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# 1 INTRODUCTION

This report presents a solution to the programming project **Evolutionary Algorithms** for the course INF421: Design and Analysis of Algorithms at École Polytechnique. Each task is developed in a section of the report which also contains the code implemented using the Python programming language.

#### 1.1 Instructions for running the project locally

The source code can be accessed on the project's Github repository. To execute it locally, clone the repository and install the project's dependencies.

```
git clone https://github.com/ArkhamKnightGPC/INF421.git pip install requirements.txt
```

All the code can be found in the code folder in the repository.

```
cd code
```

Now, to generate the scatter plots for the empiric runtime analysis of the **OneMax** and **LeadingOnes** benchmark functions (as described in task 2) we run the EmpiricRunTimes.py file. The generated plots are saved in the plots folder.

```
python EmpiricRunTimes.py
```

Similarly to generate the plot for the theoretical bound of the **OneMax** benchmark function, we run the dpOneMaxTheoreticalBound.py file. The generated plot is saved in the plots folder.

```
python dpOneMaxTheoreticalBound.py
```

Finally, to generate a series of plots for the empirical diversity tests of the  $(\mu + \lambda)$  GA, we run the GAtests.py file. The generated plots are saved in a folder inside the plots folder.

```
python GAtests.py
```

Unit tests are also provided in the unit\_tests folder.



# TASK 1: INDIVIDUALS AND BENCHMARK FUNCTIONS

Write code that allows to use individuals as well as the three functions OneMax, LeadingOnes, and Jump\_k. For individuals, do not use libraries but implement a data type that fully utilizes the memory. That is, do not store each bit value of an individual in a byte but in an actual bit.

The code for this project was developed with a strong respect for the SOLID design principles. For task1, taking the **single responsability principle** into account, three classes were developed: one defining the **Individual** data type, one providing implementations for the Benchmark Functions and one implementing the (1 + 1) EA.

#### 2.1 Individuals

Firstly, it is important to observe that the basic data types in Python use 1 byte of memory. Therefore, using a boolean variable for each bit value of an individual will not fully utilize memory. To do this, we use an array of integers, we each integer in the array represents 32 bit values.

The Individual class also provides auxiliary functions that will be used later on:

- a get function to retrieve a single bit;
- a set function to set a bit value to 1;
- a reset function to set a bit value to 0;
- a flip function to change the value of a single bit;
- a count function returning the number of bits equal to 1

```
class Individual:

"""

Represents candidate solutions x = (x1, ..., xn)

"""

def __init__(self, size):

"""

Constructor for new Individual

"""

# Number of integers necessary to represent all xi's

necessary_integers = (size + 31) // 32
```



```
11
           self.size = size
           self.bits = [0] * necessary_integers
12
13
      def get(self, idx):
14
           Get bit at index idx
16
17
           test_bit = self.bits[idx // 32] & (1 << (idx % 32))
           return 1 if test_bit > 0 else 0
19
20
      def set(self, idx):
21
22
23
           Set bit at index idx to 1
24
           self.bits[idx // 32] |= (1 << (idx % 32))
25
27
      def reset(self, idx):
28
           Set bit at index idx to 0
29
30
           self.bits[idx // 32] &= ~(1 << (idx % 32))
31
32
      def flip(self, idx):
33
           Flip bit at index idx
35
36
           bit_i = self.get(idx)
37
           if bit_i == 0:
               self.set(idx)
39
           else:
40
               self.reset(idx)
41
42
      def count(self):
43
44
           Count number of bits equal to 1
45
46
           result = 0
47
           for i in range(self.size):
48
               bit_i = self.get(i)
49
               result += bit_i
50
           return result
```

#### 2.2 Benchmark Functions

```
from Individual import Individual

def OneMax(individual):
    """

Returns the number of 1s of the input
```



```
return individual.count()
7
8
  def LeadingOnes(individual):
9
      Returns the length of the longest consecutive prefix of 1s
11
12
      n = individual.size
13
      result = 0
14
      for i in range(n):
          prefix_product = 1
16
          for j in range(0, i + 1):
17
18
               prefix_product *= individual.get(j)
           result += prefix_product
19
      return result
20
21
22
  def JumpK(individual, k):
23
      Analog to OneMax but penalizes individuals with a number of ones in n-k
24
     +1, \ldots, n-1
      n = individual.size
26
      one_max_x = OneMax(individual)
27
      if one_max_x <= n - k or one_max_x == n:</pre>
           return k + one_max_x
      return n - one_max_x
```

#### 2.3 (1+1) EA

Since the all-1s bit string is the unique global optimum of all three functions, we use a direct comparison all-1s bit string as our termination condition. A possible alternative is to impose a maximum number of iterations, but since we are interested in measuring performance against benchmark functions it is more interesting to let the EA reach the optimal solution. As mentioned in the statement the mutation rate adopted is  $p = \frac{1}{n}$ .

```
1 from Individual import Individual
 import numpy as np
3 import random
  def generateRandomOffspring(x, p):
      Generate a copy of x flipping each bit independently with probability p
      0.00
      n = x.size
9
      y = Individual(n)
10
      for idx in range(n):
11
          xi = x.get(idx)
12
          rand_var = np.random.uniform(0, 1)
13
         #bit idx in y is 1 if and only if
14
```



```
mutated_to_one = (rand_var 
     with probability p)
         stayed_one = (rand_var >= p and xi == 1) #did not mutate, was
16
     already 1 (probability 1-p)
        if (mutated_to_one or stayed_one):
17
             y.set(idx)
18
     return y
19
21 def EvolutionaryAlgorithm(f, n):
22
     (1+1) Evolutionary Algorithm
23
     0.00
24
     t = 0
     Pt = generateRandomOffspring(Individual(n), 0.5) # random initial
26
     solution
28
     while Pt.count() < n:</pre>
         y = generateRandomOffspring(Pt, 1 / n)
29
         if f(y) > f(Pt): # we pick solution that maximizes f
30
             Pt = y
         t += 1
33
     return Pt
34
```



# TASK 2: RUNTIME ANALYSIS

#### 3.1 Theoretical run time upper bounds

Prove mathematically (preferably rather tight) upper bounds on the expected run time of the (1+1) EA on OneMax and on LeadingOnes.

The method used for the proofs in this task is the classical fitness levels method (1).

Let  $(P^t)_{t\geq 0}$  represent the sequence of individuals in the population at each iteration of the algorithm, where

$$P^t = (P_1^t, \dots, P_n^t) \in \{0, 1\}^n \quad \forall t \ge 0.$$

We observe that  $(P^t)_{t\geq 0}$  is a markov chain with state space  $E=\{0,1\}^n$ .

We consider a fitness-based partition of the state space  $E = \bigcup_{i \in [0..n]} A_i$  where

$$\forall i \in [0..n] \quad A_i = \{x \in E \mid f(x) = i\}.$$

In the (1+1) EA we note that  $f(P^t) \geq f(P^{t-1})$   $\forall t \geq 1$ . Thus,  $(P^t)_{t\geq 0}$  is a non-decreasing level process.

Our strategy is to compute  $\forall i \in [0..n-1]$  the probability  $p_i$  of leaving level  $A_i$ . Then, the expected number of iterations to leave level  $A_i$  is  $\frac{1}{p_i}$ .

Thus, we have the upper bound

$$\sum_{i=1}^{n-1} \frac{1}{p_i}$$

for the expected run time.

#### 3.1.1 • Theoretical bound for OneMax

Let  $\mathcal{P}(m, i, j)$  denote the probabilty of leaving level  $A_i$  and arriving at level  $A_j$  in an iteration for a problem size of m bits. We impose the following constraints

$$1 < m < n; \quad 0 < i < m; \quad i < j < m.$$
 (1)

Let's discuss these coefficients can be calculated using dynamic programming. We start by defining the base cases (m, 0, j). We have

$$\mathcal{P}(m,0,j) = \frac{\binom{m}{j} p^j (1-p)^{m-j}}{\sum_{k=0}^m \binom{m}{k} p^k (1-p)^{m-k}}$$
(2)

Now, we formulate our recurrence  $\forall i \geq 1$ .



$$\mathcal{P}(m,i,j) = p\mathcal{P}(m-1,i-1,j) + (1-p)\mathcal{P}(m-1,i-1,j-1)$$
(3)

To calculate  $p_i$  using our coefficients, we have

$$p_i = \sum_{j=i+1}^n \mathcal{P}(n, i, j). \tag{4}$$

We have a  $O(n^3)$  algorithm to compute the coefficients  $p_i$  and thus the upper bound for the run time. Plotting the run time estimates for different values of n in figure 1, we conclude that the run time complexity is  $O(n \log(n))$ .

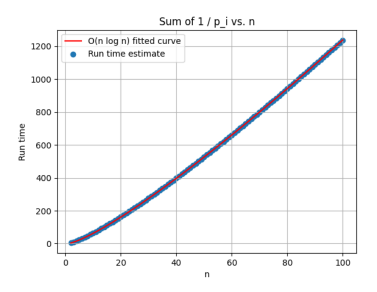


Figure 1:  $O(n \log(n))$  fit for theoretical runtime curve

```
import numpy as np
 from math import comb
  import matplotlib.pyplot as plt
  from scipy.optimize import curve_fit
  def calculate_coefficients(n, p):
      P = [[[0.0] * (n + 1) for _ in range(n + 1)] for _ in range(n + 1)]
      # Base case: m
      for m in range(1, n + 1):
10
          for j in range(m + 1):
11
              denominator = sum(comb(m, k) * (p ** k) * ((1 - p) ** (m - k))
12
     for k in range(m + 1))
              P[m][0][j] = comb(m, j) * (p ** j) * ((1 - p) ** (m - j)) /
13
     denominator
14
      # Recurrence relation
```



```
for m in range (2, n + 1):
          for i in range (1, m + 1):
17
              for j in range(i, m + 1):
18
                   P[m][i][j] = p * P[m - 1][i - 1][j] + (1 - p) * P[m - 1][i - p]
19
      1][j - 1]
20
      # Calculate p_i
21
      p_i = [0.0] * (n + 1)
      for i in range(n):
          for j in range(i + 1, n + 1):
24
              p_{i[i]} += P[n][i][j]
      return p_i
29 # Set up values for n
30 n_values = list(range(2, 101)) # Change the range as needed
32 # Calculate the sum for each n
33 sum_values = []
34 for n in n_values:
      coefficients = calculate_coefficients(n, 1 / n)
      sum_val = sum(1 / p if p != 0 else 0 for p in coefficients) # Include i
      = 0
      sum_values.append(sum_val)
_{39} # Plot the sum as a function of n
40 plt.plot(n_values, sum_values)
41 plt.xlabel('n')
42 plt.ylabel('Sum')
43 plt.title('Sum of 1 / p_i vs. n')
44 plt.grid(True)
45 plt.savefig('../plots/OneMaxTheoreticalRunTime.png')
47 #function for fitting
48 def fit_function(y, a):
     return a * y * np.log(y)
51 # Perform curve fitting
popt, pcov = curve_fit(fit_function, n_values, sum_values, maxfev=10000) #
     maxfev increased for more iterations
54 # Plot the data and fitted curve
55 plt.figure()
56 plt.scatter(n_values, sum_values, label='Run time estimate')
57 plt.plot(n_values, fit_function(n_values), *popt), 'r^-', label='0(n_values)
      log n) fitted curve')
58 plt.xlabel('n')
59 plt.ylabel('Run time')
60 plt.legend()
61 plt.grid(True)
62 plt.savefig('../plots/OneMaxTheoreticalRunTimeFit.png')
```



#### 3.1.2 • Theoretical bound for LeadingOnes

For the LeadingOnes function,  $\forall i \in [0..n-1]$ , a necessary and sufficient condition for a mutation to leave level  $A_i$  is to keep bits  $P_1^t, ..., P_i^t$  unchanged and to flip the bit  $P_{i+1}^t$ .

$$p_i = \frac{1}{n} \left( \frac{n-1}{n} \right)^i \tag{5}$$

Now, to estimate the run time

$$\sum_{i=0}^{n-1} \frac{1}{p_i} = \sum_{i=0}^{n-1} n \left( \frac{n}{n-1} \right)^i \tag{6}$$

$$= n \sum_{i=0}^{n-1} \left( 1 + \frac{1}{n-1} \right)^i \tag{7}$$

$$\leq n \sum_{i=0}^{n-1} \left( 1 + \frac{1}{n-1} \right)^{n-1} \tag{8}$$

$$\leq n \sum_{i=0}^{n-1} e 
\tag{9}$$

$$=en^2\tag{10}$$

so the expected time complexity is  $O(n^2)$ .

#### 3.2 Empirical run time estimates

Complement your theoretical bounds with empirical results and compare them.

In order to estimate the expected runtime empirically, we use the **law of large numbers**. Let  $k \in \mathbb{N}$  and  $T_1, ..., T_k$  represented run time measurements. We suppose the run times are independent and identically distributed. Then, by the law of large numbers,

$$\frac{T_1 + \dots + T_k}{k} \xrightarrow[k \to +\infty]{} \mathbb{E}[T_1] \tag{11}$$

For practical reasons, we adopt k = 30.

The empirical results were plotted in a scatter plot together with a curve representing the theoretical run time bound. Analysing the plots in figures 2 and 3, we conclude that for OneMax we observe indeed that the run time is a linear function of the problem size and for LeadingOnes we observe a quadratic curve. Thus, the theoretical run time estimates are accurate and fit well with the empirical results.



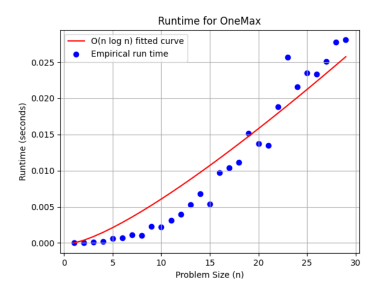


Figure 2: Empiric runtime analysis for OneMax

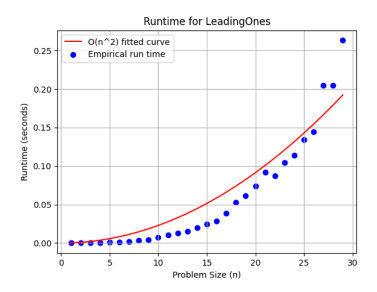


Figure 3: Empiric runtime analysis for LeadingOnes



#### 3.3 Empirical run time tests for the $(\mu + 1)$ EA

Furthermore, run empirical tests for the  $(\mu + 1)$  EA on OneMax with various, self-chosen values of  $\mu$ . Visualize the expected run time. What do you see? What  $\mu$  would you recommend?

In your report, do not forget to add a brief discussion about the parameter choices you made yourself, especially the number of tries for each value of  $\mu$  you chose.

The code for the  $(\mu + 1)$  EA is very similar to the (1 + 1) EA with the following changes:

- termination condition must be checked for all  $\mu$  individuals in the population;
- to determine if offspring is accepted, we must compare it to the least fit individual in the population.

There are two major computational costs associated with using  $\mu > 1$ :

- Sorting the population by fitness in order to eliminate the least fit individual, which must be done at each iteration in the algorithm description.
- Computing the fitness values of the entire population at each iteration.

In order to minimize the impact of these two operations, we use the **priority queue** data structure to store the population (implemented as a min heap in the **heapq** module in python).

In the priority queue, we store pairs (f(x), x) where x represents an individual in the population. This way we compute the fitness of each element only once when adding it to queue and then we can retrieve this value in the future without having to compute it again. And we can remove least fit individual by looking at the top of the min heap.

An important point is that we create a comparator for the **Individual** data type in order to use this approach. Since this is merely to break ties between individuals with same fitness, and we don't distinguish between elements with same fitness in the  $(\mu + 1)$  EA the exact content of this comparator is not important.

```
def __lt__(self, other):
    if(self.size < other.size):
        return True
    else:
        return False</pre>
```

Below is the code for the  $(\mu + 1)EA$ , which was added to EA.py.

```
from Individual import Individual import numpy as np
```



```
3 import random
4 import heapq
6 def generateRandomOffspring(x, p):
      Generate a copy of x flipping each bit independently with probability p
      0.00
9
      n = x.size
10
      y = Individual(n)
11
      for idx in range(n):
12
          xi = x.get(idx)
          rand_var = np.random.uniform(0, 1)
14
          #bit idx in y is 1 if and only if
          mutated_to_one = (rand_var 
16
     with probability p)
          stayed_one = (rand_var >= p and xi == 1) #did not mutate, was
     already 1 (probability 1-p)
          if (mutated_to_one or stayed_one):
18
              y.set(idx)
19
20
      return y
21
22 def EvolutionaryAlgorithm(f, n):
23
      (1+1) Evolutionary Algorithm
      0.00
      t = 0
26
      Pt = generateRandomOffspring(Individual(n), 0.5) # random initial
     solution
      while Pt.count() < n:</pre>
29
          y = generateRandomOffspring(Pt, 1 / n)
31
          if f(y) > f(Pt): # we pick solution that maximizes f
32
          t += 1
33
      return Pt
36
def EvolutionaryAlgorithm2(f, n, mu):
38
      (mu + 1) Evolutionary Algorithm
      11 11 11
40
      t = 0
41
      Pt = []
42
      most_fit_individual = (0, Individual(n)) #used to store solution
43
44
      for _ in range(mu):# we must create a random initial population of size
     mu
          individual = generateRandomOffspring(Individual(n), 0.5)
46
          fitness = f(individual)
47
          pair_fitness_individual = (fitness, individual) #we create pair (
48
     fitness, individual)
49
```



```
if(fitness > most_fit_individual[0]):
              most_fit_individual = pair_fitness_individual
51
52
          Pt.append(pair_fitness_individual)
53
      heapq.heapify(Pt) # we transform population into a priority queue
55
56
      while True:
          if (most_fit_individual[1].count() == n):
              return most_fit_individual[1]
60
61
          offspring = generateRandomOffspring(random.choice(Pt)[1], 1 / n)
62
          fitness = f(offspring)
63
          pair_fitness_offspring = (fitness, offspring)
          if(fitness > most_fit_individual[0]):
66
              most_fit_individual = pair_fitness_offspring
67
68
          heapq.heappushpop(Pt, pair_fitness_offspring) #we add offspring to
     the priority queue and pop least fit individual
70
```

In testing, the problem size was fixed n = 30. We tested  $\mu$  from 1 to 30. Same as before, in order to plot the run times we performed an average over 30 different execution for each value of  $\mu$ .

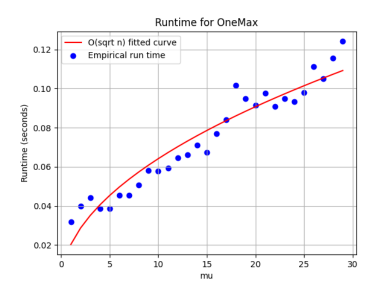


Figure 4: Run time for  $\mu \in [1..30]$ 

In figure 4, we observe that the run time is an increasing function of  $\mu$ . In order to study this dependency, different fit functions were experimented, and we conclude that the run time is  $O(\sqrt{\mu})$  for a fixed n.



#### 3.4 Code for empirical run time plots

```
import BenchmarkFunctions
2 import EA
3 import time
4 import matplotlib.pyplot as plt
5 import numpy as np
6 from scipy.optimize import curve_fit
7 from statistics import mean
9 def PlotOneMaxRunTime():
      0.00
10
      Generates scatter plot for empirical run time analysis using (1+1) EA
11
     and the OneMax benchmark function
12
      nvals = range(1, 30)
13
      run_times = []
14
15
      for n in nvals:
16
17
          number_of_trials = 30
          current_trial = 1
19
          trial_run_times = []
20
21
          while(current_trial <= number_of_trials):</pre>
23
               start_time = time.process_time() #we measure start time before
24
     running the EA
               solution = EA. Evolutionary Algorithm (Benchmark Functions. One Max, n
               end_time = time.process_time() #we measure end time after
26
     running the EA
27
               trial_run_times.append(end_time - start_time)
               current trial += 1
29
          run_times.append(mean(trial_run_times))
32
      #function for fitting
33
      def fit_function(y, a):
34
          return a * y * np.log(y) #Theoretical analysis points to n log(n)
     complexity
36
      # Perform curve fitting
      popt, pcov = curve_fit(fit_function, nvals, run_times, maxfev=10000) #
38
     maxfev increased for more iterations
39
      # we make a scatter plot for run times
41
      plt.figure()
      plt.scatter(nvals, run_times, color='blue', marker='o', label='Empirical
42
```



```
run time')
      plt.plot(nvals, fit_function(np.array(nvals), *popt), 'r-', label='0(n
43
     log n) fitted curve')
      plt.xlabel('Problem Size (n)')
44
      plt.ylabel('Runtime (seconds)')
      plt.title('Runtime for OneMax')
46
      plt.legend()
47
      plt.grid(True)
      #save plot in a png
50
      plt.savefig('../plots/OneMaxRunTime.png')
51
      return
  def PlotLeadingOnesRunTime():
54
      Generates scatter plot for empirical run time analysis using (1+1) EA
     and the LeadingOnes benchmark function
      0.00
      nvals = range(1, 30)
58
      run_times = []
59
60
      for n in nvals:
61
62
          number_of_trials = 30
          current_trial = 1
64
          trial_run_times = []
65
          while(current_trial < number_of_trials):</pre>
68
               start_time = time.process_time() #we measure start time before
69
     running the EA
               solution = EA. Evolutionary Algorithm (Benchmark Functions.
     LeadingOnes, n)
               end_time = time.process_time() #we measure end time after
71
     running the EA
72
               trial_run_times.append(end_time - start_time) #we use average of
73
     measurements as theestimator
               current_trial += 1
75
          run_times.append(mean(trial_run_times)) # we use average of
     measurements as theestimator
      #function for fitting
78
      def fit_function(y, a):
79
          return a * y * y #Theoretical analysis points to n^2 complexity
80
      # Perform curve fitting
82
      popt, pcov = curve_fit(fit_function, nvals, run_times, maxfev=10000) #
83
     maxfev increased for more iterations
84
      # we make a scatter plot for run times
85
```



```
plt.figure()
       plt.scatter(nvals, run_times, color='blue', marker='o', label='Empirical
87
       run time')
       plt.plot(nvals, fit_function(np.array(nvals), *popt), 'r-', label='0(n
88
      ^2) fitted curve')
       plt.xlabel('Problem Size (n)')
89
       plt.ylabel('Runtime (seconds)')
90
       plt.title('Runtime for LeadingOnes')
91
       plt.legend()
       plt.grid(True)
93
94
       #save plot in a png
95
       plt.savefig('../plots/LeadingOnesRunTime.png')
       return
97
  def PlotMuPlusOneEAOneMax():
99
100
       Generates scatter plot for empirical run time analysis using (mu + 1) EA
       and the OneMax benchmark function
       n = 30 #we keep n constant and vary only mu
103
       mu_vals = range(1, 30)
104
       run_times = []
105
       for mu in mu_vals:
107
108
           number_of_trials = 30
109
           current_trial = 1
110
           trial_run_times = []
           while(current_trial < number_of_trials):</pre>
113
                start_time = time.process_time() #we measure start time before
      running the EA
                solution = EA. Evolutionary Algorithm 2 (Benchmark Functions. One Max,
116
      n, mu)
                end_time = time.process_time() #we measure end time after
117
      running the EA
118
                trial_run_times.append(end_time - start_time) # we use average of
119
      measurements as theestimator
               current_trial += 1
120
121
           run_times.append(mean(trial_run_times)) # we use average of
      measurements as theestimator
123
       #function for fitting
       def fit_function(y, a):
           return a * np.sqrt(y) #Theoretical analysis points to n log(n)
126
      complexity
127
      # Perform curve fitting
128
```



```
popt, pcov = curve_fit(fit_function, mu_vals, run_times, maxfev=10000) #
       maxfev increased for more iterations
130
       # we make a scatter plot
131
       plt.figure()
       plt.scatter(mu_vals, run_times, color='blue', marker='o', label='
133
      Empirical run time')
       plt.plot(mu_vals, fit_function(np.array(mu_vals), *popt), 'r-', label='0
      (sqrt n) fitted curve')
       plt.xlabel('mu')
       plt.ylabel('Runtime (seconds)')
136
       plt.title('Runtime for OneMax')
137
138
       plt.grid(True)
       plt.legend()
139
140
       #save plot in a png
       plt.savefig('../plots/OneMaxRunTime2.png')
142
       return
143
144
145 PlotOneMaxRunTime()
146 PlotLeadingOnesRunTime()
147 PlotMuPlusOneEAOneMax()
```



# 4 TASK 3

#### 4.1 PLATEAU OF THE JUMP\_K BENCHMARK FUNCTION

Let  $k \in [1..n]$ . Assume that you start the (1+1) EA on the plateau of Jump\_k. Prove mathematically the expected number of iterations until the global optimum is created for the first time via standard bit mutation.

Let x be an individual such that  $\mathtt{OneMax}(x) = n - k$ . The probability that x mutates to the global optimum is

$$\mathfrak{p} = p^k (1-p)^{n-k} \tag{12}$$

Thus, the expected number of iterations until the global optimum is created (assuming  $p = \frac{1}{n}$ ) is

$$\frac{1}{\mathfrak{p}} = \frac{1}{p^k (1-p)^{n-k}} = \frac{n^n}{(n-1)^{n-k}} \tag{13}$$

It is important to observe that the probability  $\mathfrak{p}$  is a decreasing function of k and thus the expected number of iterations is an increasing function of k. In other words, the higher the value of k, the less likely the population is to leave the plateau.



# 5 TASK 4

#### 5.1 Implementation $(\mu + \lambda)$ GA

Implement the  $(\mu + \lambda)$  GA such that it can be run on pseudo-boolean functions.

Let's discuss some implementation details of the  $(\mu + \lambda)$  genetic algorithm. The structure is similar to that of the  $(\mu+1)$  EA, we keep the population in a priority queue of pairs (f(x), x) and at each iteration we check the fittest individual in the priority queue to evaluate the termination condition.

The process of generating the offspring at each iteration is, however, very different. We use a for loop to generate  $\lambda$  different individuals in the offspring. At each iteration, a uniform random variable branch\_decider determines if the new individual will be generated by recombination (with probability  $p_c$ ) or by the standard EA procedure.

The offspring is stored in an array and, after all  $\lambda$  individuals have been generated, we perform the **heappushpop** operation (which adds and right after removes least fit element in the priority queue) from the python **heapq** module to each of them. At the end, the population has the  $\mu$  fittest individuals generated so far. This is more efficient than storing population and offspring in an array and performing a sort operation.

```
1 import random
2 import heapq
 from Individual import Individual
  from EA import generateRandomOffspring
  def GeneticAlgorithm(f, n, k, mu, lamb, pc):
      (mu + lambda) Genetic Algorithm with recombination rate pc for Task4
8
      0.00
9
      t = 0
10
      Pt = []
11
      most_fit_individual = (0, Individual(n)) #used to store solution
13
      for _ in range(mu): # we must create a random initial population of size
14
          individual = generateRandomOffspring(Individual(n), 0.5)
          fitness = f(individual, k)
16
          pair_fitness_individual = (fitness, individual) #we create pair (
17
     fitness, individual)
          if (fitness > most_fit_individual[0]):
18
               most_fit_individual = pair_fitness_individual
19
          Pt.append(pair_fitness_individual)
20
21
```



```
heapq.heapify(Pt) # we transform population into a priority queue
22
23
      while True:
24
25
          if (most_fit_individual[1].count() == n):
              return most_fit_individual
28
          offspring = []
          for _ in range(lamb): #in each iteration, we will generate an
31
     individual in the offspring
              branch_decider = random.uniform(0, 1)
32
33
               if(branch_decider < pc): #we perform recombination with</pre>
34
     probability pc
                   individual1 = random.choice(Pt)[1]
                   individual2 = random.choice(Pt)[1]
36
                   new_individual = Individual(n)
37
                   for i in range(n):
38
                       #for each bit, we chose uniformly at random between bit
     value in individual1 and individual2
                       bit_value = random.choice([individual1.get(i),
40
     individual2.get(i)])
                       if(bit_value == 1): #if chosen value is 1, we set bit in
      offspring
                           new_individual.set(i)
42
               else: #else we do normal EA iteration
43
                   offspring.append(generateRandomOffspring(random.choice(Pt)
     [1], 1 / n)
45
          while(len(offspring) > 0):
               candidate_individual = offspring.pop(0)
47
              fitness = f(candidate_individual, k)
48
              candidate_pair = (fitness, candidate_individual)
49
50
              if(fitness > most_fit_individual[0]):
51
                   #we update most fit individual if necessary
                   most_fit_individual = candidate_pair
53
54
              #we add candidate_individual to the priority queue and pop least
      fit individual
              heapq.heappushpop(Pt, candidate_pair)
56
57
          t += 1
```



# 6 TASK 5

For at least three values of n (at least 100, preferably far larger), of k (do not go larger than 6 or 7 here, but larger than 1), of  $\mu$  (at least  $\lfloor \ln(n) \rfloor$ ), of  $\lambda$  (containing the value 1), and of  $p_c$  (containing the value 1), measure the diversity of the  $(\mu + \lambda)$  GA on Jump\_k, initialized such that the initial population  $P^{(0)}$  only contains individuals on the plateau, chosen uniformly at random. Stop the algorithm once the diversity for the maximum distance is sufficiently high (or after a maximum number of iterations).

For each parameter combination, pick one of the runs and visualize the diversity of the population over time. Please also mark the iteration when the optimum was found for the first time.

Please do not forget to briefly discuss your parameter choices, especially how many times you ran each setup. Furthermore, state whether plots for identical setups (of which you only show one in the report) are qualitatively the same. What do you see? Is there a correlation between some of the parameters and the number of iterations required in order to get to a certain level of diversity?

## 6.1 Modifications to $(\mu + \lambda)$ GA

Firstly, let's discuss the necessary modifications to the  $(\mu + \lambda)$  GA that are described in the problem statement. Instead of starting with a population picked uniformly at random over  $\{0,1\}^n$ , we pick individuals in the plateau  $\{x \text{ such that } \mathtt{OneMax}(x) = n - k\}$ . Then, we modify the termination condition to stop the algorithm once a fourth of pairs  $(x,y) \in P^t \times P^t$  is  $\geq 2k$  (or after a maximum number of iterations).

We create a function to generate individuals on the plateau uniformly at random. Then we add functions to compute the Hamming distance between two individuals, the diversity in the population for the maximum distance 2k and a boolean function for the termination condition.

```
import random
import heapq
import numpy as np
from itertools import combinations
from Individual import Individual
from EA import generateRandomOffspring

def GenerateOnPlateau(n, k):
    """

    Creates individual with k ones. Positions are chosen uniformly at random
    """
    random_permutation = np.random.permutation(range(0, n)) #we pick first n
    -k numbers in a random permutation of [1..n]
```



```
individual = Individual(n)
13
      for i in range(n-k):
14
          bit_idx = random_permutation[i]
15
          individual.set(bit_idx)
16
      return individual
17
18
19 def HammingDistance(individual1, individual2):
      n = individual1.size
20
      dist = 0
21
      for i in range(n):
22
          bit1 = individual1.get(i)
23
          bit2 = individual2.get(i)
24
          dist += (bit1 + bit2)%2
      return dist
26
  def Diversity(Pt, k):
29
      diversity = 0
      #we compute hamming distance between all pairs of individuals in the
30
     population
      for individual1 in Pt:
31
          for individual2 in Pt:
               dist = HammingDistance(individual1[1], individual2[1])
33
               if(dist == 2*k):
34
                   diversity += 1 #add to diversity pairs with maximum distance
      return diversity/2 #we divide by 2 because each pair is counted twice
37
38 def GeneticAlgorithm2(f, n, k, mu, lamb, pc, max_iter = 50):
      (mu + lambda) Genetic Algorithm with recombination rate pc for Task5
40
      0.00
41
      t = 0
42
      Pt = []
      diversities = []
44
      found_optimum = -1 #used to store time we find optimum
45
46
      for _ in range(mu): # we must create a random initial population of size
47
     mu
          individual = GenerateOnPlateau(n, k)
48
          fitness = f(individual, k)
49
          pair_fitness_individual = (fitness, individual) #we create pair (
     fitness, individual)
          Pt.append(pair_fitness_individual)
51
52
      heapq.heapify(Pt) # we transform population into a priority queue
53
54
      while True:
          diversity = Diversity(Pt, k) #we compute diversity of population
          diversities.append(diversity)
58
          total_pairs = (n*(n-1))/2
50
          if diversity >= total_pairs/4: #if diversity >= a fourth of total
     pairs, we break
```



```
return diversities, found_optimum
62
          if(t >= max_iter): #if we reach max iterations, we also break
63
              return diversities, found_optimum
64
          t += 1
66
          offspring = []
67
          for _ in range(lamb): #in each iteration, we will generate an
     individual in the offspring
              branch_decider = random.uniform(0, 1)
70
71
72
               if(branch_decider < pc): #we perform recombination with</pre>
     probability pc
                   individual1 = random.choice(Pt)[1]
73
                   individual2 = random.choice(Pt)[1]
                   new_individual = Individual(n)
75
                   for i in range(n):
                       #for each bit, we chose uniformly at random between bit
     value in individual1 and individual2
                       bit_value = random.choice([individual1.get(i),
     individual2.get(i)])
                       if(bit_value == 1): #if chosen value is 1, we set bit in
      offspring
                           new_individual.set(i)
80
                   offspring.append(new_individual)
81
               else: #else we do normal EA iteration
82
                   offspring.append(generateRandomOffspring(random.choice(Pt)
     [1], 1 / n))
84
          while(len(offspring) > 0):
               candidate_individual = offspring.pop(0)
              fitness = f(candidate_individual, k)
87
88
               if(candidate_individual.count() == n and found_optimum == -1):
89
                   #optimum must come from the offspring at some point
                   found_optimum = t
91
92
               #we add candidate_individual to the priority queue and pop least
      fit individual
              heapq.heappushpop(Pt, (fitness, candidate_individual))
```

### 6.2 Empirical tests for the $(\mu + \lambda)$ GA

We used the following parameter values for testing:

```
test_values = [(300, 2, 10, 5, 0.5), #base values n=300, k=4, mu=10, lambda=5, pc=0.5

(300, 3, 10, 5, 0.5), (300, 4, 10, 5, 0.5), #testing influence of k
```



```
(300, 2, 20, 5, 0.5), (300, 2, 30, 5, 0.5), #testing influence of mu

(300, 2, 10, 7, 0.5), (300, 2, 10, 10, 0.5), #testing influence of lambda

(300, 2, 10, 5, 0), (300, 2, 10, 5, 1)] #testing influence of pc
```

The Github repository for the project contains three complete series of tests each containing all 9 plots.

The maximum number of iterations set for testing was max\_iter = 100 to allow for better visualisation. For certain setups, we observe that the global optimum is not found. The setups used were empirically chosen because they generated the optimum solution rather quickly.

In particular, when the recombination rate  $p_c$  was 0, the algorithm never found a solution for max\_iter = 100. Which supports the idea that recombination is essential to finding the solution for the Jump\_k function. The number of iterations required also increases rapidly in function of k. The plots stick to smaller values in order to find solutions for max\_iter = 100, we show an example with k = 4 where the solution was not found.

In this report, we show only one series of plots as indicated in the problem statement. Please check the project repository for the full set of tests.

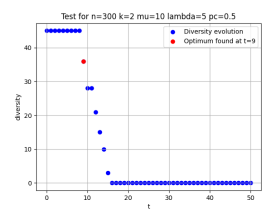
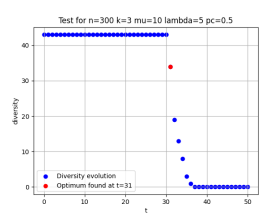


Figure 5: Test for n = 300, k = 2,  $\mu = 10$ ,  $\lambda = 5$ ,  $p_c = 0.5$ 

In this first setup, we observe that diversity is a decreasing function of the number of iterations. The recombination (or crossover) procedure decreases the diversity in the initially random individuals because we select each bit from values already existent in the population. Using biology terms, it is the EA procedure that will introduce *mutations* in the population while the crossover simply *selects* the already existing characteristics.





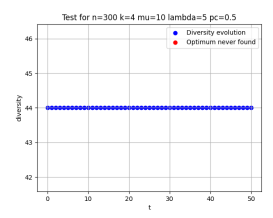
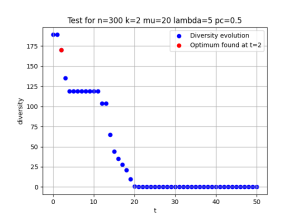


Figure 6: Varying value of k

In these plots we observe that the higher the value of k, the higher the number of iterations required to find the solution. This is logical because as we observed in task 3, the probability of leaving the plateau using the EA procedure is a decreasing function of k.



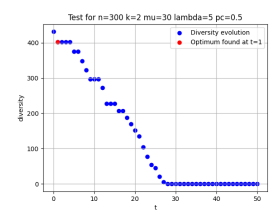
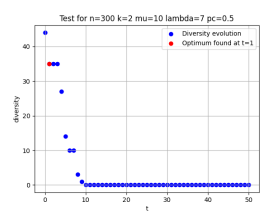


Figure 7: Varying value of  $\mu$ 

In these plots we observe that the higher the value of  $\mu$ , the faster we find the solution. We also observe that diversity falls more consistently on a larger population instead of the sudden drops we observe in smaller populations.





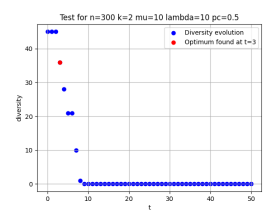
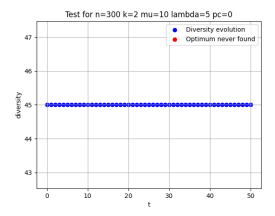


Figure 8: Varying value of lambda

In these plots we observe that the higher the value of  $\lambda$ , the faster we find the solution. This is effect is more significant than the one we observed for  $\mu$ . This logical because the more work we do per iteration, the less iterations it should take to find the solution. We observe very sudden diverse drops between iterations.



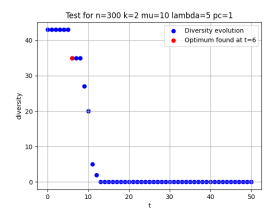


Figure 9: Varying value of  $p_c$ 

In these plots we observe that the higher the value of  $\lambda$ , the faster we find the solution. The recombination (or crossover) procedure is essential to finding a solution. For  $p_c = 0$ , we observe that diversity stays constant and no solution is found.

#### 6.3 Tests with identical setups

In this section, let's compare plots with identical parameters in the three series presented in the Github repository for the project.



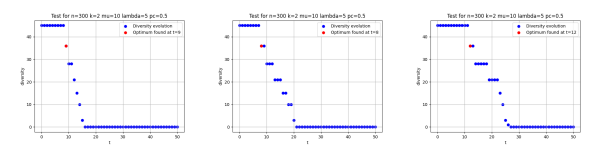


Figure 10: Tests for  $n = 300, k = 2, \mu = 10, \lambda = 5, p_c = 0.5$ 

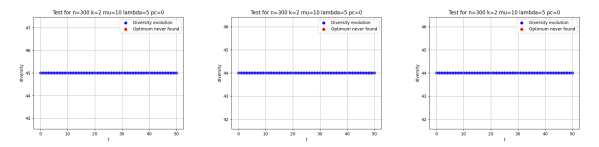


Figure 11: Tests for  $n = 300, k = 2, \mu = 10, \lambda = 5, p_c = 0$ 

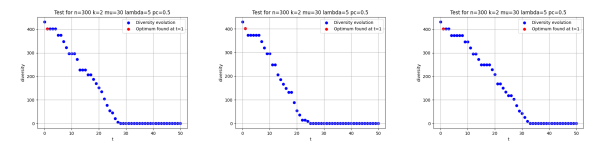


Figure 12: Tests for  $n = 300, k = 4, \mu = 15, \lambda = 1, p_c = 0.5$ 

We observe that even with multiple tests, the scenario where  $p_c = 0$  never generates the optimal solution even for a small value of k.

In general, we observe that though curves may vary in different tests runs due to the fact that the  $(\mu+\lambda)$  GA is a randomized algorithm, they still maintain similar shapes. In conclusion, a good choice of parameters tends to show good (or bad) results regardless of the random aspect of the algorithm.

## 6.4 Code used for tests on the $(\mu + \lambda)$ GA



```
1 import time
2 import numpy as np
3 from statistics import mean
4 import matplotlib.pyplot as plt
5 import GA_task5
6 import BenchmarkFunctions
  def PlotGeneticAlgorithmDiversities():
      Generates scatter plot for empirical run time analysis using (mu +
     lambda) GA and the Jumpk benchmark function
      0.00
11
12
      #we set parameter values for our tests
      test_values = [(300, 2, 10, 5, 0.5), #base values n=300, k=4, mu=10,
13
     lambda=5, pc=0.5
              (300, 3, 10, 5, 0.5), (300, 4, 10, 5, 0.5), #testing influence
     of k
              (300, 2, 20, 5, 0.5), (300, 2, 30, 5, 0.5), #testing influence
     of mu
              (300, 2, 10, 7, 0.5), (300, 2, 10, 10, 0.5), #testing influence
16
     of lambda
              (300, 2, 10, 5, 0), (300, 2, 10, 5, 1)] #testing influence of pc
17
18
      plot_number=1
20
      for n, k, mu, lamb, pc in test_values:
21
22
          diversities, found_optimum = GA_task5.GeneticAlgorithm2(
     BenchmarkFunctions.JumpK, n, k, mu, lamb, pc)
24
          # we make a scatter plot
          plt.figure()
          plt.scatter(range(len(diversities)), diversities, color='blue',
     marker='o', label='Diversity evolution')
          if (found_optimum > -1):
28
              plt.scatter(found_optimum, diversities[found_optimum], color='
     red', marker='o', label=f'Optimum found at t={found_optimum}')
30
              plt.scatter([], [], color='red', marker='o', label=f'Optimum
     never found')
          plt.xlabel('t')
32
          plt.ylabel('diversity')
33
          plt.title(f'Test for n={n} k={k} mu={mu} lambda={lamb} pc={pc}')
35
          plt.grid(True)
          plt.legend()
36
          #save plot in a png
          plt.savefig(f'../plots/GAplots3/GAtest{plot_number}.png')
          plot_number += 1
40
41
42 PlotGeneticAlgorithmDiversities()
```



# **REFERENCES**

[1] Doerr, B., Kötzing, T. Lower Bounds from Fitness Levels Made Easy. Algorithmica (2022). https://doi.org/10.1007/s00453-022-00952-w