Q1. Write a Python program to perform a Z-test for comparing a sample mean to a known population mean and interpret the results.

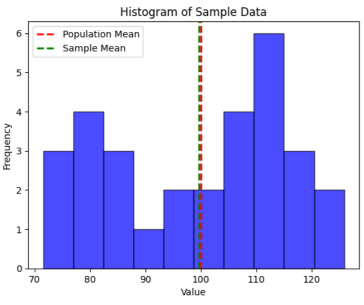
```
Ans:-
```

```
import numpy as np
import scipy.stats as stats
# Function to perform a one-sample Z-test
def z_test(sample, population_mean, population_std, alpha=0.05):
   # Calculate sample mean and sample size
    sample_mean = np.mean(sample)
   n = len(sample)
   # Calculate the Z-score
   z_score = (sample_mean - population_mean) / (population_std / np.sqrt(n))
   # Calculate the p-value (two-tailed)
   p_value = 2 * (1 - stats.norm.cdf(abs(z_score)))
   # Print results
   print(f"Sample Mean: {sample_mean:.2f}")
   print(f"Population Mean: {population_mean:.2f}")
   print(f"Z-Score: {z_score:.2f}")
    print(f"P-Value: {p_value:.4f}")
   # Interpret the results
    if p_value < alpha:</pre>
       print("Reject the null hypothesis: There is a significant difference between the sample mean and the population mean.")
    else:
       print("Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.")
# Example usage
if __name__ == "
                 _main
    # Sample data
   sample_data = [20, 22, 19, 24, 21, 23, 20, 22, 19, 21]
   # Known population parameters
   population mean = 21 # Known population mean
   population_std = 2  # Known population standard deviation
   # Perform the Z-test
    z_test(sample_data, population_mean, population_std)
    Sample Mean: 21.10
     Population Mean: 21.00
     Z-Score: 0.16
     P-Value: 0.8744
     Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.
Q2. Simulate random data to perform hypothesis testing and calculate the corresponding P-value using Python?
Ans:-
import numpy as np
import scipy.stats as stats
import matplotlib.pyplot as plt
# Function to perform a one-sample Z-test
def z_test(sample, population_mean, population_std, alpha=0.05):
   # Calculate sample mean and sample size
   sample_mean = np.mean(sample)
   n = len(sample)
   # Calculate the Z-score
   z_score = (sample_mean - population_mean) / (population_std / np.sqrt(n))
   # Calculate the p-value (two-tailed)
   p_value = 2 * (1 - stats.norm.cdf(abs(z_score)))
   # Print results
   print(f"Sample Mean: {sample_mean:.2f}")
   print(f"Population Mean: {population_mean:.2f}")
   print(f"Z-Score: {z_score:.2f}")
   print(f"P-Value: {p_value:.4f}")
    # Interpret the results
   if p_value < alpha:</pre>
       print("Reject the null hypothesis: There is a significant difference between the sample mean and the population mean.")
       print("Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.")
```

```
def simulate_data(sample_size, population_mean, population_std):
    # Generate random sample data from a normal distribution
    sample\_data = np.random.normal(loc=population\_mean, scale=population\_std, size=sample\_size)
    return sample_data
# Example usage
if __name__ == "__main__":
   # Parameters for simulation
    sample_size = 30  # Size of the sample
   population_mean = 100 # Known population mean
   population_std = 15  # Known population standard deviation
   # Simulate random data
    sample_data = simulate_data(sample_size, population_mean, population_std)
   # Perform the Z-test
   z_test(sample_data, population_mean, population_std)
   # Optional: Visualize the sample data
   plt.hist(sample_data, bins=10, alpha=0.7, color='blue', edgecolor='black')
   plt.axvline(population_mean, color='red', linestyle='dashed', linewidth=2, label='Population Mean')
   plt.axvline(np.mean(sample_data), color='green', linestyle='dashed', linewidth=2, label='Sample Mean')
   plt.title('Histogram of Sample Data')
    plt.xlabel('Value')
    plt.ylabel('Frequency')
   plt.legend()
    plt.show()
→ Sample Mean: 99.60
     Population Mean: 100.00
     Z-Score: -0.14
```

P-Value: 0.8849

Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.



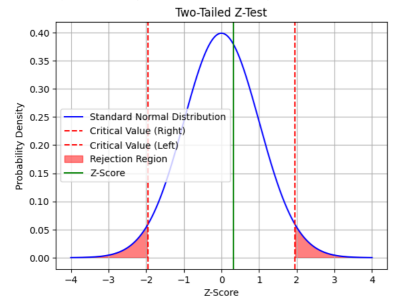
Q3.Implement a one-sample Z-test using Python to compare the sample mean with the population mean?

```
import numpy as np
import scipy.stats as stats
# Function to perform a one-sample Z-test
def one_sample_z_test(sample, population_mean, population_std, alpha=0.05):
   # Calculate sample mean and sample size
   sample_mean = np.mean(sample)
   n = len(sample)
   # Calculate the Z-score
   z_score = (sample_mean - population_mean) / (population_std / np.sqrt(n))
   # Calculate the p-value (two-tailed)
   p_value = 2 * (1 - stats.norm.cdf(abs(z_score)))
   # Print results
   print(f"Sample Mean: {sample_mean:.2f}")
   print(f"Population Mean: {population_mean:.2f}")
   print(f"Z-Score: {z_score:.2f}")
   print(f"P-Value: {p_value:.4f}")
    # Interpret the results
    if n value < alnha:
```

```
print("Reject the null hypothesis: There is a significant difference between the sample mean and the population mean.")
    else:
        print("Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.")
# Example usage
if __name__ == "__main__":
    # Sample data (you can modify this data)
    sample_data = [102, 98, 101, 105, 99, 100, 97, 104, 103, 101]
    # Known population parameters
    population_mean = 100 # Known population mean
    population_std = 10  # Known population standard deviation
    # Perform the one-sample Z-test
    one_sample_z_test(sample_data, population_mean, population_std)
→ Sample Mean: 101.00
     Population Mean: 100.00
     Z-Score: 0.32
     P-Value: 0.7518
     Fail to reject the null hypothesis: There is no significant difference between the sample mean and the population mean.
Q4.Perform a two-tailed Z-test using Python and visualize the decision region on a plot?
Ans:-
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import norm
# Function to perform a two-tailed Z-test
def two_tailed_z_test(sample, population_mean, population_std, alpha=0.05):
    # Calculate sample mean and sample size
    sample_mean = np.mean(sample)
    n = len(sample)
    # Calculate the Z-score
    z_score = (sample_mean - population_mean) / (population_std / np.sqrt(n))
    # Calculate critical values
    critical_val1 = norm.ppf(1 - alpha / 2)
    critical_val2 = norm.ppf(alpha / 2)
    # Print results
    print(f"Sample Mean: {sample_mean:.2f}")
   print(f"Population Mean: {population_mean:.2f}")
    print(f"Z-Score: {z_score:.2f}")
    # Decision based on Z-score
    if abs(z_score) > critical_val1:
       print("Reject the null hypothesis: There is a significant difference.")
    else:
        print("Fail to reject the null hypothesis: There is no significant difference.")
   # Visualization
    x = np.linspace(-4, 4, 1000)
    y = norm.pdf(x, 0, 1)
    plt.plot(x, y, label='Standard Normal Distribution', color='blue')
   plt.axvline(x=critical_val1, color='red', linestyle='--', label='Critical Value (Right)')
plt.axvline(x=critical_val2, color='red', linestyle='--', label='Critical Value (Left)')
    plt.fill\_between(x, y, where=(x <= critical\_val2) \mid (x >= critical\_val1), color='red', alpha=0.5, label='Rejection Region')
    plt.axvline(x=z_score, color='green', linestyle='-', label='Z-Score')
    plt.title('Two-Tailed Z-Test')
    plt.xlabel('Z-Score')
    plt.ylabel('Probability Density')
    plt.legend()
    plt.grid()
    plt.show()
# Example usage
if __name__ == "__main__":
    # Sample data (you can modify this data)
    sample_data = [102, 98, 101, 105, 99, 100, 97, 104, 103, 101]
    # Known population parameters
    population_mean = 100  # Known population mean
    population_std = 10  # Known population standard deviation
    # Perform the two-tailed Z-test
    two_tailed_z_test(sample_data, population_mean, population_std)
```

Sample Mean: 101.00 Population Mean: 100.00 Z-Score: 0.32

Fail to reject the null hypothesis: There is no significant difference.



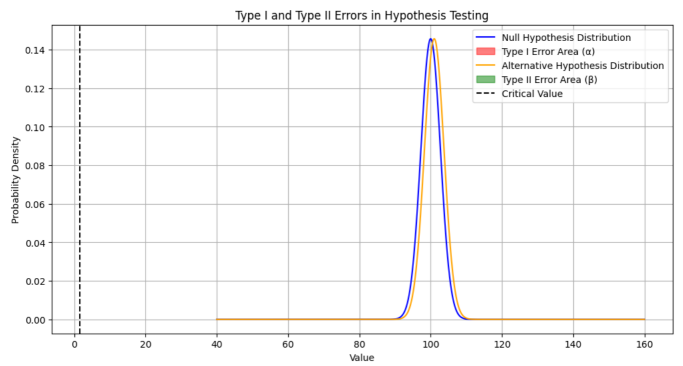
Q5.Create a Python function that calculates and visualizes Type 1 and Type 2 errors during hypothesis testing?

```
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import norm
def visualize_errors(population_mean, population_std, sample_size, alpha=0.05):
   # Define the null and alternative hypotheses
   null_hypothesis_mean = population_mean
   alternative_hypothesis_mean = population_mean + 1  # Shift the mean for the alternative hypothesis
   # Calculate critical value for Type I error
   critical_value = norm.ppf(1 - alpha)
   # Generate x values for the normal distributions
   x = np.linspace(population_mean - 4 * population_std, population_mean + 4 * population_std, 1000)
   # Calculate the probability density functions for the null and alternative hypotheses
   null_pdf = norm.pdf(x, null_hypothesis_mean, population_std / np.sqrt(sample_size))
   alternative\_pdf = norm.pdf(x, alternative\_hypothesis\_mean, population\_std / np.sqrt(sample\_size))
   # Calculate Type I error area (alpha)
   type_1_error_area = norm.cdf(critical_value)
   # Calculate Type II error area (beta)
   beta = norm.cdf(critical_value, loc=alternative_hypothesis_mean, scale=population_std / np.sqrt(sample_size))
   # Plotting
   plt.figure(figsize=(12, 6))
   # Plot null hypothesis distribution
   plt.plot(x, null_pdf, label='Null Hypothesis Distribution', color='blue')
   plt.fill_between(x, null_pdf, where=(x <= critical_value), color='red', alpha=0.5, label='Type I Error Area (α)')
   # Plot alternative hypothesis distribution
   plt.plot(x, alternative_pdf, label='Alternative Hypothesis Distribution', color='orange')
   plt.fill\_between(x, alternative\_pdf, where=(x <= critical\_value), color='green', alpha=0.5, label='Type II Error Area (\beta)')
   # Add critical value line
   plt.axvline(x=critical_value, color='black', linestyle='--', label='Critical Value')
   # Add labels and legend
   plt.title('Type I and Type II Errors in Hypothesis Testing')
   plt.xlabel('Value')
   plt.ylabel('Probability Density')
   plt.legend()
   plt.grid()
   plt.show()
   \ensuremath{\text{\#}} Print Type I and Type II error probabilities
   print(f"Type I Error (α): {type_1_error_area:.4f}")
   print(f"Type II Error (β): {beta:.4f}")
```

```
# Example usage
if __name__ == "__main__":
    population_mean = 100  # Known population mean
    population_std = 15  # Known population standard deviation
    sample_size = 30  # Sample size
    alpha = 0.05  # Significance level

    visualize_errors(population_mean, population_std, sample_size, alpha)
```





Type I Error ( $\alpha$ ): 0.9500 Type II Error ( $\beta$ ): 0.0000

Q6. Write a Python program to perform an independent T-test and interpret the results?

```
Ans:-
```

```
import numpy as np
import scipy.stats as stats
# Function to perform an independent two-sample T-test
def independent_t_test(group1, group2, alpha=0.05):
    # Calculate means and standard deviations
   mean1 = np.mean(group1)
   mean2 = np.mean(group2)
    std1 = np.std(group1, ddof=1) # Sample standard deviation
    std2 = np.std(group2, ddof=1) # Sample standard deviation
   n1 = len(group1)
   n2 = len(group2)
   # Calculate the T-statistic and p-value
    t_statistic, p_value = stats.ttest_ind(group1, group2)
    # Print results
   print(f"Group 1 Mean: {mean1:.2f}, Standard Deviation: {std1:.2f}, Sample Size: {n1}")
   print(f"Group 2 Mean: {mean2:.2f}, Standard Deviation: {std2:.2f}, Sample Size: {n2}")
   print(f"T-Statistic: {t_statistic:.2f}")
   print(f"P-Value: {p_value:.4f}")
    # Interpret the results
    if p_value < alpha:</pre>
       print("Reject the null hypothesis: There is a significant difference between the two group means.")
    else:
       print("Fail to reject the null hypothesis: There is no significant difference between the two group means.")
# Example usage
if __name__ == "
                 _main__":
    # Sample data for two independent groups
    group1 = [23, 21, 18, 25, 30, 22, 19, 24, 20, 26]
   group2 = [30, 32, 29, 35, 31, 28, 33, 34, 30, 29]
    # Perform the independent T-test
    independent_t_test(group1, group2)
→ Group 1 Mean: 22.80, Standard Deviation: 3.61, Sample Size: 10
     Group 2 Mean: 31.10, Standard Deviation: 2.33, Sample Size: 10
     T-Statistic: -6.10
```

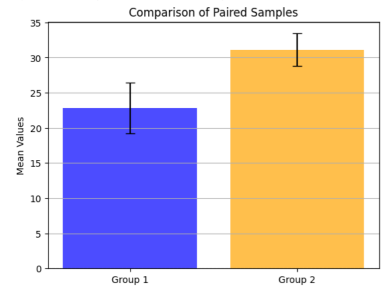
P-Value: 0.0000 Reject the null hypothesis: There is a significant difference between the two group means.

Q7.Perform a paired sample T-test using Python and visualize the comparison results?

Ans:-

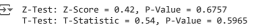
```
import numpy as np
import scipy.stats as stats
import matplotlib.pyplot as plt
# Function to perform a paired sample T-test
def paired_t_test(group1, group2, alpha=0.05):
    # Calculate the T-statistic and p-value
   t_statistic, p_value = stats.ttest_rel(group1, group2)
   # Print results
   print(f"T-Statistic: {t_statistic:.2f}")
   print(f"P-Value: {p_value:.4f}")
   # Interpret the results
    if p_value < alpha:
       print("Reject the null hypothesis: There is a significant difference between the two paired samples.")
    else:
       print("Fail to reject the null hypothesis: There is no significant difference between the two paired samples.")
    return t_statistic, p_value
# Function to visualize the comparison results
def visualize_comparison(group1, group2):
   # Calculate means and standard deviations
    means = [np.mean(group1), np.mean(group2)]
   stds = [np.std(group1, ddof=1), np.std(group2, ddof=1)]
   labels = ['Group 1', 'Group 2']
   # Create a bar plot with error bars
    x = np.arange(len(labels))
   plt.bar(x, means, yerr=stds, capsize=5, color=['blue', 'orange'], alpha=0.7)
   plt.xticks(x, labels)
   plt.ylabel('Mean Values')
   plt.title('Comparison of Paired Samples')
   plt.axhline(0, color='black', linewidth=0.8, linestyle='--')
   plt.grid(axis='y')
   plt.show()
# Example usage
if __name__ == "__main__":
    # Sample data for two paired groups
   group1 = [23, 21, 18, 25, 30, 22, 19, 24, 20, 26]
   group2 = [30, 32, 29, 35, 31, 28, 33, 34, 30, 29]
   # Perform the paired sample T-test
   t_statistic, p_value = paired_t_test(group1, group2)
   # Visualize the comparison results
    visualize_comparison(group1, group2)
    T-Statistic: -6.56
```

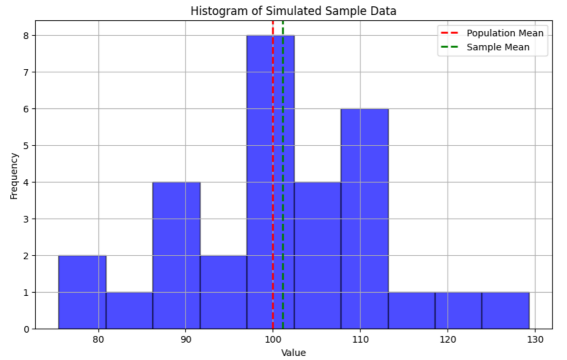
P-Value: 0.0001
Reject the null hypothesis: There is a significant difference between the two paired samples.



Q8. Simulate data and perform both Z-test and T-test, then compare the results using Python?

```
import numpy as np
import scipy.stats as stats
import matplotlib.pyplot as plt
# Function to perform a one-sample Z-test
def z_test(sample, population_mean, population_std, alpha=0.05):
    sample_mean = np.mean(sample)
    n = len(sample)
    z_score = (sample_mean - population_mean) / (population_std / np.sqrt(n))
    p_value = 2 * (1 - stats.norm.cdf(abs(z_score)))
    return z_score, p_value
# Function to perform a one-sample T-test
def t_test(sample, population_mean, alpha=0.05):
    t_statistic, p_value = stats.ttest_1samp(sample, population_mean)
    return t_statistic, p_value
# Function to simulate data
def simulate_data(sample_size, population_mean, population_std):
    return np.random.normal(loc=population_mean, scale=population_std, size=sample_size)
# Main function to run the tests and compare results
def main():
    # Parameters for simulation
    sample\_size = 30
   population_mean = 100
   population_std = 15
    # Simulate data
    sample_data = simulate_data(sample_size, population_mean, population_std)
    # Perform Z-test
    z_score, z_p_value = z_test(sample_data, population_mean, population_std)
    # Perform T-test
    t_statistic, t_p_value = t_test(sample_data, population_mean)
   # Print results
    print(f"Z-Test: Z-Score = {z_score:.2f}, P-Value = {z_p_value:.4f}")
   print(f"T-Test: T-Statistic = {t_statistic:.2f}, P-Value = {t_p_value:.4f}")
    # Visualization
    plt.figure(figsize=(10, 6))
    plt.hist(sample_data, bins=10, alpha=0.7, color='blue', edgecolor='black')
   plt.axvline(population_mean, color='red', linestyle='dashed', linewidth=2, label='Population Mean')
    plt.axvline(np.mean(sample_data), color='green', linestyle='dashed', linewidth=2, label='Sample Mean')
    plt.title('Histogram of Simulated Sample Data')
    plt.xlabel('Value')
    plt.ylabel('Frequency')
    plt.legend()
    plt.grid()
    plt.show()
# Run the main function
if __name__ == "__main__":
    main()
```





Q9. Write a Python function to calculate the confidence interval for a sample mean and explain its significance.

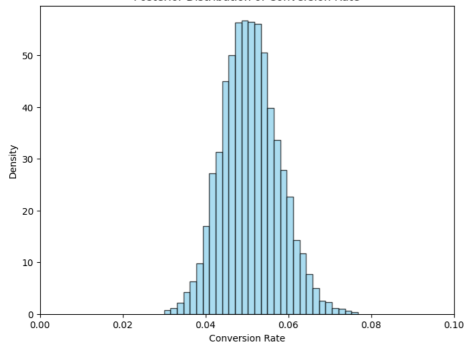
Ans:-

```
import numpy as np
import scipy.stats as stats
def confidence_interval(sample, confidence_level=0.95):
   Calculate the confidence interval for the sample mean.
   Parameters:
   sample (array-like): The sample data.
    confidence_level (float): The confidence level (default is 0.95 for 95% confidence).
   Returns:
    tuple: The lower and upper bounds of the confidence interval.
   # Calculate sample mean and standard error
    sample_mean = np.mean(sample)
    sample_std = np.std(sample, ddof=1) # Sample standard deviation
   n = len(sample) # Sample size
   standard_error = sample_std / np.sqrt(n)
   # Calculate the critical value for the t-distribution
   critical_value = stats.t.ppf((1 + confidence_level) / 2, df=n-1)
   # Calculate the margin of error
   margin_of_error = critical_value * standard_error
   # Calculate the confidence interval
   lower_bound = sample_mean - margin_of_error
   upper_bound = sample_mean + margin_of_error
   return lower_bound, upper_bound
# Example usage
if __name__ == "__main__":
   # Sample data
    sample_data = [23, 21, 18, 25, 30, 22, 19, 24, 20, 26]
   # Calculate the confidence interval
    ci = confidence_interval(sample_data, confidence_level=0.95)
   print(f"95\% \ Confidence \ Interval \ for \ the \ Sample \ Mean: \ \{ci[0]:.2f\} \ to \ \{ci[1]:.2f\}")
⇒ 95% Confidence Interval for the Sample Mean: 20.21 to 25.39
```

Q10.Write a Python program to calculate the margin of error for a given confidence level using sample data.

```
import numpy as np
import scipy.stats as stats
def margin_of_error(sample, confidence_level=0.95):
    Calculate the margin of error for a given confidence level.
    Parameters:
    sample (array-like): The sample data.
    confidence_level (float): The confidence level (default is 0.95 for 95% confidence).
    float: The margin of error.
    # Calculate sample standard deviation and sample size
    sample_std = np.std(sample, ddof=1) # Sample standard deviation
    n = len(sample) # Sample size
    # Calculate the standard error
    standard_error = sample_std / np.sqrt(n)
    # Calculate the critical value for the t-distribution
    critical_value = stats.t.ppf((1 + confidence_level) / 2, df=n-1)
    # Calculate the margin of error
    margin_of_error = critical_value * standard_error
    return margin_of_error
# Example usage
if __name__ == "
                 _main__":
    # Sample data
    sample_data = [23, 21, 18, 25, 30, 22, 19, 24, 20, 26]
   # Calculate the margin of error
    me = margin of error(sample data, confidence level=0.95)
   print(f"Margin of Error at 95% Confidence Level: {me:.2f}")
→ Margin of Error at 95% Confidence Level: 2.59
Q11.plement a Bayesian inference method using Bayes' Theorem in Python and explain the process.
Ans:-
import numpy as np
import matplotlib.pyplot as plt
from scipy.stats import beta
# Observed data
num_visitors = 1000 # Total number of visitors to the website
num_conversions = 50 # Number of conversions (desired actions)
# Prior hyperparameters for the Beta distribution
prior_alpha = 1 # Shape parameter
prior_beta = 1
                # Shape parameter
# Update the prior with the observed data to get the posterior parameters
posterior_alpha = prior_alpha + num_conversions
posterior_beta = prior_beta + (num_visitors - num_conversions)
# Generate samples from the posterior Beta distribution
posterior_samples = np.random.beta(posterior_alpha, posterior_beta, size=10000)
# Plot the posterior distribution
plt.figure(figsize=(8, 6))
plt.hist(posterior_samples, bins=30, density=True, color='skyblue', edgecolor='black', alpha=0.7)
plt.title('Posterior Distribution of Conversion Rate')
plt.xlabel('Conversion Rate')
plt.ylabel('Density')
plt.xlim(0, 0.1) # Limiting x-axis to focus on conversion rates close to zero
plt.show()
# Calculate summary statistics
mean_conversion_rate = posterior_alpha / (posterior_alpha + posterior_beta)
mode\_conversion\_rate = (posterior\_alpha - 1) \ / \ (posterior\_alpha + posterior\_beta - 2) \ \# \ Mode \ of \ the \ Beta \ distribution
print("Mean conversion rate:", mean_conversion_rate)
print("Mode conversion rate:", mode_conversion_rate)
```

## Posterior Distribution of Conversion Rate



Mean conversion rate: 0.05089820359281437

Mode conversion rate: 0.05

P-value: 0.010343173196618245 Degrees of Freedom: 2 Expected Frequencies: [[25. 25.] [17.5 17.5]

Q12.Perform a Chi-square test for independence between two categorical variables in Python?

```
Ans:-
```

```
import numpy as np
import pandas as pd
import scipy.stats as stats
# Sample data: A contingency table
data = {
    'Category A': [30, 10, 20],
    'Category B': [20, 25, 15]
# Create a DataFrame
df = pd.DataFrame(data, index=['Group 1', 'Group 2', 'Group 3'])
# Display the contingency table
print("Contingency Table:")
print(df)
# Perform the Chi-square test
chi2_statistic, p_value, dof, expected = stats.chi2_contingency(df)
# Print the results
print("\nChi-square Statistic:", chi2_statistic)
print("P-value:", p_value)
print("Degrees of Freedom:", dof)
print("Expected Frequencies:")
print(expected)
# Interpret the results
alpha = 0.05
if p_value < alpha:</pre>
   print("\nReject the null hypothesis: There is a significant association between the two categorical variables.")
    print("\nFail to reject the null hypothesis: There is no significant association between the two categorical variables.")
→ Contingency Table:
              Category A
                         Category B
                      30
                                  20
     Group 1
                      10
                                  25
     Group 2
     Group 3
                      20
                                  15
     Chi-square Statistic: 9.142857142857144
```

Reject the null hypothesis: There is a significant association between the two categorical variables.

Q13.Write a Python program to calculate the expected frequencies for a Chi-square test based on observed data?

```
Ans:-
```

```
import numpy as np
import pandas as pd
def calculate_expected_frequencies(observed):
   Calculate the expected frequencies for a Chi-square test based on observed data.
   Parameters:
   observed (array-like): The observed data in a contingency table.
   Returns:
   np.ndarray: The expected frequencies.
   # Convert observed data to a DataFrame if it's not already
   if not isinstance(observed, pd.DataFrame):
       observed = pd.DataFrame(observed)
   # Calculate row totals, column totals, and grand total
   column_totals = observed.sum(axis=0).values # Column totals
   grand_total = observed.values.sum() # Grand total
   # Calculate expected frequencies
   expected = (row_totals * column_totals) / grand_total
   return expected
# Example usage
if __name__ == "__main__":
   # Sample observed data (contingency table)
   observed_data = np.array([[30, 10, 20],
                             [20, 25, 15]])
   # Create a DataFrame for better visualization
   observed_df = pd.DataFrame(observed_data, columns=['Category A', 'Category B', 'Category C'], index=['Group 1', 'Group 2'])
   print("Observed Frequencies:")
   print(observed_df)
   # Calculate expected frequencies
   expected_frequencies = calculate_expected_frequencies(observed_data)
   # Display expected frequencies
   expected_df = pd.DataFrame(expected_frequencies, columns=['Category A', 'Category B', 'Category C'], index=['Group 1', 'Group 2'])
   print("\nExpected Frequencies:")
   print(expected_df)
→ Observed Frequencies:
             Category A Category B Category C
    Group 1
                    30
                               10
                                           20
    Group 2
                    20
                                25
                                           15
     Expected Frequencies:
             Category A Category B Category C
     Group 1
                  25.0
                            17.5
    Group 2
                  25.0
                              17.5
```

Q14.Perform a goodness-of-fit test using Python to compare the observed data to an expected distribution.

```
import numpy as np
import pandas as pd
import scipy.stats as stats
import matplotlib.pyplot as plt

def chi_square_goodness_of_fit(observed, expected):
    """
    Perform a Chi-square goodness-of-fit test.

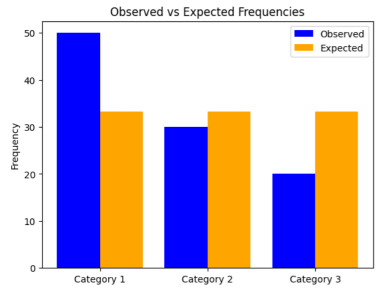
    Parameters:
    observed (array-like): The observed frequencies.
    expected (array-like): The expected frequencies.

    Returns:
    chi2_statistic (float): The Chi-square statistic.
    p_value (float): The p-value of the test.
    """

# Calculate the Chi-square statistic and p-value
```

```
chi2_statistic, p_value = stats.chisquare(observed, expected)
    return chi2_statistic, p_value
# Example usage
if __name__ == "_
                  main ":
    # Sample observed data (e.g., counts of outcomes)
    observed_data = np.array([50, 30, 20]) # Observed frequencies for three categories
    # Expected frequencies for a uniform distribution
    total_observed = observed_data.sum()
    num_categories = len(observed_data)
    expected_data = np.full(num_categories, total_observed / num_categories) # Equal expected frequencies
    # Perform the Chi-square goodness-of-fit test
    chi2_statistic, p_value = chi_square_goodness_of_fit(observed_data, expected_data)
    # Print the results
   print("Observed Frequencies:", observed_data)
print("Expected Frequencies:", expected_data)
print("Chi-square Statistic:", chi2_statistic)
    print("P-value:", p_value)
    # Interpret the results
    alpha = 0.05
    if p_value < alpha:</pre>
        print("\nReject the null hypothesis: The observed data does not fit the expected distribution.")
    else:
        print("\nFail to reject the null hypothesis: The observed data fits the expected distribution.")
    # Optional: Visualize the observed vs expected frequencies
    categories = ['Category 1', 'Category 2', 'Category 3']
    x = np.arange(len(categories))
    plt.bar(x - 0.2, observed\_data, width=0.4, label='Observed', color='blue')
    plt.bar(x + 0.2, expected_data, width=0.4, label='Expected', color='orange')
    plt.xticks(x, categories)
    plt.ylabel('Frequency')
    plt.title('Observed vs Expected Frequencies')
    plt.legend()
    plt.show()
Observed Frequencies: [50 30 20]
     Expected Frequencies: [33.3333333 33.3333333 33.33333333]
     Chi-square Statistic: 14.0
     P-value: 0.0009118819655545164
```

Reject the null hypothesis: The observed data does not fit the expected distribution.

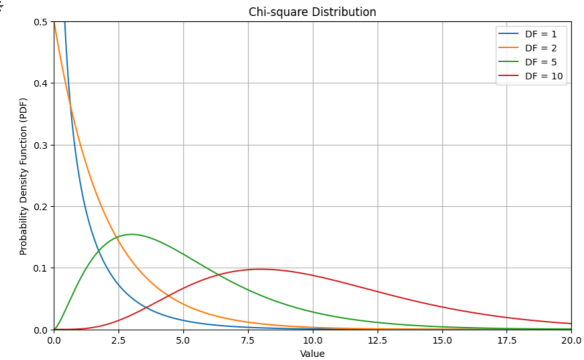


Q15.Create a Python script to simulate and visualize the Chi-square distribution and discuss its characteristics.

```
import numpy as np
import matplotlib.pyplot as plt
import scipy.stats as stats

def plot_chi_square_distribution(degrees_of_freedom, x_range):
    """
    Plot the Chi-square distribution for given degrees of freedom.
    Parameters:
```

```
degrees_of_freedom (list): List of degrees of freedom to plot.
    x_range (tuple): Range of x values for the plot.
    x = np.linspace(x_range[0], x_range[1], 1000)
    plt.figure(figsize=(10, 6))
    for df in degrees_of_freedom:
        # Calculate the Chi-square PDF
        pdf = stats.chi2.pdf(x, df)
        plt.plot(x, pdf, label=f'DF = {df}')
    plt.title('Chi-square Distribution')
    plt.xlabel('Value')
    plt.ylabel('Probability Density Function (PDF)')
   plt.legend()
    plt.grid()
    plt.xlim(x_range)
    plt.ylim(0, 0.5)
    plt.show()
# Example usage
if __name__ == "__main__":
   degrees_of_freedom = [1, 2, 5, 10] # Different degrees of freedom to visualize
    x_range = (0, 20) # Range of x values for the plot
    plot_chi_square_distribution(degrees_of_freedom, x_range)
\overline{\mathbf{x}}
                                                    Chi-square Distribution
```



Q16.Implement an F-test using Python to compare the variances of two random samples.

```
import numpy as np
import scipy.stats as stats

def f_test(sample1, sample2, alpha=0.05):
    """
    Perform an F-test to compare the variances of two samples.

    Parameters:
    sample1 (array-like): The first sample data.
    sample2 (array-like): The second sample data.
    alpha (float): The significance level (default is 0.05).

    Returns:
    f_statistic (float): The F-statistic.
    p_value (float): The p-value of the test.
    """

# Calculate the variances of the samples
    var1 = np.var(sample1, ddof=1) # Sample variance
    var2 = np.var(sample2, ddof=1) # Sample variance

# Calculate the F-statistic
    f_statistic = var1 / var2
```

```
# Calculate the degrees of freedom
    df1 = len(sample1) - 1 # Degrees of freedom for sample1
    df2 = len(sample2) - 1  # Degrees of freedom for sample2
    # Calculate the p-value
   p_value = 1 - stats.f.cdf(f_statistic, df1, df2)
    return f_statistic, p_value
# Example usage
if __name__ == "
                 __main__":
    # Sample data
    sample1 = np.array([20, 22, 19, 24, 21, 23, 20, 22, 19, 21])
    sample2 = np.array([30, 32, 29, 35, 31, 28, 33, 34, 30, 29])
   # Perform the F-test
   f_statistic, p_value = f_test(sample1, sample2)
    # Print the results
    print("F-Statistic:", f_statistic)
    print("P-Value:", p_value)
    # Interpret the results
    alpha = 0.05
    if p_value < alpha:</pre>
        print("\nReject the null hypothesis: The variances of the two samples are significantly different.")
        print("\nFail to reject the null hypothesis: The variances of the two samples are not significantly different.")
F-Statistic: 0.5092024539877301
     P-Value: 0.8354277292129488
     Fail to reject the null hypothesis: The variances of the two samples are not significantly different.
Q17. Write a Python program to perform an ANOVA test to compare means between multiple groups and interpret the results.
Ans:-
import numpy as np
import pandas as pd
import scipy.stats as stats
import matplotlib.pyplot as plt
def perform_anova(*groups):
    Perform a one-way ANOVA test to compare means between multiple groups.
   Parameters:
    *groups: Variable number of group samples (arrays).
    f statistic (float): The F-statistic.
    p_value (float): The p-value of the test.
   # Perform the ANOVA test
    f_statistic, p_value = stats.f_oneway(*groups)
    return f_statistic, p_value
def plot_groups(groups):
   Plot the boxplot of the groups for visual comparison.
    Parameters:
    groups: List of group samples (arrays).
   plt.figure(figsize=(10, 6))
    plt.boxplot(groups, labels=[f'Group {i+1}' for i in range(len(groups))])
    plt.title('Boxplot of Groups')
    plt.ylabel('Values')
   plt.grid()
   plt.show()
# Example usage
if __name__ == "__main__":
    # Sample data for three groups
    group1 = np.array([23, 21, 18, 25, 30])
    group2 = np.array([30, 32, 29, 35, 31])
   group3 = np.array([22, 24, 20, 23, 21])
    # Perform the ANOVA test
    f_statistic, p_value = perform_anova(group1, group2, group3)
    # Print the results
```

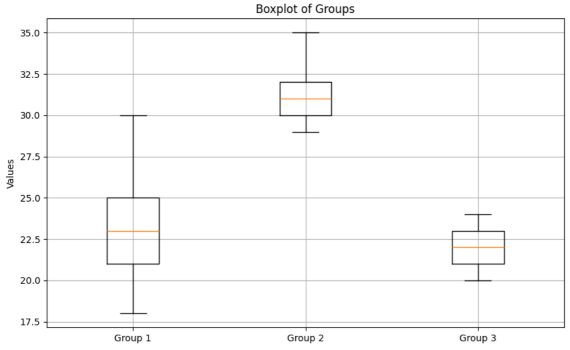
```
print("F-Statistic:", f_statistic)
print("P-Value:", p_value)

# Interpret the results
alpha = 0.05
if p_value < alpha:
    print("\nReject the null hypothesis: At least one group mean is significantly different.")
else:
    print("\nFail to reject the null hypothesis: There is no significant difference between group means.")

# Plot the groups for visual comparison
plot_groups([group1, group2, group3])</pre>
# F-Statistic: 13 729537366548039
```

F-Statistic: 13.729537366548039 P-Value: 0.0007910539293303842

Reject the null hypothesis: At least one group mean is significantly different.
<ipython-input-17-343a11efb66b>:29: MatplotlibDeprecationWarning: The 'labels' parameter of boxplot() has been renamed 'tick\_labels' since Matplot plt.boxplot(groups, labels=[f'Group {i+1}' for i in range(len(groups))])



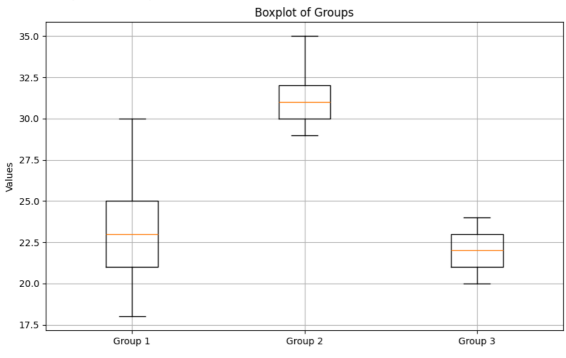
Q18.Perform a one-way ANOVA test using Python to compare the means of different groups and plot the results.

```
import numpy as np
import pandas as pd
import scipy.stats as stats
import matplotlib.pyplot as plt
def perform_anova(*groups):
   Perform a one-way ANOVA test to compare means between multiple groups.
   Parameters:
    *groups: Variable number of group samples (arrays).
   Returns:
    f_statistic (float): The F-statistic.
   p\_value (float): The p-value of the test.
   # Perform the ANOVA test
    f_statistic, p_value = stats.f_oneway(*groups)
   return f_statistic, p_value
def plot_groups(groups, group_labels):
   Plot the boxplot of the groups for visual comparison.
   Parameters:
    groups: List of group samples (arrays).
   group_labels: List of labels for each group.
   plt.figure(figsize=(10, 6))
   plt.boxplot(groups, labels=group_labels)
    plt.title('Boxplot of Groups')
   plt.ylabel('Values')
```

```
plt.grid()
    plt.show()
# Example usage
if __name__ == "__main__":
   # Sample data for three groups
    group1 = np.array([23, 21, 18, 25, 30])
   group2 = np.array([30, 32, 29, 35, 31])
   group3 = np.array([22, 24, 20, 23, 21])
   # Perform the ANOVA test
   f_statistic, p_value = perform_anova(group1, group2, group3)
   # Print the results
   print("F-Statistic:", f_statistic)
   print("P-Value:", p_value)
    # Interpret the results
    alpha = 0.05
    if p_value < alpha:</pre>
       print("\nReject the null hypothesis: At least one group mean is significantly different.")
    else:
       print("\nFail to reject the null hypothesis: There is no significant difference between group means.")
    # Plot the groups for visual comparison
    plot_groups([group1, group2, group3], ['Group 1', 'Group 2', 'Group 3'])
    F-Statistic: 13.729537366548039
```

P-Value: 0.0007910539293303842

Reject the null hypothesis: At least one group mean is significantly different. <ipython-input-18-584463ee2571>:30: MatplotlibDeprecationWarning: The 'labels' parameter of boxplot() has been renamed 'tick\_labels' since Matplot plt.boxplot(groups, labels=group\_labels)



Q19. Write a Python function to check the assumptions (normality, independence, and equal variance) for ANOVA.

```
import numpy as np
import pandas as pd
import scipy.stats as stats
import matplotlib.pyplot as plt
import seaborn as sns
{\tt def\ check\_anova\_assumptions(*groups):}
    Check the assumptions for ANOVA: normality, independence, and equal variance.
    Parameters:
    *groups: Variable number of group samples (arrays).
    # Check Normality using Shapiro-Wilk test
    print("Normality Test Results:")
    for i, group in enumerate(groups):
        stat, p_value = stats.shapiro(group)
```

```
print(f"Group {i+1}: W-statistic = {stat:.4f}, p-value = {p_value:.4f}")
        if p_value < 0.05:
           print(f"Group \ \{i+1\} \ does \ not \ follow \ a \ normal \ distribution \ (reject \ H0).\ \ ")
           print(f"Group {i+1} follows a normal distribution (fail to reject H0).\n")
   # Check Equal Variance using Levene's test
    stat, p_value = stats.levene(*groups)
   print("Equal Variance Test (Levene's Test):")
   print(f"Levene's W-statistic = {stat:.4f}, p-value = {p_value:.4f}")
   if p_value < 0.05:
       print("The variances are significantly different (reject H0).\n")
    else:
       print("The variances are equal (fail to reject H0).\n")
   # Visualize the data distribution
   plt.figure(figsize=(12, 6))
   for i, group in enumerate(groups):
        sns.kdeplot(group, label=f'Group {i+1}', fill=True)
   plt.title('Kernel Density Estimation of Groups')
   plt.xlabel('Values')
   plt.ylabel('Density')
   plt.legend()
   plt.grid()
   plt.show()
   # Boxplot for visual inspection of equal variance
   plt.figure(figsize=(10, 6))
   plt.boxplot(groups, labels=[f'Group {i+1}' for i in range(len(groups))])
   plt.title('Boxplot of Groups')
   plt.ylabel('Values')
   plt.grid()
   plt.show()
# Example usage
if __name__ == "__main__":
   # Sample data for three groups
    group1 = np.random.normal(loc=20, scale=5, size=30)
   group2 = np.random.normal(loc=22, scale=5, size=30)
    group3 = np.random.normal(loc=21, scale=5, size=30)
   # Check ANOVA assumptions
    check_anova_assumptions(group1, group2, group3)
```

Normality Test Results:

Group 1: W-statistic = 0.9838, p-value = 0.9157

Group 1 follows a normal distribution (fail to reject H0).

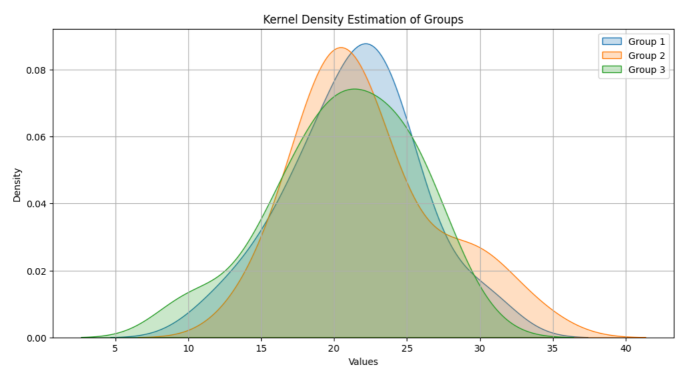
Group 2: W-statistic = 0.9456, p-value = 0.1287

Group 2 follows a normal distribution (fail to reject H0).

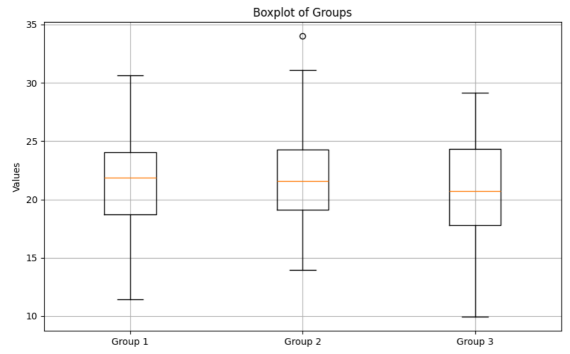
Group 3: W-statistic = 0.9666, p-value = 0.4510

Group 3 follows a normal distribution (fail to reject H0).

Equal Variance Test (Levene's Test): Levene's W-statistic = 0.1427, p-value = 0.8672 The variances are equal (fail to reject H0).



<ipython-input-19-4a75d65e6e31>:46: MatplotlibDeprecationWarning: The 'labels' parameter of boxplot() has been renamed 'tick\_labels' since Matplot
plt.boxplot(groups, labels=[f'Group {i+1}' for i in range(len(groups))])



Q20.Perform a two-way ANOVA test using Python to study the interaction between two factors and visualize the results.

Ans:-Step 1: Import Required Libraries

python

import pandas as pd import numpy as np

import statsmodels.api as sm import statsmodels.formula.api as smf

```
import seaborn as sns
import matplotlib.pyplot as plt
```

### Step 2: Create or Load Your Data

For demonstration, let's create a sample dataset. In practice, you would load your data from a CSV or other source.

```
# Create a sample dataset
np.random.seed(42)
data = {
    'Factor1': np.repeat(['A', 'B', 'C'], 20),
    'Factor2': np.tile(np.repeat(['X', 'Y'], 10), 3),
    'Response': np.random.normal(loc=0, scale=1, size=60)
df = pd.DataFrame(data)
Step 3: Perform Two-Way ANOVA
# Fit the model
model = smf.ols('Response ~ C(Factor1) * C(Factor2)', data=df).fit()
# Perform ANOVA
anova_table = sm.stats.anova_lm(model, typ=2)
print(anova_table)
                                        df
                                                        PR(>F)
                              sum sa
     C(Factor1)
                            0.580882
                                       2.0 0.398223 0.673468
                            1.275222 1.0 1.748451 0.191645
     C(Factor2)
                                      2.0 5.113208 0.009255
     C(Factor1):C(Factor2)
                            7,458570
                           39.384553 54.0
     Residual
                                                 NaN
                                                           NaN
```

## Step 4: Check for Significant Interactions

The ANOVA table will show you the p-values for the main effects and their interaction. If the p-value for the interaction term is less than 0.05, it indicates a significant interaction between the two factors.

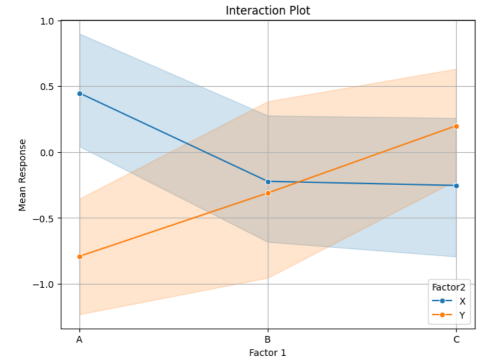
### Step 5: Visualize the Results

You can visualize the interaction using an interaction plot.

```
import pandas as pd
import numpy as np
import statsmodels.api as sm
import statsmodels.formula.api as smf
import seaborn as sns
import matplotlib.pyplot as plt
# Create a sample dataset
np.random.seed(42)
data = {
    'Factor1': np.repeat(['A', 'B', 'C'], 20),
    'Factor2': np.tile(np.repeat(['X', 'Y'], 10), 3),
    'Response': np.random.normal(loc=0, scale=1, size=60)
df = pd.DataFrame(data)
# Fit the model
model = smf.ols('Response ~ C(Factor1) * C(Factor2)', data=df).fit()
# Perform ANOVA
anova_table = sm.stats.anova_lm(model, typ=2)
print(anova_table)
# Create interaction plot using lineplot
plt.figure(figsize=(8, 6))
sns.lineplot(data=df, x='Factor1', y='Response', hue='Factor2', estimator='mean', marker='o')
plt.title('Interaction Plot')
plt.xlabel('Factor 1')
plt.ylabel('Mean Response')
plt.grid()
plt.show()
```



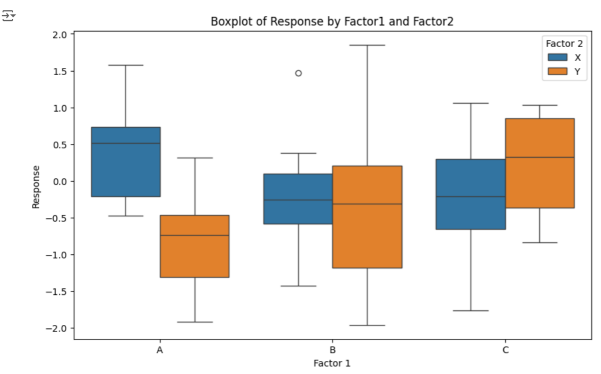
C(Factor1):C(Factor2) 7.458570 2.0 5.113208 0.009255 Residual 39.384553 54.0 NaN NaN



Step 6: Boxplot for Group Comparison

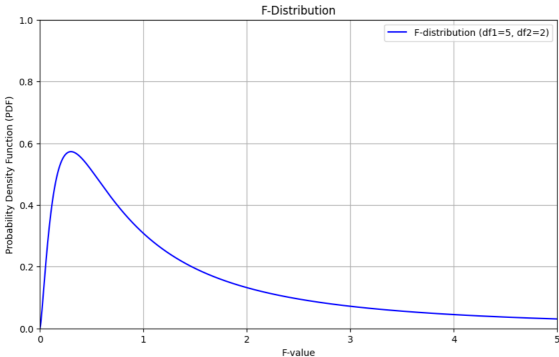
Additionally, you can create boxplots to visualize the distribution of the response variable across the different groups.

```
# Boxplot
plt.figure(figsize=(10, 6))
sns.boxplot(x='Factor1', y='Response', hue='Factor2', data=df)
plt.title('Boxplot of Response by Factor1 and Factor2')
plt.xlabel('Factor 1')
plt.ylabel('Response')
plt.legend(title='Factor 2')
plt.show()
```



Q21.Write a Python program to visualize the F-distribution and discuss its use in hypothesis testing.

```
import numpy as np
import matplotlib.pyplot as plt
import scipy.stats as stats
def plot_f_distribution(degrees_of_freedom1, degrees_of_freedom2):
    Plot the F-distribution for given degrees of freedom.
    degrees_of_freedom1 (int): Degrees of freedom for the numerator.
    {\tt degrees\_of\_freedom2\ (int):\ Degrees\ of\ freedom\ for\ the\ denominator.}
    x = np.linspace(0, 5, 1000) # Range of x values
    pdf = stats.f.pdf(x, degrees_of_freedom1, degrees_of_freedom2) # F-distribution PDF
    plt.figure(figsize=(10, 6))
    plt.plot(x, pdf, label=f'F-distribution (df1={degrees_of_freedom1}, df2={degrees_of_freedom2})', color='blue')
    plt.title('F-Distribution')
    plt.xlabel('F-value')
   plt.ylabel('Probability Density Function (PDF)')
    plt.grid()
    plt.legend()
   plt.xlim(0, 5)
    plt.ylim(0, 1)
    plt.show()
# Example usage
if __name__ == "__main__":
    # Define degrees of freedom
    df1 = 5  # Degrees of freedom for the numerator
   df2 = 2 # Degrees of freedom for the denominator
    # Plot the F-distribution
    plot_f_distribution(df1, df2)
\overline{2}
```



Q22.Perform a one-way ANOVA test in Python and visualize the results with boxplots to compare group means.

```
import numpy as np
import pandas as pd
import scipy.stats as stats
import seaborn as sns
import matplotlib.pyplot as plt

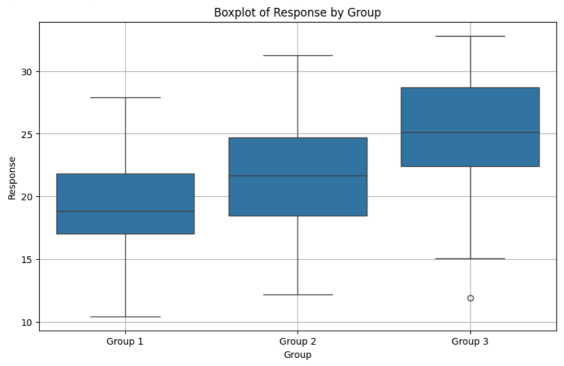
# Create a sample dataset
np.random.seed(42)
group1 = np.random.normal(loc=20, scale=5, size=30) # Group 1
group2 = np.random.normal(loc=22, scale=5, size=30) # Group 2
group3 = np.random.normal(loc=25, scale=5, size=30) # Group 3

# Combine the groups into a DataFrame
data = {
    'Group': ['Group 1'] * 30 + ['Group 2'] * 30 + ['Group 3'] * 30,
```

```
'Response': np.concatenate([group1, group2, group3])
}
df = pd.DataFrame(data)
# Perform the one-way ANOVA test
f_statistic, p_value = stats.f_oneway(df[df['Group'] == 'Group 1']['Response'],
                                       df[df['Group'] == 'Group 2']['Response'],
                                       df[df['Group'] == 'Group 3']['Response'])
# Print the results
print("F-Statistic:", f_statistic)
print("P-Value:", p_value)
# Interpret the results
alpha = 0.05
if p_value < alpha:</pre>
   print("\nReject the null hypothesis: At least one group mean is significantly different.")
else:
    print("\nFail to reject the null hypothesis: There is no significant difference between group means.")
\# Visualize the results with boxplots
plt.figure(figsize=(10, 6))
sns.boxplot(x='Group', y='Response', data=df)
plt.title('Boxplot of Response by Group')
plt.xlabel('Group')
plt.ylabel('Response')
plt.grid()
plt.show()
    F-Statistic: 12.397871901825155
```

P-Value: 1.8305780660638664e-05

Reject the null hypothesis: At least one group mean is significantly different.



Q23. Simulate random data from a normal distribution, then perform hypothesis testing to evaluate the means.

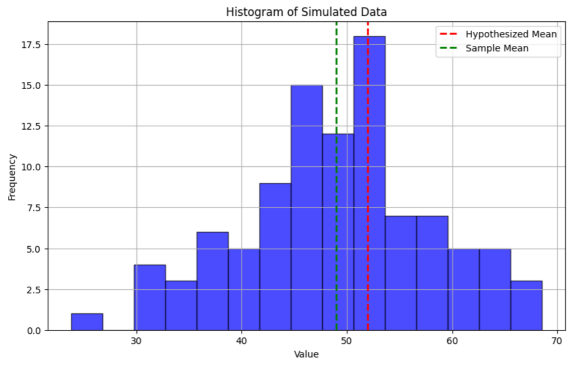
Ans:-Python Code for Simulating Data and Performing Hypothesis Testing

```
import numpy as np
import scipy.stats as stats
import matplotlib.pyplot as plt
# Set parameters for the normal distribution
np.random.seed(42) # For reproducibility
mean = 50
                     # True mean of the population
std dev = 10
                     # Standard deviation of the population
sample_size = 100
                     # Number of samples to simulate
\hbox{\tt\# Simulate random data from a normal distribution}\\
data = np.random.normal(loc=mean, scale=std_dev, size=sample_size)
# Perform a one-sample t-test
population_mean = 52  # Hypothesized population mean
t_statistic, p_value = stats.ttest_1samp(data, population_mean)
# Print the results
```

```
print("Sample Mean:", np.mean(data))
print("T-Statistic:", t_statistic)
print("P-Value:", p_value)
# Interpret the results
alpha = 0.05
if p_value < alpha:</pre>
    print("\nReject the null hypothesis: The sample mean is significantly different from the population mean.")
else:
    print("\nFail to reject the null hypothesis: The sample mean is not significantly different from the population mean.")
# Visualize the data
plt.figure(figsize=(10, 6))
plt.hist(data, bins=15, alpha=0.7, color='blue', edgecolor='black')
plt.axvline(population_mean, color='red', linestyle='dashed', linewidth=2, label='Hypothesized Mean')
plt.axvline(np.mean(data), color='green', linestyle='dashed', linewidth=2, label='Sample Mean')
plt.title('Histogram of Simulated Data')
plt.xlabel('Value')
plt.ylabel('Frequency')
plt.legend()
plt.grid()
plt.show()
```

Sample Mean: 48.96153482605907 T-Statistic: -3.3457066775666733 P-Value: 0.0011606436334217142

Reject the null hypothesis: The sample mean is significantly different from the population mean.



Q24.Perform a hypothesis test for population variance using a Chi-square distribution and interpret the results.

Ans:-

import numpy as np

```
import scipy.stats as stats

# Sample data
np.random.seed(42)  # For reproducibility
data = np.random.normal(loc=50, scale=10, size=30)  # Simulated data

# Hypothesized population variance
sigma_squared_0 = 100  # Hypothesized variance (10^2)

# Calculate sample variance and sample size
n = len(data)
sample_variance = np.var(data, ddof=1)  # Sample variance with Bessel's correction

# Calculate the Chi-square statistic
chi_square_statistic = (n - 1) * sample_variance / sigma_squared_0

# Degrees of freedom
df = n - 1

# Significance level
alpha = 0.05
```

```
# Critical values for two-tailed test
critical_value_lower = stats.chi2.ppf(alpha / 2, df)
critical_value_upper = stats.chi2.ppf(1 - alpha / 2, df)
# Print results
print("Sample Variance:", sample_variance)
print("Chi-square Statistic:", chi_square_statistic)
print("Critical Value Lower:", critical_value_lower)
print("Critical Value Upper:", critical_value_upper)
# Interpret the results
if \ chi\_square\_statistic \ < \ critical\_value\_lower \ or \ chi\_square\_statistic \ > \ critical\_value\_upper:
   print("\nReject the null hypothesis: The population variance is significantly different from the hypothesized variance.")
    print("\nFail to reject the null hypothesis: The population variance is not significantly different from the hypothesized variance.")
→ Sample Variance: 81.00115693656518
     Chi-square Statistic: 23.490335511603902
     Critical Value Lower: 16.04707169536489
     Critical Value Upper: 45.72228580417452
     Fail to reject the null hypothesis: The population variance is not significantly different from the hypothesized variance.
Q25.Write a Python script to perform a Z-test for comparing proportions between two datasets or groups.
Ans:-
import numpy as np
import scipy.stats as stats
def z_test_proportions(success_a, n_a, success_b, n_b):
    Perform a Z-test for comparing proportions between two groups.
    Parameters:
    success a (int): Number of successes in group A.
    n_a (int): Total number of observations in group A.
    success_b (int): Number of successes in group B.
   n_b (int): Total number of observations in group B.
    Returns:
    z_statistic (float): The Z-statistic.
    p_value (float): The p-value of the test.
    # Calculate proportions
    p_a = success_a / n_a
    p_b = success_b / n_b
    # Calculate the pooled proportion
    p_pool = (success_a + success_b) / (n_a + n_b)
    # Calculate the standard error
    standard\_error = np.sqrt(p\_pool * (1 - p\_pool) * (1/n_a + 1/n_b))
    # Calculate the Z-statistic
    z_statistic = (p_a - p_b) / standard_error
    # Calculate the p-value (two-tailed)
    p value = 2 * (1 - stats.norm.cdf(abs(z statistic)))
    return z_statistic, p_value
# Example usage
if __name__ == "_
                 main
    # Data for group A
    success_a = 30 # Number of successes in group A
   n_a = 100
                  # Total observations in group A
    # Data for group B
    success_b = 20 # Number of successes in group B
                   # Total observations in group B
    n b = 80
    # Perform the Z-test for proportions
    z_statistic, p_value = z_test_proportions(success_a, n_a, success_b, n_b)
    # Print the results
    print("Z-Statistic:", z_statistic)
   print("P-Value:", p_value)
    # Interpret the results
    alpha = 0.05
    if p_value < alpha:
        print("\nReject the null hypothesis: There is a significant difference between the proportions.")
```

```
print("\nFail to reject the null hypothesis: There is no significant difference between the proportions.")
₹ Z-Statistic: 0.7442084075352506
     P-Value: 0.45675040242316944
     Fail to reject the null hypothesis: There is no significant difference between the proportions.
Q26.Implement an F-test for comparing the variances of two datasets, then interpret and visualize the results.
Ans:-
import numpy as np
import pandas as pd
import scipy.stats as stats
import matplotlib.pyplot as plt
import seaborn as sns
def f_test(sample1, sample2):
   Perform an F-test to compare the variances of two samples.
   Parameters:
    sample1 (array-like): The first sample data.
   sample2 (array-like): The second sample data.
    Returns:
   f_statistic (float): The F-statistic.
   p_value (float): The p-value of the test.
   # Calculate the variances of the samples
   var1 = np.var(sample1, ddof=1) # Sample variance
   var2 = np.var(sample2, ddof=1) # Sample variance
   # Calculate the F-statistic
   f_statistic = var1 / var2
   # Calculate the degrees of freedom
   df1 = len(sample1) - 1  # Degrees of freedom for sample1
   df2 = len(sample2) - 1 # Degrees of freedom for sample2
   # Calculate the p-value
   p_value = 1 - stats.f.cdf(f_statistic, df1, df2)
   return f_statistic, p_value
# Example usage
if __name__ == "__main__":
   # Generate sample data
   np.random.seed(42) # For reproducibility
   sample1 = np.random.normal(loc=50, scale=10, size=30) # Sample 1
   sample2 = np.random.normal(loc=55, scale=15, size=30) # Sample 2
   # Perform the F-test
    f_statistic, p_value = f_test(sample1, sample2)
   # Print the results
   print("F-Statistic:", f_statistic)
   print("P-Value:", p_value)
   # Interpret the results
   alpha = 0.05
    if p_value < alpha:</pre>
       print("\nReject the null hypothesis: The variances of the two samples are significantly different.")
       print("\nFail to reject the null hypothesis: The variances of the two samples are not significantly different.")
   # Visualize the results with boxplots
   plt.figure(figsize=(10, 6))
    sns.boxplot(data=[sample1, sample2], palette="Set2")
   plt.title('Boxplot of Two Samples')
```

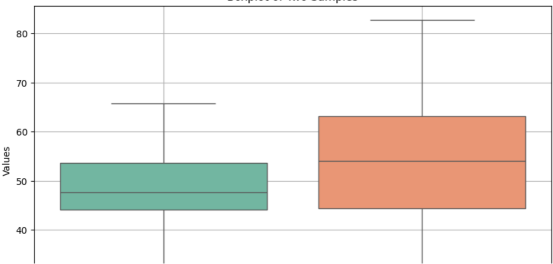
plt.xlabel('Sample')
plt.ylabel('Values')

plt.grid()
plt.show()

plt.xticks([0, 1], ['Sample 1', 'Sample 2'])

Fail to reject the null hypothesis: The variances of the two samples are not significantly different.

# **Boxplot of Two Samples**



Q27.Perform a Chi-square test for goodness of fit with simulated data and analyze the results.

Ans:-Python Code for Chi-Square Goodness of Fit Test

```
import numpy as np
import pandas as pd
import scipy.stats as stats
{\tt import\ matplotlib.pyplot\ as\ plt}
{\tt def\ chi\_square\_goodness\_of\_fit(observed,\ expected):}
   Perform a Chi-square goodness of fit test.
    observed (array-like): Observed frequencies.
    expected (array-like): Expected frequencies.
    Returns:
    chi2_statistic (float): The Chi-square statistic.
   p_value (float): The p-value of the test.
    chi2_statistic, p_value = stats.chisquare(f_obs=observed, f_exp=expected)
    return chi2_statistic, p_value
# Simulate categorical data
np.random.seed(42) # For reproducibility
categories = ['A', 'B', 'C', 'D']
```