(range(1,6))
plt.show()
```



Selecting K = 2 to maximize accuracy and minimize variance:

```
knn = neighbors.KNeighborsClassifier(n_neighbors = 2)
knn.fit(X_train_red,y_train)
pred = knn.predict(X_test_red)
from sklearn.metrics import mean_squared_error, confusion_matrix, classification_report
cm = pd.DataFrame(confusion_matrix(y_test, pred), index=['No CKD', 'CKD'], columns=['No CKD',
sensitivity = cm.iloc[1,1]/(cm.iloc[1,0]+cm.iloc[1,1])
print('Sensitivity: ', sensitivity)
print('Accuracy: ', round(metrics.accuracy_score(y_test, pred),2))
```

Sensitivity: 0.9714285714285714

Accuracy: 0.98

Now using a decision tree using the original data with dummy variables for categorical variables (all variables, unscaled).

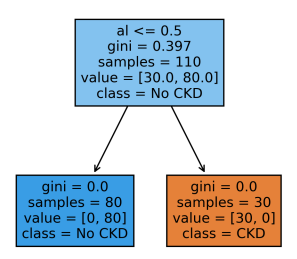
```
from sklearn.tree import DecisionTreeClassifier, DecisionTreeRegressor, plot_tree
for col in cat_vars.drop('class'):
    X_train[col] = pd.Categorical(X_train[col]).codes
    X_test[col] = pd.Categorical(X_test[col]).codes

cs_dt = DecisionTreeClassifier(
    max_depth = 15,
    random_state=1
)
cs_dt.fit(X_train, y_train)
```

DecisionTreeClassifier(max\_depth=15, random\_state=1)

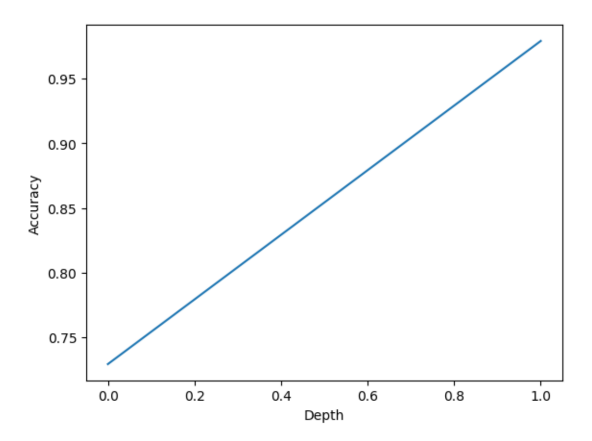
```
## Decision Tree
from sklearn.tree import DecisionTreeClassifier, DecisionTreeRegressor, plot_tree
cs_dt = DecisionTreeClassifier(
    max_depth = 15,
    random_state=1
    )
cs_dt.fit(X_train, y_train)
fig, axes = plt.subplots(
    nrows = 1,ncols = 1,figsize = (4,4), dpi=300
    )
plot_tree(
    cs_dt,
    max_depth= 5,
    feature_names = X_train.columns.tolist(),
    class_names=['CKD', 'No CKD'],
    filled = True
    )
```

```
[Text(0.5, 0.75, 'al <= 0.5\ngini = 0.397\nsamples = 110\nvalue = [30.0, 80.0]\nclass = No CKD Text(0.25, 0.25, 'gini = 0.0\nsamples = 80\nvalue = [0, 80]\nclass = No CKD'),
```



Using a pruning tree to maximize accuaracy:

```
plt.ylabel('Accuracy')
plt.show()
```



Here, it appears the variable albumin is such a prominent variable in detecting chronic kidney disease that no other variables are considered in the process.

```
cs_dt = DecisionTreeClassifier(
    max_depth = 1,
    random_state=1
)

cs_dt.fit(X_train, y_train)

pred = cs_dt.predict(X_test)

cm = pd.DataFrame(confusion_matrix(y_test, pred), index=['No CKD', 'CKD'], columns=['No CKD',
    cm.index.name = 'True'

cm.columns.name = 'Predicted'

sensitivity = cm.iloc[1,1]/(cm.iloc[1,0]+cm.iloc[1,1])
```

```
print('Sensitivity : ', sensitivity)
print('Accuracy : ', cs_dt.score(X_test, y_test))
```

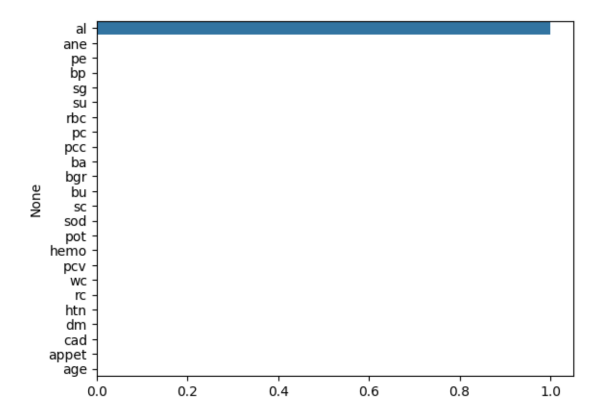
Sensitivity: 1.0

Accuracy: 0.979166666666666

We can see KNN using scaled variables blood glucose random, sodium, hemoglobin, packed cell volume, and white blood cell count gives an accuracy of 0.98 and a sensitivity of 0.97 in the classification of chronic kidney disease and no kidney disease. The decision tree, which had access to all variables, unscaled, only used albumin and had an accuracy of 0.98 and a sensitivity of 1, with one case of no chronic kidney disease being labelled a chronic kidney disease. Overall, the decision tree performed better despite only using one variable.

13. We re-train the interpretable classifier (decision tree) using all the data and analyze and interpret the significance of the predictor variables.

```
## Decision Tree
for col in cat_vars.drop('class'):
    x[str(col)] = pd.Categorical(x[str(col)]).codes
from sklearn.tree import DecisionTreeClassifier, DecisionTreeRegressor, plot_tree
cs_dt_best = DecisionTreeClassifier(
    max_depth = 1,
    random_state=0
    )
cs_dt_best.fit(x, y)
fea_imp = cs_dt_best.feature_importances_
sorted_indices = fea_imp.argsort()[::-1]
sorted_feature_names = X_train.columns[sorted_indices]
sorted_importances = fea_imp[sorted_indices]
sns.barplot(x = sorted_importances, y = sorted_feature_names)
plt.show()
```



From this plot, we can see albumin is the only important variable in determining an individual's risk of chronic kidney disease according to the decision tree. Specifically, if albumin is 0, the patient is very unlikely to have chronic kidney disease, otherwise they likely do have ckd. This could be due to the sample size and/or test train split. Using random forests or dropping albumin may reveal other important variables.

14. Now split the sub-groups identified in question 7 to improve the decision tree accuracy.

```
xtrain1 = []
xtrain2 = []
ytrain1 = []
ytrain2 = []
for i in range(len(X_train)):
    if cluster_labels_km1[i] == 0:
        xtrain1.append(X_train.iloc[i])
        ytrain1.append(y_train.iloc[i])
    else:
```

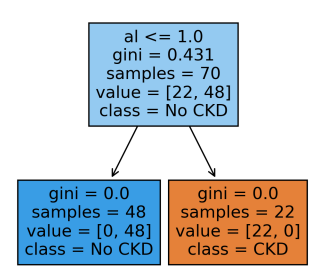
```
xtrain2.append(X_train.iloc[i])
ytrain2.append(y_train.iloc[i])

xtest1 = []
xtest2 = []
ytest1 = []
ytest2 = []
for i in range(len(X_test)):
    if cluster_labels_km1[i] == 0:
        xtest1.append(X_test.iloc[i])
        ytest1.append(y_test.iloc[i])
    else:
        xtest2.append(X_test.iloc[i])
```

```
## Using decsion tree
x1_dt = DecisionTreeClassifier(
    max_depth = 1,
    random_state=1
)
x1_dt.fit(xtrain1,ytrain1)
fig, axes = plt.subplots(
    nrows = 1,ncols = 1,figsize = (4,4), dpi=300
    )
plot_tree(
    x1_dt,
    max_depth= 1,
    feature_names = X_train.columns.tolist(),
    class_names=['CKD', 'No CKD'],
    filled = True
    )
```

 $[Text(0.5, 0.75, 'al \le 1.0 \neq 0.431 \le 70 \le = 70]$ 

```
Text(0.25, 0.25, 'gini = 0.0\nsamples = 48\nvalue = [0, 48]\nclass = No CKD'),
Text(0.75, 0.25, 'gini = 0.0\nsamples = 22\nvalue = [22, 0]\nclass = CKD')]
```



```
x1_dt.fit(xtrain1, ytrain1)
pred = x1_dt.predict(xtest1)
cm = pd.DataFrame(confusion_matrix(ytest1, pred), index=['No CKD', 'CKD'], columns=['No CKD',
cm.index.name = 'True'
cm.columns.name = 'Predicted'
sensitivity = cm.iloc[1,1]/(cm.iloc[1,0]+cm.iloc[1,1])
print('Sensitivity : ', sensitivity)
print('Accuracy : ', x1_dt.score(xtest1, ytest1))

Sensitivity : 1.0
Accuracy : 1.0

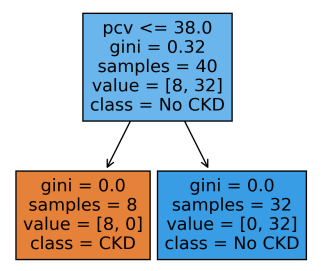
## Using decsion tree
x2_dt = DecisionTreeClassifier(
    max_depth = 1,
    random_state=1
```

```
x2_dt.fit(xtrain2,ytrain2)
fig, axes = plt.subplots(
    nrows = 1,ncols = 1,figsize = (4,4), dpi=300
)
plot_tree(
    x2_dt,
    max_depth= 1,
    feature_names = X_train.columns.tolist(),
    class_names=['CKD', 'No CKD'],
    filled = True
)
```

```
[Text(0.5, 0.75, 'pcv <= 38.0\ngini = 0.32\nsamples = 40\nvalue = [8, 32]\nclass = No CKD'),

Text(0.25, 0.25, 'gini = 0.0\nsamples = 8\nvalue = [8, 0]\nclass = CKD'),

Text(0.75, 0.25, 'gini = 0.0\nsamples = 32\nvalue = [0, 32]\nclass = No CKD')]
```



```
x2_dt.fit(xtrain2, ytrain2)
pred = x2_dt.predict(xtest2)
cm = pd.DataFrame(confusion_matrix(ytest2, pred), index=['No CKD', 'CKD'], columns=['No CKD',
```

```
cm.index.name = 'True'

cm.columns.name = 'Predicted'

sensitivity = cm.iloc[1,1]/(cm.iloc[1,0]+cm.iloc[1,1])

print('Sensitivity : ', sensitivity)

print('Accuracy : ', x2_dt.score(xtest2, ytest2))
```

Sensitivity: 1.0

Accuracy: 1.0

The accuracy and sensitivity for the first and second subgroup is 1. In both question 12 (without sub-group considerations), the accuracy is 0.98 and the sensitivity is 1. Therefore, the accuracy is slightly improved after first splitting the data based on K-means clustering, with K=2.

- 15. Entirely individual submission.
- 16. Here is the link to the public GitHub repository https://github.com/seebachl/3DA3-Asgmt-6.

### **Bibliography**

Rubini, L., Soundarapandian P., and P. Eswaran. 2015. "Chronic Kidney Disease." UCI Machine Learning Repository.