```
import pandas as pd
import numpy as np
import plotly.express as px
import plotly.graph_objects as go
import plotly.io as pio
from pprint import pprint
import seaborn as sns
```

patient_data = pd.read_csv('PatientData.csv')
patient_data.head()

| | | Patient_Status | Т | RR |
|-------------|---|----------------|-----------|-----------|
| | 0 | HEALTHY | 35.821266 | 19.142584 |
| | 1 | HEALTHY | 36.353128 | 20.759505 |
| | 2 | SICK | 37.352552 | 22.507282 |
| | 3 | HEALTHY | 38.645271 | 20.855217 |
| | 4 | SICK | 38.987791 | 20.886224 |

(Q1) You decide to fit a 2D Bayesian classifier to your data, where x=[T,RR], SICK is the 'positive' class, and we assume that $p(x|0_1)$ $N(\mu\iota,\Sigma_1)$. Use unbiased estimators to estimate the 2D mean and covariance matrix for each class-conditional distribution. Report your two estimated mean vectors and covariance matrices.

Q1 Answer:

Mean: [38.39730213 21.97277948]

Covariance Matrix: [[3.36436875 0.08317142] [0.08317142 0.79046862]]

$$\operatorname{Cov}(X,Y) = rac{1}{n-1} \sum_{i=1}^n (X_i - ar{X})(Y_i - ar{Y})$$

For the covariance, an unbiased estimator adjusts the denominator from n to n-1. This adjustment corrects for the bias that arises when estimating population variance or covariance from a sample.

```
# Separate data into two classes: Healthy and Sick
healthy data = patient data[patient data['Patient Status'] == 'HEALTHY'][['T', 'RR']]
sick_data = patient_data[patient_data['Patient_Status'] == 'SICK'][['T', 'RR']]
# Calculate the unbiased estimators for mean and covariance for each class
mean_healthy = healthy_data.mean().values
cov_healthy = healthy_data.cov().values
mean_sick = sick_data.mean().values
cov_sick = sick_data.cov().values
print("Unbiased Estimators for Healthy Class:")
print("Mean:", mean_healthy)
print("Covariance Matrix:\n", cov_healthy)
print("\nUnbiased Estimators for Sick Class:")
print("Mean:", mean_sick)
print("Covariance Matrix:\n", cov_sick)

→ Unbiased Estimators for Healthy Class:
    Mean: [36.77292777 20.03604645]
    Covariance Matrix:
     [[ 2.11598617e+00 -2.32805447e-04]
     [-2.32805447e-04 2.97857263e-01]]
    Unbiased Estimators for Sick Class:
```

(Q2) Based on historical effluent testing data, we know that approximately 20% of guests will have MF. Use Bayes' theorem to compute the posterior probability that a patient with a temperature of 37.5 degrees and a respiration rate of 26 is sick (numeric answer, show equation).

Q2 Answer:

Bayes' Theorem

$$P(w_1|x)=rac{P(x|w_1)P(w_1)}{P(x)}$$

Where:

- $P(w_1|x)$ is the posterior probability that the patient is **SICK** given the observed data x.
- $P(x|w_1)$ is the likelihood of observing the data x given that the patient is **SICK**.
- $P(w_1)$ is the prior probability that a patient is **SICK**, which we know to be 20% (or 0.2).
- P(x) is the evidence, which is the total probability of observing the data x.

The observed data x is represented by:

$$x = [T, RR] = [37.5, 26]$$

We need to calculate the posterior probability $P(w_1|x)$, where w_1 represents the **SICK** class.

The likelihood $p(x|w_1)$ is modeled as a bivariate normal (Gaussian) distribution.

$$p(x|w_1) = rac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \mathrm{exp}igg(-rac{1}{2}(x-\mu)^T \Sigma^{-1}(x-\mu)igg)$$

Where:

- μ is the mean vector for the **SICK** class.
- Σ is the covariance matrix for the **SICK** class.
- d is the dimensionality of the data (in this case, d=2).
- ullet x is the observed data point.

We already have:

- Mean vector for SICK: $\mu_{SICK} = [38.40, 21.97]$
- Covariance matrix for SICK:

$$\Sigma_{SICK} = egin{bmatrix} 3.364 & 0.083 \ 0.083 & 0.790 \end{bmatrix}$$

```
from scipy.stats import multivariate_normal

#[T, RR]
x = np.array([37.5, 26])

mean_sick_vector = mean_sick
cov_sick_matrix = cov_sick

# Prior 'SICK'
from scipy.stats import multivariate_normal

#[T, RR]
x = np.array([37.5, 26])

mean_sick_vector = mean_sick
cov_sick_matrix = cov_sick
```

Prior 'SICK'
P_sick = 0.2
likelihood_sick = multivariate_normal.pdf(x, mean=mean_sick_vector, cov=cov_sick_matrix)
print(f'Likelihood of observing the data x given that the patient is SICK:{likelihood_sick}')

Likelihood of observing the data x given that the patient is SICK:2.640732348188037e-06

The p(x) is computed by considering both classes (**HEALTHY** and **SICK**):

$$p(x) = p(x|w_1)P(w_1) + p(x|w_0)P(w_0)$$

Where:

- w_0 represents the **HEALTHY** class.
- $P(w_0) = 1 P(w_1) = 0.8$

mean_healthy_vector = mean_healthy
cov_healthy_matrix = cov_healthy

Prior 'HEALTHY'
P_healthy = 0.8

likelihood_healthy = multivariate_normal.pdf(x, mean=mean_healthy_vector, cov=cov_healthy_matrix)
print("Likelihood of observing the data x given that the patient is HEALTHY:", likelihood_healthy)

p(x)
evidence = (likelihood_sick * P_sick) + (likelihood_healthy * P_healthy)
print("Evidence (total probability of observing the data x):", evidence)

Likelihood of observing the data x given that the patient is HEALTHY: 2.0719974466511435e-27 Evidence (total probability of observing the data x): 5.281464696376074e-07

Now, we can use Bayes' theorem to calculate the posterior probability that the patient is SICK:

$$P(w_1|x)=rac{P(x|w_1)P(w_1)}{P(x)}$$

posterior_sick = (likelihood_sick * P_sick) / evidence
print("Posterior Probability that the patient is SICK given the observed data x:", posterior_sick)

Posterior Probability that the patient is SICK given the observed data x: 1.0

In General, the general formula for Bayes' theorem:

$$P(\text{sick}|x) = \frac{p(x|\text{sick})P(\text{sick})}{p(x|\text{sick})P(\text{sick}) + p(x|\text{healthy})P(\text{healthy})}$$

where $p(x|\omega_i)$ is the multivariate Gaussian density:

$$p(x|\omega_i) = rac{1}{(2\pi)^{d/2}|\Sigma_i|^{1/2}} \mathrm{exp}igg(-rac{1}{2}(x-\mu_i)^T\Sigma_i^{-1}(x-\mu_i)igg)$$

Substituting our case's specific prior probabilities:

$$P(\operatorname{sick}|x) = \frac{p(x|\operatorname{sick})(0.2)}{p(x|\operatorname{sick})(0.2) + p(x|\operatorname{healthy})(0.8)}$$

(Q3) If that guest (T=37.5) wanted to fool the health sensor and held their breath (i.e., RR=0), would it work? Answer this question by computing the posterior probability of the guest being sick with T=37.5 and R=0. Explain your answer by plotting the likelihood of the guest being sick, the likelihood of the guest being healthy, along with the posterior probability of the guest being sick, as a function of RR (for fixed T). (numeric answer, graph, and ~25 words of explanation)

Q3 Answer:

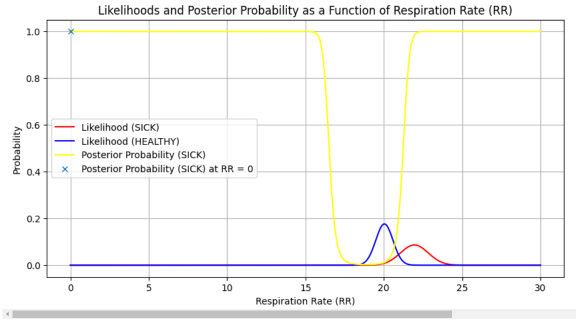
$$P(w=sick|x_1:T=37.5,x_2:RR=0)$$

$$P(\mathrm{sick}|x) = \frac{p(x|\mathrm{sick})P(\mathrm{sick})}{p(x|\mathrm{sick})P(\mathrm{sick}) + p(x|\mathrm{healthy})P(\mathrm{healthy})}$$

The posterior probability of the guest being SICK with T=37.5 and RR=0 is approximately 1.

```
import matplotlib.pyplot as plt
rr values = np.linspace(0, 30, 300)
t_value = 37.5
likelihood_sick_values = []
likelihood_healthy_values = []
posterior_sick_values = []
for rr in rr_values: #checking every possible values between 0-30 for RR
      P(sick|x)=p(x|sick)P(sick)p(x|sick)P(sick)+p(x|healthy)P(healthy)
      likelihood_sick_rr = p(x|sick)
      likelihood_healthy_rr = p(x|healthy)
      evidence\_rr = p(x|sick)P(sick)+p(x|healthy)P(healthy)
                                                            --> P(sick|x)=p(x|sick)P(sick)p(x|sick)P(sick)+p(x|healthy)P(healthy)
      posterior_sick_rr = p(x|sick)P(sick)/evidence_rr
    x_rr = np.array([t_value, rr])
    likelihood_sick_rr = multivariate_normal.pdf(x_rr, mean=mean_sick_vector, cov=cov_sick_matrix)
    likelihood_healthy_rr = multivariate_normal.pdf(x_rr, mean=mean_healthy_vector, cov=cov_healthy_matrix)
    evidence_rr = (likelihood_sick_rr * P_sick) + (likelihood_healthy_rr * P_healthy)
    posterior_sick_rr = (likelihood_sick_rr * P_sick) / evidence_rr
    likelihood sick values.append(likelihood sick rr)
    likelihood_healthy_values.append(likelihood_healthy_rr)
    posterior_sick_values.append(posterior_sick_rr)
# Plotting the results
plt.figure(figsize=(10, 5))
plt.plot(rr_values, likelihood_sick_values, label='Likelihood (SICK)', color='red')
plt.plot(rr_values, likelihood_healthy_values, label='Likelihood (HEALTHY)', color='blue')
plt.plot(rr_values, posterior_sick_values, label='Posterior Probability (SICK)', color='yellow')
# Calculate the posterior probability for RR = 0
x_r_0 = np.array([t_value, 0])
likelihood sick rr \theta = multivariate normal.pdf(x rr \theta, mean=mean sick vector, cov=cov sick matrix)
likelihood\_healthy\_rr\_0 = multivariate\_normal.pdf(x\_rr\_0, mean=mean\_healthy\_vector, cov=cov\_healthy\_matrix)
evidence_rr_0 = (likelihood_sick_rr_0 * P_sick) + (likelihood_healthy_rr_0 * P_healthy)
posterior_sick_rr_0 = (likelihood_sick_rr_0 * P_sick) / evidence_rr_0
print("Posterior Probability that the patient is SICK given RR = 0:", posterior_sick rr_0)
plt.plot(0, posterior\_sick\_rr\_0, 'x', label='Posterior Probability (SICK) at RR = 0')
plt.xlabel('Respiration Rate (RR)')
plt.ylabel('Probability')
plt.title('Likelihoods and Posterior Probability as a Function of Respiration Rate (RR)')
plt.legend()
plt.grid(True)
plt.show()
```

 \rightarrow Posterior Probability that the patient is SICK given RR = 0: 1.0



Posterior Probability that the patient is SICK given RR = 0: 1.0

The graph shows that the posterior probability of being sick remains high across a range of respiration rates, indicating that holding one's breath does not significantly reduce the probability of being identified as sick.



- Q4) For the Bayesian classifier in Q2, compute the probability that each of the 200 guests has MF, given their observed T and RR. Do not report the posterior probability for each patient. Instead, plot an ROC and a P-R curve for your classifier over these 200 patients.
 - For the ROC plot, include the AUC-ROC in the title.
 - For the P-R curve, include the average precision (across all recall values) in the title.

Q4 Answer:

We assume everyone is sick, So we calculate this posterior for all guests. Same as we did in Q2.

```
from sklearn.metrics import roc_curve, auc, precision_recall_curve, average_precision_score

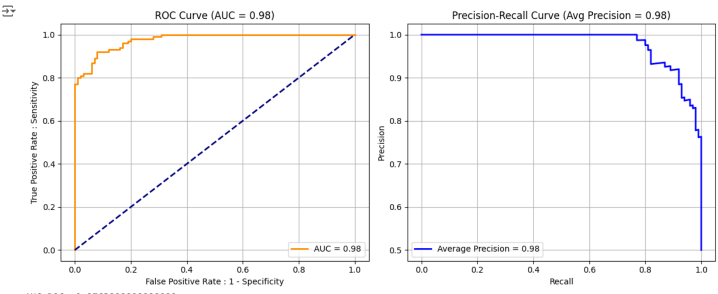
X = patient_data[['T', 'RR']].values
y_true = (patient_data['Patient_Status'] == 'SICK').astype(int).values

posterior_probabilities = []
posterior_probabilities = np.ones(X.shape[0])

for idx, x_point in enumerate(X): #Going through all the guests

    likelihood_sick_point = multivariate_normal.pdf(x_point, mean=mean_sick_vector, cov=cov_sick_matrix)
    likelihood_healthy_point = multivariate_normal.pdf(x_point, mean=mean_healthy_vector, cov=cov_healthy_matrix)
    evidence_point = (likelihood_sick_point * P_sick) + (likelihood_healthy_point * P_healthy)
    posterior_sick_point = (likelihood_sick_point * P_sick) / evidence_point
    posterior_probabilities[idx] = posterior_sick_point
```

```
fpr, tpr, thresh_roc = roc_curve(y_true, posterior_probabilities)
roc_auc = auc(fpr, tpr)
# PrRl curve
precision, recall, _ = precision_recall_curve(y_true, posterior_probabilities)
avg_precision = average_precision_score(y_true, posterior_probabilities)
plt.figure(figsize=(12, 5))
plt.subplot(1, 2, 1)
plt.plot(fpr, tpr, color='darkorange', lw=2, label=f'AUC = \{roc\_auc:.2f\}')
plt.plot([0, 1], [0, 1], color='navy', lw=2, linestyle='--')
plt.xlabel('False Positive Rate : 1 - Specificity')
plt.ylabel('True Positive Rate : Sensitivity')
plt.title(f'ROC Curve (AUC = {roc_auc:.2f})')
plt.legend(loc='lower right')
plt.grid(True)
# Plot Precision-Recall curve
plt.subplot(1, 2, 2)
plt.plot(recall, precision, color='blue', lw=2, label=f'Average Precision = {avg_precision:.2f}')
plt.xlabel('Recall')
plt.ylabel('Precision')
plt.title(f'Precision-Recall Curve (Avg Precision = {avg_precision:.2f})')
plt.legend(loc='lower left')
plt.grid(True)
plt.tight_layout()
plt.show()
print("AUC-ROC:", roc_auc)
print("Average Precision:", avg_precision)
```



▼.

Q5) Every time you stop a truly HEALTHY person from entering the gondola, they get very angry. Therefore, your boss says that "no more than 5% of HEALTHY guests

should be banned from using the gondola". i) What is the maximum sensitivity we can

achieve? ii) What is the maximum precision that we can achieve at the same sensitivity as part i? iii) Report a confusion matrix for this decision threshold.

∨ Q5 Answer:

i)

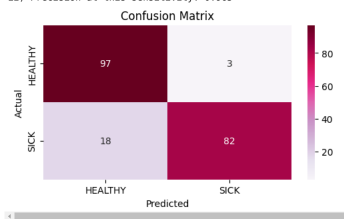
- Every time you stop a truly healthy person from entering the gondola, they become very angry:
 - Meaning that our **False Positive Rate (FPR)** must be ≤ 5%
- Therefore, your boss says that "no more than 5% of HEALTHY guests should be banned from using the gondola".

$$FPR = rac{FP}{FP + TN}$$

Our goal is to have $FPR \leq 0.05$.

```
# maximum sensitivity (TPR) where FPR <= 0.05</pre>
max_sensitivity = tpr[fpr <= 0.05].max()</pre>
optimal idx = np.where((fpr \le 0.05) \& (tpr == max sensitivity))[0][0]
optimal_threshold = thresh_roc[optimal_idx]
# y_pred for this thresh
y_pred = (posterior_probabilities >= optimal_threshold).astype(int)
# Calculate confusion matrix
conf_matrix = confusion_matrix(y_true, y_pred)
\# Calculate precision at this threshold TP / TP + FP
confusion_matrix = [
              # Row 0: Actual HEALTHY
    [TN, FP],
    [FN, TP]
                 # Row 1: Actual SICK
]
precision = conf_matrix[1,1] / (conf_matrix[1,1] + conf_matrix[0,1])
print(f"i) Maximum achievable sensitivity while maintaining FPR ≤ 5%: {max sensitivity:.3f}")
print(f"ii) Precision at this sensitivity: {precision:.3f}")
plt.figure(figsize=(6, 3))
sns.heatmap(conf_matrix, annot=True, fmt='d', cmap='PuRd', xticklabels=['HEALTHY', 'SICK'], yticklabels=['HEALTHY', 'SICK'])
plt.xlabel('Predicted')
plt.ylabel('Actual')
plt.title('Confusion Matrix')
plt.show()
```

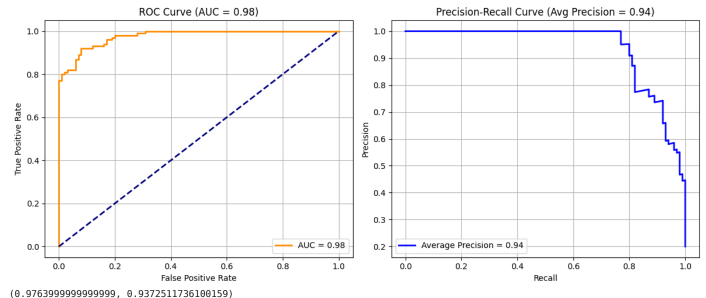
 \implies i) Maximum achievable sensitivity while maintaining FPR $\le 5\%$: 0.820 ii) Precision at this sensitivity: 0.965



Q6) To account for the fact that the class imbalance in the deployment environment (1:4) is very different from the class imbalance among your 200 training samples (1:1), you decide to add 3 additional copies of each HEALTHY guest to your dataset leading to 500 samples in total. Without 'retraining' your classifier, report the ROC and P-R curves for this new test set, along with ROCAUC and average precision. Briefly discuss what changed, what didn't, and why. (75 words)

Q6 Answer:

```
healthy data augmented = pd.concat([healthy data] * 4, ignore index=True)
sick_data_augmented = sick_data.copy()
healthy_data_augmented['Patient_Status'] = 'HEALTHY'
sick data augmented['Patient Status'] = 'SICK'
augmented_data = pd.concat([healthy_data_augmented, sick_data_augmented], ignore_index=True)
X_augmented = augmented_data[['T', 'RR']].values
y_augmented_true = (augmented_data['Patient_Status'] == 'SICK').astype(int).values
posterior probabilities augmented = []
for x_point in X_augmented:
    likelihood sick point = multivariate normal.pdf(x point, mean=mean sick vector, cov=cov sick matrix)
    likelihood_healthy_point = multivariate_normal.pdf(x_point, mean=mean_healthy_vector, cov=cov_healthy_matrix)
    evidence_point = (likelihood_sick_point * P_sick) + (likelihood_healthy_point * P_healthy)
    posterior_sick_point = (likelihood_sick_point * P_sick) / evidence_point
    posterior_probabilities_augmented.append(posterior_sick_point)
posterior_probabilities_augmented = np.array(posterior_probabilities_augmented)
fpr_augmented, tpr_augmented, _ = roc_curve(y_augmented_true, posterior_probabilities_augmented)
roc_auc_augmented = auc(fpr_augmented, tpr_augmented)
precision_augmented, recall_augmented, _ = precision_recall_curve(y_augmented_true, posterior_probabilities_augmented)
avg precision augmented = average precision score(y augmented true, posterior probabilities augmented)
plt.figure(figsize=(12, 5))
plt.subplot(1, 2, 1)
plt.plot(fpr_augmented, tpr_augmented, color='darkorange', lw=2, label=f'AUC = {roc_auc_augmented:.2f}')
plt.plot([0, 1], [0, 1], color='navy', lw=2, linestyle='--')
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title(f'ROC Curve (AUC = {roc_auc_augmented:.2f})')
plt.legend(loc='lower right')
plt.grid(True)
plt.subplot(1, 2, 2)
plt.plot(recall_augmented, precision_augmented, color='blue', lw=2, label=f'Average Precision = {avg_precision_augmented:.2f}')
plt.xlabel('Recall')
plt.ylabel('Precision')
plt.title(f'Precision-Recall Curve (Avg Precision = {avg_precision_augmented:.2f})')
plt.legend(loc='lower left')
plt.grid(True)
plt.tight_layout()
plt.show()
roc_auc_augmented, avg_precision_augmented
```



- The ROC AUC stayed relatively high (close to 0.98) because it measures the ability of the classifier to separate the two classes without being influenced by the class proportions. However, average precision is directly impacted by the change in class balance.
- The average precision actually decreased from 0.98 to 0.94. This drop in precision occurred because the dataset now contains a higher
 proportion of HEALTHY guests, making it more challenging for the classifier to precisely identify the SICK cases without mistakenly
 labeling more HEALTHY guests as sick. The classifier's performance in distinguishing between classes was impacted by the change in
 class distribution, hence the reduction in average precision.

lacksquare

Q7) <time passes...> Due to hard-working scientists and engineers, we now have a partially effective vaccine for MF. We now estimate that only one guest out of 100 will have MF this season. Without calculating anything, what do you expect the P-R curve to look like and why? (40 words)

Q7 Answer:

In an imbalanced scenario where only 1% of guests are SICK, the **precision** tends to decrease because there are significantly more opportunities for false positives (healthy guests being classified incorrectly as sick). This lowers the ratio of true positives to false positives, which directly impacts precision.

On the other hand, **recall** is defined as the ratio of true positives to all actual positives, and it generally remains more stable because it's not influenced by the increased number of healthy guests.

Thus, I expect the P-R curve to show a lower precision for the same recall levels compared to the original dataset. The rarity of the positive class makes it more challenging for the model to maintain a high precision.



Q8) Discuss how, in this assignment, we have committed both methodological errors of i) "Testing on the training set" and ii) "Training on the testing set". (~100 words)

Q8 Answer:

• i) Testing on the Training Set:

We used the same data for both training and testing our Bayesian classifier. This means that the classifier was evaluated on data it had already seen during "training," resulting in overly optimistic performance metrics that may not generalize well to new data.

• ii) Training on the Testing Set:

When augmenting the dataset to match the deployment environment (adding copies of healthy guests), we modified the testing set without retraining the classifier. This causes inconsistencies, as the classifier wasn't trained on the updated class distribution, leading to biased evaluations that don't reflect real-world performance.