Interpretation

June 14, 2025

1 Predicting LOS of patients during their stay in the Intensive Care Unit in the MIMIC-III

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Link: GitHub

1.1 Objective

Accurately forecasting the Length of Stay (LOS) of patients in the Intensive Care Unit (ICU) is critical for resource allocation, clinical decision-making, and improving patient outcomes. This document outlines the development and optimization of a machine learning pipeline to predict ICU LOS using the MIMIC-III (Medical Information Mart for Intensive Care) dataset, a rich but challenging clinical database.

1.2 Challenges

- Skewed LOS distribution: Most patients stay 22 days, but a long tail of extended stays complicates modeling.
- Sparse, irregular time-series data: Clinical measurements (e.g., lab values, vitals) are recorded at varying frequencies.
- High-dimensional features: Thousands of potential predictors require careful selection to avoid overfitting

1.3 Why This Matters

- Clinical utility: Earlier, accurate LOS predictions can aid ICU bed management and discharge planning.
- Methodological insights: Demonstrates how to handle real-world clinical data constraints (missingness, skewness, high dimensionality).

2 Disease choice

We aimed to identify a disease that would provide:

- Sufficient sample size: Enough ICU cases for robust statistical analysis
- Clinical relevance: A condition with significant impact on ICU resource utilization

• Temporal predictability: Clear patterns of clinical measurements within the critical first 24 hours of ICU admission

After querying the MIMIC-III database for conditions meeting these criteria, we selected Pneumonia (ICD-9 code 482.83) based on:

- High prevalence in ICU populations
- Well-defined clinical markers for monitoring
- Consistent patterns of clinical deterioration that often require ICU admission

2.1 Loading relational MIMIC-III tables

We extracted and merged data from multiple MIMIC-III relational tables:

- PATIENTS table:
 - Demographic information (age, gender, ethnicity)
- ADMISSIONS table:
 - Hospital admission/discharge timestamps
 - Primary and secondary diagnoses
- ICUSTAYS table:
 - ICU admission/discharge timestamps
 - Length of stay calculation
- CHARTEVENTS table (first 24 hours):
 - Vital signs (heart rate, blood pressure, SpO)
 - Respiratory parameters (ventilator settings, FiO)
 - Neurological assessments (GCS scores)
- DIAGNOSES ICD table:
 - Confirmation of primary pneumonia diagnosis
- D ITEMS table:
 - Item description

```
[2]: import pandas as pd
import matplotlib.pyplot as plt

data_path = "./data_sets/"

df_patients= pd.read_csv(data_path + 'PATIENTS.csv.gz')
 df_admissions = pd.read_csv(data_path + 'ADMISSIONS.csv.gz')
 df_icustays = pd.read_csv(data_path + 'ICUSTAYS.csv.gz')
 df_diagnoses = pd.read_csv(data_path + 'DIAGNOSES_ICD.csv.gz', low_memory=False)
```

The integration of MIMIC-III's relational tables was carefully engineered to ensure data accuracy, temporal consistency, and clinical validity for our pneumonia cohort.

```
[3]: # Filter for Pneumonia (ICD-9 = 48283)
    df_diagnoses = df_diagnoses[df_diagnoses['ICD9_CODE'] ==__
     df_admissions_merged = pd.merge(
        df diagnoses,
        df_admissions[['SUBJECT_ID', 'HADM_ID', 'ADMITTIME', 'ETHNICITY']],
        on=['SUBJECT ID', 'HADM ID'],
        how='left'
    )
    print('Number of hospital admissions: ', df admissions merged.shape[0])
    df_icustays_merged = pd.merge(
        df_admissions_merged,
        df_icustays[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'INTIME', 'LOS']],
        on=['SUBJECT_ID', 'HADM_ID'],
        how='left'
    )
    print('Number of icu stays: ', df icustays merged.shape[0])
    df icustays patients merged = pd.merge(
        df_icustays_merged,
        df_patients[['SUBJECT_ID', 'GENDER', 'DOB']],
        on='SUBJECT_ID',
        how='left'
    df_icustays_patients_merged.head(3)
```

```
Number of hospital admissions: 264
Number of icu stays: 307
```

```
[3]:
       SUBJECT_ID HADM_ID ICD9_CODE
                                               ADMITTIME
                                                                      ETHNICITY \
                              48283 2146-08-29 01:18:00
                    178393
    0
              114
                                                          UNKNOWN/NOT SPECIFIED
    1
              339
                    112625
                              48283 2187-04-20 23:10:00 BLACK/AFRICAN AMERICAN
    2
              285
                   165312
                              48283 2152-09-21 22:47:00
                                                              HISPANIC OR LATINO
       ICUSTAY_ID
                                           LOS GENDER
                               INTIME
                                                                       DOB
                                                    M 2098-05-09 00:00:00
    0
           258626 2146-08-29 17:59:00
                                        1.8132
           221278 2187-04-20 23:10:50 31.7018
                                                    F 2120-07-17 00:00:00
    1
    2
           238023 2152-09-21 22:48:50 28.6880
                                                    M 2107-05-16 00:00:00
```

```
[4]: df_chart_events = pd.read_csv(data_path + 'd_pneumonia.csv', low_memory=False)
    # Merge df merged with CHARTEVENTS (left join to preserve all ICU stays)
    df_merged_with_chartevents = pd.merge(
        df_icustays_patients_merged,
        df_chart_events[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'ITEMID', |
      on=['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID'],
        how='left',
        indicator=True # Adds a column '_merge' to show match status
    )
    has_chartevents = df_merged_with_chartevents['_merge'] == 'both'
    no_chartevents = df_merged_with_chartevents['_merge'] == 'left_only'
    icu_with_data = df_merged_with_chartevents[has_chartevents]['ICUSTAY_ID'].
      →unique()
    icu_without_data = df_merged_with_chartevents[no_chartevents]['ICUSTAY_ID'].

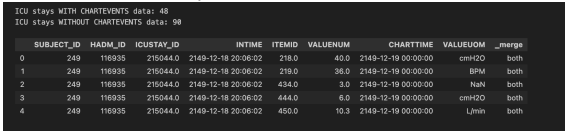
unique()
    print(f"ICU stays WITH CHARTEVENTS data: {len(icu_with_data)}")
    print(f"ICU stays WITHOUT CHARTEVENTS data: {len(icu_without_data)}")
    print(df_merged_with_chartevents.shape)
    df_merged_with_chartevents.head(2)
    ICU stays WITH CHARTEVENTS data: 145
    ICU stays WITHOUT CHARTEVENTS data: 162
    (986958, 14)
[4]:
       SUBJECT_ID HADM_ID ICD9_CODE
                                               ADMITTIME
                                                                      ETHNICITY \
                    178393
                               48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
    0
              114
              114
                    178393
                               48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
    1
       ICUSTAY_ID
                                INTIME
                                          LOS GENDER
                                                                      DOB
                                                                          \
           258626 2146-08-29 17:59:00 1.8132
                                                   M 2098-05-09 00:00:00
    0
                                                   M 2098-05-09 00:00:00
    1
           258626 2146-08-29 17:59:00 1.8132
       ITEMID VALUENUM
                                   CHARTTIME _merge
    0
        211.0
                   72.0 2146-08-30 10:00:00
                                              both
                  102.0 2146-08-30 10:00:00
        581.0
                                              both
```

The number icu stays which have chart events is of 145, this being enough to have a decent data frame.

An inner join in the last merge could've saved this step, but now we need to remove the icu stays without chart events.

```
[]: df_final = df_merged_with_chartevents[df_merged_with_chartevents['_merge'] ==__
      df final = df final.drop(columns=[' merge'])
    print(f"Shape of the original DataFrame: {df_merged_with_chartevents.shape}")
    print(f"Shape of the final DataFrame (with data): {df_final.shape}")
    df_final.head(3)
    Shape of the original DataFrame: (986958, 14)
    Shape of the final DataFrame (with data): (986796, 13)
[]:
       SUBJECT_ID
                   HADM_ID ICD9_CODE
                                                 ADMITTIME
                                                                        ETHNICITY
    0
               114
                    178393
                               48283
                                      2146-08-29 01:18:00
                                                           UNKNOWN/NOT SPECIFIED
               114
                    178393
                               48283
                                      2146-08-29 01:18:00
                                                           UNKNOWN/NOT SPECIFIED
    1
    2
                               48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
               114
                    178393
       ICUSTAY_ID
                                INTIME
                                           LOS GENDER
                                                                       DOB \
    0
            258626
                   2146-08-29 17:59:00
                                        1.8132
                                                    M 2098-05-09 00:00:00
           258626
                   2146-08-29 17:59:00
                                        1.8132
                                                    M 2098-05-09 00:00:00
    1
                   2146-08-29 17:59:00 1.8132
                                                    M 2098-05-09 00:00:00
            258626
              VALUENUM
       ITEMID
                                   CHARTTIME
    0
        211.0
                   72.0
                         2146-08-30 10:00:00
        581.0
                   102.0
                         2146-08-30 10:00:00
    1
        618.0
                    20.0
                         2146-08-30 10:00:00
```

All of this steps were also done for bronchitis, but we were left with only 48 icu stays with chart events which is by far insuficient as a dataframe to train and validate a model.



3 Statistical analysis and visualization of patients with Pneumonia

3.1 Dataframe cleaning preparation for the analysis

- Since the time window of 24 hours was chosen for prediction, all chart events past that time will be discarded
- There are 4 very far outliers with LOS above 60 days and they will be discarded since they are not enough to prove helpful for the model training
- Based on the fact that 75% of the admissions have a LOS under 21 days and 87.2% have a LOS under 30 days, entries with more than that will also be discarded

- Not all ICU stays have all the item values present
 - The two possible approaces of dealing with that missing data are imputing 0 in their place or leaving them as null
 - Since XGBoost, the model planned to be trained, works well with missing values, these will be left as null
 - In order to mark their missing as a relevant feature, a count of the number of their appearances will be added for each item, 0 meaning a missing value

We need to take out all the events that happen outside of the first 24h windows of a patient's stay to be able to visualize a patient 24h in the ICU and its relation with items and also the simplify the dataframe.

```
[6]: import seaborn as sns

df_final['INTIME'] = pd.to_datetime(df_final['INTIME'])

df_final['CHARTTIME'] = pd.to_datetime(df_final['CHARTTIME'])

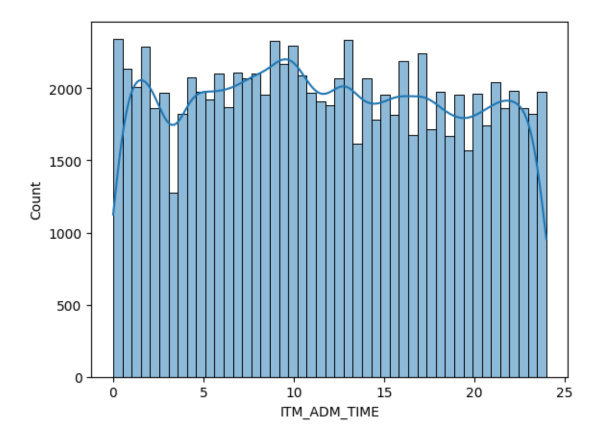
df_final['ITM_ADM_TIME'] = (df_final['CHARTTIME'] - df_final['INTIME']).dt.

$\int \text{total_seconds()} / 3600

df_final = df_final[(df_final['ITM_ADM_TIME'] <= 24) &_\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\text{\t
```

(92430, 14)

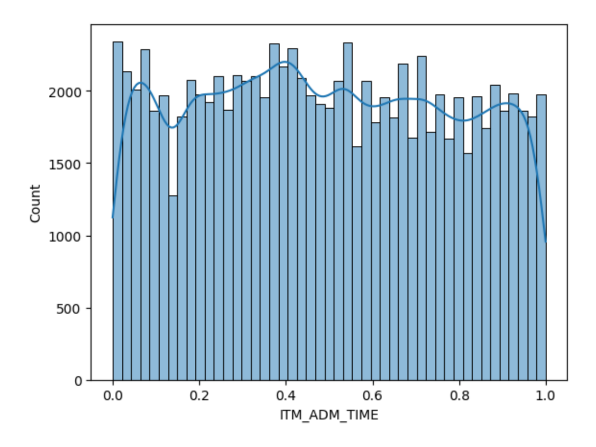
[6]: <Axes: xlabel='ITM_ADM_TIME', ylabel='Count'>



We normalize it to a time interval of (0, 1) to make any possible model easier to be trained and to be able to interpret results.

```
[7]: df_final['ITM_ADM_TIME'] = df_final['ITM_ADM_TIME'] / 24.0
sns.histplot(df_final['ITM_ADM_TIME'], kde=True)
```

[7]: <Axes: xlabel='ITM_ADM_TIME', ylabel='Count'>



3.2 Patients overview

Three patients were chosen with little, medium and high LOS in order to visualize different trends.

A sorted CSV of the ICU stays has been generated using an sql query which is saved to the csv norm_pneumonia.csv to be able to chose said three examples.

```
# Prepare LOWESS smoothed values
  x = filtered_limited["ITM_ADM_TIME"]
  y = filtered_limited["VALUENUM"]
  lowess_smoothed = lowess(y, x, frac=0.3, return_sorted=True)
  lowess_x = lowess_smoothed[:, 0]
  lowess_y = lowess_smoothed[:, 1]
  # Bootstrap for confidence interval
  bootstraps = []
  for _ in range(100):
      sample = filtered_limited.sample(frac=1, replace=True)
      smoothed = lowess(sample["VALUENUM"], sample["ITM_ADM_TIME"], frac=0.3,_
→return_sorted=True)
      bootstraps.append(np.interp(lowess_x, smoothed[:, 0], smoothed[:, 1]))
  bootstraps = np.array(bootstraps)
  ci_lower = np.percentile(bootstraps, 2.5, axis=0)
  ci_upper = np.percentile(bootstraps, 97.5, axis=0)
  # Plot
  plt.figure(figsize=(10, 7))
  sns.set(style="whitegrid")
  sns.scatterplot(
      data=filtered_limited,
      x="ITM_ADM_TIME",
      y="VALUENUM",
      hue="ITEMID",
      palette="Blues",
      alpha=0.8
  )
  # Add LOWESS trend line
  plt.plot(lowess_x, lowess_y, color="blue", label="Trend")
  # Add ribbon (confidence interval)
  plt.fill_between(lowess_x, ci_lower, ci_upper, color="blue", alpha=0.2, u
⇔label="95% CI")
  # Adjust legend
  plt.legend(title="ITEMID", loc='center left', bbox_to_anchor=(1, 0.5))
  plt.title(f"ICU_{icu_id}", fontsize=16)
  plt.xlabel("Time")
  plt.ylabel("Values")
  plt.tight_layout()
  plt.show()
```

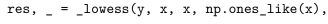
3.2.1 Small LOS

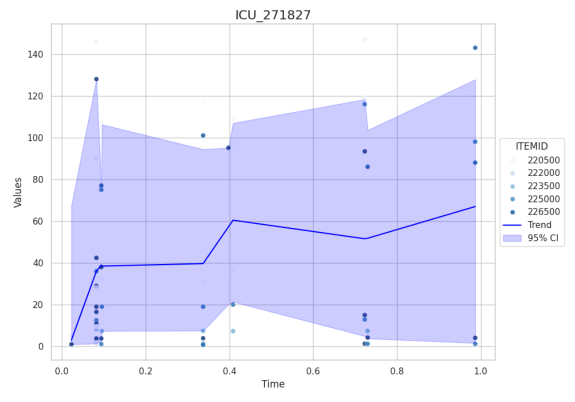
In this example we can see a slight increase of LOESS over time, but the trend is unstable do to sparse data and high variability.

Dense concentration of measurements in early stages suggest early implication of doctors that might suggest the shorter LOS.

```
[9]: import numpy as np plot_icu_data(icu_id=271827)
```

/home/cox/.local/lib/python3.10/site-packages/statsmodels/nonparametric/smoothers_lowess.py:226: RuntimeWarning: invalid value encountered in divide

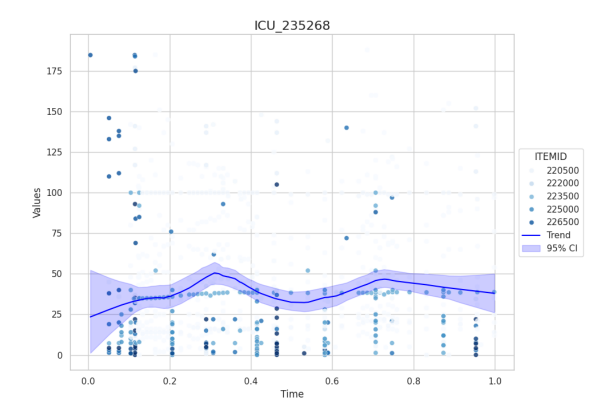




3.2.2 Medium LOS

Here, LOESS trend display a well formed curve and a consistent trajectory. Also, the evenly spread of items over the ICU stay suggest a stable and systematic mmonitoring over this time period.

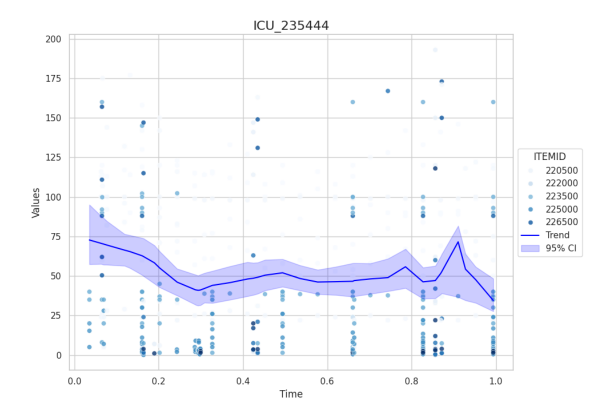
```
[10]: plot_icu_data(icu_id=235268)
```



3.2.3 High LOS

This example ilustrates a slight downward slope, possibly due to treatment effects and the spike at 0.9 suggests a check-up.

[11]: plot_icu_data(icu_id=235444)



4 Training and validation of a model to predict LOS

First of all our dataframe needs some processing for the model to be able to digest the data accordingly.

4.1 Data transformation

- The age of the patients is computed from the ICU admission time minus their birth date
 - Patients with ages greater than 89 years old are entered in the database to 300 in order to protect confidentiality
 - All the patients in this case will have their age moved to 91.4, the average of their group
- Categorical features of patients are encoded
 - Gender is label encoded, while ethnicity is encoded using one hot encoding for better model interpretation
- The distribution of the target variable, LOS, is right-skewed
 - In order to have that data better interpretable by a wider range of models, a log transformation will be applied on it ### Processing of DOB, INTIME and ADMITTIME To be able to make use of the dates, we transform them to pandas datetime format and create meaningful features out of them. These are the age, which is the difference between the time of admission in the ICU minus the date of birth, and the hospital time, which is the time of admission in the ICU minus the hospital admission time.

In this dataset, patients older than 89 years old have been mapped to 300 years old out of privacy reasons. We need their actual age so it has meaning for the future model. This is done by assigning the average of this age group, 91.4, to all of them.

```
[13]: df_final.loc[df_final['AGE'] > 89, 'AGE'] = 91.4
```

To make predictions on the LOS on patients at the 24 hour mark of their stay in ICU, the patients that stayed **more** than that, are not useful and will be discarded.

```
[14]: df_final = df_final[df_final['LOS'] > 1]
df_final.head(3)
```

```
[14]:
         SUBJECT_ID HADM_ID ICD9_CODE
                                                 ADMITTIME
                                                                         ETHNICITY
                      178393
                                 48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
      0
                114
                114
                      178393
                                 48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
      1
      2
                                 48283 2146-08-29 01:18:00 UNKNOWN/NOT SPECIFIED
                114
                      178393
                                            LOS GENDER
                                                              DOB ITEMID VALUENUM \
         ICUSTAY_ID
                                 INTIME
      0
             258626 2146-08-29 17:59:00
                                         1.8132
                                                     M 2098-05-09
                                                                     211.0
                                                                                72.0
      1
             258626 2146-08-29 17:59:00
                                         1.8132
                                                     M 2098-05-09
                                                                    581.0
                                                                               102.0
             258626 2146-08-29 17:59:00 1.8132
                                                     M 2098-05-09
                                                                    618.0
                                                                                20.0
                  CHARTTIME
                             ITM_ADM_TIME
                                                 AGE
                                                      HOSP_TIME
      0 2146-08-30 10:00:00
                                 0.667361 48.306639
                                                            0.0
      1 2146-08-30 10:00:00
                                 0.667361 48.306639
                                                            0.0
      2 2146-08-30 10:00:00
                                 0.667361 48.306639
                                                            0.0
```

4.1.1 Ethnicity and gender processing

Categorical features of patients are to be encoded.

Within this disease's sub data set there are different ethnicities, those that have less than 5 repre-

sentatives will be grouped together in an OTHER ethnicity.

```
[15]: unique_ethnicities = df_final['ETHNICITY'].unique()
      print(unique_ethnicities)
      ethnicity_counts = df_final['ETHNICITY'].value_counts()
      print(ethnicity_counts)
      rare_ethnicities = ['HISPANIC/LATINO - PUERTO RICAN', 'WHITE - RUSSIAN',
                           'HISPANIC/LATINO - DOMINICAN', 'ASIAN - CHINESE',
                           'UNABLE TO OBTAIN', 'PATIENT DECLINED TO ANSWER']
      df_final['ETHNICITY'] = df_final['ETHNICITY'].replace(rare_ethnicities, 'OTHER')
      ethnicity_counts = df_final['ETHNICITY'].value_counts()
      print(ethnicity_counts)
     ['UNKNOWN/NOT SPECIFIED' 'BLACK/AFRICAN AMERICAN' 'HISPANIC OR LATINO'
      'ASIAN' 'WHITE' 'OTHER' 'WHITE - RUSSIAN'
      'HISPANIC/LATINO - PUERTO RICAN' 'ASIAN - CHINESE' 'UNABLE TO OBTAIN'
      'PATIENT DECLINED TO ANSWER']
     ETHNICITY
     WHITE
                                        77730
     BLACK/AFRICAN AMERICAN
                                         5450
     UNKNOWN/NOT SPECIFIED
                                         2309
     HISPANIC OR LATINO
                                         1810
     ASIAN
                                         1268
     OTHER
                                          883
     WHITE - RUSSIAN
                                          813
     UNABLE TO OBTAIN
                                          753
     PATIENT DECLINED TO ANSWER
                                          400
     ASIAN - CHINESE
                                          376
     HISPANIC/LATINO - PUERTO RICAN
                                          215
     Name: count, dtype: int64
     ETHNICITY
     WHITE
                                77730
     BLACK/AFRICAN AMERICAN
                                 5450
     OTHER
                                 3440
     UNKNOWN/NOT SPECIFIED
                                 2309
     HISPANIC OR LATINO
                                 1810
     ASIAN
                                 1268
     Name: count, dtype: int64
```

The gender is encoded using a Label Encoder since in this data set it has 2 values.

But in the case of ethnicities, using a Label Encoder to encode these values would create an ordinal relationship between them, which will make them lose their meaning. Instead, the etnicities are one hot encoded, with the first column being dropped to prevent multicollinearity.

```
[16]: from sklearn.preprocessing import LabelEncoder
      gender_encoder = LabelEncoder()
      df_final['GENDER'] = gender_encoder.fit_transform(df_final['GENDER'])
      print("Gender Classes:", gender_encoder.classes_)
      df_final = pd.get_dummies(
          df_final,
          columns=['ETHNICITY'],
          prefix='ETH',
          drop_first=True
      df_final.head(3)
     Gender Classes: ['F' 'M']
「16]:
        SUBJECT_ID HADM_ID ICD9_CODE
                                                 ADMITTIME ICUSTAY_ID \
                114
                     178393
                                 48283 2146-08-29 01:18:00
                                                                258626
                114
                      178393
                                 48283 2146-08-29 01:18:00
                                                                258626
      1
      2
                114
                      178393
                                48283 2146-08-29 01:18:00
                                                                258626
                     INTIME
                               LOS GENDER
                                                   DOB ITEMID
                                                                VALUENUM \
      0 2146-08-29 17:59:00 1.8132
                                          1 2098-05-09
                                                         211.0
                                                                    72.0
      1 2146-08-29 17:59:00 1.8132
                                          1 2098-05-09
                                                         581.0
                                                                   102.0
      2 2146-08-29 17:59:00 1.8132
                                                         618.0
                                          1 2098-05-09
                                                                    20.0
                  CHARTTIME ITM_ADM_TIME
                                                 AGE HOSP_TIME \
      0 2146-08-30 10:00:00
                                 0.667361 48.306639
                                                            0.0
      1 2146-08-30 10:00:00
                                 0.667361 48.306639
                                                            0.0
      2 2146-08-30 10:00:00
                                0.667361 48.306639
                                                            0.0
        ETH_BLACK/AFRICAN AMERICAN ETH_HISPANIC OR LATINO ETH_OTHER \
      0
                              False
                                                      False
                                                                 False
      1
                              False
                                                      False
                                                                 False
      2
                              False
                                                      False
                                                                 False
        ETH_UNKNOWN/NOT SPECIFIED ETH_WHITE
      0
                              True
                                        False
                                        False
      1
                              True
      2
                              True
                                        False
[52]: df_icustays_patients_merged.describe().T
```

```
[52]:
                                                                            min
                   count
                                                     mean
      SUBJECT_ID
                   307.0
                                            33228.550489
                                                                          114.0
      HADM ID
                                           150245.521173
                   307.0
                                                                       100395.0
      ICUSTAY_ID
                   307.0
                                           252298.661238
                                                                       200387.0
                                                           2100-12-10 12:50:48
      INTIME
                     307
                          2150-10-01 05:11:23.130293248
      LOS
                   307.0
                                                15.420087
                                                                         0.0705
      DOB
                     307
                          2080-01-13 13:12:42.214983680
                                                           1805-12-07 00:00:00
      AGE
                   307.0
                                                65.235295
                                                                      19.350445
                                               25%
                                                                                50%
                                                                                     \
      SUBJECT_ID
                                                                           25474.0
                                          11215.0
      HADM_ID
                                         125328.0
                                                                          152895.0
      ICUSTAY_ID
                                         230449.5
                                                                          253214.0
                   2125-01-24 21:55:23.500000256
      INTIME
                                                    2150-04-22 17:40:51.000000512
      LOS
                                          4.42235
                                                                            11.9535
      DOB
                             2056-08-04 00:00:00
                                                               2088-08-09 00:00:00
      AGE
                                        57.278234
                                                                         67.956194
                                              75%
                                                                     max
                                                                                    std
      SUBJECT ID
                                          49267.5
                                                                 99715.0
                                                                          28758.748753
                                                                199845.0
      HADM ID
                                         174117.5
                                                                          27775.541971
      ICUSTAY ID
                                                                          28329.125649
                                         275776.0
                                                                299728.0
      INTIME
                   2174-08-08 07:02:47.000000512
                                                    2207-12-13 20:57:20
                                                                                    NaN
      LOS
                                           21.613
                                                                 84.0409
                                                                              13.732065
      DOB
                             2110-02-18 00:00:00
                                                    2173-02-23 00:00:00
                                                                                    NaN
      AGE
                                        76.819302
                                                                    91.4
                                                                              15.791862
```

- Based on the fact that 75% of the admissions have a LOS under 21 days and 87.2% have a LOS under 30 days, entries with more than that will also be discarded
- Not all ICU stays have all the item values present
 - The two possible approaces of dealing with that missing data are imputing 0 in their place or leaving them as null
 - Since XGBoost, the model planned to be trained, works well with missing values, these will be left as null
 - In order to mark their missing as a relevant feature, a count of the number of their appearances will be added for each item, 0 meaning a missing value

```
[17]: print('Initial size: ', df_final.shape[0])
df_final = df_final[df_final['LOS'] <= 30]
print('Current size: ', df_final.shape[0])</pre>
```

Initial size: 92007 Current size: 87861

Without removal of the outliers, the LOS distribution was right skewed, which would've suggested using the log of the LOS. But after the elimination, log becomes left skewed.

```
[18]: import matplotlib.pyplot as plt

df_final_log = df_icustays_patients_merged.copy()

df_final_log['LOG_LOS'] = np.log(df_final_log['LOS'])

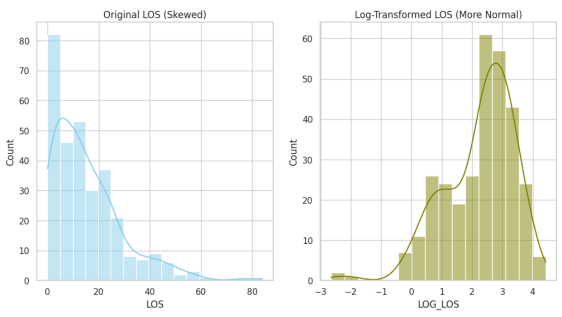
fig, axes = plt.subplots(1, 2, figsize=(12, 6))
 fig.suptitle('Comparison of LOS and Log-Transformed LOS Distributions')

sns.histplot(df_final_log['LOS'], kde=True, ax=axes[0], color='skyblue')
 axes[0].set_title('Original LOS (Skewed)')

sns.histplot(df_final_log['LOG_LOS'], kde=True, ax=axes[1], color='olive')
 axes[1].set_title('Log-Transformed LOS (More Normal)')

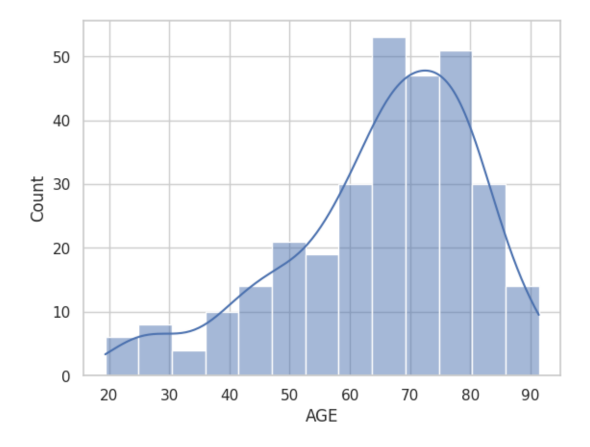
# plt.tight_layout()
 plt.show()
```

Comparison of LOS and Log-Transformed LOS Distributions



Age distribution looks good.

[19]: <Axes: xlabel='AGE', ylabel='Count'>



4.2 Feature Engineering

- The time-stamps of each chart event will be normalized as starting from the addmission time
- Given the time-stamped nature of the item entries, the item values for each ICU stay will be aggregated into the following 5 features per item:
 - average
 - standard deviation

- trend
- range
- count
- Since not all ICU stays have the same items in their respective events, only the items with more than 78% appearance will be selected as features
 - This criteria means selecting the first 32 items, multiplied by the number of 5 metrics for each, giving 160 features
 - This sorting is performed by an SQL query, that saves the results into items_appearance_pneumonia.csv ### Items aggregation We need to introduce the items into our dataframe, we do that by calculating a set of aggregate features for the top 32 most frequent items in our disease.

```
[20]:
              SUBJECT_ID
                          HADM_ID
                                    ICUSTAY_ID
                                                 HOSP_TIME
                                                                   AGE
                                                                        GENDER
                     114
                           178393
                                        258626
                                                  0.000000
                                                             48.306639
                                                                              1
      13145
                     285
                           165312
                                        238023
                                                  0.000000
                                                            45.353183
                                                                              1
      21486
                     605
                                        248569
                                                  0.000000
                                                            91.400000
                                                                              0
                           115545
      46380
                    3482
                           192399
                                        202786
                                                  0.002738
                                                             55.577687
                                                                              1
      48072
                    2090
                           138877
                                        256557
                                                  0.000000
                                                            76.605065
                                                                              1
             ETH BLACK/AFRICAN AMERICAN ETH HISPANIC OR LATINO
                                                                     ETH OTHER
      0
                                    False
                                                              False
                                                                          False
```

```
13145
                              False
                                                          True
                                                                     False
21486
                              False
                                                         False
                                                                     False
46380
                              False
                                                         False
                                                                     False
48072
                              False
                                                         False
                                                                     False
```

```
ETH_UNKNOWN/NOT SPECIFIED
                                    ETH_WHITE
                                                    LOS
0
                              True
                                        False
                                                 1.8132
13145
                             False
                                        False
                                                28.6880
21486
                             False
                                        False
                                                16.8598
46380
                             False
                                          True
                                                 4.9694
48072
                             False
                                          True
                                                10.0869
```

```
[21]: import pandas as pd import numpy as np

# The 'features_df' (for mapping names) is now a separate, optional argument.
```

```
def create_feature_matrix(df_events, df_static, item_ids, features_df=None,_
 →agg_list=None):
    Creates a feature matrix by combining static data with aggregated ____
 \hookrightarrow time-series features.
    This version correctly accepts a list of ITEMIDs and robustly flattens \Box
 \hookrightarrow MultiIndex columns.
    Args:
        df_events (pd.DataFrame): Time-series data with 'SUBJECT_ID', 'ITEMID', \( \)
 → 'VALUENUM', 'CHARTTIME'.
        df_static (pd.DataFrame): Static patient data with 'SUBJECT_ID'.
        item_ids (list): A Python list of ITEMIDs to use for feature generation.
        features df (pd.DataFrame, optional): For mapping ITEMID to readable ⊔
 ⇔names.
                                                  Must have 'ITEMID' and
 → 'item_name' columns. Defaults to None.
        agg\_list (list, optional): Aggregations to compute. Defaults to \sqcup
 →['mean', 'std', 'count', 'range', 'trend'].
    Returns:
        pd.DataFrame: Combined \ static + aggregated \ features, \ with \ NaN \ filled \ as_{\sqcup}
 \hookrightarrow 0.
    11 11 11
    if agg_list is None:
        agg_list = ['mean', 'std', 'count', 'range', 'trend']
    # 1. Filter relevant ITEMIDs from the provided list
    df filtered = df events[df events['ITEMID'].isin(item ids)].copy()
    df_filtered['CHARTTIME'] = pd.to_datetime(df_filtered['CHARTTIME'])
    # 2. Define aggregation calculations
    def calculate_aggregated_features(group):
        vals = group['VALUENUM']
        count = vals.count()
        features = {}
        for agg in agg_list:
            features[agg] = 0
        if count == 0:
            return pd.Series(features)
        if 'count' in agg_list: features['count'] = count
        if 'mean' in agg_list: features['mean'] = vals.mean()
        if 'std' in agg_list: features['std'] = vals.std() if count > 1 else 0
        if 'range' in agg_list: features['range'] = vals.max() - vals.min() if
 ⇔count > 1 else 0
        if 'trend' in agg_list and count > 1:
```

```
group = group.sort_values('CHARTTIME')
          time_in_hours = (group['CHARTTIME'] - group['CHARTTIME'].iloc[0]).
⇔dt.total_seconds() / 3600.0
          valid indices = time in hours.notna() & vals.notna()
          if valid indices.sum() > 1:
              slope = np.polyfit(time in hours[valid indices],
⇔vals[valid_indices], 1)[0]
              features['trend'] = slope if np.isfinite(slope) else 0
      return pd.Series(features)
  # 3. Compute aggregations
  item stats = (
      df_filtered.groupby(['SUBJECT_ID', 'ITEMID'])
      .apply(calculate_aggregated_features)
      .unstack(fill_value=0)
  )
  # 4. Flatten MultiIndex columns
  if isinstance(item_stats.columns, pd.MultiIndex):
      if features_df is not None:
          # Ensure data types match for mapping
          features_df['ITEMID'] = features_df['ITEMID'].astype(int)
          itemid_to_label = dict(zip(features_df['ITEMID'],__

¬features_df['item_name']))

          item stats.columns = [
              f"{agg}_{itemid_to_label.get(int(itemid), f'ITEM_{itemid}')}"
              for agg, itemid in item stats.columns
      else: # If no features_df is provided, use default names
          item_stats.columns = [
              f"{agg}_ITEM_{itemid}" for agg, itemid in item_stats.columns
          1
  item_stats = item_stats.reset_index()
  # 5. Merge with static data
  model_data = pd.merge(
      df static,
      item_stats,
      on='SUBJECT_ID',
      how='left'
  ).fillna(0)
  return model_data
```

The top features have been achieved by quering the dataset and exporting it to $items_appearance_pneumonia$.

```
[22]: top_features_df = pd.read_csv(data_path + 'items_appearance_pneumonia.csv')
top_features_df.head()
```

```
[22]:
         rank
                 ITEMID
                                 item_name VALUEUOM stay_count
            1 220645.0
                            Sodium (serum)
                                              mEq/L
                                                             135
      1
            2 220615.0
                                Creatinine
                                              mg/dL
                                                             135
      2
            3 220602.0 Chloride (serum)
                                              mEq/L
                                                             135
      3
            4 225624.0
                                       BUN
                                              mg/dL
                                                             135
            5 227443.0
                              HCO3 (serum)
                                              mEq/L
                                                             135
```

The number of items is: 32

For each of these items the mean, standard deviation, count, trend and range are calculated and inserted as features in the dataframe.

(126, 172)

/tmp/ipykernel_1368/3013210344.py:53: DeprecationWarning: DataFrameGroupBy.apply operated on the grouping columns. This behavior is deprecated, and in a future version of pandas the grouping columns will be excluded from the operation. Either pass `include_groups=False` to exclude the groupings or explicitly select the grouping columns after groupby to silence this warning.

.apply(calculate_aggregated_features)

```
[24]:
        SUBJECT_ID HADM_ID ICUSTAY_ID HOSP_TIME
                                                          AGE
                                                               GENDER
      0
                114
                     178393
                                 258626
                                               0.0 48.306639
                                                                    1
                                    ETH_HISPANIC OR LATINO ETH_OTHER \
        ETH_BLACK/AFRICAN AMERICAN
      0
                             False
                                                     False
                                                                False
        ETH_UNKNOWN/NOT SPECIFIED ... trend_Calcium non-ionized \
      0
                             True
                                                            0.0
        trend_Phosphorous trend_TCO2 (calc) Arterial trend_SpO2 Desat Limit \
```

```
0
                      0.0
                                                   0.0
                                                                          0.0
        trend_Anion_gap_trend_Potassium_(serum) trend_HCO3 (serum) \
                    0.0
      0
                                              0.0
        trend_Platelet Count trend_Prothrombin time trend_INR
                                                 0.0
                                                            0.0
                         0.0
      [1 rows x 172 columns]
     Also, for a better prediction we add the SOFA score, which is widely used in real life.
[25]: # Define SOFA-related ITEMIDs
      sofa_itemids = {
          'RESPIRATION': [220224, 223835], # PaO, FiO
          'COAGULATION': [828],
                                        # Platelets
          'LIVER': [225690],
                                           # Bilirubin
          'CARDIOVASCULAR': [220052],
                                            # MAP
          'CNS': [198],
                                            # GCS
          'RENAL': [220615]
                                # Creatinine
      }
      # 1. Flatten the list of ITEMIDs
      all_sofa_ids = [item for sublist in sofa_itemids.values() for item in sublist]
      # 2. Apply the filter directly
      df_sofa_components = df_chart_events[df_chart_events['ITEMID'].
       ⇔isin(all_sofa_ids)]
      print(f"Found {len(df_sofa_components)} SOFA-related measurements.")
      df_sofa_components.head(1)
     Found 67420 SOFA-related measurements.
[25]:
        ROW ID
                SUBJECT_ID HADM_ID ICUSTAY_ID ITEMID
                                                                    CHARTTIME \
      0 170754
                       1709
                             127294
                                          207018 220052 2118-01-04 12:15:00
                  STORETIME
                                CGID VALUE VALUENUM VALUEUOM WARNING ERROR \
      0 2118-01-04 14:22:00 19783.0 124.0
                                                124.0
                                                                     0.0
                                                                            0.0
                                                          mmHg
       RESULTSTATUS STOPPED
      0
                NaN
                        NaN
[26]: # Extract PaO and FiO (assuming FiO is in %, e.g., 50% = 0.5)
      df_pao2 = df_sofa_components[df_sofa_components['ITEMID'] == 220224]
      df_fio2 = df_sofa_components[df_sofa_components['ITEMID'] == 223835]
```

df pao2

```
# Merge and calculate PaO /FiO ratio
      df_respiration = pd.merge(
         df_pao2[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'CHARTTIME', 'VALUENUM']],
         df_fio2[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'CHARTTIME', 'VALUENUM']],
          on=['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'CHARTTIME'],
          suffixes=('_PAO2', '_FIO2')
      )
      df_respiration['PA02_FI02_RATIO'] = df_respiration['VALUENUM_PA02'] / __
      ⇔(df_respiration['VALUENUM_FIO2'] / 100)
      # Assign SOFA Respiration Subscore (0-4)
      df_respiration['SOFA_RESPIRATION'] = pd.cut(
         df respiration['PAO2_FIO2_RATIO'],
         bins=[0, 100, 200, 300, 400, float('inf')],
         labels=[4, 3, 2, 1, 0],
         right=False
      )
      df respiration.head(2)
                                                    CHARTTIME VALUENUM_PAO2 \
[26]:
        SUBJECT_ID HADM_ID ICUSTAY_ID
              3482
                     192399
                                  202786 2152-11-18 20:00:00
                                                                       106.0
      1
              12110
                      105928
                                  212945 2172-03-02 09:00:00
                                                                       154.0
        VALUENUM_FIO2 PAO2_FIO2_RATIO SOFA_RESPIRATION
      0
                  30.0
                             353.333333
      1
                  50.0
                             308.000000
                                                       1
[27]: | df_platelets = df_sofa_components[df_sofa_components['ITEMID'] == 828].copy()
      df_platelets['SOFA_COAGULATION'] = pd.cut(
         df_platelets['VALUENUM'],
         bins=[0, 20, 50, 100, 150, float('inf')],
         labels=[4, 3, 2, 1, 0],
         right=False
      df_platelets.head(3)
[27]:
                ROW ID
                       SUBJECT_ID HADM_ID ICUSTAY_ID ITEMID \
      941838 35644178
                               285
                                    165312
                                                 238023
                                                            828
      942333 35661979
                               285
                                     165312
                                                 238023
                                                            828
      942532 35667689
                               285
                                     165312
                                                 238023
                                                            828
                                                           CGID VALUE VALUENUM \
                       CHARTTIME
                                             STORETIME
      941838 2152-09-26 01:03:00 2152-09-26 02:08:00 15331.0 181.0
                                                                           181.0
```

```
942333 2152-10-12 01:42:00 2152-10-12 02:39:00 15331.0 260.0
                                                                           260.0
      942532 2152-10-18 02:29:00 2152-10-18 03:48:00
                                                        15331.0 333.0
                                                                           333.0
             VALUEUOM WARNING ERROR RESULTSTATUS
                                                     STOPPED SOFA_COAGULATION
      941838
                  NaN
                           NaN
                                  NaN
                                             Final NotStopd
                  NaN
                                                                            0
      942333
                           NaN
                                  NaN
                                             Final NotStopd
      942532
                  NaN
                           NaN
                                  NaN
                                             Final NotStopd
                                                                            0
[28]: df_bilirubin = df_sofa_components[df_sofa_components['ITEMID'] == 225690].copy()
      df bilirubin['SOFA LIVER'] = pd.cut(
          df bilirubin['VALUENUM'],
          bins=[0, 1.2, 2.0, 6.0, 12.0, float('inf')],
          labels=[0, 1, 2, 3, 4],
          right=False
      )
      df bilirubin.head(3)
[28]:
          ROW ID
                  SUBJECT ID HADM ID ICUSTAY ID ITEMID
                                                                      CHARTTIME \
                         1709
                                            207018 225690 2118-01-04 12:45:00
      33
          170787
                                127294
      269 171808
                         1709
                                127294
                                            207018 225690 2118-01-06 04:50:00
      333 171233
                         1709 127294
                                            207018 225690 2118-01-05 03:08:00
                     STORETIME
                                   CGID VALUE VALUENUM VALUEUOM WARNING ERROR \
          2118-01-04 14:39:00 20889.0
                                           1.9
                                                            mg/dL
                                                                              0.0
      33
                                                     1.9
                                                                       1.0
                                           2.5
                                                     2.5
      269 2118-01-06 05:27:00 20889.0
                                                            mg/dL
                                                                       1.0
                                                                              0.0
      333 2118-01-05 04:03:00 20889.0
                                           2.3
                                                     2.3
                                                            mg/dL
                                                                       1.0
                                                                              0.0
          RESULTSTATUS STOPPED SOFA_LIVER
      33
                  {\tt NaN}
                           NaN
                                        1
                  NaN
                                        2
      269
                           NaN
                  {\tt NaN}
                           NaN
                                        2
      333
[29]: df vasopressors = df sofa components[df sofa components['ITEMID'] == 220052].
       ⇔copy()
      # Assign SOFA Cardiovascular Subscore
      df_vasopressors['SOFA_CARDIOVASCULAR'] = pd.cut(
          df_vasopressors['VALUENUM'],
          bins=[0, 70, float('inf')],
          labels=[1, 0], # 1 if MAP < 70, else 0
          right=False
      )
      df_vasopressors['SOFA_CARDIOVASCULAR'].value_counts()
```

```
[29]: SOFA_CARDIOVASCULAR
           25909
      0
      1
            7634
      Name: count, dtype: int64
[30]: df_gcs = df_sofa_components[df_sofa_components['ITEMID'] == 198].copy()
      df_gcs['SOFA_CNS'] = pd.cut(
          df gcs['VALUENUM'],
          bins=[0, 6, 9, 12, 14, 16],
          labels=[4, 3, 2, 1, 0],
          right=False
      )
      df_gcs.head(2)
                        SUBJECT_ID HADM_ID ICUSTAY_ID
[30]:
                ROW ID
      940886
              35647251
                               285
                                     165312
                                                 238023
                                                            198
             35654790
                               285
                                     165312
                                                 238023
      940923
                                                            198
                        CHARTTIME
                                             STORETIME
                                                           CGID VALUE VALUENUM \
      940886 2152-09-28 03:00:00 2152-09-28 05:36:00
                                                        14997.0
                                                                   9.0
                                                                             9.0
      940923 2152-10-04 07:00:00 2152-10-04 06:45:00
                                                        17048.0
                                                                  11.0
                                                                             11.0
             VALUEUOM WARNING ERROR RESULTSTATUS
                                                     STOPPED SOFA_CNS
      940886
               points
                           NaN
                                  NaN
                                               NaN
                                                    NotStopd
                                                                     2
      940923
               points
                           NaN
                                  NaN
                                               {\tt NaN}
                                                    NotStopd
                                                                     2
     We aggregate all the SOFA scores into one.
[31]: df_creatinine = df_sofa_components[df_sofa_components['ITEMID'] == 220615].
       →copy()
      df_creatinine['SOFA_RENAL'] = pd.cut(
          df_creatinine['VALUENUM'],
          bins=[0, 1.2, 2.0, 3.5, 5.0, float('inf')],
          labels=[0, 1, 2, 3, 4],
          right=False
      )
      df_creatinine.head()
[31]:
           ROW_ID
                   SUBJECT_ID HADM_ID
                                        ICUSTAY_ID ITEMID
                                                                      CHARTTIME \
      23
           170777
                         1709
                                127294
                                            207018 220615 2118-01-04 12:45:00
      114 170225
                         1709
                                127294
                                            207018 220615 2118-01-03 05:23:00
      259 171798
                         1709
                                127294
                                            207018 220615 2118-01-06 04:50:00
                         1709
                                            207018 220615 2118-01-05 03:08:00
      323 171223
                                127294
      553 173781
                         1709
                                127294
                                            207018 220615 2118-01-10 22:59:00
```

```
CGID VALUE VALUENUM VALUEUOM WARNING ERROR \
                     STORETIME
                                                                             0.0
      23
          2118-01-04 14:39:00 20889.0
                                          7.1
                                                    7.1
                                                           mg/dL
                                                                      1.0
                                           6.6
                                                    6.6
      114 2118-01-03 07:01:00 20889.0
                                                                      1.0
                                                                             0.0
                                                           mg/dL
      259 2118-01-06 05:27:00 20889.0
                                          4.8
                                                    4.8
                                                           mg/dL
                                                                      1.0
                                                                             0.0
      323 2118-01-05 04:03:00 20889.0
                                          6.0
                                                    6.0
                                                           mg/dL
                                                                      1.0
                                                                             0.0
      553 2118-01-10 23:55:00 20889.0
                                          6.7
                                                    6.7
                                                           mg/dL
                                                                      1.0
                                                                             0.0
         RESULTSTATUS STOPPED SOFA RENAL
      23
                  {\tt NaN}
                          NaN
      114
                  NaN
                          NaN
                                       4
                          NaN
      259
                  NaN
                                       3
      323
                  {\tt NaN}
                          NaN
      553
                  NaN
                          NaN
[32]: # Merge all subscores
      df_sofa_scores = pd.concat([
         df_respiration[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'SOFA_RESPIRATION']],
         df_platelets[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'SOFA_COAGULATION']],
         df_bilirubin[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'SOFA_LIVER']],
         df_vasopressors[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', "
       df_gcs[['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID', 'SOFA_CNS']],
         df creatinine[['SUBJECT ID', 'HADM ID', 'ICUSTAY ID', 'SOFA RENAL']]
      ])
      df_sofa_scores = df_sofa_scores.fillna(0)
      df_sofa_scores.head(2)
[32]:
        SUBJECT_ID HADM_ID ICUSTAY_ID SOFA_RESPIRATION SOFA_COAGULATION \
                                  202786
              3482
                     192399
      0
             12110
                                 212945
                                                       1
                                                                        0
      1
                     105928
       SOFA_LIVER SOFA_CARDIOVASCULAR SOFA_CNS SOFA_RENAL
      0
                0
                                    0
                                             0
                                                        0
                0
                                    0
                                             0
                                                        0
      1
[33]: # 1. First ensure each component has only one score per ICU stay by taking the
      →worst (max) score
      sofa_components = ['SOFA_RESPIRATION', 'SOFA_COAGULATION', 'SOFA_LIVER',
                         'SOFA_CARDIOVASCULAR', 'SOFA_CNS', 'SOFA_RENAL']
      df_sofa_scores_aggregated = df_sofa_scores.groupby(
          ['SUBJECT ID', 'HADM ID', 'ICUSTAY ID']
      )[sofa_components].max().reset_index()
      # 2. Verify uniqueness
      print(f"Before aggregation: {len(df_sofa_scores)} records")
```

```
print(f"After aggregation: {len(df_sofa_scores_aggregated)} unique ICU stays")
      print("Duplicate check:", df_sofa_scores_aggregated.
       duplicated(subset=['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID']).any())
      # 3. Now merge with main data
      df model full with sofa = pd.merge(
          left=model data full,
          right=df_sofa_scores_aggregated,
          how='left',
          on=['SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID'],
          validate='one_to_one' # This will now pass
      )
      # 4. Calculate total SOFA (handle missing components as 0)
      df_model_full_with_sofa['total_sofa'] =__
       df_model_full_with_sofa[sofa_components].fillna(0).sum(axis=1)
     Before aggregation: 37829 records
     After aggregation: 146 unique ICU stays
     Duplicate check: False
[34]: df model full with sofa = df model full with sofa.drop(columns=sofa components)
      df_model_full_with_sofa.head()
[34]:
         SUBJECT_ID HADM_ID ICUSTAY_ID HOSP_TIME
                                                           AGE GENDER
      0
                114
                      178393
                                  258626
                                           0.000000 48.306639
                                                                      1
      1
                285
                      165312
                                  238023
                                           0.000000 45.353183
                                                                      1
                                                                      0
      2
                605
                                  248569
                                           0.000000 91.400000
                      115545
      3
               3482
                                                                      1
                      192399
                                  202786
                                           0.002738 55.577687
      4
               2090
                      138877
                                  256557
                                           0.000000 76.605065
                                                                      1
         ETH_BLACK/AFRICAN AMERICAN ETH_HISPANIC OR LATINO ETH_OTHER \
      0
                              False
                                                      False
                                                                  False
      1
                              False
                                                       True
                                                                 False
      2
                              False
                                                      False
                                                                 False
      3
                                                      False
                                                                 False
                              False
      4
                              False
                                                      False
                                                                 False
         ETH_UNKNOWN/NOT SPECIFIED ... trend_Phosphorous
      0
                              True ...
                                                0.000000
      1
                             False ...
                                                0.000000
      2
                             False ...
                                                0.089292
      3
                             False ...
                                                0.000000
      4
                             False ...
                                               -0.021077
         trend_TCO2 (calc) Arterial trend_SpO2 Desat Limit trend_Anion gap \
                                               0.000000e+00
      0
                           0.000000
                                                                     0.000000
```

```
1
                      0.000000
                                           0.000000e+00
                                                                  0.000000
2
                     -0.073447
                                          -7.002831e-16
                                                                  0.137920
3
                      0.243856
                                          -5.129622e-02
                                                                  0.000000
4
                     -0.016348
                                           1.739162e-15
                                                                  0.075682
   trend_Potassium (serum)
                             trend_HCO3 (serum)
                                                   trend_Platelet Count
0
                   0.000000
                                        0.000000
                                                               0.000000
1
                   0.000000
                                        0.000000
                                                               0.000000
2
                  -0.124548
                                        0.433000
                                                               0.082418
3
                   0.000000
                                        0.000000
                                                               0.000000
4
                  -0.031053
                                       -0.258931
                                                              -1.545801
   trend Prothrombin time
                            trend INR total sofa
0
                  0.000000
                             0.000000
                  0.000000
                             0.00000
                                                  0
1
2
                  0.000000
                             0.000000
                                                  3
3
                  0.000000
                             0.000000
                                                  0
4
                  0.055427
                                                  4
                             0.013857
```

[5 rows x 173 columns]

- 160 features, plus the encoded categorical ones, for each entry being too many for such a small number of entries (about 140), careful feature selection is necessary
- The filtering of the features will be performed in three steps (proven useful by iterative experiments):
 - 1. Filtering out the features with a variance lower than 1%
 - 2. Building a *correlation matrix* on the remaining features and determine the pairs with correlation higher than 0.9
 - Out of these pairs, only one of them is worth keeping, so the other one is dropped
 - 3. Training a dummy XGBoost model in order to extract the permutation importance of the features
 - Drop the features with a contribution to the RMS less than 1%

4.2.1 Dropping features with less than 1% variance

Said features that do not change are not usefull in our training.

```
selector.fit(features_df)
      kept_cols = features_df.columns[selector.get_support()]
      print(f"Original number of features: {len(feature_cols)}")
      print(f"Number of features kept: {len(kept_cols)}")
      final cols to keep = non numerical feature cols + kept cols.tolist()
      df_selected = model_data_full[final_cols_to_keep]
      print("\nShape of the new DataFrame:", df_selected.shape)
      print("\nFirst 5 rows of the new DataFrame with selected features:")
      df_selected.head(3)
     Original number of features: 169
     Number of features kept: 154
     Shape of the new DataFrame: (126, 158)
     First 5 rows of the new DataFrame with selected features:
[35]:
         SUBJECT ID HADM ID ICUSTAY ID
                                              LOS
                                                         AGE GENDER \
      0
                114
                      178393
                                  258626
                                           1.8132 48.306639
                                                                    1
                285
                                  238023 28.6880 45.353183
                                                                    1
      1
                      165312
      2
                                  248569 16.8598 91.400000
                605
                      115545
                                                                    0
         ETH_BLACK/AFRICAN AMERICAN ETH_HISPANIC OR LATINO ETH_OTHER \
      0
                              False
                                                      False
                                                                 False
                              False
                                                       True
                                                                 False
      1
      2
                                                                 False
                              False
                                                      False
         ETH_UNKNOWN/NOT SPECIFIED ... trend_Resp Alarm - High \
      0
                              True ...
                                                  0.000000e+00
                                                  0.000000e+00
      1
                             False ...
      2
                                                 -4.747817e-16
                             False ...
         trend_Arterial Base Excess trend_BUN trend_TCO2 (calc) Arterial \
      0
                           0.000000
                                      0.000000
                                                                  0.000000
                                      0.000000
                                                                  0.000000
      1
                           0.000000
      2
                          -0.118072
                                      1.967882
                                                                 -0.073447
         trend_Sp02 Desat Limit trend_Anion gap trend_HCO3 (serum) \
                   0.000000e+00
                                         0.00000
                                                               0.000
      0
      1
                   0.000000e+00
                                         0.00000
                                                               0.000
                                                               0.433
      2
                  -7.002831e-16
                                         0.13792
```

features_df = model_data_full[feature_cols]

```
trend_Platelet Count trend_Prothrombin time total_sofa
      0
                     0.000000
                                                  0.0
                                                  0.0
      1
                     0.000000
                                                                 0
                                                  0.0
                                                                 3
                     0.082418
      [3 rows x 158 columns]
[36]: dropped_cols_mask = ~selector.get_support()
      dropped_cols = features_df.columns[dropped_cols_mask]
      # Dropped columns
      print(len(dropped cols))
      dropped_cols
     15
[36]: Index(['HOSP_TIME', 'std_PH (Arterial)', 'range_PH (Arterial)',
             'trend_Hemoglobin', 'trend_02 saturation pulseoxymetry',
             'trend_Creatinine', 'trend_Magnesium',
             'trend_02 Saturation Pulseoxymetry Alarm - High',
             'trend_02 Saturation Pulseoxymetry Alarm - Low', 'trend_PH (Arterial)',
             'trend_Resp Alarm - Low', 'trend_Calcium non-ionized',
             'trend_Phosphorous', 'trend_Potassium (serum)', 'trend_INR'],
```

Droppin features with very low variance results in dropping 15 features.

4.2.2 Dropping highly correlated features

dtype='object')

By dropping highly correlated features, we keep item value importance and greatly decrease multicollinearity.

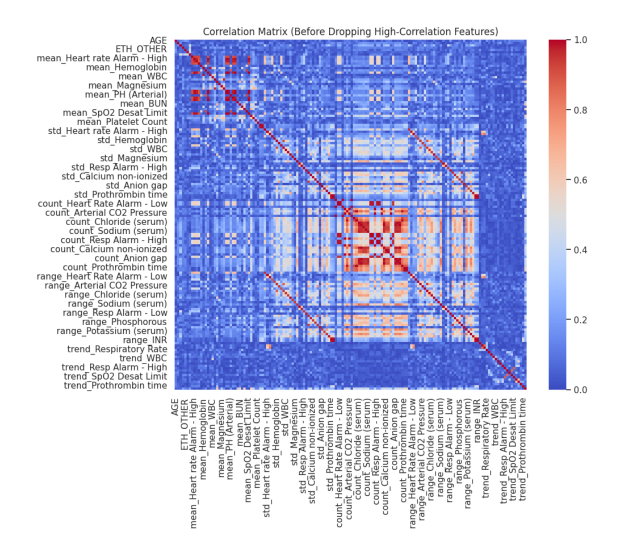
```
[37]: X = df_selected.drop(columns=non_numerical_feature_cols)

corr_matrix = X.corr().abs()

print("Generating heatmap for features before correlation-based removal...")

plt.figure(figsize=(10, 8))
    sns.heatmap(corr_matrix, cmap='coolwarm', annot=False)
    plt.title('Correlation Matrix (Before Dropping High-Correlation Features)')
    plt.show()
```

Generating heatmap for features before correlation-based removal...



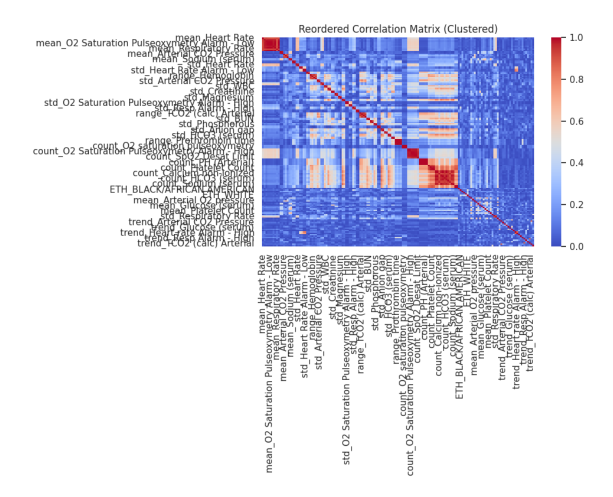
We tried to debug the correlation matrix because we thought it would eliminate way too many features. But, it turns out clusters are bigger than 2, so for example in a cluster of 10, we eliminate 9 features resulting in only 1 remaining.

```
[39]: import networkx as nx
      upper = corr matrix.where(np.triu(np.ones(corr matrix.shape), k=1).astype(bool))
      # Step 3: Extract highly correlated feature pairs (threshold > 0.9)
      high_corr_pairs = upper.stack()[upper.stack() > 0.9]
      print(f"Number of high-correlation feature pairs (corr > 0.9):
       →{len(high_corr_pairs)}")
      df_reduced = df_selected.drop(columns=['HADM_ID', 'ICUSTAY_ID', 'LOS', |
       ⇔'SUBJECT_ID'])
      # Step 4: Build graph and find connected components (clusters)
      G = nx.Graph()
      G.add_edges_from(high_corr_pairs.index.tolist())
      clusters = list(nx.connected_components(G))
      print(f"Number of correlated feature clusters: {len(clusters)}")
      # Step 5: Debug cluster sizes
      for i, group in enumerate(clusters):
          print(f"Cluster {i+1}: size = {len(group)} → dropping {len(group) - 1}")
      # Step 6: Compute features to drop (keep first in each group)
      to_drop_groups = [list(group)[1:] for group in clusters if len(group) > 1]
      to_drop = [item for sublist in to_drop_groups for item in sublist]
      print(f"Total number of features to drop: {len(to_drop)}")
      print("Features being dropped:", to_drop)
      # Step 7: Drop the features
      df_reduced = df_reduced.drop(columns=to_drop)
      # Step 8: Reorder features for visual inspection
      ordered_features = [feat for group in clusters for feat in sorted(group)]
      remaining features = [f for f in df reduced.columns if f not in_
       ⇔ordered_features]
      ordered_features += sorted(remaining_features)
      reordered_corr = corr_matrix.loc[ordered_features, ordered_features]
      # Step 9: Plot the reordered correlation matrix
      plt.figure(figsize=(10, 8))
      sns.heatmap(reordered_corr, cmap='coolwarm', annot=False)
      plt.title("Reordered Correlation Matrix (Clustered)")
```

```
plt.show()
Number of high-correlation feature pairs (corr > 0.9): 171
Number of correlated feature clusters: 38
Cluster 1: size = 10 → dropping 9
Cluster 2: size = 2 → dropping 1
Cluster 3: size = 3 → dropping 2
Cluster 4: size = 2 → dropping 1
Cluster 5: size = 2 → dropping 1
Cluster 6: size = 2 → dropping 1
Cluster 7: size = 2 → dropping 1
Cluster 8: size = 2 → dropping 1
Cluster 9: size = 2 \rightarrow \text{dropping } 1
Cluster 10: size = 4 → dropping 3
Cluster 11: size = 2 → dropping 1
Cluster 12: size = 2 → dropping 1
Cluster 13: size = 2 → dropping 1
Cluster 14: size = 2 → dropping 1
Cluster 15: size = 2 → dropping 1
Cluster 16: size = 2 → dropping 1
Cluster 17: size = 2 → dropping 1
Cluster 18: size = 2 → dropping 1
Cluster 19: size = 2 \rightarrow \text{dropping } 1
Cluster 20: size = 2 → dropping 1
Cluster 21: size = 2 \rightarrow \text{dropping } 1
Cluster 22: size = 2 → dropping 1
Cluster 23: size = 4 \rightarrow \text{dropping } 3
Cluster 24: size = 2 → dropping 1
Cluster 25: size = 2 → dropping 1
Cluster 26: size = 2 → dropping 1
Cluster 27: size = 2 → dropping 1
Cluster 28: size = 2 → dropping 1
Cluster 29: size = 2 → dropping 1
Cluster 30: size = 2 → dropping 1
Cluster 31: size = 2 \rightarrow \text{dropping } 1
Cluster 32: size = 4 → dropping 3
Cluster 33: size = 3 → dropping 2
Cluster 34: size = 7 → dropping 6
Cluster 35: size = 5 → dropping 4
Cluster 36: size = 4 → dropping 3
Cluster 37: size = 11 → dropping 10
Cluster 38: size = 2 → dropping 1
Total number of features to drop: 73
Features being dropped: ['mean_Resp Alarm - Low', 'mean_Sp02 Desat Limit',
'mean_02 Saturation Pulseoxymetry Alarm - High', 'mean_02 Saturation
Pulseoxymetry Alarm - Low', 'mean_02 saturation pulseoxymetry', 'mean_Heart
```

plt.tight_layout()

Rate', 'mean_Heart rate Alarm - High', 'mean_Respiratory Rate', 'mean_Heart Rate Alarm - Low', 'mean_Hematocrit (serum)', 'mean_Arterial CO2 Pressure', 'mean_PH (Arterial)', 'mean_Sodium (serum)', 'mean_Prothrombin time', 'std_Heart Rate', 'std_Heart rate Alarm - High', 'range_Heart Rate Alarm - Low', 'std_Arterial 02 pressure', 'std Hematocrit (serum)', 'std Hemoglobin', 'range Hematocrit (serum)', 'range_Arterial CO2 Pressure', 'range_O2 saturation pulseoxymetry', 'std_WBC', 'range_Chloride (serum)', 'range_Creatinine', 'std_Glucose (serum)', 'range_Magnesium', 'range_Sodium (serum)', 'std_02 Saturation Pulseoxymetry Alarm - High', 'range O2 Saturation Pulseoxymetry Alarm - Low', 'range Resp Alarm - High', 'std_Resp Alarm - Low', 'std_TCO2 (calc) Arterial', 'range_Arterial Base Excess', 'range_TCO2 (calc) Arterial', 'range BUN', 'std_Calcium non-ionized', 'std_Phosphorous', 'range_Sp02 Desat Limit', 'range_Anion gap', 'std_Potassium (serum)', 'std_HCO3 (serum)', 'range_Platelet Count', 'std_INR', 'range_Prothrombin time', 'std_Prothrombin time', 'count_Heart Rate', 'count_02 saturation pulseoxymetry', 'count_02 Saturation Pulseoxymetry Alarm - High', 'count Resp Alarm - Low', 'count Heart Rate Alarm -Low', 'count_Resp Alarm - High', 'count_02 Saturation Pulseoxymetry Alarm -Low', 'count_Sp02 Desat Limit', 'count_Arterial CO2 Pressure', 'count_Arterial O2 pressure', 'count_PH (Arterial)', 'count_TCO2 (calc) Arterial', 'count_Hematocrit (serum)', 'count_Platelet Count', 'count_Hemoglobin', 'count Anion gap', 'count Creatinine', 'count Phosphorous', 'count Sodium (serum)', 'count Chloride (serum)', 'count Magnesium', 'count BUN', 'count_Potassium (serum)', 'count_Glucose (serum)', 'count_HCO3 (serum)', 'count_INR']



Elimination by correlation, resulting 73 features eliminated.

ICUSTAY_ID

[40]:

SUBJECT_ID HADM_ID

178393

114

LOS

258626 1.8132 48.306639

AGE

GENDER

1

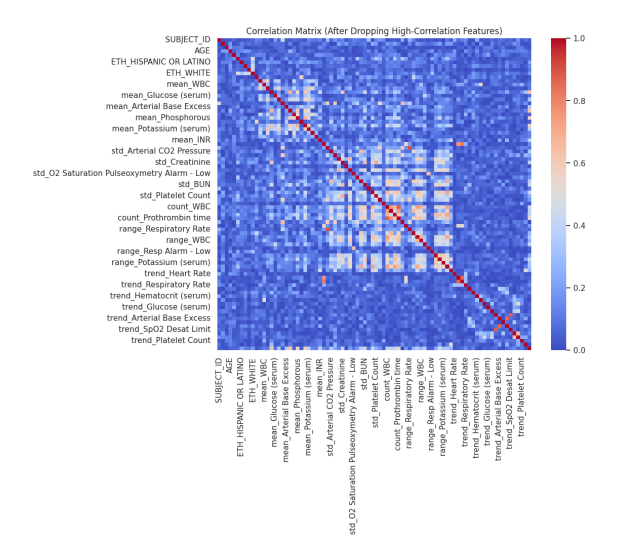
```
ETH HISPANIC OR LATINO
                                                         ETH_OTHER
  ETH_BLACK/AFRICAN AMERICAN
0
                        False
                                                 False
                                                             False
  ETH_UNKNOWN/NOT SPECIFIED
                                  trend_Resp Alarm - High
0
                                                       0.0
                        True
  trend_Arterial Base Excess
                                trend_BUN
                                           trend_TCO2 (calc) Arterial
0
                                      0.0
                           0.0
                                                                   0.0
  trend_Sp02 Desat Limit trend_Anion gap
                                             trend HCO3 (serum)
0
                      0.0
                                        0.0
                                                             0.0
   trend_Platelet Count
                         trend_Prothrombin time
0
                    0.0
                                             0.0
```

[1 rows x 85 columns]

Steps of eliminating the highly correlated features: - Isolate Features (X): We start with df variance selected, the DataFrame that has already been filtered for low variance. We create X which contains only the feature columns from this set. - Correlation Matrix: We compute the correlation matrix for X and take the absolute value, as we're interested in the strength of the correlation, not its direction (positive or negative). - Upper Triangle: The line np.triu(...) creates a mask for the upper triangle of the matrix (everything above the main diagonal). We do this because a correlation matrix is symmetrical (corr(A,B) is the same as corr(B,A)), and we only need to check each pair of features once to avoid redundancy. - Find Columns to Drop: We iterate through the columns of our upper-triangle view. If any value in a column is greater than our threshold (0.9), we add that column's name to our to drop list. This effectively keeps one feature from each highly correlated pair and flags the other for removal. - Create Final DataFrame: We drop the columns in the to drop list from df variance selected to produce our final, cleaned dataset df final selected.

```
[41]: final features = df final selected elimcorr.drop(columns='LOS')
      # Calculate the new correlation matrix
      final_corr_matrix = final_features.corr().abs()
      print("\nGenerating heatmap for the final set of features...")
      plt.figure(figsize=(10, 8))
      sns.heatmap(final_corr_matrix, cmap='coolwarm', annot=False, vmin=0, vmax=1)
      plt.title('Correlation Matrix (After Dropping High-Correlation Features)')
      plt.show()
```

Generating heatmap for the final set of features...



4.2.3 Permutation importance

This filter will discard features that contribute less than 1% to the RMSE.

```
Parameters:
   - df: DataFrame containing features and target
   - target_column: Name of the target column
  - group_column: Name of the column used for grouping (optional)
  - threshold: Minimum relative importance to keep a feature (default: 0.01)
   - n_splits: Number of splits for GroupKFold (default: 5)
  - n_repeats: Number of repeats for permutation importance (default: 5)
  Returns:
   - DataFrame with selected features and target column
  # Prepare data
  y = df[target_column]
  X = df.drop(columns=[target_column, 'HADM_ID', 'SUBJECT_ID', 'ICUSTAY_ID', L

  'LOS'])
  groups = df[group_column] if group_column else np.arange(len(df))
  # Initialize model and cross-validator
  model = XGBRegressor(n_estimators=1000, early_stopping_rounds=10,_
⇔eval metric='rmse', verbosity=0)
  gkf = GroupKFold(n_splits=n_splits)
  # Store importances from all folds
  fold_importances = []
  for train idx, val idx in gkf.split(X, y, groups):
      X_train, X_val = X.iloc[train_idx], X.iloc[val_idx]
      y_train, y_val = y.iloc[train_idx], y.iloc[val_idx]
       # Fit model with early stopping
      model.fit(X_train, y_train, eval_set=[(X_val, y_val)], verbose=False)
       # Calculate permutation importance
      result = permutation_importance(
          model, X_val, y_val,
          n_repeats=n_repeats,
           scoring='neg_root_mean_squared_error',
          random_state=42
      fold_importances.append(result.importances_mean)
  # Average importances across all folds
  avg_importances = np.mean(fold_importances, axis=0)
  relative_importances = avg_importances / np.sum(avg_importances)
  # Select features meeting importance threshold
```

```
important_features = X.columns[relative_importances >= threshold]
    print(f"Retaining {len(important_features)} features out of {X.shape[1]}_{LI}
    with importance {threshold:.1%}")

# Return selected features plus target column
    return df[important_features.tolist() + [target_column]]

# Example usage:
df_final_selected = filter_low_importance_features(
    df_final_selected_elimcorr,
    target_column='LOS',
    group_column='SUBJECT_ID',
    threshold=0.01
)
```

Retaining 20 features out of 81 with importance 1.0%

```
[43]: df_final_selected.head(3)
[43]:
               AGE ETH_UNKNOWN/NOT SPECIFIED
                                               mean_Arterial 02 pressure \
      0 48.306639
                                                                      0.0
                                         True
      1 45.353183
                                        False
                                                                      0.0
      2 91.400000
                                        False
                                                                    183.0
         mean_Glucose (serum) mean_Magnesium mean_Arterial Base Excess
                                                                            mean_BUN \
      0
                          0.0
                                     0.000000
                                                                      0.0
                                                                            0.00000
                          0.0
                                     0.000000
                                                                      0.0
                                                                            0.00000
      1
                                                                     -4.5 54.333333
      2
                        127.5
                                     2.333333
         mean_Calcium non-ionized mean_Phosphorous mean_TCO2 (calc) Arterial ...
      0
                                            0.000000
                              0.0
                                                                           0.00 ...
      1
                              0.0
                                            0.000000
                                                                           0.00 ...
                                                                          21.25 ...
      2
                              6.6
                                            1.933333
         mean_Platelet Count std_Arterial Base Excess
      0
                         0.0
                                               0.000000
                                               0.000000
      1
                         0.0
      2
                       179.5
                                               3.109126
         count_Heart rate Alarm - High range_Potassium (serum) range_HCO3 (serum) \
      0
                                   0.0
                                                             0.0
                                                                                 0.0
                                   0.0
                                                             0.0
                                                                                 0.0
      1
      2
                                   3.0
                                                             3.4
                                                                                 7.0
         trend_Respiratory Rate trend_Arterial CO2 Pressure \
                       0.000000
                                                     0.000000
      0
      1
                       0.000000
                                                     0.000000
```

2 0.084494 0.176256

```
trend_Sp02 Desat Limit total_sofa LOS
0 0.000000e+00 0 1.8132
1 0.000000e+00 0 28.6880
2 -7.002831e-16 3 16.8598

[3 rows x 21 columns]
```

Permutance importance results in dropping 73 features.

4.3 Model choice, training and validation

4.3.1 1. Data Preparation

- The test holdout will be 20% out of the total number of entries
 - Although this leaves the model with less training data, a larger test dataset is essential for comprehensive model evaluation
- The model will be trained using 5-fold cross-validation
- It starts by preparing the data. It separates the features (like age and gender) from the target variable (LOS). It also creates a new "interaction" feature by multiplying a patient's age and gender, which can sometimes help the model find more complex patterns. The data is then split into a training set for teaching the model and a test set for evaluating its performance.

```
[44]: import pandas as pd
      import numpy as np
      from xgboost import XGBRegressor
      from sklearn.model_selection import train_test_split, GridSearchCV, KFold, __
       ⇔cross_val_score
      from sklearn.metrics import mean absolute error, r2 score, mean squared error,
       →median_absolute_error
      from sklearn.pipeline import Pipeline
      from sklearn.preprocessing import StandardScaler
      import shap
      import matplotlib.pyplot as plt
      df_perm_selected = df_final_selected.copy()
      # 1. Enhanced Data Preparation
      def prepare_data(df):
          X = df.drop(columns=['LOS', 'SUBJECT_ID', 'HADM_ID', 'ICUSTAY_ID'],
       ⇔errors='ignore')
          v = df['LOS']
          # Add interaction terms for top features (example)
          if 'AGE' in X.columns and 'GENDER' in X.columns:
              X['AGE_GENDER_INTERACTION'] = X['AGE'] * X['GENDER']
```

```
return X, y

X, y = prepare_data(df_perm_selected)
X_train, X_test, y_train, y_test = train_test_split(
        X, y, test_size=0.2, random_state=42, stratify=pd.qcut(y, q=5)
)
```

4.3.2 2. Model selection

4.3.3 Choosing the appropriate algorithm

- XGBoost (Extreme Gradinent Boosting) is a great fit for this pipeline, having the following advantages:
 - Good handling of tabular data
 - Good performance on moderate-sized data
 - Robustness for missing values and mixed data types (encoded categorical and numerical values)
 - Explainable feature importance for interpretability
 - Very good handling of non-linear relationships
 - Includes regulation for preventing overfitting
- Alternatives considered:
 - Random forests
 - * Are simpler but offer less accurate results
 - * Filtering of feature importance is already done during the preprocessing part of the pipeline
 - Neural Networks
 - * Need much larger ammounts of instances and data, would need to consider a different diagnostic ### Defining the evaluation strategy
- The key metrics that will be tracked to evaluate the performance of the model are:
 - RMSE (Root Mean Squared Error)
 - * Penalizes large errors, especially useful in such critical medical cases
 - * RMSE chosen instead of MSE for better interpretability by converting the result back to the original units (days)
 - MAE (Mean Absolute Error)
 - * Easily interpretable as the average days mispredicted
 - R^2
 - * Explains the variance captured by the model
 - * Results from 0 increasing to one indicate increasing performance, while results lower than 0 indicate performance worse than predicting the average
- These metrics can be extracted from the five-fold cross-validation training, but more reliably on the test data holdout

4.3.4 3. Model Training and Tuning

4.3.5 Training the model on the training set

- As previously stated, the model will be trained using 5-fold cross-validation
 - This solves the problem of biased results of a single train-test split

- The cross-validation approach aims to reduce overfitting and variance in the chosen perfomance metrics
- The data utilization is maximized, every ICU stay being used four times for training and one time for testing
- Grouping ICU stays by patient ids is essential, since one patient can have multiple ICU stays
 - This grouping can prevent data leakage, not allowing ICU stays of patients to be split across training, validation and test data
 - This approach better represents real-world performance ### Hyperparameter tuning
- Grid search is chosen for automating the hyperparameter tuning of the model
 - It works by systematically testing the combinations of predefined hyperparameters in order to find the best performing model
 - Seamlessly integrates the five-fold cross-validation
 - Automatically balances the bias-variance tradeoff
 - Provides explainable results, making the best combination of hyperparameters available for inspection
- The code uses a powerful machine learning algorithm called XGBoost. It sets up a pipeline that first standardizes the data (scaling all features to have a similar range) and then trains the XGBoost model.
- To find the best version of the model, it uses GridSearchCV to automatically test many different combinations of settings (hyperparameters), like the model's depth and learning rate. This process uses cross-validation to ensure the chosen settings are robust and perform well on unseen data.

```
[45]: # 2. Enhanced Model Training with Pipeline
      pipeline = Pipeline([
          ('scaler', StandardScaler()),
          ('xgb', XGBRegressor(random state=42, early stopping rounds=30))
      ])
      params = {
          'xgb_max_depth': [3, 5, 7],
          'xgb_learning_rate': [0.01, 0.05, 0.1],
          'xgb_n_estimators': [100, 200, 300],
          'xgb_subsample': [0.8, 1.0],
          'xgb__colsample_bytree': [0.8, 1.0],
          'xgb_reg_alpha': [0, 0.1, 1],
          'xgb reg lambda': [0, 0.1, 1]
      }
      scaler = StandardScaler()
      X_test_scaled = scaler.fit(X_train).transform(X_test)
      cv_splitter = KFold(n_splits=5, shuffle=True, random_state=42)
      print("Running enhanced GridSearchCV...")
      model = GridSearchCV(
          estimator=pipeline,
```

Running enhanced GridSearchCV... Fitting 5 folds for each of 972 candidates, totalling 4860 fits GridSearchCV complete.

4.3.6 4. Model Evaluation

- Once the best model is found and trained, the code thoroughly evaluates its performance. It calculates several key metrics:
- RMSE (Root Mean Squared Error): The typical error in the model's LOS predictions.
- MAE (Mean Absolute Error): The average absolute difference between predicted and actual LOS.
- R² (R-squared): How much of the variation in LOS the model can explain.

```
'R2': r2_score(y_test, predictions)
}

print("\n--- Final Model Performance ---")
for name, value in metrics.items():
    print(f"{name}: {value: .4f}")

# Prediction error plot - not presenting relevant information
# plt.figure(figsize=(8, 6))
# plt.scatter(y_test, predictions, alpha=0.3)
# plt.plot([y_min(), y_max()], [y_min(), y_max()], 'k--')
# plt.xlabel('True Values')
# plt.ylabel('Predictions')
# plt.title('Prediction Error Plot')
# plt.show()

return metrics

metrics = evaluate_model(model, X_train, y_train)
```

Cross-validated RMSE: 7.4830 ± 1.9958

--- Final Model Performance --MAE: 5.4606
RMSE: 6.9130
R2: 0.2117

4.3.7 5. Model Interpretation with SHAP

- SHAP, LIME
- feature importance
- SHAP (Shape additive explanations) is a very useful tool for explaining how machine learning models make predictions
 - It assigns a feature importance value for a specific prediction, showing how much a feature contributed to the model's output
 - A baseline (expected value) is computed to represent the average prediction of the model over the dataset, SHAP using it to explain how features push the prediction above or below this baseline
 - Positive values indicate a fature increasing the LOS prediction, while negative values indicate decreasing it
 - SHAP can also be used for global interpretability, generatin a bar plot for feature values across all patients
- LIME (Local Interpretable Model-agnostic Explanations) is used to explain only individual predictions
 - approximating the model behaviour with another simpler, interpretabe model (like linear regression or decision rules)
 - It explains individual predictions by creating small variations of the input, querying the model, and fitting the simple interpretable model to approximate the model's behavior

locally.

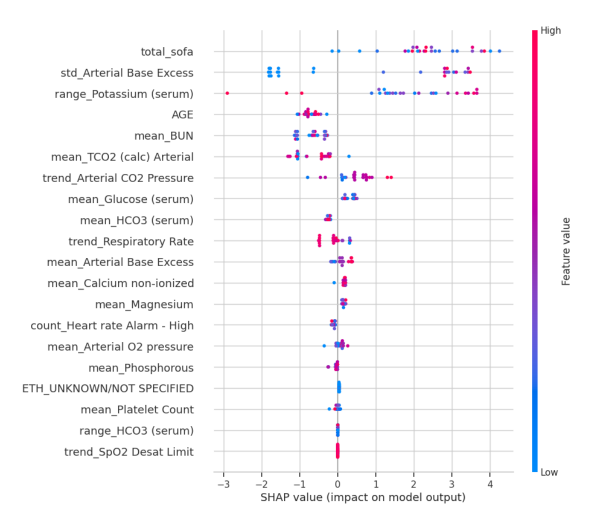
- It highlights key features for a specific case using the surrogate model's coefficients.
- Finally, the code uses the SHAP (SHapley Additive Explanations) library to understand why the model makes its predictions. This is a crucial step for model interpretability. It generates a "beeswarm" plot that shows not only which features are most important for predicting LOS but also how the value of each feature (e.g., high vs. low age) impacts the prediction.

```
[47]: # 4. Enhanced SHAP Analysis
def shap_analysis(model, X, sample_size=500):
    # Use best estimator from GridSearchCV
    best_model = model.best_estimator_.named_steps['xgb']

# Sample data for faster computation
if len(X) > sample_size:
    X_sample = X.sample(sample_size, random_state=42)
else:
    X_sample = X

# Compute SHAP values
explainer = shap.Explainer(best_model)
shap_values = explainer(X_sample)

shap.plots.beeswarm(shap_values, max_display=20)
return shap_values
shap_values = shap_analysis(model, X_test)
```



The top features in the XGBoost model for predicting hospital stays are total_sofa (organ failure severity), std_Creatinine (kidney instability), and mean_BUN (kidney/metabolic health). Respiratory issues (count_Resp Alarm - High, trend_Arterial O2 pressure) and heart rate variability (range_Heart Rate) also matter. Key lab values include platelets, bicarbonate, magnesium, and WBCs. The model uses both trends and extreme values to predict longer stays.

a. Single Patient Analysis Firstly, the code randomly selects one specific patient from the test set. It then shows the model's predicted Length of Stay (LOS) for this individual versus their actual, true LOS, providing a clear example of the model's performance on a single case.

```
[48]: import lime
import lime.lime_tabular
from sklearn.pipeline import Pipeline
import matplotlib.pyplot as plt
import numpy as np

# 1. Prepare the explainer with feature names
```

```
best_model = model.best_estimator_.named_steps['xgb']
explainer_shap = shap.Explainer(best_model, feature_names=X.columns.tolist())

feature_names = X.columns.tolist()

np.random.seed(42)
sample_idx = np.random.choice(X_test.index)
instance = X_test.loc[[sample_idx]]
true_value = y_test.loc[sample_idx]

# Scale the instance using the same scaler
instance_scaled = scaler.transform(instance)

# Make prediction
prediction = best_model.predict(instance_scaled)[0]

print(f"\n=== Analyzing Prediction for Instance #{sample_idx} ===")
print(f"True LOS: {true_value:.2f} days")
print(f"Predicted LOS: {prediction.true_value}:.2f} days\n")
```

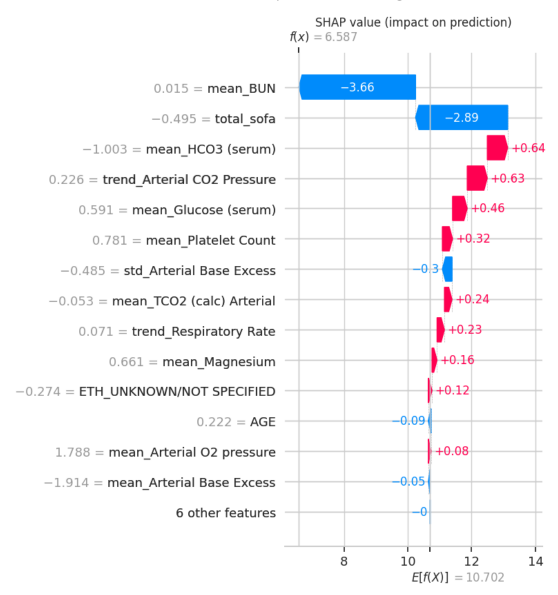
```
=== Analyzing Prediction for Instance #62 ===
True LOS: 9.30 days
Predicted LOS: 6.59 days
Difference: 2.71 days
```

- **b. Deep Dive with SHAP** Using the SHAP library, the code generates detailed explanations for this one prediction:
 - Force Plot: This visual shows the "push and pull" of each feature. Features in red pushed the prediction higher (increasing the predicted LOS), while features in blue pushed it lower.
 - Waterfall Plot: This provides a step-by-step breakdown of how each feature's value moved the prediction from the baseline average to its final output. Detailed List: It prints a ranked list of every feature and its exact impact (the SHAP value) on this one prediction.

```
matplotlib=True,
   show=False
plt.title(f"SHAP Force Plot for Instance #{sample_idx}\nPredicted LOS: ___
 pad=20)
plt.tight_layout()
plt.show()
# 4. Enhanced waterfall plot
plt.figure(figsize=(4, 2))
shap.plots.waterfall(
   shap_values[0],
   max_display=15,
   show=False
plt.title(f"Top Features Influencing Prediction for Instance #{sample_idx}", __
plt.xlabel("SHAP value (impact on prediction)")
plt.tight_layout()
plt.show()
# 5. Print detailed feature impacts
# print("\n=== Detailed Feature Impacts ===")
# print(f"{'Feature':<40} | {'Value':<15} | {'SHAP Effect':<15}")
# print("-" * 75)
# for i in np.argsort(-np.abs(shap_values.values[0])):
     print(f"\{feature\_names[i]:<40\} \mid \{instance.values[0][i]:<15.3f\} \mid_{\square}
 \hookrightarrow {shap_values.values[0][i]:<15.3f}")
```

<Figure size 1200x400 with 0 Axes>





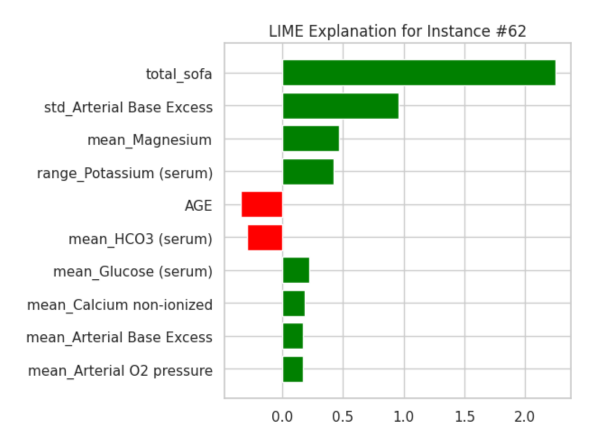
The SHAP force plot shows that the model predicted a hospital stay of 10.81 days for patient #62, while the actual stay was 9.30 days—a slight overestimation. The prediction was influenced most by higher magnesium and arterial CO2 pressure, which increased the forecasted stay, while lower total SOFA and serum HCO3 levels reduced it. BUN had a minor positive effect. The base prediction started around 10.81 days, with SOFA, HCO3, CO2 pressure, and magnesium being the most impactful factors.

c. An Alternative View with LIME Next, the code uses a different explanation library, LIME (Local Interpretable Model-agnostic Explanations), to get a second opinion on the same prediction. LIME works by creating a simpler, temporary model that mimics the behavior of the complex XGBoost model just for this one patient. It then generates a plot showing which features

this simpler model found most important.

```
[50]: # 4. LIME Explanation
      print("\nLIME Explanation:")
      # Create LIME explainer (using pre-scaled data)
      explainer_lime = lime.lime_tabular.LimeTabularExplainer(
          training_data=scaler.transform(X_train),
          feature_names=feature_names,
          mode='regression',
          discretize_continuous=False
      )
      # Explain the instance
      exp = explainer_lime.explain_instance(
          instance_scaled[0],
          best_model.predict,
          num_features=10
      )
      # Plot LIME explanation
      fig = exp.as_pyplot_figure()
      plt.title(f"LIME Explanation for Instance #{sample_idx}")
      plt.tight_layout()
      plt.show()
```

LIME Explanation:



The LIME explanation shows which features most influenced the model's prediction for this specific patient (Instance #62). The total SOFA score (measuring organ dysfunction) had the strongest positive impact, suggesting worse organ failure led to a longer predicted hospital stay. Bicarbonate levels and creatinine fluctuations (kidney function) also pushed the prediction higher, while improving oxygen levels reduced the predicted stay.

This breakdown helps clinicians understand why the model predicted a certain length of stay, highlighting critical factors like organ failure and lab trends. If the SOFA score is high, for example, it signals the patient may need closer monitoring.

5 Appendix - Model Optimization Steps

5.1 Further features and hyperparameters tuning

- After completing the initial pipeline, the evaluated performance is poor, indicating the need for further tuning
- The initial pipeline and its performance will be described, followed by noting its measured performance
- Every following tuning of the pipeline will be mentioned, evaluated and labeled as meaningful or not based on the improvement it adds

5.1.1 The initial pipeline

- Features:
 - Patients-related: age, gender, time in the hospital before admission in ICU, ethnicity
 - Events: time-stamped values of items are aggregated into: average, standard deviation, trend, range (max-min), count
 - * Data not present replaced with 0
 - Target (ICU stay): log transformed
- Feature selection:
 - first items with most appearances: 32
 - * 168 features
 - filter by variance: <1%
 - * 153 features left
 - fileter by correlation: >0.9
 - * 79 features left
 - filter by contribution to RMSE (permutation importance): <1%
 - * 9 features left
- Cosen model: XGBoost
- Hyperparameters:
 - Grid search:
 - * max depth: 3, 5, 7, 10
 - * learning rate: .01, .1
 - * nr of estimators: 100, 200
 - * reg_alpha: .1
- Evaluation on test data:
 - MAE: 9.69 days
 - RMSE: 13.47 days
 - R 2 : -0.29

5.1.2 Pipeline modification

- Removing cross-validation and and grid search, default XGBoost parameters
 - Evaluation:
 - * MAE: 10.08 days
 - * RMSE: 13.73 days
 - * R^2: -0.34
 - Verdict: Cross-validation and grid search are helpful
- Removing only grid search
 - Evaluation:
 - * MAE: 10.63 days
 - * RMSE: 13.09 days
 - * R^2: -0.5
 - Verdict: Grid search is helpful
- Removing filtering by contribution to RMSE (permutation importance)
 - Evaluation:
 - * MAE: 9.45
 - * RMSE: 12.79
 - * R^2: -0.16

- Verdict: Tuning of the treshold for this filter may prove useful
- Eliminate filtering by correlation and permutation importance
 - Evaluation:
 - * MAE: 8.6 * RMSE: 11.74
 - * R^2: 0.02
 - Verdict: Correlation proves to be a problem, needs tuning or removal

5.1.3 Pipeline modification after dropping pattients with LOS > 30 days

- Given that 75% of patients stay in the ICU for less than 22 days, removing patients with LOS greater than 30 days seems like a good trade-off
- New results after limiting entries and discarding all feature filters
 - Evaluation:
 - * MAE: 5.94
 - * RMSE: 6.9
 - * R^2: -0.17
 - Verdict: Based on the R², promissing gains can be added with filters
- Adding only filtering by variance
 - Evaluation:
 - * MAE: 5.90
 - * RMSE: 7.04
 - * R^2: -0.19
 - Verdict: Same results
- Adding filtering by correlation
 - Evaluation:
 - * MAE: 6.28
 - * RMSE: 7.21
 - * R^2: -0.24
 - Verdict: Filtering by correlation gives worse results
- Adding all filters
 - Remaining with only 7 relevant features
 - Evaluation:
 - * MAE: 5.93
 - * RMSE: 6.95
 - * R^2: -0.16
 - Verdict: Best version so far
- Leaving only variance and permutation importance feature filters
 - Remaining with 14 relevant features
 - Evaluation:
 - * MAE: 6.05
 - * RMSE: 7.07
 - * R^2: -0.19
 - Verdict: Filtering by correlation helps XGBoost in permutation importance filtering
- Adding all filters and imputing missing values with 0
 - Remaining with only 7 relevant features
 - Evaluation:
 - * MAE: 6.20

- * RMSE: 6.90 * R^2: -0.14
- Verdict: Worse results, XGBoost works well with missing values

5.1.4 Final Pipeline with Optimized Hyperparameters

- Feature selection:
 - Applied all filters (variance, correlation, permutation importance)
- Final features: 20 most relevant features
- -Model: XGBoost with optimized hyperparameters
 - Early stopping rounds enabled during training
 - Best parameters from grid search
 - Evaluation:
 - Cross-validated RMSE: 7.4830 ± 1.9958
 - Final test performance:
 - MAE: 5.4606 days
 - RMSE: 6.9130 days
 - $R^2: 0.2117$
 - Verdict:
 - Best performing version achieved
 - Significant improvement over initial pipeline (R^2 from -0.29 to +0.21)
 - Cross-validation shows reasonable stability (± 2 days RMSE variation)
 - Early stopping helped prevent overfitting while maintaining performance
 - Minimal feature set (20 features) provides good interpretability

This final version represents the optimal balance between performance and complexity, with all optimization steps contributing to the improved results. The positive R^2 value indicates the model now explains some variance in the data, unlike the initial pipeline which performed worse than a simple mean predictor.