3. Plotting for Exploratory data analysis (EDA)

→ (3.1) Basic Terminology

- · What is EDA?
- · Data-point/vector/Observation
- Data-set.
- Feature/Variable/Input-variable/Dependent-varibale
- Label/Indepdendent-variable/Output-varible/Class/Class-label/Response label
- Vector: 2-D, 3-D, 4-D,.... n-D

Q. What is a 1-D vector: Scalar

▼ Iris Flower dataset

Toy Dataset: Iris Dataset: [https://en.wikipedia.org/wiki/Iris_flower_data_set]

- · A simple dataset to learn the basics.
- 3 flowers of Iris species. [see images on wikipedia link above]
- 1936 by Ronald Fisher.
- Petal and Sepal: http://terpconnect.umd.edu/~petersd/666/html/iris_with_labels.jpg
- Objective: Classify a new flower as belonging to one of the 3 classes given the 4 features.
- Importance of domain knowledge.
- Why use petal and sepal dimensions as features?
- Why do we not use 'color' as a feature?

```
import pandas as pd
import seaborn as sns
import matplotlib.pyplot as plt
import numpy as np

'''downlaod iris.csv from https://raw.githubusercontent.com/uiuc-cse/data-fa14/gh-pages/data/
#Load Iris.csv into a pandas dataFrame.
iris = pd.read_csv("iris.csv")
```

```
# (Q) how many data-points and features?
print (iris.shape)
     (150, 5)
#(Q) What are the column names in our dataset?
print (iris.columns)
     Index(['sepal_length', 'sepal_width', 'petal_length', 'petal_width',
            'species'],
           dtype='object')
#(Q) How many data points for each class are present?
#(or) How many flowers for each species are present?
iris["species"].value counts()
# balanced-dataset vs imbalanced datasets
#Iris is a balanced dataset as the number of data points for every class is 50.
     virginica
                   50
     setosa
                   50
     versicolor
                   50
     Name: species, dtype: int64
```

→ (3.2) 2-D Scatter Plot

```
#2-D scatter plot:
#ALWAYS understand the axis: labels and scale.
iris.plot(kind='scatter', x='sepal_length', y='sepal_width');
plt.show()
#cannot make much sense out it.
#What if we color the points by thier class-label/flower-type.
```

```
# 2-D Scatter plot with color-coding for each flower type/class.

# Here 'sns' corresponds to seaborn.

sns.set_style("whitegrid");

sns.FacetGrid(iris, hue="species", size=4) \
    .map(plt.scatter, "sepal_length", "sepal_width") \
    .add_legend();

plt.show();

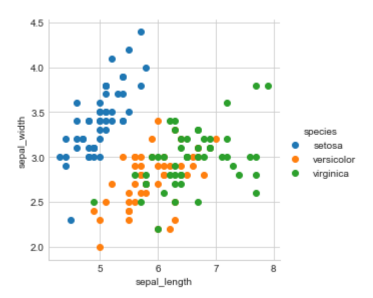
# Notice that the blue points can be easily seperated

# from red and green by drawing a line.

# But red and green data points cannot be easily seperated.

# Can we draw multiple 2-D scatter plots for each combination of features?

# How many cobinations exist? 4C2 = 6.
```



Observation(s):

- 1. Using sepal_length and sepal_width features, we can distinguish Setosa flowers from others.
- 2. Seperating Versicolor from Viginica is much harder as they have considerable overlap.

3D Scatter plot

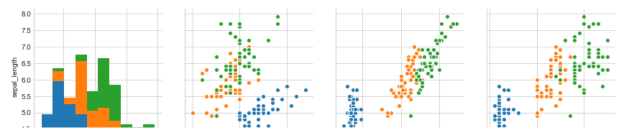
https://plot.ly/pandas/3d-scatter-plots/

Needs a lot to mouse interaction to interpret data.

What about 4-D, 5-D or n-D scatter plot?

→ (3.3) Pair-plot

```
# pairwise scatter plot: Pair-Plot
# Dis-advantages:
##Can be used when number of features are high.
##Cannot visualize higher dimensional patterns in 3-D and 4-D.
#Only possible to view 2D patterns.
plt.close();
sns.set_style("whitegrid");
sns.pairplot(iris, hue="species", size=3);
plt.show()
# NOTE: the diagnol elements are PDFs for each feature. PDFs are expalined below.
```



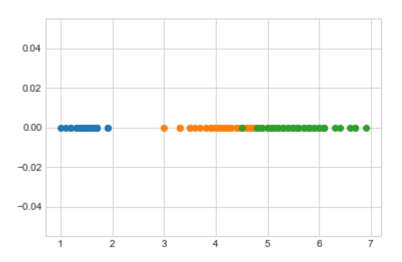
Observations

- 1. petal_length and petal_width are the most useful features to identify various flower types.
- 2. While Setosa can be easily identified (linearly seperable), Virnica and Versicolor have some overlap (almost linearly seperable).
- 3. We can find "lines" and "if-else" conditions to build a simple model to classify the flower types.

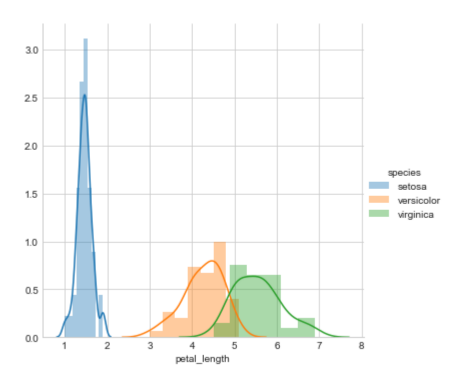
→ (3.4) Histogram, PDF, CDF

```
# What about 1-D scatter plot using just one feature?
#1-D scatter plot of petal-length
import numpy as np
iris_setosa = iris.loc[iris["species"] == "setosa"];
iris_virginica = iris.loc[iris["species"] == "virginica"];
iris_versicolor = iris.loc[iris["species"] == "versicolor"];
#print(iris_setosa["petal_length"])
plt.plot(iris_setosa["petal_length"], np.zeros_like(iris_setosa['petal_length']), 'o')
plt.plot(iris_versicolor["petal_length"], np.zeros_like(iris_versicolor['petal_length']), 'o'
plt.plot(iris_virginica["petal_length"], np.zeros_like(iris_virginica['petal_length']), 'o')

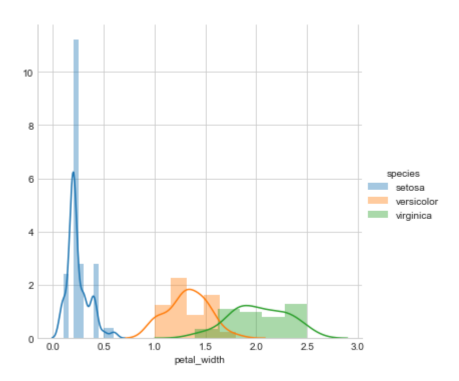
plt.show()
#Disadvantages of 1-D scatter plot: Very hard to make sense as points
#are overlapping a lot.
#Are there better ways of visualizing 1-D scatter plots?
```



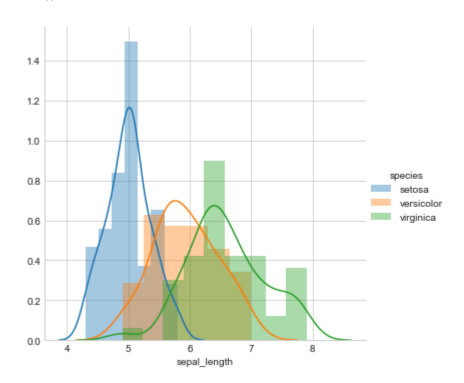
```
sns.FacetGrid(iris, hue="species", size=5) \
    .map(sns.distplot, "petal_length") \
    .add_legend();
plt.show();
```



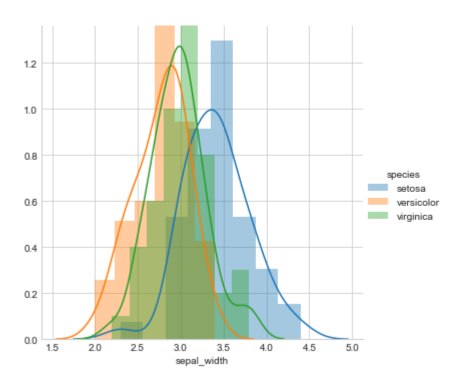
```
sns.FacetGrid(iris, hue="species", size=5) \
    .map(sns.distplot, "petal_width") \
    .add_legend();
plt.show();
```



```
sns.FacetGrid(iris, hue="species", size=5) \
    .map(sns.distplot, "sepal_length") \
    .add_legend();
plt.show();
```



```
sns.FacetGrid(iris, hue="species", size=5) \
    .map(sns.distplot, "sepal_width") \
    .add_legend();
plt.show();
```



How to compute PDFs using counts/frequencies of data points in each window.

```
# How window width effects the PDF plot.
# Interpreting a PDF:
## why is it called a density plot?
## Why is it called a probability plot?
## for each value of petal length, what does the value on y-axis mean?
# Notice that we can write a simple if..else condition as if(petal_length) < 2.5 then flower
# Using just one feature, we can build a simple "model" suing if..else... statements.
# Disadv of PDF: Can we say what percentage of versicolor points have a petal length of less
# Do some of these plots look like a bell-curve you studied in under-grad?
# Gaussian/Normal distribution.
# What is "normal" about normal distribution?
# e.g: Hieghts of male students in a class.
# One of the most frequent distributions in nature.
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 5?
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of petal_length
counts, bin edges = np.histogram(iris setosa['petal length'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges);
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf);
plt.plot(bin edges[1:], cdf)
counts, bin_edges = np.histogram(iris_setosa['petal_length'], bins=20,
                                 density = True)
pdf = counts/(sum(counts))
plt.plot(bin_edges[1:],pdf);
plt.show();
```

```
0.24
                                      0.28 0.14 0.08
     [ 0.02 0.02 0.04 0.14
                                                          0.
                                                                0.04]
     [ 1.
             1.09
                   1.18 1.27
                                1.36
                                       1.45
                                             1.54
                                                   1.63
                                                          1.72 1.81 1.9 ]
      1.0
      0.8
      0.6
      0.4
      0.2
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 1.6?
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of petal_length
counts, bin_edges = np.histogram(iris_setosa['petal_length'], bins=10,
                                   density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
#compute CDF
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
plt.show();
     [ 0.02
             0.02 0.04
                          0.14
                                0.24
                                       0.28
                                             0.14
                                                   0.08
                                                          0.
                                                                0.04]
     [ 1.
             1.09
                   1.18
                          1.27
                                1.36
                                       1.45
                                             1.54
                                                   1.63
                                                         1.72
                                                                1.81 1.9 ]
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
          1.1
               1.2
                     1.3
                          1.4
                               1.5
                                    1.6
                                         1.7
                                               1.8
                                                    1.9
```

```
# Plots of CDF of petal_length for various types of flowers.
# Misclassification error if you use petal length only.
counts, bin edges = np.histogram(iris setosa['petal length'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
# virginica
counts, bin_edges = np.histogram(iris_virginica['petal_length'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges)
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin edges[1:], cdf)
#versicolor
counts, bin_edges = np.histogram(iris_versicolor['petal_length'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges)
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin edges[1:], cdf)
plt.show();
```

```
[ 0.02 0.02 0.04
                   0.14
                         0.24
                               0.28
                                     0.14
                                           0.08
                                                 0.
                                                       0.04]
                               1.45
       1.09 1.18
                   1.27
                                     1.54
                                                       1.81
                         1.36
                                           1.63
                                                 1.72
                                                             1.9 ]
                                                 0.02
 0.02
       0.1
             0.24
                   0.08
                         0.18
                               0.16
                                     0.1
                                           0.04
                                                       0.061
       4.74 4.98
                   5.22
                         5.46
                               5.7
                                     5.94
                                           6.18
                                                 6.42
                                                       6.66 6.9 1
 4.5
 0.02 0.04 0.06
                                                 0.14
                   0.04
                         0.16
                               0.14
                                     0.12
                                           0.2
                                                       0.08]
       3.21
            3.42
                         3.84
                               4.05
                                     4.26
                                           4.47 4.68
                                                      4.89 5.1 ]
 3.
                   3.63
```

(3.5) Mean, Variance and Std-dev

```
#Mean, Variance, Std-deviation,
print("Means:")
print(np.mean(iris_setosa["petal_length"]))
#Mean with an outlier.
print(np.mean(np.append(iris_setosa["petal_length"],50)));
print(np.mean(iris_virginica["petal_length"]))
print(np.mean(iris_versicolor["petal_length"]))
print("\nStd-dev:");
print(np.std(iris_setosa["petal_length"]))
print(np.std(iris_virginica["petal_length"]))
print(np.std(iris_versicolor["petal_length"]))
```

```
Means:
1.464
2.41568627451
5.552
4.26
Std-dev:
0.171767284429
0.546347874527
0.465188133985
```

→ (3.6) Median, Percentile, Quantile, IQR, MAD

```
#Median, Quantiles, Percentiles, IQR.
print("\nMedians:")
print(np.median(iris_setosa["petal_length"]))
#Median with an outlier
print(np.median(np.append(iris_setosa["petal_length"],50)));
print(np.median(iris_virginica["petal_length"]))
print(np.median(iris_versicolor["petal_length"]))
```

```
print("\nQuantiles:")
print(np.percentile(iris setosa["petal length"],np.arange(0, 100, 25)))
print(np.percentile(iris virginica["petal length"],np.arange(0, 100, 25)))
print(np.percentile(iris versicolor["petal length"], np.arange(0, 100, 25)))
print("\n90th Percentiles:")
print(np.percentile(iris setosa["petal length"],90))
print(np.percentile(iris virginica["petal length"],90))
print(np.percentile(iris_versicolor["petal_length"], 90))
from statsmodels import robust
print ("\nMedian Absolute Deviation")
print(robust.mad(iris_setosa["petal_length"]))
print(robust.mad(iris virginica["petal length"]))
print(robust.mad(iris versicolor["petal length"]))
     Medians:
     1.5
     1.5
     5.55
     4.35
     Quantiles:
            1.4
                    1.5
                            1.575]
     [ 1.
             5.1
     [ 4.5
                     5.55
                            5.875]
     [ 3.
                   4.35 4.6 1
             4.
     90th Percentiles:
     1.7
     6.31
     4.8
     Median Absolute Deviation
     0.148260221851
     0.667170998328
     0.518910776477
```

→ (3.7) Box plot and Whiskers

```
#Box-plot with whiskers: another method of visualizing the 1-D scatter plot more intuitivey.

# The Concept of median, percentile, quantile.

# How to draw the box in the box-plot?

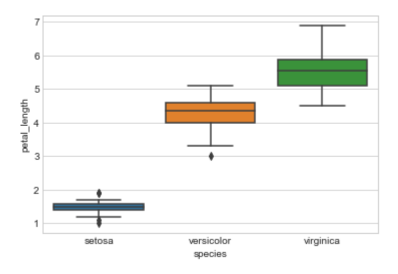
# How to draw whiskers: [no standard way] Could use min and max or use other complex statisti

# IQR like idea.
```

#NOTE: IN the plot below, a technique call inter-quartile range is used in plotting the whisk #Whiskers in the plot below donot correposnd to the min and max values.

#Box-plot can be visualized as a PDF on the side-ways.

sns.boxplot(x='species',y='petal_length', data=iris)
plt.show()

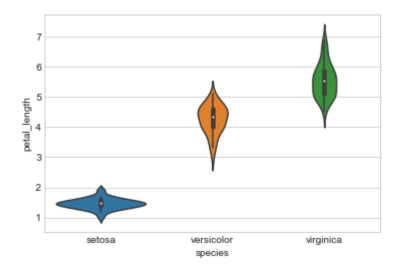


→ (3.8) Violin plots

 $\mbox{\#}$ A violin plot combines the benefits of the previous two plots $\mbox{\#}\mbox{and}$ simplifies them

Denser regions of the data are fatter, and sparser ones thinner #in a violin plot

sns.violinplot(x="species", y="petal_length", data=iris, size=8)
plt.show()



→ (3.9) Summarizing plots in english

- Exaplain your findings/conclusions in plain english
- Never forget your objective (the probelm you are solving). Perform all of your EDA aligned with your objectives.

(3.10) Univariate, bivariate and multivariate analysis.

```
Def: Univariate, Bivariate and Multivariate analysis.

File "<ipython-input-20-f25211abae88>", line 3
    Def: Univariate, Bivariate and Multivariate analysis.

SyntaxError: invalid syntax

SEARCH STACK OVERFLOW
```

→ (3.11) Multivariate probability density, contour plot.

```
#2D Density plot, contors-plot
sns.jointplot(x="petal_length", y="petal_width", data=iris_setosa, kind="kde");
plt.show();
```

→ (3.12) Exercise:

"#Cancer Survival dataset"

- Download Haberman Cancer Survival dataset from Kaggle. You may have to create a Kaggle account to donwload data. (https://www.kaggle.com/gilsousa/habermans-survival-data-set)
- 2. Perform a similar alanlaysis as above on this dataset with the following sections:
- High level statistics of the dataset: number of points, numer of features, number of classes, data-points per class.
- Explain our objective.
- Perform Univaraite analysis(PDF, CDF, Boxplot, Voilin plots) to understand which features are useful towards classification.
- Perform Bi-variate analysis (scatter plots, pair-plots) to see if combinations of features are useful in classfication.

 Write your observations in english as crisply and unambigously as possible. Always quantify your results.

```
iris virginica SW = iris virginica.iloc[:,1]
iris_versicolor_SW = iris_versicolor.iloc[:,1]
from scipy import stats
stats.ks_2samp(iris_virginica_SW, iris_versicolor_SW)
x = stats.norm.rvs(loc=0.2, size=10)
stats.kstest(x,'norm')
x = stats.norm.rvs(loc=0.2, size=100)
stats.kstest(x,'norm')
x = stats.norm.rvs(loc=0.2, size=1000)
stats.kstest(x,'norm')
import pandas as pd
import seaborn as sns
import matplotlib.pyplot as plt
import numpy as np
from google.colab import files
uploaded=files.upload()
haberman1 = pd.read csv("haberman1.csv")
     Choose Files haberman1.csv

    haberman1.csv(n/a) - 3432 bytes, last modified: 4/18/2021 - 100% done

     Saving haberman1.csv to haberman1.csv
# (Q) how many data-points and features?
print (haberman1.shape)
     (306, 4)
#(Q) What are the column names in our dataset?
print (haberman1.columns)
```

```
Index(['age', 'year', 'nodes', 'status'], dtype='object')
```

#(Q) How many data points for each class are present?

```
haberman1["status"].value_counts()
# balanced-dataset vs imbalanced datasets
#haberman1 is a balanced dataset as the number of data points for every class is 50.
```

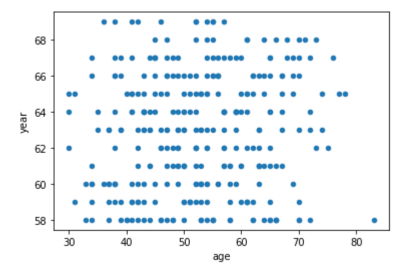
225
 81

Name: status, dtype: int64

(3.2) 2-D Scatter Plot

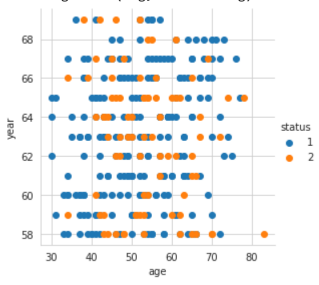
```
#2-D scatter plot:
#ALWAYS understand the axis: labels and scale.
haberman1.plot(kind='scatter', x='age', y='year');
plt.show()
```

#cannot make much sense out it.



- # DUL TEU and green data points cannot be easily seperated.
- # Can we draw multiple 2-D scatter plots for each combination of features?
- # How many cobinations exist? 4C2 = 6.

/usr/local/lib/python3.7/dist-packages/seaborn/axisgrid.py:316: UserWarning: The `size` warnings.warn(msg, UserWarning)



Observation(s):

Using sepal_length and sepal_width features, we can distinguish Setosa flowers from others. Seperating Versicolor from Viginica is much harder as they have considerable overlap.

3-D SCATTER PLOT

https://plot.ly/pandas/3d-scatter-plots/

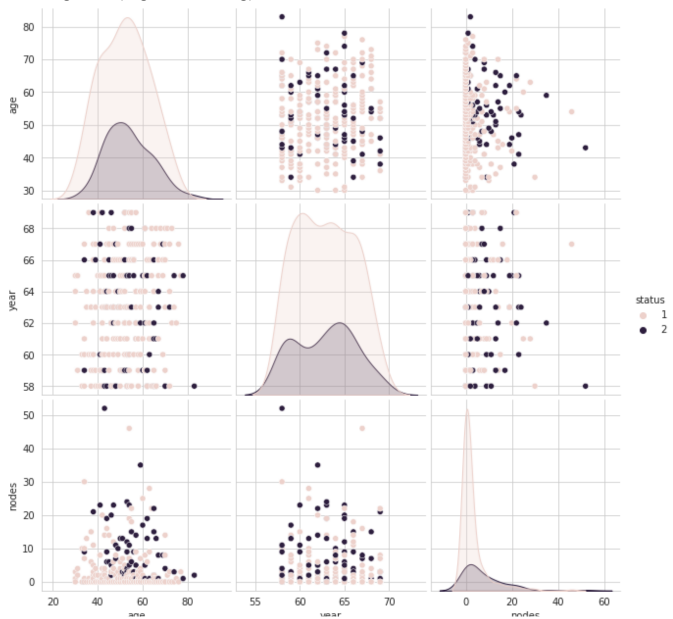
Needs a lot to mouse interaction to interpret data.

What about 4-D, 5-D or n-D scatter plot?

3D PAIR PLOT

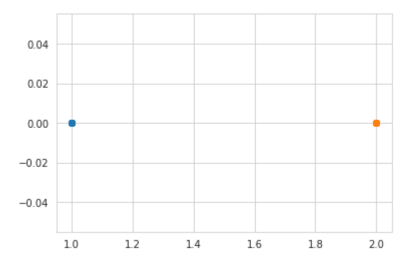
```
# pairwise scatter plot: Pair-Plot
# Dis-advantages:
##Can be used when number of features are high.
##Cannot visualize higher dimensional patterns in 3-D and 4-D.
#Only possible to view 2D patterns.
plt.close();
sns.set_style("whitegrid");
sns.pairplot(haberman1, hue="status", size=3);
plt.show()
# NOTE: the diagnol elements are PDFs for each feature. PDFs are expalined below.
```

/usr/local/lib/python3.7/dist-packages/seaborn/axisgrid.py:1969: UserWarning: The `size` warnings.warn(msg, UserWarning)



(3.4) Histogram, PDF, CDF

```
# What about 1-D scatter plot using just one feature?
#1-D scatter plot of petal-length
import numpy as np
haberman1_1 = haberman1.loc[haberman1["status"] == 1];
haberman1_2 = haberman1.loc[haberman1["status"] == 2];
#print(haberman1_1["status"])
plt.plot(haberman1_1["status"], np.zeros_like(haberman1_1['status']), 'o')
plt.plot(haberman1_2["status"], np.zeros_like(haberman1_2['status']), 'o')
plt.show()
#Disadvantages of 1-D scatter plot: Very hard to make sense as points
#are overlapping a lot.
#Are there better ways of visualizing 1-D scatter plots?
```

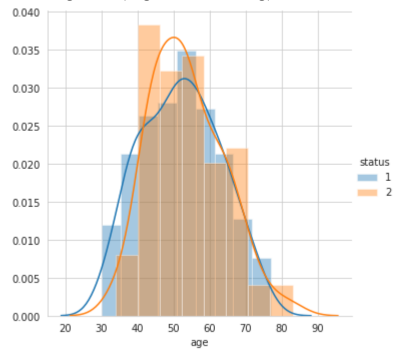


```
sns.FacetGrid(haberman1, hue="status", size=5) \
    .map(sns.distplot, "age") \
    .add_legend();
plt.show();
```

/usr/local/lib/python3.7/dist-packages/seaborn/axisgrid.py:316: UserWarning: The `size` warnings.warn(msg, UserWarning)

/usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `di warnings.warn(msg, FutureWarning)

/usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `diwarnings.warn(msg, FutureWarning)

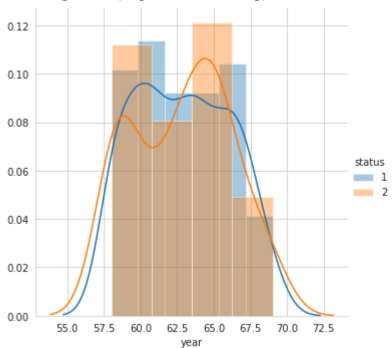


```
sns.FacetGrid(haberman1, hue="status", size=5) \
    .map(sns.distplot, "year") \
    .add_legend();
plt.show();
```

```
/usr/local/lib/python3.7/dist-packages/seaborn/axisgrid.py:316: UserWarning: The `size` warnings.warn(msg, UserWarning)
```

/usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `di warnings.warn(msg, FutureWarning)

/usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `di warnings.warn(msg, FutureWarning)



```
sns.FacetGrid(haberman1, hue="status", size=5) \
    .map(sns.distplot, "nodes") \
    .add_legend();
plt.show();
```

```
/usr/local/lib/python3.7/dist-packages/seaborn/axisgrid.py:316: UserWarning: The `size`
       warnings.warn(msg, UserWarning)
     /usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `di
       warnings.warn(msg, FutureWarning)
     /usr/local/lib/python3.7/dist-packages/seaborn/distributions.py:2557: FutureWarning: `di
       warnings.warn(msg, FutureWarning)
# Histograms and Probability Density Functions (PDF) using KDE
# How to compute PDFs using counts/frequencies of data points in each window.
# How window width effects the PDF plot.
# Interpreting a PDF:
## why is it called a density plot?
## Why is it called a probability plot?
## for each value of petal length, what does the value on y-axis mean?
# Notice that we can write a simple if..else condition as if(petal length) < 2.5 then flower
# Using just one feature, we can build a simple "model" suing if..else... statements.
# Disadv of PDF: Can we say what percentage of versicolor points have a petal length of less
# Do some of these plots look like a bell-curve you studied in under-grad?
# Gaussian/Normal distribution.
# What is "normal" about normal distribution?
# e.g: Hieghts of male students in a class.
# One of the most frequent distributions in nature.
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 5?
# How to construct a CDF?
# How to read a CDF?
import numpy as np
#Plot CDF of petal_length
counts, bin edges = np.histogram(haberman1['year'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges);
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf);
plt.plot(bin edges[1:], cdf)
counts, bin edges = np.histogram(haberman1['year'], bins=20,
                                 density = True)
pdf = counts/(sum(counts))
plt.plot(bin_edges[1:],pdf);
```

```
prt.snow();
```

```
[0.20588235 0.09150327 0.08496732 0.0751634 0.09803922 0.10130719 0.09150327 0.09150327 0.08169935 0.07843137] [58. 59.1 60.2 61.3 62.4 63.5 64.6 65.7 66.8 67.9 69. ]
```

```
1.0 -

0.8 -

0.6 -

0.4 -

0.2 -

0.0 -

60 62 64 66 68
```

```
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 5?
# How to construct a CDF?
# How to read a CDF?
import numpy as np
#Plot CDF of petal length
counts, bin edges = np.histogram(haberman1['nodes'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges);
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf);
plt.plot(bin edges[1:], cdf)
counts, bin edges = np.histogram(haberman1['nodes'], bins=20,
                                 density = True)
pdf = counts/(sum(counts))
plt.plot(bin_edges[1:],pdf);
plt.show();
```

```
[0.77124183 0.09803922 0.05882353 0.02614379 0.02941176 0.00653595
      0.00326797 0.
                             0.00326797 0.00326797]
            5.2 10.4 15.6 20.8 26. 31.2 36.4 41.6 46.8 52.
     [ 0.
      1.0
      0.8
      0.6
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 5?
# How to construct a CDF?
# How to read a CDF?
import numpy as np
#Plot CDF of petal_length
counts, bin edges = np.histogram(haberman1['age'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges);
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf);
plt.plot(bin edges[1:], cdf)
counts, bin edges = np.histogram(haberman1['age'], bins=20,
                                  density = True)
pdf = counts/(sum(counts))
plt.plot(bin_edges[1:],pdf);
plt.show();
     [0.05228758 0.08823529 0.1503268 0.17320261 0.17973856 0.13398693
      0.13398693 0.05882353 0.02287582 0.00653595]
     [30.
           35.3 40.6 45.9 51.2 56.5 61.8 67.1 72.4 77.7 83.
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
                        50
                                 60
                                         70
                40
                                                 80
```

```
# Need for Cumulative Distribution Function (CDF)
# We can visually see what percentage of versicolor flowers have a
# petal length of less than 5?
# How to construct a CDF?
# How to read a CDF?
import numpy as np
#Plot CDF of petal_length
counts, bin edges = np.histogram(haberman1['status'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges);
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf);
plt.plot(bin edges[1:], cdf)
counts, bin edges = np.histogram(haberman1['status'], bins=20,
                                  density = True)
pdf = counts/(sum(counts))
plt.plot(bin_edges[1:],pdf);
plt.show();
     [0.73529412 0.
                                                                0.
                             0.
      0.
                             0.
                                         0.264705881
                 0.
     [1.
          1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2.
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
                1.2
                         1.4
                                  1.6
                                           1.8
                                                    2.0
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of nodes
counts, bin edges = np.histogram(haberman1['nodes'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
```

```
print(bin_edges)
#compute CDF
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
     [0.77124183 0.09803922 0.05882353 0.02614379 0.02941176 0.00653595
      0.00326797 0.
                             0.00326797 0.00326797]
     [ 0.
            5.2 10.4 15.6 20.8 26. 31.2 36.4 41.6 46.8 52. ]
     [<matplotlib.lines.Line2D at 0x7f8cae22ea90>]
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
                       20
                                30
               10
                                         40
                                                  50
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of status
counts, bin edges = np.histogram(haberman1['status'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
#compute CDF
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
```

```
[0.73529412 0.
                             0.
                                                    0.
                                                               0.
      0.
                             0.
                                        0.264705881
                 0.
     [1. 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2. ]
     [<matplotlib.lines.Line2D at 0x7f8cae109090>]
      1.0
      0.8
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of age
counts, bin_edges = np.histogram(haberman1['age'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
#compute CDF
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
     [0.05228758 0.08823529 0.1503268 0.17320261 0.17973856 0.13398693
      0.13398693 0.05882353 0.02287582 0.00653595]
     [30. 35.3 40.6 45.9 51.2 56.5 61.8 67.1 72.4 77.7 83. ]
     [<matplotlib.lines.Line2D at 0x7f8cae089950>]
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
                                60
                                        70
                                                 80
              40
                       50
# How to construct a CDF?
# How to read a CDF?
#Plot CDF of year
counts, bin edges = np.histogram(haberman1['year'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
```

```
print(pat);
print(bin_edges)
#compute CDF
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
     [0.20588235 0.09150327 0.08496732 0.0751634 0.09803922 0.10130719
      0.09150327 0.09150327 0.08169935 0.07843137]
           59.1 60.2 61.3 62.4 63.5 64.6 65.7 66.8 67.9 69.
     [<matplotlib.lines.Line2D at 0x7f8cadff8150>]
      1.0
      0.8
      0.6
      0.4
      0.2
                      62
                               64
                                       66
              60
                                               68
# Plots of CDF of petal length for various ages of cancer patients.
# Misclassification error if you use status only.
counts, bin_edges = np.histogram(haberman1['status'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges)
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
counts, bin edges = np.histogram(haberman1['status'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
```

cdf = np.cumsum(pdf)

plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)

```
plt.show();
     [0.73529412 0.
                                                                 0.
                              0.
                                          0.
                                                      0.
      0.
                              0.
                                          0.264705881
                  0.
          1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2. ]
     [0.73529412 0.
                              0.
                                                                 0.
                                          0.
      0.
                  0.
                              0.
                                          0.26470588]
     [1.
          1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2. ]
      1.0
      0.8
      0.6
      0.4
      0.2
      0.0
                        1.4
                                  1.6
                                            1.8
               1.2
                                                     2.0
# Plots of CDF of petal length for various ages of cancer patients.
# Misclassification error if you use age only.
counts, bin_edges = np.histogram(haberman1['age'], bins=10,
                                   density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
# virginica
```

counts, bin edges = np.histogram(haberman1['status'], bins=10,

pdf = counts/(sum(counts))

plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)

print(pdf);

print(bin_edges)
cdf = np.cumsum(pdf)

density = True)

```
[0.05228758 0.08823529 0.1503268 0.17320261 0.17973856 0.13398693
      0.13398693 0.05882353 0.02287582 0.00653595]
     [30. 35.3 40.6 45.9 51.2 56.5 61.8 67.1 72.4 77.7 83. ]
     [0.73529412 0.
                             0.
                                        0.
                                                               0.
      0.
                 0.
                             0.
                                        0.264705881
     [1.
          1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2. ]
     [<matplotlib.lines.Line2D at 0x7f8cadf71b50>]
      1.0
      0.8
      0.6
      0.4
# Plots of CDF of petal length for various ages of cancer patients.
# Misclassification error if you use status only.
counts, bin_edges = np.histogram(haberman1['nodes'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
counts, bin_edges = np.histogram(haberman1['status'], bins=10,
                                  density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges)
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
```

```
[0.77124183 0.09803922 0.05882353 0.02614379 0.02941176 0.00653595
      0.00326797 0.
                            0.00326797 0.003267971
            5.2 10.4 15.6 20.8 26. 31.2 36.4 41.6 46.8 52.
     [0.73529412 0.
                            0.
                                       0.
                                                              0.
      0.
                 0.
                            0.
                                        0.264705881
     [1. 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2. ]
     [<matplotlib.lines.Line2D at 0x7f8cadf11390>]
      1.0
# Plots of CDF of petal length for various ages of cancer patients.
# Misclassification error if you use status only.
counts, bin edges = np.histogram(haberman1['year'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin_edges)
cdf = np.cumsum(pdf)
plt.plot(bin edges[1:],pdf)
plt.plot(bin_edges[1:], cdf)
counts, bin_edges = np.histogram(haberman1['status'], bins=10,
                                 density = True)
pdf = counts/(sum(counts))
print(pdf);
print(bin edges)
cdf = np.cumsum(pdf)
plt.plot(bin_edges[1:],pdf)
plt.plot(bin edges[1:], cdf)
```

[0.20588235 0.09150327 0.08496732 0.0751634 0.09803922 0.10130719

```
(3.5) Mean, Variance and Std-dev
```

```
A 26/7AE001
#Mean, Variance, Std-deviation,
print("Means:")
print(np.mean(haberman1["status"]))
#Mean with an outlier.
print(np.mean(np.append(haberman1["status"],50)));
print(np.mean(haberman1["status"]))
print("\nStd-dev:");
print(np.std(haberman1["status"]))
print(np.std(haberman1["age"]))
     Means:
     1,2647058823529411
     1.4234527687296417
     1.2647058823529411
     Std-dev:
     0.4411764705882353
     10.78578520363183
```

(3.6) Median, Percentile, Quantile, IQR, MAD

```
#Median, Quantiles, Percentiles, IQR.
print("\nMedians:")
print(np.median(haberman1["year"]))
#Median with an outlier
print(np.median(np.append(haberman1["year"],50)));
print(np.median(haberman1["year"]))

print("\nQuantiles:")
print(np.percentile(haberman1["year"],np.arange(0, 100, 25)))

print("\n90th Percentiles:")
print(np.percentile(haberman1["year"],90))

from statsmodels import robust
print ("\nMedian Absolute Deviation")
print(robust.mad(haberman1["year"]))
```

Medians:

```
63.0
     63.0
     63.0
     Quantiles:
     [58.
            60.
                  63.
                        65.75]
     90th Percentiles:
     67.0
     Median Absolute Deviation
     4.447806655516806
     /usr/local/lib/python3.7/dist-packages/statsmodels/tools/ testing.py:19: FutureWarning:
       import pandas.util.testing as tm
#Median, Quantiles, Percentiles, IQR.
print("\nMedians:")
print(np.median(haberman1["age"]))
#Median with an outlier
print(np.median(np.append(haberman1["age"],50)));
print(np.median(haberman1["age"]))
print("\nQuantiles:")
print(np.percentile(haberman1["age"],np.arange(0, 100, 25)))
print("\n90th Percentiles:")
print(np.percentile(haberman1["age"],90))
from statsmodels import robust
print ("\nMedian Absolute Deviation")
print(robust.mad(haberman1["age"]))
     Medians:
     52.0
     52.0
     52.0
     Quantiles:
     [30.
            44.
                  52.
                        60.751
     90th Percentiles:
     67.0
     Median Absolute Deviation
     11.860817748044816
#Median, Quantiles, Percentiles, IQR.
print("\nMedians:")
print(np.median(haberman1["nodes"]))
```

```
#Median with an outlier
print(np.median(np.append(haberman1["nodes"],50)));
print(np.median(haberman1["nodes"]))
print("\nQuantiles:")
print(np.percentile(haberman1["nodes"],np.arange(0, 100, 25)))
print("\n90th Percentiles:")
print(np.percentile(haberman1["nodes"],90))
from statsmodels import robust
print ("\nMedian Absolute Deviation")
print(robust.mad(haberman1["nodes"]))
     Medians:
     1.0
     1.0
     1.0
     Ouantiles:
     [0. 0. 1. 4.]
     90th Percentiles:
     13.0
     Median Absolute Deviation
     1.482602218505602
#Median, Quantiles, Percentiles, IQR.
print("\nMedians:")
print(np.median(haberman1["status"]))
#Median with an outlier
print(np.median(np.append(haberman1["status"],50)));
print(np.median(haberman1["status"]))
print("\nQuantiles:")
print(np.percentile(haberman1["status"],np.arange(0, 100, 25)))
print("\n90th Percentiles:")
print(np.percentile(haberman1["status"],90))
from statsmodels import robust
print ("\nMedian Absolute Deviation")
print(robust.mad(haberman1["status"]))
```

Medians:

1.0 1.0

1.0

Quantiles:

[1. 1. 1. 2.]

90th Percentiles:

2.0

Median Absolute Deviation

0.0

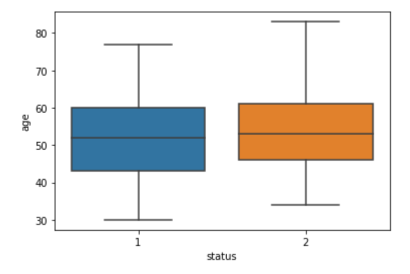
(3.7) Box plot and Whiskers

#Box-plot with whiskers: another method of visualizing the 1-D scatter plot more intuitivey.

- # The Concept of median, percentile, quantile.
- # How to draw the box in the box-plot?
- # How to draw whiskers: [no standard way] Could use min and max or use other complex statisti
- # IQR like idea.

#NOTE: IN the plot below, a technique call inter-quartile range is used in plotting the whisk #Whiskers in the plot below donot correposed to the min and max values.

#Box-plot can be visualized as a PDF on the side-ways.



#Box-plot with whiskers: another method of visualizing the 1-D scatter plot more intuitivey.

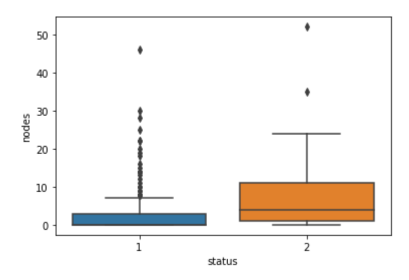
- # The Concept of median, percentile, quantile.
- # How to draw the box in the box-plot?
- # How to draw whiskers: [no standard way] Could use min and max or use other complex statisti
- # IQR like idea.

#NOTE: IN the plot below, a technique call inter-quartile range is used in plotting the whisk https://colab.research.google.com/drive/1hrjzy2UViascSCxzm7e68C8Sze5g8IPO#scrollTo=-n6MmDDTgnxV 34/39

#Whiskers in the plot below donot correposnd to the min and max values.

#Box-plot can be visualized as a PDF on the side-ways.

sns.boxplot(x='status',y='nodes', data=haberman1)
plt.show()



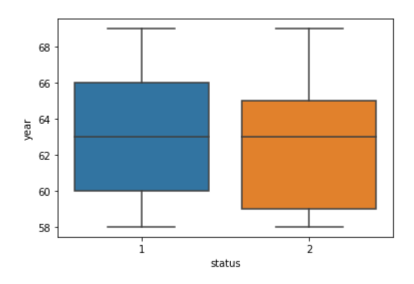
#Box-plot with whiskers: another method of visualizing the 1-D scatter plot more intuitivey.

- # The Concept of median, percentile, quantile.
- # How to draw the box in the box-plot?
- # How to draw whiskers: [no standard way] Could use min and max or use other complex statisti
- # IOR like idea.

#NOTE: IN the plot below, a technique call inter-quartile range is used in plotting the whisk #Whiskers in the plot below do not correposed to the min and max values.

#Box-plot can be visualized as a PDF on the side-ways.

sns.boxplot(x='status',y='year', data=haberman1)
plt.show()

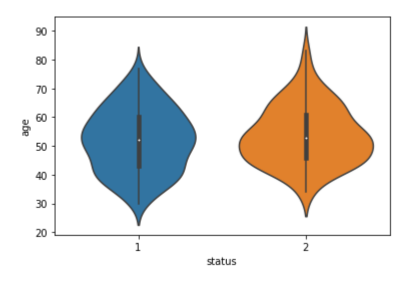


(3.8) Violin plots

A violin plot combines the benefits of the previous two plots # and simplifies them

Denser regions of the data are fatter, and sparser ones thinner #in a violin plot

sns.violinplot(x="status", y="age", data=haberman1, size=8)
plt.show()



A violin plot combines the benefits of the previous two plots #and simplifies them

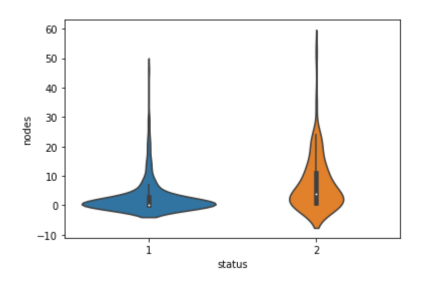
Denser regions of the data are fatter, and sparser ones thinner
#in a violin plot

sns.violinplot(x="status", y="year", data=haberman1, size=8)
plt.show()

A violin plot combines the benefits of the previous two plots # and simplifies them

Denser regions of the data are fatter, and sparser ones thinner
#in a violin plot

sns.violinplot(x="status", y="nodes", data=haberman1, size=8)
plt.show()



(3.10) Univariate, bivariate and multivariate analysis.

Def: Univariate, Bivariate and Multivariate analysis.

File <u>"<ipython-input-46-3bcb4bd9ab1d>"</u>, line 2
Def: Univariate, Bivariate and Multivariate analysis.

SyntaxError: invalid syntax

SEARCH STACK OVERFLOW

(3.11) Multivariate probability density, contour plot.

```
#2D Density plot, contors-plot
sns.jointplot(x="age", y="nodes", data=haberman1, kind="kde");
plt.show();
```

