Proseminar: High Performance Computing Assignment 02

1 Exercise 1

1.1 Write a sequential application pi_seq in C or C++ that computes π for a given number of samples (command line argument). Test your application for various, large sample sizes to verify the correctness of your implementation.

```
#include <stdio.h>
2 #include <stdlib.h>
3 #include <math.h>
  double calculate_pi(int num_points) {
      int points_in_circle = 0;
6
      for (int count = 0; count < num_points; count++) {</pre>
           double random_x = (double)rand() / RAND_MAX;
9
           double random_y = (double)rand() / RAND_MAX;
10
           if (random_x * random_x + random_y * random_y <= 1.0) {</pre>
12
13
               points_in_circle++;
14
      }
15
16
       return 4.0 * points_in_circle / num_points;
17
18 }
19
int main(int argc, char *argv[]) {
      int total_points = atoi(argv[1]);
21
22
       double pi_approx = calculate_pi(total_points);
23
24
       printf("Approximation of Pi: %f\n", pi_approx);
25
26
       return 0;
27
28 }
```

Listing 1: Sequential Pi Approximation

1.2 Consider a parallelization strategy using MPI. Which communication pattern(s) would you choose and why?

As there are no dependencies while generating random points and checking if they lie in the circle, we can do that individually per rank. So we can just split the problem size N into $N \div n_{ranks}$ chunks. At the end one core have to retrieve the number of points in circle from all other cores, build a sum out of it and calculate π with it.

For this we would use MPI_Reduce(...) as it can collect a variable from all ranks and directly use operations as addition by giving reduce the operation type MPI_SUM.

1.3 Implement your chosen parallelization strategy as a second application pi_mpi. Run it with varying numbers of ranks and sample sizes and verify its correctness by comparing the output to pi_seq.

```
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#include <mpi.h>
```

```
6 void calculate_pi(int num_points, int rank, int numProcs) {
       // WTime Initialization
       double starttime, endtime;
       starttime = MPI_Wtime();
9
10
       // Define how many points each rank generates and generate them
       int points_for_rank = round(num_points / numProcs);
       int points_in_circle_in_rank = 0;
12
       int points_in_circle_global = 0;
13
14
15
       for (int count = 0; count < points_for_rank; count++) {</pre>
           double random_x = (double)rand() / RAND_MAX;
16
           double random_y = (double)rand() / RAND_MAX;
17
18
           if (random_x * random_x + random_y * random_y <= 1.0) {</pre>
19
20
                points_in_circle_in_rank++;
21
       }
22
       // Gather results from all ranks
23
       MPI_Reduce(&points_in_circle_in_rank, &points_in_circle_global, 1, MPI_INT, MPI_SUM, 0,
24
       MPI_COMM_WORLD);
25
       if (rank == 0) {
26
           double pi_approx = 4.0 * points_in_circle_global / num_points;
27
           endtime = MPI_Wtime();
28
           \label{lem:printf("WTime: \%f seconds n", endtime - starttime);} printf("Approximation of Pi: \%f n", pi_approx);
29
30
31
32 }
33
int main(int argc, char *argv[]) {
       // MPI Initialization
35
36
       MPI_Init(&argc, &argv);
       int rank, numProcs;
37
       MPI_Comm_rank(MPI_COMM_WORLD, &rank);
38
       MPI_Comm_size(MPI_COMM_WORLD, &numProcs);
39
40
41
       int total_points = atoi(argv[1]);
42
43
       if (rank == 0) {
           printf("Total Points used for Approximation: %i \n", total_points);
44
45
46
47
       calculate_pi(total_points, rank, numProcs);
48
       MPI_Finalize();
49
50
       return 0;
51 }
```

Listing 2: Parallel Pi Approximation

1.4 Discuss the effects and implications of your parallelization.

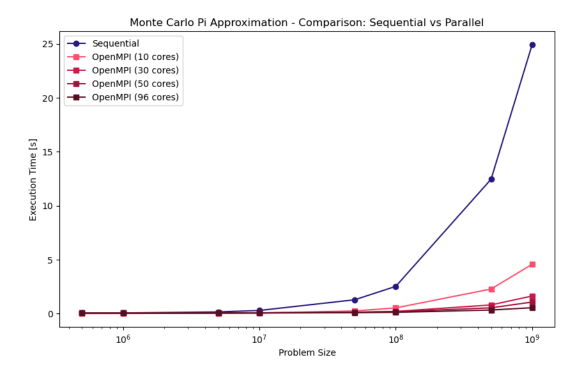


Figure 1: Monte Carlo Pi Approximation - Runtime Comparison

When running the experiment with a problem size below 10^7 the sequential code and the parallelized code have around the same runtimes. If more than 10^7 points are used the execution time of the the sequential program grows rapidly. By adding more cores the highest problem size (10^9) gets faster computed. The execution time for 10^9 points is 5x faster with 96 cores instead of just 10.

2 Exercise 2

2.1 A sequential implementation of a 1-D heat stencil is available in heat_stencil_1D_seq.c. Read the code and make sure you understand what happens. See the Wikipedia article on Stencil Codes for more information.

Iterative Stencil Loops

- 2.2 Consider a parallelization strategy using MPI. Which communication pattern(s) would you choose and why? Are there additional changes required in the code beyond calling MPI functions? If so, elaborate!
 - Boundary-Exchange-Pattern (using MPI_Sendrecv), because exchanging boundary values between processes is essential for correct calculations in each section.
 - Broadcast-Pattern (with MPI_Bcast), to efficiently distribute initial data to all processes.
 - Gather-Pattern (with MPI_Gather), to collect results from individual processes at the end of the computation.

Additional changes that need to be made include implementing functionality that splits the problem into chunks that can be distributed via MPI. This also includes additional error handling to ensure that the input fits our strategy.

2.3 Implement your chosen parallelization strategy as a second application heat_stencil_1D_mpi. Run it with varying numbers of ranks and problem sizes and verify its correctness by comparing the output to heat_stencil_1D_seq.

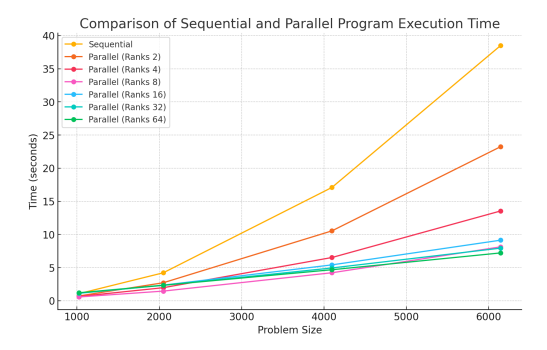
```
#include <stdio.h>
#include <stdlib.h>
3 #include <mpi.h>
5 typedef double value_t;
7 #define RESOLUTION 120
9 // -- vector utilities --
typedef value_t *Vector;
Vector createVector(int N);
void releaseVector(Vector m);
void printTemperature(Vector m, int N);
17 // -- simulation code ---
18
int main(int argc, char **argv) {
    // 'parsing' optional input parameter = problem size
20
    int N = 2000;
21
    if (argc > 1) {
     N = atoi(argv[1]);
23
24
    int T = N * 500;
25
26
    // MPI Initialization
27
    MPI_Init(&argc, &argv);
28
    int rank, numProcs;
29
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
30
    MPI_Comm_size(MPI_COMM_WORLD, &numProcs);
31
32
    // Time measurement variables
33
    double starttime, endtime;
34
35
    // Calculate chunk size and handle remainder
36
    int chunkSize = N / numProcs;
37
    int remainder = N % numProcs;
38
39
    // Determine start and end index for each rank
40
41
    int startIdx = rank * chunkSize + (rank < remainder ? rank : remainder);</pre>
    int endIdx = startIdx + chunkSize + (rank < remainder ? 1 : 0);</pre>
42
43
    int localSize = endIdx - startIdx;
44
    // Debug information
45
    printf("[DEBUG] Rank: %i, localSize: %i, startIdx: %i, endIdx: %i \n", rank, localSize, startIdx,
46
       endIdx):
    // Create a buffer for storing temperature fields
48
    Vector A = createVector(N);
49
50
    // Counts and displacements for MPI_Gatherv
51
    int *counts = NULL;
52
    int *displs = NULL;
53
    if (rank == 0) {
55
      counts = malloc(sizeof(int) * numProcs);
      displs = malloc(sizeof(int) * numProcs);
56
      for (int i = 0; i < numProcs; i++) {</pre>
57
        int tempStartIdx = i * chunkSize + (i < remainder ? i : remainder);</pre>
58
59
        int tempEndIdx = tempStartIdx + chunkSize + (i < remainder ? 1 : 0);</pre>
        counts[i] = tempEndIdx - tempStartIdx;
60
61
        displs[i] = tempStartIdx;
62
```

```
}
63
64
65
     int source_x = N / 4;
66
67
     if (rank == 0) {
       printf("Computing heat-distribution for room size N=%d for T=%d timesteps\n", N, T);
68
69
       // ----- setup -----
70
       // Set up initial conditions in A
71
       for (int i = 0; i < N; i++) {</pre>
72
        A[i] = 273; // temperature is 0 C everywhere (273 K)
73
74
75
       // Heat source in one corner
76
77
       A[source_x] = 273 + 60;
78
       printf("Initial:\t");
79
       printTemperature(A, N);
80
      printf("\n");
81
82
83
     // Distribute A to all processes
84
     MPI_Bcast(A, N, MPI_DOUBLE, O, MPI_COMM_WORLD);
85
     printf("[DEBUG] Broadcasted A to every rank for computation. \n");
86
87
88
     // Start time measurement before the computation begins
     starttime = MPI_Wtime();
89
90
     // ----- compute -----
91
92
     // Create buffers for the computation and initialize with A
93
94
     Vector B = createVector(localSize);
     Vector C = createVector(localSize);
95
     for (int i = startIdx; i < endIdx; i++) {</pre>
      B[i - startIdx] = A[i];
97
98
99
     // Create variables for the neighbours
100
     value_t rightNeighbour = 0;
101
     value_t leftNeighbour = 0;
     // For each time step
104
     for (int t = 0; t < T; t++) {
106
       // Exchange boundary values with neighboring processes
       if (rank > 0) {
        MPI_Sendrecv(&B[0], 1, MPI_DOUBLE, rank - 1, 0, &leftNeighbour, 1, MPI_DOUBLE, rank - 1, 1,
108
       MPI_COMM_WORLD, MPI_STATUS_IGNORE);
109
       } else {
         leftNeighbour = B[0];
110
111
112
       if (rank < numProcs - 1) {</pre>
         MPI_Sendrecv(&B[localSize - 1], 1, MPI_DOUBLE, rank + 1, 1, &rightNeighbour, 1, MPI_DOUBLE,
113
       rank + 1, 0, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
114
       } else {
         rightNeighbour = B[localSize - 1];
115
116
117
       // Propagate the temperature
118
       for (int i = startIdx; i < endIdx; i++) {</pre>
119
         // Heat source remains constant
120
         if (i == source_x) {
121
           C[i - startIdx] = B[i - startIdx];
122
           continue;
124
125
         // Current temperature
126
         value_t tc = B[i - startIdx];
127
```

```
// Temperatures of adjacent cells
129
         value_t tl = (i == startIdx) ? leftNeighbour : B[i - 1 - startIdx];
130
         value_t tr = (i == endIdx - 1) ? rightNeighbour : B[i - startIdx + 1];
131
132
133
         // Compute new temperature
         C[i - startIdx] = tc + 0.2 * (tl + tr + (-2 * tc));
134
135
136
       // Swap buffers
137
       Vector H = B;
138
       B = C;
139
       C = H;
140
141
       // Print temperature at intervals
142
143
       if (t % 10000 == 0) {
         MPI_Gatherv(B, localSize, MPI_DOUBLE, A, counts, displs, MPI_DOUBLE, 0, MPI_COMM_WORLD);
144
145
         if (rank == 0) {
146
           printf("Step t=%d:\t", t);
147
           printTemperature(A, N);
148
           printf("\n");
149
150
       }
151
     }
152
153
     // Gather final results from all processes
154
     MPI_Gatherv(B, localSize, MPI_DOUBLE, A, counts, displs, MPI_DOUBLE, O, MPI_COMM_WORLD);
155
156
     releaseVector(B);
157
158
     releaseVector(C);
159
160
     // End time measurement after computation
     endtime = MPI_Wtime();
161
     // ----- check -----
163
164
     int success = 1;
165
     if (rank == 0) {
166
      printf("Final:\t\t");
167
       printTemperature(A, N);
168
       printf("\n");
169
170
       for (int i = 0; i < N; i++) {</pre>
171
         value_t temp = A[i];
172
         if (273 <= temp && temp <= 273 + 60)
           continue;
174
         success = 0:
175
176
         break;
177
178
179
       printf("Verification: %s\n", (success) ? "OK" : "FAILED");
180
181
       // Print the time
       printf("WTime: %f seconds\n", endtime - starttime);
182
183
184
       // Free counts and displacements arrays
       free(counts);
185
       free(displs);
187
188
     // ----- cleanup -----
189
190
     releaseVector(A);
191
192
     // Finalize MPI
193
194
     MPI_Finalize();
     return (success) ? EXIT_SUCCESS : EXIT_FAILURE;
195
196 }
```

```
197
198 Vector createVector(int N) {
return malloc(sizeof(value_t) * N);
200 }
202 void releaseVector(Vector m) {
203 free(m);
204 }
205
void printTemperature(Vector m, int N) \{
   const char *colors = " .-:=+*^X#%0";
207
     const int numColors = 12;
208
209
    // Boundaries for temperature (for simplicity hard-coded)
210
211
    const value_t max = 273 + 30;
    const value_t min = 273 + 0;
212
213
     // Set the 'render' resolution
214
     int W = RESOLUTION;
215
216
     // Step size in each dimension
217
     int sW = N / W;
219
     // Room
220
     // Left wall
221
     printf("X");
222
     // Actual room
223
     for (int i = 0; i < W; i++) {</pre>
224
      // Get max temperature in this tile
225
226
      value_t max_t = 0;
      for (int x = sW * i; x < sW * i + sW; x++) {</pre>
227
228
        max_t = (max_t < m[x]) ? m[x] : max_t;
229
       value_t temp = max_t;
231
       // Pick the 'color'
232
       int c = ((temp - min) / (max - min)) * numColors;
233
       c = (c >= numColors) ? numColors - 1 : ((c < 0) ? 0 : c);
234
235
       // Print the average temperature
236
      printf("%c", colors[c]);
237
238
    // Right wall
239
     printf("X");
240
241 }
```

Listing 3: Parallel propagation



2.4 Discuss the effects and implications of your parallelization.

1. Performance Improvements

With MPI, the problem is split into parts, and each process (rank) works on one part. This way, the work is shared between cores or nodes, which makes the simulation faster compared to the sequential version.

2. Data Communication Overhead

The parallel version needs processes to exchange boundary information using MPI_Sendrecv. This adds communication overhead, especially as the number of processes increases. Also, collective functions like MPI_Bcast and MPI_Gather make all processes wait for each other, which can slow down the program if not managed well.

3. Scalability Considerations

Each process handles the same amount of work. But if the problem size does not divide evenly between processes (e.g., N%numProcs $\neq 0$), the program stops. More processes can also increase communication overhead, especially if there is too much communication compared to the actual computation.

4. Potential Bottlenecks

Global synchronization using MPI_Bcast and MPI_Gather can cause delays because all processes must wait for the slowest one to finish its work. This may lead to idle time, reducing efficiency.