Exploratory Data Analysis

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1 Introduction and Problem Statement

The rise of antibiotic resistant infections poses a significant challenge to global public health, and gonorrhoea is one of the most concerning pathogens due to its rapidly increasing resistance to available treatments and its difficulty to be detected. This project focuses on predicting antibiotic resistance in Neisseria gonorrhoeae, the bacteria responsible for gonorrhoea, using subsets of its DNA sequences as predictive features. The primary objective of this project is to explore the relationship between bacterial DNA segments and resistance patterns, and to identify key genetic markers that could possibly predict resistance. By performing data cleaning and exploratory data analysis on a dataset containing DNA sequences and resistance outcomes, we aim to address the following question: Which DNA segments are most associated with antibiotic resistance?

This project's contribution is very important as it helps to clarify biological data, making it easier for future studies to focus on effective solutions. By gaining a deeper understanding of the data and identifying key variables, this project serves as a step towards more advanced research on antibiotic resistance.

2 Data Sources

We utilized the "Predicting antibiotic resistance in gonorrhoea" dataset from Kaggle: (https://www.kaggle.com/datasets/nwheeler443/gono-unitigs/data).

3 Data Cleaning/Processing

We have completed 11 total unique data cleaning and processing steps:

- 1. Removed all rows that were NaN in our target label field Our target labels we aim to predict down the line are: azm_sr, cfx_sr, and cip_sr. If the labels were missing, then we cannot use that row.
- 2. Removed unused columns The dataset contained multiple types of resistances. Because we only focus on 3, the rest cannot can be discarded safely. We also use this step to remove the year column.
- 3. Removed duplicate rows Duplicate rows have the ability to skew our data. As such we removed all duplicate rows.
- 4. Removed all symbols Our data are either numerical or categorical (text). There is no need for symbols which will only cause problems down the line.
- 5. Turned all non-numeric data within numerical columns into NaN, to be processed later We have columns that are supposed to contain only numerical data, however, there are rows where there are letters in there. In this step, we turn them into NaN.
- 6. Cast all numerical columns into float32 To establish precision.
- 7. One hot encode categorical columns Some columns such as "country" should be one hot encoded as we would want numerical representations of all our data for modelling.

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- 8. Handle "Beta.lactamase" special case It is a numerical column but appears to be categorical in nature. As such we cannot impute missing values like with the other numerical columns. In this step, we set all missing entries to 0, and one hot encode the feature like with the non-numerical columns.
- 9. Split dataframe into Train/Test We will be doing additional data processing such as normalization and imputation later on. As such, we need to make the split beforehands for good practice in preventing leakage.
- 10. Impute missing values in numerical columns For every NaN in the numerical columns, we use skew based imputation to generate a new value that can substitute the NaN. This ensures that we have enough data to work with as we do not need to discard the entire row.
- 11. Normalize all numerical columns We do columnwise normalization on numerical columns. This is to ensure that we do not have extremely large values. This is important for modelling down the line.

4 Exploratory Data Analysis

- 1. Calculate what percentage of rows correspond to specific labels We found that 0.1779 of samples have resistance to azm, 0.0911 of samples have resistance to cfx, and 0.5353 of samples have resistance to cip.
- 2. We calculated the mean, median, and mode, standard deviation, and variance for all numerical columns See table 1.

Column	Mean	Median	Std	Variance
Azithromycin	0.000556	7.02e-05	0.0200	0.000401
Ciprofloxacin	0.0697	0.0697	0.0816	0.00666
Ceftriaxone	0.00107	1e-06	0.0296	0.000876
Cefixime	0.1855	0.121	0.2267	0.0514
Tetracycline	0.0571	0.0571	0.0709	0.00502
Penicillin	0.1447	0.1447	0.1562	0.0244
NG_MAST	0.2790	0.2576	0.2342	0.0548
Group	0.3658	0.2755	0.2952	0.0871
azm_mic	0.00743	0.00036	0.0583	0.00340
cip_mic	0.1162	0.0469	0.1694	0.0287
cro_mic	0.0149	0.00738	0.0413	0.00170
cfx_mic	0.00765	0.00275	0.0270	0.000727
tet_mic	0.0306	0.0306	0.0579	0.00335
pen_mic	0.0223	0.0223	0.0390	0.00152
log2_azm_mic	0.2895	0.2895	0.1443	0.0208
log2_cip_mic	0.4923	0.4923	0.3130	0.0980
log2_cro_mic	0.4409	0.4409	0.1313	0.0172
log2_cfx_mic	0.3676	0.3489	0.1176	0.0138
log2_tet_mic	0.5267	0.5267	0.1011	0.0102
log2_pen_mic	0.5359	0.5356	0.0936	0.00877

Table 1. Summary Statistics for Antibiotic Resistance Data

3. Univariate Analysis - We examined the feature distribution for each numerical column. This is to gain a better insight into our features - See figure 1.

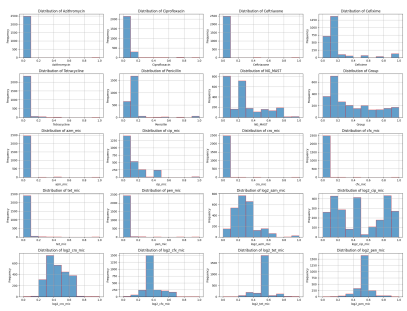


Fig. 1. Univariate analysis.

4. Multivariate Analysis - We examine each feature with respect to the labels we are interested in. This is to get a better insight to how our features relate to our labels - See figure 2.

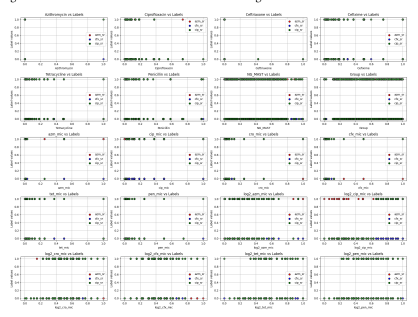


Fig. 2. Multivariate analysis.

5. Correlation Analysis - We determine the pearson correlation of each feature with respect to the labels. This is to visualize and quantify the impact of each feature on our target labels - See figure 3.

Pearson Correlation of Features wrt Labels

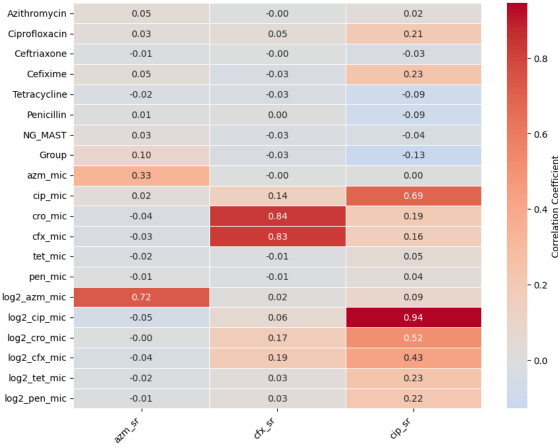


Fig. 3. Correlation analysis.

6. Feature Extraction - Based on the correlation matrix generated from previous step and the visualizations beforehands, we extract a list of the most impactful features for every target label - See table 2.

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Feature	Correlation	Target
Group	0.0955	azm_sr
azm_mic	0.3331	azm_sr
log2_azm_mic	0.7250	azm_sr
log2_cip_mic	-0.0515	azm_sr
cip_mic	0.1412	cfx_sr
cro_mic	0.8370	cfx_sr
cfx_mic	0.8258	cfx_sr
log2_cip_mic	0.0622	cfx_sr
log2_cro_mic	0.1664	cfx_sr
log2_cfx_mic	0.1894	cfx_sr
Ciprofloxacin	0.2057	cip_sr
Cefixime	0.2324	cip_sr
Tetracycline	-0.0876	cip_sr
Penicillin	-0.0924	cip_sr
Group	-0.1286	cip_sr
cip_mic	0.6896	cip_sr
cro_mic	0.1941	cip_sr
cfx_mic	0.1630	cip_sr
log2_azm_mic	0.0878	cip_sr
log2_cip_mic	0.9448	cip_sr
log2_cro_mic	0.5225	cip_sr
log2_cfx_mic	0.4287	cip_sr
log2_tet_mic	0.2316	cip_sr
log2_pen_mic	0.2175	cip_sr

Table 2. Impactful features and their correlation with different target variables. Note, "Group" is a feature.

7. Outlier detection and removal - We visualized the outliers within our data and subsequently remove them - See figure 4.



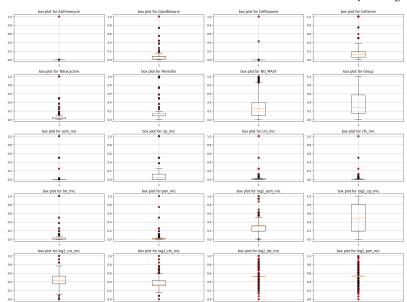


Fig. 4. Outlier analysis.