Predicting Chub Density based on river environment

github: https://github.com/bendavidhayes/ML_Final_Project.git

Project Topic

This Notebook analyzes how the environment impacts fish life, specifically a fish called chub.

Goal: The objective of the analysis is to develop a model to accurately predict the amount of fish in a river section, based on environmental factors.

Data Description

The Data used in this analysis was provided by the University of Natural Resources and Life Sciences in Vienna, Austria.

It was collected in a variety of rivers in Austria. For each observation a section of a river was contained using nets. All fish in the containment were caught, weighed and then released. The target in this analysis is the biomass of all Chub caught in a section, measured in kg/ha.

target:

• Chub_BIO: Chub Biomass (kg/ha)

features:

- Fischregion
- CL1_Artificial surfaces: Proportion of Area surrounding the catchment area that is artificial, e.g. Buildings, streets.
- SSI: Standard Sinuousity Index (higher values reflect more curves in the river)
- BW_MORPH: Riverbank width morphology index
- PHB_SAPRO: phytobenthic condition
- MZB_SAPRO: saprobic condition
- TROPH_ZUST: trophic condition
- ZUST_SCHAD = ZUST_SCHADST_EU_2015: pollutant grading according to EU standard
- ZUST_BIO_7 = ZUST_BIOLOGIE_STOFFE_2015: grading of biological condition
- LAUFENTWICKLUNG: development of river run
- SUBSTRATZUSAMMENSETZUNG: composition of river substrate
- FIA: Fish Index Austria
- Chub_corridor_m: length of river system, that is accessible to fish
- 5J_AMP_50_REL_MAXNAT: hydrological variable
- 5J_MAFR_CNT: hydrological variable

Import Libraries & Data

```
In [1]: #import libraries
        import pandas as pd
        import matplotlib.pyplot as plt
        import numpy as np
        import math
        import scipy as sp
        import scipy.stats as stats
        import seaborn as sns
In [2]: #read csv
        csv_file_path = 'C:/Users/benda/cyprinid_2023.csv'
        biomass = pd.read_csv(csv_file_path, delimiter=';', decimal=',')
        # labels of features and targets to include in analysis
        columns to keep = ["Chub BIO",
                           "Fischregion",
                           "CL1_Artificial surfaces",
                           "SSI",
                           "BW MORPH",
                           "PHB SAPRO",
                           "MZB_SAPRO"
                           "TROPH_ZUST",
                           "ZUST_SCHAD = ZUST_SCHADST_EU_2015",
                           "ZUST BIO 7 = ZUST BIOLOGIE STOFFE 2015",
                           "LAUFENTWICKLUNG",
                           "SUBSTRATZUSAMMENSETZUNG",
                           "FIA",
                           "Chub_corridor_m",
                           "5J_AMP_50_REL_MAXNAT",
                           "5J MAFR CNT",
        biomass = biomass[columns to keep] # select features for analysis
        # transform FIA into float64
        biomass['FIA'] = biomass['FIA'].str.replace(',', '.', regex=True).astype(float)
        # collaps variables in feature "Fischregion"
        "Epipotamal mittel 1", "Epipotamal mittel 2"]
        replacement = ["Hyporhithral", "Hyporhithral",
                       "Epipomatal", "Epipomatal", "Epipomatal",
"Epipomatal", "Epipomatal"]
        biomass["Fischregion"].replace(to replace, replacement, inplace=True)
```

EDA & Data Cleaning

Since the EDA informs the Data Cleaning Steps needed I decided to combine the two. This helps avoids showing the same visuals for different purposes twice.

```
In [3]: biomass.info() # show basic info about data
```

<class 'pandas.core.frame.DataFrame'>
RangeIndex: 115 entries, 0 to 114
Data columns (total 16 columns):
 # Column

#	Column	Non-Null Count	Dtype
0	Chub_BIO	115 non-null	float64
1	Fischregion	115 non-null	object
2	CL1_Artificial surfaces	115 non-null	float64
3	SSI	115 non-null	float64
4	BW_MORPH	53 non-null	float64
5	PHB_SAPRO	115 non-null	object
6	MZB_SAPRO	115 non-null	float64
7	TROPH_ZUST	115 non-null	object
8	ZUST_SCHAD = ZUST_SCHADST_EU_2015	115 non-null	int64
9	<pre>ZUST_BIO_7 = ZUST_BIOLOGIE_STOFFE_2015</pre>	115 non-null	int64
10	LAUFENTWICKLUNG	91 non-null	float64
11	SUBSTRATZUSAMMENSETZUNG	87 non-null	float64
12	FIA	115 non-null	float64
13	Chub_corridor_m	114 non-null	float64
14	5J_AMP_50_REL_MAXNAT	115 non-null	float64
15	5J_MAFR_CNT	115 non-null	int64
1.6	(1 164/40) 1 164/2) 1 1 1/2)		

dtypes: float64(10), int64(3), object(3)

memory usage: 14.5+ KB

The data set has 115 observations and 16 features (including the target). There are 10 quantitative variables, 3 ordinal, and 3 qualitative. Four features have missing values.

Summary Statistics

In [4]: pd.set_option('display.max_columns', 150) # increase max columns shown in output
biomass.describe() # summary statistics of quantitative variables

Out[4]:		Chub_BIO	CL1_Artificial surfaces	SSI	BW_MORPH	MZB_SAPRO	ZUST_SCHAD = ZUST_SCHADST_EU_2015
	count	115.000000	115.000000	115.000000	53.000000	115.000000	115.000000
	mean	36.168235	0.062406	1.077643	0.142628	1.782609	1.017391
	std	56.564671	0.015785	0.113820	0.135535	0.146306	0.186501
	min	0.000000	0.033827	1.000000	0.019885	1.250000	1.000000
	25%	0.000000	0.051312	1.008264	0.064863	1.750000	1.000000
	50%	7.768000	0.063618	1.041361	0.099541	1.750000	1.000000
	75%	45.880000	0.071134	1.108768	0.168475	1.750000	1.000000
	max	326.084000	0.087296	1.913400	0.879331	2.000000	3.000000

In [5]: biomass.head()

Out[5]:		Chub_BIO	Fischregion	CL1_Artificial surfaces	SSI	BW_MORPH	PHB_SAPRO	MZB_SAPRO	TROPI
	0	0.00	Hyporhithral	0.070794	1.007143	0.062240	I-II B	1.50	
	1	0.00	Hyporhithral	0.051312	1.041361	0.168475	I-II B	1.75	
	2	0.00	Hyporhithral	0.040292	1.121244	0.080595	I-II B	1.50	
	3	0.00	Hyporhithral	0.040292	1.042523	0.063849	I-II B	1.75	
	4	0.01	Hyporhithral	0.040292	1.129526	0.096156	I-II B	1.75	
•									•

Correlation

In [6]: biomass.corr()

C:\Users\benda\AppData\Local\Temp\ipykernel_2312\2084939813.py:1: FutureWarning: T he default value of numeric_only in DataFrame.corr is deprecated. In a future vers ion, it will default to False. Select only valid columns or specify the value of numeric_only to silence this warning.

biomass.corr()

Out[6]:

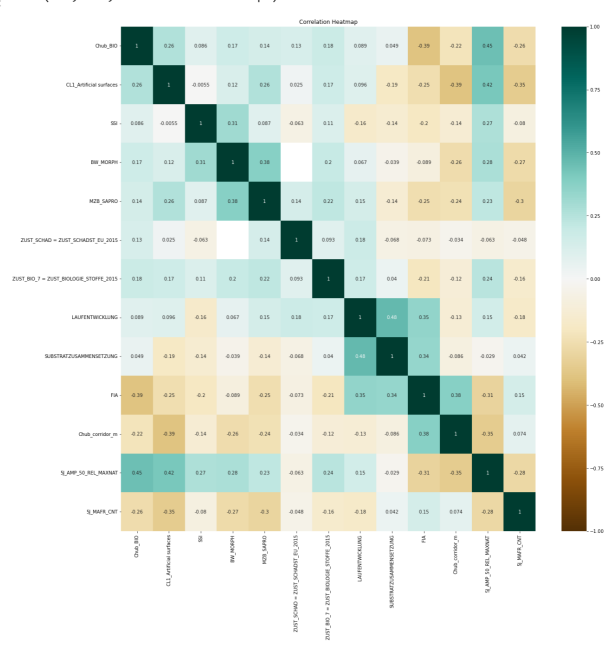
	Chub_BIO	CL1_Artificial surfaces	SSI	BW_MORPH	MZB_SAPRO	ZU
Chub_BIO	1.000000	0.261430	0.085997	0.165504	0.143203	
CL1_Artificial surfaces	0.261430	1.000000	-0.005526	0.122175	0.259991	
SSI	0.085997	-0.005526	1.000000	0.308774	0.086686	
BW_MORPH	0.165504	0.122175	0.308774	1.000000	0.379584	
MZB_SAPRO	0.143203	0.259991	0.086686	0.379584	1.000000	
ZUST_SCHAD = ZUST_SCHADST_EU_2015	0.131053	0.024699	-0.063399	NaN	0.139773	
ZUST_BIO_7 = ZUST_BIOLOGIE_STOFFE_2015	0.182150	0.166062	0.112231	0.200858	0.221852	
LAUFENTWICKLUNG	0.088531	0.095971	-0.164914	0.067082	0.145032	
SUBSTRATZUSAMMENSETZUNG	0.048613	-0.189610	-0.141465	-0.039393	-0.140083	
FIA	-0.386005	-0.251162	-0.195856	-0.089231	-0.251307	
Chub_corridor_m	-0.220738	-0.387683	-0.136852	-0.260197	-0.238651	
5J_AMP_50_REL_MAXNAT	0.448390	0.421268	0.270942	0.277298	0.225730	
5J_MAFR_CNT	-0.261384	-0.351328	-0.079804	-0.270984	-0.299024	

In [7]: # plot the correlation using a heatmap
plt.figure(figsize=(20, 20))
heatmap = sns.heatmap(biomass.corr(), vmin=-1, vmax=1,cmap='BrBG', annot=True)
heatmap.set_title("Correlation Heatmap")

e default value of numeric only in DataFrame.corr is deprecated. In a future versi on, it will default to False. Select only valid columns or specify the value of nu meric_only to silence this warning.

heatmap = sns.heatmap(biomass.corr(), vmin=-1, vmax=1,cmap='BrBG', annot=True) Text(0.5, 1.0, 'Correlation Heatmap')

Out[7]:



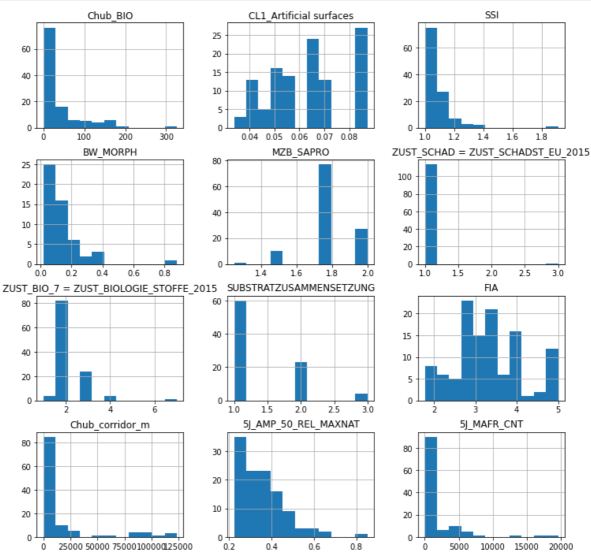
Important Features: Based on the correlation matrix, it looks like the most important features in predicting: 'Chub_BIO' are

- '5J_AMP_50_REL_MAXNAT'(corr=0.45),
- 'FIA'(corr=-0.39),
- 'CL1_Artificial surfaces' (corr=0.26), and
- 5J_MAFR_CNT (corr=-0.26)

Collinearity: The two features "SUBSTRATZUSAMMENSETZUNG" and "LAUFENTWICKLUNG" are very collinear, with a correlation between them of 0.48. Since "LAUFENTWICKLUNG" also has a stronger correlation with the target, I will keep it, and remove the other feature.

Distribution of Data

In [9]: # plot numerical variables as histogram
bm1.hist(bins=11, figsize=(12, 12));



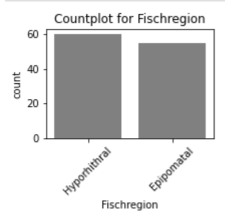
It's clear that "ZUST_SCHAD = ZUST_SCHADST_EU_2015" can be removed. Since almost all observations are identical, it will add no value to the model.

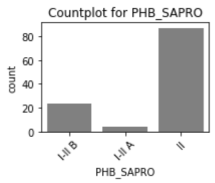
```
In [10]: bm2 = bm1.drop("ZUST_SCHAD = ZUST_SCHADST_EU_2015", axis=1)
```

Note on logarithmic transformation: A lot of the features have a strong right skew. They might need to be logarithmically transformed, if more advanced linear methods are applied. However since not all models need normally distributed data, I will wait until we know which type of model to develop to decide whether to transform them.

Vizualize categorical data

plt.title(f'Countplot for {column}') # Set the title of the plot
plt.xticks(rotation=45) # Rotate x-axis labels for better readability
plt.show()







Impute missing values

In [12]:	<pre>bm2.isna().sum() # check for missing values</pre>				
Out[12]:	Chub_BIO	0			
out[12].	Fischregion	0			
	CL1_Artificial surfaces	0			
	SSI	0			
	BW_MORPH	62			
	PHB_SAPRO	0			
	MZB_SAPRO	0			
	TROPH_ZUST	0			
	<pre>ZUST_BIO_7 = ZUST_BIOLOGIE_STOFFE_2015</pre>	0			
	SUBSTRATZUSAMMENSETZUNG	28			
	FIA	0			
	Chub_corridor_m	1			
	5J_AMP_50_REL_MAXNAT	0			
	5J_MAFR_CNT	0			
	dtype: int64				

Features with missing values:
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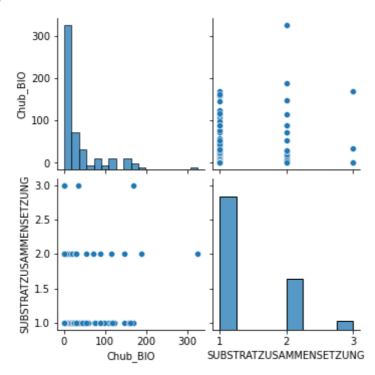
"Chub_corridor_m" is missing a single value. We'll just remove that row.

The other two features with missing values can be removed because:

- "BW_MORPH" has nearly 50% missing
- "SUBSTRATZUSAMMENSETZUNG has 25% missing values and little correlation to the target. Since it is also collinear to another feature, some of the information will be retained in the model anyways. But before removing it, we'll check if there is any nonlinear correlation to the target.

```
In [13]: sns.pairplot(bm2.loc[:,["Chub_BIO", "SUBSTRATZUSAMMENSETZUNG"]])
```

Out[13]: <seaborn.axisgrid.PairGrid at 0x1a1ba117c10>



Since the pairplot also shows little correlation with the target, I will remove the feature "SUBSTRATZUSAMMENSETZUNG".

```
In [14]: # remove the two features
bm3 = bm2.drop(["BW_MORPH", "SUBSTRATZUSAMMENSETZUNG"], axis=1)
# remove the row, where "Chub_corridor_m" has a missing value
bm3 = bm3.loc[biomass.Chub_corridor_m.notna(),:]
```

Recode Categorical Variables

to improve linear regression the categorical variables will be encoded as binary dummy variables.

```
In [15]: # create list of categorical columns
    categorical_data = bm3.select_dtypes(include='object')
    categorical_labels = list(categorical_data.columns)
    print(categorical_labels)

['Fischregion', 'PHB_SAPRO', 'TROPH_ZUST']
```

```
bm4 = pd.get_dummies(bm3)
In [17]:
           # Look at new features
           bm4.head()
Out[17]:
                         CL1_Artificial
                                                                               ZUST_BIO_7 =
              Chub BIO
                                                                                              FIA Chub_corrid
                                             SSI MZB SAPRO
                                                                ZUST_BIOLOGIE_STOFFE_2015
                              surfaces
                    0.00
                              0.070794 1.007143
                                                          1.50
                                                                                              5.0
                                                                                                        9429.46
           1
                    0.00
                              0.051312 1.041361
                                                          1.75
                                                                                           2
                                                                                              5.0
                                                                                                       30268.86
           2
                    0.00
                              0.040292 1.121244
                                                          1.50
                                                                                           2
                                                                                              4.0
                                                                                                       79230.11
           3
                    0.00
                              0.040292 1.042523
                                                          1.75
                                                                                           2
                                                                                              5.0
                                                                                                       79230.11
                    0.01
           4
                              0.040292 1.129526
                                                          1.75
                                                                                              5.0
                                                                                                       79230.11
```

Summary of EDA

Few features seem to be normally distributed; this means that transformations could improve linear models, and even become necessary enable use of more complex linear models like Ridge Regression or Lasso.

3 features were removed:

- 2 because of large amounts of missing data
- 1 because nearly all values were identical

Analysis Strategy: Since the aim is to develop the most accurate prediction, I will compare the estimated test error of multiple base models using cross validation on the training set. This will give an idea on which model to choose for finetuning: I will finetune model with the best score (Mean Squared Error). After training this final model, I will calculate the real test error on the test set to evaluate performance on new data.

Model training

Split Data

```
In [19]: bm5 = bm4.copy()
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```

```
X = bm5.drop("Chub_BIO", axis=1)
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_statest
```

The training set has 91 samples and 18 attributes.

The test set has 23 samples and 18 attributes.

Multilinear Regression

```
In [20]: # estimate the test error of the multilinear model
lm = LinearRegression()
    -cross_val_score(lm, X_train, y_train, scoring='neg_mean_squared_error', cv=10).sum
Out[20]: 2813.5259668306426
```

Decision Tree

```
In [21]: # estimate the test error of a basic Decision Tree Model
    dt = DTR(random_state=42)
        -cross_val_score(dt, X_train, y_train, scoring='neg_mean_squared_error', cv=10).sum
Out[21]: 3318.7657664888893
```

Random Forest

```
In [22]: # estimate the test error of a Random Forest Model
    rf = RFR(random_state=42)
    -cross_val_score(rf, X_train, y_train, scoring='neg_mean_squared_error', cv=10).sum
Out[22]: 2460.1840555830495
```

Gradient Boosting

```
In [23]: # estimate the test error of a Gradient Boosting Model
gb = GBR(random_state=42)
    -cross_val_score(gb, X_train, y_train, scoring='neg_mean_squared_error', cv=10).sum
Out[23]:
2685.8869596836685
```

AdaBoost

```
In [24]: # estimate the test error using an AdaBoost Model
    ada = ADA(random_state=42)
    -cross_val_score(ada, X_train, y_train, scoring='neg_mean_squared_error', cv=10).su

Out[24]:
2940.48751384636
```

Finetuning Best Model

Since the Random Forest Model gives the best estimated test error, I will fine tune this model using Grid Search. Also since we are using Random Forest there is no need to transform the exponentially distributed features and target. There is also no need to

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Results and Analysis

```
In [27]: # best parameters for Random Forest Model
         clf.best estimator
Out[27]:
                                       RandomForestRegressor
         RandomForestRegressor(max_depth=5, max_features='log2', n_estimators=300)
In [28]: print("Mean Squared Error of best Random Forest Estimator is: ", -clf.best_score_)
         Mean Squared Error of best Random Forest Estimator is: 2267.762333010481
In [29]: # calculate test error
         y_pred = clf.predict(X_test)
In [30]: print("Mean Squared Error of tuned Random Forest on test data: ", mean_squared_error
         Mean Squared Error of tuned Random Forest on test data: 2317.8045507310608
In [31]: rf.fit(X_train, y_train)
          y pred = rf.predict(X test)
         print("Mean Squared Error of basic Random Forest on test data: ", mean_squared_error
         Mean Squared Error of basic Random Forest on test data: 3482.92131482647
         Feature Importance Random Forests are not intuitive for Inferrence of feature importance. I
         will normalize the data and run a linear model to find the most important features. I decided
         to normalize rather than standardize the data, since the distribution of most columns is not
         normal.
```

```
In [32]: # Normalize data
bm5_norm = Normalizer().fit_transform(bm5)
X_norm = bm5_norm[:,1:]
y_norm = bm5_norm[:,0]

In [33]: # feature importance
lm = LinearRegression()
lm.fit(X_norm, y_norm)

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coef_labels.sort_values()
```

```
TROPH ZUST om
                                                    -156.231931
Out[33]:
         SSI
                                                    -123.166781
         FIA
                                                     -78.237445
         ZUST BIO 7 = ZUST BIOLOGIE STOFFE 2015
                                                     -31.393109
         PHB_SAPRO_II
                                                     -26.870764
         Chub_corridor_m
                                                      -0.113990
         5J MAFR CNT
                                                      -0.078099
         PHB SAPRO I-II B
                                                      60.832188
         TROPH ZUST me2
                                                      61.251178
         TROPH_ZUST_mt
                                                      75.415045
         TROPH ZUST ot
                                                      84.232325
         Fischregion Hyporhithral
                                                      96.061458
         Fischregion_Epipomatal
                                                     103.723130
         TROPH_ZUST_me1
                                                     135.117972
         5J_AMP_50_REL_MAXNAT
                                                     150.501993
         PHB SAPRO I-II A
                                                     165.823164
         MZB SAPRO
                                                     178.528449
         CL1 Artificial surfaces
                                                     181.111743
         dtype: float64
```

This shows which features have the highest impact on the model: the features that have the largest absolute coefficients:

- CL1_Artificial surfaces
- MZB_SAPRO
- PHB_SAPRO_I-II A
- TROPH_ZUST_om
- 5J_AMP_50_REL_MAXNAT

Only two of these features showed significance when looking at the correlation heat map. This as well as the scatterplots above, suggest that the relationships between the other features and the target are more complex (interactions, polynomials). Using polynomial regression could improve upon the feature analysis.

Discussion

As expected more complex models performed better than linear regression. Hyperparameter tuning gave us the final best parameters for RandomForest:

- max_depth=4
- max_features='sqrt'
- n_estimators=200

This model improved upon training error compared to the basic Random Forest Model by nearly 10%. Although the test error of the final model is 15% higher than the training error, the test error is lower than training errors of all other models.

However this difference between training and test error suggests the model is overfitting. There are 3 ways to improve this:

- 1. Feature Selection
- 2. Test more hyperparameters on RF
- 3. use a simpler model: linear regression. As expected Linear base model might have Loading [MathJax]/jax/output/CommonHTML/fonts/TeX/fontdata.js ta distributions. Transforming, and normalizing

	performance.
In []:	

data would improve performance. Also using polynomial regression could improve