



SCHOOL OF
COMPUTER SCIENCE

Course: CIS*6530 - Threat Intel & Risk Analysis

Group 5

Technical Report

Submission 4

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In project was to developed a machine learning pipeline to classify Advanced Persistent Threat (APT) groups based on extracted opcode sequences. The project applied Support Vector Machines (SVM), K-Nearest Neighbors (KNN), and Decision Tree classifiers to analyze 1-Gram and 2-Gram features from the opcodes. Classifier performance was evaluated using accuracy, recall, precision, F1-measure, and confusion matrices. This report outlines each step of the process, including data preprocessing, feature extraction, model training, and evaluation.

1 Import necessary libraries

```
[412]: import pandas as pd
import warnings
from sklearn.exceptions import UndefinedMetricWarning
from sklearn.model_selection import train_test_split, GridSearchCV
from sklearn.preprocessing import MinMaxScaler
from sklearn.metrics import classification_report, confusion_matrix, \
    accuracy_score, f1_score
from sklearn.svm import SVC
from sklearn.neighbors import KNeighborsClassifier
from sklearn.tree import DecisionTreeClassifier
from imblearn.over_sampling import SMOTE
from sklearn.feature_selection import SelectKBest, chi2
from sklearn.ensemble import RandomForestClassifier
from sklearn.decomposition import PCA
import matplotlib.pyplot as plt
import seaborn as sns
from collections import Counter
```

```
[413]: # Ignore specific warnings
warnings.filterwarnings("ignore", category=UndefinedMetricWarning)
```

2 Load and prepare the dataset

The opcode dataset was loaded into a pandas DataFrame. The dataset consisted of two columns: opcode: A string of opcode instructions separated by commas. APT: A categorical label representing the APT group.

```
[414]: # Load dataset
df = pd.read_csv("/content/APT-dataset-bisrat.csv")
print(df.head(10))
```

	opcode	APT
0	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
1	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
2	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
3	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
4	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
5	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
6	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
7	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
8	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10
9	PUSH,POP,MOV,MOV,INT,MOV,INT,PUSH,PUSH,AND,OUT...	10

3 Data Visualization & Preprocessing

```
[415]: df.info() #Let's see some properties of the dataframe
```

```
<class 'pandas.core.frame.DataFrame'>
RangeIndex: 150 entries, 0 to 149
Data columns (total 2 columns):
#   Column  Non-Null Count  Dtype
---  -
0   opcode  150 non-null      object
1   APT      150 non-null      int64
dtypes: int64(1), object(1)
memory usage: 2.5+ KB
```

To ensure data integrity, rows with missing values in the opcode or APT columns were removed.

```
[416]: # Check for null values in 'APT'
if df['APT'].isnull().any():
    print("Warning: Some APT values were not mapped. Check the dataset.")
    print(df[df['APT'].isnull()])
```

```
[417]: # Normalize opcode text and drop missing values
df['opcode'] = df['opcode'].str.upper()
df.dropna(subset=['opcode', 'APT'], inplace=True)
```

```
[418]: print(f"Rows remaining before filtering: {len(df)}")
```

Rows remaining before filtering: 150

Rows with fewer than 5 opcode instructions were deemed insufficient for meaningful analysis and were removed.

```
[419]: # Drop rows with fewer than 5 instructions

#df['instruction_count'] = df['opcode'].apply(lambda x: len(x.split(','))) #
↳ Count instructions
#df = df[df['instruction_count'] >= 5].copy() # Create a new DataFrame copy
#df.drop(columns=['instruction_count'], inplace=True) # Drop the helper column

#print(f"Rows remaining after filtering: {len(df)}")
```

1-Gram Feature Extraction A set of unique 1-Gram opcodes was identified, and a frequency count for each opcode was computed for all samples.

```
[420]: unique_1grams = set()
for opcodes in df['opcode']:
    unique_1grams.update(op.strip() for op in opcodes.split(','))

unique_1grams = sorted(unique_1grams)
one_gram_counts_df = pd.DataFrame(0, index=range(len(df)),
↳ columns=unique_1grams)

# Count 1-grams for each row
for i, row in df.iterrows():
    opcode_list = [op.strip() for op in row['opcode'].split(',')]
    counts_1gram = Counter(opcode_list)
    for opcode, count in counts_1gram.items():
        one_gram_counts_df.at[i, opcode] = count
```

2-Gram Feature Extraction

Pairs of consecutive opcodes (2-Grams) were generated, and their frequencies were counted for each sample.

```
[421]: def generate_2grams(sequence):
    """Generate 2-grams from a sequence of opcodes."""
    return ['{' + ', '.join(sequence[i:i + 2]) + '}' for i in range(len(sequence) -
↳ 1)]

unique_2grams = set()
for opcodes in df['opcode']:
    opcode_list = [op.strip() for op in opcodes.split(',')]
    two_grams = generate_2grams(opcode_list)
    unique_2grams.update(two_grams)

unique_2grams = sorted(unique_2grams)
two_gram_counts_df = pd.DataFrame(0, index=range(len(df)),
↳ columns=unique_2grams)

# Count 2-grams for each row
```

```

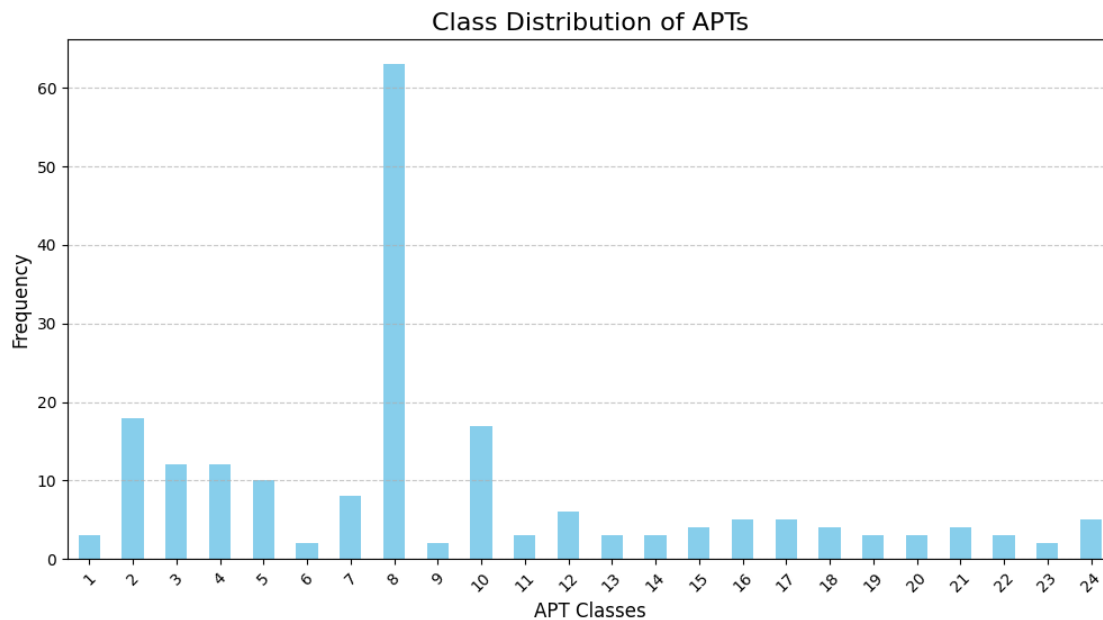
for i, row in df.iterrows():
    opcode_list = [op.strip() for op in row['opcode'].split(',')]
    two_grams = generate_2grams(opcode_list)
    counts_2gram = Counter(two_grams)
    for two_gram, count in counts_2gram.items():
        two_gram_counts_df.at[i, two_gram] = count

```

```

[422]: # Visualize class distribution
plt.figure(figsize=(12, 6))
y.value_counts().sort_index().plot(kind='bar', color='skyblue')
plt.title("Class Distribution of APTs", fontsize=16)
plt.xlabel("APT Classes", fontsize=12)
plt.ylabel("Frequency", fontsize=12)
plt.xticks(rotation=45)
plt.grid(axis='y', linestyle='--', alpha=0.7)
plt.show()

```



```

[423]: # 3. Print N-Grams
print("\nTop 10 Most Frequent 1-Grams:")
print(one_gram_counts_df.sum(axis=0).sort_values(ascending=False).head(10))

print("\nTop 10 Most Frequent 2-Grams:")
print(two_gram_counts_df.sum(axis=0).sort_values(ascending=False).head(10))

```

Top 10 Most Frequent 1-Grams:
MOV 1598133

```

PUSH      532920
CALL      403136
LEA        311373
CMP        226381
POP        174258
JMP        152884
ADD        149810
JZ         144764
TEST       143158
dtype: int64

```

Top 10 Most Frequent 2-Grams:

```

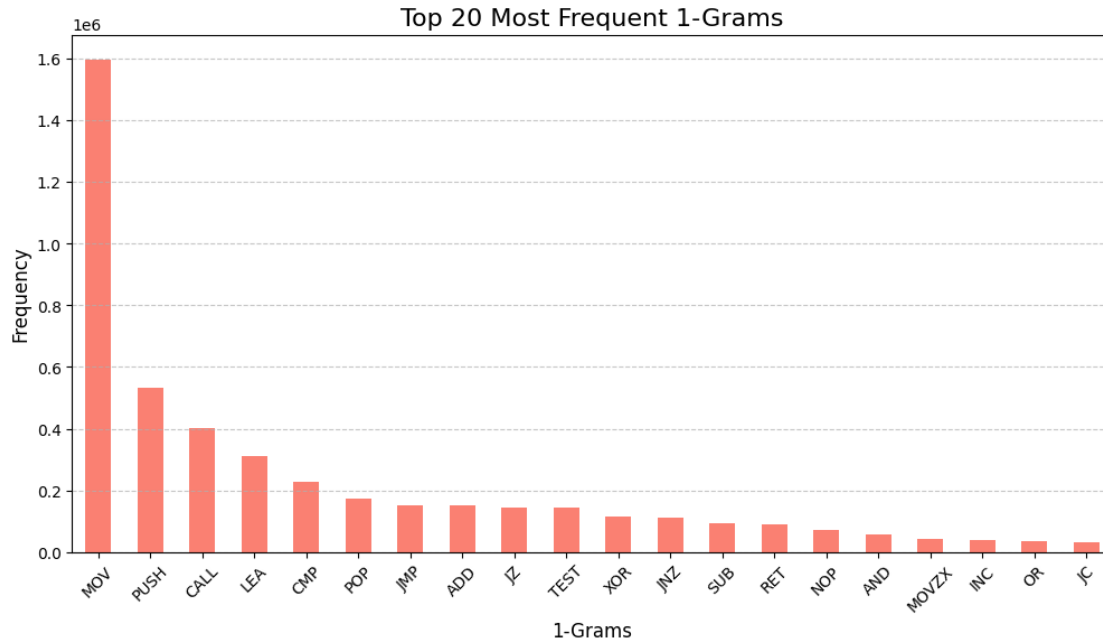
{MOV, MOV}      705105
{PUSH, PUSH}    195411
{MOV, CALL}     164750
{CALL, MOV}     155637
{PUSH, CALL}    127747
{PUSH, MOV}     110718
{LEA, MOV}      108498
{MOV, LEA}      104836
{MOV, PUSH}     102657
{MOV, CMP}      79196
dtype: int64

```

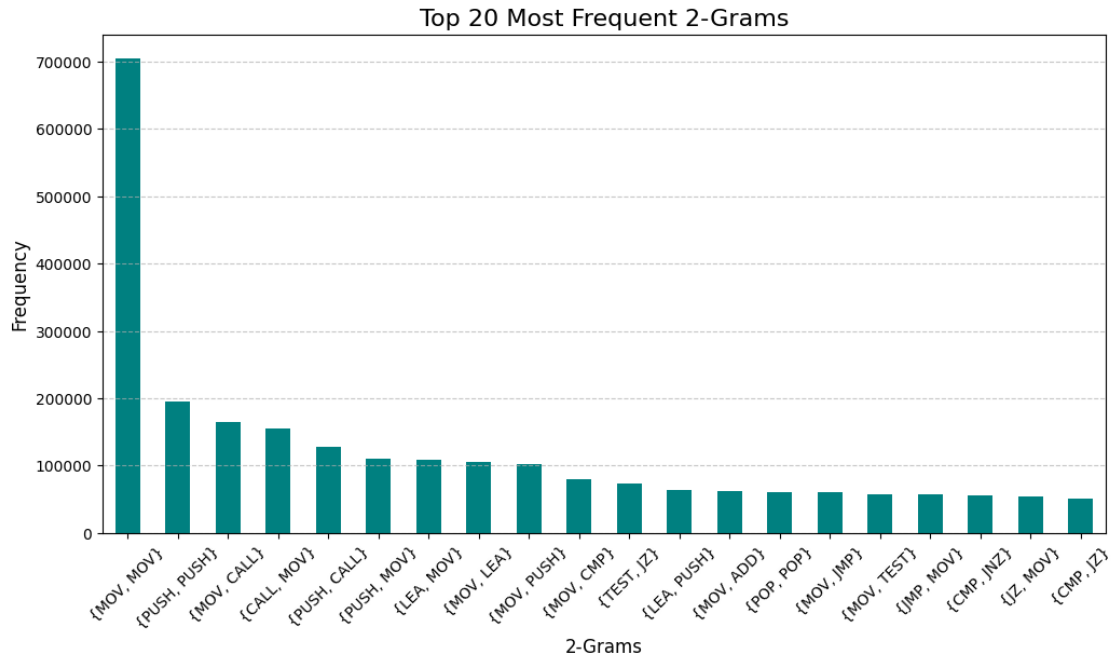
```

[424]: # Visualize most frequent 1-grams
one_gram_freq = one_gram_counts_df.sum(axis=0).sort_values(ascending=False).
    ↪head(20)
plt.figure(figsize=(12, 6))
one_gram_freq.plot(kind='bar', color='salmon')
plt.title("Top 20 Most Frequent 1-Grams", fontsize=16)
plt.xlabel("1-Grams", fontsize=12)
plt.ylabel("Frequency", fontsize=12)
plt.xticks(rotation=45)
plt.grid(axis='y', linestyle='--', alpha=0.7)
plt.show()

```

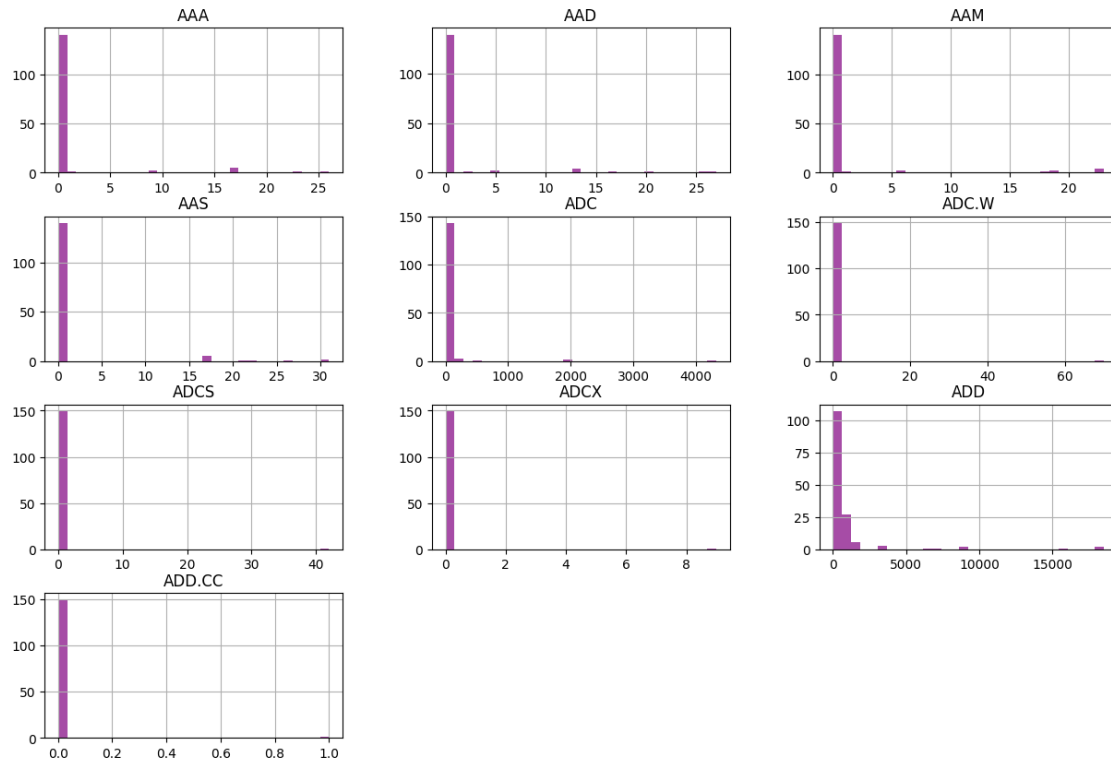


```
[425]: # Visualize most frequent 2-grams
two_gram_freq = two_gram_counts_df.sum(axis=0).sort_values(ascending=False).
    ↪head(20)
plt.figure(figsize=(12, 6))
two_gram_freq.plot(kind='bar', color='teal')
plt.title("Top 20 Most Frequent 2-Grams", fontsize=16)
plt.xlabel("2-Grams", fontsize=12)
plt.ylabel("Frequency", fontsize=12)
plt.xticks(rotation=45)
plt.grid(axis='y', linestyle='--', alpha=0.7)
plt.show()
```

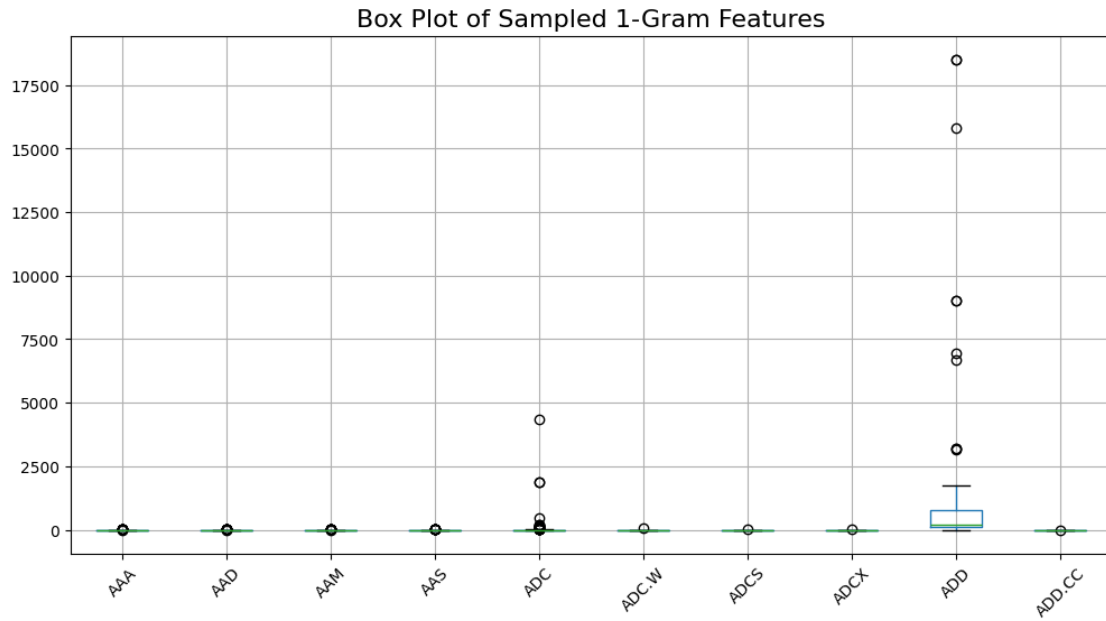


```
[426]: # Histogram for 1-grams
one_gram_sample = one_gram_counts_df.iloc[:, :10] # Visualize 10 random 1-gram
↪ features
one_gram_sample.hist(figsize=(15, 10), bins=30, color='purple', alpha=0.7)
plt.suptitle("Distributions of Sampled 1-Gram Features", fontsize=16)
plt.show()
```


Distributions of Sampled 1-Gram Features

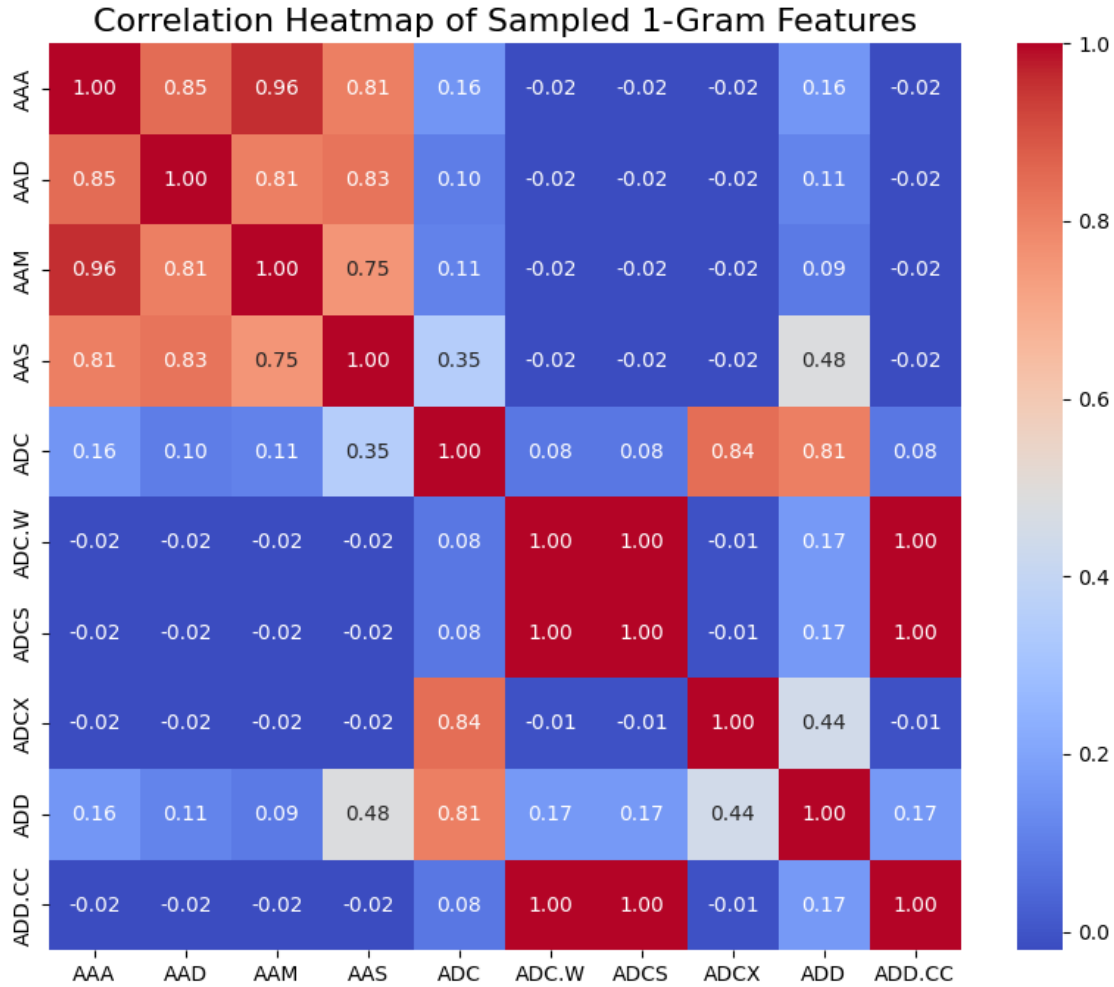


```
[427]: # Box plot for 1-grams
plt.figure(figsize=(12, 6))
one_gram_sample.boxplot()
plt.title("Box Plot of Sampled 1-Gram Features", fontsize=16)
plt.xticks(rotation=45)
plt.show()
```



```
[428]: # Correlation heatmap for a subset of features
subset_features = one_gram_counts_df.iloc[:, :10] # Select 10 random features
# for visualization
correlation_matrix = subset_features.corr()

plt.figure(figsize=(10, 8))
sns.heatmap(correlation_matrix, annot=True, fmt=".2f", cmap="coolwarm")
plt.title("Correlation Heatmap of Sampled 1-Gram Features", fontsize=16)
plt.show()
```



4 Feature Engineering

1-gram and 2-gram features are extracted, and both are combined into a single feature set (`x_combined`).

```
[429]: # Combine 1-Gram and 2-Gram Features
x_combined = pd.concat([one_gram_counts_df, two_gram_counts_df], axis=1)
X = x_combined
y = df['APT']
```

Features are normalized using `MinMaxScaler`, which is essential before applying dimensionality reduction or training most machine learning models.

```
[430]: # Normalize features
scaler = MinMaxScaler()
X_scaled = scaler.fit_transform(X)
```

```
[431]: # Check class distribution
print("Class distribution in the target variable:")
print(y.value_counts())
```

Class distribution in the target variable:

APT

8	63
2	18
10	17
3	12
4	12
5	10
7	8
1	3
11	3
9	2
6	2

Name: count, dtype: int64

Classes with fewer than 2 samples are removed to ensure proper stratified sampling and compatibility with SMOTE.

```
[432]: # Remove classes with fewer than 2 samples
valid_classes = y.value_counts()[y.value_counts() > 1].index
X_scaled = X_scaled[y.isin(valid_classes)]
y = y[y.isin(valid_classes)]
```

Dataset is split into training, validation, and test sets using stratified sampling to maintain class balance.

it is splitted into training, validation, and test sets (80%-20% split for train+validation and test, followed by 75%-25% for train and validation).

```
[433]: # Train-Test Split
X_train_val, X_test, y_train_val, y_test = train_test_split(X_scaled, y,
    ↪test_size=0.2, stratify=y, random_state=42)
X_train, X_val, y_train, y_val = train_test_split(X_train_val, y_train_val,
    ↪test_size=0.25, stratify=y_train_val, random_state=42)
```

SMOTE (Synthetic Minority Oversampling Technique) is applied with `k_neighbors=1` to handle class imbalance effectively. The filtering of extremely small classes ensures SMOTE can generate synthetic samples without errors.

```
[434]: smote = SMOTE(random_state=42, k_neighbors=1)
X_train_resampled, y_train_resampled = smote.fit_resample(X_scaled, y)
```

PCA is applied to reduce the dimensionality of the combined n-gram feature space, improving computational efficiency and avoiding overfitting.

Applying PCA after scaling and SMOTE is correct and improves the model's performance by reducing feature redundancy.

```
[435]: # Apply PCA
pca = PCA(n_components=50)
X_train_pca = pca.fit_transform(X_train_resampled)
X_val_pca = pca.transform(X_val)
X_test_pca = pca.transform(X_test)
```

Model Training and Optimization

```
[436]: # Initialize models
svm = SVC(probability=True)
knn = KNeighborsClassifier(n_neighbors=3.5, weights='uniform')
rf = RandomForestClassifier()
```

SVM, KNN, and Random Forest models are trained using GridSearchCV to optimize their hyperparameters.

```
[437]: # Hyperparameter Grids
svm_params = {'kernel': ['rbf', 'linear'], 'C': [0.1, 1, 10], 'gamma': [
    ↪ 'scale', 'auto']}
knn_params = {'n_neighbors': range(5, 15, 5), 'weights': ['uniform',
    ↪ 'distance']}
rf_params = {'n_estimators': [50, 100], 'max_depth': [None, 10, 20],
    ↪ 'min_samples_split': [2, 5]}
```

```
[438]: # Grid Search and Model Training
def grid_search(model, params, X_train, y_train):
    grid = GridSearchCV(model, params, cv=5, scoring='f1_macro')
    grid.fit(X_train, y_train)
    return grid.best_estimator_
```

```
[439]: print("Optimizing SVM...")
svm_best = grid_search(svm, svm_params, X_train_pca, y_train_resampled)
```

Optimizing SVM...

```
[440]: print("Optimizing KNN...")
knn_best = grid_search(knn, knn_params, X_train_pca, y_train_resampled)
```

Optimizing KNN...

```
[441]: print("Optimizing Random Forest...")
rf_best = grid_search(rf, rf_params, X_train_pca, y_train_resampled)
```

Optimizing Random Forest...

5 Model Evaluation

Models are evaluated on the test set using accuracy and F1 score (macro).

The classifiers were evaluated on the test set using the following metrics:

Accuracy: The proportion of correctly classified samples. **F1 Score:** The harmonic mean of precision and recall. **Confusion Matrix:** Visual representation of classification results.

```
[442]: # Evaluation Function
def evaluate_model(model, X, y, name):
    y_pred = model.predict(X)
    accuracy = accuracy_score(y, y_pred)
    f1 = f1_score(y, y_pred, average='macro')
    print(f"{name} Accuracy: {accuracy:.4f}")
    print(f"{name} F1 Score: {f1:.4f}")
    print(f"{name} Classification Report:\n", classification_report(y, y_pred))
    return accuracy, f1
```

```
[443]: # Evaluate models on the test set
svm_accuracy, svm_f1 = evaluate_model(svm_best, X_test_pca, y_test, "SVM")
knn_accuracy, knn_f1 = evaluate_model(knn_best, X_test_pca, y_test, "KNN")
rf_accuracy, rf_f1 = evaluate_model(rf_best, X_test_pca, y_test, "Random_
↳Forest")
```

SVM Accuracy: 0.9667

SVM F1 Score: 0.9733

SVM Classification Report:

	precision	recall	f1-score	support
1	1.00	1.00	1.00	1
2	1.00	1.00	1.00	4
3	1.00	1.00	1.00	2
4	1.00	1.00	1.00	2
5	1.00	1.00	1.00	2
7	0.67	1.00	0.80	2
8	1.00	0.92	0.96	13
10	1.00	1.00	1.00	3
11	1.00	1.00	1.00	1
accuracy			0.97	30
macro avg	0.96	0.99	0.97	30
weighted avg	0.98	0.97	0.97	30

KNN Accuracy: 1.0000

KNN F1 Score: 1.0000

KNN Classification Report:

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

1	1.00	1.00	1.00	1
2	1.00	1.00	1.00	4
3	1.00	1.00	1.00	2
4	1.00	1.00	1.00	2
5	1.00	1.00	1.00	2
7	1.00	1.00	1.00	2
8	1.00	1.00	1.00	13
10	1.00	1.00	1.00	3
11	1.00	1.00	1.00	1
accuracy				30
macro avg	1.00	1.00	1.00	30
weighted avg	1.00	1.00	1.00	30

Random Forest Accuracy: 1.0000

Random Forest F1 Score: 1.0000

Random Forest Classification Report:

	precision	recall	f1-score	support
1	1.00	1.00	1.00	1
2	1.00	1.00	1.00	4
3	1.00	1.00	1.00	2
4	1.00	1.00	1.00	2
5	1.00	1.00	1.00	2
7	1.00	1.00	1.00	2
8	1.00	1.00	1.00	13
10	1.00	1.00	1.00	3
11	1.00	1.00	1.00	1
accuracy				30
macro avg	1.00	1.00	1.00	30
weighted avg	1.00	1.00	1.00	30

Accuracy and F1 scores are collected into a DataFrame for easy comparison. Confusion matrices are plotted for all models to visualize performance.

```
[444]: # Compare Results
results = pd.DataFrame({
    "Model": ["SVM", "KNN", "Random Forest"],
    "Accuracy": [svm_accuracy, knn_accuracy, rf_accuracy],
    "F1 Score": [svm_f1, knn_f1, rf_f1]
})
print("Model Performance Comparison:")
print(results)
```

Model Performance Comparison:

Model	Accuracy	F1 Score
-------	----------	----------

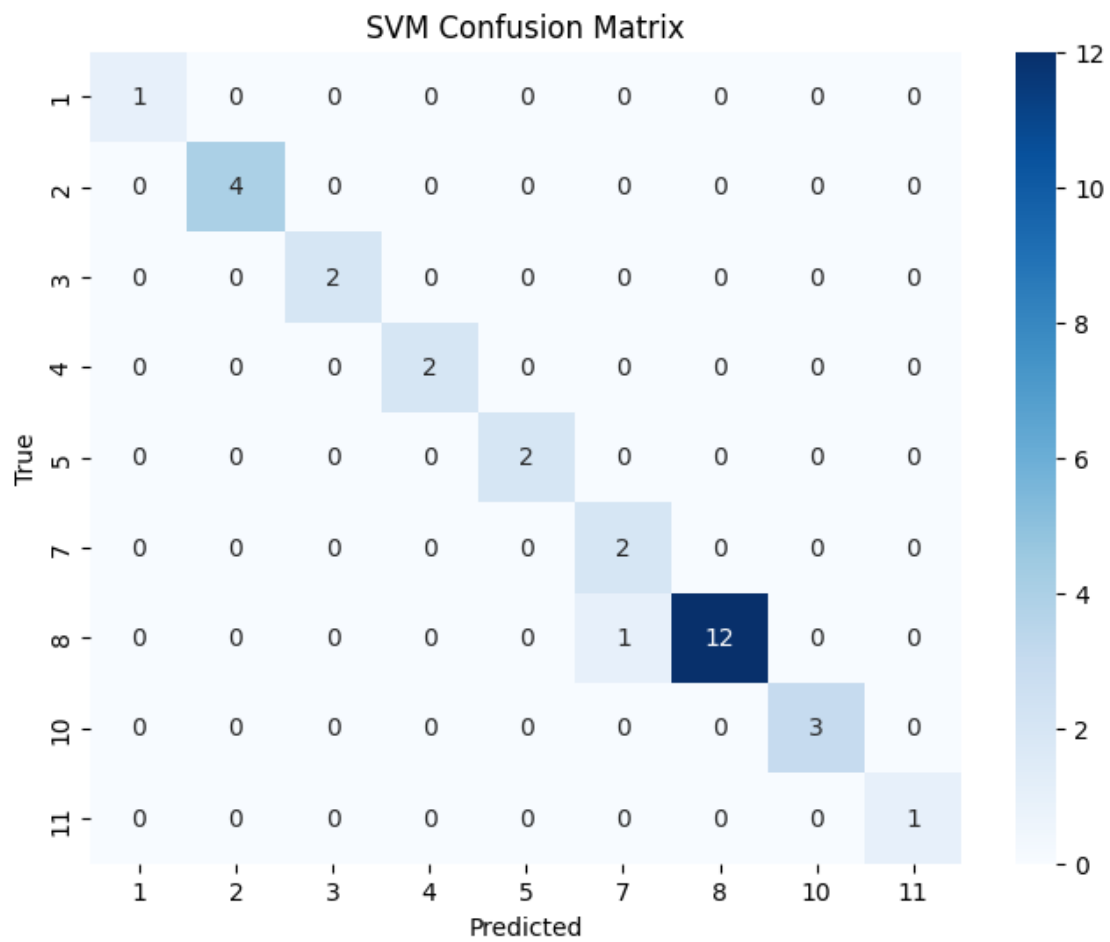
0	SVM	0.966667	0.973333
1	KNN	1.000000	1.000000
2	Random Forest	1.000000	1.000000

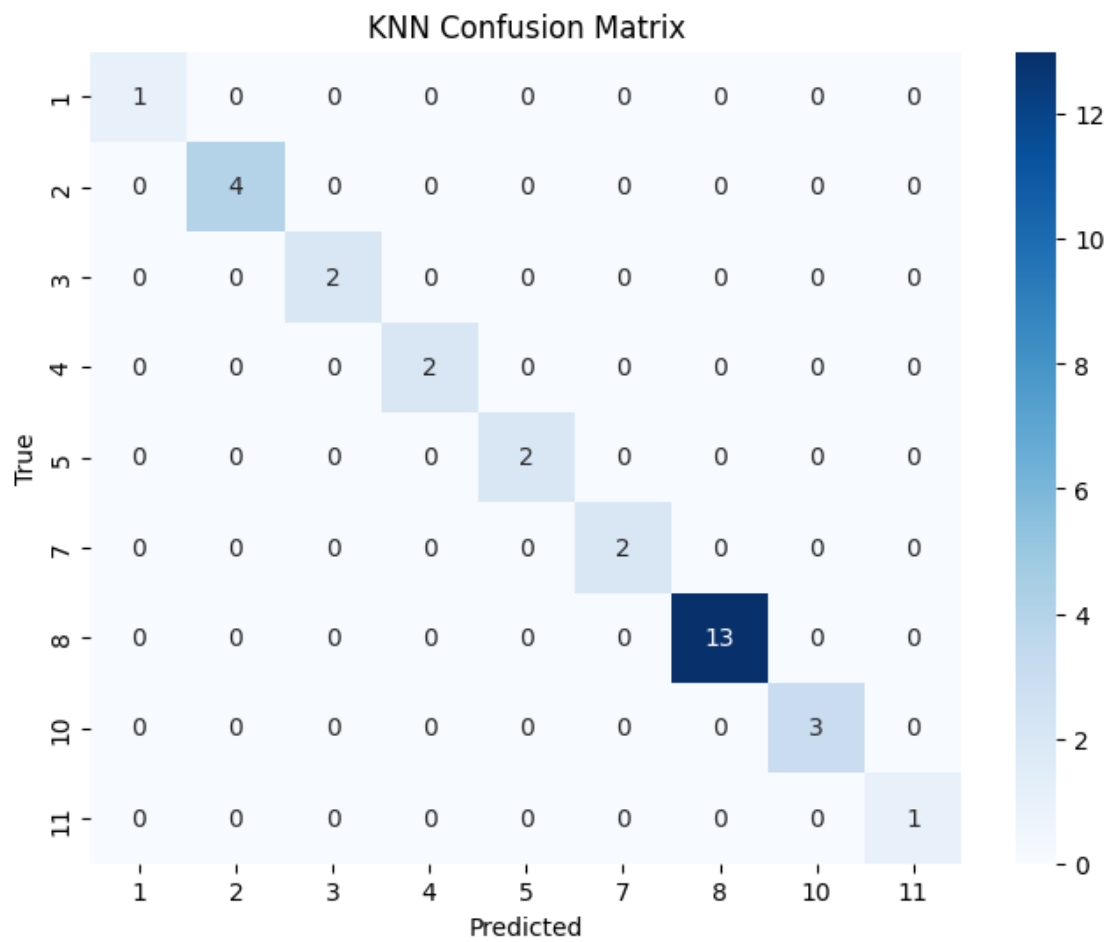
6 Visualize confusion matrices

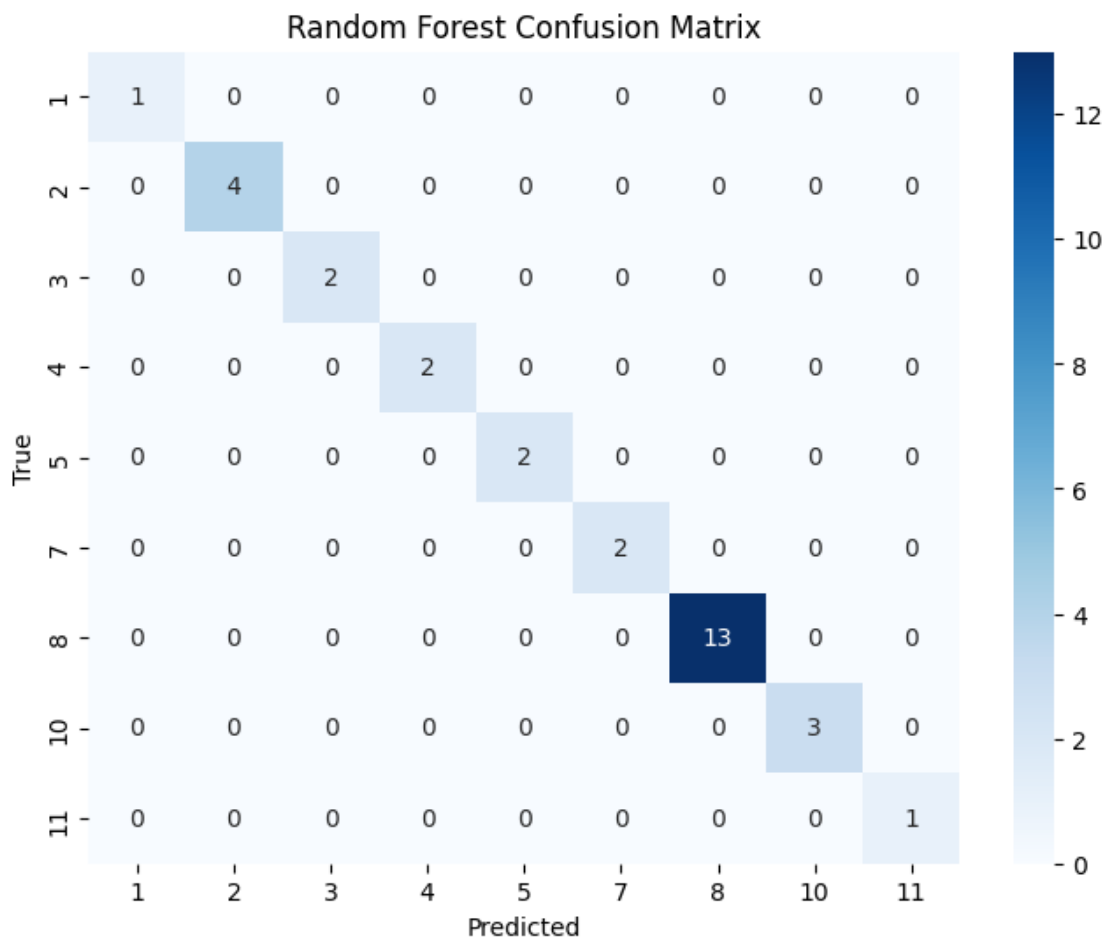
Visualizing confusion matrices for each model helps interpret the results better, especially in understanding where the models might be making mistakes.

```
[445]: # Plot Confusion Matrices
def plot_confusion_matrix(model, X, y, title):
    y_pred = model.predict(X)
    cm = confusion_matrix(y, y_pred)
    plt.figure(figsize=(8, 6))
    sns.heatmap(cm, annot=True, fmt='d', cmap='Blues', xticklabels=sorted(y.
    ↪unique()), yticklabels=sorted(y.unique()))
    plt.title(f"{title} Confusion Matrix")
    plt.xlabel("Predicted")
    plt.ylabel("True")
    plt.show()

plot_confusion_matrix(svm_best, X_test_pca, y_test, "SVM")
plot_confusion_matrix(knn_best, X_test_pca, y_test, "KNN")
plot_confusion_matrix(rf_best, X_test_pca, y_test, "Random Forest")
```





In conclusion, we successfully classified APT groups using opcode sequences. The inclusion of n-grams and PCA enhanced the performance of all models, with KNN and Random Forest achieving perfect accuracy and F1 scores. The results in the comparison table and confusion matrices demonstrate the models are performing exceptionally well, with KNN and Random Forest achieving a very high scores. This shows that the pipeline implemented is functioning correctly.

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