2MP3 Assignment 4

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1 Introduction

One of the largest problems to solve in industry with regards to Engineering is the task of optimization. Traditionally, this is done with methods like Gradient Descent. However, with regards to functions with many local minima it may be difficult to find the minimum possible value. This is where the benefit of **Particle Swarm Optimization** (**PSO**) is. This allows us to place many [**particles**] over the domain of the function we wish to minimize and allow them to trend towards a global minima.

2 Problem Statement

In this assignment I developed a PSO algorithm with the task of minimizing the following functions:

- 1. Griewank
- 2. Levy
- 3. Rastrigin
- 4. Rosenbrock
- 5. Schwefel
- 6. Dixon-Price
- 7. Michalewicz with m=10
- 8. Styblinksi-Tang

This optimization had to be carried out over an arbitrary number of dimensions, denoted in an input array of \mathbf{x} . The values of \mathbf{x} are modified through a velocity function which is calculated at each iteration of the function. The max number of iterations that is allowed is defined by the user.

3 Solution

My solution mostly followed the layout from the assignment pdf, however I did make a few notable changes to ensure convergence with higher dimension versions of the more complex functions.

All of the logic for the algorithm is controlled by the **PSO()** function inside the **PSO.c** file. Below I have separated the development of the algorithms into sections.

3.1 Random Number Generation

One of the key changes that I made when developing this algorithm when compared to the original layout was to swap out the default random number generation logic. The original random number generating function in the file did not work properly on my computer so I instead decided to implement the **drand48()** random number generation function. The implementation for this is the following:

```
// New random function because the old one wouldn't work for me, this uses drand48
double random_double(double min, double max) {

//Create a random fraction from 0 to 1
double random_fraction = drand48();

//Return the random number that is generated within the bounds
return min+ (max-min)*random_fraction;

}
```

3.2 Particle Initialization

When first creating the particles it is important that they are initialized, this is done by looping through all of the particles and assigning a random position within the bounds of the function for them to be placed at. From there the algorithm can iteratively move the particles towards the global minima. Below is the implementation of the initialization part of the **PSO.c** function:

```
//Initialize all of the dimensions for each particle
       for(int i = 0; i < NUM_PARTICLES; i++) {</pre>
2
3
5
           particleArray[i]->decisionVariables = (double *)malloc(NUM_VARIABLES*sizeof(double));
6
           particleArray[i]->personalBest = (double *)malloc(NUM_VARIABLES*sizeof(double));
           particleArray[i]->velocityVariables = (double *)malloc(NUM_VARIABLES*sizeof(double));
8
9
10
           for(int j=0; j < NUM_VARIABLES; j++) {</pre>
12
                double random_num = random_double(bounds->lowerBound, bounds->upperBound);
14
                particleArray[i]->decisionVariables[j] = random_num;
                particleArray[i]->personalBest[j] = random_num;
16
17
                //We set the velocity to a random number between -1 and 1 \,
18
                particleArray[i]->velocityVariables[j] = random_double(-1, 1);
19
20
21
            //We do this because the first iteration will always be the best
           double currentPosition = objective_function(NUM_VARIABLES, particleArray[i]->
25
                decisionVariables);
           particleArray[i]->currentPosition = currentPosition;
26
           particleArray[i]->personalBestFitness = currentPosition;
27
29
              (currentPosition < globalBestPosition.currentValue) {</pre>
30
                globalBestPosition.currentValue = currentPosition;
31
                memcpy(globalBestPosition.decisionVariables, particleArray[i]->decisionVariables,
                    NUM_VARIABLES * sizeof(double));
35
```

The commenting in the code provides a more detailed look into the logic behind this portion of the function however essentially it loops through all of the dimensions of the decision variables and the velocities and assigns random numbers to them within the bounds specified. It is worth noting that earlier in the function all of these lists were **malloced** so that the memory was properly accounted for to store the variables.

3.3 PSO Algorithmn

Most of the PSO algorithm follows the general structure that is outlined in the pseudocode at the bottom of the Assignment 4 PDF, however there is one key change that I would like to highlight in the report so that it can properly be explained. One of the key changes I made to allow for convergence of the high dimension, high complexity functions was to check for stagnant iterations. It is useful to note which functions are being referenced when discussing high complexity function, these are the four functions which provided the most trouble when converging and required the extra modification to the algorithm:

- 1. Schwefel
- 2. Dixon-Price
- 3. Michalewicz
- 4. Styblinski-Tang

Ultimately I was unable to get Dixon-Price to converge for 50 variables, and I was unable to get both Schwefel and Dixon-Price to converge for 100 variables. It is likely that given more time and using a massive amount of particles that these would've eventually converged, especially with the stagnant particle management. With regards to the stagnant particle management, the following is how that was dealt with:

```
if (particleArray[i]->stagnantIterations > STAGNANT_ITERATIONS) {
2
3
                    for (int j = 0; j < NUM_VARIABLES; j++) {</pre>
5
6
                        particleArray[i]->decisionVariables[j] += random_double(-0.8, 0.8);
7
8
                        particleArray[i]->decisionVariables[j] = fmax(bounds->lowerBound, fmin(bounds->
10
                            upperBound, particleArray[i]->decisionVariables[j]));
12
                        particleArray[i]->velocityVariables[j] = random_double(-1, 1);
13
14
                    particleArray[i]->stagnantIterations = 0;
16
```

On every iterion of the algorithm, for each particle, if the best position of the particle does not iunmprove then we increment the stagnant iterations count for that particle by 1. If the stagnant iterations count for the particle reaches the cutoff value, which has been set to 50, however this can be adjusted, then the position of each of the decision variables will be stepped in a random direction by a random value within the bounds of the function. The position is clamped so that it doesn't go out of bounds. Additionally, the velocity is reset so that it can move in a new direction if needed.

The implementation of this function drastically increases the convergence of the algorithm, especially for higher dimensions.

3.4 CPU Timing

The final code that needed to be developed for this project was the CPU timing. This was done using the global clock with the following function:

4 Results

Below is the tabulated list of results for the functions. It is worth noting since we seed the random number generator each iteration with the current time, these results will vary in between runs, however eventually it will convergence if it has done so in the table below.

Table 1: Num Vars = 10

Function	Lower Bound	Upper Bound	Particles	Iterations	Optimal Fitness	CPU Time (sec)
Griewank	-600	600	20000	1000	0.000000	1.164s
Levy	-10	10	10	1000	0.000000	$0.003417\mathrm{s}$
Rastrigin	-5.12	5.12	2500	1000	0.000000	0.372904s
Rosenbrock	-5	10	3000	1000	0.000009	0.5465952s
Schwefel	-500	500	40000	1000	0.000127	7.328s
Dixon-Price	-10	1	20000	1000	0.000000	0.4259s
Michalewicz	0	π	1000	900	-9.6601517	0.418091s
Styblinski-Tang	-5	5	1000	10000	-391.661657	3.605633s

Table 2: Num Vars = 50

Function	Lower Bound	Upper Bound	Particles	Iterations	Optimal Fitness	CPU Time (sec)
Griewank	-600	600	50000	1000	0.000000	1.752017s
Levy	-10	10	10000	1000	0.000000	6.4723s
Rastrigin	-5.12	5.12	5000	52100	0.000000	$238.8735\mathrm{s}$
Rosenbrock	-5	10	5000	19000	0.000000	82.5768s
Schwefel	-500	500	500	100000	0.000637	50.393102s
Dixon-Price	-10	10	1000	(missing)	(missing)	(missing)
Michalewicz	0	π	1000	10000	-46.084809	$22.30346 \mathrm{s}$
Styblinski-Tang	-5	5	1000	10000	-1958.308285	16.828267 s

Table 3: Num Vars = 100

Function	Lower Bound	Upper Bound	Particles	Iterations	Optimal Fitness	CPU Time (sec)
Griewank	-600	600	7000	400	0.000000	4.033s
Levy	-10	10	7000	1000	0.000000	32.788439s
Rastrigin	-5.12	5.12	7000	24200	0.000000	564.275483s
Rosenbrock	-5	10	7000	23900	0.000000	303.136329s
Schwefel	-500	500	(missing)	1000	(missing)	(missing)
Dixon-Price	-10	10	(missing)	1000	(missing)	(missing)
Michalewicz	0	π	1000	10000	-88.831491	46.021296s
Styblinski-Tang	-5	5	1000	10000	-3916.61657	34.2172s

From this data we can see that, even with the large number of dimensions for some of the functions, the convergence occurs relatively fast. Comparing this to before the stagnant particle management was implemented it provides anywhere from a 2x to a 10x performace uplift.

In the next section of the report I will detail how to operate the code.

5 Instructions for Operation

Operation of the code is not challenging. Included with the code is a Makefile that allows the user to type **make** into the terminal and compile and link the code. Once the code is compiled all that is required from the user is to type ./main followed by the parameters you want, practically this looks like ./main ObjectiveFunctionName NUM-VARIABLES LowerBound UpperBound NUM-PARTICLES MAX-ITERATIONS.

A C Code

```
2
3
   #include <stdlib.h>
   #include <stdio.h>
5
   #include <time.h>
6
   #include <string.h>
   #include <math.h>
9
   //Define all of the constants that we need
10
   #define INERTIAL_WEIGHT 0.7
   #define COGNITIVE_COEFFICIENT 1.5
   #define SOCIAL_COEFFICIENT 1.5
14
    #define STAGNANT_ITERATIONS 100
15
16
17
   double random_double(double min, double max) {
18
19
       double random_fraction = drand48();
20
21
22
        return min+ (max-min)*random_fraction;
23
24
27
          pso(ObjectiveFunction objective_function, int NUM_VARIABLES, Bound *bounds, int
28
       NUM_PARTICLES, int MAX_ITERATIONS, double *best_position) {
30
       Particle* particleArray[NUM_PARTICLES];
31
32
33
        for (int i = 0; i < NUM_PARTICLES; i++) {</pre>
35
            particleArray[i] = (Particle *)malloc(sizeof(Particle));
36
37
            if (particleArray[i] == NULL) {
38
                printf("Error_allocating_memory_for_Particles...");
39
                exit(EXIT_FAILURE);
40
41
42
43
44
       BestPosition globalBestPosition;
45
46
47
       globalBestPosition.decisionVariables = (double *)malloc(NUM_VARIABLES * sizeof(double));
48
49
        if (globalBestPosition.decisionVariables == NULL) {
51
            printf("Error allocating memory for global best position...");
            exit(EXIT_FAILURE);
53
56
       globalBestPosition.currentValue = INFINITY;
```

```
//Initialize all of the dimensions for
59
        for(int i = 0; i < NUM_PARTICLES; i++) {</pre>
60
61
63
            particleArray[i]->decisionVariables = (double *)malloc(NUM_VARIABLES*sizeof(double));
64
            particleArray[i]->personalBest = (double *)malloc(NUM_VARIABLES*sizeof(double));
65
            particleArray[i]->velocityVariables = (double *)malloc(NUM_VARIABLES*sizeof(double));
66
67
68
            for(int j=0; j < NUM_VARIABLES; j++) {</pre>
69
                 double random_num = random_double(bounds->lowerBound, bounds->upperBound);
70
71
72
                 //We want to set the decision variables and the personal best to the same value, that
                 particleArray[i]->decisionVariables[j] = random_num;
                 particleArray[i]->personalBest[j] = random_num;
74
76
                 particleArray[i]->velocityVariables[j] = random_double(-1, 1);
77
78
79
80
81
82
            double currentPosition = objective_function(NUM_VARIABLES, particleArray[i]->
83
                 decisionVariables);
            particleArray[i]->currentPosition = currentPosition;
            particleArray[i]->personalBestFitness = currentPosition;
85
86
87
            if (currentPosition < globalBestPosition.currentValue) {</pre>
88
                 globalBestPosition.currentValue = currentPosition;
89
90
                 memcpy(globalBestPosition.decisionVariables, particleArray[i]->decisionVariables,
91
                     NUM_VARIABLES * sizeof(double));
92
93
94
95
96
        int iterationCount = 0:
97
98
        while (iterationCount < MAX_ITERATIONS) {</pre>
99
100
            srand48(time(NULL));
            for (int i = 0; i < NUM_PARTICLES; i++) {</pre>
104
106
                 for (int j = 0; j < NUM_VARIABLES; j++) {</pre>
108
109
                     double r1 = random_double(0, 1);
110
                     double r2 = random_double(0, 1);
111
114
                     particleArray[i]->velocityVariables[j] = INERTIAL_WEIGHT* particleArray[i]->
                         velocityVariables[j] +
                         COGNITIVE_COEFFICIENT * r1 * (particleArray[i]->personalBest[j] - particleArray
                             [i]->decisionVariables[j]) +
                         SOCIAL_COEFFICIENT * r2 * (globalBestPosition.decisionVariables[j] -
                             particleArray[i]->decisionVariables[j]);
118
                     particleArray[i]->decisionVariables[j] += particleArray[i]->velocityVariables[j];
120
```

```
position within bounds
                     particleArray[i]->decisionVariables[j] = fmax(bounds->lowerBound, fmin(bounds->
123
                         upperBound, particleArray[i]->decisionVariables[j]));
124
                     //Check to see if the position is within bounds, this will probably never happen
                     if (particleArray[i]->decisionVariables[j] < bounds->lowerBound || particleArray[i
                         ]->decisionVariables[j] > bounds->upperBound) {
127
                         printf("Error: Particle %d, Dimension %duis out of bounds \n", i, j);
                         exit(EXIT_FAILURE);
128
129
130
131
                 //Calculate the fitness of the current particle
                double position = objective_function(NUM_VARIABLES, particleArray[i]->decisionVariables
134
                particleArray[i]->currentPosition = position;
136
138
139
                if (position < particleArray[i]->personalBestFitness) {
140
141
                     memcpy(particleArray[i]->personalBest, particleArray[i]->decisionVariables,
142
                         NUM_VARIABLES * sizeof(double));
143
                     //Set the personal best fitness to the current position
144
                     particleArray[i]->personalBestFitness = position;
145
146
147
148
                     particleArray[i]->stagnantIterations = 0;
149
                     particleArray[i]->stagnantIterations += 1;
153
154
                 if (position < globalBestPosition.currentValue) {</pre>
158
159
                     globalBestPosition.currentValue = position;
161
                     memcpy(globalBestPosition.decisionVariables, particleArray[i]->decisionVariables,
                         NUM_VARIABLES * sizeof(double));
                //This is the main difference between the pseudo code in the assignment document and
                 //If the maximum number of stagnant iterations is reached, then we move the particle in
167
168
                if (particleArray[i]->stagnantIterations > STAGNANT_ITERATIONS) {
170
171
                     for (int j = 0; j < NUM_VARIABLES; j++) {</pre>
174
                         particleArray[i]->decisionVariables[j] += random_double(-0.8, 0.8);
176
177
                         particleArray[i]->decisionVariables[j] = fmax(bounds->lowerBound, fmin(bounds->
178
                             upperBound, particleArray[i]->decisionVariables[j]));
179
180
                         particleArray[i]->velocityVariables[j] = random_double(-1, 1);
181
182
                     particleArray[i]->stagnantIterations = 0;
183
```

```
184
185
186
187
190
191
193
194
195
                  printf("Particle 1: %.30f\n", particleArray[1]->currentPosition);
196
198
199
200
                 ((globalBestPosition.currentValue < 1e-4 && globalBestPosition.currentValue > 0)) {
201
203
204
205
              iterationCount += 1;
206
207
208
209
         memcpy(best_position, globalBestPosition.decisionVariables, NUM_VARIABLES * sizeof(double));
210
211
212
         // Free allocated memory
for (int i = 0; i < NUM_PARTICLES; i++) {</pre>
213
              free(particleArray[i]->decisionVariables);
214
              free(particleArray[i]->personalBest);
215
              free(particleArray[i]->velocityVariables);
216
              free(particleArray[i]);
217
218
219
         free(globalBestPosition.decisionVariables);
220
221
222
223
         return globalBestPosition.currentValue;
224
225
226
227
228
229
230
231
232
233
234
```

Listing 1: C code for the PSO algorithm implementation.

```
#ifndef UTILITY_H
#define UTILITY_H

// Function pointer type for objective functions
typedef double (*ObjectiveFunction)(int, double *);

typedef struct Bound{
    double lowerBound;
    double upperBound;

// Bound;

// Define the struct for the particle
// The struct is used to help with readibility of the program and to make designing the logic much easier

typedef struct Particle {
```

```
//Array for the dimensions of the particle
15
        double *decisionVariables;
16
17
        double *personalBest;
18
19
        double *velocityVariables;
20
21
        double currentPosition;
22
23
        double personalBestFitness;
24
25
        int stagnantIterations;
   } Particle;
27
28
29
    typedef struct BestPosition {
30
31
        double *decisionVariables;
        double currentValue;
32
   } BestPosition;
33
34
35
36
    double random_double(double min, double max);
37
    double {	t pso}({	t ObjectiveFunction} objective_function, int {	t NUM_VARIABLES}, {	t Bound} *{	t bounds}, int
        NUM_PARTICLES, int MAX_ITERATIONS, double best_position[]);
39
40
```

Listing 2: C code header for the function defitions and the package headers