

# Medical Text Transcription Classification ML model with XAI Integration

July 17, 2024

```
[1]: # Import necessary libraries
import os # For interacting with the operating system
import re # For regular expressions
import warnings # For managing warnings
import random # For generating random numbers

import numpy as np # For numerical operations
import pandas as pd # For data manipulation and analysis
import matplotlib # For plotting and visualization
import matplotlib.pyplot as plt # For plotting and visualization
import seaborn as sns # For statistical data visualization

from wordcloud import WordCloud # For generating word clouds

# Natural Language Toolkit (NLTK) libraries for text processing
from nltk.tokenize import WhitespaceTokenizer, word_tokenize # For tokenizing
↳text
from nltk.stem import WordNetLemmatizer # For lemmatizing text
from nltk.corpus import stopwords # For accessing stop words

# Scikit-learn libraries for machine learning and data processing
from sklearn import preprocessing # For preprocessing data
from sklearn.feature_extraction.text import CountVectorizer, TfidfVectorizer #
↳For text vectorization
from sklearn.decomposition import PCA, TruncatedSVD # For dimensionality
↳reduction
from sklearn.ensemble import RandomForestClassifier # For random forest
↳classification
from sklearn.svm import SVC # For support vector classification
from xgboost import XGBClassifier # For XGBoost classification
from sklearn.neighbors import KNeighborsClassifier # For k-nearest neighbors
↳classification
from sklearn.linear_model import LogisticRegression # For logistic regression
from sklearn.tree import DecisionTreeClassifier # For decision tree
↳classification
```

```

from sklearn.metrics import classification_report, confusion_matrix, roc_curve,
    ↪ auc, roc_auc_score # For model evaluation metrics
from sklearn.preprocessing import LabelEncoder, StandardScaler, label_binarize
    ↪ # For preprocessing data
from sklearn.model_selection import train_test_split, KFold # For splitting
    ↪ data and cross-validation

# Imbalanced-learn library for handling imbalanced datasets
from imblearn.over_sampling import SMOTE # For synthetic minority
    ↪ over-sampling technique

# PyTorch libraries for deep learning
import torch # For PyTorch operations
from torch.utils.data import TensorDataset, DataLoader, RandomSampler,
    ↪ SequentialSampler # For PyTorch data handling
from transformers import BertTokenizer, BertForSequenceClassification,
    ↪ RobertaTokenizer, RobertaForSequenceClassification, AdamW,
    ↪ get_linear_schedule_with_warmup # For using BERT, BioBERT, ClinicalBERT and
    ↪ RoBERTa models

```

```

[2]: # Mount Google Drive to access files
from google.colab import drive
drive.mount('/content/drive')

# Read a CSV file from the specified path in Google Drive
df = pd.read_csv("/content/drive/My Drive/dataset/mtsamples.csv")

# Display the first few rows of the DataFrame
df.head()

```

Mounted at /content/drive

```

[2]: Unnamed: 0      description \
0      0      A 23-year-old white female presents with comp...
1      1      Consult for laparoscopic gastric bypass.
2      2      Consult for laparoscopic gastric bypass.
3      3      2-D M-Mode. Doppler.
4      4      2-D Echocardiogram

      medical_specialty      sample_name \
0      Allergy / Immunology      Allergic Rhinitis
1      Bariatrics      Laparoscopic Gastric Bypass Consult - 2
2      Bariatrics      Laparoscopic Gastric Bypass Consult - 1
3      Cardiovascular / Pulmonary      2-D Echocardiogram - 1
4      Cardiovascular / Pulmonary      2-D Echocardiogram - 2

      transcription \

```

```

0 SUBJECTIVE:, This 23-year-old white female pr...
1 PAST MEDICAL HISTORY:, He has difficulty climb...
2 HISTORY OF PRESENT ILLNESS: , I have seen ABC ...
3 2-D M-MODE: , ,1. Left atrial enlargement wit...
4 1. The left ventricular cavity size and wall ...

```

keywords

```

0 allergy / immunology, allergic rhinitis, aller...
1 bariatrics, laparoscopic gastric bypass, weigh...
2 bariatrics, laparoscopic gastric bypass, heart...
3 cardiovascular / pulmonary, 2-d m-mode, dopple...
4 cardiovascular / pulmonary, 2-d, doppler, echo...

```

```

[3]: # Calculate the count of non-null entries for each column in the dataframe
non_null_counts = df.count()

```

```

# Set the size of the figure for the plot
plt.figure(figsize=(12, 6))

# Create a bar plot of the non-null counts
ax = non_null_counts.plot(kind='bar')

# Set the title of the plot
plt.title('Non-null Entries per Column')

# Label the x-axis
plt.xlabel('Columns')

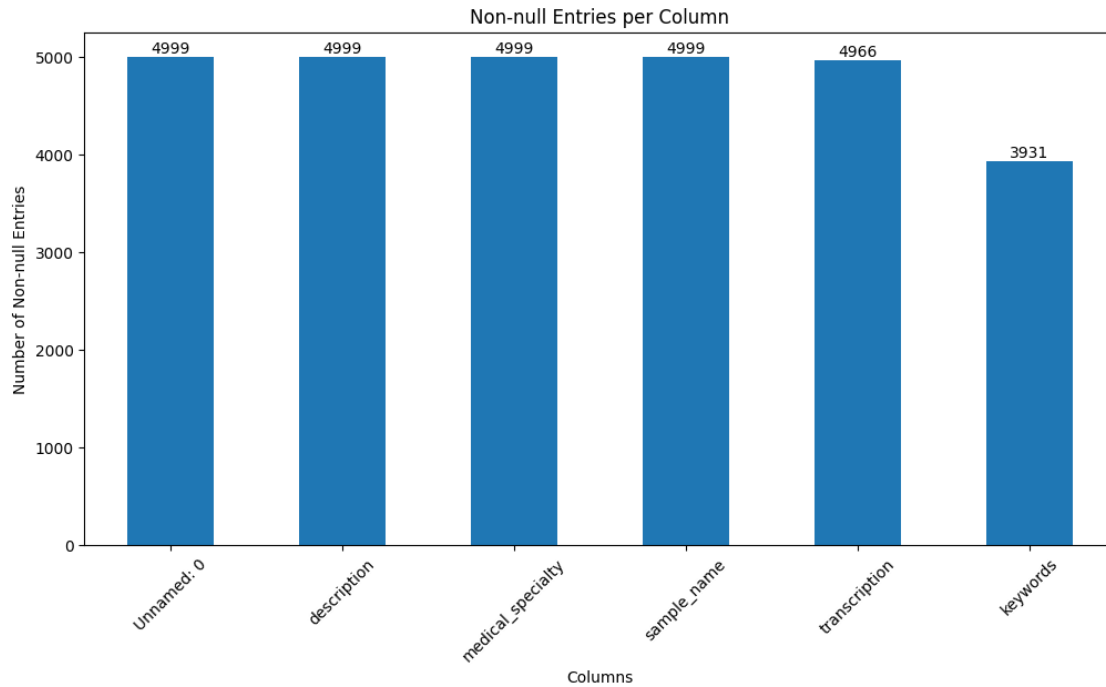
# Label the y-axis
plt.ylabel('Number of Non-null Entries')

# Rotate the x-axis labels for better readability
plt.xticks(rotation=45)

# Add labels to each bar in the plot
for i in ax.containers:
    ax.bar_label(i)

# Display the plot
plt.show()

```



```
[4]: # Calculate the number of null entries in each column of the dataframe
null_counts = df.isnull().sum()

# Set the figure size for the plot
plt.figure(figsize=(12, 6))

# Create a bar plot of the null counts per column
ax = null_counts.plot(kind='bar')

# Set the title of the plot
plt.title('Null Entries per Column')

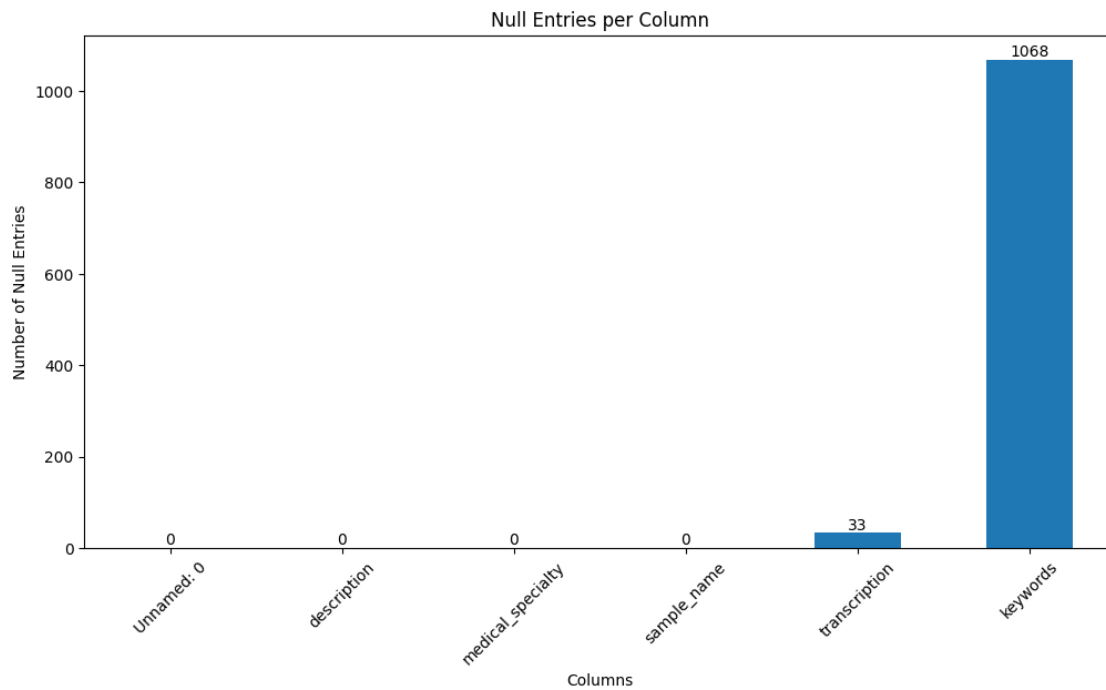
# Set the label for the x-axis
plt.xlabel('Columns')

# Set the label for the y-axis
plt.ylabel('Number of Null Entries')

# Rotate the x-axis labels for better readability
plt.xticks(rotation=45)

# Add labels to the bars in the plot
for i in ax.containers:
    ax.bar_label(i)
```

```
# Display the plot
plt.show()
```



```
[5]: def trim(df):
    """
    Cleans the DataFrame by performing the following operations:
    - Strips whitespace from column names
    - Drops duplicate rows
    - Converts column names to lowercase
    - Replaces spaces in column names with underscores
    - Strips whitespace from categorical data (object dtype columns)
    """

    # Strip whitespace from column names
    df.columns = df.columns.str.strip()
    # Drop duplicate rows
    df = df.drop_duplicates()
    # Convert column names to lowercase
    df.columns = df.columns.str.lower()
    # Replace spaces in column names with underscores
    df.columns = df.columns.str.replace(' ', '_')

    # Select columns with object dtype (categorical data)
    df_obj = df.select_dtypes(['object'])
    # Strip whitespace from categorical data
```

```

df[df_obj.columns] = df_obj.apply(lambda x: x.str.strip())

# Print status messages
print("All column names have been stripped, converted to lowercase, and_
↳spaces replaced with underscores.")
print("Duplicate rows have been dropped.")
print("Whitespace has been stripped from categorical data.")
return df

# Set pandas option to display max column width
pd.set_option('display.max_colwidth', 255)

# Drop the 'Unnamed: 0' column from the DataFrame
df.drop('Unnamed: 0', axis=1, inplace=True)

# Apply the trim function to clean the DataFrame
df = trim(df)

def vc(df, column, r=False):
    """
    Computes value counts and percentages for a specified column.

    Parameters:
    df (pd.DataFrame): The DataFrame to analyze.
    column (str): The column name to compute value counts for.
    r (bool): If True, returns the resulting DataFrame. If False, prints and_
↳displays it.

    Returns:
    pd.DataFrame: Value counts and percentages if r is True.
    """
    # Compute value counts and convert to DataFrame
    vc_df = df.reset_index().groupby([column]).size().to_frame('count')
    # Compute percentages
    vc_df['percentage (%)'] = vc_df['count'].div(sum(vc_df['count'])).mul(100)
    # Sort by percentage in descending order
    vc_df = vc_df.sort_values(by=['percentage (%)'], ascending=False)

    if r:
        return vc_df
    else:
        # Print status message and display DataFrame
        print(f'STATUS: Value counts of "{column}"...')
        display(vc_df)

def shape(df, df_name):
    """

```

*Prints the dimensions (number of rows and columns) of the DataFrame.*

*Parameters:*

*df (pd.DataFrame): The DataFrame whose dimensions are to be printed.*

*df\_name (str): The name of the DataFrame to be printed in the message.*

*"""*

*# Print dimensions of the DataFrame*

*print(f'STATUS: Dimension of "{df\_name}" = {df.shape}')*

*# Display the first 3 rows of the DataFrame*

*df.head(3)*

All column names have been stripped, converted to lowercase, and spaces replaced with underscores.

Duplicate rows have been dropped.

Whitespace has been stripped from categorical data.

```
[5]:                                     description \
0  A 23-year-old white female presents with complaint of allergies.
1                                     Consult for laparoscopic gastric bypass.
2                                     Consult for laparoscopic gastric bypass.

      medical_specialty                                     sample_name \
0  Allergy / Immunology                                     Allergic Rhinitis
1          Bariatrics  Laparoscopic Gastric Bypass Consult - 2
2          Bariatrics  Laparoscopic Gastric Bypass Consult - 1

      transcription \
0  SUBJECTIVE:, This 23-year-old white female presents with complaint of
allergies. She used to have allergies when she lived in Seattle but she thinks
they are worse here. In the past, she has tried Claritin, and Zyrtec. Both
worked for short time b...
1  PAST MEDICAL HISTORY:, He has difficulty climbing stairs, difficulty with
airline seats, tying shoes, used to public seating, and lifting objects off the
floor. He exercises three times a week at home and does cardio. He has
difficulty walking two b...
2  HISTORY OF PRESENT ILLNESS: , I have seen ABC today. He is a very pleasant
gentleman who is 42 years old, 344 pounds. He is 5'9". He has a BMI of 51. He
has been overweight for ten years since the age of 33, at his highest he was 358
pounds, at hi...

                                     keywords
0  allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergic,
1  bariatrics, laparoscopic gastric bypass, weight loss programs, gastric
bypass, atkin's diet, weight watcher's, body weight, laparoscopic gastric,
```

weight loss, pounds, months, weight, laparoscopic, band, loss, diets, overweight, lost  
 2 bariatrics, laparoscopic gastric bypass, heart attacks, body weight, pulmonary embolism, potential complications, sleep study, weight loss, gastric bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart, pounds, weight,

```
[6]: # Print the dimensions of the dataframe (number of rows and columns)
      print(df.shape)
```

(4999, 5)

```
[8]: # Select specific columns from the DataFrame
      df = df[['transcription', 'keywords', 'medical_specialty']]

      # Display the first few rows of the DataFrame to verify the selection
      df.head()
```

```
[8]:      transcription \
0  SUBJECTIVE:, This 23-year-old white female presents with complaint of allergies. She used to have allergies when she lived in Seattle but she thinks they are worse here. In the past, she has tried Claritin, and Zyrtec. Both worked for short time b...
1  PAST MEDICAL HISTORY:, He has difficulty climbing stairs, difficulty with airline seats, tying shoes, used to public seating, and lifting objects off the floor. He exercises three times a week at home and does cardio. He has difficulty walking two b...
2  HISTORY OF PRESENT ILLNESS: , I have seen ABC today. He is a very pleasant gentleman who is 42 years old, 344 pounds. He is 5'9". He has a BMI of 51. He has been overweight for ten years since the age of 33, at his highest he was 358 pounds, at hi...
3  2-D M-MODE: , ,1. Left atrial enlargement with left atrial diameter of 4.7 cm.,2. Normal size right and left ventricle.,3. Normal LV systolic function with left ventricular ejection fraction of 51%.,4. Normal LV diastolic function.,5. No pericard..
4  1. The left ventricular cavity size and wall thickness appear normal. The wall motion and left ventricular systolic function appears hyperdynamic with estimated ejection fraction of 70% to 75%. There is near-cavity obliteration seen. There also ap...
```

```
      keywords \
0
allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays, rhinitis, nasal, erythematous, allegra, sprays, allergic,
1      bariatrics, laparoscopic gastric bypass, weight loss programs, gastric bypass, atkin's diet, weight watcher's, body weight, laparoscopic gastric, weight loss, pounds, months, weight, laparoscopic, band,
```



```

loss, diets, overweight, lost
2          bariatrics, laparoscopic gastric bypass, heart attacks,
body weight, pulmonary embolism, potential complications, sleep study, weight
loss, gastric bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass,
heart, pounds, weight,
3 cardiovascular / pulmonary, 2-d m-mode, doppler, aortic valve, atrial
enlargement, diastolic function, ejection fraction, mitral, mitral valve,
pericardial effusion, pulmonary valve, regurgitation, systolic function,
tricuspid, tricuspid valve, normal lv
4 cardiovascular / pulmonary, 2-d, doppler, echocardiogram, annular, aortic
root, aortic valve, atrial, atrium, calcification, cavity, ejection fraction,
mitral, obliteration, outflow, regurgitation, relaxation pattern, stenosis,
systolic function, tric...

```

```

          medical_specialty
0      Allergy / Immunology
1          Bariatrics
2          Bariatrics
3 Cardiovascular / Pulmonary
4 Cardiovascular / Pulmonary

```

##Exploratory Data Analysis

```

[9]: # Create a copy of the DataFrame to avoid modifying the original DataFrame
df_1 = df.copy()

```

```

[10]: # Calculate the counts of unique values in the 'medical_specialty' column of
      ↪the DataFrame 'df_1'
value_counts = df_1['medical_specialty'].value_counts()

# Print the resulting counts
print(value_counts)

```

```

medical_specialty
Surgery          1103
Consult - History and Phy.    516
Cardiovascular / Pulmonary    372
Orthopedic        355
Radiology         273
General Medicine   259
Gastroenterology   230
Neurology          223
SOAP / Chart / Progress Notes  166
Obstetrics / Gynecology    160
Urology           158
Discharge Summary    108
ENT - Otolaryngology    98
Neurosurgery        94

```

Hematology - Oncology	90
Ophthalmology	83
Nephrology	81
Emergency Room Reports	75
Pediatrics - Neonatal	70
Pain Management	62
Psychiatry / Psychology	53
Office Notes	51
Podiatry	47
Dermatology	29
Cosmetic / Plastic Surgery	27
Dentistry	27
Letters	23
Physical Medicine - Rehab	21
Sleep Medicine	20
Endocrinology	19
Bariatrics	18
IME-QME-Work Comp etc.	16
Chiropractic	14
Rheumatology	10
Diets and Nutritions	10
Speech - Language	9
Autopsy	8
Lab Medicine - Pathology	8
Allergy / Immunology	7
Hospice - Palliative Care	6

Name: count, dtype: int64

#Analyze Categorical Variable

```
[11]: # Calculate the count of each unique value in the 'medical_specialty' column of
      ↪ the dataframe df_1
value_counts = df_1['medical_specialty'].value_counts()

# Set the size of the plot
plt.figure(figsize=(12, 8))

# Create a bar plot with the index (medical specialties) on the x-axis and the
↪ counts on the y-axis
plt.bar(value_counts.index, value_counts)

# Add a title to the plot
plt.title('Distribution of Medical Specialties')

# Label the x-axis
plt.xlabel('Medical Specialty')

# Label the y-axis
```

```

plt.ylabel('Count')

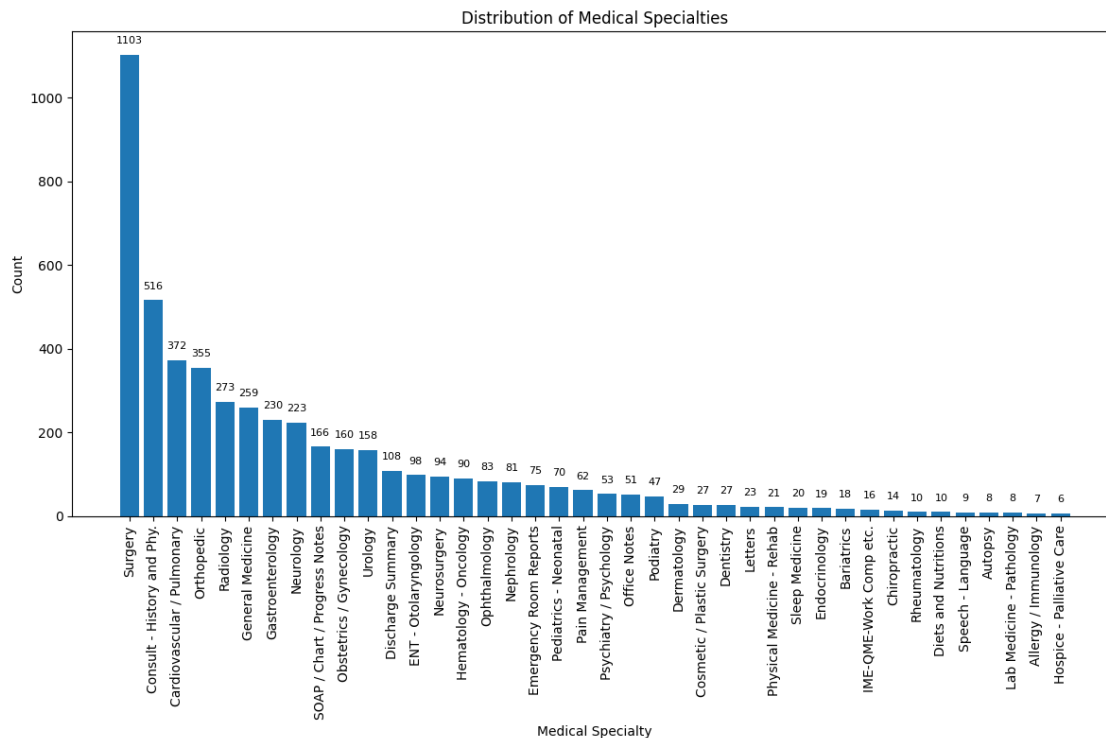
# Add the count labels on top of each bar
for i, count in enumerate(value_counts):
    # Position the text slightly above the bar
    plt.text(i, count + 0.02 * max(value_counts), str(count), ha='center',
    ↪va='bottom', fontsize=8)

# Rotate the x-axis labels by 90 degrees for better readability
plt.xticks(rotation=90)

# Adjust the plot to ensure everything fits without overlap
plt.tight_layout()

# Display the plot
plt.show()

```



Word cloud of the medical specialty column

```

[12]: # Convert the 'medical_specialty' column in the dataframe to string type
df_1['medical_specialty'] = df_1['medical_specialty'].astype(str)

```



```

    # Filter the DataFrame to only include rows with the current medical_
↪specialty
    df_filter = df_1.loc[df_1['medical_specialty'] == medical_specialty]
    # Calculate the total word count for the current specialty by summing the_
↪word counts of all transcriptions
    word_count_temp = df_filter['transcription'].str.split().str.len().sum()
    # Append the current medical specialty and its word count to their_
↪respective lists
    medical_specialty_list.append(medical_specialty)
    word_count_list.append(word_count_temp)

# Create a new DataFrame with the collected medical specialties and word counts
word_count_df = pd.DataFrame({'Medical Specialty': medical_specialty_list,
↪'Word Count': word_count_list})
# Ensure the 'Word Count' column is of integer type
word_count_df['Word Count'] = word_count_df['Word Count'].astype('int')
# Sort the DataFrame by word count in descending order
word_count_df = word_count_df.sort_values('Word Count', ascending=False)
# Reset the index of the DataFrame and drop the old index
word_count_df.reset_index(drop=True)

```

```

[13]:

```

	Medical Specialty	Word Count
0	Surgery	526754
1	Consult - History and Phy.	287961
2	Orthopedic	198489
3	Cardiovascular / Pulmonary	160867
4	General Medicine	120978
5	Neurology	110677
6	Gastroenterology	80347
7	Radiology	74969
8	Obstetrics / Gynecology	72589
9	Urology	63419
10	SOAP / Chart / Progress Notes	59558
11	Neurosurgery	54233
12	Discharge Summary	43103
13	Psychiatry / Psychology	42972
14	ENT - Otolaryngology	42032
15	Emergency Room Reports	41854
16	Hematology - Oncology	36953
17	Nephrology	35443
18	Pediatrics - Neonatal	30724
19	Ophthalmology	30354
20	Podiatry	22468
21	IME-QME-Work Comp etc.	17418
22	Cosmetic / Plastic Surgery	14791
23	Pain Management	14496
24	Dentistry	13745

25	Chiropractic	13006
26	Office Notes	12713
27	Endocrinology	12119
28	Dermatology	11790
29	Letters	11511
30	Autopsy	9474
31	Physical Medicine - Rehab	9237
32	Bariatrics	7012
33	Sleep Medicine	5868
34	Rheumatology	4880
35	Hospice - Palliative Care	4057
36	Diets and Nutritions	3967
37	Speech - Language	3631
38	Allergy / Immunology	3132
39	Lab Medicine - Pathology	1828

```
[14]: # Initialize empty lists to store medical specialties and their corresponding
      ↪ word counts
medical_specialty_list = []
word_count_list = []

# Loop through each unique medical specialty in the DataFrame 'df_1'
for medical_specialty in df_1['medical_specialty'].unique():
    # Filter the DataFrame to only include rows with the current medical
    ↪ specialty
    df_filter = df_1.loc[df_1['medical_specialty'] == medical_specialty]
    # Calculate the total word count for the current specialty by summing the
    ↪ word counts of all transcriptions
    word_count_temp = df_filter['transcription'].str.split().str.len().sum()
    # Append the current medical specialty and its word count to their
    ↪ respective lists
    medical_specialty_list.append(medical_specialty)
    word_count_list.append(word_count_temp)

# Create a new DataFrame with the collected medical specialties and word counts
word_count_df = pd.DataFrame({'Medical Specialty': medical_specialty_list,
    ↪ 'Word Count': word_count_list})
# Ensure the 'Word Count' column is of integer type
word_count_df['Word Count'] = word_count_df['Word Count'].astype('int')
# Sort the DataFrame by word count in descending order
word_count_df = word_count_df.sort_values('Word Count', ascending=False)
# Reset the index of the DataFrame and drop the old index
word_count_df = word_count_df.reset_index(drop=True)

# Import matplotlib for plotting
import matplotlib.pyplot as plt
```

```

# Set the figure size for the plot
plt.figure(figsize=(10, 6))

# Create a horizontal bar plot
bars = plt.barh(y=word_count_df['Medical Specialty'], width=word_count_df['Word_
Count'], color='skyblue')

# Add word counts as text on the bars
for bar in bars:
    width = bar.get_width()
    plt.annotate(f'{int(width)}',
                 xy=(width, bar.get_y() + bar.get_height() / 2),
                 xytext=(5, 0),
                 textcoords='offset points',
                 ha='left',
                 va='center')

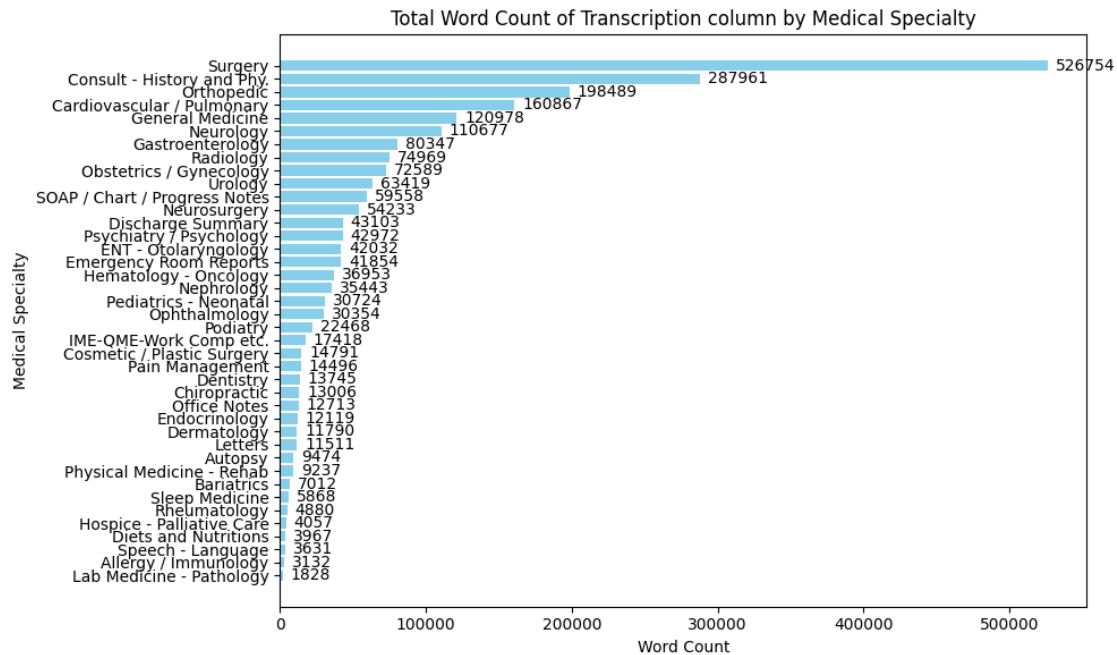
# Label the axes and title the plot
plt.xlabel('Word Count')
plt.ylabel('Medical Specialty')
plt.title('Total Word Count of Transcription column by Medical Specialty')

# Invert y-axis to have the specialty with the highest word count at the top
plt.gca().invert_yaxis()

# Adjust spacing between subplots
plt.tight_layout()

# Display the plot
plt.show()

```



Word cloud of the transcription column

```
[15]: # Convert the 'transcription' column to string type to ensure compatibility for
      ↪ text processing
df_1['transcription'] = df_1['transcription'].astype(str)

# Concatenate all the strings in the 'transcription' column into a single text
text = ' '.join(df_1['transcription'])

# Generate a word cloud with the specified background color using the
      ↪ concatenated text
wordcloud = WordCloud(background_color='white').generate(text)

# Plot the word cloud
plt.figure(figsize=(10, 10))
plt.imshow(wordcloud, interpolation='bilinear')
plt.axis('off')
plt.show()
```





```

# Convert 'Word Count' column to integer type
word_count_df['Word Count'] = word_count_df['Word Count'].astype('int')

# Sort DataFrame by 'Word Count' in descending order
word_count_df = word_count_df.sort_values('Word Count', ascending=False)

# Reset index of the sorted DataFrame
word_count_df = word_count_df.reset_index(drop=True)

# Create a horizontal bar plot
plt.figure(figsize=(10, 6))
bars = plt.barh(y=word_count_df['Medical Specialty'], width=word_count_df['Word_
Count'], color='skyblue')

# Annotate each bar with its value
for bar in bars:
    width = bar.get_width()
    plt.annotate(f'{int(width)}',
                xy=(width, bar.get_y() + bar.get_height() / 2),
                xytext=(5, 0),
                textcoords='offset points',
                ha='left',
                va='center')

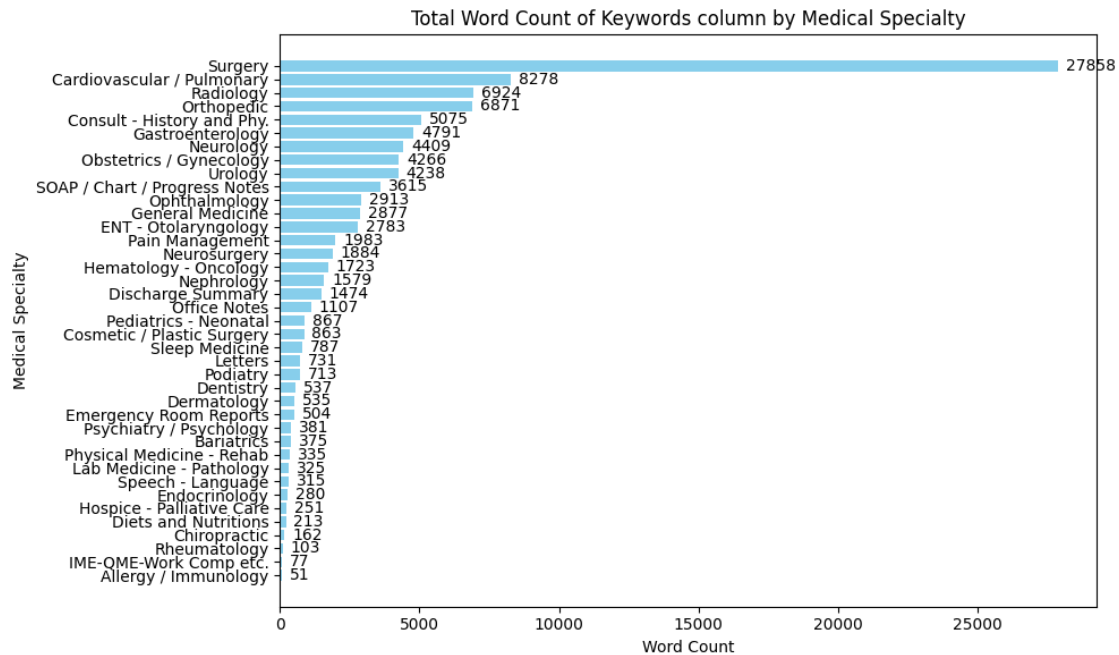
# Add labels and title to the plot
plt.xlabel('Word Count')
plt.ylabel('Medical Specialty')
plt.title('Total Word Count of Keywords column by Medical Specialty')

# Invert y-axis to display highest word count at the top
plt.gca().invert_yaxis()

# Adjust layout to prevent cropping of labels
plt.tight_layout()

# Display the plot
plt.show()

```



Word cloud of the keywords column

```
[17]: # Convert the 'keywords' column in df_1 to string type
df_1['keywords'] = df_1['keywords'].astype(str)

# Join all values in the 'keywords' column into a single string separated by
↳ spaces
text = ' '.join(df_1['keywords'])

# Generate a WordCloud object with white background
wordcloud = WordCloud(background_color='white').generate(text)

# Plot the WordCloud
plt.figure(figsize=(10, 10)) # Set the size of the plot
plt.imshow(wordcloud, interpolation='bilinear') # Display the WordCloud with
↳ bilinear interpolation
plt.axis('off') # Turn off axis labels
plt.show() # Display the plot
```



```
docs/stable/user_guide/indexing.html#returning-a-view-versus-a-copy
df['keywords'] = df['keywords'].fillna('')
```

```
[21]: # Print the shape of the DataFrame 'df'
print(f'data shape is: {df.shape}')

# Check and display the count of null values in each column of 'df', sorted in
↳descending order
df.isnull().sum().sort_values(ascending=False)
```

```
data shape is: (4966, 3)
```

```
[21]: transcription      0
      keywords          0
      medical_specialty  0
      dtype: int64
```

```
#Concatenation and Subsetting
```

```
[22]: # Concatenate 'keywords' and 'transcription' columns and store the result in a
↳new 'transcription' column
df['transcription'] = df['keywords'] + df['transcription']
df.head()
```

```
<ipython-input-22-625e380a33d0>:2: SettingWithCopyWarning:
A value is trying to be set on a copy of a slice from a DataFrame.
Try using .loc[row_indexer,col_indexer] = value instead
```

See the caveats in the documentation: [https://pandas.pydata.org/pandas-docs/stable/user\\_guide/indexing.html#returning-a-view-versus-a-copy](https://pandas.pydata.org/pandas-docs/stable/user_guide/indexing.html#returning-a-view-versus-a-copy)

```
df['transcription'] = df['keywords'] + df['transcription']
```

```
[22]:      transcription \
0  allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergic,SUBJECTIVE:, This
23-year-old white female presents with complaint of allergies. She used to have
allergies when she l...
1  bariatrics, laparoscopic gastric bypass, weight loss programs, gastric
bypass, atkin's diet, weight watcher's, body weight, laparoscopic gastric,
weight loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostPAST MEDICAL HISTOR...
2  bariatrics, laparoscopic gastric bypass, heart attacks, body weight,
pulmonary embolism, potential complications, sleep study, weight loss, gastric
bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart, pounds,
weight,HISTORY OF PRESE...
3  cardiovascular / pulmonary, 2-d m-mode, doppler, aortic valve, atrial
enlargement, diastolic function, ejection fraction, mitral, mitral valve,
pericardial effusion, pulmonary valve, regurgitation, systolic function,
```

```

tricuspid, tricuspid valve, normal...
4 cardiovascular / pulmonary, 2-d, doppler, echocardiogram, annular, aortic
root, aortic valve, atrial, atrium, calcification, cavity, ejection fraction,
mitral, obliteration, outflow, regurgitation, relaxation pattern, stenosis,
systolic function, tric...

```

```

keywords \
0
allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergic,
1
    bariatrics, laparoscopic gastric bypass, weight loss
programs, gastric bypass, atkin's diet, weight watcher's, body weight,
laparoscopic gastric, weight loss, pounds, months, weight, laparoscopic, band,
loss, diets, overweight, lost
2
    bariatrics, laparoscopic gastric bypass, heart attacks,
body weight, pulmonary embolism, potential complications, sleep study, weight
loss, gastric bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass,
heart, pounds, weight,
3 cardiovascular / pulmonary, 2-d m-mode, doppler, aortic valve, atrial
enlargement, diastolic function, ejection fraction, mitral, mitral valve,
pericardial effusion, pulmonary valve, regurgitation, systolic function,
tricuspid, tricuspid valve, normal lv
4 cardiovascular / pulmonary, 2-d, doppler, echocardiogram, annular, aortic
root, aortic valve, atrial, atrium, calcification, cavity, ejection fraction,
mitral, obliteration, outflow, regurgitation, relaxation pattern, stenosis,
systolic function, tric...

```

```

    medical_specialty
0 Allergy / Immunology
1 Bariatrics
2 Bariatrics
3 Cardiovascular / Pulmonary
4 Cardiovascular / Pulmonary

```

```

[23]: # Select only the 'transcription' and 'medical_specialty' columns from the
      ↪ DataFrame 'df'
df = df[['transcription', 'medical_specialty']]
df.head()

```

```

[23]: transcription \
0 allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergic,SUBJECTIVE:, This
23-year-old white female presents with complaint of allergies. She used to have
allergies when she l...
1 bariatrics, laparoscopic gastric bypass, weight loss programs, gastric
bypass, atkin's diet, weight watcher's, body weight, laparoscopic gastric,
weight loss, pounds, months, weight, laparoscopic, band, loss, diets,

```

```

overweight, lostPAST MEDICAL HISTOR...
2 bariatrics, laparoscopic gastric bypass, heart attacks, body weight,
pulmonary embolism, potential complications, sleep study, weight loss, gastric
bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart, pounds,
weight,HISTORY OF PRESE...
3 cardiovascular / pulmonary, 2-d m-mode, doppler, aortic valve, atrial
enlargement, diastolic function, ejection fraction, mitral, mitral valve,
pericardial effusion, pulmonary valve, regurgitation, systolic function,
tricuspid, tricuspid valve, normal...
4 cardiovascular / pulmonary, 2-d, doppler, echocardiogram, annular, aortic
root, aortic valve, atrial, atrium, calcification, cavity, ejection fraction,
mitral, obliteration, outflow, regurgitation, relaxation pattern, stenosis,
systolic function, tric...

        medical_specialty
0      Allergy / Immunology
1                Bariatrics
2                Bariatrics
3 Cardiovascular / Pulmonary
4 Cardiovascular / Pulmonary

```

```

[24]: # Print the shape of the DataFrame 'df'
print(f'data shape is: {df.shape}')

# Check and display the count of null values in each column of 'df', sorted in_
↳descending order
df.isnull().sum().sort_values(ascending=False)

```

data shape is: (4966, 2)

```

[24]: transcription      0
      medical_specialty  0
      dtype: int64

```

Word count of the transcription column by medical specialty after concatenation

```

[25]: # Initialize lists to store medical specialties and their corresponding word_
↳counts
medical_specialty_list = []
word_count_list = []

# Iterate over unique values in 'medical_specialty' column
for medical_specialty in df['medical_specialty'].unique():
    # Filter DataFrame for the current medical specialty
    df_filter = df.loc[df['medical_specialty'] == medical_specialty]

```

```

    # Calculate total word count for transcriptions in the current medical_
↪specialty
    word_count_temp = df_filter['transcription'].str.split().str.len().sum()

    # Append medical specialty and its word count to lists
    medical_specialty_list.append(medical_specialty)
    word_count_list.append(word_count_temp)

# Create a DataFrame from the lists
word_count_df = pd.DataFrame({'Medical Specialty': medical_specialty_list,
↪'Word Count': word_count_list})

# Convert 'Word Count' to integer type
word_count_df['Word Count'] = word_count_df['Word Count'].astype('int')

# Sort DataFrame by 'Word Count' in descending order
word_count_df = word_count_df.sort_values('Word Count', ascending=False)

# Reset index for the DataFrame
word_count_df = word_count_df.reset_index(drop=True)

# Display the resulting DataFrame (note: printing the result is not shown in_
↪the original code)
word_count_df

```

```

[25]:

```

	Medical Specialty	Word Count
0	Surgery	552159
1	Consult - History and Phy.	292816
2	Orthopedic	205064
3	Cardiovascular / Pulmonary	168780
4	General Medicine	123718
5	Neurology	114924
6	Gastroenterology	84407
7	Radiology	81645
8	Obstetrics / Gynecology	76246
9	Urology	67294
10	SOAP / Chart / Progress Notes	63033
11	Neurosurgery	56038
12	ENT - Otolaryngology	44516
13	Discharge Summary	44501
14	Psychiatry / Psychology	43334
15	Emergency Room Reports	42329
16	Hematology - Oncology	38616
17	Nephrology	36959
18	Ophthalmology	33188
19	Pediatrics - Neonatal	31553
20	Podiatry	23139



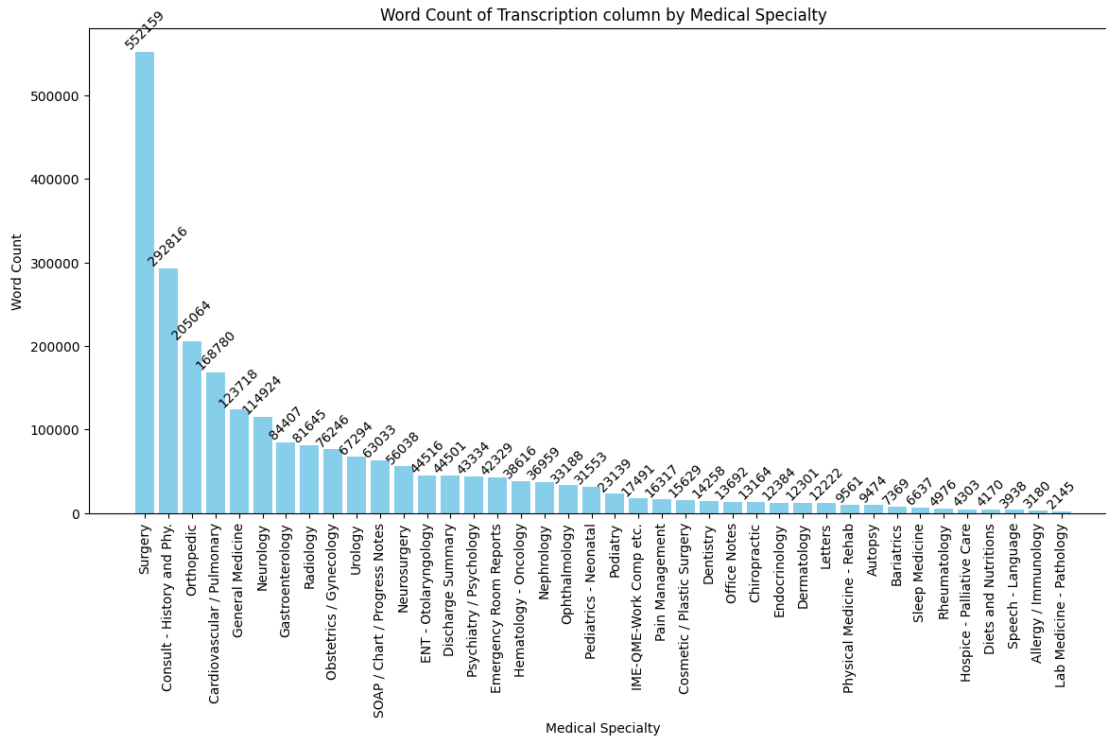
21	IME-QME-Work Comp etc.	17491
22	Pain Management	16317
23	Cosmetic / Plastic Surgery	15629
24	Dentistry	14258
25	Office Notes	13692
26	Chiropractic	13164
27	Endocrinology	12384
28	Dermatology	12301
29	Letters	12222
30	Physical Medicine - Rehab	9561
31	Autopsy	9474
32	Bariatrics	7369
33	Sleep Medicine	6637
34	Rheumatology	4976
35	Hospice - Palliative Care	4303
36	Diets and Nutritions	4170
37	Speech - Language	3938
38	Allergy / Immunology	3180
39	Lab Medicine - Pathology	2145

```
[26]: # Plotting the bar graph
plt.figure(figsize=(12, 8)) # Set the figure size
bars = plt.bar(word_count_df['Medical Specialty'], word_count_df['Word Count'],
               color='skyblue') # Create bars for each medical specialty
plt.xlabel('Medical Specialty') # Label for the x-axis
plt.ylabel('Word Count') # Label for the y-axis
plt.title('Word Count of Transcription column by Medical Specialty') # Title
of the plot
plt.xticks(rotation=90) # Rotate x-axis labels for better visibility

# Adding the word count on top of each bar with rotation
for bar in bars:
    yval = bar.get_height() # Get the height (word count) of each bar
    plt.text(
        bar.get_x() + bar.get_width()/2, # x-coordinate for placing the text
        at the center of the bar
        yval, # y-coordinate for placing the text (same as bar height)
        int(yval), # Text to display (converted to integer for neatness)
        ha='center', # Horizontal alignment of the text
        va='bottom', # Vertical alignment of the text
        rotation=45 # Rotate the text for better alignment
    )

plt.tight_layout() # Adjust layout to prevent clipping of labels

# Show the plot
plt.show()
```



```
[27]: # Calculate the total word count of all transcriptions before data preprocessing
total_word_count = df['transcription'].str.split().str.len().sum()

# Print the total word count
print(f'The word count of all transcriptions before data preprocessing is:␣
↪{int(total_word_count)}')
```

The word count of all transcriptions before data preprocessing is: 2407470

## 0.1 Converting to Lowercase

```
[28]: def lower(df, attribute):
    # Apply lowercase conversion using lambda function to the specified␣
    ↪attribute column
    df[attribute] = df[attribute].apply(lambda x: str(x).lower() if pd.
    ↪notnull(x) else x)
    return df

# Example usage:
# Apply lowercase conversion to 'transcription' column of DataFrame 'df'
df = lower(df, 'transcription')

# Display the first 3 rows of the modified DataFrame
```

```
df.head(3)
```

```
[28]: transcription \
0 allergy / immunology, allergic rhinitis, allergies, asthma, nasal sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergic,subjective:, this
23-year-old white female presents with complaint of allergies. she used to have
allergies when she l...
1 bariatrics, laparoscopic gastric bypass, weight loss programs, gastric
bypass, atkin's diet, weight watcher's, body weight, laparoscopic gastric,
weight loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostpast medical histor...
2 bariatrics, laparoscopic gastric bypass, heart attacks, body weight,
pulmonary embolism, potential complications, sleep study, weight loss, gastric
bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart, pounds,
weight,history of prese...

    medical_specialty
0 Allergy / Immunology
1 Bariatrics
2 Bariatrics
```

##Removing Punctuation and Numbers

```
[29]: # Ignore warnings (optional)
warnings.filterwarnings('ignore')

def remove_punc_num(df, attribute):
    # Ensure the attribute column is treated as string
    df[attribute] = df[attribute].astype(str)

    # Remove standalone digits
    df[attribute] = df[attribute].apply(lambda x: re.sub(r'\b\d+\b', '', x))

    # Remove alphanumeric sequences with digits
    df[attribute] = df[attribute].apply(lambda x: re.sub(r'\w*\d\w*', '', x))

    # Remove punctuation
    df[attribute] = df[attribute].apply(lambda x: re.sub(r'[\w\s]', '', x))

    # Remove extra spaces and convert to lowercase
    df[attribute] = df[attribute].apply(lambda x: " ".join(re.
↵findall(r'\b\w+\b', x.lower()))))

    # Display the cleaned DataFrame
    print("Head of DataFrame after cleaning:")
    print(df.head(3)) # Display the first 3 rows of the DataFrame
```

```
return df
```

```
# Call the function to clean the 'transcription' column of DataFrame 'df'  
df_cleaned = remove_punc_num(df, 'transcription')
```

Head of DataFrame after cleaning:

```
transcription \  
0 allergy immunology allergic rhinitis allergies asthma nasal sprays rhinitis  
nasal erythematous allegra sprays allergicsubjective this yearold white female  
presents with complaint of allergies she used to have allergies when she lived  
in seattle but sh...  
1 bariatrics laparoscopic gastric bypass weight loss programs gastric bypass  
atkins diet weight watchers body weight laparoscopic gastric weight loss pounds  
months weight laparoscopic band loss diets overweight lostpast medical history  
he has difficulty...  
2 bariatrics laparoscopic gastric bypass heart attacks body weight pulmonary  
embolism potential complications sleep study weight loss gastric bypass  
anastomosis loss sleep laparoscopic gastric bypass heart pounds weighthistory of  
present illness i have ...
```

```
medical_specialty  
0 Allergy / Immunology  
1 Bariatrics  
2 Bariatrics
```

#Tokenization

```
[30]: # Initialize WhitespaceTokenizer  
tk = WhitespaceTokenizer()  
  
def tokenise(df, attribute):  
    # Tokenize each row of the specified attribute column using  
    ↪ WhitespaceTokenizer  
    df['tokenised'] = df.apply(lambda row: tk.tokenize(str(row[attribute])),  
    ↪ axis=1)  
    return df  
  
# Call the tokenise function to tokenize the 'transcription' column of  
    ↪ DataFrame 'df'  
df = tokenise(df, 'transcription')  
  
# Create a copy of the DataFrame for experimentation  
df_experiment = df.copy()  
  
# Display the first 3 rows of the DataFrame after tokenization  
df.head(3)
```

```
[30]: transcription \
0 allergy immunology allergic rhinitis allergies asthma nasal sprays rhinitis
nasal erythematous allegra sprays allergicsubjective this yearold white female
presents with complaint of allergies she used to have allergies when she lived
in seattle but sh...
1 bariatrics laparoscopic gastric bypass weight loss programs gastric bypass
atkins diet weight watchers body weight laparoscopic gastric weight loss pounds
months weight laparoscopic band loss diets overweight lostpast medical history
he has difficulty...
2 bariatrics laparoscopic gastric bypass heart attacks body weight pulmonary
embolism potential complications sleep study weight loss gastric bypass
anastomosis loss sleep laparoscopic gastric bypass heart pounds weighthistory of
present illness i have ...
```

```
medical_specialty \
0 Allergy / Immunology
1 Bariatrics
2 Bariatrics
```

```
tokenised
0 [allergy, immunology, allergic, rhinitis, allergies, asthma, nasal, sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergicsubjective, this,
yearold, white, female, presents, with, complaint, of, allergies, she, used, to,
have, allergies, whe...
1 [bariatrics, laparoscopic, gastric, bypass, weight, loss, programs, gastric,
bypass, atkins, diet, weight, watchers, body, weight, laparoscopic, gastric,
weight, loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostpast, medi...
2 [bariatrics, laparoscopic, gastric, bypass, heart, attacks, body, weight,
pulmonary, embolism, potential, complications, sleep, study, weight, loss,
gastric, bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart,
pounds, weighthistory...
```

#Stop Words Removal

```
[31]: import nltk

# Download NLTK stopwords dataset
nltk.download('stopwords')
```

```
[nltk_data] Downloading package stopwords to /root/nltk_data...
[nltk_data] Unzipping corpora/stopwords.zip.
```

```
[31]: True
```

```
[32]: # Showing the list of the English stop words, it has a number of 179 stop words
↳ in this list
```

```
stop = stopwords.words('english')
print(f"There are {len(stop)} stop words \n")
print(stop)
```

There are 179 stop words

```
['i', 'me', 'my', 'myself', 'we', 'our', 'ours', 'ourselves', 'you', "you're",
"you've", "you'll", "you'd", 'your', 'yours', 'yourself', 'yourselves', 'he',
'him', 'his', 'himself', 'she', "she's", 'her', 'hers', 'herself', 'it', "it's",
'its', 'itself', 'they', 'them', 'their', 'theirs', 'themselves', 'what',
'which', 'who', 'whom', 'this', 'that', "that'll", 'these', 'those', 'am', 'is',
'are', 'was', 'were', 'be', 'been', 'being', 'have', 'has', 'had', 'having',
'do', 'does', 'did', 'doing', 'a', 'an', 'the', 'and', 'but', 'if', 'or',
'because', 'as', 'until', 'while', 'of', 'at', 'by', 'for', 'with', 'about',
'against', 'between', 'into', 'through', 'during', 'before', 'after', 'above',
'below', 'to', 'from', 'up', 'down', 'in', 'out', 'on', 'off', 'over', 'under',
'again', 'further', 'then', 'once', 'here', 'there', 'when', 'where', 'why',
'how', 'all', 'any', 'both', 'each', 'few', 'more', 'most', 'other', 'some',
'such', 'no', 'nor', 'not', 'only', 'own', 'same', 'so', 'than', 'too', 'very',
's', 't', 'can', 'will', 'just', 'don', "don't", 'should', "should've", 'now',
'd', 'll', 'm', 'o', 're', 've', 'y', 'ain', 'aren', "aren't", 'couldn',
"couldn't", 'didn', "didn't", 'doesn', "doesn't", 'hadn', "hadn't", 'hasn',
"hasn't", 'haven', "haven't", 'isn', "isn't", 'ma', 'mightn', "mightn't",
'mustn', "mustn't", 'needn', "needn't", 'shan', "shan't", 'shouldn',
"shouldn't", 'wasn', "wasn't", 'weren', "weren't", 'won', "won't", 'wouldn',
"wouldn't"]
```

```
[33]: # Function to remove stop words
def remove_stop_words(df, attribute):
    stop = set(stopwords.words('english')) # Use set for faster lookup
    df['stop_words_removal'] = df[attribute].apply(lambda x: [word for word in x
↪ if word.lower() not in stop])
    return df

# Remove stop words from the 'tokenised' column
df = remove_stop_words(df, 'tokenised')
df.head(2)
```

```
[33]: transcription \
0 allergy immunology allergic rhinitis allergies asthma nasal sprays rhinitis
nasal erythematous allegra sprays allergicsubjective this yearold white female
presents with complaint of allergies she used to have allergies when she lived
in seattle but sh...
1 bariatrics laparoscopic gastric bypass weight loss programs gastric bypass
atkins diet weight watchers body weight laparoscopic gastric weight loss pounds
months weight laparoscopic band loss diets overweight lostpast medical history
he has difficulty...
```

```

    medical_specialty \
0 Allergy / Immunology
1          Bariatrics

```

```

    tokenised \
0 [allergy, immunology, allergic, rhinitis, allergies, asthma, nasal, sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergicsubjective, this,
yearold, white, female, presents, with, complaint, of, allergies, she, used, to,
have, allergies, whe...
1 [bariatrics, laparoscopic, gastric, bypass, weight, loss, programs, gastric,
bypass, atkins, diet, weight, watchers, body, weight, laparoscopic, gastric,
weight, loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostpast, medi...

```

```

stop_words_removal
0 [allergy, immunology, allergic, rhinitis, allergies, asthma, nasal, sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergicsubjective, yearold,
white, female, presents, complaint, allergies, used, allergies, lived, seattle,
thinks, worse, pas...
1 [bariatrics, laparoscopic, gastric, bypass, weight, loss, programs, gastric,
bypass, atkins, diet, weight, watchers, body, weight, laparoscopic, gastric,
weight, loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostpast, medi...

```

#Lemmatization

```

[34]: import nltk

# Download NLTK datasets
nltk.download('punkt')
nltk.download('wordnet')

# Initialize WordNetLemmatizer
wordnet_lemmatizer = WordNetLemmatizer()

# Function to lemmatize text
def lemmatize_text(input_text):
    lemmatized_words = []
    words = word_tokenize(input_text)
    lemmatized_words.extend([wordnet_lemmatizer.lemmatize(word) for word in
↪words])
    return lemmatized_words

# Function to lemmatize a column of text
def lemmatize(df, input_column, output_column):

```

```

    df[output_column] = df[input_column].apply(lambda x: ', '.
↪join(lemmatize_text(' '.join(x))))
    return df

# Assuming 'df' is your DataFrame and 'stop_words_removal' is the column with
↪the text to be lemmatized
df = lemmatize(df, 'stop_words_removal', 'lemmatized_transcription')

```

```

[nltk_data] Downloading package punkt to /root/nltk_data...
[nltk_data] Unzipping tokenizers/punkt.zip.
[nltk_data] Downloading package wordnet to /root/nltk_data...

```

```
[35]: df.head(2)
```

```

[35]:      transcription \
0  allergy immunology allergic rhinitis allergies asthma nasal sprays rhinitis
nasal erythematous allegra sprays allergicsubjective this yearold white female
presents with complaint of allergies she used to have allergies when she lived
in seattle but sh...
1  bariatrics laparoscopic gastric bypass weight loss programs gastric bypass
atkins diet weight watchers body weight laparoscopic gastric weight loss pounds
months weight laparoscopic band loss diets overweight lostpast medical history
he has difficulty...

      medical_specialty \
0  Allergy / Immunology
1           Bariatrics

      tokenised \
0  [allergy, immunology, allergic, rhinitis, allergies, asthma, nasal, sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergicsubjective, this,
yearold, white, female, presents, with, complaint, of, allergies, she, used, to,
have, allergies, whe...
1  [bariatrics, laparoscopic, gastric, bypass, weight, loss, programs, gastric,
bypass, atkins, diet, weight, watchers, body, weight, laparoscopic, gastric,
weight, loss, pounds, months, weight, laparoscopic, band, loss, diets,
overweight, lostpast, medi...

stop_words_removal \
0  [allergy, immunology, allergic, rhinitis, allergies, asthma, nasal, sprays,
rhinitis, nasal, erythematous, allegra, sprays, allergicsubjective, yearold,
white, female, presents, complaint, allergies, used, allergies, lived, seattle,
thinks, worse, pas...
1  [bariatrics, laparoscopic, gastric, bypass, weight, loss, programs, gastric,
bypass, atkins, diet, weight, watchers, body, weight, laparoscopic, gastric,
weight, loss, pounds, months, weight, laparoscopic, band, loss, diets,

```



```
overweight, lostpast, medi...
```

```
lemmatized_transcription
```

```
0  allergy, immunology, allergic, rhinitis, allergy, asthma, nasal, spray,
rhinitis, nasal, erythematous, allegra, spray, allergicsubjective, yearold,
white, female, present, complaint, allergy, used, allergy, lived, seattle,
think, worse, past, tried, c...
1  bariatrics, laparoscopic, gastric, bypass, weight, loss, program, gastric,
bypass, atkins, diet, weight, watcher, body, weight, laparoscopic, gastric,
weight, loss, pound, month, weight, laparoscopic, band, loss, diet, overweight,
lostpast, medical, h...
```

```
[36]: # Drop columns 'transcription', 'tokenised', 'stop_words_removal'
df = df.drop(['transcription', 'tokenised', 'stop_words_removal'], axis=1)

# Display the first 2 rows of the modified DataFrame
df.head(2)
```

```
[36]:      medical_specialty \
0  Allergy / Immunology
1      Bariatrics

lemmatized_transcription
0  allergy, immunology, allergic, rhinitis, allergy, asthma, nasal, spray,
rhinitis, nasal, erythematous, allegra, spray, allergicsubjective, yearold,
white, female, present, complaint, allergy, used, allergy, lived, seattle,
think, worse, past, tried, c...
1  bariatrics, laparoscopic, gastric, bypass, weight, loss, program, gastric,
bypass, atkins, diet, weight, watcher, body, weight, laparoscopic, gastric,
weight, loss, pound, month, weight, laparoscopic, band, loss, diet, overweight,
lostpast, medical, h...
```

```
[37]: # Rename the column 'lemmatized_transcription' to 'preprocessed_transcription'
df.rename(columns={'lemmatized_transcription': 'preprocessed_transcription'},
          inplace=True)

# Display the first few rows of the DataFrame to verify the rename operation
df.head()
```

```
[37]:      medical_specialty \
0      Allergy / Immunology
1      Bariatrics
2      Bariatrics
3  Cardiovascular / Pulmonary
4  Cardiovascular / Pulmonary

preprocessed_transcription
```

0 allergy, immunology, allergic, rhinitis, allergy, asthma, nasal, spray, rhinitis, nasal, erythematous, allegra, spray, allergicsubjective, yearold, white, female, present, complaint, allergy, used, allergy, lived, seattle, think, worse, past, tried, c...

1 bariatrics, laparoscopic, gastric, bypass, weight, loss, program, gastric, bypass, atkins, diet, weight, watcher, body, weight, laparoscopic, gastric, weight, loss, pound, month, weight, laparoscopic, band, loss, diet, overweight, lostpast, medical, h...

2 bariatrics, laparoscopic, gastric, bypass, heart, attack, body, weight, pulmonary, embolism, potential, complication, sleep, study, weight, loss, gastric, bypass, anastomosis, loss, sleep, laparoscopic, gastric, bypass, heart, pound, weighthistory, pr...

3 cardiovascular, pulmonary, mmode, doppler, aortic, valve, atrial, enlargement, diastolic, function, ejection, fraction, mitral, mitral, valve, pericardial, effusion, pulmonary, valve, regurgitation, systolic, function, tricuspid, tricuspid, valve, nor...

4 cardiovascular, pulmonary, doppler, echocardiogram, annular, aortic, root, aortic, valve, atrial, atrium, calcification, cavity, ejection, fraction, mitral, obliteration, outflow, regurgitation, relaxation, pattern, stenosis, systolic, function, tricu...

```
[38]: # Calculate word count for each preprocessed_transcription
df['word_count'] = df['preprocessed_transcription'].apply(lambda x: len(str(x).
    ↪split()))

# Aggregate word counts by category (assuming the category column is
    ↪'medical_specialty')
word_counts_by_category = df.groupby('medical_specialty')['word_count'].sum().
    ↪reset_index()

# Sort the DataFrame by word counts in descending order
word_counts_by_category = word_counts_by_category.sort_values(by='word_count',
    ↪ascending=False)

# Plotting the bar graph
plt.figure(figsize=(12, 8))
bars = plt.bar(word_counts_by_category['medical_specialty'],
    ↪word_counts_by_category['word_count'], color='skyblue')
plt.xlabel('Medical Specialty')
plt.ylabel('Word Count')
plt.title('Word Count of Preprocessed Transcription by Medical Specialty')
plt.xticks(rotation=90)

# Adding the word count on top of each bar with rotation
for bar in bars:
    yval = bar.get_height()
    plt.text(
```

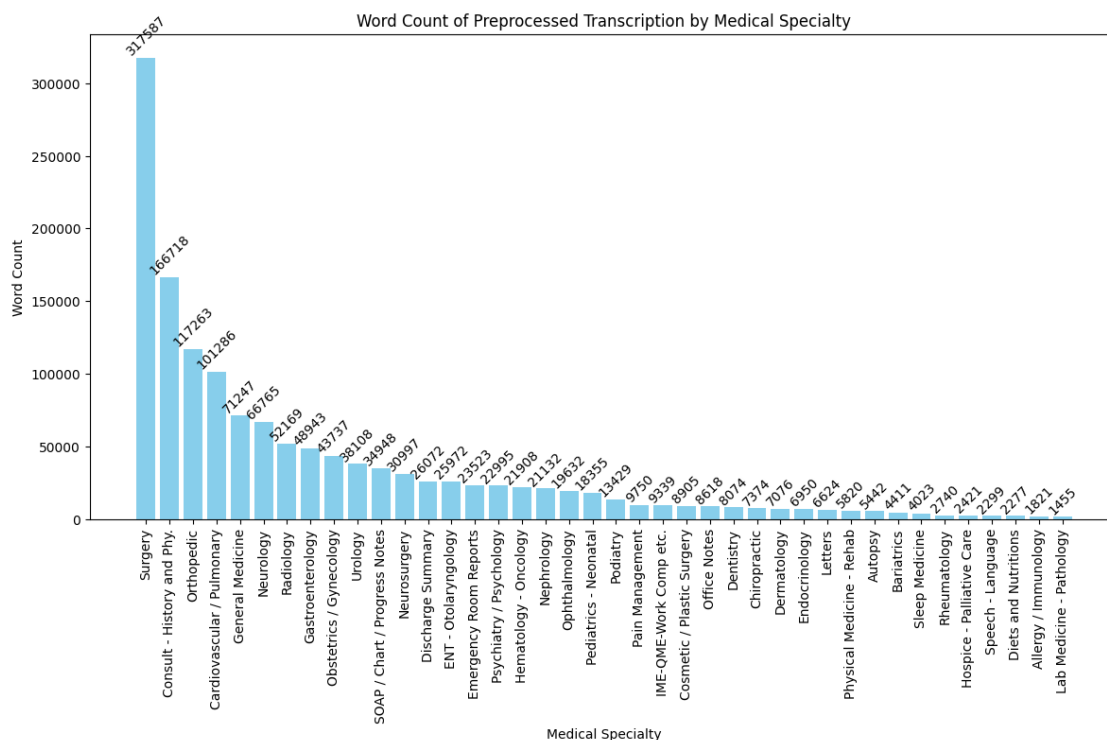
```

        bar.get_x() + bar.get_width() / 2,
        yval,
        int(yval),
        ha='center',
        va='bottom',
        rotation=45
    )

plt.tight_layout()

# Show the plot
plt.show()

```



##Applying Domian Knowledge

```

[39]: filtered_data_categories = df.copy()

# Strip whitespace from 'medical_specialty' column values

filtered_data_categories['medical_specialty'] =_
    ↪filtered_data_categories['medical_specialty'].apply(lambda x: x.strip())

# List of categories to exclude

```

```

categories_to_exclude = [
    'Surgery', 'SOAP / Chart / Progress Notes', 'Office Notes',
    'Consult - History and Phy.', 'Emergency Room Reports',
    'Discharge Summary', 'Pain Management', 'General Medicine', 'Letters'
]

# Exclude specified categories
for category in categories_to_exclude:
    mask = filtered_data_categories['medical_specialty'] == category
    filtered_data_categories = filtered_data_categories[~mask]

# Adjusting categories
filtered_data_categories['medical_specialty'] =
    ↪ filtered_data_categories['medical_specialty'].replace({
        'Neurosurgery': 'Neurology',
        'Nephrology': 'Urology'
    })

# Group by 'medical_specialty' and get the size of each group
grouped_data = filtered_data_categories.groupby('medical_specialty').size()

# Sort the groups by size in descending order
sorted_grouped_data = grouped_data.sort_values(ascending=False)

i = 1
print('=====Reduced Categories=====')
for catName, count in sorted_grouped_data.items():
    print(f'Cat: {i} {catName} : {count}')
    i += 1
print('=====Reduced Categories=====')

# Select relevant columns and drop rows with NaN values in
    ↪ 'preprocessed_transcription'
df = filtered_data_categories[['preprocessed_transcription',
    ↪ 'medical_specialty']]
df = df.dropna(subset=['preprocessed_transcription'])
print(f'data shape is: {df.shape}')

```

```

=====Reduced Categories=====
Cat: 1 Cardiovascular / Pulmonary : 371
Cat: 2 Orthopedic : 355
Cat: 3 Neurology : 317
Cat: 4 Radiology : 273
Cat: 5 Urology : 237
Cat: 6 Gastroenterology : 224
Cat: 7 Obstetrics / Gynecology : 155

```

```

Cat: 8 ENT - Otolaryngology : 96
Cat: 9 Hematology - Oncology : 90
Cat: 10 Ophthalmology : 83
Cat: 11 Pediatrics - Neonatal : 70
Cat: 12 Psychiatry / Psychology : 53
Cat: 13 Podiatry : 47
Cat: 14 Dermatology : 29
Cat: 15 Cosmetic / Plastic Surgery : 27
Cat: 16 Dentistry : 27
Cat: 17 Physical Medicine - Rehab : 21
Cat: 18 Sleep Medicine : 20
Cat: 19 Endocrinology : 19
Cat: 20 Bariatrics : 18
Cat: 21 IME-QME-Work Comp etc. : 16
Cat: 22 Chiropractic : 14
Cat: 23 Diets and Nutritions : 10
Cat: 24 Rheumatology : 10
Cat: 25 Speech - Language : 9
Cat: 26 Lab Medicine - Pathology : 8
Cat: 27 Autopsy : 8
Cat: 28 Allergy / Immunology : 7
Cat: 29 Hospice - Palliative Care : 6
=====Reduced Categories=====
data shape is: (2620, 2)

```

Count of medical specialties after applying domain knowledge

```

[40]: # Set the figure size
plt.figure(figsize=(10, 6))

# Create the count plot using Seaborn
ax = sns.countplot(y='medical_specialty', # Plotting against
    ↪ 'medical_specialty' on the y-axis
                  data=filtered_data_categories, # Using data from
    ↪ 'filtered_data_categories'
                  order=filtered_data_categories['medical_specialty'].
    ↪ value_counts().index) # Order by value counts

# Set plot title and labels
plt.title('Count of Medical Specialties')
plt.xlabel('Count')
plt.ylabel('Medical Specialty')

# Annotate each bar with its count
for p in ax.patches:
    ax.annotate(f'{int(p.get_width())}', # Text to display (count of each bar)
                (p.get_width(), p.get_y() + p.get_height() / 2), # Position to
    ↪ annotate

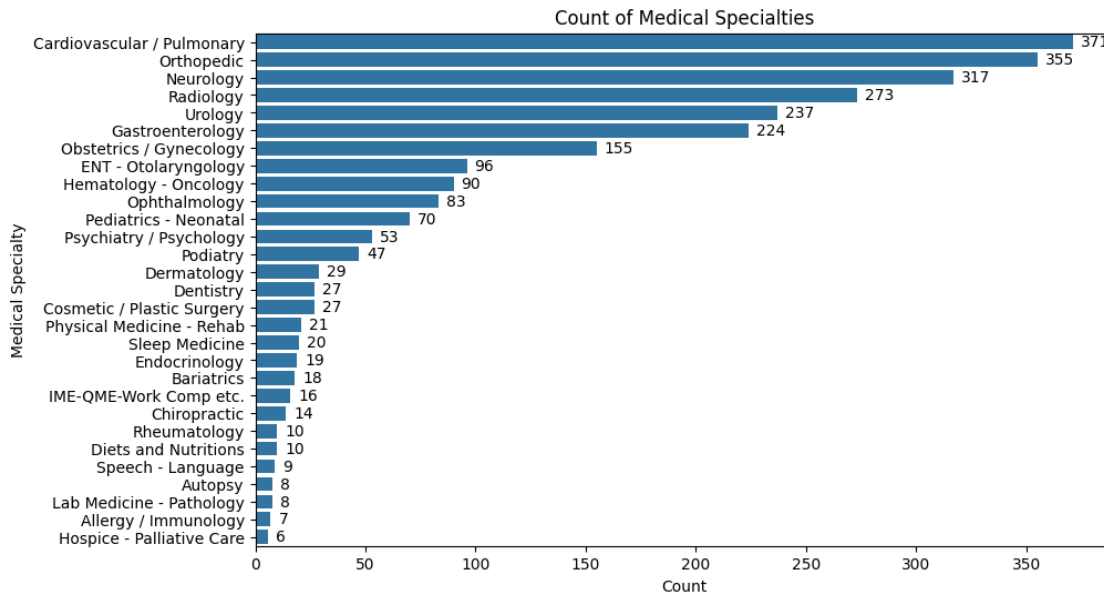
```

```

ha='left', va='center', # Alignment of the text
xytext=(5, 0), # Offset of the text from the annotation point
textcoords='offset points') # Offset in points

# Display the plot
plt.show()

```



### ##Category Filtering and Adjustment

```

[41]: # Count the occurrences of each medical specialty
counts = df['medical_specialty'].value_counts()

# Filter the DataFrame to keep only rows where 'medical_specialty' appears 50
↳ times or more
df = df[~df['medical_specialty'].isin(counts[counts < 50].index)]

```

Finalized medical specialty and respective count

```

[42]: # Check the value counts again to see the cleaned data
print(df['medical_specialty'].value_counts())

```

```

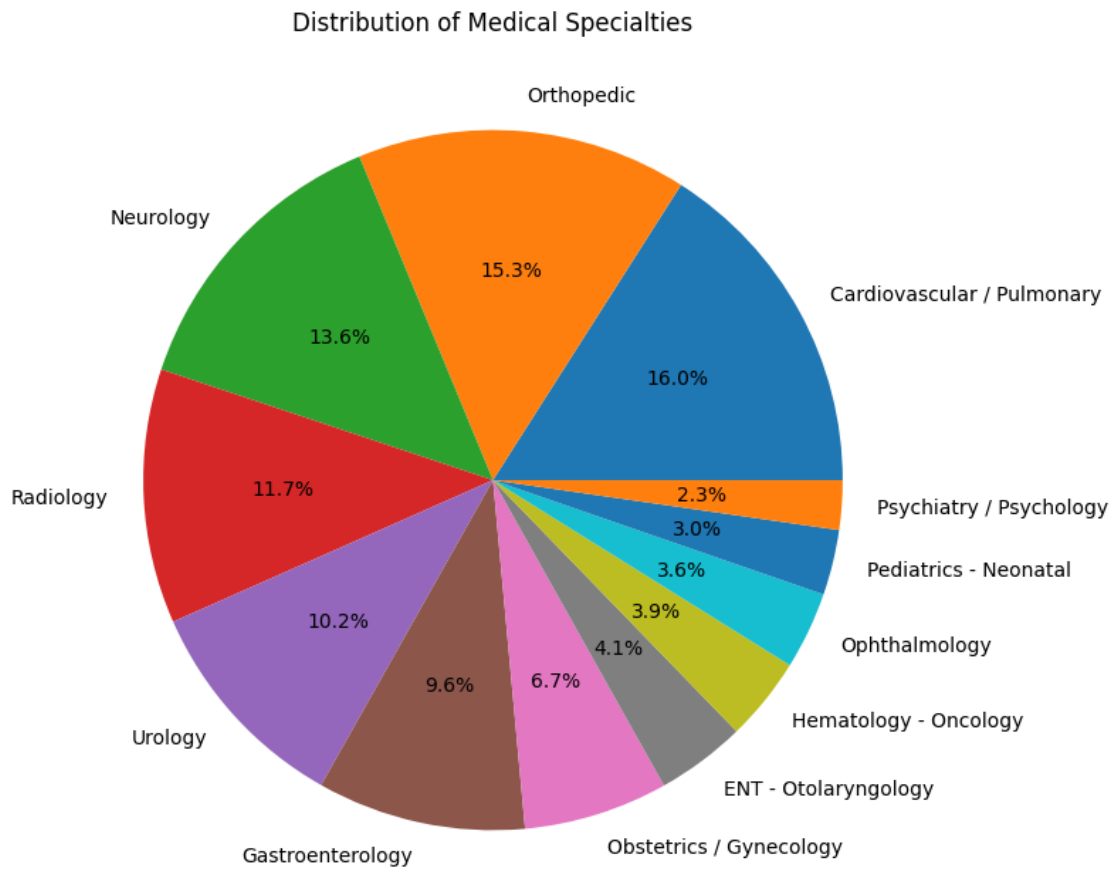
medical_specialty
Cardiovascular / Pulmonary    371
Orthopedic                    355
Neurology                     317
Radiology                     273
Urology                       237
Gastroenterology              224

```

Obstetrics / Gynecology	155
ENT - Otolaryngology	96
Hematology - Oncology	90
Ophthalmology	83
Pediatrics - Neonatal	70
Psychiatry / Psychology	53

Name: count, dtype: int64

```
[43]: # Plotting a pie chart
df['medical_specialty'].value_counts().plot(kind='pie', autopct='%1.1f%%',
figsize=(10, 8))
plt.title('Distribution of Medical Specialties')
plt.ylabel('')
plt.show()
```



Bar graph of finalized medical specialty distribution

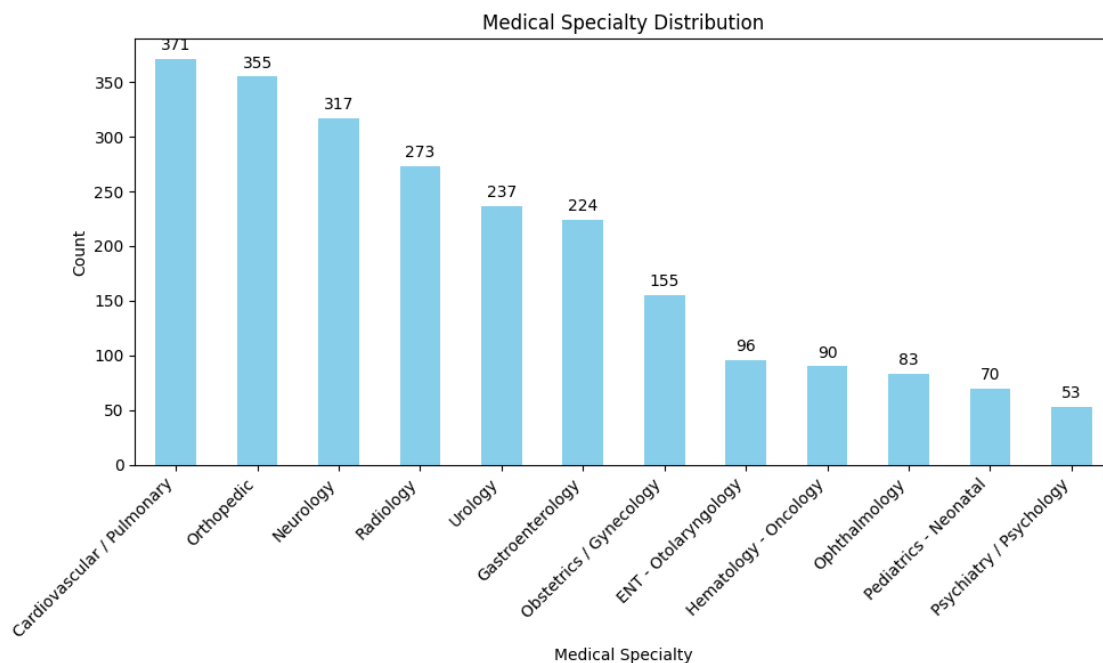
```
[44]: # Calculate the counts of each medical specialty
medical_specialty_counts = df['medical_specialty'].value_counts()

# Plotting the bar graph
plt.figure(figsize=(10, 6)) # Set the figure size
bars = medical_specialty_counts.plot(kind='bar', color='skyblue') # Plotting a
    ↪ bar chart with skyblue color

plt.title('Medical Specialty Distribution') # Adding a title to the plot
plt.xlabel('Medical Specialty') # Labeling the x-axis
plt.ylabel('Count') # Labeling the y-axis
plt.xticks(rotation=45, ha='right') # Rotating x-axis labels for better
    ↪ readability

# Adding the counts on top of each bar
for bar in bars.patches:
    yval = bar.get_height()
    plt.text(bar.get_x() + bar.get_width()/2, yval + 5, round(yval),
    ↪ ha='center', va='bottom')

plt.tight_layout()
plt.show() # Displaying the plot
```



```
[45]: # Calculate total word count of preprocessed transcriptions
```



```

total_word_count_preprocessing = df['preprocessed_transcription'].str.split().
    ↪str.len().sum()

# Print the total word count after preprocessing
print(f'The word count of transcription after Data Preprocessing is:␣
    ↪{int(total_word_count_preprocessing)}')

# Calculate percentage reduction in word count after preprocessing
percentage_reduction = round((total_word_count -
    ↪total_word_count_preprocessing) / total_word_count * 100, 2)

# Print the percentage reduction
print(f'{percentage_reduction}% less word')

```

The word count of transcription after Data Preprocessing is: 629262  
 73.86% less word

```

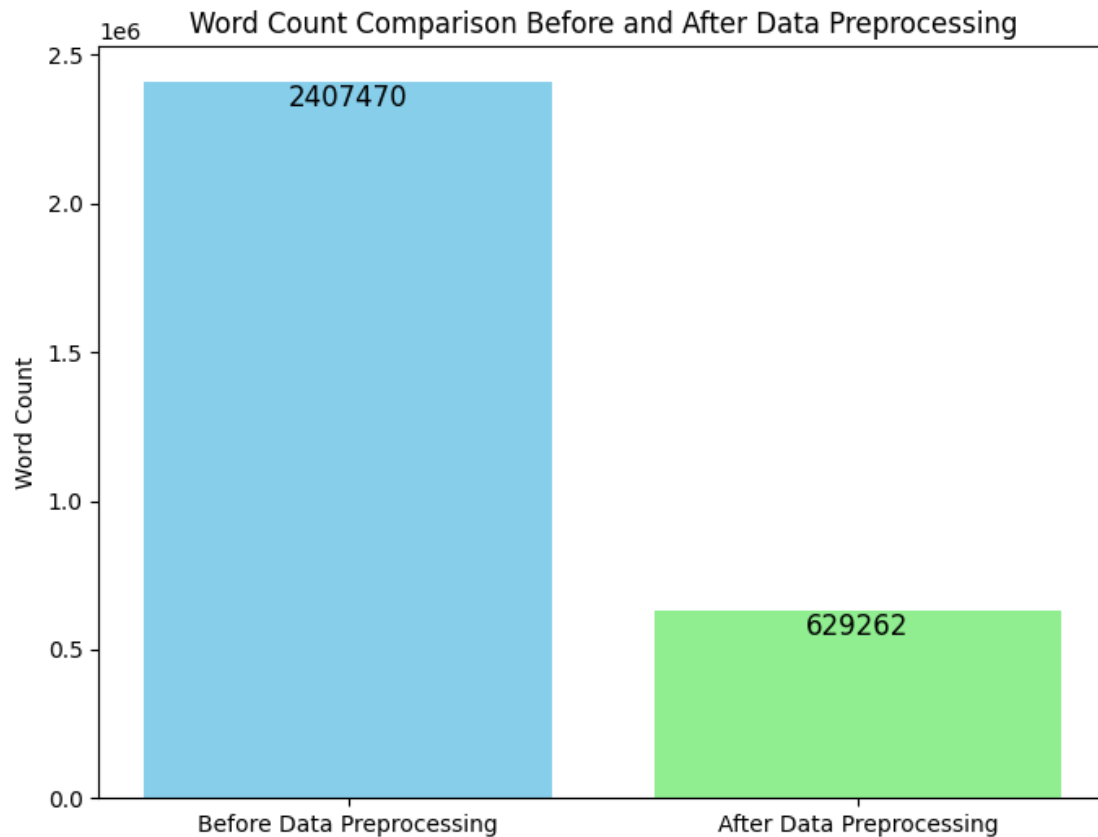
[46]: # Word counts before and after normalization
word_counts = [total_word_count, total_word_count_preprocessing]
labels = ['Before Data Preprocessing', 'After Data Preprocessing']

# Plotting
plt.figure(figsize=(8, 6))
bars = plt.bar(labels, word_counts, color=['skyblue', 'lightgreen'])
plt.title('Word Count Comparison Before and After Data Preprocessing')
plt.ylabel('Word Count')

# Adding the word count on top of each bar
for bar, count in zip(bars, word_counts):
    plt.text(bar.get_x() + bar.get_width()/2,
             bar.get_height() - 100000, # Adjusting the height of the text
             f'{count}',
             ha='center',
             va='bottom',
             fontsize=12,
             color='black')

plt.show()

```



#Label Encoding and Flattening

```
[47]: from sklearn import preprocessing

# Initialize LabelEncoder
le = preprocessing.LabelEncoder()

# Fit LabelEncoder with unique values from 'medical_specialty' and transform to
# numerical labels
le.fit(df['medical_specialty'])

# Encode 'medical_specialty' column with numerical labels
df['encoded_target'] = le.transform(df['medical_specialty'])

# Display the first few rows of the DataFrame with encoded labels
df.head()
```

```
[47]: preprocessed_transcription \
3    cardiovascular, pulmonary, mmode, doppler, aortic, valve, atrial,
enlargement, diastolic, function, ejection, fraction, mitral, mitral, valve,
```

pericardial, effusion, pulmonary, valve, regurgitation, systolic, function, tricuspid, tricuspid, valve, nor...

4 cardiovascular, pulmonary, doppler, echocardiogram, annular, aortic, root, aortic, valve, atrial, atrium, calcification, cavity, ejection, fraction, mitral, obliteration, outflow, regurgitation, relaxation, pattern, stenosis, systolic, function, tricu...

7 cardiovascular, pulmonary, echocardiogram, cardiac, function, doppler, echocardiogram, multiple, view, aortic, valve, coronary, artery, descending, aorta, great, vessel, heart, hypertrophy, interatrial, septum, intracardiac, pericardial, effusion, tri...

9 cardiovascular, pulmonary, ejection, fraction, lv, systolic, function, cardiac, chamber, regurgitation, tricuspid, normal, lv, systolic, function, normal, lv, systolic, ejection, fraction, estimated, normal, lv, lv, systolic, systolic, function, funct...

11 cardiovascular, pulmonary, study, doppler, tricuspid, regurgitation, heart, pressure, stenosis, ventricular, heart, ventricle, tricuspid, regurgitation, study, mild, aortic, stenosis, widely, calcified, minimally, restricted, mild, left, ventricular,...

	medical_specialty	encoded_target
3	Cardiovascular / Pulmonary	0
4	Cardiovascular / Pulmonary	0
7	Cardiovascular / Pulmonary	0
9	Cardiovascular / Pulmonary	0
11	Cardiovascular / Pulmonary	0

```
[48]: df.shape
```

```
[48]: (2324, 3)
```

```
[49]: # function to flatten one list
def flat_list(unflat_list):
    flattened = [item for sublist in unflat_list for item in sublist]
    return flattened

def to_list(df, attribute):
    # Select the normalised transcript column
    df_transcription = df[[attribute]]
    # To convert the attribute into list format, but it has inner list. So it
    ↪ cannot put into the CountVectorizer
    unflat_list_transcription = df_transcription.values.tolist()
    # Let's use back the function defined above, "flat_list", to flatten the
    ↪ list
    flat_list_transcription = flat_list(unflat_list_transcription)
    return flat_list_transcription
flat_list_transcription = to_list(df, 'preprocessed_transcription')
```

```
#Baseline Models (Traditional ML) Development
#ML model with Bag of Words Representation and PCA
```

```
[50]: # Step 1: Bag of Words Representation
vectorizer = CountVectorizer()
X = vectorizer.fit_transform(df['preprocessed_transcription'])

# Step 2: Dimension Reduction with PCA
pca = PCA(n_components=100) # You can adjust the number of components as needed
X_pca = pca.fit_transform(X.toarray())

# Step 3: Handling Imbalanced Data with SMOTE
smote = SMOTE()
X_resampled, y_resampled = smote.fit_resample(X_pca, df['encoded_target'])

# Step 4: Splitting Data into Train and Test Sets
X_train, X_test, y_train, y_test = train_test_split(X_resampled, y_resampled,
    ↪test_size=0.2, random_state=42)

# list of category names from the "medical_specialty" column
category_names = df['medical_specialty'].unique().tolist()

# Step 5: Model Building and Evaluation
classifiers = {
    "Random Forest": RandomForestClassifier(),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(),
    "Logistic Regression": LogisticRegression()
}

for name, classifier in classifiers.items():
    print(f"Training and evaluating {name}...")
    classifier.fit(X_train, y_train)
    y_pred = classifier.predict(X_test)
    print(f"Results for {name}:")
    print(classification_report(y_test, y_pred, target_names=category_names))
    print("\n" + "="*60 + "\n")
```

Training and evaluating Random Forest...

Results for Random Forest:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.65	0.69	0.67	74
Neurology	0.91	0.94	0.92	77
Urology	0.83	0.76	0.79	76
Radiology	0.82	0.87	0.85	79
Psychiatry / Psychology	0.43	0.44	0.43	66

Pediatrics - Neonatal	0.84	0.93	0.88	74
Orthopedic	0.93	1.00	0.96	55
Ophthalmology	0.72	0.67	0.69	79
Obstetrics / Gynecology	0.78	0.80	0.79	71
Hematology - Oncology	0.97	0.99	0.98	73
Gastroenterology	0.28	0.24	0.26	87
ENT - Otolaryngology	0.84	0.79	0.81	80
accuracy			0.75	891
macro avg	0.75	0.76	0.75	891
weighted avg	0.74	0.75	0.75	891

=====

Training and evaluating Support Vector Machine...

Results for Support Vector Machine:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.77	0.73	0.75	74
Neurology	0.91	0.83	0.87	77
Urology	0.88	0.78	0.83	76
Radiology	0.77	0.89	0.82	79
Psychiatry / Psychology	0.54	0.55	0.54	66
Pediatrics - Neonatal	0.89	0.89	0.89	74
Orthopedic	0.95	1.00	0.97	55
Ophthalmology	0.66	0.65	0.65	79
Obstetrics / Gynecology	0.81	0.77	0.79	71
Hematology - Oncology	0.93	0.95	0.94	73
Gastroenterology	0.44	0.54	0.49	87
ENT - Otolaryngology	0.91	0.79	0.85	80
accuracy			0.77	891
macro avg	0.79	0.78	0.78	891
weighted avg	0.78	0.77	0.78	891

=====

Training and evaluating XGBoost...

Results for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.69	0.69	0.69	74
Neurology	0.93	0.92	0.93	77
Urology	0.91	0.79	0.85	76
Radiology	0.83	0.94	0.88	79
Psychiatry / Psychology	0.43	0.45	0.44	66

Pediatrics - Neonatal	0.86	0.93	0.90	74
Orthopedic	0.96	1.00	0.98	55
Ophthalmology	0.70	0.62	0.66	79
Obstetrics / Gynecology	0.79	0.83	0.81	71
Hematology - Oncology	0.97	0.99	0.98	73
Gastroenterology	0.37	0.33	0.35	87
ENT - Otolaryngology	0.81	0.82	0.82	80
accuracy			0.77	891
macro avg	0.77	0.78	0.77	891
weighted avg	0.77	0.77	0.77	891

=====

Training and evaluating Logistic Regression...

Results for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.74	0.73	0.73	74
Neurology	0.92	0.90	0.91	77
Urology	0.82	0.86	0.84	76
Radiology	0.80	0.87	0.84	79
Psychiatry / Psychology	0.52	0.47	0.49	66
Pediatrics - Neonatal	0.86	0.92	0.89	74
Orthopedic	0.96	1.00	0.98	55
Ophthalmology	0.72	0.66	0.69	79
Obstetrics / Gynecology	0.81	0.79	0.80	71
Hematology - Oncology	0.95	1.00	0.97	73
Gastroenterology	0.50	0.48	0.49	87
ENT - Otolaryngology	0.84	0.84	0.84	80
accuracy			0.79	891
macro avg	0.79	0.79	0.79	891
weighted avg	0.78	0.79	0.78	891

=====

#ML model with TF-IDF Word Representation with PCA

```
[51]: # Step 1: TF-IDF Word Representation

# Initialize TF-IDF vectorizer
tfidf_vectorizer = TfidfVectorizer()

# Apply TF-IDF vectorization to the text data
```

```

X_tfidf = tfidf_vectorizer.fit_transform(flat_list_transcription)

# Step 2: PCA for Dimension Reduction

# Initialize PCA with 95% variance explained
pca = PCA(n_components=0.95)

# Perform PCA on the TF-IDF transformed data
X_pca = pca.fit_transform(X_tfidf.toarray())

# Calculate explained variance ratio and cumulative explained variance
explained_variance = pca.explained_variance_ratio_
cumulative_explained_variance = explained_variance.cumsum()

# Visualizing PCA results

# Create a dataframe with the first two principal components and labels
pca_df = pd.DataFrame(data=X_pca, columns=[f'PC{i+1}' for i in range(X_pca.
    ↪shape[1])])
pca_df['label'] = df['medical_specialty'] # assuming 'medical_specialty' is
    ↪the label

# Plot the first two principal components
plt.figure(figsize=(14, 8))
sns.scatterplot(
    x='PC1', y='PC2',
    hue='label',
    palette=sns.color_palette("hsv", len(pca_df['label'].unique())),
    data=pca_df,
    legend="full",
    alpha=0.6
)
plt.title('PCA - First two principal components')
plt.show()

# Plot the explained variance ratio with secondary y-axis for cumulative
    ↪explained variance
fig, ax1 = plt.subplots(figsize=(12, 6))

color = 'tab:blue'
ax1.set_xlabel('Principal components')
ax1.set_ylabel('Individual explained variance ratio', color=color)
ax1.bar(range(1, len(explained_variance) + 1), explained_variance, alpha=0.7,
    ↪align='center', color=color)
ax1.tick_params(axis='y', labelcolor=color)

ax2 = ax1.twinx() # instantiate a second axes that shares the same x-axis

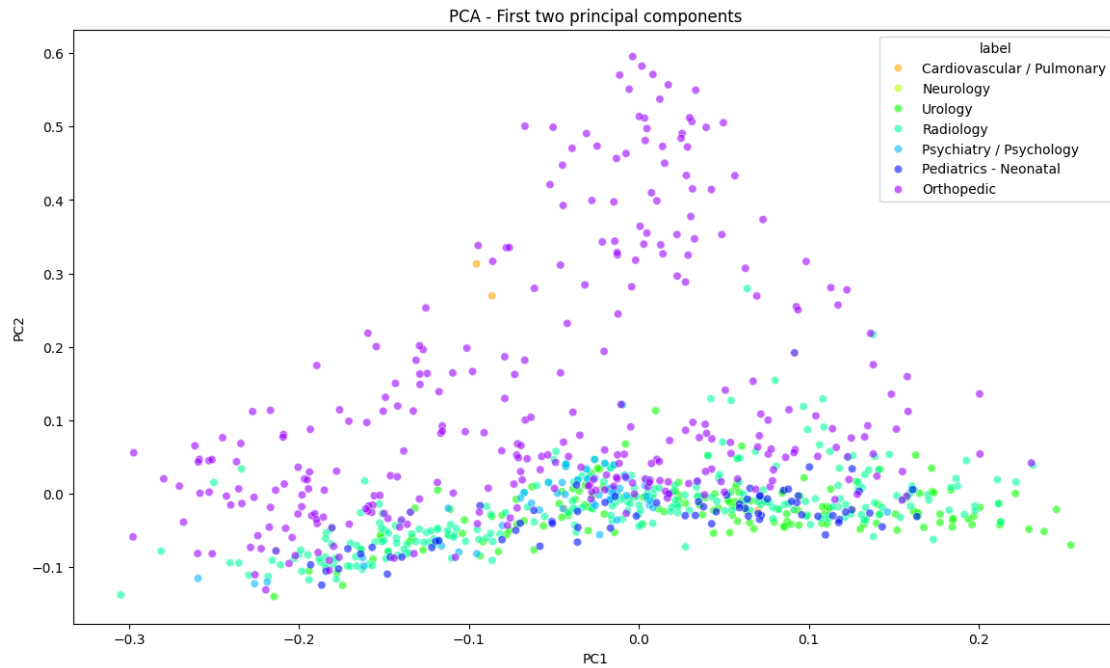
```

```

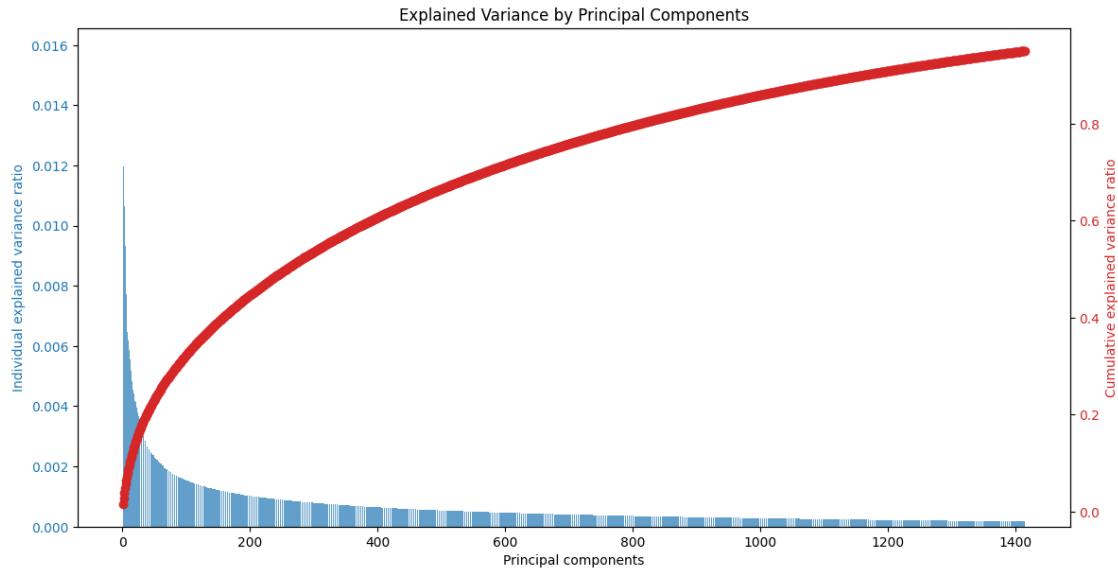
color = 'tab:red'
ax2.set_ylabel('Cumulative explained variance ratio', color=color)
ax2.plot(range(1, len(cumulative_explained_variance) + 1),
         cumulative_explained_variance, color=color, marker='o')
ax2.tick_params(axis='y', labelcolor=color)

fig.tight_layout()
plt.title('Explained Variance by Principal Components')
plt.show()

```







```
[52]: # Step 3: Handling Imbalanced Data with SMOTE

# Apply SMOTE for oversampling
smote = SMOTE()
X_resampled, y_resampled = smote.fit_resample(X_pca, df['medical_specialty'])
    ↪ # assuming 'medical_specialty' is the label

# Step 4: Splitting Data into Train and Test Sets (80% train, 20% test)

# Split data into train and test sets
X_train, X_test, y_train, y_test = train_test_split(X_resampled, y_resampled,
    ↪ test_size=0.2, random_state=42)

# Step 5: Building and Evaluating Classifiers

# Define classifiers to evaluate
classifiers = {
    'Random Forest': RandomForestClassifier(),
    'Support Vector Machine': SVC(),
    'XGBoost': XGBClassifier(),
    'Logistic Regression': LogisticRegression()
}

# Initialize LabelEncoder
label_encoder = LabelEncoder()

# Encode categorical labels
y_train_encoded = label_encoder.fit_transform(y_train)
```

```

y_test_encoded = label_encoder.transform(y_test)

# Train and evaluate each classifier
for name, clf in classifiers.items():
    clf.fit(X_train, y_train_encoded)
    y_pred = clf.predict(X_test)
    print(f"Classifier: {name}")
    print(classification_report(y_test_encoded, y_pred,
    ↪target_names=label_encoder.classes_))
    print("\n" + "="*60 + "\n")

```

Classifier: Random Forest

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.64	0.70	0.67	74
ENT - Otolaryngology	0.94	0.95	0.94	77
Gastroenterology	0.85	0.83	0.84	76
Hematology - Oncology	0.89	0.89	0.89	79
Neurology	0.49	0.41	0.45	66
Obstetrics / Gynecology	0.82	0.95	0.88	74
Ophthalmology	0.98	1.00	0.99	55
Orthopedic	0.72	0.70	0.71	79
Pediatrics - Neonatal	0.85	0.87	0.86	71
Psychiatry / Psychology	1.00	0.97	0.99	73
Radiology	0.26	0.24	0.25	87
Urology	0.88	0.89	0.88	80
accuracy			0.77	891
macro avg	0.78	0.78	0.78	891
weighted avg	0.77	0.77	0.77	891

=====

Classifier: Support Vector Machine

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.66	0.76	0.70	74
ENT - Otolaryngology	0.93	0.88	0.91	77
Gastroenterology	0.89	0.84	0.86	76
Hematology - Oncology	0.89	0.92	0.91	79
Neurology	0.54	0.55	0.54	66
Obstetrics / Gynecology	0.84	0.92	0.88	74
Ophthalmology	0.96	1.00	0.98	55
Orthopedic	0.79	0.71	0.75	79
Pediatrics - Neonatal	0.87	0.92	0.89	71
Psychiatry / Psychology	1.00	0.97	0.99	73

Radiology	0.29	0.25	0.27	87
Urology	0.88	0.89	0.88	80
accuracy			0.79	891
macro avg	0.79	0.80	0.80	891
weighted avg	0.79	0.79	0.79	891

=====

Classifier: XGBoost

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.67	0.76	0.71	74
ENT - Otolaryngology	0.92	0.94	0.93	77
Gastroenterology	0.87	0.88	0.88	76
Hematology - Oncology	0.88	0.89	0.88	79
Neurology	0.51	0.45	0.48	66
Obstetrics / Gynecology	0.86	0.91	0.88	74
Ophthalmology	0.98	1.00	0.99	55
Orthopedic	0.74	0.70	0.72	79
Pediatrics - Neonatal	0.85	0.94	0.89	71
Psychiatry / Psychology	0.99	0.97	0.98	73
Radiology	0.40	0.36	0.38	87
Urology	0.91	0.88	0.89	80
accuracy			0.80	891
macro avg	0.80	0.81	0.80	891
weighted avg	0.79	0.80	0.79	891

=====

Classifier: Logistic Regression

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.73	0.82	0.78	74
ENT - Otolaryngology	0.93	0.96	0.94	77
Gastroenterology	0.89	0.88	0.89	76
Hematology - Oncology	0.90	0.94	0.92	79
Neurology	0.64	0.58	0.61	66
Obstetrics / Gynecology	0.86	0.95	0.90	74
Ophthalmology	1.00	1.00	1.00	55
Orthopedic	0.79	0.71	0.75	79
Pediatrics - Neonatal	0.90	0.90	0.90	71
Psychiatry / Psychology	1.00	0.99	0.99	73
Radiology	0.42	0.39	0.41	87
Urology	0.90	0.93	0.91	80

accuracy			0.83	891
macro avg	0.83	0.84	0.83	891
weighted avg	0.82	0.83	0.83	891

=====

#AUC and ROC of TF-IDF Word Representation with PCA ( Logistic Regression)

```
[53]: # Number of classes
n_classes = len(label_encoder.classes_)

# Binarize the output
y_test_binarized = label_binarize(y_test_encoded, classes=np.arange(n_classes))
y_pred_binarized = label_binarize(y_pred, classes=np.arange(n_classes))

# Compute ROC curve and ROC area for each class
fpr = dict()
tpr = dict()
roc_auc = dict()
for i in range(n_classes):
    fpr[i], tpr[i], _ = roc_curve(y_test_binarized[:, i], y_pred_binarized[:, i])
    roc_auc[i] = auc(fpr[i], tpr[i])

# Compute micro-average ROC curve and ROC area
fpr["micro"], tpr["micro"], _ = roc_curve(y_test_binarized.ravel(),
    y_pred_binarized.ravel())
roc_auc["micro"] = auc(fpr["micro"], tpr["micro"])

# Plot all ROC curves
plt.figure(figsize=(14, 10))

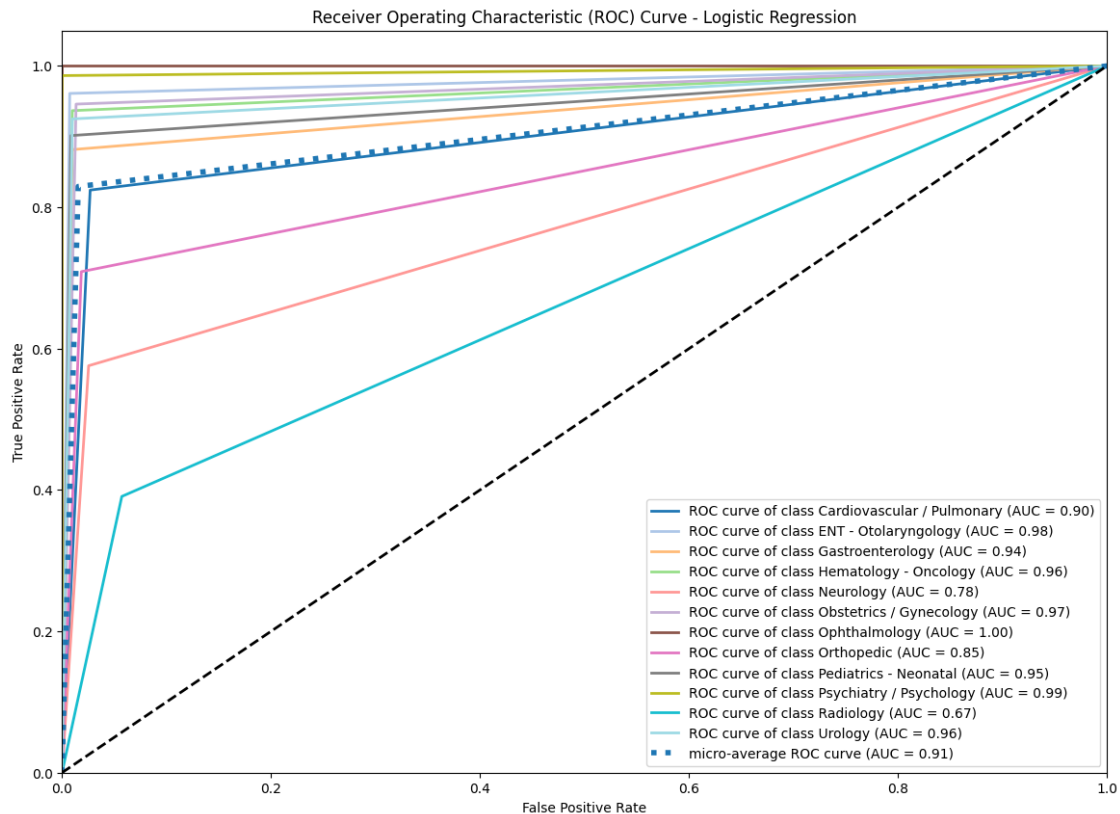
colors = plt.cm.get_cmap('tab20', n_classes)

for i, color in zip(range(n_classes), colors.colors):
    plt.plot(fpr[i], tpr[i], color=color, lw=2,
        label=f'ROC curve of class {label_encoder.classes_[i]} (AUC = {roc_auc[i]:0.2f})')

plt.plot(fpr["micro"], tpr["micro"], linestyle=':', linewidth=4,
        label=f'micro-average ROC curve (AUC = {roc_auc["micro"]:0.2f})')

plt.plot([0, 1], [0, 1], 'k--', lw=2)
plt.xlim([0.0, 1.0])
plt.ylim([0.0, 1.05])
```

```
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('Receiver Operating Characteristic (ROC) Curve - Logistic Regression')
plt.legend(loc="lower right")
plt.show()
```



#ML model with Bag-of-n-grams (1,1) Word Representation and TruncatedSVD

```
[54]: # Step 1: N-gram Representation
vectorizer = CountVectorizer(ngram_range=(1, 1))
X = vectorizer.fit_transform(flat_list_transcription)
y = df['encoded_target']

# Step 2: Imbalanced Data Handling
smote = SMOTE(random_state=42)
X_resampled, y_resampled = smote.fit_resample(X, y)

# Step 3: Dimensionality Reduction with TruncatedSVD
svd = TruncatedSVD(n_components=100)
X_svd = svd.fit_transform(X_resampled)
```

```

# Step 4: Splitting Data
X_train, X_test, y_train, y_test = train_test_split(X_svd, y_resampled,
    ↪test_size=0.2, random_state=42)

# Extract unique category names from the "medical_specialty" column
category_names = df['medical_specialty'].unique()

# Step 5: Model Building
models = {
    "Random Forest": RandomForestClassifier(),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(),
    "Logistic Regression": LogisticRegression()
}

for name, model in models.items():
    model.fit(X_train, y_train)
    y_pred = model.predict(X_test)
    report = classification_report(y_test, y_pred, target_names=category_names)
    print(f"Classification Report for {name}:\n{report}")
    print("\n" + "="*60 + "\n")

```

Classification Report for Random Forest:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.61	0.69	0.65	74
Neurology	0.91	0.96	0.94	77
Urology	0.82	0.78	0.80	76
Radiology	0.74	0.80	0.77	79
Psychiatry / Psychology	0.42	0.38	0.40	66
Pediatrics - Neonatal	0.82	0.84	0.83	74
Orthopedic	0.96	1.00	0.98	55
Ophthalmology	0.69	0.65	0.67	79
Obstetrics / Gynecology	0.81	0.90	0.85	71
Hematology - Oncology	0.96	0.99	0.97	73
Gastroenterology	0.25	0.22	0.23	87
ENT - Otolaryngology	0.82	0.75	0.78	80
accuracy			0.74	891
macro avg	0.73	0.75	0.74	891
weighted avg	0.73	0.74	0.73	891

=====  
Classification Report for Support Vector Machine:

	precision	recall	f1-score	support
--	-----------	--------	----------	---------

Cardiovascular / Pulmonary	0.65	0.72	0.68	74
Neurology	0.92	0.86	0.89	77
Urology	0.86	0.67	0.76	76
Radiology	0.72	0.90	0.80	79
Psychiatry / Psychology	0.59	0.55	0.57	66
Pediatrics - Neonatal	0.80	0.74	0.77	74
Orthopedic	0.98	1.00	0.99	55
Ophthalmology	0.75	0.62	0.68	79
Obstetrics / Gynecology	0.54	0.82	0.65	71
Hematology - Oncology	0.93	0.95	0.94	73
Gastroenterology	0.40	0.40	0.40	87
ENT - Otolaryngology	0.93	0.70	0.80	80
accuracy			0.73	891
macro avg	0.76	0.74	0.74	891
weighted avg	0.75	0.73	0.74	891

=====

#### Classification Report for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.62	0.72	0.66	74
Neurology	0.91	0.95	0.93	77
Urology	0.82	0.79	0.81	76
Radiology	0.81	0.82	0.82	79
Psychiatry / Psychology	0.48	0.45	0.47	66
Pediatrics - Neonatal	0.83	0.85	0.84	74
Orthopedic	0.96	1.00	0.98	55
Ophthalmology	0.76	0.71	0.73	79
Obstetrics / Gynecology	0.83	0.85	0.84	71
Hematology - Oncology	0.97	0.99	0.98	73
Gastroenterology	0.28	0.28	0.28	87
ENT - Otolaryngology	0.83	0.74	0.78	80
accuracy			0.75	891
macro avg	0.76	0.76	0.76	891
weighted avg	0.75	0.75	0.75	891

=====

#### Classification Report for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.71	0.77	0.74	74

Neurology	0.90	0.92	0.91	77
Urology	0.85	0.88	0.86	76
Radiology	0.82	0.91	0.86	79
Psychiatry / Psychology	0.58	0.56	0.57	66
Pediatrics - Neonatal	0.88	0.89	0.89	74
Orthopedic	0.98	1.00	0.99	55
Ophthalmology	0.79	0.72	0.75	79
Obstetrics / Gynecology	0.90	0.87	0.89	71
Hematology - Oncology	0.91	0.99	0.95	73
Gastroenterology	0.46	0.40	0.43	87
ENT - Otolaryngology	0.88	0.81	0.84	80
accuracy			0.80	891
macro avg	0.80	0.81	0.81	891
weighted avg	0.80	0.80	0.80	891

=====

#ML model with Bag-of-n-grams (1,2) Word Representation and TruncatedSVD

```
[55]: # Step 1: N-gram Representation
vectorizer = CountVectorizer(ngram_range=(1, 2))
X = vectorizer.fit_transform(flat_list_transcription)
y = df['encoded_target']

# Step 2: Imbalanced Data Handling
smote = SMOTE(random_state=42)
X_resampled, y_resampled = smote.fit_resample(X, y)

# Step 3: Dimensionality Reduction with TruncatedSVD
svd = TruncatedSVD(n_components=100)
X_svd = svd.fit_transform(X_resampled)

# Step 4: Splitting Data
X_train, X_test, y_train, y_test = train_test_split(X_svd, y_resampled,
    ↪test_size=0.2, random_state=42)

# Extract unique category names from the "medical_specialty" column
category_names = df['medical_specialty'].unique()

# Step 5: Model Building
models = {
    "Random Forest": RandomForestClassifier(),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(),
    "Logistic Regression": LogisticRegression()
```



```

}

for name, model in models.items():
    model.fit(X_train, y_train)
    y_pred = model.predict(X_test)
    report = classification_report(y_test, y_pred, target_names=category_names)
    print(f"Classification Report for {name}:\n{report}")
    print("\n" + "="*60 + "\n")

```

Classification Report for Random Forest:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.56	0.66	0.61	74
Neurology	0.92	0.92	0.92	77
Urology	0.81	0.76	0.78	76
Radiology	0.81	0.82	0.82	79
Psychiatry / Psychology	0.49	0.42	0.46	66
Pediatrics - Neonatal	0.81	0.86	0.84	74
Orthopedic	0.95	1.00	0.97	55
Ophthalmology	0.71	0.68	0.70	79
Obstetrics / Gynecology	0.76	0.83	0.79	71
Hematology - Oncology	0.94	0.99	0.96	73
Gastroenterology	0.26	0.23	0.25	87
ENT - Otolaryngology	0.82	0.76	0.79	80
accuracy			0.74	891
macro avg	0.74	0.75	0.74	891
weighted avg	0.73	0.74	0.73	891

=====

Classification Report for Support Vector Machine:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.65	0.72	0.68	74
Neurology	0.93	0.86	0.89	77
Urology	0.85	0.70	0.77	76
Radiology	0.72	0.86	0.79	79
Psychiatry / Psychology	0.62	0.56	0.59	66
Pediatrics - Neonatal	0.81	0.77	0.79	74
Orthopedic	0.92	1.00	0.96	55
Ophthalmology	0.71	0.63	0.67	79
Obstetrics / Gynecology	0.56	0.80	0.66	71
Hematology - Oncology	0.91	0.95	0.93	73
Gastroenterology	0.42	0.40	0.41	87
ENT - Otolaryngology	0.90	0.69	0.78	80

accuracy			0.74	891
macro avg	0.75	0.74	0.74	891
weighted avg	0.75	0.74	0.74	891

=====

Classification Report for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.61	0.69	0.65	74
Neurology	0.92	0.90	0.91	77
Urology	0.81	0.82	0.81	76
Radiology	0.81	0.81	0.81	79
Psychiatry / Psychology	0.52	0.45	0.48	66
Pediatrics - Neonatal	0.82	0.85	0.83	74
Orthopedic	0.95	1.00	0.97	55
Ophthalmology	0.72	0.70	0.71	79
Obstetrics / Gynecology	0.82	0.82	0.82	71
Hematology - Oncology	0.96	0.99	0.97	73
Gastroenterology	0.30	0.29	0.29	87
ENT - Otolaryngology	0.79	0.76	0.78	80
accuracy			0.75	891
macro avg	0.75	0.76	0.75	891
weighted avg	0.74	0.75	0.74	891

=====

Classification Report for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.70	0.82	0.76	74
Neurology	0.90	0.94	0.92	77
Urology	0.81	0.86	0.83	76
Radiology	0.84	0.91	0.87	79
Psychiatry / Psychology	0.56	0.53	0.55	66
Pediatrics - Neonatal	0.88	0.89	0.89	74
Orthopedic	0.98	1.00	0.99	55
Ophthalmology	0.76	0.71	0.73	79
Obstetrics / Gynecology	0.83	0.82	0.82	71
Hematology - Oncology	0.88	0.99	0.93	73
Gastroenterology	0.49	0.38	0.43	87
ENT - Otolaryngology	0.92	0.81	0.86	80
accuracy			0.80	891

macro avg	0.80	0.80	0.80	891
weighted avg	0.79	0.80	0.79	891

=====

#ML model with Bag-of-n-grams (2,2) Word Representation and TruncatedSVD

```
[56]: # Step 1: N-gram Representation
vectorizer = CountVectorizer(ngram_range=(2, 2))
X = vectorizer.fit_transform(flat_list_transcription)
y = df['encoded_target']

# Step 2: Imbalanced Data Handling
smote = SMOTE(random_state=42)
X_resampled, y_resampled = smote.fit_resample(X, y)

# Step 3: Dimensionality Reduction with TruncatedSVD
svd = TruncatedSVD(n_components=100, random_state=42)
X_svd = svd.fit_transform(X_resampled)

# Step 4: Splitting Data
X_train, X_test, y_train, y_test = train_test_split(X_svd, y_resampled,
    ↪test_size=0.2, random_state=42)

# Extract unique category names from the "medical_specialty" column
# Assuming df['medical_specialty'] contains the actual category names
    ↪corresponding to y
category_names = df['medical_specialty'].unique()

# Step 5: Model Building
models = {
    "Random Forest": RandomForestClassifier(random_state=42),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(random_state=42),
    "Logistic Regression": LogisticRegression(random_state=42)
}

for name, model in models.items():
    model.fit(X_train, y_train)
    y_pred = model.predict(X_test)
    report = classification_report(y_test, y_pred, target_names=category_names)
    print(f"Classification Report for {name}:\n{report}")
    print("\n" + "="*60 + "\n")
```

Classification Report for Random Forest:

	precision	recall	f1-score	support
--	-----------	--------	----------	---------

Cardiovascular / Pulmonary	0.61	0.69	0.65	74
Neurology	0.86	0.90	0.88	77
Urology	0.63	0.68	0.65	76
Radiology	0.73	0.77	0.75	79
Psychiatry / Psychology	0.47	0.41	0.44	66
Pediatrics - Neonatal	0.80	0.82	0.81	74
Orthopedic	0.88	0.95	0.91	55
Ophthalmology	0.71	0.63	0.67	79
Obstetrics / Gynecology	0.76	0.83	0.79	71
Hematology - Oncology	0.80	0.96	0.87	73
Gastroenterology	0.31	0.28	0.29	87
ENT - Otolaryngology	0.76	0.53	0.62	80
accuracy			0.69	891
macro avg	0.69	0.70	0.70	891
weighted avg	0.69	0.69	0.69	891

=====

#### Classification Report for Support Vector Machine:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.58	0.64	0.61	74
Neurology	0.85	0.68	0.75	77
Urology	0.49	0.26	0.34	76
Radiology	0.65	0.42	0.51	79
Psychiatry / Psychology	0.53	0.42	0.47	66
Pediatrics - Neonatal	0.84	0.51	0.64	74
Orthopedic	0.58	0.76	0.66	55
Ophthalmology	0.58	0.56	0.57	79
Obstetrics / Gynecology	0.21	0.90	0.34	71
Hematology - Oncology	0.85	0.30	0.44	73
Gastroenterology	0.34	0.18	0.24	87
ENT - Otolaryngology	0.86	0.31	0.46	80
accuracy			0.48	891
macro avg	0.61	0.50	0.50	891
weighted avg	0.61	0.48	0.50	891

=====

#### Classification Report for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.64	0.69	0.66	74

Neurology	0.91	0.90	0.90	77
Urology	0.68	0.68	0.68	76
Radiology	0.80	0.81	0.81	79
Psychiatry / Psychology	0.40	0.38	0.39	66
Pediatrics - Neonatal	0.78	0.84	0.81	74
Orthopedic	0.94	0.93	0.94	55
Ophthalmology	0.67	0.65	0.66	79
Obstetrics / Gynecology	0.77	0.85	0.81	71
Hematology - Oncology	0.79	0.97	0.87	73
Gastroenterology	0.31	0.30	0.31	87
ENT - Otolaryngology	0.80	0.56	0.66	80
accuracy			0.70	891
macro avg	0.71	0.71	0.71	891
weighted avg	0.70	0.70	0.70	891

=====

#### Classification Report for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.72	0.72	0.72	74
Neurology	0.91	0.94	0.92	77
Urology	0.60	0.49	0.54	76
Radiology	0.78	0.80	0.79	79
Psychiatry / Psychology	0.53	0.47	0.50	66
Pediatrics - Neonatal	0.84	0.73	0.78	74
Orthopedic	0.81	0.85	0.83	55
Ophthalmology	0.74	0.68	0.71	79
Obstetrics / Gynecology	0.52	0.80	0.63	71
Hematology - Oncology	0.63	0.95	0.76	73
Gastroenterology	0.48	0.33	0.39	87
ENT - Otolaryngology	0.81	0.62	0.70	80
accuracy			0.69	891
macro avg	0.70	0.70	0.69	891
weighted avg	0.70	0.69	0.68	891

=====

#ML model with Bag-of-n-grams (2,3) Word Representation and TruncatedSVD

```
[57]: # Step 1: N-gram Representation
vectorizer = CountVectorizer(ngram_range=(2, 3))
X = vectorizer.fit_transform(flat_list_transcription)
```

```

y = df['encoded_target']

# Step 2: Imbalanced Data Handling
smote = SMOTE(random_state=42)
X_resampled, y_resampled = smote.fit_resample(X, y)

# Step 3: Dimensionality Reduction with TruncatedSVD
svd = TruncatedSVD(n_components=100)
X_svd = svd.fit_transform(X_resampled)

# Step 4: Splitting Data
X_train, X_test, y_train, y_test = train_test_split(X_svd, y_resampled,
    ↪test_size=0.2, random_state=42)

# Extract unique category names from the "medical_specialty" column
# Create a mapping from encoded target to category name
encoded_target_to_category = dict(zip(df['encoded_target'],
    ↪df['medical_specialty']))

# Ensure the category names are in the correct order of the encoded targets
category_names = [encoded_target_to_category[i] for i in
    ↪sorted(encoded_target_to_category)]

# Step 5: Model Building
models = {
    "Random Forest": RandomForestClassifier(),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(),
    "Logistic Regression": LogisticRegression()
}

for name, model in models.items():
    model.fit(X_train, y_train)
    y_pred = model.predict(X_test)
    report = classification_report(y_test, y_pred, target_names=category_names)
    print(f"Classification Report for {name}: \n{report}")
    print("\n" + "="*60 + "\n")

```

Classification Report for Random Forest:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.59	0.66	0.62	74
ENT - Otolaryngology	0.84	0.86	0.85	77
Gastroenterology	0.58	0.63	0.60	76
Hematology - Oncology	0.68	0.70	0.69	79
Neurology	0.43	0.39	0.41	66
Obstetrics / Gynecology	0.77	0.72	0.74	74

Ophthalmology	0.87	0.95	0.90	55
Orthopedic	0.69	0.62	0.65	79
Pediatrics - Neonatal	0.78	0.79	0.78	71
Psychiatry / Psychology	0.75	0.95	0.84	73
Radiology	0.28	0.24	0.26	87
Urology	0.64	0.53	0.58	80
accuracy			0.66	891
macro avg	0.66	0.67	0.66	891
weighted avg	0.65	0.66	0.65	891

=====

Classification Report for Support Vector Machine:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.56	0.61	0.58	74
ENT - Otolaryngology	0.83	0.52	0.64	77
Gastroenterology	0.35	0.28	0.31	76
Hematology - Oncology	0.65	0.38	0.48	79
Neurology	0.53	0.42	0.47	66
Obstetrics / Gynecology	0.83	0.41	0.55	74
Ophthalmology	0.51	0.65	0.58	55
Orthopedic	0.62	0.57	0.59	79
Pediatrics - Neonatal	0.19	0.90	0.31	71
Psychiatry / Psychology	0.76	0.26	0.39	73
Radiology	0.30	0.11	0.17	87
Urology	0.88	0.26	0.40	80
accuracy			0.44	891
macro avg	0.58	0.45	0.46	891
weighted avg	0.59	0.44	0.45	891

=====

Classification Report for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.69	0.73	0.71	74
ENT - Otolaryngology	0.86	0.87	0.86	77
Gastroenterology	0.59	0.67	0.63	76
Hematology - Oncology	0.78	0.75	0.76	79
Neurology	0.41	0.39	0.40	66
Obstetrics / Gynecology	0.82	0.80	0.81	74
Ophthalmology	0.98	0.96	0.97	55
Orthopedic	0.67	0.63	0.65	79

Pediatrics - Neonatal	0.75	0.82	0.78	71
Psychiatry / Psychology	0.79	0.95	0.86	73
Radiology	0.35	0.32	0.33	87
Urology	0.76	0.60	0.67	80
accuracy			0.70	891
macro avg	0.70	0.71	0.70	891
weighted avg	0.70	0.70	0.70	891

=====

#### Classification Report for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.72	0.69	0.70	74
ENT - Otolaryngology	0.88	0.88	0.88	77
Gastroenterology	0.59	0.45	0.51	76
Hematology - Oncology	0.67	0.71	0.69	79
Neurology	0.51	0.45	0.48	66
Obstetrics / Gynecology	0.79	0.70	0.74	74
Ophthalmology	0.76	0.80	0.78	55
Orthopedic	0.71	0.62	0.66	79
Pediatrics - Neonatal	0.38	0.83	0.52	71
Psychiatry / Psychology	0.72	0.79	0.75	73
Radiology	0.42	0.26	0.32	87
Urology	0.80	0.59	0.68	80
accuracy			0.64	891
macro avg	0.66	0.65	0.64	891
weighted avg	0.66	0.64	0.64	891

=====

#ML model with Bag-of-n-grams (3,3) Word Representation and TruncatedSVD

```
[58]: # Step 1: N-gram Representation
vectorizer = CountVectorizer(ngram_range=(3, 3))
X = vectorizer.fit_transform(flat_list_transcription)
y = df['encoded_target']

# Step 2: Imbalanced Data Handling
smote = SMOTE(random_state=42)
X_resampled, y_resampled = smote.fit_resample(X, y)

# Step 3: Dimensionality Reduction with TruncatedSVD
```



```

svd = TruncatedSVD(n_components=100, random_state=42)
X_svd = svd.fit_transform(X_resampled)

# Step 4: Splitting Data
X_train, X_test, y_train, y_test = train_test_split(X_svd, y_resampled,
    ↪test_size=0.2, random_state=42)

# Extract unique category names from the "medical_specialty" column
# Create a mapping from encoded target to category name
encoded_target_to_category = dict(zip(df['encoded_target'],
    ↪df['medical_specialty']))

# Ensure the category names are in the correct order of the encoded targets
category_names = [encoded_target_to_category[i] for i in
    ↪sorted(encoded_target_to_category)]

# Step 5: Model Building
models = {
    "Random Forest": RandomForestClassifier(random_state=42),
    "Support Vector Machine": SVC(),
    "XGBoost": XGBClassifier(random_state=42),
    "Logistic Regression": LogisticRegression(random_state=42)
}

for name, model in models.items():
    model.fit(X_train, y_train)
    y_pred = model.predict(X_test)
    report = classification_report(y_test, y_pred, target_names=category_names)
    print(f"Classification Report for {name}:\n{report}")
    print("\n" + "="*60 + "\n")

```

Classification Report for Random Forest:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.49	0.46	0.47	74
ENT - Otolaryngology	0.65	0.57	0.61	77
Gastroenterology	0.43	0.38	0.40	76
Hematology - Oncology	0.63	0.58	0.61	79
Neurology	0.35	0.35	0.35	66
Obstetrics / Gynecology	0.67	0.62	0.64	74
Ophthalmology	0.63	0.87	0.73	55
Orthopedic	0.56	0.58	0.57	79
Pediatrics - Neonatal	0.67	0.48	0.56	71
Psychiatry / Psychology	0.46	0.90	0.61	73
Radiology	0.20	0.16	0.18	87
Urology	0.63	0.42	0.51	80

accuracy			0.52	891
macro avg	0.53	0.53	0.52	891
weighted avg	0.53	0.52	0.51	891

=====

#### Classification Report for Support Vector Machine:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.57	0.41	0.47	74
ENT - Otolaryngology	0.30	0.18	0.23	77
Gastroenterology	0.29	0.13	0.18	76
Hematology - Oncology	0.44	0.05	0.09	79
Neurology	0.44	0.21	0.29	66
Obstetrics / Gynecology	1.00	0.18	0.30	74
Ophthalmology	0.22	0.22	0.22	55
Orthopedic	0.55	0.46	0.50	79
Pediatrics - Neonatal	0.12	0.92	0.21	71
Psychiatry / Psychology	0.78	0.10	0.17	73
Radiology	0.25	0.05	0.08	87
Urology	0.72	0.16	0.27	80
accuracy			0.25	891
macro avg	0.47	0.25	0.25	891
weighted avg	0.48	0.25	0.25	891

=====

#### Classification Report for XGBoost:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.49	0.46	0.48	74
ENT - Otolaryngology	0.73	0.57	0.64	77
Gastroenterology	0.43	0.42	0.42	76
Hematology - Oncology	0.64	0.59	0.62	79
Neurology	0.34	0.30	0.32	66
Obstetrics / Gynecology	0.67	0.59	0.63	74
Ophthalmology	0.67	0.89	0.77	55
Orthopedic	0.57	0.65	0.61	79
Pediatrics - Neonatal	0.63	0.46	0.54	71
Psychiatry / Psychology	0.45	0.92	0.61	73
Radiology	0.21	0.18	0.20	87
Urology	0.63	0.41	0.50	80
accuracy			0.53	891
macro avg	0.54	0.54	0.53	891

weighted avg	0.54	0.53	0.52	891
--------------	------	------	------	-----

=====

#### Classification Report for Logistic Regression:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.65	0.58	0.61	74
ENT - Otolaryngology	0.62	0.39	0.48	77
Gastroenterology	0.46	0.30	0.37	76
Hematology - Oncology	0.48	0.14	0.22	79
Neurology	0.49	0.32	0.39	66
Obstetrics / Gynecology	0.76	0.43	0.55	74
Ophthalmology	0.40	0.49	0.44	55
Orthopedic	0.71	0.56	0.62	79
Pediatrics - Neonatal	0.26	0.41	0.32	71
Psychiatry / Psychology	0.25	0.99	0.40	73
Radiology	0.43	0.26	0.33	87
Urology	0.61	0.29	0.39	80
accuracy			0.42	891
macro avg	0.51	0.43	0.43	891
weighted avg	0.51	0.42	0.42	891

=====

#Advanced Model Development

#BioBERT ('dmis-lab/biobert-v1.1')

```
[59]: # Set up device for PyTorch
device = torch.device("cuda" if torch.cuda.is_available() else "cpu")

# Split data into features and target
X = df['preprocessed_transcription']
y = df['medical_specialty']

# Encode target labels
label_encoder = LabelEncoder()
y_encoded = label_encoder.fit_transform(y)

# Load BioBERT tokenizer
tokenizer = BertTokenizer.from_pretrained('dmis-lab/biobert-v1.1',
do_lower_case=True)
```

```

# Tokenize and encode sequences
encoded_data = tokenizer(X.tolist(), padding=True, truncation=True,
    ↳max_length=128, return_tensors='pt')

# Split data into train and test sets
train_inputs, test_inputs, train_labels, test_labels =
    ↳train_test_split(encoded_data['input_ids'],

    ↳y_encoded,

    ↳random_state=42,

    ↳test_size=0.2,

    ↳stratify=y_encoded)

# Apply SMOTE to balance the dataset
smote = SMOTE(random_state=42)
train_inputs_resampled, train_labels_resampled = smote.
    ↳fit_resample(train_inputs, train_labels)

# Convert data to PyTorch tensors
train_inputs_tensor = torch.tensor(train_inputs_resampled)
test_inputs_tensor = torch.tensor(test_inputs)
train_labels_tensor = torch.tensor(train_labels_resampled)
test_labels_tensor = torch.tensor(test_labels)

# Create DataLoader for train and test sets
train_data = TensorDataset(train_inputs_tensor, train_labels_tensor)
train_sampler = RandomSampler(train_data)
train_dataloader = DataLoader(train_data, sampler=train_sampler, batch_size=32)

test_data = TensorDataset(test_inputs_tensor, test_labels_tensor)
test_sampler = SequentialSampler(test_data)
test_dataloader = DataLoader(test_data, sampler=test_sampler, batch_size=32)

# Load BioBERT model for sequence classification
model = BertForSequenceClassification.from_pretrained('dmis-lab/biobert-v1.1',
    ↳num_labels=len(label_encoder.classes_))
model.to(device)

# Set up optimizer and scheduler
optimizer = AdamW(model.parameters(), lr=2e-5, eps=1e-8)
epochs = 4
total_steps = len(train_dataloader) * epochs

```

```

scheduler = get_linear_schedule_with_warmup(optimizer, num_warmup_steps=0,
↪num_training_steps=total_steps)

# Train the model
model.train()
for epoch in range(epochs):
    total_loss = 0
    for batch in train_dataloader:
        batch = tuple(t.to(device) for t in batch)
        inputs = {'input_ids': batch[0],
                  'labels': batch[1]}
        optimizer.zero_grad()
        outputs = model(**inputs)
        loss = outputs.loss
        total_loss += loss.item()
        loss.backward()
        torch.nn.utils.clip_grad_norm_(model.parameters(), 1.0)
        optimizer.step()
        scheduler.step()

    avg_train_loss = total_loss / len(train_dataloader)
    print(f'Epoch {epoch + 1}/{epochs}, Average Training Loss:
↪{avg_train_loss}')

# Evaluate the model
model.eval()
predictions, true_labels = [], []
for batch in test_dataloader:
    batch = tuple(t.to(device) for t in batch)
    inputs = {'input_ids': batch[0],
              'labels': batch[1]}
    with torch.no_grad():
        outputs = model(**inputs)
    logits = outputs.logits
    logits = logits.detach().cpu().numpy()
    label_ids = inputs['labels'].cpu().numpy()
    predictions.append(logits)
    true_labels.append(label_ids)

predictions = np.concatenate(predictions, axis=0)
true_labels = np.concatenate(true_labels, axis=0)
predicted_labels = np.argmax(predictions, axis=1)
class_names = label_encoder.classes_

# Print classification report with method name included
print("Classification Report for BioBERT:")

```

```
print(classification_report(true_labels, predicted_labels,
↪target_names=class_names))
```

```
tokenizer_config.json: 0%|          | 0.00/49.0 [00:00<?, ?B/s]
vocab.txt: 0%|          | 0.00/213k [00:00<?, ?B/s]
special_tokens_map.json: 0%|          | 0.00/112 [00:00<?, ?B/s]
config.json: 0%|          | 0.00/462 [00:00<?, ?B/s]
pytorch_model.bin: 0%|          | 0.00/433M [00:00<?, ?B/s]
```

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:

['classifier.bias', 'classifier.weight']

You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

We strongly recommend passing in an `attention\_mask` since your input\_ids may be padded. See <https://huggingface.co/docs/transformers/troubleshooting#incorrect-output-when-padding-tokens-arent-masked>.

Epoch 1/4, Average Training Loss: 1.6693773828446865

Epoch 2/4, Average Training Loss: 0.620629906787404

Epoch 3/4, Average Training Loss: 0.4752154874482325

Epoch 4/4, Average Training Loss: 0.4098977278252797

Classification Report for BioBERT:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.95	0.95	0.95	74
ENT - Otolaryngology	1.00	1.00	1.00	19
Gastroenterology	0.96	1.00	0.98	45
Hematology - Oncology	0.88	0.83	0.86	18
Neurology	0.84	0.90	0.87	63
Obstetrics / Gynecology	0.88	0.97	0.92	31
Ophthalmology	1.00	1.00	1.00	17
Orthopedic	0.90	0.93	0.92	71
Pediatrics - Neonatal	0.92	0.86	0.89	14
Psychiatry / Psychology	0.88	0.64	0.74	11
Radiology	0.98	0.85	0.91	55
Urology	0.98	0.98	0.98	47
accuracy			0.93	465
macro avg	0.93	0.91	0.92	465
weighted avg	0.93	0.93	0.93	465

#AUC and ROC Plot of BioBERT Model

```
[60]: # Compute ROC curve and ROC area for each class
fpr = dict()
```

```

tpr = dict()
roc_auc = dict()
n_classes = len(class_names) # Number of classes

for i in range(n_classes):
    fpr[i], tpr[i], _ = roc_curve((true_labels == i), predictions[:, i])
    roc_auc[i] = auc(fpr[i], tpr[i])

# Compute micro-average ROC curve and ROC area
fpr["micro"], tpr["micro"], _ = roc_curve(label_binarize(true_labels,
↪classes=np.arange(n_classes)).ravel(),
                                           predictions.ravel())
roc_auc["micro"] = auc(fpr["micro"], tpr["micro"])

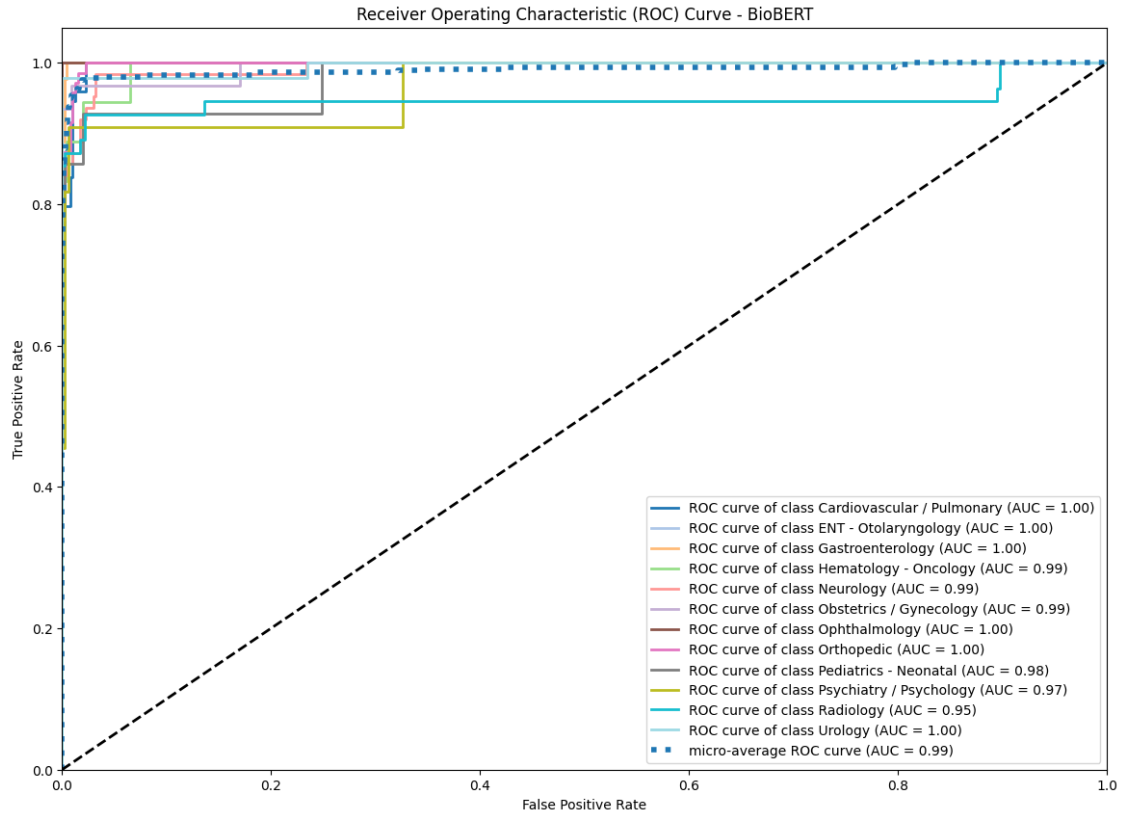
# Plot all ROC curves
plt.figure(figsize=(14, 10))
colors = plt.cm.get_cmap('tab20', n_classes)

for i, color in zip(range(n_classes), colors.colors):
    plt.plot(fpr[i], tpr[i], color=color, lw=2,
             label=f'ROC curve of class {class_names[i]} (AUC = {roc_auc[i]:0.
↪2f})')

plt.plot(fpr["micro"], tpr["micro"], linestyle=':', linewidth=4,
        label=f'micro-average ROC curve (AUC = {roc_auc["micro"]:0.2f})')

plt.plot([0, 1], [0, 1], 'k--', lw=2)
plt.xlim([0.0, 1.0])
plt.ylim([0.0, 1.05])
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('Receiver Operating Characteristic (ROC) Curve - BioBERT')
plt.legend(loc="lower right")
plt.show()

```

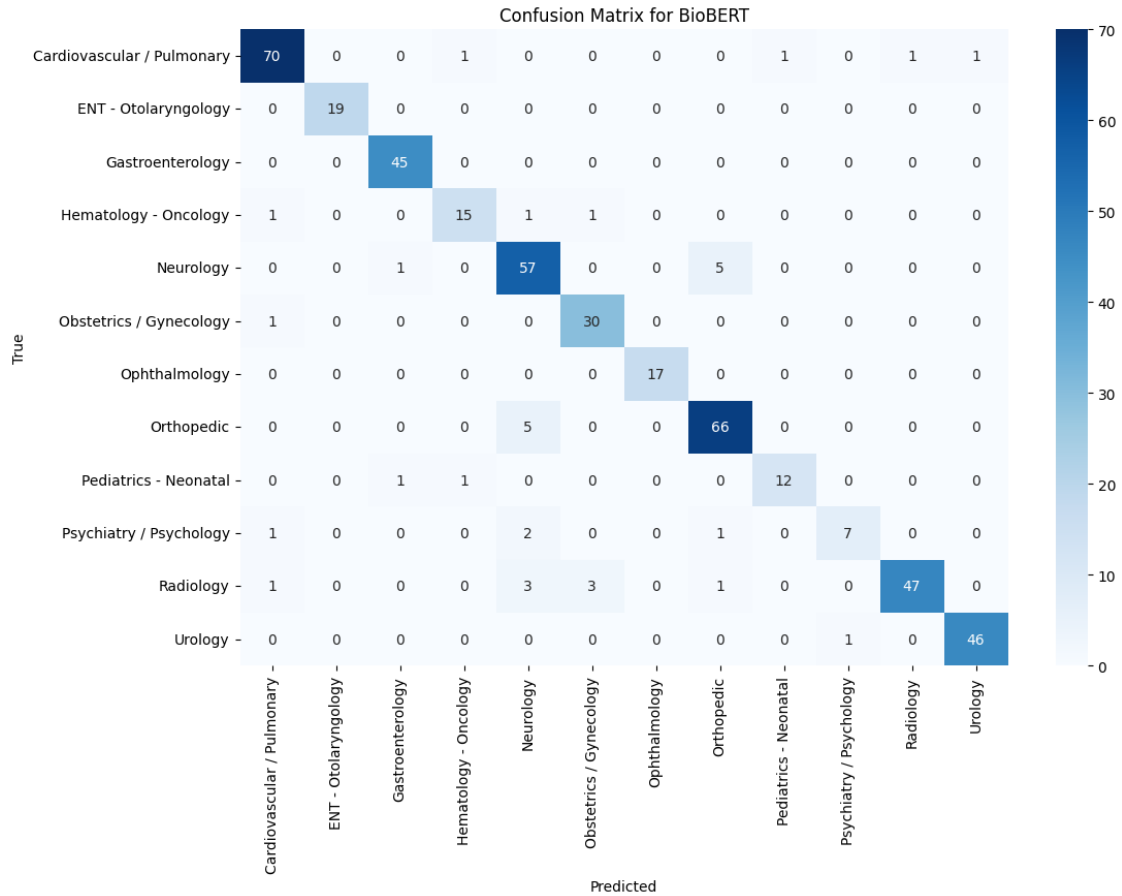


#Confusion Matrix of BioBERT Model

```
[61]: # Calculate the confusion matrix
conf_matrix = confusion_matrix(true_labels, predicted_labels)

# Plot the confusion matrix
plt.figure(figsize=(12, 8))
sns.heatmap(conf_matrix, annot=True, fmt='d', cmap='Blues',
            xticklabels=class_names, yticklabels=class_names)
plt.xlabel('Predicted')
plt.ylabel('True')
plt.title('Confusion Matrix for BioBERT')
plt.show()
```





### #K-Fold Cross-Validation of BioBERT Model

```
[62]: kfold = KFold(n_splits=5, shuffle=True, random_state=42)
kfold_scores = []

for fold, (train_index, val_index) in enumerate(kfold.
    ↪split(encoded_data['input_ids'], y_encoded)):
    print(f'Fold {fold + 1}/{kfold.n_splits}')

    # Split data into train and validation sets
    train_inputs_fold = encoded_data['input_ids'][train_index]
    train_labels_fold = y_encoded[train_index]
    val_inputs_fold = encoded_data['input_ids'][val_index]
    val_labels_fold = y_encoded[val_index]

    # Convert to PyTorch tensors
    train_inputs_tensor = torch.tensor(train_inputs_fold)
    val_inputs_tensor = torch.tensor(val_inputs_fold)
    train_labels_tensor = torch.tensor(train_labels_fold)
```

```

val_labels_tensor = torch.tensor(val_labels_fold)

# Create DataLoader for train and validation sets
train_data_fold = TensorDataset(train_inputs_tensor, train_labels_tensor)
train_sampler_fold = RandomSampler(train_data_fold)
train_dataloader_fold = DataLoader(train_data_fold,
↪sampler=train_sampler_fold, batch_size=32)

val_data_fold = TensorDataset(val_inputs_tensor, val_labels_tensor)
val_sampler_fold = SequentialSampler(val_data_fold)
val_dataloader_fold = DataLoader(val_data_fold, sampler=val_sampler_fold,
↪batch_size=32)

# Reload model for each fold
model = BertForSequenceClassification.from_pretrained('dmis-lab/biobert-v1.
↪1', num_labels=len(label_encoder.classes_))
model.to(device)
optimizer = AdamW(model.parameters(), lr=2e-5, eps=1e-8)
scheduler = get_linear_schedule_with_warmup(optimizer, num_warmup_steps=0,
↪num_training_steps=total_steps)

# Training loop
model.train()
for epoch in range(epochs):
    total_loss = 0
    for batch in train_dataloader_fold:
        batch = tuple(t.to(device) for t in batch)
        inputs = {'input_ids': batch[0],
                  'labels': batch[1]}
        optimizer.zero_grad()
        outputs = model(**inputs)
        loss = outputs.loss
        total_loss += loss.item()
        loss.backward()
        torch.nn.utils.clip_grad_norm_(model.parameters(), 1.0)
        optimizer.step()
        scheduler.step()

    avg_train_loss = total_loss / len(train_dataloader_fold)
    print(f'Epoch {epoch + 1}/{epochs}, Average Training Loss:
↪{avg_train_loss}')

# Evaluation loop
model.eval()
predictions, true_labels = [], []
for batch in val_dataloader_fold:
    batch = tuple(t.to(device) for t in batch)

```

```

        inputs = {'input_ids': batch[0],
                  'labels': batch[1]}
        with torch.no_grad():
            outputs = model(**inputs)
            logits = outputs.logits
            logits = logits.detach().cpu().numpy()
            label_ids = inputs['labels'].cpu().numpy()
            predictions.append(logits)
            true_labels.append(label_ids)

    predictions = np.concatenate(predictions, axis=0)
    true_labels = np.concatenate(true_labels, axis=0)
    predicted_labels = np.argmax(predictions, axis=1)

    # Calculate metrics for this fold
    fold_report = classification_report(true_labels, predicted_labels,
    ↪target_names=class_names, output_dict=True)
    kfold_scores.append(fold_report)

    # Print fold metrics
    print("Classification Report for Fold:")
    print(classification_report(true_labels, predicted_labels,
    ↪target_names=class_names))
    print("\n" + "="*60 + "\n")

# Aggregate K-Fold results
avg_metrics = {
    'precision': np.mean([fold['weighted avg']['precision'] for fold in
    ↪kfold_scores]),
    'recall': np.mean([fold['weighted avg']['recall'] for fold in
    ↪kfold_scores]),
    'f1-score': np.mean([fold['weighted avg']['f1-score'] for fold in
    ↪kfold_scores])
}

print("Average Metrics across all Folds:")
print(avg_metrics)

```

Fold 1/5

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:

['classifier.bias', 'classifier.weight']

You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

Epoch 1/4, Average Training Loss: 1.893898162801387

Epoch 2/4, Average Training Loss: 0.6543802331059666

Epoch 3/4, Average Training Loss: 0.32252542409351315

Epoch 4/4, Average Training Loss: 0.1966225555387594

Classification Report for Fold:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.94	0.99	0.96	74
ENT - Otolaryngology	1.00	1.00	1.00	14
Gastroenterology	1.00	0.91	0.95	44
Hematology - Oncology	0.83	0.90	0.86	21
Neurology	0.82	0.86	0.84	63
Obstetrics / Gynecology	0.94	0.97	0.95	32
Ophthalmology	1.00	1.00	1.00	16
Orthopedic	0.91	0.92	0.92	65
Pediatrics - Neonatal	1.00	0.73	0.84	11
Psychiatry / Psychology	0.93	0.81	0.87	16
Radiology	0.97	0.94	0.96	70
Urology	0.97	0.97	0.97	39
accuracy			0.93	465
macro avg	0.94	0.92	0.93	465
weighted avg	0.93	0.93	0.93	465

=====

Fold 2/5

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:

['classifier.bias', 'classifier.weight']

You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

Epoch 1/4, Average Training Loss: 1.7774644362724434

Epoch 2/4, Average Training Loss: 0.6324314500315714

Epoch 3/4, Average Training Loss: 0.31388682139627005

Epoch 4/4, Average Training Loss: 0.2173106681744931

Classification Report for Fold:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.97	0.99	0.98	70
ENT - Otolaryngology	0.95	0.95	0.95	21
Gastroenterology	0.95	0.97	0.96	36
Hematology - Oncology	0.93	0.87	0.90	15
Neurology	0.91	0.89	0.90	65
Obstetrics / Gynecology	0.91	0.97	0.94	33
Ophthalmology	1.00	1.00	1.00	19
Orthopedic	0.97	0.96	0.97	73
Pediatrics - Neonatal	0.81	0.81	0.81	16

Psychiatry / Psychology	0.86	0.75	0.80	8
Radiology	0.95	0.94	0.95	66
Urology	0.98	1.00	0.99	43
accuracy			0.95	465
macro avg	0.93	0.92	0.93	465
weighted avg	0.95	0.95	0.95	465

=====

Fold 3/5

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:

['classifier.bias', 'classifier.weight']

You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

Epoch 1/4, Average Training Loss: 1.8254361991154946

Epoch 2/4, Average Training Loss: 0.6619001639596487

Epoch 3/4, Average Training Loss: 0.33570041166523756

Epoch 4/4, Average Training Loss: 0.21859307675543477

Classification Report for Fold:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.96	0.93	0.95	74
ENT - Otolaryngology	0.93	1.00	0.96	26
Gastroenterology	0.96	0.93	0.94	54
Hematology - Oncology	0.88	0.88	0.88	17
Neurology	0.89	0.96	0.92	68
Obstetrics / Gynecology	0.92	0.96	0.94	23
Ophthalmology	0.94	1.00	0.97	15
Orthopedic	0.95	0.95	0.95	66
Pediatrics - Neonatal	0.81	0.76	0.79	17
Psychiatry / Psychology	0.71	0.71	0.71	7
Radiology	0.94	0.88	0.91	50
Urology	0.96	0.94	0.95	48
accuracy			0.93	465
macro avg	0.90	0.91	0.91	465
weighted avg	0.93	0.93	0.93	465

=====

Fold 4/5

Some weights of BertForSequenceClassification were not initialized from the

model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:  
['classifier.bias', 'classifier.weight']  
You should probably TRAIN this model on a down-stream task to be able to use it  
for predictions and inference.

Epoch 1/4, Average Training Loss: 1.7640899692551564  
Epoch 2/4, Average Training Loss: 0.6257501029867237  
Epoch 3/4, Average Training Loss: 0.29930050014439274  
Epoch 4/4, Average Training Loss: 0.19988448640047493  
Classification Report for Fold:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.91	0.96	0.94	75
ENT - Otolaryngology	1.00	0.89	0.94	19
Gastroenterology	0.93	0.97	0.95	38
Hematology - Oncology	0.95	0.78	0.86	23
Neurology	0.84	0.92	0.88	64
Obstetrics / Gynecology	0.96	0.93	0.95	28
Ophthalmology	1.00	1.00	1.00	16
Orthopedic	0.93	0.90	0.91	71
Pediatrics - Neonatal	0.79	0.85	0.81	13
Psychiatry / Psychology	0.82	0.82	0.82	11
Radiology	0.97	0.89	0.93	44
Urology	0.97	0.97	0.97	63
accuracy			0.92	465
macro avg	0.92	0.91	0.91	465
weighted avg	0.93	0.92	0.92	465

=====

Fold 5/5

Some weights of BertForSequenceClassification were not initialized from the  
model checkpoint at dmis-lab/biobert-v1.1 and are newly initialized:  
['classifier.bias', 'classifier.weight']  
You should probably TRAIN this model on a down-stream task to be able to use it  
for predictions and inference.

Epoch 1/4, Average Training Loss: 1.677667657197532  
Epoch 2/4, Average Training Loss: 0.5729492114762128  
Epoch 3/4, Average Training Loss: 0.3026561723169634  
Epoch 4/4, Average Training Loss: 0.20843907527751843  
Classification Report for Fold:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.92	0.92	0.92	78
ENT - Otolaryngology	0.94	0.94	0.94	16

Gastroenterology	0.98	0.98	0.98	52
Hematology - Oncology	0.81	0.93	0.87	14
Neurology	0.83	0.84	0.83	57
Obstetrics / Gynecology	0.97	0.95	0.96	39
Ophthalmology	0.94	0.94	0.94	17
Orthopedic	0.88	0.96	0.92	80
Pediatrics - Neonatal	0.75	0.46	0.57	13
Psychiatry / Psychology	1.00	0.64	0.78	11
Radiology	0.90	0.86	0.88	43
Urology	0.98	1.00	0.99	44
accuracy			0.91	464
macro avg	0.91	0.87	0.88	464
weighted avg	0.91	0.91	0.91	464

=====

Average Metrics across all Folds:

```
{'precision': 0.9287861952034742, 'recall': 0.9277039302929182, 'f1-score': 0.927142847241272}
```

#Misclassification Analysis of BioBERT Model

```
[63]: # Calculate the number of entries for each class
class_counts = np.bincount(true_labels)

# Calculate the number of misclassifications for each class
misclassified_counts = np.bincount(true_labels[true_labels !=
    ↪ predicted_labels], minlength=len(class_names))

# Calculate accuracy for each class
accuracy_per_class = (class_counts - misclassified_counts) / class_counts * 100

# Create a DataFrame to display the results
misclassification_analysis_df = pd.DataFrame({
    'Medical Specialties': class_names,
    'Number of Entries': class_counts,
    'Number of BioBERT Misclassifications Errors': misclassified_counts,
    'Accuracy (%)': accuracy_per_class
})

# Display the DataFrame
print(misclassification_analysis_df)
```

	Medical Specialties	Number of Entries \
0	Cardiovascular / Pulmonary	78
1	ENT - Otolaryngology	16

2	Gastroenterology	52
3	Hematology - Oncology	14
4	Neurology	57
5	Obstetrics / Gynecology	39
6	Ophthalmology	17
7	Orthopedic	80
8	Pediatrics - Neonatal	13
9	Psychiatry / Psychology	11
10	Radiology	43
11	Urology	44

	Number of BioBERT Misclassifications Errors	Accuracy (%)
0	6	92.307692
1	1	93.750000
2	1	98.076923
3	1	92.857143
4	9	84.210526
5	2	94.871795
6	1	94.117647
7	3	96.250000
8	7	46.153846
9	4	63.636364
10	6	86.046512
11	0	100.000000

#Number of BioBERT Misclassifications by True Category

```
[64]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()
```



```

    return probs

# Select misclassified examples
misclassified_indices = np.where(predicted_labels != true_labels)[0]
misclassified_texts = df['preprocessed_transcription'].
    ↪iloc[misclassified_indices].tolist()
misclassified_true_labels = true_labels[misclassified_indices]
misclassified_predicted_labels = predicted_labels[misclassified_indices]

# Convert misclassified labels to their category names
misclassified_true_categories = [class_names[label] for label in
    ↪misclassified_true_labels]
misclassified_predicted_categories = [class_names[label] for label in
    ↪misclassified_predicted_labels]

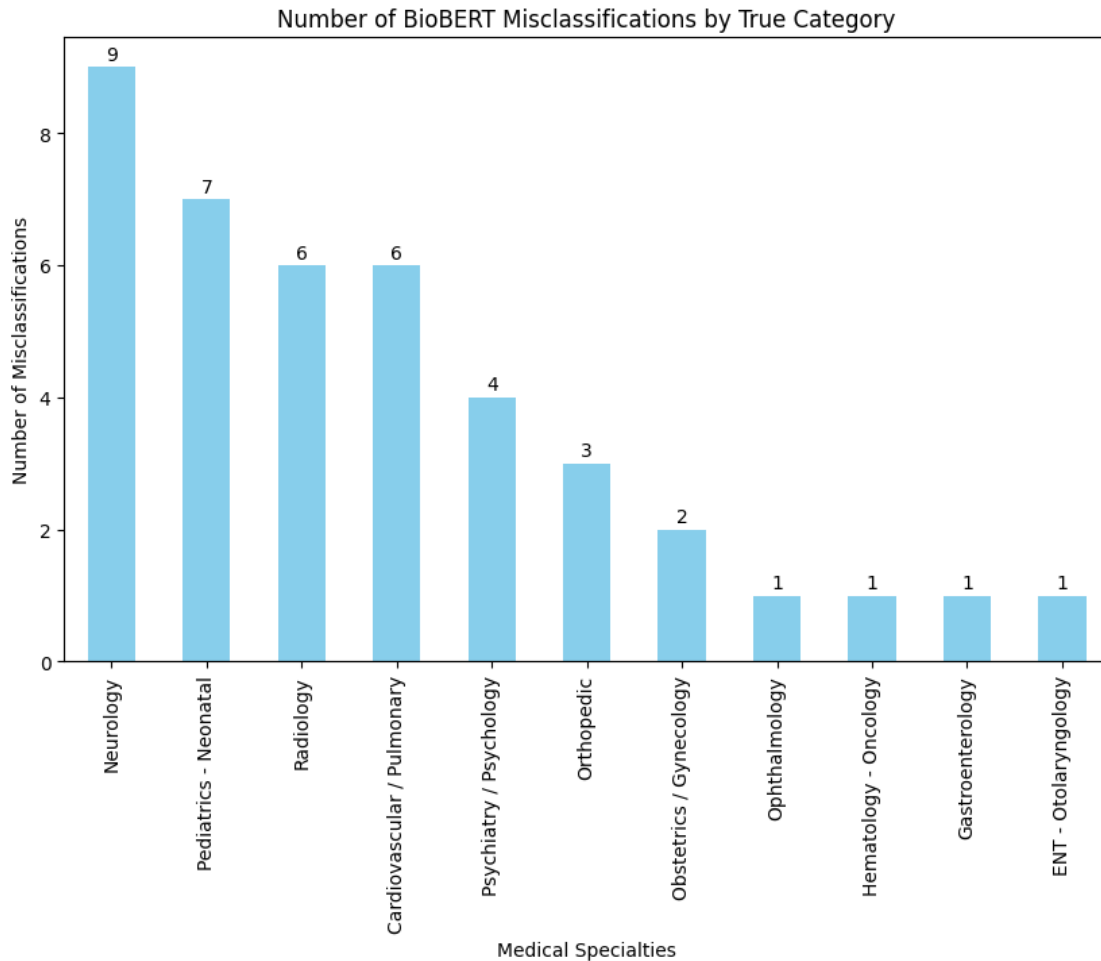
# Bar graph of misclassified categories
misclassified_counts = pd.Series(misclassified_true_categories).value_counts()

# Plotting
plt.figure(figsize=(10, 6))
bars = misclassified_counts.plot(kind='bar', color='skyblue')
plt.title('Number of BioBERT Misclassifications by True Category')
plt.xlabel('Medical Specialties')
plt.ylabel('Number of Misclassifications')

# Add numbers on top of the bars
for bar in bars.patches:
    plt.text(
        bar.get_x() + bar.get_width() / 2,
        bar.get_height() + 0.05,
        int(bar.get_height()),
        ha='center',
        va='bottom',
        color='black',
        fontsize=10
    )

plt.show()

```



#Explainable AI (XAI) Technique

#Integration of XAI with BioBERT Model

```
[65]: pip install shap
```

Collecting shap

Downloading shap-0.46.0-cp310-cp310-manylinux\_2\_12\_x86\_64.manylinux2010\_x86\_64.manylinux\_2\_17\_x86\_64.manylinux2014\_x86\_64.whl (540 kB)

540.1/540.1

kB 7.3 MB/s eta 0:00:00

Requirement already satisfied: numpy in /usr/local/lib/python3.10/dist-packages (from shap) (1.25.2)

Requirement already satisfied: scipy in /usr/local/lib/python3.10/dist-packages (from shap) (1.11.4)

Requirement already satisfied: scikit-learn in /usr/local/lib/python3.10/dist-packages (from shap) (1.2.2)

```

Requirement already satisfied: pandas in /usr/local/lib/python3.10/dist-packages
(from shap) (2.0.3)
Requirement already satisfied: tqdm>=4.27.0 in /usr/local/lib/python3.10/dist-
packages (from shap) (4.66.4)
Requirement already satisfied: packaging>20.9 in /usr/local/lib/python3.10/dist-
packages (from shap) (24.1)
Collecting slicer==0.0.8 (from shap)
  Downloading slicer-0.0.8-py3-none-any.whl (15 kB)
Requirement already satisfied: numba in /usr/local/lib/python3.10/dist-packages
(from shap) (0.58.1)
Requirement already satisfied: cloudpickle in /usr/local/lib/python3.10/dist-
packages (from shap) (2.2.1)
Requirement already satisfied: llvmlite<0.42,>=0.41.0dev0 in
/usr/local/lib/python3.10/dist-packages (from numba->shap) (0.41.1)
Requirement already satisfied: python-dateutil>=2.8.2 in
/usr/local/lib/python3.10/dist-packages (from pandas->shap) (2.8.2)
Requirement already satisfied: pytz>=2020.1 in /usr/local/lib/python3.10/dist-
packages (from pandas->shap) (2023.4)
Requirement already satisfied: tzdata>=2022.1 in /usr/local/lib/python3.10/dist-
packages (from pandas->shap) (2024.1)
Requirement already satisfied: joblib>=1.1.1 in /usr/local/lib/python3.10/dist-
packages (from scikit-learn->shap) (1.4.2)
Requirement already satisfied: threadpoolctl>=2.0.0 in
/usr/local/lib/python3.10/dist-packages (from scikit-learn->shap) (3.5.0)
Requirement already satisfied: six>=1.5 in /usr/local/lib/python3.10/dist-
packages (from python-dateutil>=2.8.2->pandas->shap) (1.16.0)
Installing collected packages: slicer, shap
Successfully installed shap-0.46.0 slicer-0.0.8

```

```

[66]: # SHAP library for explainable AI
      import shap # For SHapley Additive exPlanations

```

*#SHAP Visualization of Correctly Classified Examples*

*#SHAP Visualization of Sample Index '0'*

```

[67]: # Define the class names
      class_names = label_encoder.classes_

      # Create a predict function for SHAP
      def predict_proba(texts):
          encoded_inputs = tokenizer.batch_encode_plus(
              texts,
              max_length=128,
              padding=True,
              truncation=True,
              return_tensors='pt'
          )

```

```

input_ids = encoded_inputs['input_ids'].to(device)
attention_mask = encoded_inputs['attention_mask'].to(device)

with torch.no_grad():
    outputs = model(input_ids=input_ids, attention_mask=attention_mask)
    logits = outputs.logits
    probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

return probs

# Select a single input from the test data
sample_index = 0
single_input = df['preprocessed_transcription'].iloc[sample_index]

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

# Explain the prediction for the selected input
shap_values = explainer([single_input])

# Map predicted labels to category names
predicted_label = np.argmax(predict_proba([single_input])[0])
predicted_category = class_names[predicted_label]

# Display true medical specialty (target variable) and predicted category
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category)

# Visualize the results
shap.initjs()
shap.plots.text(shap_values[0])

```

```
0%|          | 0/498 [00:00<?, ?it/s]
```

```
PartitionExplainer explainer: 2it [00:15, 15.50s/it]
```

```
True Medical Specialty:
Cardiovascular / Pulmonary
```

```
Predicted Category:
Cardiovascular / Pulmonary
```

```
<IPython.core.display.HTML object>
```

```
<IPython.core.display.HTML object>
```

#SHAP Visualization of Sample Index '1'

```
[68]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

    return probs

# Select a single input from the test data
sample_index = 1
single_input = df['preprocessed_transcription'].iloc[sample_index]

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

# Explain the prediction for the selected input
shap_values = explainer([single_input])

# Map predicted labels to category names
predicted_label = np.argmax(predict_proba([single_input])[0])
predicted_category = class_names[predicted_label]

# Display true medical specialty (target variable) and predicted category
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category)

# Visualize the results
shap.initjs()
```

```
shap.plots.text(shap_values[0])
```

Token indices sequence length is longer than the specified maximum sequence length for this model (525 > 512). Running this sequence through the model will result in indexing errors

```
0%|          | 0/498 [00:00<?, ?it/s]
```

PartitionExplainer explainer: 2it [00:20, 20.50s/it]

True Medical Specialty:  
Cardiovascular / Pulmonary

Predicted Category:  
Cardiovascular / Pulmonary

<IPython.core.display.HTML object>

<IPython.core.display.HTML object>

#SHAP Visualization of Sample Index '825'

```
[69]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

    return probs

# Select a single input from the test data
sample_index = 825
single_input = df['preprocessed_transcription'].iloc[sample_index]
```

```

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

# Explain the prediction for the selected input
shap_values = explainer([single_input])

# Map predicted labels to category names
predicted_label = np.argmax(predict_proba([single_input])[0])
predicted_category = class_names[predicted_label]

# Display true medical specialty (target variable) and predicted category
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category)

# Visualize the results
shap.initjs()
shap.plots.text(shap_values[0])

```

0%| | 0/498 [00:00<?, ?it/s]

PartitionExplainer explainer: 2it [00:13, 13.13s/it]

True Medical Specialty:  
Orthopedic

Predicted Category:  
Orthopedic

<IPython.core.display.HTML object>

<IPython.core.display.HTML object>

#SHAP Visualization of Sample Index '116'

```

[70]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

```

```

input_ids = encoded_inputs['input_ids'].to(device)
attention_mask = encoded_inputs['attention_mask'].to(device)

with torch.no_grad():
    outputs = model(input_ids=input_ids, attention_mask=attention_mask)
    logits = outputs.logits
    probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

return probs

# Select a single input from the test data
sample_index = 116
single_input = df['preprocessed_transcription'].iloc[sample_index]

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

# Explain the prediction for the selected input
shap_values = explainer([single_input])

# Map predicted labels to category names
predicted_label = np.argmax(predict_proba([single_input])[0])
predicted_category = class_names[predicted_label]

# Display true medical specialty (target variable) and predicted category
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category)

# Visualize the results
shap.initjs()
shap.plots.text(shap_values[0])

```

```
0%|          | 0/498 [00:00<?, ?it/s]
```

```
PartitionExplainer explainer: 2it [00:11, 11.57s/it]
```

```
True Medical Specialty:
```

```
Urology
```

```
Predicted Category:
```

```
Urology
```

```
<IPython.core.display.HTML object>
```

```
<IPython.core.display.HTML object>
```



## 1 SHAP Visualization of Misclassified Sample Index '263'

```
[71]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

    return probs

# Select a single input from the test data
sample_index = 263
single_input = df['preprocessed_transcription'].iloc[sample_index]

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

# Explain the prediction for the selected input
shap_values = explainer([single_input])

# Map predicted labels to category names
predicted_label = np.argmax(predict_proba([single_input])[0])
predicted_category = class_names[predicted_label]

# Display true medical specialty (target variable) and predicted category
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category)

# Visualize the results
shap.initjs()
```

```
shap.plots.text(shap_values[0])
```

```
0%|          | 0/498 [00:00<?, ?it/s]
```

True Medical Specialty:

Radiology

Predicted Category:

Orthopedic

<IPython.core.display.HTML object>

<IPython.core.display.HTML object>

#Perturbation Testing of SHAP XAI Technique

#Removing the Most Important Words Identified by SHAP

```
[72]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

    return probs

# Select a single input from the test data
sample_index = 1
single_input = df['preprocessed_transcription'].iloc[sample_index]

# Words to remove based on SHAP output with commas included
words_to_remove = [
    "cardiovascular,", "pulmonary,", "doppler,", "echocardiogram,", "annular,",
    "aortic,", "root,", "aortic,", "valve,", "atrial,", "atrium,",
    ↪ "calcification,"
```

```

        "cavity,", "ejection,", "fraction,", "mitral,", "obliteration,", "outflow,",
        "regurgitation,", "relaxation,", "pattern,", "stenosis,", "systolic,",
        ↪ "function,",
        "tricuspid,", "valve,", "ventricular,", "ventricular,", "cavity,", "wall,",
        "motion,", "pulmonary,", "left,", "ventricular,", "cavity,", "size,",
        ↪ "wall,",
        "thickness,", "appear,", "normal,", "wall,", "motion,", "left,",
        ↪ "ventricular,",
        "systolic,", "function,", "appears,", "hyperdynamic,", "estimated,",
        ↪ "ejection,",
        "fraction,", "near,", "cavity,", "obliteration,"
    ]

    # Remove the words from the input text
    modified_input = single_input
    for word in words_to_remove:
        modified_input = modified_input.replace(word, '')

    # SHAP Explainer expects list of strings for transformer models
    explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

    # Explain the prediction for the modified input
    shap_values_modified = explainer([modified_input])

    # Map predicted labels to category names for modified input
    predicted_label_modified = np.argmax(predict_proba([modified_input])[0])
    predicted_category_modified = class_names[predicted_label_modified]

    # Display true medical specialty (target variable) and predicted category for
    ↪ modified input
    print("\nAfter Removing the Most Important Words:")
    print("True Medical Specialty:")
    print(df['medical_specialty'].iloc[sample_index])
    print("\nPredicted Category:")
    print(predicted_category_modified)

    # Visualize the results for modified input
    shap.initjs()
    shap.plots.text(shap_values_modified[0])

```

```
0%|          | 0/498 [00:00<?, ?it/s]
```

After Removing the Most Important Words:  
 True Medical Specialty:  
 Cardiovascular / Pulmonary

Predicted Category:  
Cardiovascular / Pulmonary  
<IPython.core.display.HTML object>  
<IPython.core.display.HTML object>  
#Adding Noisy Words

```
[73]: # Define the class names
class_names = label_encoder.classes_

# Create a predict function for SHAP
def predict_proba(texts):
    encoded_inputs = tokenizer.batch_encode_plus(
        texts,
        max_length=128,
        padding=True,
        truncation=True,
        return_tensors='pt'
    )

    input_ids = encoded_inputs['input_ids'].to(device)
    attention_mask = encoded_inputs['attention_mask'].to(device)

    with torch.no_grad():
        outputs = model(input_ids=input_ids, attention_mask=attention_mask)
        logits = outputs.logits
        probs = torch.nn.functional.softmax(logits, dim=1).cpu().numpy()

    return probs

# Select a single input from the test data
sample_index = 1
single_input = df['preprocessed_transcription'].iloc[sample_index]

# Words to remove based on SHAP output with commas included
words_to_remove = [
    "cardiovascular,", "pulmonary,", "doppler,", "echocardiogram,", "annular,",
    "aortic,", "root,", "aortic,", "valve,", "atrial,", "atrium,",
    ↪ "calcification,",
    "cavity,", "ejection,", "fraction,", "mitral,", "obliteration,", "outflow,",
    "regurgitation,", "relaxation,", "pattern,", "stenosis,", "systolic,",
    ↪ "function,",
    "tricuspid,", "valve,", "ventricular,", "ventricular,", "cavity,", "wall,",
    "motion,", "pulmonary,", "left,", "ventricular,", "cavity,", "size,",
    ↪ "wall,",
    "thickness,", "appear,", "normal,", "wall,", "motion,", "left,",
    ↪ "ventricular,"
```

```

    "systolic,", "function,", "appears,", "hyperdynamic,", "estimated,",
    ↪ "ejection,",
    "fraction,", "near,", "cavity,", "obliteration,"
]

# Remove the words from the input text
modified_input = single_input
for word in words_to_remove:
    modified_input = modified_input.replace(word, '')

# Define a function to add noise to the text
def add_noise(text):
    # Domain-specific irrelevant terms
    irrelevant_terms = ["aspirin", "antibiotic", "bandage", "nausea",
    ↪ "headache"]

    # Split text into words
    words = text.split()

    # List to store noisy words
    added_noisy_words = []

    # Introduce random noise: add irrelevant terms and slight misspellings
    noisy_words = []
    for word in words:
        if random.random() < 0.1: # 10% chance to add an irrelevant term
            new_word = random.choice(irrelevant_terms)
            noisy_words.append(new_word)
            added_noisy_words.append(new_word)
        if random.random() < 0.1: # 10% chance to slightly alter the word
            new_word = word[:-1] + random.choice('abcdefghijklmnopqrstuvwxyz')
            noisy_words.append(new_word)
            added_noisy_words.append(new_word)
        else:
            noisy_words.append(word)

    # Join words back into a string
    noisy_text = ' '.join(noisy_words)

    return noisy_text, added_noisy_words

# Add noise to the modified input
noisy_modified_input, added_noisy_words = add_noise(modified_input)

# SHAP Explainer expects list of strings for transformer models
explainer = shap.Explainer(predict_proba, tokenizer, output_names=class_names)

```

```

# Explain the prediction for the noisy modified input
shap_values_noisy_modified = explainer([noisy_modified_input])

# Map predicted labels to category names for noisy modified input
predicted_label_noisy_modified = np.
    ↪argmax(predict_proba([noisy_modified_input])[0])
predicted_category_noisy_modified = class_names[predicted_label_noisy_modified]

# Display true medical specialty (target variable) and predicted category for
    ↪noisy modified input
print("\nAfter Adding Noise to the Input:")
print("True Medical Specialty:")
print(df['medical_specialty'].iloc[sample_index])
print("\nPredicted Category:")
print(predicted_category_noisy_modified)

# Display the noisy words added
print("\nNoisy Words Added:")
print(added_noisy_words)

# Visualize the results for noisy modified input
shap.initjs()
shap.plots.text(shap_values_noisy_modified[0])

```

After Adding Noise to the Input:

True Medical Specialty:

Cardiovascular / Pulmonary

Predicted Category:

Cardiovascular / Pulmonary

Noisy Words Added:

```
['alsor', 'increasedq', 'nausea', 'antibiotic', 'consistentp', 'nausea',
'wellx', 'headache', 'bandage', 'mmp', 'supportivey', 'structuref', 'nausea',
'nausea', 'aspirin', 'aspirin', 'missedc', 'studz']
```

<IPython.core.display.HTML object>

<IPython.core.display.HTML object>

#ClinicalBERT ('emilyalsentzer/Bio\_ClinicalBERT')

```

[74]: # Set up device for PyTorch
device = torch.device("cuda" if torch.cuda.is_available() else "cpu")

# Split data into features and target
X = df['preprocessed_transcription']
y = df['medical_specialty']

```

```

# Encode target labels
label_encoder = LabelEncoder()
y_encoded = label_encoder.fit_transform(y)

# Load Clinical BERT tokenizer
tokenizer = BertTokenizer.from_pretrained('emilyalsentzer/Bio_ClinicalBERT',
    ↪do_lower_case=True)

# Tokenize and encode sequences
encoded_data = tokenizer(X.tolist(), padding=True, truncation=True,
    ↪max_length=128, return_tensors='pt')

# Split data into train and test sets
train_inputs, test_inputs, train_labels, test_labels =
    ↪train_test_split(encoded_data['input_ids'],
    ↪y_encoded,
    ↪random_state=42,
    ↪test_size=0.2,
    ↪stratify=y_encoded)

# Apply SMOTE to balance the dataset
smote = SMOTE(random_state=42)
train_inputs_resampled, train_labels_resampled = smote.
    ↪fit_resample(train_inputs, train_labels)

# Convert data to PyTorch tensors
train_inputs_tensor = torch.tensor(train_inputs_resampled)
test_inputs_tensor = torch.tensor(test_inputs)
train_labels_tensor = torch.tensor(train_labels_resampled)
test_labels_tensor = torch.tensor(test_labels)

# Create DataLoader for train and test sets
train_data = TensorDataset(train_inputs_tensor, train_labels_tensor)
train_sampler = RandomSampler(train_data)
train_dataloader = DataLoader(train_data, sampler=train_sampler, batch_size=32)

test_data = TensorDataset(test_inputs_tensor, test_labels_tensor)
test_sampler = SequentialSampler(test_data)
test_dataloader = DataLoader(test_data, sampler=test_sampler, batch_size=32)

# Load Clinical BERT model for sequence classification

```

```

model = BertForSequenceClassification.from_pretrained('emilyalsentzer/
↳Bio_ClinicalBERT', num_labels=len(label_encoder.classes_))
model.to(device)

# Set up optimizer and scheduler
optimizer = AdamW(model.parameters(), lr=2e-5, eps=1e-8)
epochs = 4
total_steps = len(train_dataloader) * epochs
scheduler = get_linear_schedule_with_warmup(optimizer, num_warmup_steps=0,↳
↳num_training_steps=total_steps)

# Train the model
model.train()
for epoch in range(epochs):
    total_loss = 0
    for batch in train_dataloader:
        batch = tuple(t.to(device) for t in batch)
        inputs = {'input_ids': batch[0],
                  'labels': batch[1]}
        optimizer.zero_grad()
        outputs = model(**inputs)
        loss = outputs.loss
        total_loss += loss.item()
        loss.backward()
        torch.nn.utils.clip_grad_norm_(model.parameters(), 1.0)
        optimizer.step()
        scheduler.step()

    avg_train_loss = total_loss / len(train_dataloader)
    print(f'Epoch {epoch + 1}/{epochs}, Average Training Loss:↳
↳{avg_train_loss}')

# Evaluate the model
model.eval()
predictions, true_labels = [], []
for batch in test_dataloader:
    batch = tuple(t.to(device) for t in batch)
    inputs = {'input_ids': batch[0],
              'labels': batch[1]}
    with torch.no_grad():
        outputs = model(**inputs)
    logits = outputs.logits
    logits = logits.detach().cpu().numpy()
    label_ids = inputs['labels'].cpu().numpy()
    predictions.append(logits)
    true_labels.append(label_ids)

```



```

predictions = np.concatenate(predictions, axis=0)
true_labels = np.concatenate(true_labels, axis=0)
predicted_labels = np.argmax(predictions, axis=1)
class_names = label_encoder.classes_

# Print classification report with method name included
print("Classification Report for ClinicalBERT:")
print(classification_report(true_labels, predicted_labels,
    ↪target_names=class_names))

```

```

vocab.txt: 0%|          | 0.00/213k [00:00<?, ?B/s]
config.json: 0%|         | 0.00/385 [00:00<?, ?B/s]
pytorch_model.bin: 0%|          | 0.00/436M [00:00<?, ?B/s]

```

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at emilyalsentzer/Bio\_ClinicalBERT and are newly initialized: ['classifier.bias', 'classifier.weight']

You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

Epoch 1/4, Average Training Loss: 1.8036292142101698

Epoch 2/4, Average Training Loss: 0.6997512030814376

Epoch 3/4, Average Training Loss: 0.5137819602553334

Epoch 4/4, Average Training Loss: 0.4420144919838224

Classification Report for ClinicalBERT:

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.94	0.91	0.92	74
ENT - Otolaryngology	1.00	0.95	0.97	19
Gastroenterology	0.92	1.00	0.96	45
Hematology - Oncology	0.94	0.89	0.91	18
Neurology	0.84	0.89	0.86	63
Obstetrics / Gynecology	0.86	0.97	0.91	31
Ophthalmology	1.00	0.94	0.97	17
Orthopedic	0.93	0.94	0.94	71
Pediatrics - Neonatal	0.79	0.79	0.79	14
Psychiatry / Psychology	0.89	0.73	0.80	11
Radiology	0.94	0.85	0.90	55
Urology	0.96	0.96	0.96	47
accuracy			0.92	465
macro avg	0.92	0.90	0.91	465
weighted avg	0.92	0.92	0.92	465

#BERT ('bert-base-uncased')

```

[75]: # Set up device for PyTorch
device = torch.device("cuda" if torch.cuda.is_available() else "cpu")

# Split data into features and target
X = df['preprocessed_transcription']
y = df['medical_specialty']

# Encode target labels
label_encoder = LabelEncoder()
y_encoded = label_encoder.fit_transform(y)

# Load BERT tokenizer
tokenizer = BertTokenizer.from_pretrained('bert-base-uncased',
    ↪do_lower_case=True)

# Tokenize and encode sequences
encoded_data = tokenizer(X.tolist(), padding=True, truncation=True,
    ↪max_length=128, return_tensors='pt')

# Split data into train and test sets
train_inputs, test_inputs, train_labels, test_labels =
    ↪train_test_split(encoded_data['input_ids'],
    ↪y_encoded,
    ↪random_state=42,
    ↪test_size=0.2,
    ↪stratify=y_encoded)

# Convert data to PyTorch tensors
train_inputs_tensor = torch.tensor(train_inputs)
test_inputs_tensor = torch.tensor(test_inputs)
train_labels_tensor = torch.tensor(train_labels)
test_labels_tensor = torch.tensor(test_labels)

# Create DataLoader for train and test sets
train_data = TensorDataset(train_inputs_tensor, train_labels_tensor)
train_sampler = RandomSampler(train_data)
train_dataloader = DataLoader(train_data, sampler=train_sampler, batch_size=32)

test_data = TensorDataset(test_inputs_tensor, test_labels_tensor)
test_sampler = SequentialSampler(test_data)
test_dataloader = DataLoader(test_data, sampler=test_sampler, batch_size=32)

```

```

# Load BERT model for sequence classification
model = BertForSequenceClassification.from_pretrained('bert-base-uncased',
    ↪ num_labels=len(label_encoder.classes_))
model.to(device)

# Set up optimizer and scheduler
optimizer = AdamW(model.parameters(), lr=2e-5, eps=1e-8)
epochs = 4
total_steps = len(train_dataloader) * epochs
scheduler = get_linear_schedule_with_warmup(optimizer, num_warmup_steps=0,
    ↪ num_training_steps=total_steps)

# Train the model
model.train()
for epoch in range(epochs):
    total_loss = 0
    for batch in train_dataloader:
        batch = tuple(t.to(device) for t in batch)
        inputs = {'input_ids': batch[0],
            'labels': batch[1]}
        optimizer.zero_grad()
        outputs = model(**inputs)
        loss = outputs.loss
        total_loss += loss.item()
        loss.backward()
        torch.nn.utils.clip_grad_norm_(model.parameters(), 1.0)
        optimizer.step()
        scheduler.step()

    avg_train_loss = total_loss / len(train_dataloader)
    print(f'Epoch {epoch + 1}/{epochs}, Average Training Loss:
    ↪ {avg_train_loss}')

# Evaluate the model
model.eval()
predictions, true_labels = [], []
for batch in test_dataloader:
    batch = tuple(t.to(device) for t in batch)
    inputs = {'input_ids': batch[0],
        'labels': batch[1]}
    with torch.no_grad():
        outputs = model(**inputs)
    logits = outputs.logits
    logits = logits.detach().cpu().numpy()
    label_ids = inputs['labels'].cpu().numpy()
    predictions.append(logits)
    true_labels.append(label_ids)

```

```

predictions = np.concatenate(predictions, axis=0)
true_labels = np.concatenate(true_labels, axis=0)
predicted_labels = np.argmax(predictions, axis=1)
class_names = label_encoder.classes_

# Print classification report with method name included
print("Classification Report for BERT:")
print(classification_report(true_labels, predicted_labels,
    ↪target_names=class_names))

```

```

tokenizer_config.json: 0%|          | 0.00/48.0 [00:00<?, ?B/s]
vocab.txt: 0%|          | 0.00/232k [00:00<?, ?B/s]
tokenizer.json: 0%|          | 0.00/466k [00:00<?, ?B/s]
config.json: 0%|          | 0.00/570 [00:00<?, ?B/s]
model.safetensors: 0%|          | 0.00/440M [00:00<?, ?B/s]

```

Some weights of BertForSequenceClassification were not initialized from the model checkpoint at bert-base-uncased and are newly initialized: ['classifier.bias', 'classifier.weight']  
You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

```

Epoch 1/4, Average Training Loss: 2.0324573436025846
Epoch 2/4, Average Training Loss: 0.8897511072583117
Epoch 3/4, Average Training Loss: 0.5055445887274661
Epoch 4/4, Average Training Loss: 0.37606183527889897
Classification Report for BERT:

```

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.94	0.97	0.95	74
ENT - Otolaryngology	1.00	0.95	0.97	19
Gastroenterology	0.91	0.96	0.93	45
Hematology - Oncology	0.79	0.83	0.81	18
Neurology	0.81	0.87	0.84	63
Obstetrics / Gynecology	0.88	0.97	0.92	31
Ophthalmology	1.00	0.94	0.97	17
Orthopedic	0.91	0.94	0.92	71
Pediatrics - Neonatal	0.92	0.79	0.85	14
Psychiatry / Psychology	1.00	0.73	0.84	11
Radiology	0.98	0.85	0.91	55
Urology	0.98	0.91	0.95	47
accuracy			0.91	465
macro avg	0.93	0.89	0.91	465
weighted avg	0.92	0.91	0.91	465

```
#RoBERTa ('roberta-base')
```

```
[76]: # Set up device for PyTorch
device = torch.device("cuda" if torch.cuda.is_available() else "cpu")

# Split data into features and target
X = df['preprocessed_transcription']
y = df['medical_specialty']

# Encode target labels
label_encoder = LabelEncoder()
y_encoded = label_encoder.fit_transform(y)

# Load RoBERTa tokenizer
tokenizer = RobertaTokenizer.from_pretrained('roberta-base')

# Tokenize and encode sequences
encoded_data = tokenizer(X.tolist(), padding=True, truncation=True,
    ↪max_length=128, return_tensors='pt')

# Split data into train and test sets
train_inputs, test_inputs, train_labels, test_labels =
    ↪train_test_split(encoded_data['input_ids'],
    ↪y_encoded,
    ↪random_state=42,
    ↪test_size=0.2,
    ↪stratify=y_encoded)

# Convert data to PyTorch tensors
train_inputs_tensor = torch.tensor(train_inputs)
test_inputs_tensor = torch.tensor(test_inputs)
train_labels_tensor = torch.tensor(train_labels)
test_labels_tensor = torch.tensor(test_labels)

# Create DataLoader for train and test sets
train_data = TensorDataset(train_inputs_tensor, train_labels_tensor)
train_sampler = RandomSampler(train_data)
train_dataloader = DataLoader(train_data, sampler=train_sampler, batch_size=32)

test_data = TensorDataset(test_inputs_tensor, test_labels_tensor)
test_sampler = SequentialSampler(test_data)
```

```

test_dataloader = DataLoader(test_data, sampler=test_sampler, batch_size=32)

# Load RoBERTa model for sequence classification
model = RobertaForSequenceClassification.from_pretrained('roberta-base',
    ↪ num_labels=len(label_encoder.classes_))
model.to(device)

# Set up optimizer and scheduler
optimizer = AdamW(model.parameters(), lr=2e-5, eps=1e-8)
epochs = 4
total_steps = len(train_dataloader) * epochs
scheduler = get_linear_schedule_with_warmup(optimizer, num_warmup_steps=0,
    ↪ num_training_steps=total_steps)

# Train the model
model.train()
for epoch in range(epochs):
    total_loss = 0
    for batch in train_dataloader:
        batch = tuple(t.to(device) for t in batch)
        inputs = {'input_ids': batch[0],
            'labels': batch[1]}
        optimizer.zero_grad()
        outputs = model(**inputs)
        loss = outputs.loss
        total_loss += loss.item()
        loss.backward()
        torch.nn.utils.clip_grad_norm_(model.parameters(), 1.0)
        optimizer.step()
        scheduler.step()

    avg_train_loss = total_loss / len(train_dataloader)
    print(f'Epoch {epoch + 1}/{epochs}, Average Training Loss:
    ↪ {avg_train_loss}')

# Evaluate the model
model.eval()
predictions, true_labels = [], []
for batch in test_dataloader:
    batch = tuple(t.to(device) for t in batch)
    inputs = {'input_ids': batch[0],
        'labels': batch[1]}
    with torch.no_grad():
        outputs = model(**inputs)
    logits = outputs.logits
    logits = logits.detach().cpu().numpy()
    label_ids = inputs['labels'].cpu().numpy()

```

```

        predictions.append(logits)
        true_labels.append(label_ids)

predictions = np.concatenate(predictions, axis=0)
true_labels = np.concatenate(true_labels, axis=0)
predicted_labels = np.argmax(predictions, axis=1)
class_names = label_encoder.classes_

# Print classification report with method name included
print("Classification Report for RoBERTa:")
print(classification_report(true_labels, predicted_labels,
    ↪target_names=class_names))

```

```

tokenizer_config.json:  0%|          | 0.00/25.0 [00:00<?, ?B/s]
vocab.json:           0%|          | 0.00/899k [00:00<?, ?B/s]
merges.txt:           0%|          | 0.00/456k [00:00<?, ?B/s]
tokenizer.json:        0%|          | 0.00/1.36M [00:00<?, ?B/s]
config.json:          0%|          | 0.00/481 [00:00<?, ?B/s]
model.safetensors:     0%|          | 0.00/499M [00:00<?, ?B/s]

```

Some weights of RobertaForSequenceClassification were not initialized from the model checkpoint at roberta-base and are newly initialized:  
['classifier.dense.bias', 'classifier.dense.weight', 'classifier.out\_proj.bias', 'classifier.out\_proj.weight']  
You should probably TRAIN this model on a down-stream task to be able to use it for predictions and inference.

```

Epoch 1/4, Average Training Loss: 1.8289603065636197
Epoch 2/4, Average Training Loss: 0.542975342374737
Epoch 3/4, Average Training Loss: 0.29224961591979204
Epoch 4/4, Average Training Loss: 0.21937284069293636
Classification Report for RoBERTa:

```

	precision	recall	f1-score	support
Cardiovascular / Pulmonary	0.95	0.96	0.95	74
ENT - Otolaryngology	1.00	1.00	1.00	19
Gastroenterology	0.94	1.00	0.97	45
Hematology - Oncology	0.89	0.89	0.89	18
Neurology	0.82	0.87	0.85	63
Obstetrics / Gynecology	0.88	0.97	0.92	31
Ophthalmology	1.00	0.94	0.97	17
Orthopedic	0.91	0.94	0.92	71
Pediatrics - Neonatal	0.92	0.79	0.85	14
Psychiatry / Psychology	0.89	0.73	0.80	11
Radiology	0.98	0.85	0.91	55
Urology	1.00	0.96	0.98	47

accuracy			0.92	465
macro avg	0.93	0.91	0.92	465
weighted avg	0.93	0.92	0.92	465