1 Exercise 1

```
1 def align(seq1, seq2, match=1, gap_penalty=1, mismatch_penalty=1):
      m, n = len(seq1), len(seq2)
      # Initialize DP matrix and traceback indicators
3
      dp = [[0] * (n + 1) for _ in range(m + 1)]
      traceback = [[None] * (n + 1) for _ in range(m + 1)]
      max\_score = 0
      max_pos = None
      # DP matrix
10
      for i in range(1, m + 1):
11
          for j in range(1, n + 1):
13
               if seq1[i - 1] == seq2[j - 1]:
                   match\_score = dp[i - 1][j - 1] + match
14
15
                   match_score = dp[i - 1][j - 1] - mismatch_penalty
16
               gap_seq1 = dp[i - 1][j] - gap_penalty
gap_seq2 = dp[i][j - 1] - gap_penalty
18
19
20
               dp[i][j] = max(0, match_score, gap_seq1, gap_seq2)
21
22
               # Store traceback direction
24
               if dp[i][j] == match_score:
                   traceback[i][j] = "DIAG"
25
               elif dp[i][j] == gap_seq1:
                   traceback[i][j] = "UP"
27
               elif dp[i][j] == gap_seq2:
28
                   traceback[i][j] = "LEFT"
30
               # Update max score position
31
               if dp[i][j] > max_score:
                   max_score = dp[i][j]
33
                   max_pos = (i, j)
34
35
36
      # Traceback to count the optimal alignment
37
      aligned_seq1 = []
      aligned_seq2 = []
38
39
      i, j = max_pos
      while dp[i][j] != 0:
41
           if traceback[i][j] == "DIAG":
43
               aligned_seq1.append(seq1[i - 1])
               aligned_seq2.append(seq2[j - 1])
44
               i -= 1
               j -= 1
46
           elif traceback[i][j] == "UP":
               aligned_seq1.append(seq1[i - 1])
               aligned_seq2.append("-")
49
               i -= 1
           elif traceback[i][j] == "LEFT":
51
               aligned_seq1.append("-")
52
               aligned_seq2.append(seq2[j - 1])
               j -= 1
54
      # Reverse the sequences (since we traceback from the end)
      aligned_seq1 = ','.join(reversed(aligned_seq1))
57
      aligned_seq2 = ''.join(reversed(aligned_seq2))
58
59
      return aligned_seq1, aligned_seq2, max_score
```

Listing 1: Dynamic Programming Smith-Waterman

Answers and Explanations

1. Correctness of the Implementation

The implementation of the Smith-Waterman algorithm correctly computes the local alignment between two sequences, taking into account matches, mismatches, and gap penalties. The function uses a dynamic programming (DP) approach to compute the optimal local alignment and score. It fills the DP matrix using the recurrence relation:

```
dp[i][j] = max(0, dp[i-1][j-1] + match/mismatch_score, dp[i-1][j] - gap_penalty, dp[i][j-1] - gap_penalty),
```

where: - Matches add the match score. - Mismatches subtract the mismatch_penalty. - Gaps subtract the gap_penalty.

Traceback begins from the cell with the maximum score and continues until reaching a score of 0, ensuring correct local alignment reconstruction.

2. Tests and Results

The function was tested using the following cases:

• Default Parameters:

- Input: align('tgcatcgagaccctacgtgac', 'actagacctagcatcgac')
- Output:

```
seq1 = agacccta-cgt-gac, seq2 = aga-cctagcatcgac, score = 8
```

This matches the expected result, demonstrating correct alignment.

• Higher Gap Penalty:

- Input: align('tgcatcgagaccctacgtgac', 'actagacctagcatcgac', gap_penalty=2)
- Output:

```
seq1 = gcatcga, seq2 = gcatcga, score = 7
```

The increased gap penalty results in a shorter alignment, as expected.

```
seq1, seq2, score = align('tgcatcgagaccctacgtgac', 'actagacctagcatcgac')
print(seq1, seq2, score)

/ 0.0s

agacccta-cgt-gac aga-cctagcatcgac 8

/ seq1, seq2, score = align('tgcatcgagaccctacgtgac', 'actagacctagcatcgac', gap_penalty=2)
print(seq1, seq2, score)

/ 0.0s

gcatcga gcatcga 7
```

Figure 1: Test Result

3. Explanation of Testing

The test cases demonstrate that the function handles default and customized scoring schemes correctly:

- With default parameters, the function identifies the optimal alignment and calculates the correct score.
- By increasing the gap penalty, the alignment length decreases, and the score adjusts accordingly, reflecting the expected behavior.

The outputs align with theoretical expectations for the Smith-Waterman algorithm and validate the function's correctness.

2 Exercise 2

This QuadTree implementation 1 organizes 2D points into a hierarchical tree structure to enable efficient k-nearest neighbor searches. The tree divides a rectangular region (defined by x_bounds and y_bounds) into four quadrants if the number of points exceeds max_points. Points are recursively assigned to child nodes, following the hint to "split into children if enough points exist." Then, the _within_distance method ensures efficient searching by checking if a point lies within a distance d (Traditional Euclidean Distance) from the node's bounds, as suggested by the hint to use distance-based containment checks. Another section is the query method recursively retrieves k-nearest neighbors by exploring only relevant child nodes. It avoids unnecessary descent into regions that cannot contain neighbors. Finally, after collecting candidates, they are sorted by Euclidean distance (use squared distance here to avoid the overlapping point calculation error), and the closest k points are returned.

```
# QuadTree Implementation
  class QuadTree:
      def __init__(self, points, x_bounds, y_bounds, max_points=4):
           self.points = points
           self.x_bounds = x_bounds
           self.y_bounds = y_bounds
           self.children = []
           self.max_points = max_points
           self._split()
10
      def _split(self):
11
           if len(self.points) <= self.max_points:</pre>
13
               return
           # Calculate midpoints and define four quadrants for splitting each region
           mid_x = (self.x_bounds[0] + self.x_bounds[1]) / 2
15
          mid_y = (self.y_bounds[0] + self.y_bounds[1]) / 2
16
17
           quadrants = [
               ([], (self.x_bounds[0], mid_x), (self.y_bounds[0], mid_y)),
18
               ([], (mid_x, self.x_bounds[1]), (self.y_bounds[0], mid_y)),
19
               ([], (self.x_bounds[0], mid_x), (mid_y, self.y_bounds[1])),
20
               ([], (mid_x, self.x_bounds[1]), (mid_y, self.y_bounds[1])),
21
          ]
           # Distribute points into appropriate quadrants and recursively create child QuadTree
23
       nodes for non-empty quadrants
24
           for point in self.points:
               x, y, _ = point
25
26
               for quadrant in quadrants:
                   if quadrant[1][0] <= x <= quadrant[1][1] and quadrant[2][0] <= y <= quadrant</pre>
27
      [2][1]: ##check 'contains
28
                        quadrant [0].append(point)
                        break
29
           self.points = []
30
           for q_points, q_x_bounds, q_y_bounds in quadrants:
               if q_points:
32
                   self.children.append(QuadTree(q_points, q_x_bounds, q_y_bounds, self.
      max_points))
34
35
      def _within_distance(self, x, y, d):
           x_min, x_max = self.x_bounds
36
           y_min, y_max = self.y_bounds
37
           closest_x = min(max(x, x_min), x_max)
          closest_y = min(max(y, y_min), y_max)
return (closest_x - x) ** 2 + (closest_y - y) ** 2 <= d ** 2</pre>
39
40
41
      def query(self, x, y, k):
42
           candidates = []
43
44
           def search(node):
               if not node.children:
45
                   candidates.extend(node.points)
47
               else:
                   for child in node.children:
48
                        if child._within_distance(x, y, float('inf')):
                            search(child)
50
51
           search(self)
           candidates.sort(key=lambda p: (p[0] - x) ** 2 + (p[1] - y) ** 2)
           return candidates[:k]
53
```

Listing 2: QuadTree implementation

¹I wrote the body part of the QuadTree implementation following the Youtube Video https://www.youtube.com/watch?v=iG100qOyZIA and our slides. Some minor problems are debugged by CHATGPT (I fell into the Infinite Splitting and sometimes got stuck when implementing the 'check-contain' process)

Then we take advantage of QuadTree data structure to do Nearest Neighbor Searching. The $knn_predict$ function queries the QuadTree to find the k-nearest neighbors for each test point. It predicts the most common class among these neighbors using a majority 'vote'.

```
# Build the QuadTree
quadtree = QuadTree(train_data, (x_min, x_max), (y_min, y_max))

# KNN prediction function
def knn_predict(quadtree, test_points, k):
    predictions = []
for x, y, _ in test_points:
    neighbors = quadtree.query(x, y, k)
    classes = [neighbor[2] for neighbor in neighbors]
    predictions.append(Counter(classes).most_common(1)[0][0])
return predictions
```

Listing 3: KNN implementation

Before implementing KNN on our rice data, we do basic preprocessing. Normalize the seven quantitative columns to a mean of 0 and standard deviation 1. Reduce the data to two dimensions using PCA.

Listing 4: Data preprocessing

The scatterplot after PCA shows that the two rice types (Cammeo and Osmancik) are reasonably well-separated in the reduced 2D space, with clusters corresponding to each type. However, there is some overlap between the clusters. The overlapping indicates that some points near the boundary will likely be misclassified, especially when the true class distributions are close in the feature space. Do a train-test split with test size = 20%.

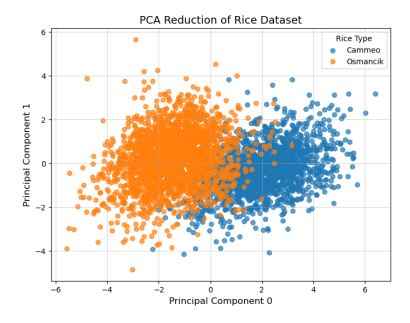


Figure 2: PCA visualization

```
5 # Build the QuadTree
 quadtree = QuadTree(train_data, (x_min, x_max), (y_min, y_max))
8 # Test KNN with k=1 and k=5
9 test_labels = test_data[:, 2]
predictions_k1 = knn_predict(quadtree, test_data, k=1)
predictions_k5 = knn_predict(quadtree, test_data, k=5)
^{13} # Confusion Matrices, this and the visualization below are assisted by CHATGPT
confusion_k1 = pd.crosstab(pd.Series(test_labels, name='Actual'), pd.Series(predictions_k1,
      name='Predicted'))
 confusion_k5 = pd.crosstab(pd.Series(test_labels, name='Actual'), pd.Series(predictions_k5,
     name='Predicted'))
16
plt.figure(figsize=(8, 6))
19 for rice_type, group in data.groupby('Class'):
      plt.scatter(group['PCO'], group['PC1'], label=rice_type, alpha=0.7)
20
22 plt.title('PCA Reduction of Rice Dataset', fontsize=14)
plt.xlabel('Principal Component 0', fontsize=12)
24 plt.ylabel('Principal Component 1', fontsize=12)
plt.legend(title='Rice Type')
plt.grid(True, alpha=0.5)
27 plt.show()
28 # Heatmaps for confusion matrices
29 plt.figure(figsize=(12, 5))
30 plt.subplot(1, 2, 1)
sns.heatmap(confusion_k1, annot=True, fmt='d', cmap='Blues', cbar=False)
plt.title('Confusion Matrix (k=1)', fontsize=14)
plt.xlabel('Predicted', fontsize=12)
plt.ylabel('Actual', fontsize=12)
35 plt.subplot(1, 2, 2)
sns.heatmap(confusion_k5, annot=True, fmt='d', cmap='Blues', cbar=False)
37 plt.title('Confusion Matrix (k=5)', fontsize=14)
plt.xlabel('Predicted', fontsize=12)
plt.ylabel('Actual', fontsize=12)
40 plt.tight_layout()
plt.show()
```

Listing 5: data preparation

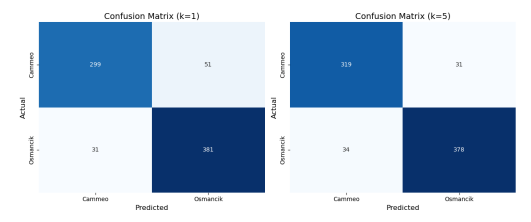


Figure 3: Confusion Matrix

As shown in figure 3, for k=1, the model correctly classifies 299 instances of Cammeo and 381 instances of Osmancik, but misclassifies 51 Cammeo instances as Osmancik and 31 Osmancik instances as Cammeo. This indicates good performance but highlights the sensitivity of k=1 to noise and boundary cases, leading to higher misclassification rates. For k=5, the model correctly classifies 319 Cammeo and 378 Osmancik instances, with 31 and 34 misclassifications, respectively. This shows that increasing k in this case improves robustness by reducing the influence of noise

3 Exercise 3

1 from mpi4py import MPI

```
2 import numpy as np
3 import matplotlib.pyplot as plt
4 import time
# Define bounds and parameters
7 \text{ xlo, ylo = -2.5, -1.5}
8 \text{ xhi, yhi} = 0.75, 1.5
9 nx, ny = 2048, 1536

10 dx, dy = (xhi - xlo) / nx, (yhi - ylo) / ny
iter_limit = 200
12 set_threshold = 2
14 def mandelbrot_test(x, y):
15
    z = 0
      c = x + y * 1j
16
      for i in range(iter_limit):
17
          z = z ** 2 + c
18
19
          if abs(z) > set_threshold:
               return i
20
      return i
22
23 def calculate_chunk(start, end):
      Calculates a chunk of the Mandelbrot set rows from 'start' to 'end'.
25
26
      local_result = np.zeros([end - start, nx])
27
28
      for i, row in enumerate(range(start, end)):
          y = row * dy + ylo
29
          for j in range(nx):
30
               x = j * dx + xlo
31
32
               local_result[i, j] = mandelbrot_test(x, y)
      return local_result
33
34
35 def calculate_serial():
36
      Serial implementation of Mandelbrot set calculation for comparison.
37
38
      result = np.zeros([ny, nx])
39
      for i in range(ny):
          y = i * dy + ylo
41
          for j in range(nx):
42
               x = j * dx + xlo
43
              result[i, j] = mandelbrot_test(x, y)
44
45
      return result
46
47 if __name__ == "__main__":
      # Initialize MPI
      comm = MPI.COMM_WORLD
49
      rank = comm.Get_rank()
50
51
      size = comm.Get_size()
52
      # Determine workload per process
53
      rows_per_process = ny // size
54
      start_row = rank * rows_per_process
55
      end_row = (rank + 1) * rows_per_process if rank != size - 1 else ny
57
      start_time = time.perf_counter()
58
      # Each process calculates its chunk
59
      local_result = calculate_chunk(start_row, end_row)
60
61
      # Gather all chunks to the root process
62
      if rank == 0:
63
          mandelbrot_set = np.zeros([ny, nx])
64
      else:
65
66
          mandelbrot set = None
67
      comm.Gather(local_result, mandelbrot_set, root=0)
68
      stop_time = time.perf_counter()
69
70
      # Perform correctness and performance comparison
71
72
      if rank == 0:
          print(f"Parallel calculation took {stop_time - start_time:.2f} seconds with {size}
73
      processes")
          # Serial calculation for comparison
75
76
           serial_start_time = time.perf_counter()
        serial_result = calculate_serial()
```

```
serial_stop_time = time.perf_counter()
78
          print(f"Serial calculation took {serial_stop_time - serial_start_time:.2f} seconds")
79
80
          # Check exact correctness, debugged by CHATGPT
          if np.array_equal(mandelbrot_set, serial_result):
82
83
              print("Parallel and serial results match")
84
              print("Mismatch between parallel and serial results.")
85
86
87
          # Plot the results
          plt.imshow(mandelbrot_seat, interpolation="nearest", cmap="Greys")
          plt.gca().set_aspect("equal")
          plt.axis("off")
90
          plt.show()
```

Listing 6: code implementation of exercise 3

The parallel implementation of the Mandelbrot set calculation was verified by comparing the results of the parallel and serial executions. For all test cases (n = 1, 2, 4), the parallel and serial outputs matched exactly, demonstrating the correctness of the parallel approach (cannot use '==' directly, otherwise will return a list of boolean values).

To run the code, the command will look like this

```
mpiexec -n 4 python /Users/yihang/Desktop/BIS_634/HW5/question3.py
```

The execution time of the parallel implementation was observed to improve as the number of processes increased. For n = 1, the parallel computation took approximately 6.76 seconds. When n = 2, the computation time decreased to 3.52 seconds, and for n = 4, it further reduced to 3.25 seconds. And the corresponding serial calculation took 7.10, 7.56, 7.42 seconds separately. These results highlight the performance benefits of distributing the workload across multiple processes.

The parallel implementation divides the rows of the Mandelbrot set among the available processes using MPI. Each process computes a subset of rows independently, and the results are gathered at the root process. This approach minimizes computation time by leveraging multiple processes and ensures correctness through careful synchronization using MPI.Gather.