Summary

Intro

- Get row with index x with df.loc[x]. Get row number x with df.iloc[x].
- Delete rows (axis=0) or columns with df.drop('col_name', axis-1).
- Sort by cols with df.sort_values(ascending=[True,False], by=['col1', 'col2']).
- Access month of datetime column with df.datetime_col.dt.month.
- Options of merging: pd.merge(vessels, segments, left_index=False, right_on='mmsi', left_on='left_col')
- Concatenate rows of a dataframe pd.concat([mb1, mb2], axis=0)
- Stacking dataframes; create one index col where the original column identifier of the data point is. df.stack(). The inverse is df.unstack()s
- Transform a table of with multiple observations to a new table showing, e.g. observations of some kind for a patient: df.pivot(index='', columns='', values='')
 - Similarly, with pivot_table we can use and aggregation function to create the table.
- With pd.crosstab(df.col1, df.col2) we can aggregate the value counts of the selected columns.
- Convert categorical data to dummy / indicator variables: pd.get_dummies(df.col).
 - this also works for multiple columns at the same time

```
categorical_cols = ['animal_type', 'intake_condition', 'intake_type', 'sex_upon_intake'
```

num_rep = pd.get_dummies(original_data, columns=categorical_cols).drop(['outcome_type']
• Bucketizing data: pd.cut(df.age, [20,40,60,80], labels=['young',

- ... ,'old'])
 - pd.qcut divides the data by quanitles
- select rows from indices with df.take(indices)

Plotting

• Always add title and ax-lables with

```
plt.xlabel('bla')
plt.ylablel('bla')
plt.title('bla')
```

- Visualize two varibles: sns.jointplot(x=df['x'], y=df['y'], kind='hex')
 - kind='reg' does a scatter plot with indicated regression direction and errorbar.
 - a similar result to kind='reg' can be obtained with sns.lmplot(x='col1',y='col2',
 data=df, hue = 'hue_col')

```
    Barplots in seaborn provide errorbars by default sns.barplot(x='labels', y='values', data=df)
```

```
• Manual errorbars with plt.errorbar(x, y, yerr=None)
```

```
- filled error area: plt.fill_between(x, y1, y2=0)
```

• Populate 4x4 subplot with histograms

```
fig, ax = plt.subplots(4,4,figsize= (8,6), sharey = True, sharex = True)
for i in range(16):
    sbplt = ax[i%4, math.floor(i/4)]
    sbplt.hist(stats_by_genre.iloc[i].values,range = [0,200],bins = 20)
    sbplt.set_title(stats_by_genre.index[i])
fig.tight_layout()
fig.text(0.4,0, "x label here")
fig.text(0,0.6, "y label here", rotation = 90)
  • another way more for a quick subplot is
fig, axs = plt.subplots(1, 3, sharey=True)
data.plot(kind='scatter', x='TV', y='sales', ax=axs[0], figsize=(16, 5), grid=True)
data.plot(kind='scatter', x='radio', y='sales', ax=axs[1], grid=True)
data.plot(kind='scatter', x='newspaper', y='sales', ax=axs[2], grid=True)
  • heatmaps with seaborn
df2 = pd.crosstab(df['col1'],df['col2'])
sns.heatmap(df2, annot=True)
  • draw a loglog pdf (eg for a power law)
array = plt.hist(df.col,bins=1000,log=True, density=True)
plt.close()
plt.loglog(array[1][1:],array[0])
```

Describing data

• Test if data is exponential ('exp') or normal ('norm') distributed statsmodels.stats.diagnostic.kstest_normal(df['col'].values, dist='norm', pvalmethod='table')

• sns.pairplot(df) is useful to describe the df as df.describe()

- Personr measures linear correlation stats.pearsonr(df['col1'],df['col2']).
 Spearmanr measures monotonic correlation, i.e. compares the rankings of the two datastreams stats.spearmanr(df['col1'],df['col2']).
- Compare if two datastreams have the same mean with stats.ttest_ind(df['col1'], df['col2']).

• Get p-value from a one sided binominal test scipy.stats.binom_test(x, n=None, p=0.5, alternative='greater') where x is the number of successful outcomes, n the number of outcomes and p the probability of success.

Regression analysis

• least square model and summary printing:

```
mod = smf.ols(formula='outcome ~ C(catogory)+ continuous, data=df)
res = mod.fit()
print(res.summary())
```

- options
 - a:b in the formula is an interaction term
 - a*b is shorthand for a+b+a:b
 - logistic regression with smf.logit(...)
- postprocessing pipeline

• A log transform on the outcome makes the model multiplicative. Don't forget to exponentiate the coefficient again befor reasoning about the result.

Observational studies

• Default: do a propensity score matching, i.e. do the analysis on paires matched on the probability of treatment. A logistic regression can predict the propensity score for all samples.

```
mod = smf.logit(formula='treatment ~ age + educ + C(black) , data=df)
res = mod.fit()

# Extract the estimated propensity scores
df['Propensity_score'] = res.predict()
```

• Networkx can calculate a maximal matching. Hence, we have to converte the propensity scores to similarity rating $= 1 - |p_1 - p_2|$. The latter is

the edgeweigth and we add an edge between between all nodes of different groups (bipartite graph).

Supervised learning

- Use entire dataset and cross validation in one step y_pred = cross_val_predict(model, X, y, cv=5)
- Impute missing values with the mean X.fillna(X.mean()).
- Use cross validation to predict precision or recall recall = cross_val_score(model, X, y, cv=10, scoring="recall")
- ROC-AUC with cross-validation

```
y_pred = cross_val_predict(logistic, X, y, cv=10, method="predict_proba")
fpr, tpr, _ = roc_curve(y, y_pred[:, 1])
auc_score = auc(fpr, tpr)
```

• an example to do cross-validation and prediction in one go

```
ridge = Ridge()
ridge_hyper = {'alpha':(0.001, 0.01, 0.1)}
ridge_cv = GridSearchCV(ridge, ridge_hyper, cv=3)
ridge_cv.fit(X_train, y_train)
```

Unsupervised learning

- show kmeans results in a scatterplot ax.scatter(X[:,0], X[:,1], c=kmean.labels_, alpha=0.6)
 - the alpha parameter is the blending value
- Criterions to find k:
 - silhouette score=(b-a)/max(a,b) where b is avg. inter cluster distance and a the avg. intra cluster distance. silhouette_score(X, labels)
 - elbow method: look at the sum of all distances to the closest cluster center. Select k at the elbow. kmeans.inertia_
- Dimensionality reduction
 - t-stochastic neighbor embedding(t-SNE): gives each datapoint a location in a lower dimensional map with high probability.

 X_reduced_tsne = TSNE(n_components=2, init='random', learning_rate='auto', random_state=42).fit_transform(X)
- PCA: X_reduced_pca = PCA(n_components=2).fit(X).transform(X)
- Remove the mean and scale to unit variance with preprocessing.StandardScaler().fit_transform(X)

Handling text

- new lines can be removed by applying " ".join(txt.split())
- get a spacy object (tokens) from text doc = nlp(txt)
- get sentences as a list doc.sents

- get wordtype of a token [(token.text, token.pos_) for token in doc]
 - or more detailed with token.tag_ instead of token.pos_
- a list of stopwords is available in spacy spacy.lang.en.stop_words.STOP_WORDS
- token.lemma_ is the lemmatized version of the token
- chuncks consist of a noun and the words describing the noun: doc.noun_chunks
- There is a sentiment analyzer for short sentences.

```
analyzer = SentimentIntensityAnalyzer()
vs = analyzer.polarity_scores(example)
print(example, '\n')
print('Negative sentiment:',vs['neg'])
print('Neutral sentiment:',vs['neu'])
print('Positive sentiment:',vs['pos'])
print('Compound sentiment:',vs['compound'])
```

- interpretation
 - positive, negative and neutral sum to one. The compound represents all three values.
- The count vectorizer transforms a collection of text into a matrix of token counts.

```
vectorizer = CountVectorizer()
X = vectorizer.fit_transform(chunks)
```

- The same exists for getting the TF-IDF matrix TfidfVectorizer()
- Print the top 20 feature representing a category:

```
coefs=clf.coef_[0]
top = np.argpartition(coefs, -20)[-20:]
print(np.array(vectorizer.get_feature_names())[top])
```

- we can also represent a sentense as vector using pretrained GloVe nlp(example).vector
- topic detection with LDA: LdaMulticore(corpus=corpus, num_topics=4, id2word=dictionary, workers=6,passes=params['passes'], random_state=params['random_state']) then show the topics with model.show_topics(num_words=5).
- A dictionary with typical words per category is provided by

```
from empath import Empath
lexicon = Empath()
```

• There is a command to analyze a text empath_features = lexicon.analyze(doc.text,categories = ["disappointment", "pain", "joy", "beauty", "affection"])

Networks

```
• nx.Graph() per default, nx.DiGraph() for graphs with self-loops,
     nx.MultiDiGraph() for graphs with parallel edges.
  • Add nodes G.add_node(id), add edges G.add_edge(u,v)
       - can add multiple at once with G.add_nodes_from(range(2,9)) and
         G.add_edges_from([(1,2),(2,3)])
  • Print number of nodes and edges with nx.info()
  • plot graph with nx.draw_spring(G, with_labels=True,
     0.8)

    or with a circular layout nx.draw_circular(karateG, with_labels=True,

         node_color='g', alpha = 0.8)
  • We can create a graph directly from a pandas edgelist with G
     =nx.from_pandas_edgelist(edges, 'Source', 'Target', edge_attr=None,
     create_using= nx.Graph()).
  • Node attributes are added
                                           nx.set_node_attributes(G,
     nodes['Role'].to_dict(), 'Role' )
  • get the sparsity of a network with nx.density(G)
  • connected components of a graph nx.connected_components(G)
       - the boolean query is nx.is connected(G)
  • the shortest path between a pair of nodes nx.shortest_path(quakerG,
     source="src", target="dst")
  • node induced subraph subgraph = G.subgraph(node_set)
  • diameter nx.diameter(G)
  • average shortest paht nx.average_shortest_path_length(G)
  • global clustering coefficient nx.transitivity(G)
  • local clustering coefficinet nx.clustering(G, ['node', ...])
  • get all degrees in a graph degrees = dict(G.degree(G.nodes()))
  • katz centrality katz = nx.katz_centrality(G)
  • betweenness centrality betweenness = nx.betweenness_centrality(G)
       - choose node colors according to the betweenness centrality:
     nx.set_node_attributes(G, nx.betweenness_centrality(G), 'betweenness')
     pos = nx.spring_layout(G)
     ec = nx.draw_networkx(G, pos, nodelist=G.nodes(),
                              node_color=[G.nodes[n]["betweenness"] for n in G.nodes()],
                              node_shape = '.', node_size=1200, font_color="white", font_weig
     plt.colorbar(nc)
    plt.axis('off')
     plt.show()
  • The Girvan Newman algorithm is implemented as in iterator in nx
comp = girvan_newman(G)
it = 0
for communities in itertools.islice(comp, 4):
    print('Iteration', it)
```

print(tuple(sorted(c) for c in communities))

- The partition with the highest modularity can be optained directly with the Louvain algorithm with partition = community_louvain.best_partition(quakerG).
- We can measure the similarity of connections in a graph with respect to a given attribute trough the attribute assortativity coefficient. nx.attribute_assortativity_coefficient(G, 'Gender')