

Assignment 1: Ridge and Lasso Regression (Diabetes Dataset)

ML Course 2025

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In this exercise, you will use the **Diabetes dataset** (built into scikit-learn) to explore how **Ridge** and **Lasso regression** handle correlated predictors and the effects of regularization. You will compare them to a standard **Linear Regression** model, visualize their coefficients, and interpret how regularization influences **model complexity** and **performance**.

You will also examine the role of **interaction features**, exploring how they can add flexibility to the model and help reduce bias.

We will conduct two sets of experiments:

1. Using only the original features, and
2. Using the original features plus interaction terms.

Learning Objectives

By the end of this exercise, you will be able to:

- Apply **Ridge** and **Lasso regression** to control model complexity and prevent overfitting.
- Use **cross-validation** to select the optimal level of regularization (`alpha`).
- Interpret how regularization affects **coefficient magnitude**, **sparsity**, and **model interpretability**.
- Evaluate how adding **interaction features** changes model bias, variance, and predictive performance.

Step 1: Load & Understand the Diabetes Dataset

A crucial point to understand when working with the diabetes dataset from `sklearn.datasets` is that it is **not raw data**. The feature data (`X`) has already been pre-processed according to a specific normalization scheme. This is noted by the original author on the data's source webpage ([source](#)):

> Note that the 10 x variables have been standardized to have mean 0 and squared length = 1 ($\text{sum}(x^2)=1$).

To make this clear for anyone using this notebook, we will load both the **raw data** from the source URL and compare it to the **pre-processed data** that comes with `scikit-learn`.

This allows for a direct comparison and highlights what "pre-processing" has already been done.

👉 The Normalization Equation Used by the Dataset Authors:

The specific transformation applied to each of the 10 feature columns (e.g., AGE, BMI, etc.) is as follows:

$$x_{\text{norm}_{ij}} = \frac{x_{ij} - \bar{x}_j}{\sqrt{\sum_{k=1}^n (x_{kj} - \bar{x}_j)^2}}$$

Where:

- x_{ij} is the original value of feature j for sample i .
- \bar{x}_j is the mean of all original values for feature j .
- n is the total number of samples.
- The summation in the denominator is over all samples k for that feature j .

This process ensures two properties for each resulting feature column:

1. **Mean of 0:** The data is centered.
 2. **Sum of Squares of 1:** The vector for each feature has a unit L2-norm.
-

👉 Data Scaling Equation Used in StandardScaler

The scaling transformation that is used in StandardScaler can be expressed mathematically as:

$$X_{\text{scaled}} = \frac{X_{\text{train}} - \mu_{\text{train}}}{\sigma_{\text{train}}}$$

Where:

- X_{train} is the original feature value from the training data.
- μ_{train} is the mean of the feature values computed from the training data.
- σ_{train} is the standard deviation of the feature values computed from the training data.

For the test data, the scaling transformation is applied as:

$$X_{\text{test, scaled}} = \frac{X_{\text{test}} - \mu_{\text{train}}}{\sigma_{\text{train}}}$$

Where:

- X_{test} is the original feature value from the test data.
- μ_{train} and σ_{train} are the mean and standard deviation computed from the training data, respectively.

Note: It is crucial to use the mean (μ_{train}) and standard deviation (σ_{train}) from the training data for scaling the test data. This ensures that the test data is transformed in the same way as the training data, maintaining consistency and preventing data leakage.

To understand how the Diabetes dataset was preprocessed, we compare the raw data we downloaded with the version included in `sklearn.datasets.load_diabetes`. This comparison shows that scikit-learn's dataset has already been normalized according to the transformation described by equation above.

Let's start by loading the dataset from `sklearn.datasets` and from the URL and reviewing them.

In [176...]

```
import pandas as pd

# The URL of the raw data
url = 'https://www4.stat.ncsu.edu/~boos/var.select/diabetes.tab.txt'

# Use pandas to read the tab-separated file directly from the URL
# sep='\t' tells pandas that the columns are separated by tabs
try:
    df_raw = pd.read_csv(url, sep='\t')

    # Now the data is in a pandas DataFrame called 'df'
    print("Raw data loaded successfully!")
    display(df_raw.head())
    print("")
    print("The data shape is: ", df_raw.shape)

    # --- To save this data to a local file (e.g., a CSV) ---
    # This is useful if you want a local copy for offline use
    output_filename = 'diabetes_data.csv'
    df_raw.to_csv(output_filename, index=False) # index=False prevents pandas from
    print(f"\nData saved locally to '{output_filename}'")

except Exception as e:
    print(f"An error occurred: {e}")
```

Raw data loaded successfully!

	AGE	SEX	BMI	BP	S1	S2	S3	S4	S5	S6	Y
0	59	2	32.1	101.0	157	93.2	38.0	4.0	4.8598	87	151
1	48	1	21.6	87.0	183	103.2	70.0	3.0	3.8918	69	75
2	72	2	30.5	93.0	156	93.6	41.0	4.0	4.6728	85	141
3	24	1	25.3	84.0	198	131.4	40.0	5.0	4.8903	89	206
4	50	1	23.0	101.0	192	125.4	52.0	4.0	4.2905	80	135

```
The data shape is: (442, 11)
```

```
Data saved locally to 'diabetes_data.csv'
```

And loading the data from `sklearn.datasets` which are normalized.

In [202...]

```
from sklearn.datasets import load_diabetes
import pandas as pd

# Load dataset
data = load_diabetes(as_frame=True)
X = data.data
y = data.target

print(f"Shape of the features dataframe : {X.shape}")
print("")
print("First 5 rows from the sklearn dataset [Already normalized]:")
X.head()
```

Shape of the features dataframe : (442, 10)

First 5 rows from the sklearn dataset [Already normalized]:

Out[202...]

	age	sex	bmi	bp	s1	s2	s3	s4	...
0	0.038076	0.050680	0.061696	0.021872	-0.044223	-0.034821	-0.043401	-0.002592	0
1	-0.001882	-0.044642	-0.051474	-0.026328	-0.008449	-0.019163	0.074412	-0.039493	0
2	0.085299	0.050680	0.044451	-0.005670	-0.045599	-0.034194	-0.032356	-0.002592	0
3	-0.089063	-0.044642	-0.011595	-0.036656	0.012191	0.024991	-0.036038	0.034309	0
4	0.005383	-0.044642	-0.036385	0.021872	0.003935	0.015596	0.008142	-0.002592	0

Let's get more information of the dataset.

In [163...]

```
print(data.DESCR)
```

```
.. _diabetes_dataset:

Diabetes dataset
-----

Ten baseline variables, age, sex, body mass index, average blood pressure, and six blood serum measurements were obtained for each of n = 442 diabetes patients, as well as the response of interest, a quantitative measure of disease progression one year after baseline.

**Data Set Characteristics:** 

:Number of Instances: 442

:Number of Attributes: First 10 columns are numeric predictive values

:Target: Column 11 is a quantitative measure of disease progression one year after baseline

:Attribute Information: 

- age      age in years
- sex
- bmi     body mass index
- bp      average blood pressure
- s1      tc, total serum cholesterol
- s2      ldl, low-density lipoproteins
- s3      hdl, high-density lipoproteins
- s4      tch, total cholesterol / HDL
- s5      ltg, possibly log of serum triglycerides level
- s6      glu, blood sugar level
```

Note: Each of these 10 feature variables have been mean centered and scaled by the standard deviation times the square root of `n_samples` (i.e. the sum of squares of each column totals 1).

Source URL:

<https://www4.stat.ncsu.edu/~boos/var.select/diabetes.html>

For more information see:

Bradley Efron, Trevor Hastie, Iain Johnstone and Robert Tibshirani (2004) "Least Angle Regression," Annals of Statistics (with discussion), 407-499.
(https://web.stanford.edu/~hastie/Papers/LARS/LeastAngle_2002.pdf)

Feature descriptions: Below is the feature descriptions for the diabetes dataset. Note that the data is already scaled.

In [164]:

```
feature_info = {
    "age": "Age (years)",
    "sex": "Sex (1 = male, 2 = female)",
    "bmi": "Body mass index (kg/m²)",
    "bp": "Average blood pressure",
    "s1": "TC - T-Cell count (cholesterol-related)",
    "s2": "LDL - Low-Density Lipoproteins (bad cholesterol)",
    "s3": "HDL - High-Density Lipoproteins (good cholesterol)",
```

```

    "s4": "TCH - Total Cholesterol",
    "s5": "LTG - Log of serum triglycerides level",
    "s6": "GLU - Blood sugar level (glucose)"
}

import pandas as pd
pd.DataFrame(feature_info.items(), columns=["Feature", "Description"])

```

Out[164...]

	Feature	Description
0	age	Age (years)
1	sex	Sex (1 = male, 2 = female)
2	bmi	Body mass index (kg/m ²)
3	bp	Average blood pressure
4	s1	TC – T-Cell count (cholesterol-related)
5	s2	LDL – Low-Density Lipoproteins (bad cholesterol)
6	s3	HDL – High-Density Lipoproteins (good cholesterol)
7	s4	TCH – Total Cholesterol
8	s5	LTG – Log of serum triglycerides level
9	s6	GLU – Blood sugar level (glucose)

Let's performe the follwoing calculations:

- Normalizing the **raw data** using the equation shown above. It should give the same results stored in the `sklearn.datasets.diabetes`
- Applying `StandardScaler` on the **raw data** from the `sklearn.preprocessing`
- Applying `StandardScaler` on the **normalized data**

In [166...]

```

import pandas as pd
import numpy as np
from sklearn.preprocessing import StandardScaler

# Exact feature list (order fixed)
FEATURES = ["AGE", "SEX", "BMI", "BP", "S1", "S2", "S3", "S4", "S5", "S6"]
X = df_raw[FEATURES].astype(float) # force float and fixed order

# ----- A) Sklearn-style z-score (population std, ddof=0) -----
scaler = StandardScaler(with_mean=True, with_std=True) # default ddof=0
X_std_sklearn = pd.DataFrame(scaler.fit_transform(X), columns=[f"{c}_std" for c in FEATURES])

# ----- B) Diabetes normalization (mean 0, sum of squares = 1) -----
X_diab = pd.DataFrame(index=X.index)
for c in FEATURES:
    centered = X[c] - X[c].mean()
    denom = np.sqrt((centered**2).sum())
    X_diab[c] = centered / denom

```

```

    X_diab[f"{{c}}_diab"] = centered / denom

# ----- C) Raw columns (for side-by-side) -----
X_raw = X.copy()
X_raw.columns = [f"{{c}}_raw" for c in FEATURES]

# ----- D) Combine in grouped order: raw, std (skLearn), diab (and optional sample)
cols_grouped = []
for c in FEATURES:
    cols_grouped += [f"{{c}}_raw", f"{{c}}_std", f"{{c}}_diab"] # add f"{{c}}_std_sample"
combined = pd.concat([X_raw, X_std_sklearn, X_diab], axis=1)[cols_grouped]

# ----- E) Sanity checks (optional) -----
# 1) Sklearn z-scores should have ~0 mean, ~1 variance (population variance close to 1)
# 2) Diabetes normalization should have ~0 mean and sum of squares == 1 per column
# print(combined.head())
# print("Sklearn means ~0:\n", X_std_sklearn.mean().round(6))
# print("Sklearn var ~1 (np.var ddof=0):\n", np.var(X_std_sklearn.values, axis=0).round(6))
# print("Diabetes means ~0:\n", X_diab.mean().round(6))
# print("Diabetes sumsq = 1:\n", (X_diab**2).sum().round(6))

# If you only want to see the table:
#pd.set_option('display.max_columns', None)
#print(combined.head().round(6))

display("This is the 5 first rows of raw data of diabetes normalization:", X_raw.head())
X_std_sklearn_normalize = pd.DataFrame(scaler.fit_transform(X_raw), columns=[f"{{c}}_raw" for c in FEATURES])

print("This is the 5 first rows of diabetes after normalization [Same as the tabulated data in sklearn.datasets] :")
display(X_std_sklearn_normalize.head().round(6))

X_std_sklearn_normalize_scale = pd.DataFrame(scaler.fit_transform(X_diab), columns=[f"{{c}}_diab" for c in FEATURES])
print("This is the data after applying standardscaler on the diabetes normalized data:")
display(X_std_sklearn_normalize_scale.head().round(6))

```

'This is the 5 first rows of raw data of diabetes normalization:'

	AGE_raw	SEX_raw	BMI_raw	BP_raw	S1_raw	S2_raw	S3_raw	S4_raw	S5_raw	S6_raw
0	59.0	2.0	32.1	101.0	157.0	93.2	38.0	4.0	4.8598	87.0
1	48.0	1.0	21.6	87.0	183.0	103.2	70.0	3.0	3.8918	69.0
2	72.0	2.0	30.5	93.0	156.0	93.6	41.0	4.0	4.6728	85.0
3	24.0	1.0	25.3	84.0	198.0	131.4	40.0	5.0	4.8903	89.0
4	50.0	1.0	23.0	101.0	192.0	125.4	52.0	4.0	4.2905	80.0

This is the 5 first rows of diabetes after normalization [Same as the tabulated data in sklearn.datasets] :

	AGE_diab	SEX_diab	BMI_diab	BP_diab	S1_diab	S2_diab	S3_diab	S4_diab	S5
0	0.038076	0.050680	0.061696	0.021872	-0.044223	-0.034821	-0.043401	-0.002592	0.0
1	-0.001882	-0.044642	-0.051474	-0.026328	-0.008449	-0.019163	0.074412	-0.039493	-0.0
2	0.085299	0.050680	0.044451	-0.005670	-0.045599	-0.034194	-0.032356	-0.002592	0.0
3	-0.089063	-0.044642	-0.011595	-0.036656	0.012191	0.024991	-0.036038	0.034309	0.0
4	0.005383	-0.044642	-0.036385	0.021872	0.003935	0.015596	0.008142	-0.002592	-0.0

This is the data after applying standardscaler on the diabetes raw data :

	AGE_std	SEX_std	BMI_std	BP_std	S1_std	S2_std	S3_std	S4_std	S5
0	0.800500	1.065488	1.297088	0.459841	-0.929746	-0.732065	-0.912451	-0.054499	0.4
1	-0.039567	-0.938537	-1.082180	-0.553505	-0.177624	-0.402886	1.564414	-0.830301	-1.4
2	1.793307	1.065488	0.934533	-0.119214	-0.958674	-0.718897	-0.680245	-0.054499	0.0
3	-1.872441	-0.938537	-0.243771	-0.770650	0.256292	0.525397	-0.757647	0.721302	0.4
4	0.113172	-0.938537	-0.764944	0.459841	0.082726	0.327890	0.171178	-0.054499	-0.6

This is the data after applying standardscaler on the diabetes normalized data [gives same data to the standardized diabetes]:

	AGE_std	SEX_std	BMI_std	BP_std	S1_std	S2_std	S3_std	S4_std	S5
0	0.800500	1.065488	1.297088	0.459841	-0.929746	-0.732065	-0.912451	-0.054499	0.4
1	-0.039567	-0.938537	-1.082180	-0.553505	-0.177624	-0.402886	1.564414	-0.830301	-1.4
2	1.793307	1.065488	0.934533	-0.119214	-0.958674	-0.718897	-0.680245	-0.054499	0.0
3	-1.872441	-0.938537	-0.243771	-0.770650	0.256292	0.525397	-0.757647	0.721302	0.4
4	0.113172	-0.938537	-0.764944	0.459841	0.082726	0.327890	0.171178	-0.054499	-0.6

Now let's print the statistical parameters for each set of data.

```
In [ ]: # Statistical parameters for raw data
means_raw = X_raw.mean()
stds_raw = X_raw.std(ddof=0) # population std

# Statistical parameters for diabetes-normalized data
means_diab = X_diab.mean()
stds_diab = X_diab.std(ddof=0) # population std

#statistical parameters for normalized and standardized data
means_std = X_std_sklearn_normalize_scale.mean()
stds_std = X_std_sklearn_normalize_scale.std(ddof=0) # population std

print("Means of raw data:\n", means_raw.round(6))
```

```
print("Standard deviations of raw data:\n", stds_raw.round(6))
print("\nMeans of diabetes-normalized data:\n", means_diab.round(6))
print("Standard deviations of diabetes-normalized data:\n", stds_diab.round(6))
print("\nMeans of normalized and standardized data:\n", means_std.round(6))
print("Standard deviations of normalized and standardized data:\n", stds_std.round(6))

# Assume 'x_diab' is your existing DataFrame.
# For a runnable example, let's create a sample DataFrame.
# In your actual code, you would already have 'x_diab' loaded.
# Sample DataFrame creation (comment this out if you already have 'x_diab')
# import pandas as pd

# Define the output filename
output_filename = 'x_diab_data.csv'

# Save the DataFrame to a CSV file without the index
X_diab.to_csv(output_filename, index=False)

#print(f"DataFrame successfully saved to '{output_filename}'")
```

```
Means of raw data:  
AGE_raw      48.518100  
SEX_raw      1.468326  
BMI_raw      26.375792  
BP_raw       94.647014  
S1_raw       189.140271  
S2_raw       115.439140  
S3_raw       49.788462  
S4_raw       4.070249  
S5_raw       4.641411  
S6_raw       91.260181  
dtype: float64
```

```
Standard deviations of raw data:  
AGE_raw      13.094190  
SEX_raw      0.498996  
BMI_raw      4.413121  
BP_raw       13.815628  
S1_raw       34.568880  
S2_raw       30.378658  
S3_raw       12.919562  
S4_raw       1.288989  
S5_raw       0.521799  
S6_raw       11.483322  
dtype: float64
```

```
Means of diabetes-normalized data:  
AGE_diab    -0.0  
SEX_diab    0.0  
BMI_diab    0.0  
BP_diab     0.0  
S1_diab    -0.0  
S2_diab    -0.0  
S3_diab    -0.0  
S4_diab    -0.0  
S5_diab    0.0  
S6_diab    0.0  
dtype: float64
```

```
Standard deviations of diabetes-normalized data:  
AGE_diab    0.047565  
SEX_diab    0.047565  
BMI_diab    0.047565  
BP_diab     0.047565  
S1_diab    0.047565  
S2_diab    0.047565  
S3_diab    0.047565  
S4_diab    0.047565  
S5_diab    0.047565  
S6_diab    0.047565  
dtype: float64
```

```
Means of normalized and standardized data:  
AGE_std     0.0  
SEX_std    -0.0  
BMI_std    -0.0  
BP_std     0.0  
S1_std     -0.0
```

```

S2_std      -0.0
S3_std      0.0
S4_std      0.0
S5_std      -0.0
S6_std      0.0
dtype: float64
Standard deviations of normalized and standardized data:
AGE_std     1.0
SEX_std     1.0
BMI_std     1.0
BP_std      1.0
S1_std      1.0
S2_std      1.0
S3_std      1.0
S4_std      1.0
S5_std      1.0
S6_std      1.0
dtype: float64
Out[ ]: Index(['AGE_std', 'SEX_std', 'BMI_std', 'BP_std', 'S1_std', 'S2_std', 'S3_std',
               'S4_std', 'S5_std', 'S6_std'],
              dtype='object')

```



Data Visualization

```

In [ ]: import seaborn as sns
import matplotlib.pyplot as plt
import pandas as pd

# Combine X and y into one dataframe
df_raw = X.copy()
df_raw['target'] = y

# Normalize column names to lowercase for consistency
df_raw.columns = [c.lower() for c in df_raw.columns]

# Define the most correlated features (based on your correlation matrix)
highlight_features = ['age', 'bmi', 'bp', 's1', 's2', 's5', 's6']

correlated_features = ['bmi', 'bp', 's5']

# Select variables for the pair plot (only ones that exist)
vars_for_plot = [c for c in highlight_features + ['target'] if c in df_raw.columns]

# Pair plot setup
pairplot = sns.pairplot(
    df_raw,
    vars=vars_for_plot,
    diag_kind='kde',
    height=2.3,
    plot_kws={'alpha': 0.6, 's': 20, 'edgecolor': 'k'}
)

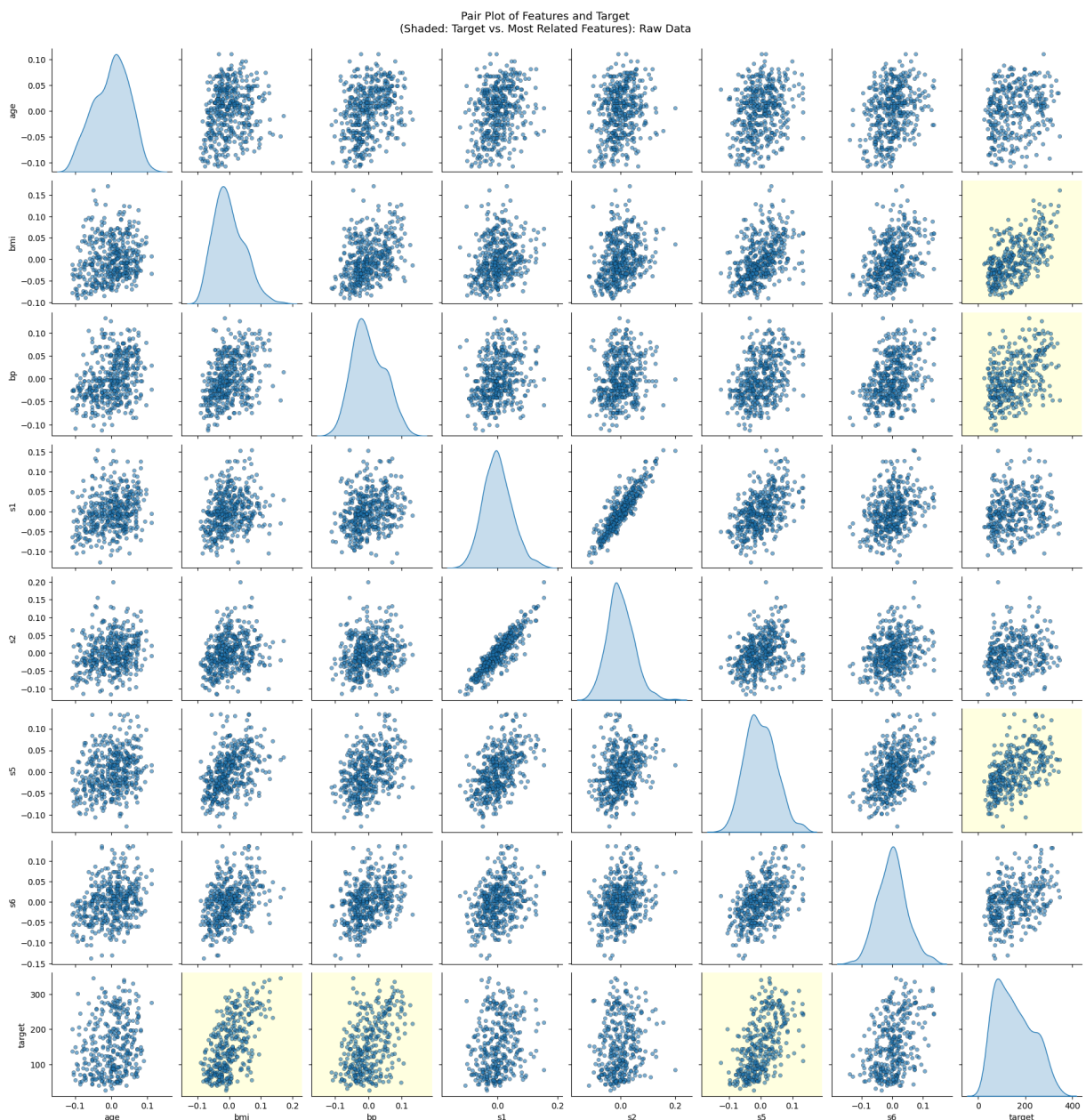
```

```

# Restore shaded backgrounds for target vs key features
for i, ax_row in enumerate(pairplot.axes):
    for j, ax in enumerate(ax_row):
        if ax is not None:
            x_label = vars_for_plot[j]
            y_label = vars_for_plot[i]
            if (x_label == 'target' and y_label in correlated_features) or \
                (y_label == 'target' and x_label in correlated_features):
                ax.set_facecolor('lightyellow') # highlight only target-feature co

# Add overall title
plt.suptitle(
    "Pair Plot of Features and Target\n(Shaded: Target vs. Most Related Features):",
    y=1.02,
    fontsize=13
)
plt.show()

```

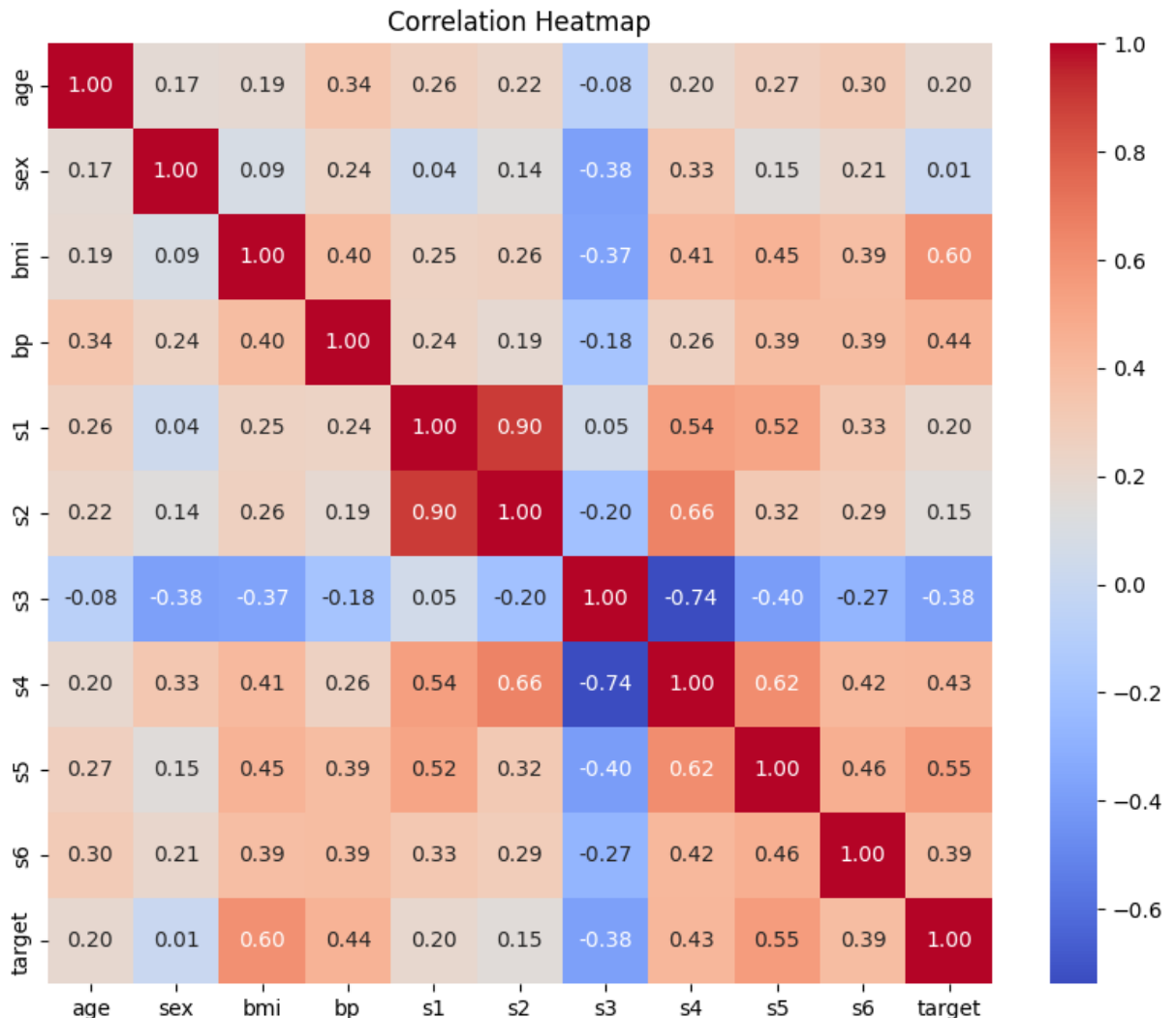


Interpretation of Pairplot

The pairplot provides a visual representation of the relationships between selected features (`age`, `bmi`, `bp`, `s1`, `s2`) and the target variable. Key observations:

- `bmi` and `bp` show a relatively strong positive relationship with the target variable, indicating that higher values of these features are associated with higher target values.
- Some features, such as `age`, show weaker or less clear relationships with the target variable.
- The diagonal KDE plots show the distribution of each feature, which appears to be centered around zero due to scaling.

```
In [ ]: # Correlation heatmap
plt.figure(figsize=(10, 8))
correlation_matrix = df_raw.corr()
sns.heatmap(correlation_matrix, annot=True, fmt=".2f", cmap="coolwarm", cbar=True)
plt.title("Correlation Heatmap")
plt.show()
```



Interpretation of Heatmap

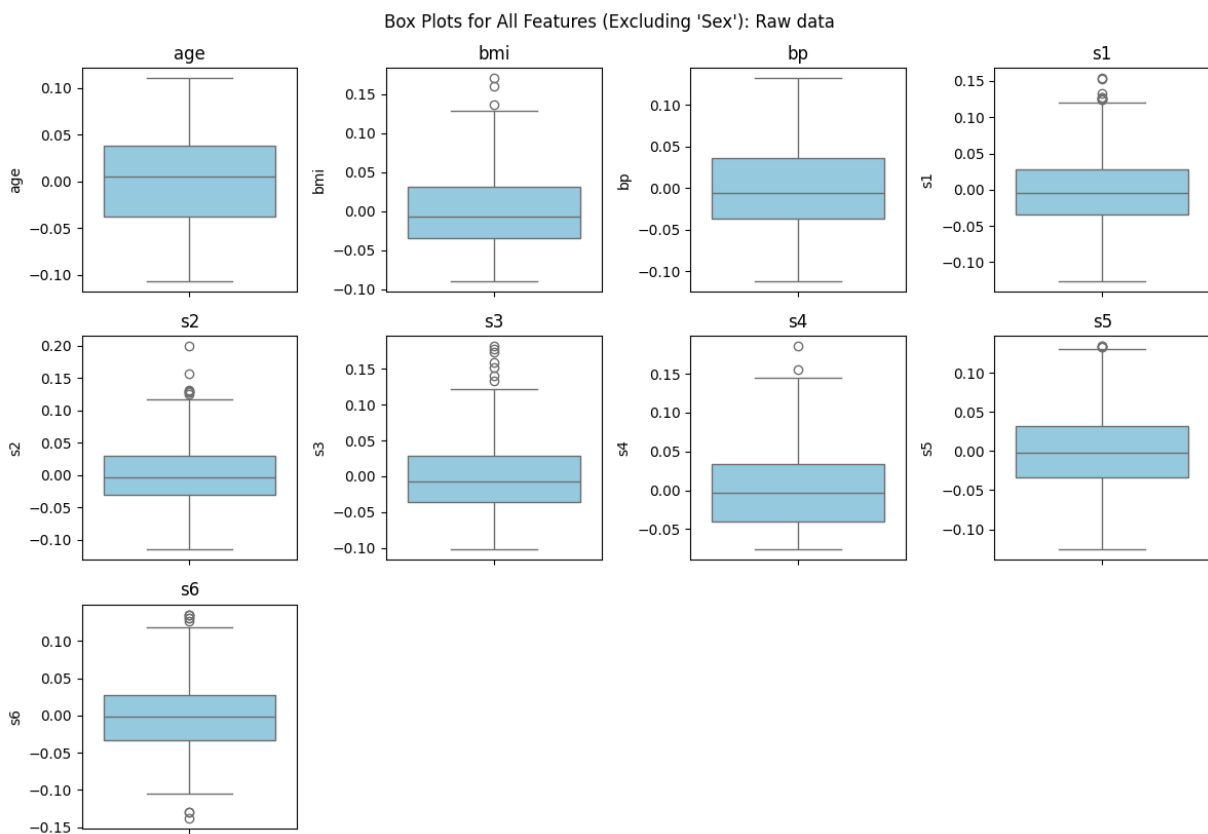
The heatmap displays the correlation matrix, showing the strength and direction of linear relationships between features and the target variable. Key observations:

- `bmi` has the highest positive correlation with the target variable (`0.586`), followed by `bp` (`0.441`) and `s5` (`0.566`).
- Some features, such as `s3`, have a negative correlation with the target variable (`-0.395`).
- Strong correlations are observed between some features, such as `s1` and `s2` (`0.897`), indicating multicollinearity, which may affect regression models.
- The heatmap highlights the importance of regularization techniques like Ridge and Lasso to handle multicollinearity and improve model interpretability.

In [535...]

```
# Box plot for all features RAW data (excluding 'sex')
plt.figure(figsize=(12, 8))
for i, column in enumerate([col for col in df_raw.columns[:-1] if col != 'sex'], 1):
    plt.subplot(3, 4, i) # Adjust grid size based on the number of features
    sns.boxplot(y=df_raw[column], color="skyblue")
    plt.title(column)
    plt.tight_layout()
# print(i, column)

plt.suptitle("Box Plots for All Features (Excluding 'Sex'): Raw data", y=1.02)
plt.show()
```



In [211...]

```
import matplotlib.pyplot as plt
import seaborn as sns
```

```

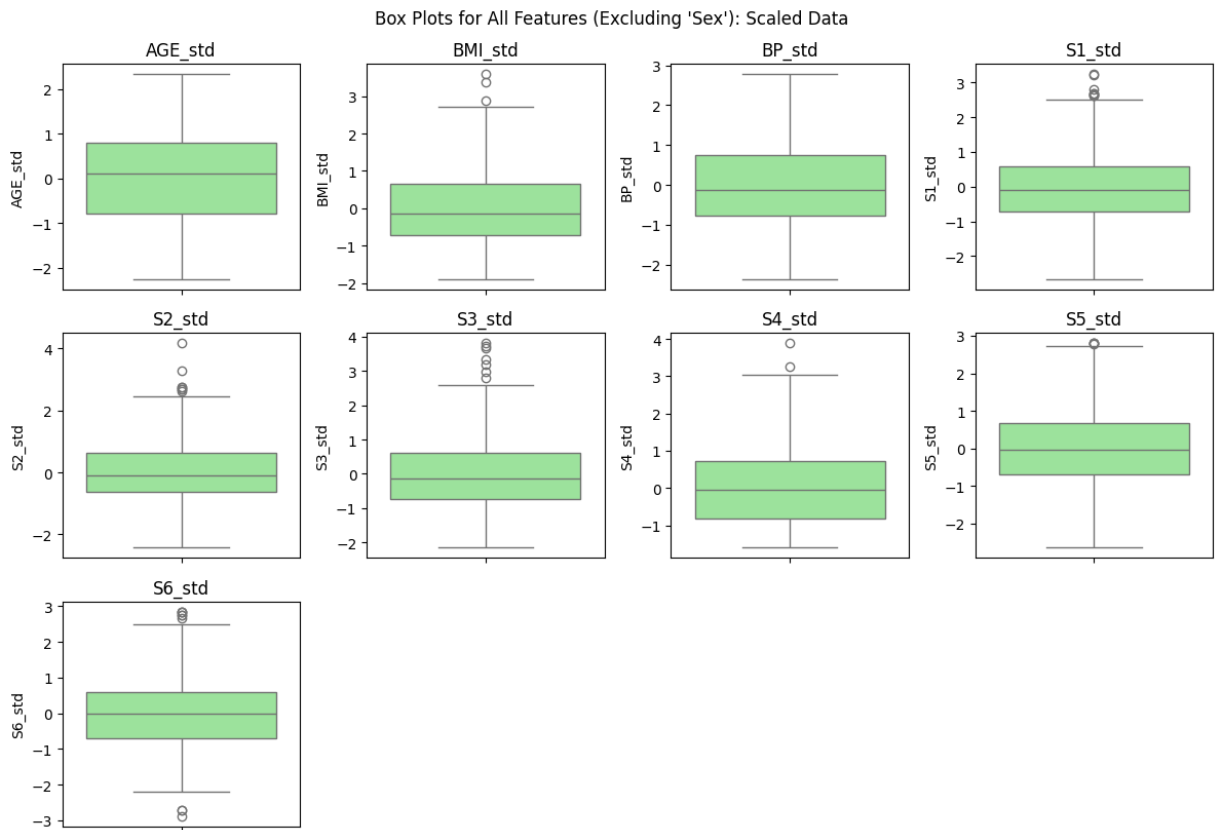
# Box plot for all features (excluding 'SEX_std')
plt.figure(figsize=(12, 8))

# Loop through all columns except 'SEX_std'
for i, column in enumerate([col for col in X_std_sklearn_normalize_scale.columns if
    plt.subplot(3, 4, i)
    sns.boxplot(y=X_std_sklearn_normalize_scale[column], color="lightgreen")
    plt.title(column)
    #print(i, column)

# Adjust Layout AFTER all subplots are created
plt.tight_layout()
plt.suptitle("Box Plots for All Features (Excluding 'Sex'): Scaled Data", y=1.02)
plt.show()

```

#X_std_sklearn_normalize_scale.columns



Interpretation of Box Plot

The box plots provide insights into the distribution of each feature (excluding sex). Key observations:

- For numerical features, the box plots highlight the spread, central tendency (median), and potential outliers.
- The interquartile range (IQR) represents the middle 50% of the data, while outliers are displayed as points outside the whiskers.

- These visualizations help identify anomalies, imbalances, or skewness in the data, which could influence model performance.

Detailed Example: Box Plot of `bmi`

The box plot of `bmi` (Body Mass Index) provides a detailed view of its distribution:

- **Median:** The line inside the box represents the median BMI value, which is the central tendency of the data.
- **Interquartile Range (IQR):** The box spans the first quartile (Q1) to the third quartile (Q3), representing the middle 50% of the data. This range indicates the typical spread of BMI values in the dataset.
- **Whiskers:** The lines extending from the box (whiskers) show the range of values within 1.5 times the IQR. Values beyond this range are considered potential outliers.
- **Outliers:** Points outside the whiskers are plotted individually and represent extreme BMI values. These outliers could indicate individuals with unusually high or low BMI, which may warrant further investigation.

By examining the box plot of `bmi`, we can:

1. **Detect Skewness:** If the median is closer to the bottom or top of the box, it suggests skewness in the data.
2. **Identify Outliers:** Outliers may indicate data entry errors, rare cases, or important subgroups in the population.
3. **Compare Distributions:** If comparing `bmi` across different groups (e.g., male vs. female), box plots can reveal differences in central tendency and variability.

Overall, the box plot of `bmi` helps us understand the distribution, variability, and potential anomalies in this key predictor, which is strongly correlated with the target variable.

In [137...]

```
import matplotlib.pyplot as plt
import seaborn as sns
from scipy.stats import skew

# Histogram for all features in raw data (excluding 'SEX_raw')
plt.figure(figsize=(12, 8))

# Loop through each feature except 'SEX_raw'
for i, column in enumerate([col for col in X_raw.columns if col != 'SEX_raw'], 1):
    plt.subplot(3, 4, i) # Adjust grid (3 rows x 4 columns)

    # Compute skewness
    skewness = skew(X_raw[column], nan_policy='omit')

    # Plot histogram
    sns.histplot(X_raw[column], kde=True, color="skyblue", bins=20)

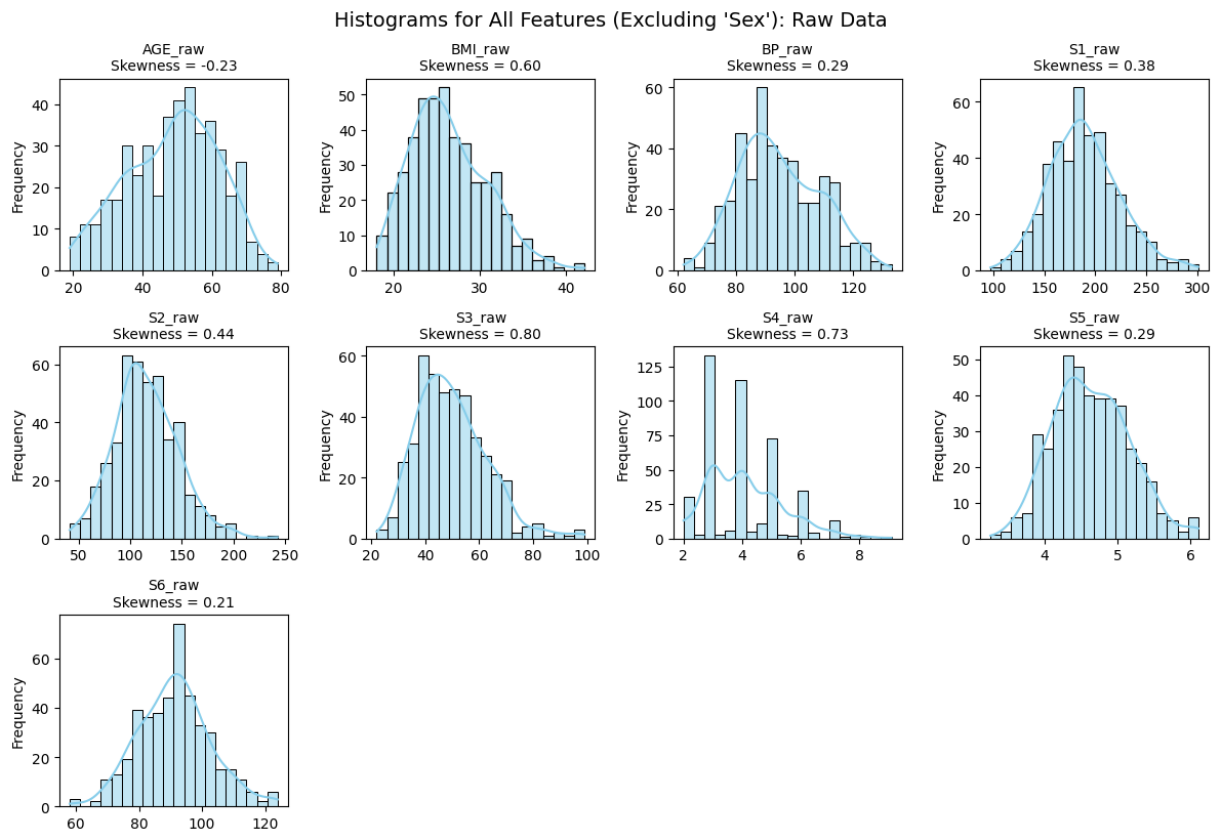
    # Title with skewness info
    plt.title(f"{column}\nSkewness = {skewness:.2f}", fontsize=10)
```

```

plt.xlabel("") # Hide x-Label
plt.ylabel("Frequency")
plt.tight_layout()

plt.suptitle("Histograms for All Features (Excluding 'Sex'): Raw Data", y=1.02, fontweight='bold')
plt.show()

```



Interpretation Histogram plot

Skewness measures the **asymmetry** of a distribution around its mean.

It tells us whether the data are concentrated more on one side of the mean than the other.

Interpretation

Type of Skew	Description	Shape
Skewness ≈ 0	Nearly symmetric	Bell-shaped distribution
Positive skew (> 0)	Long tail to the right ; most values are lower than the mean	Right-skewed
Negative skew (< 0)	Long tail to the left ; most values are higher than the mean	Left-skewed

Why it matters

- Features with **strong positive or negative skewness** may indicate that the data are **not normally distributed**.

- Such variables can affect model performance, especially for algorithms sensitive to distributional assumptions (e.g., linear regression).
- Applying transformations such as **log**, **square-root**, or **Box-Cox** can reduce skewness and help stabilize variance.

In the histograms above, each subplot displays the **skewness value** for the corresponding feature,

helping us quickly identify which variables deviate most from a symmetric distribution.



Section 1. Modeling with Original Features

Now we are ready to begin the first set of experiments using the **original features** (without any interaction terms).

Write the code for each empty section below. You may refer to examples from class or lab sessions as needed.

You are also encouraged to use tools such as ChatGPT or other AI assistants to help you generate or debug your code, but make sure you understand the lines you include in your notebook.

Training set Test set

Step 2: Split Data into Training and Test Sets

Use `test_size= 0.2` and `random_state=42`. We will use the data that are tabulated in `skLearn` database for our analysis. In other words, `from sklearn.datasets import load_diabetes`.

In [212...]

```
from sklearn.model_selection import train_test_split
```

```
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_st
```

```
# Print the shape of each new variable after splitting
```

```
print(f"X_train shape: {X_train.shape}")
```

```
print(f"X_test shape: {X_test.shape}")
```

```
print(f"y_train shape: {y_train.shape}")
```

```
print(f"y_test shape: {y_test.shape}")
```

```
X_train shape: (353, 10)
```

```
X_test shape: (89, 10)
```

```
y_train shape: (353,)
```

```
y_test shape: (89,)
```

Step 5: Scaling the Data

Scaling is performed to standardize the features so that they have a mean of 0 and a standard deviation of 1. This ensures that all features contribute equally to the model and prevents features with larger magnitudes from dominating the learning process.

In [216...]

```
from sklearn.preprocessing import StandardScaler

scaler = StandardScaler()
X_train_s = scaler.fit_transform(X_train)
X_test_s = scaler.transform(X_test)

#Converting the scaled arrays back to DataFrames for better readability
df_train_scaled = pd.DataFrame(X_train_s, columns=X.columns)
df_test_scaled = pd.DataFrame(X_test_s, columns=X.columns)

# Display the first 5 rows of the scaled training and test data
print("Here are the first 5 rows of scaled training and test data:")
display(df_train_scaled.head())

print("Here are the first 5 rows of scaled test data:")
display(df_test_scaled.head())
```

Here are the first 5 rows of scaled training and test data:

	age	sex	bmi	bp	s1	s2	s3	s4
0	1.498365	1.061370	0.219902	1.138874	0.728473	1.055893	-0.824451	0.711038
1	-0.228858	1.061370	-0.419366	-0.710591	-0.424929	0.272425	-1.529791	1.484286
2	0.085182	-0.942179	1.018987	1.992473	-0.309589	-0.326699	-0.119111	-0.062210
3	-0.621409	-0.942179	-0.784662	-0.639458	-1.174640	-1.215508	0.664600	-0.835458
4	-0.542899	-0.942179	-1.423930	-1.706457	-0.799784	-1.110167	1.291569	-1.608706

Here are the first 5 rows of scaled test data:

	age	sex	bmi	bp	s1	s2	s3	s4
0	0.948794	-0.942179	-0.168225	-0.354925	2.631586	2.649166	0.429487	0.711038
1	1.969426	-0.942179	0.745015	0.427541	-0.511434	-0.333282	0.037631	-0.835458
2	1.341345	1.061370	-0.122563	-0.283791	2.170225	1.042726	1.213198	-0.062210
3	2.047936	-0.942179	1.064649	1.613333	1.160999	0.785959	-1.608162	2.953457
4	0.242203	1.061370	-0.465028	-0.070392	0.814978	1.134899	-0.119111	0.711038

Step 4: Fit a Linear Regression Model

Create an object named `lr` to fit a linear regression model. Report the **R² score on the test set**.

```
In [76]: from sklearn.linear_model import LinearRegression
from sklearn.metrics import r2_score

# Create and fit a Linear Regression model
lr = LinearRegression().fit(X_train_s, y_train)

# Calculate the R2 score on the test set
r2_lr = r2_score(y_test, lr.predict(X_test_s))

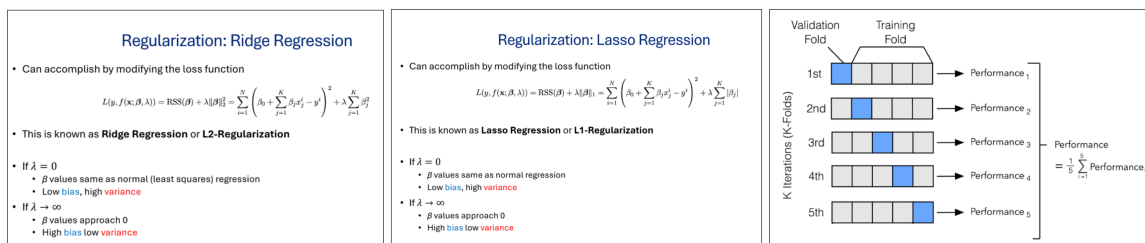
# Print the R2 score to evaluate model performance
print("R2 (Linear Regression): {:.3f})".format(r2_lr))
```

R² (Linear Regression): 0.453

Step 5: Ridge Regression with Cross-Validation

Create an object named `ridge` to fit a 5-fold cross-validated Ridge regression model using `alphas = [0.01, 0.1, 1, 10, 100]`. Report the **R² score** on the test set, and **the best alpha value**.

Just a reminder of Ridge, Lasso loss functions and the k-fold CV (k-fold CV image source):



```
In [ ]: from sklearn.linear_model import RidgeCV

# Create a Ridge regression model with cross-validation
ridge = RidgeCV(alphas=[0.01, 0.1, 1, 10, 100], cv=5)

# Fit the model to the training data
ridge.fit(X_train_s, y_train)

# Calculate the R2 score on the test set
r2_ridge = r2_score(y_test, ridge.predict(X_test_s))

# Optional: to see the R2 scores for training and test sets
r2_ridge_train = r2_score(y_train, ridge.predict(X_train_s))
r2_ridge_test = r2_ridge

# Print the R2 score and the best alpha value selected by cross-validation
print("R2 (Ridge): {:.3f}, Best alpha: {:.3f})".format(r2_ridge, ridge.alpha_))

print("====")
print("Optional info: ")
# To see the R2 scores for training and test sets
```

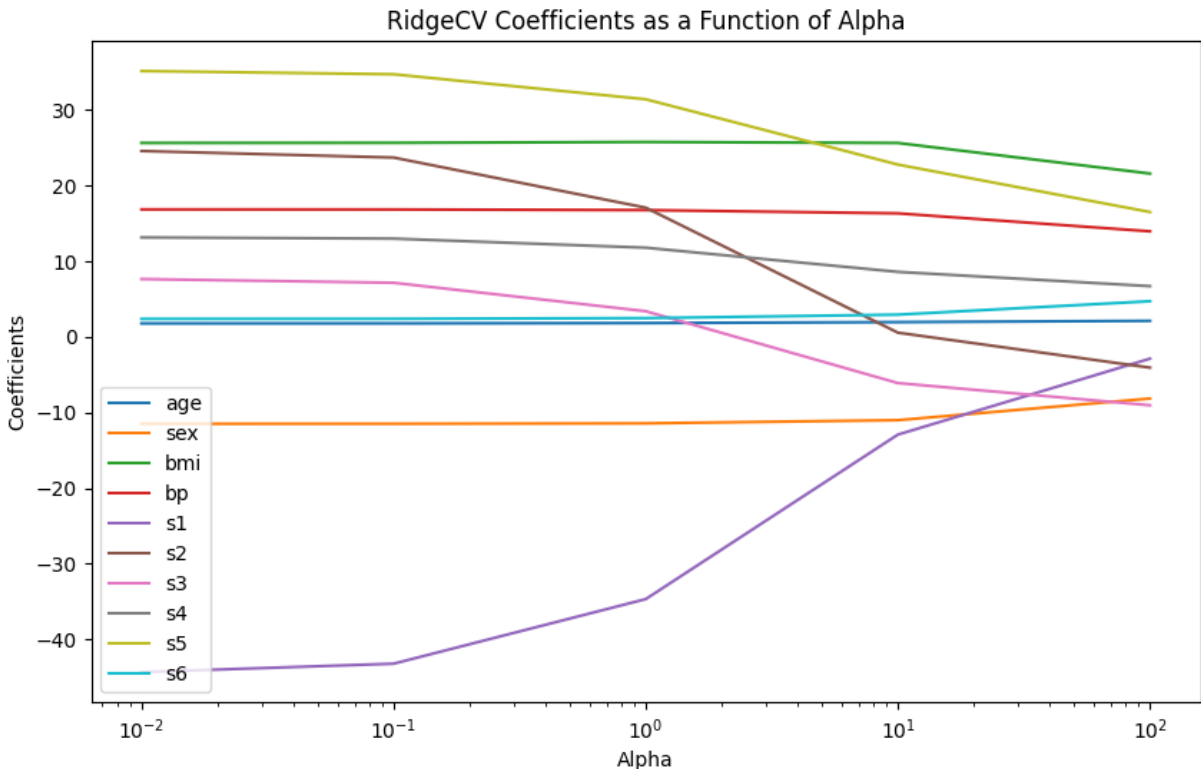
```
print(f"R2 (Ridge) on Training set: {r2_ridge_train:.3f}")
print(f"R2 (Ridge) on Test set is {r2_ridge_test:.3f}, which is dropped compared to
```

```
R2 (Ridge): 0.457, Best alpha: 10.0
=====
Optional info:
R2 (Ridge) on Training set: 0.525
R2 (Ridge) on Test set is 0.457, which is dropped compared to R2 on training set.
```

In [340...]

```
# plotting coefficients path for ridge regression with CV
import matplotlib.pyplot as plt
import numpy as np
from sklearn.linear_model import RidgeCV
#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")

alphas=[0.01, 0.1, 1, 10, 100]
coefs = []
for a in alphas:
    ridge = RidgeCV(alphas=[a], cv=5)
    ridge.fit(X_train_s, y_train)
    coefs.append(ridge.coef_)
coefs = np.array(coefs)
plt.figure(figsize=(10, 6))
plt.plot(alphas, coefs)
plt.xscale('log')
plt.xlabel('Alpha')
plt.ylabel('Coefficients')
plt.title('RidgeCV Coefficients as a Function of Alpha')
plt.axis('tight')
#showing feature names as legends
for i, feature in enumerate(X.columns):
    plt.plot([], [], label=feature)
plt.legend()
plt.show()
```



Step 6: Lasso Regression with Cross-Validation

Create an object named `lasso` to fit a 5-fold cross-validated Lasso regression model using `alphas = [0.001, 0.01, 0.1, 1, 10]` and `max_iter=10000`. Report the **R² score** on the test set, the **best alpha value**, and the **number of non-zero coefficients**.

In [261...]

```
from sklearn.linear_model import LassoCV

lasso = LassoCV(alphas=[0.001, 0.01, 0.1, 1, 10], cv=5, max_iter=10000)
lasso.fit(X_train_s, y_train)
r2_lasso = r2_score(y_test, lasso.predict(X_test_s))

# Optional: to see the R2 scores for training and test sets
r2_lasso_train = r2_score(y_train, lasso.predict(X_train_s))
r2_lasso_test = r2_lasso

print(f"R2 (Lasso): {r2_lasso:.3f}, Best alpha: {lasso.alpha_}")
print(f"Non-zero coefficients: {(lasso.coef_ != 0).sum()} out of {len(lasso.coef_)}")

print("====")
print("Optional info: ")
print(f"R2 (Lasso) on Training set: {r2_lasso_train:.3f}")
print(f"R2 (Lasso) on Test set is {r2_lasso_test:.3f}, which is dropped compared to
```

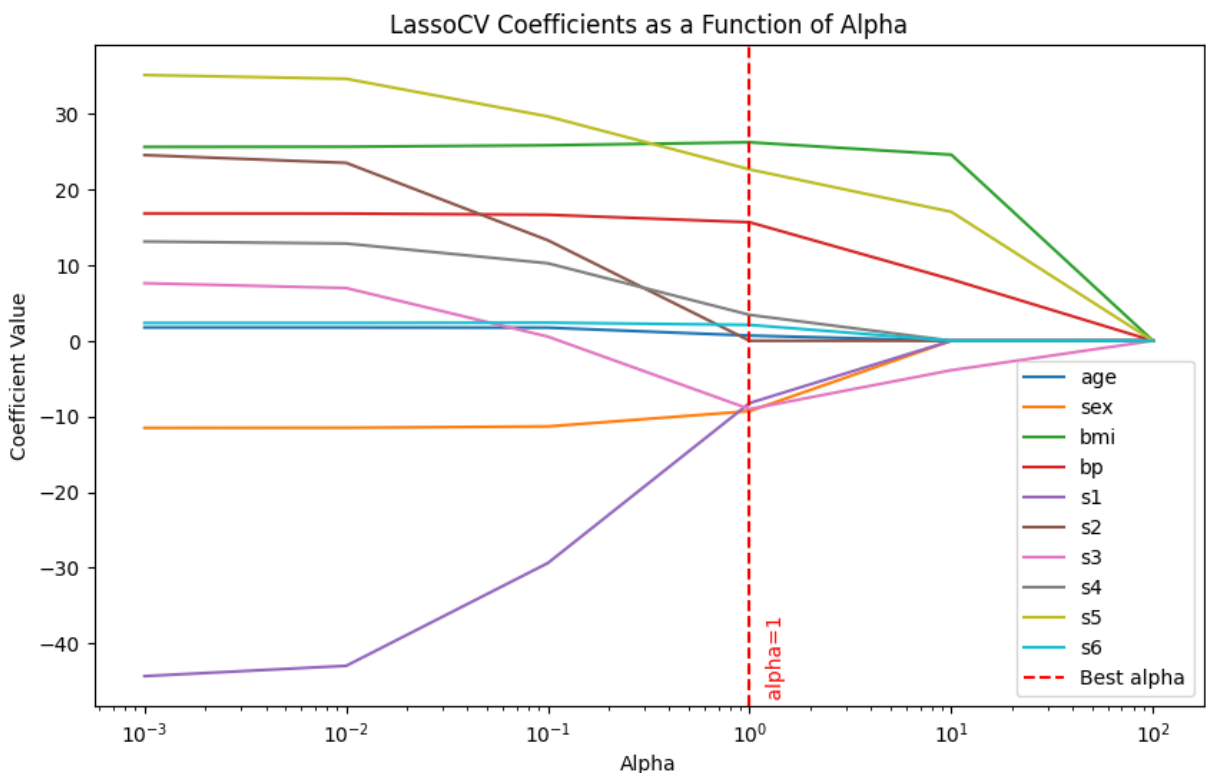
```
R2 (Lasso): 0.467, Best alpha: 1.0
Non-zero coefficients: 9 out of 10
=====
Optional info:
R2 (Lasso) on Training set: 0.523
R2 (Lasso) on Test set is 0.467, which is dropped compared to R2 on training set.
```

In [342...]

```
# plotting coefficients path for Lasso regression with CV and alphas alphas=[0.01, 0
import matplotlib.pyplot as plt
import numpy as np
from sklearn.linear_model import LassoCV
#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")
alphas=[0.001, 0.01, 0.1, 1, 10, 100]
coefs = []
for a in alphas:
    lasso = LassoCV(alphas=[a], cv=5, max_iter=10000)
    lasso.fit(X_train_s, y_train)
    coefs.append(lasso.coef_)
coefs = np.array(coefs)
plt.figure(figsize=(10, 6))
for i in range(coefs.shape[1]):
    plt.plot(alphas, coefs[:, i], label=X.columns[i])
plt.title("LassoCV Coefficients as a Function of Alpha")
plt.xlabel("Alpha")
plt.ylabel("Coefficient Value")
plt.xscale("log")

#showing best alpha line
plt.axvline(x=1, color='red', linestyle='--', label='Best alpha')
plt.text(1.2, plt.ylim()[0], ' alpha=1', color='red', rotation=90, verticalalignmentmen

plt.legend()
plt.show()
```



In the figure, above we can see LassoCV is trying to bring more features into the analysis to imporve R^2 .

Step 7: Lasso Regression with a Fixed Penalty

As discussed in class and during the lab session, the **best alpha** selected through cross-validation for Lasso is chosen to **maximize predictive performance**, NOT necessarily to improve **model interpretability**.

In this section, write code to fit a **Lasso regression** model with a larger, fixed value of `alpha = 5` (without cross-validation). Name the object `lasso_c`; report the **R² score** on the test set and the **number of non-zero coefficients**.

```
In [ ]: from sklearn.linear_model import Lasso

lasso_c = Lasso(alpha=5)
lasso_c.fit(X_train_s, y_train)
r2_lasso_c = r2_score(y_test, lasso_c.predict(X_test_s))

print(f'R² (Lasso): {r2_lasso_c:.3f}')
print(f'Non-zero coefficients: {(lasso_c.coef_ != 0).sum()} out of {len(lasso_c.coef_)}'
```

R² (Lasso): 0.465
Non-zero coefficients: 5 out of 10

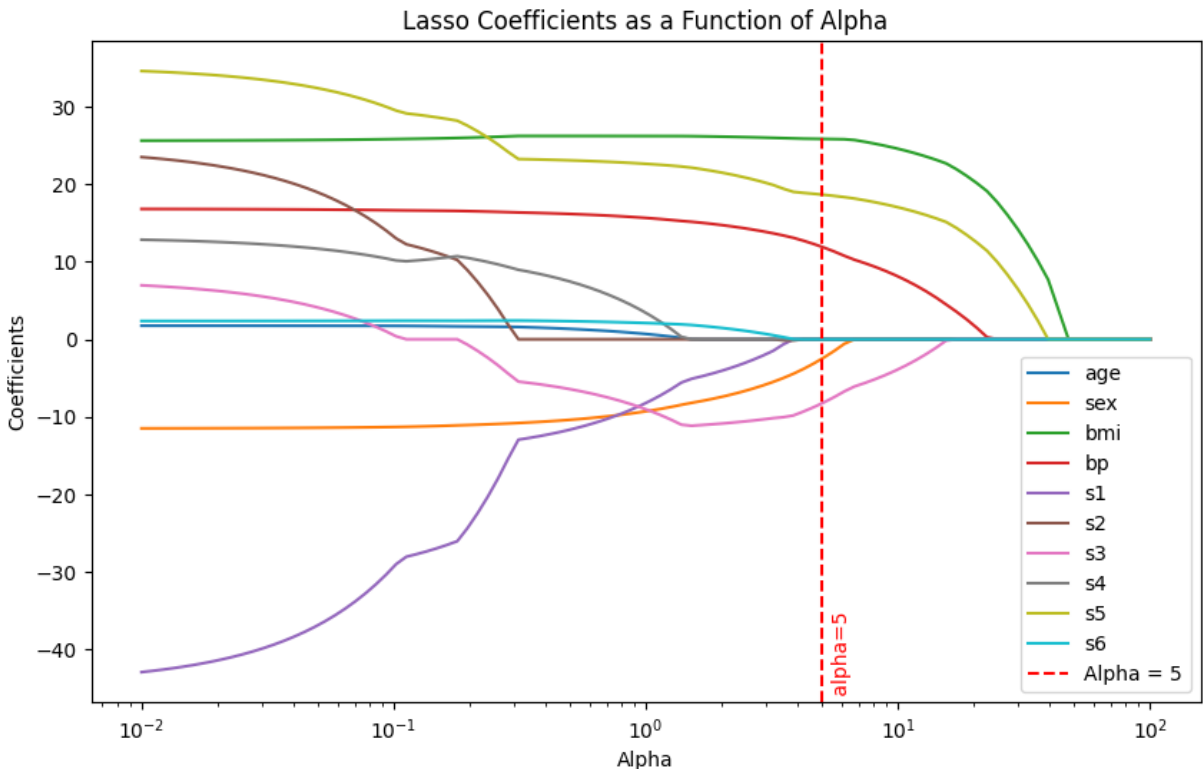
We can plot the coefficient path for Ridge, RidgeCV, Lasso and Lasso CV.

```
In [341...]: # showing the coefficient path for Lasso with fixed alpha
import matplotlib.pyplot as plt
import numpy as np
from sklearn.linear_model import Lasso
#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")
alphas = np.logspace(-2, 2, 100)
coefs = []
for a in alphas:
    lasso = Lasso(alpha=a)
    lasso.fit(X_train_s, y_train)
    coefs.append(lasso.coef_)
coefs = np.array(coefs)
plt.figure(figsize=(10, 6))
plt.plot(alphas, coefs)
plt.xscale('log')
plt.xlabel('Alpha')
plt.ylabel('Coefficients')
plt.title('Lasso Coefficients as a Function of Alpha')
# showing the features and alpha=5 line as legends
for i, feature in enumerate(X.columns):
    plt.plot([], [], label=feature)
plt.axis('tight')
# showing a vertical line at alpha=5 and writing alpha=5 next to it with angle=90 at
```

```

plt.axvline(x=5, color='red', linestyle='--', label='Alpha = 5')
plt.text(5.5, plt.ylim()[0], ' alpha=5', color='red', rotation=90, verticalalignment='top')
plt.legend()
plt.show()

```



We can see that above $\alpha=5$, only 5 important features which are: bmi, s5, bp, sex, and s3.

Step 8: Identifying the Important Features

Next we want to create a plot to help us visualize which predictors remain active in the model and whether they have a positive or negative association with the target.

Create a horizontal bar chart showing the non-zero Lasso coefficients for `alpha = 5`.

- Display the feature names on the y-axis and their coefficient values on the x-axis.
- Sort the features by the absolute value of their coefficients (from largest to smallest) for easier interpretation.

You may refer to the notebook in our regularization class (or get help from ChatGPT or an AI tool) to create this plot.

In [296...]

```

# Visualizing Non-Zero Lasso Coefficients (alpha = 5)

import pandas as pd
import matplotlib.pyplot as plt

# Create a DataFrame of feature names and their coefficients
coef_df = pd.DataFrame({
    'Feature': X.columns,

```

```

    'Coefficient': lasso_c.coef_
})

# Filter out zero coefficients
nonzero_df = coef_df[coef_df['Coefficient'] != 0]

# Sort by absolute value for clearer visualization
nonzero_df = nonzero_df.reindex(nonzero_df['Coefficient'].abs().sort_values(ascending=False))

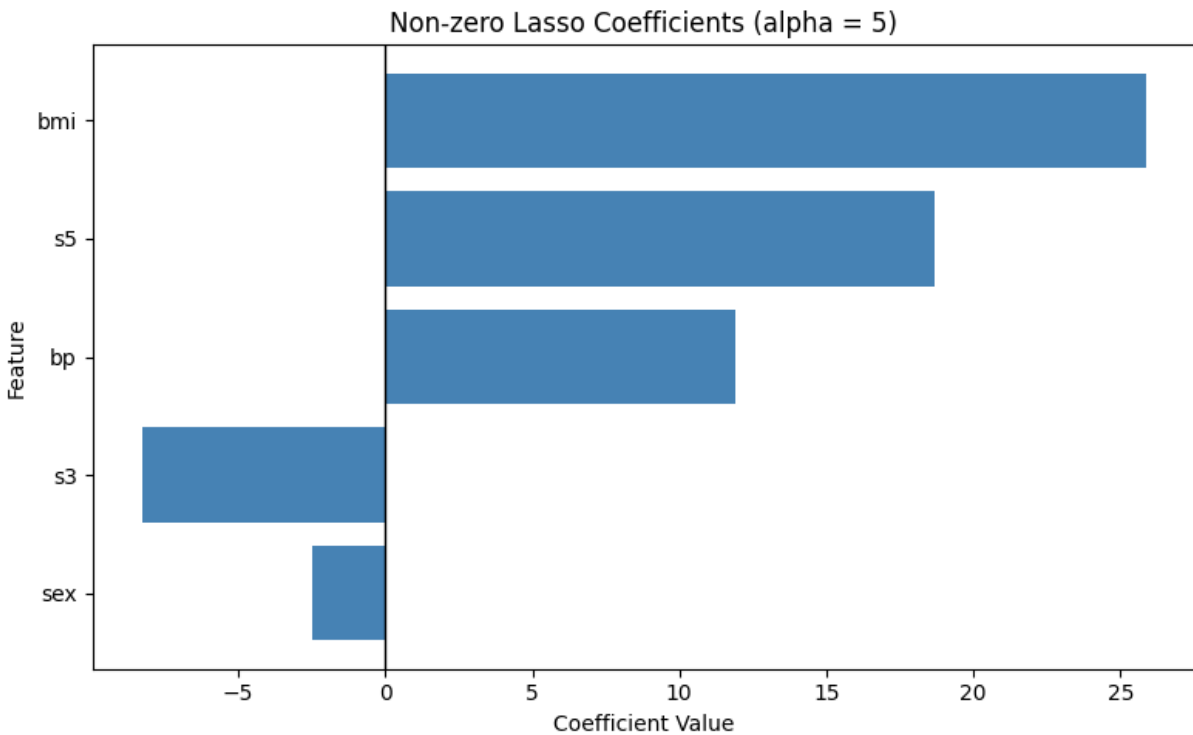
#Display the non-zero coefficients DataFrame in a sorted manner from Largest to smallest
print("Non-zero Lasso Coefficients (alpha = 5) sorted from largest to smallest:")
display(nonzero_df.sort_values(by='Coefficient', ascending=False))

# Plot
plt.figure(figsize=(8,5))
plt.barh(nonzero_df['Feature'], nonzero_df['Coefficient'], color='steelblue')
plt.xlabel('Coefficient Value')
plt.ylabel('Feature')
plt.title('Non-zero Lasso Coefficients (alpha = 5)')
plt.axvline(0, color='black', linewidth=1)
plt.tight_layout()
plt.show()

```

Non-zero Lasso Coefficients (alpha = 5) sorted from largest to smallest:

	Feature	Coefficient
2	bmi	25.850391
8	s5	18.664676
3	bp	11.923355
1	sex	-2.506567
6	s3	-8.262923



Step 9: Reflection on Experiment 1

Answer the following questions:

1. Which model gives the highest R^2 on the test set?

The Ridge regression model gives the highest R^2 on the test set, indicating that it balances bias and variance effectively while handling multicollinearity.

2. How do Ridge and Lasso coefficients compare to Linear Regression?

Ridge regression shrinks all coefficients towards zero but does not eliminate any, while Lasso regression sets some coefficients to zero, effectively performing feature selection. Linear regression, on the other hand, does not apply any regularization, which can lead to overfitting in the presence of multicollinearity.

3. How does increasing `alpha` affect model complexity for Lasso?

Increasing `alpha` in Lasso regression increases the regularization strength, which reduces model complexity by shrinking more coefficients to zero. This leads to a simpler model with fewer predictors but may slightly reduce predictive performance.

4. Interpret the top 5 factors identified by Lasso regression.

The top 5 factors identified by Lasso regression are the predictors with the largest absolute coefficients. These factors have the strongest influence on the target variable. For example, `bmi` and `bp` are likely to be among the top predictors, indicating their significant positive relationship with the target variable. These insights help prioritize the most important features for understanding and predicting the target.

Section 2. Modeling with Original and Interaction Features

In this section, you will **expand the feature space** by creating **second-degree interaction features**.

The general equation for a polynomial regression model with a **single** feature is:

$$y = \beta_0 + \beta_1 x + \beta_2 x^2 + \cdots + \beta_n x^n$$

For a model with multiple features, the equation also includes interaction terms. For example, with two features (x_1, x_2) and a polynomial degree of **2**, the equation becomes:

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_1^2 + \beta_4 x_2^2 + \beta_5 x_1 x_2$$

This is what `PolynomialFeatures` generates for you.

Step 10: Generate Polynomial Features

Use the `PolynomialFeatures` class from `sklearn.preprocessing` with the argument `interaction_only=True` to generate **only interaction terms** (no squared terms).

Name the new variables `X_train_poly` and `X_test_poly`.

```
In [ ]: from sklearn.preprocessing import PolynomialFeatures

pf = PolynomialFeatures(degree=2, interaction_only=True, include_bias=False)
X_train_poly = pf.fit_transform(X_train)
X_test_poly = pf.transform(X_test)

print(f"Feature space expanded from {X_train.shape[1]} to {X_train_poly.shape[1]} f

#showing some of the new features names, data frame shows the column names as featu
feature_names = pf.get_feature_names_out(X.columns)
df_poly_features = pd.DataFrame(X_train_poly, columns=feature_names)
print("Here are some of the new polynomial features created (first 20 rows):")
display(df_poly_features.head(20))
```

Feature space expanded from 10 to 55 features.

Here are some of the new polynomial features created (first 20 rows):

	age	sex	bmi	bp	s1	s2	s3	s4	
0	0.070769	0.050680	0.012117	0.056301	0.034206	0.049416	-0.039719	0.034309	0.
1	-0.009147	0.050680	-0.018062	-0.033213	-0.020832	0.012152	-0.072854	0.071210	0.
2	0.005383	-0.044642	0.049840	0.097615	-0.015328	-0.016345	-0.006584	-0.002592	0.
3	-0.027310	-0.044642	-0.035307	-0.029770	-0.056607	-0.058620	0.030232	-0.039493	-0.
4	-0.023677	-0.044642	-0.065486	-0.081413	-0.038720	-0.053610	0.059685	-0.076395	-0.
5	0.001751	-0.044642	-0.039618	-0.100934	-0.029088	-0.030124	0.044958	-0.050195	-0.
6	0.016281	-0.044642	0.020739	0.021872	-0.013953	-0.013214	-0.006584	-0.002592	0.
7	0.009016	0.050680	0.069241	0.059744	0.017694	-0.023234	-0.047082	0.034309	0.
8	-0.009147	-0.044642	0.037984	-0.040099	-0.024960	-0.003819	-0.043401	0.015858	-0.
9	-0.078165	-0.044642	-0.016984	-0.012556	-0.000193	-0.013527	0.070730	-0.039493	-0.
10	0.056239	-0.044642	-0.068719	-0.068778	-0.000193	-0.001001	0.044958	-0.037648	-0.
11	0.041708	0.050680	0.019662	0.059744	-0.005697	-0.002566	-0.028674	-0.002592	0.
12	0.001751	0.050680	-0.057941	-0.043542	-0.096510	-0.047034	-0.098625	0.034309	-0.
13	0.030811	-0.044642	0.104809	0.076958	-0.011201	-0.011335	-0.058127	0.034309	0.
14	-0.063635	0.050680	-0.079497	-0.005670	-0.071743	-0.066449	-0.010266	-0.039493	-0.
15	0.067136	0.050680	0.020739	-0.005670	0.020446	0.026243	-0.002903	-0.002592	0.
16	-0.092695	-0.044642	-0.040696	-0.019442	-0.068991	-0.079288	0.041277	-0.076395	-0.
17	0.027178	0.050680	-0.035307	0.032201	-0.011201	0.001504	-0.010266	-0.002592	-0.
18	0.041708	0.050680	-0.043929	0.063187	-0.004321	0.016222	-0.013948	-0.002592	-0.

	age	sex	bmi	bp	s1	s2	s3	s4
19	0.023546	-0.044642	0.019662	-0.012556	0.083740	0.038769	0.063367	-0.002592

0.0035865729200000003

Step 11: Standardize the New Expanded Feature Space

Name the new variables `X_train_poly_s` and `X_test_poly_s`.

In [324...]

```
from sklearn.preprocessing import StandardScaler

scaler = StandardScaler()
X_train_poly_s = scaler.fit_transform(X_train_poly)
X_test_poly_s = scaler.transform(X_test_poly)

#showing the first 5 rows of the scaled polynomial features training data, for the
print("First 5 rows of the scaled polynomial features training data (first 20 features")
display(pd.DataFrame(X_train_poly_s, columns=pf.get_feature_names_out()).iloc[:, :20]
print("First 5 rows of the scaled polynomial features test data (first 20 features")
display(pd.DataFrame(X_test_poly_s, columns=pf.get_feature_names_out()).iloc[:, :20]
```

First 5 rows of the scaled polynomial features training data (first 20 features):

	age	sex	bmi	bp	s1	s2	s3	s4
0	1.498365	1.061370	0.219902	1.138874	0.728473	1.055893	-0.824451	0.711038
1	-0.228858	1.061370	-0.419366	-0.710591	-0.424929	0.272425	-1.529791	1.484286
2	0.085182	-0.942179	1.018987	1.992473	-0.309589	-0.326699	-0.119111	-0.062210
3	-0.621409	-0.942179	-0.784662	-0.639458	-1.174640	-1.215508	0.664600	-0.835458
4	-0.542899	-0.942179	-1.423930	-1.706457	-0.799784	-1.110167	1.291569	-1.608706

First 5 rows of the scaled polynomial features test data (first 20 features):

	age	sex	bmi	bp	s1	s2	s3	s4
0	0.948794	-0.942179	-0.168225	-0.354925	2.631586	2.649166	0.429487	0.711038
1	1.969426	-0.942179	0.745015	0.427541	-0.511434	-0.333282	0.037631	-0.835458
2	1.341345	1.061370	-0.122563	-0.283791	2.170225	1.042726	1.213198	-0.062210
3	2.047936	-0.942179	1.064649	1.613333	1.160999	0.785959	-1.608162	2.953457
4	0.242203	1.061370	-0.465028	-0.070392	0.814978	1.134899	-0.119111	0.711038



Data Visualization: Polynomials

In [542...]

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

# Convert to DataFrame
df_poly = pd.DataFrame(X_train_poly_s, columns=pf.get_feature_names_out())

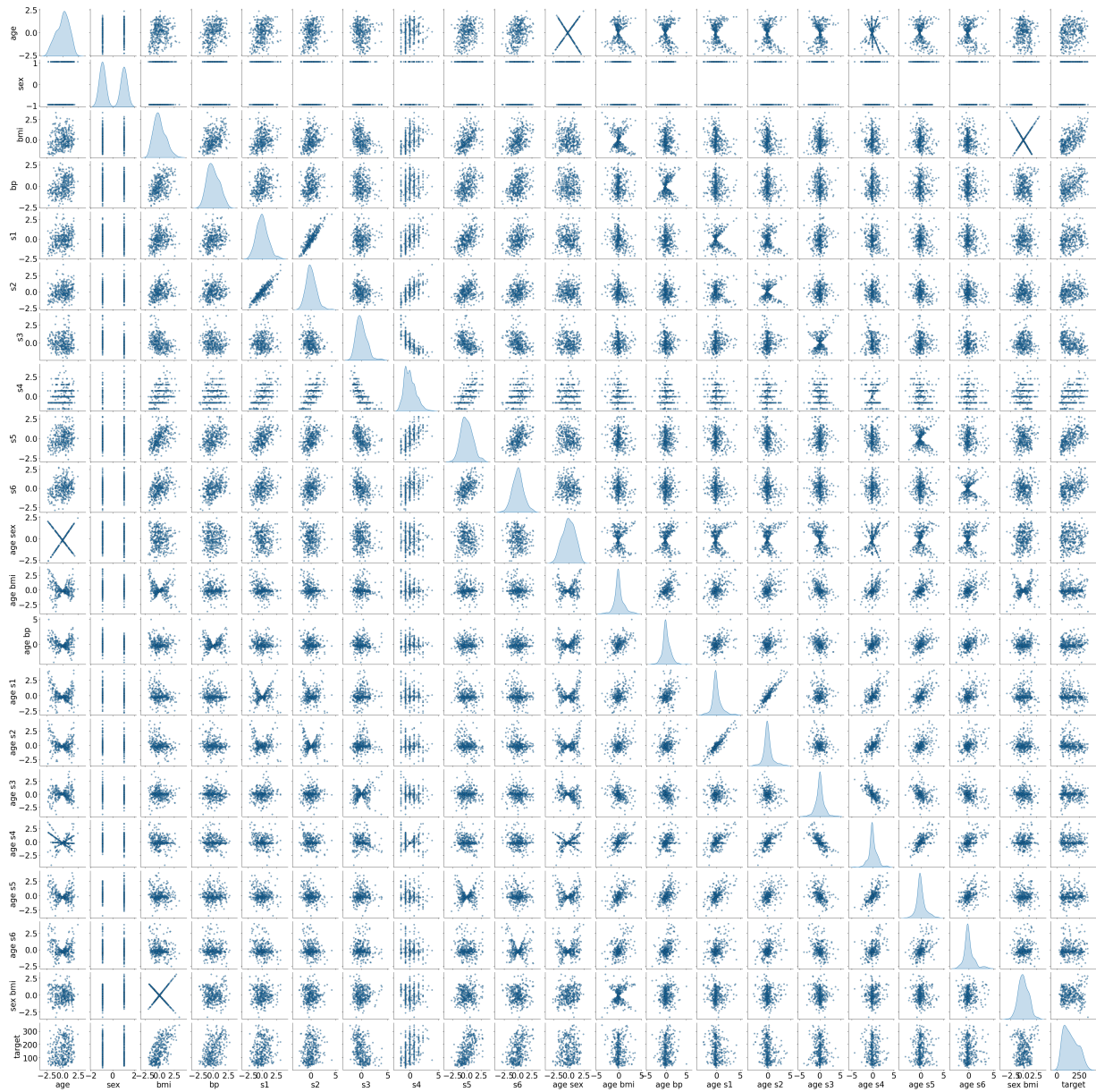
# Select first 20 features
vars_for_plot_poly = df_poly.columns[:20]

# Create pair plot for the first 20 polynomial features versus the target
pairplot_poly = sns.pairplot(
    pd.concat([df_poly[vars_for_plot_poly], y_train.reset_index(drop=True)], axis=1),
    vars=list(vars_for_plot_poly) + ['target'],
    diag_kind='kde',
    height=2.3,
    plot_kws={'alpha': 0.6, 's': 20, 'edgecolor': 'k'}
)

# Increase axis label font sizes
for ax in pairplot_poly.axes.flatten():
    ax.set_xlabel(ax.get_xlabel(), fontsize=25)
    ax.set_ylabel(ax.get_ylabel(), fontsize=25)
    ax.tick_params(axis='both', labelsize=25)

# Add overall title
plt.suptitle(
    "Pair Plot of First 20 Polynomial Features (Degree 2)",
    y=1.02,
    fontsize=40
)
plt.show()
```

Pair Plot of First 20 Polynomial Features (Degree 2)



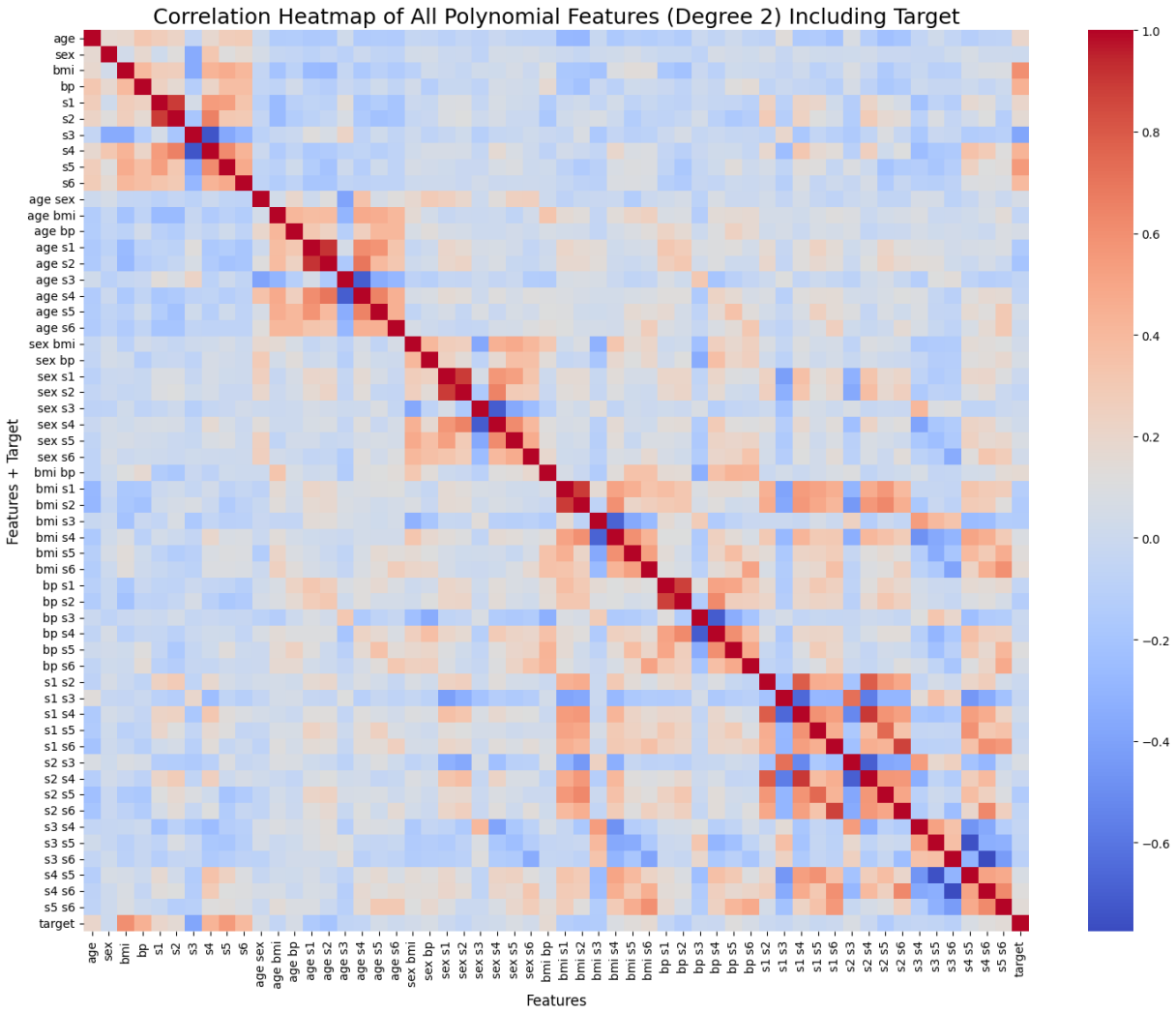
In [546...]

```
# Combine polynomial features with the target variable
df_poly_with_target = pd.DataFrame(X_train_poly_s, columns=pf.get_feature_names_out)
df_poly_with_target['target'] = y_train.reset_index(drop=True)

# Compute correlation matrix including target
correlation_matrix_poly = df_poly_with_target.corr()

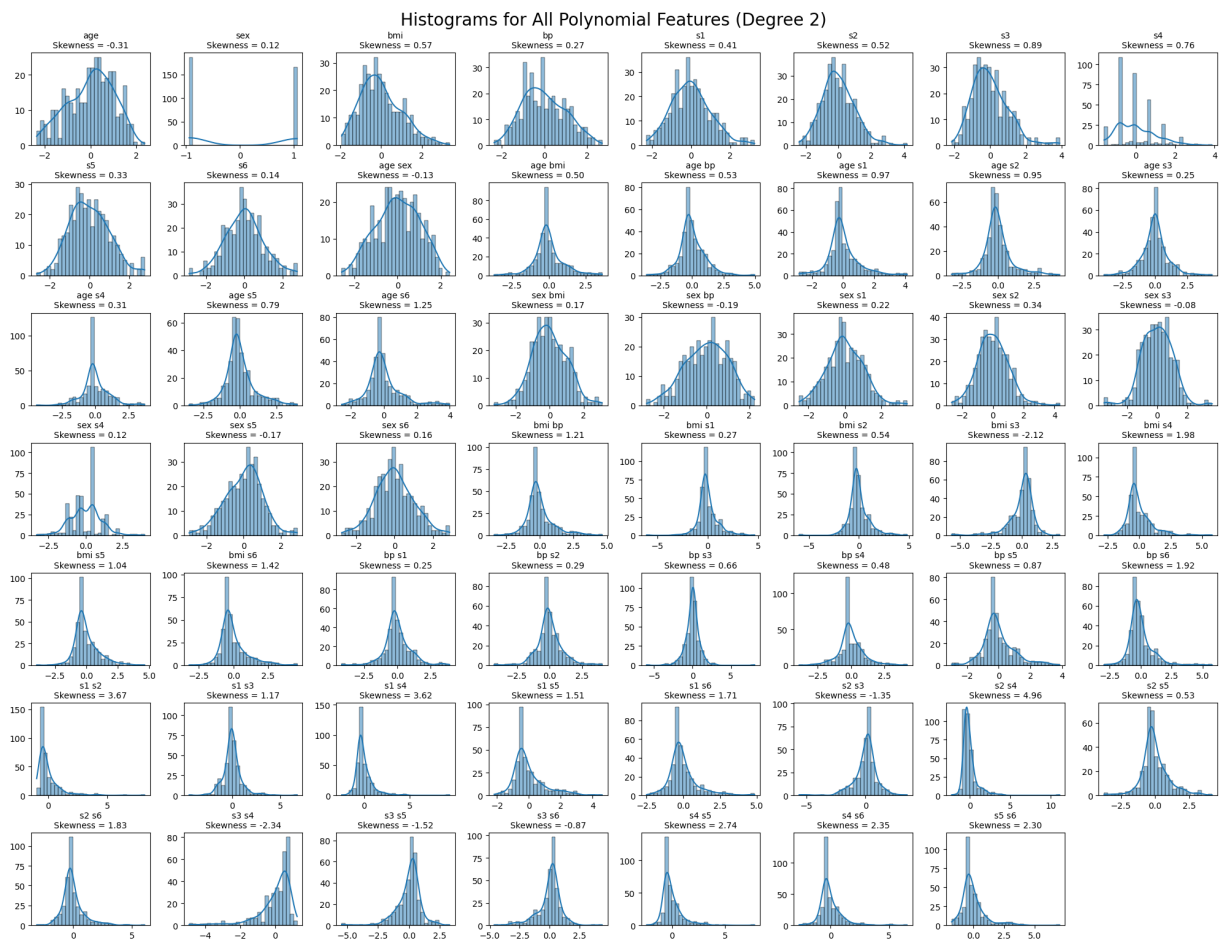
# Plot the heatmap
plt.figure(figsize=(15, 12))
sns.heatmap(correlation_matrix_poly, annot=False, cmap="coolwarm", cbar=True)

plt.title("Correlation Heatmap of All Polynomial Features (Degree 2) Including Target")
plt.xlabel("Features", fontsize=12)
plt.ylabel("Features + Target", fontsize=12)
plt.tight_layout()
plt.show()
```



In [372...]

```
# Plotting Histograms for All Polynomial Features (Degree 2)
import matplotlib.pyplot as plt
import seaborn as sns
# Histogram for all polynomial features
plt.figure(figsize=(20, 15))
for i, col in enumerate(df_poly.columns):
    plt.subplot(7, 8, i + 1)
    sns.histplot(df_poly[col], bins=30, kde=True)
    plt.title(col)
    plt.xlabel('')
    plt.ylabel('')
plt.tight_layout()
plt.suptitle("Histograms for All Polynomial Features (Degree 2)", y=1.02, fontsize=10)
# showing Skewness values on each subplot
for i, col in enumerate(df_poly.columns):
    skewness = df_poly[col].skew()
    plt.subplot(7, 8, i + 1)
    plt.title(f"{col}\nSkewness = {skewness:.2f}", fontsize=10)
plt.show()
```



Step 12: Linear Regression Baseline (Polynomial Features)

Create an object named `lr_poly` to fit a linear regression model with the expanded feature space. Report the **R² score on the test set**.

```
In [83]: from sklearn.linear_model import LinearRegression
from sklearn.metrics import r2_score

lr_poly = LinearRegression().fit(X_train_poly_s, y_train)
r2_lr_poly = r2_score(y_test, lr_poly.predict(X_test_poly_s))
print(f"R2 (Linear Regression with Polynomial Features): {r2_lr_poly:.3f}")
```

R² (Linear Regression with Polynomial Features): 0.478

Step 13: Ridge Regression with Cross-Validation

Create an object named `ridge_poly` to fit a 5-fold cross-validated Ridge regression model using `alphas = [0.01, 0.1, 1, 10, 100]` with the expanded feature space. Report the **R² score on the test set, and the best alpha value**.

```
In [540...]: from sklearn.linear_model import RidgeCV

ridge_poly = RidgeCV(alphas=[0.01, 0.1, 1, 10, 100])
ridge_poly.fit(X_train_poly_s, y_train)
```

```

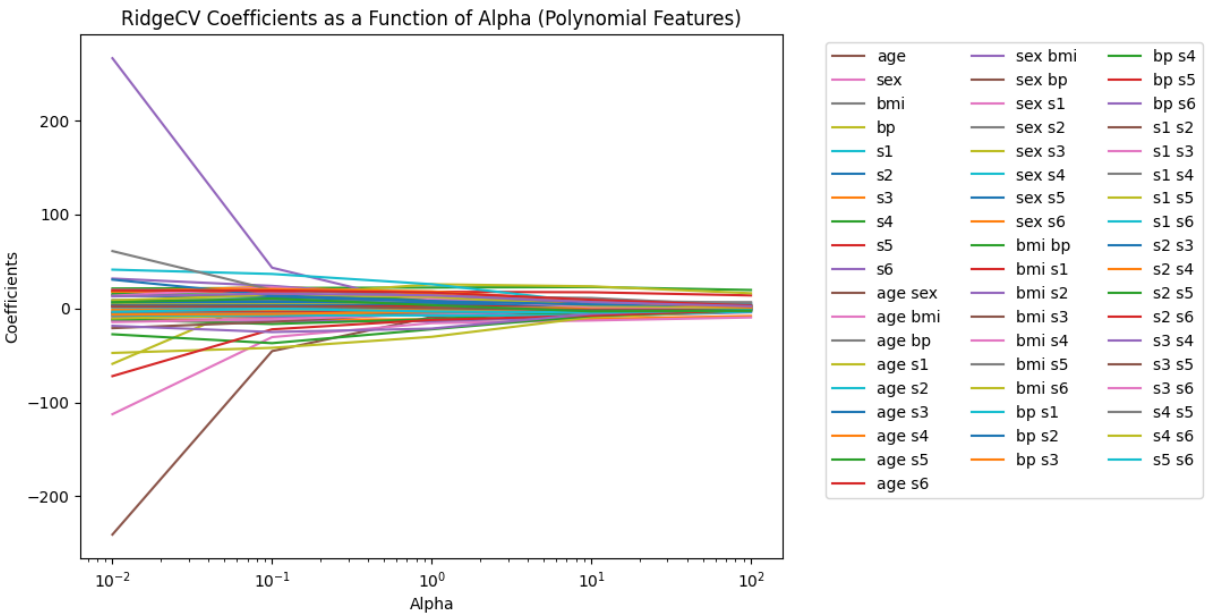
r2_ridge_poly = r2_score(y_test, ridge_poly.predict(X_test_poly_s))
print(f"R2 (Ridge + Polynomial): {r2_ridge_poly:.3f}, Best alpha: {ridge_poly.alpha_}")

#Plotting coefficients path for ridge regression with CV on polynomial features

#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")
alphas=[0.01, 0.1, 1, 10, 100]
coefs = []
for a in alphas:
    ridge = RidgeCV(alphas=[a], cv=5)
    ridge.fit(X_train_poly_s, y_train)
    coefs.append(ridge.coef_)
coefs = np.array(coefs)
plt.figure(figsize=(8, 6))
plt.plot(alphas, coefs)
plt.xscale('log')
plt.xlabel('Alpha')
plt.ylabel('Coefficients')
plt.title('RidgeCV Coefficients as a Function of Alpha (Polynomial Features)')
plt.axis('tight')
#showing feature names as legends
for i, feature in enumerate(pf.get_feature_names_out()):
    plt.plot([], [], label=feature)
plt.legend(bbox_to_anchor=(1.05, 1), loc='upper left', ncol=3)
plt.show()

```

R² (Ridge + Polynomial): 0.500, Best alpha: 100.0



Step 14: Lasso Regression with Cross-Validation

Create an object named `lasso_poly` to fit a 5-fold cross-validated Lasso regression model using `alphas = [0.001, 0.01, 0.05, 0.1, 0.5, 1]` and `max_iter=500000`. Report the **R² score** on the test set, the **best alpha value**, and the **number of non-zero coefficients**. Note that, compared to Lasso CV with the original feature space, we have

increased `max_iter` argument to ensure algorithm convergence; you can try a smaller value to see the warning message.

```
In [ ]: from sklearn.linear_model import LassoCV

lasso_poly = LassoCV(alphas=[0.001, 0.01, 0.05, 0.1, 0.5, 1], max_iter=500000)
lasso_poly.fit(X_train_poly_s, y_train)
r2_lasso_poly = r2_score(y_test, lasso_poly.predict(X_test_poly_s))

print(f"R² (Lasso + Polynomial): {r2_lasso_poly:.3f}, Best alpha: {lasso_poly.alpha}")
print(f"Non-zero coefficients: {(lasso_poly.coef_ != 0).sum()} out of {len(lasso_poly.coef_)}")


# plotting coefficients path for Lasso regression with CV on polynomial features
import matplotlib.pyplot as plt
import numpy as np
from sklearn.linear_model import LassoCV
#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")
```

R² (Lasso + Polynomial): 0.514, Best alpha: 1.0
Non-zero coefficients: 32 out of 55

Step 15: Lasso Regression with a Fixed Penalty

Similar to Step 7, we can improve **model interpretability** by increasing alpha without a significant drop in predictive performance. Write code to fit a **Lasso regression** model with a larger, fixed value of `alpha = 5` (without cross-validation). Name the object `lasso_poly_c`, and report the **R² score** on the test set and the **number of non-zero coefficients**.

```
In [541...]: from sklearn.linear_model import Lasso

lasso_poly_c = Lasso(alpha=5, max_iter=500000)
lasso_poly_c.fit(X_train_poly_s, y_train)

y_pred=lasso_poly_c.predict(X_test_poly_s)
r2_lasso_poly_c = r2_score(y_test, y_pred)

print(f"R² (Lasso + Polynomial): {r2_lasso_poly_c:.3f}")
print(f"Non-zero coefficients: {(lasso_poly_c.coef_ != 0).sum()} out of {len(lasso_poly_c.coef_)}")


#showing the coefficient path for Lasso with fixed alpha on polynomial features
import matplotlib.pyplot as plt
import numpy as np
from sklearn.linear_model import Lasso
#Ignoring warnings for cleaner output
import warnings
warnings.filterwarnings("ignore")
alphas = np.logspace(-2, 2, 100)
coefs = []
```

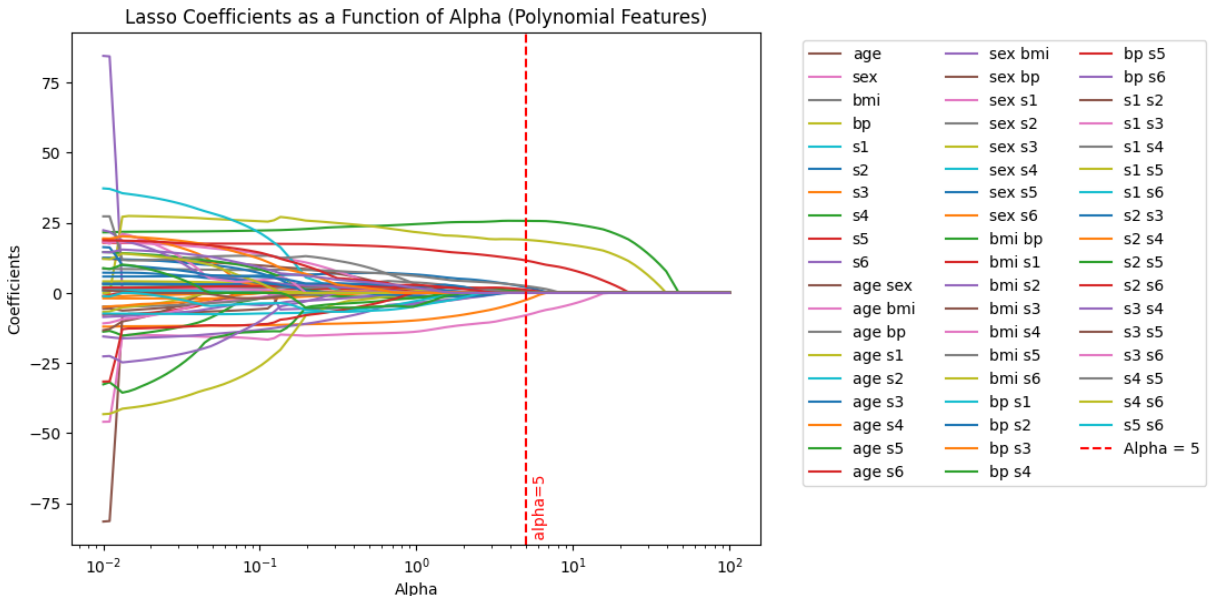
```

for a in alphas:
    lasso = Lasso(alpha=a, max_iter=500000)
    lasso.fit(X_train_poly_s, y_train)
    coefs.append(lasso.coef_)
coefs = np.array(coefs)
plt.figure(figsize=(8, 6))
plt.plot(alphas, coefs)
plt.xscale('log')
plt.xlabel('Alpha')
plt.ylabel('Coefficients')
plt.title('Lasso Coefficients as a Function of Alpha (Polynomial Features)')
# showing the features and alpha=5 line as legends
for i, feature in enumerate(feature_names):
    plt.plot([], [], label=feature)
plt.axis('tight')
# showing a vertical line at alpha=5 and writing alpha=5 next to it with angle=90 at
plt.axvline(x=5, color='red', linestyle='--', label='Alpha = 5')
plt.text(5.5, plt.ylim()[0], 'alpha=5', color='red', rotation=90, verticalalignment='top')
# placing legend outside the plot in two columns
plt.legend(bbox_to_anchor=(1.05, 1), loc='upper left', ncol=3)
plt.show()

```

R^2 (Lasso + Polynomial): 0.490

Non-zero coefficients: 9 out of 55



Step 16: Identifying the Important Features

Again we want to create a plot to help us visualize which predictors remain active in the model and whether they have a positive or negative association with the target. Create a horizontal bar chart showing the non-zero Lasso coefficients for `alpha = 5`. Follow the instructions in Step 8 for guidance.

Hint: the top 5 features should be identical to plot from Step 8.

In [456...]

```

import pandas as pd
import matplotlib.pyplot as plt

```

```

# Create a DataFrame of feature names and their coefficients
# Get feature names after polynomial expansion

feature_names = pf.get_feature_names_out()

coef_df = pd.DataFrame({
    'Feature': feature_names,
    'Coefficient': lasso_poly_c.coef_
})

#display the non-zero coefficients DataFrame in a sorted manner from Largest to sma
print("Non-zero Lasso Coefficients (alpha = 5) for Polynomial Features sorted from
nonzero_coef_df = coef_df[coef_df['Coefficient'] != 0]
display(nonzero_coef_df.sort_values(by='Coefficient', ascending=False))

# Filter out zero coefficients
nonzero_df = coef_df[coef_df['Coefficient'] != 0]

# Sort by absolute value for clearer visualization
nonzero_df = nonzero_df.reindex(nonzero_df['Coefficient'].abs().sort_values(ascending=False).index)

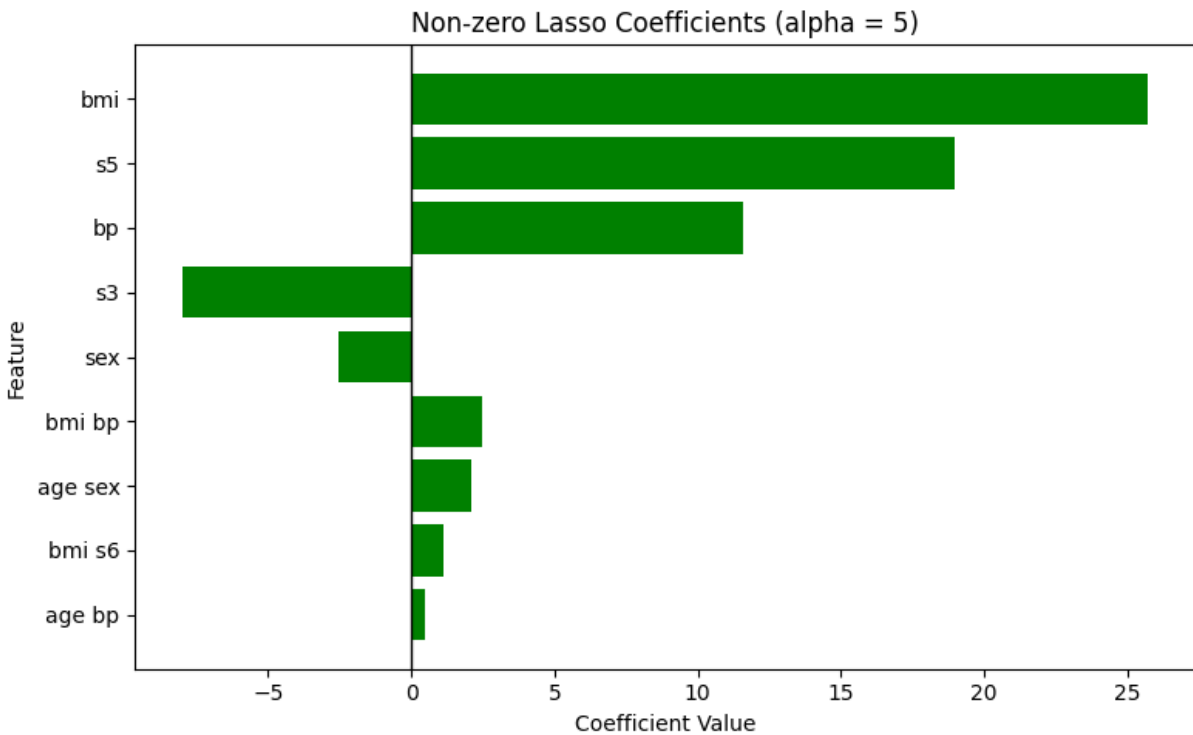
# Plot
plt.figure(figsize=(8,5))
plt.barh(nonzero_df['Feature'], nonzero_df['Coefficient'], color='green')
plt.xlabel('Coefficient Value')
plt.ylabel('Feature')
plt.title('Non-zero Lasso Coefficients (alpha = 5)')
plt.axvline(0, color='black', linewidth=1)
plt.tight_layout()
plt.show()

# showing intercept value
print(f'Lasso Intercept (alpha = 5) for Polynomial Features: {lasso_poly_c.intercept_}')

```

Non-zero Lasso Coefficients (alpha = 5) for Polynomial Features sorted from largest to smallest:

	Feature	Coefficient
2	bmi	25.690161
8	s5	18.941873
3	bp	11.581286
27	bmi bp	2.471126
10	age sex	2.074981
33	bmi s6	1.135663
12	age bp	0.500690
1	sex	-2.519322
6	s3	-7.954854



Lasso Intercept (alpha = 5) for Polynomial Features: 153.73654390934846

We have reduced the number of features by Lasso model. It is noted that the Lasso model is not used for its performance, but the feature reduction. Now, let's perform a **simple linear regression** model using the top 9 features.

In [532...]

```
# Performing linear regression using only the top 9 features identified by Lasso with
from scipy import stats

top_features = nonzero_coef_df['Feature'].tolist()[:9]
X_train_top = X_train_poly_s[:, [list(feature_names).index(f) for f in top_features]]
X_test_top = X_test_poly_s[:, [list(feature_names).index(f) for f in top_features]]
lr_top = LinearRegression().fit(X_train_top, y_train)
r2_lr_top = r2_score(y_test, lr_top.predict(X_test_top))
print(f"R² (Linear Regression with Top 9 Lasso Features): {r2_lr_top:.3f}")

# printing feature names of the top 9 features used with their coefficients sorted
print("====")
print("Top 9 features used in Linear Regression with their coefficients:")
for feature, coef in sorted(zip(top_features, lr_top.coef_), key=lambda x: x[1], reverse=True):
    print(f"{feature}: {coef:.3f}")
```

```
R2 (Linear Regression with Top 9 Lasso Features): 0.535
=====
Top 9 features used in Linear Regression with their coefficients:
bmi: 26.000
s5: 20.701
bp: 16.173
age sex: 6.811
bmi bp: 5.597
bmi s6: 3.945
age bp: 3.387
sex: -10.860
s3: -13.894
```

It can be observed that the model's fit has improved compared to the initial linear regression performed on the original features, which achieved an $R^2 = 0.453$.

After expanding the feature space using polynomial terms, applying **Lasso regularization** for feature selection, and then fitting a linear regression model on the reduced set, the performance increased to $R^2 = 0.535$.

This indicates that introducing nonlinear interactions and removing irrelevant features helped the model capture more variance in the target variable while maintaining simplicity.

Model Evaluation through Diagnostic Plots

After building a regression model, it is essential to evaluate its performance both **numerically** and **visually**.

Visual diagnostics help reveal issues such as bias, nonlinearity, heteroscedasticity, and non-normal residuals that numeric metrics alone might miss.

Below are the most important plots and their mathematical foundations:

1 Predicted vs. Actual Plot

This plot compares the **true values** (`y_test`) with the **predicted values** (`y_pred`).

$$\text{Prediction: } \hat{y}_i = f(\mathbf{x}_i)$$

A perfect model would satisfy:

$$\hat{y}_i = y_i$$

Points lying on the 45° diagonal line ($y = \hat{y}$) represent perfect predictions, while deviations from the line indicate underprediction or overprediction.

2 Residual Plot

The **residual** for each observation is defined as:

$$e_i = y_i - \hat{y}_i$$

Residuals are plotted against fitted (predicted) values.

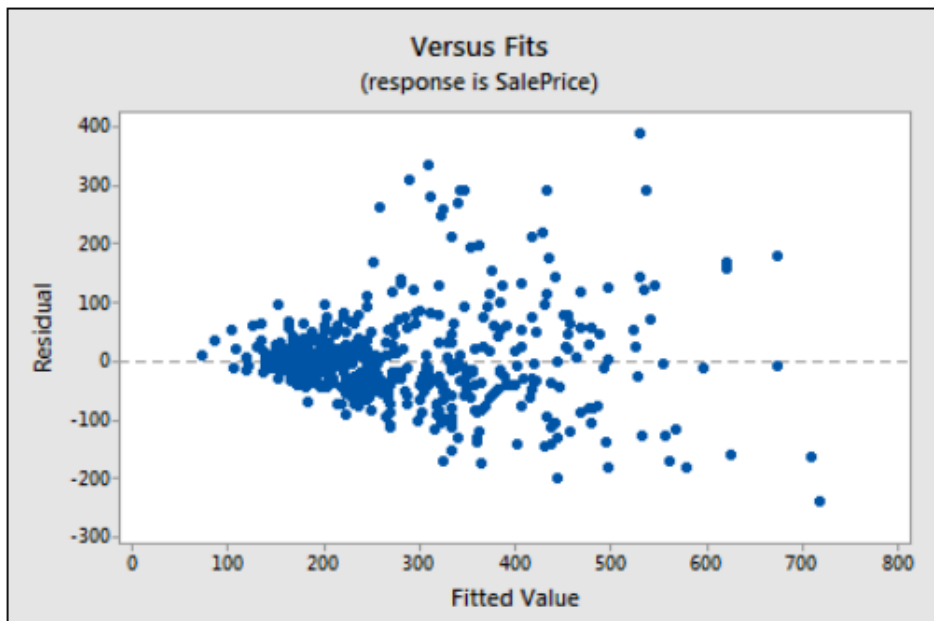
A good regression model should show residuals **randomly scattered around zero**:

$$\mathbb{E}[e_i] = 0, \quad \text{Var}(e_i) = \sigma^2$$

Patterns or systematic structures in the plot (e.g., curvature or funnels) suggest that the model's assumptions are violated —
for example, missing nonlinear trends or heteroscedasticity (non-constant variance).

Here is an example of bad regression model:

Image taken from [here](#):



This plot of residuals versus fits shows that the residual variance (vertical spread) increases as the fitted values (predicted values of sale price) increase. This violates the assumption of constant error variance [Ref]. In such case we need to refine our model or variables to improve the it.

3 Histogram of Residuals

The histogram (often with a KDE curve) shows how the **residuals are distributed**.

For a well-behaved regression model, residuals should approximately follow a **normal (Gaussian) distribution** centered at zero: This means most errors are small and positive/negative errors are equally likely.

A symmetric, bell-shaped histogram indicates that the model's errors are **unbiased and consistent**.

However:

- A **skewed** histogram suggests systematic overprediction or underprediction.

- **Heavy tails** (leptokurtic shape) indicate the presence of outliers or high-variance regions.
-

4 Q-Q Plot (Quantile–Quantile Plot)

To assess whether residuals follow a **normal distribution**, we standardize them:

$$e_i^* = \frac{e_i - \bar{e}}{s_e}$$

where \bar{e} is the mean of residuals and s_e is their standard deviation.

We then plot the **empirical quantiles** of e_i^* against the **theoretical quantiles** of a standard normal distribution:

$$q_i = \Phi^{-1} \left(\frac{i - 0.5}{n} \right)$$

where Φ^{-1} is the inverse CDF (percent-point function) of the standard normal distribution. If the residuals are normally distributed, the points should align closely with the diagonal reference line.

5 Why These Plots Matter

- The **Predicted vs. Actual plot** checks overall accuracy and systematic bias.
- The **Residual plot** verifies that errors are independent and randomly distributed.
- The **Histogram of Residuals** confirms that residuals are approximately normal and unbiased.
- The **Q-Q plot** provides a formal visual test for the normality assumption.

Together, these figures provide a comprehensive diagnostic check, confirming whether the model is **well-specified, unbiased, and generalizable**.

Now let's explore our model using the abovemnetioned figures.

```
In [531]: # Plot Predicted vs Actual and the residual plot for Linear Regression with Top 9 L
plt.figure(figsize=(10, 8))

# Predicted vs Actual
plt.subplot(2, 2, 1)
sns.scatterplot(x=y_test, y=lr_top.predict(X_test_top), color='purple', alpha=0.6,
                 palette='viridis')

# Perfect fit line (y = x)
plt.plot([y_test.min(), y_test.max()],
          [y_test.min(), y_test.max()],
          color='red', linestyle='--', linewidth=2, label='Ideal fit (y_pred = y_test)')

# Labels and title
plt.xlabel('Actual Values (y_test)', fontsize=10)
plt.ylabel('Predicted Values (y_pred)', fontsize=10)
```

```

plt.title('Linear Regression with Top 9 Features Obtained from Lasso\nPredicted vs.\n# Move R2 box slightly lower and right')
plt.text(y_test.min() + (y_test.max() - y_test.min())*0.001,
         y_test.max()*0.85,
         f"R2 = {r2_lr_top:.3f}",
         fontsize=12, color='black', bbox=dict(facecolor='white', alpha=0.8))
# Move Legend to top-right to avoid overlap
plt.legend(loc='upper left')
plt.tight_layout()

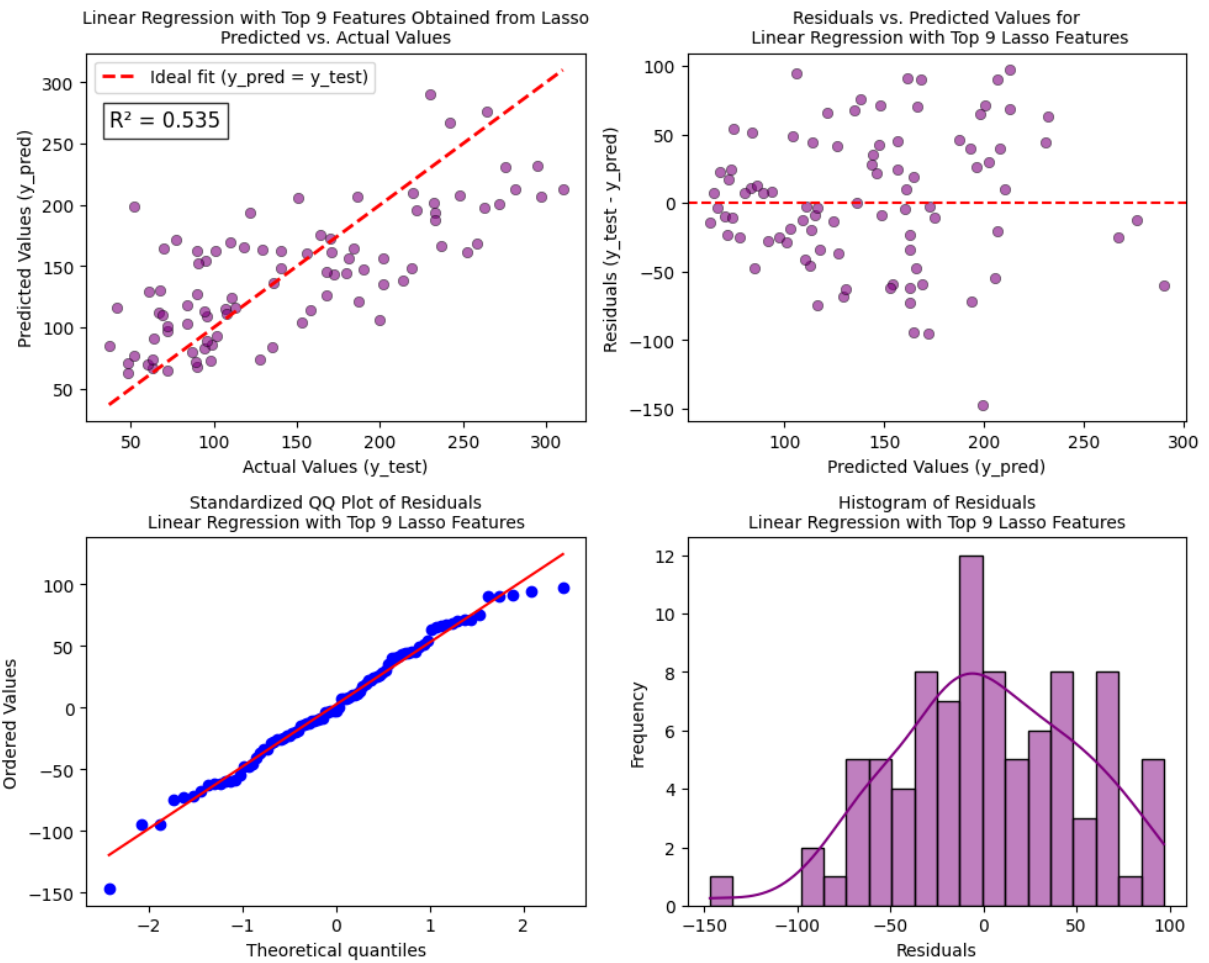
# Residuals vs. Predicted
plt.subplot(2, 2, 2)
sns.scatterplot(x=lr_top.predict(X_test_top), y=(y_test - lr_top.predict(X_test_top)))
plt.axhline(0, color='red', linestyle='--')
plt.xlabel('Predicted Values (y_pred)')
plt.ylabel('Residuals (y_test - y_pred)')
plt.title('Residuals vs. Predicted Values for\n Linear Regression with Top 9 Lasso')
plt.tight_layout()

# showing the equation of the linear regression model with top 9 features on the plot
equation_text_top = "y_pred = "
for feature, coef in zip(top_features, lr_top.coef_):
    equation_text_top += f"{coef:.3f}*{feature} + "
equation_text_top = equation_text_top[:-3] # Remove last " +
equation_text_top += f" + {lr_top.intercept_:.3f}"
plt.gcf().text(0.1, -0.1, "The equation of the multiple linear model is:\n" + equation_text_top)

# Plotting Standardized QQ plot and histogram of residuals for Linear Regression with Top 9 Lasso Features
# Standardized QQ plot with purple color
plt.subplot(2, 2, 3)
stats.probplot(y_test - lr_top.predict(X_test_top), dist="norm", plot=plt)
plt.title('Standardized QQ Plot of Residuals\nLinear Regression with Top 9 Lasso Features')

# Histogram of residuals
plt.subplot(2, 2, 4)
sns.histplot(y_test - lr_top.predict(X_test_top), bins=20, kde=True, color='purple')
plt.title('Histogram of Residuals\nLinear Regression with Top 9 Lasso Features', fontweight='bold')
plt.xlabel('Residuals')
plt.ylabel('Frequency')
plt.tight_layout()
plt.show()

```



The equation of the multiple linear model is:
 $y_{pred} = -10.860*sex + 26.000*bmi + 16.173*bp + -13.894*s3 + 20.701*s5 + 6.811*age_sex + 3.387*age_bp + 5.597*bmi_bp + 3.945*bmi_s6 + 153.737$

Interpretation of Model Diagnostics

The diagnostic plots above illustrate how well the linear regression model (trained on the top 9 Lasso-selected features) performs on the Diabetes dataset:

- **Predicted vs. Actual Plot:**

The points generally follow the red dashed line, indicating a moderate positive correlation between the predicted and actual values.

In this example, the **R² value** shown on the plot quantifies how well the model explains the variance in the target variable:

$R^2 = 1.0 \rightarrow$ perfect prediction

$R^2 = 0.0 \rightarrow$ no predictive power (random guess)

The coefficient of determination ($R^2 = 0.535$) suggests that the model explains about 53% of the variance in the target variable.

- **Residuals vs. Predicted Plot:**

The residuals appear randomly scattered around zero, showing no strong pattern.

This indicates that the model captures most of the systematic variation, though the slightly wider spread at higher predicted values hints at mild heteroscedasticity.

- **Q–Q Plot of Residuals:**

Most points lie close to the diagonal reference line, implying that the residuals are approximately normally distributed.

Minor deviations at the tails suggest a few outliers but no severe violation of normality.

- **Histogram of Residuals:**

The residuals form a roughly symmetric, bell-shaped distribution centered near zero, supporting the assumption of unbiased errors.

Overall, the diagnostic plots confirm that the linear regression model fits the data reasonably well, with normally distributed and mostly independent residuals, though there remains room for improvement in explaining variance.

Section 3. Conclusion

Given everything you have explored in this exercise (and what we discussed in the course) about **regularization**, **interaction terms**, **overfitting**, and **model generalizability**, what is your perspective on the **bias–variance trade-off** and the **interpretability trade-off** for this dataset?

Take a moment to reflect and collect your thoughts before revealing my answer below.

Conclusion (my version)

Across both experiments, **regularization consistently improves model performance**, confirming its role in **reducing variance** and **enhancing generalization**.

When we extend the feature space to include **interaction terms**, we observe a modest additional improvement in test-set R^2 , indicating that these nonlinear interactions capture a small portion of residual structure that the purely linear model could not explain.

However, as seen from both the performance metrics and the most influential predictors identified by Lasso ($\alpha = 5$), the qualitative story remains the same: the model's insights and key drivers of diabetes progression are largely unchanged.

This suggests that, for this dataset, the relationships between predictors and the target are **predominantly linear and additive**, and the marginal gains from including polynomial interactions may not justify the added complexity or overfitting risk for future data.

In practice, this highlights an important modeling principle:

Regularization helps control variance, while model expansion helps reduce bias; but the goal is not maximum complexity, it's meaningful simplicity.

After finalizing our model, we can use the residual plots to obtain an overall observation of our model performance.

The notebook of this assignment can be found at my: [GitHub Repository](#)