# KU LEUVEN

# GENETIC ALGORITHMS

# TSP

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### 1 Implementation

### 1.1 Representation

The original code employed adjacency representation. In adjacency representation a tour is represented as a list of n cities where city j is listed in position i if and only if the tour leads from city i to city j. Thus, the list:

(76853421)

represents the tour:

3-8-1-7-2-6-4-5

1

In our implementation, we have decided to use path representation. Path representation is the most natural way of representing a tour. This can be easily seen with the following example. The list:

 $(1\ 2\ 7\ 5\ 6\ 3\ 4)$ 

represents the path

1-2-7-5-6-3-4

It was also the simplest representation possible for this problem, and it was easy to implement, as the tour was first encoded in path representation, and then translated to adjancency representation using the function path2adj. Finally, it was translated again to path representation to be used in the plots with the function adj2path.

#### 1.2 Crossover

The new representation required a new crossover operator. The available options were:

- Partially Matched Crossover (PMX)
- Order Crossover (OX)
- Cyclic Crossover (CX)
- Edge Recombination Crossover (ERX)

The one selected is Order Crossover. This crossover exploits a property of the path representation, that the order of the cities (not their positions) are important. It construct an offspring by choosing a subtour of one parent and preserving the relative order of cities of the other parent. Let's consider the following tours:

$$\begin{pmatrix}
1 & 2 & 3 & 4 & 5 & 6 & 7 & 8 \\
2 & 4 & 6 & 8 & 7 & 5 & 3 & 1
\end{pmatrix}$$

<sup>&</sup>lt;sup>1</sup>This explanation can be found in the slides about Traveling Salesman Problem

If we choose the cut point between the second and the third city, and the second cut point between the fifth and the sixth, we have:

$$(1\ 2 - 3\ 4\ 5 - 6\ 7\ 8)$$
  
 $(2\ 4 - 6\ 8\ 7 - 5\ 3\ 1)$ 

And the offspring is created copying the segments between the cut points, and then, starting from the second cut points of one parent, the rest of the cities are copied in the order in which they appear in the other parent, giving:

$$(8\ 7 - 3\ 4\ 5 - 1\ 2\ 6)$$
  
 $(4\ 5 - 6\ 8\ 7 - 1\ 2\ 3)$ 

### 1.3 Mutation

We were also asked to implement a new mutation operator. For this task, multiple and different options where available, such as Exchange Mutation, Scramble Mutation, Displacement Mutation or Insertion Mutation. As the crossover operator chosen exploits the order property aforementioned, we decided that the mutation should break this order in some way to avoid reaching a local optima due to the lack of diversity in the order.

The mutation operator selected for this problem is insertion mutation due to its simplicity and effectiveness. This mutation works in the following way. Imagine that we have the path:

Take the 2 out of the sequence,

and reinsert the 2 at a randomly chosen position:

#### 1.4 Fitness Function

The fitness function has been changed as the original one worked with adjacency representation. The simplest way to adapt the old function to the new representation was using path2adj, converting the path to adjacency representation, and then computing the fitness in the same way as it was computed before.

# 2 Experiments

In order to properly study and select the different configurations, it is necessary to analyze the behaviour of the algorithm, when we modify one parameter at a time. An automated test was developed for this purpose, provided in the annex, with which we run a set of tests, changing the number of individuals, the number of generations, the percentage of elitism, and the balance of percentage of mutation and crossover. Each test, in each city is run a total of 5 times, and the average is what is taken into account.

After that, a set of tests, created from the results obtained in the general set, is made, by selecting specific values. Both the general, and the specific tests will serve as a medium to compare both representations, in order to look for differences (or lack thereof)

### 2.1 General tests

For the general tests, the default values are:

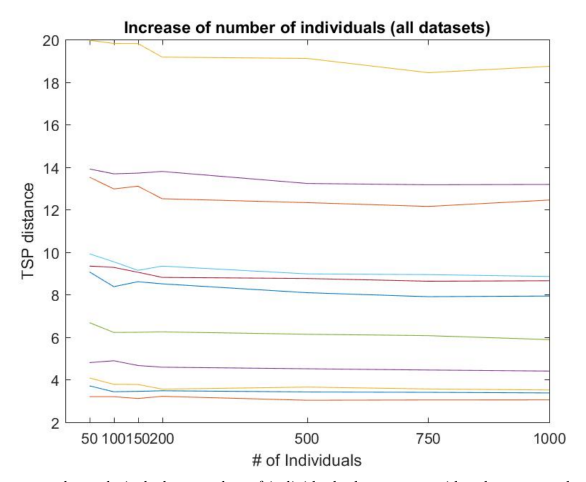
- Number of individuals 50
- Number of generations 50
- Elitism 0.05
- Crossover 0.95
- Mutation 0.05
- Stop percentage condition 0.95
- Detection of loops on

And, the different values for each modified parameter are

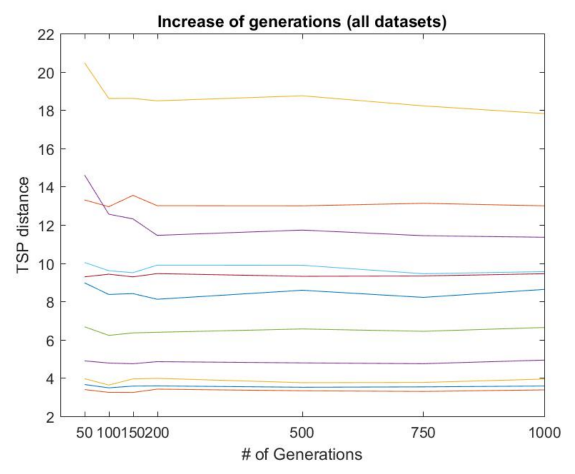
- Number of individuals [50,100,150,200,500,750,1000]
- Number of individuals [50,100,150,200,500,750,1000]
- Elitism percentage [0,0.05,0.1,0.2,0.5,0.75,1]
- Crossover—Mutation balance [1-0.0.95-0.05,0.9-0.1,0.75-0.25,0.5-0.5,0.25-0.75,0-1]

### 2.1.1 Adjacency representation

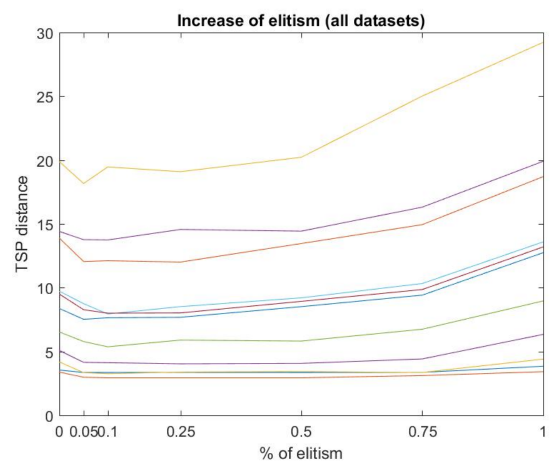
With the provided representation, adjacency representation, the results of the general tests are:



As expected, a relatively low number of individuals does not provide adecuate results, as can be observed by looking at the start of the graph. Almost all datasets start in global maximum, and only a couple in a local max. However, the mayority of them have one of their lowest point when the number of individuals is 200 (except for the highest dataset, which has it's lowest point at 750) and from that point, it stays constant, or even raises, as happens with the highest, and third highest dataset, thus it can be said that any number of individuals higher than 200 would not be benefitial, and actually be just cumbersome when it comes to computational cost.

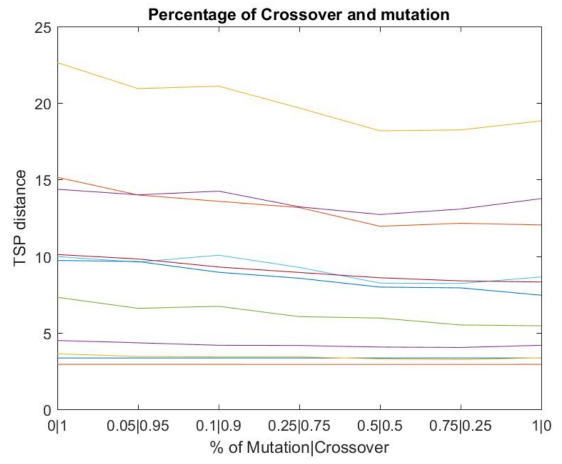


Once again, it is to be expected that a low quantity of generations will not yield good results. But that is not the only thing the number of individuals and generations have in common, since it seems like 200 is one of the best options for the number of generations. There are some differences, for instance, at lower quantities of generations, there is more fluctuation, and more datasets are positive towards higher number of generations. In the specific tests this will be further studied, whether 200, or a higher number is better, and whether the higher associated computational cost is worth.



When it comes to the percentage of elitism, the results are much more clear. With no exceptions, the lowest value for every single one of the datasets is between 0.05 and 0.1, any higher, or any lower, and the distance skyrockets, having the highest distances values at elitism = 1.

This phenomena has an easy explanation, the higher the elitism, the more likely it is that the algorithm will stay at a local maxima.

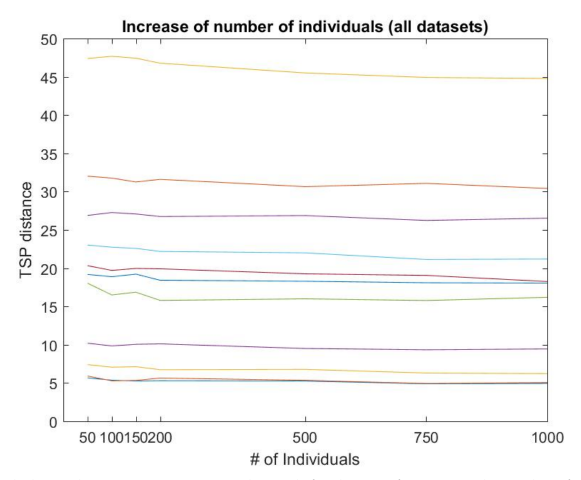


The test reflects perfectly the balance between exploitation and exploration, the overall result shows that the best performance comes when mutation has a value of 0.5, and crossover 0.5.

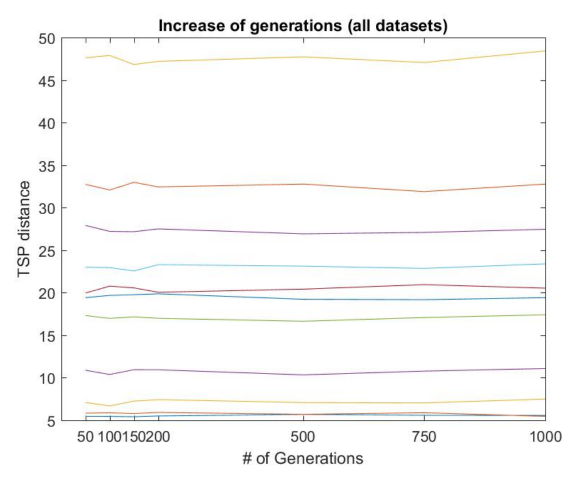
Any value of mutation higher than 0.5, and the results start to worsen, because there is too much exploitation, and too litle exploration. Any value of mutation lower than 0.5, and the results, most of the cases, are far worse. This leads to the conclusion that 0.5 is the candidate for the specific tests, alghouth the nature of the result makes it necessary to test other values, since, we think these results are a bit odd, hence we will further study in the specific tests, order to make a conclussion

### 2.1.2 Path representation

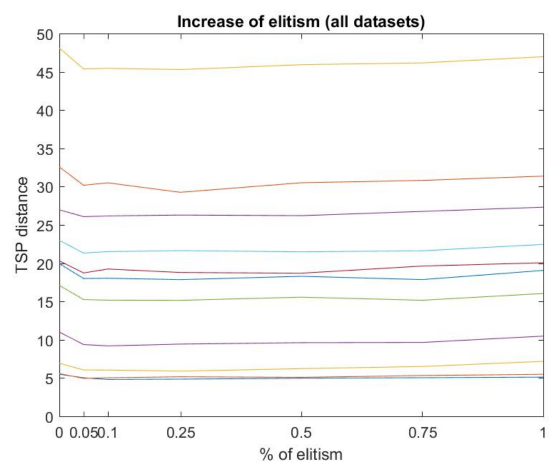
As explained in the implementation section, the representation we implemented is path representation. The results after executing the same tests as before are



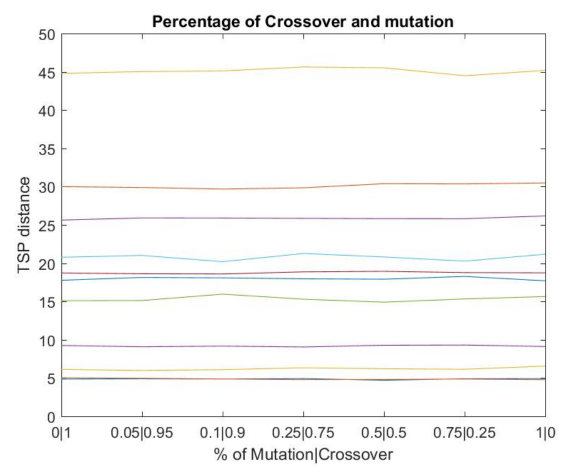
Similarly to adjacency representation, the result for the test of increasing the number of individuals has a generally located minimum local at 200, although for some cases, the distance becomes constant at 100. The only noticeable difference is that it is more stable at lower quantities of individuals, and the values for the distances when the number of individuals is very high (750,100) does not increase, rather it seems to keep ever so slightly decreasing.



Once again, as expected, there is a number of individuals from which the change in the result is null. That point, as can be observed, is 200 individuals, the same as the other representation. But then again, there are differences, two main ones, the first, the change from 50 to 100 individuals is not so apparent, and the values obtained from 750 individuals forward is actually worse in some cases, if not the same, while with the other representation there were some cases in which it improved.



From what can be observed, there is no doubt that 0.05 is the best percentage of elitism that can be choosen with this representation. The results are somewhat similar to the previous representation, but the slope at the latest values (0.5 forward) is not so steep



The result of this test is daunting. It was repeated, in case it was somehow erroneous, but the same graph was obtained. With our representation, there seems to be no effect whatsoever on the exploitation vs exploration dilemma. It does not matter, apparently, the percentage of mutation or crossover

### 2.2 Specific tests

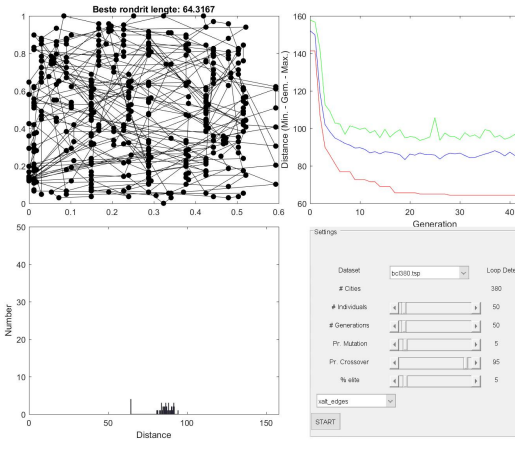
The specific tests are separeted in 2 phases.

First phase, study the impact of the best parameter values seen in the general tests, for each representation individually. The second phase corresponds to the comparison between the representation results, first with the base case, and then with the best result obtained. All the tests have been made with the benchmark dataset called bcl380.tsp.

### 2.2.1 Adjacency representation

The base values are the same as with the general case.

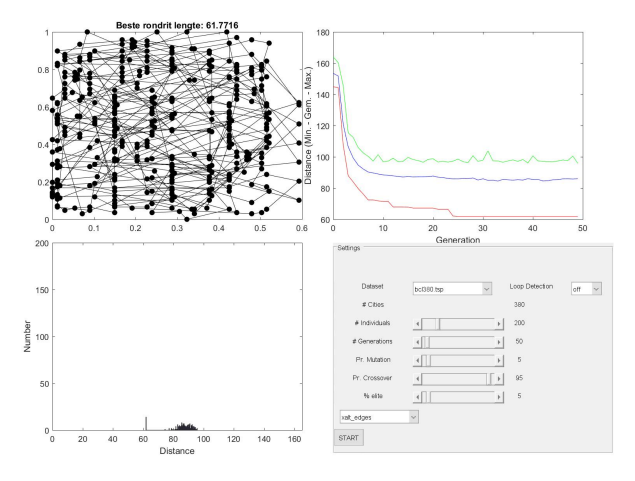
[Number of individuals, generations, percentage of elitism, of crossover and mutation] = [50,50,5%,95%,5%]



The result obtained is The test was executed in 12.66 sec

200 was the number of individuals that was deemed to be apropiate. The test, keeping all other values, but changing just the number of individuals is

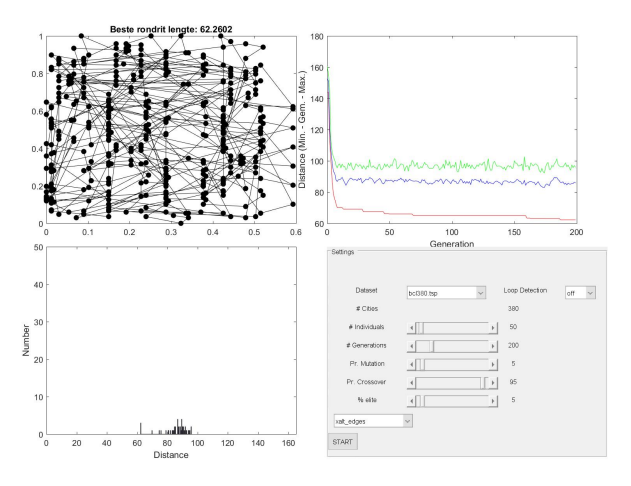
[Nind, gens, elitism, crossover, mutation] = [200,50,5%,95%,5%]



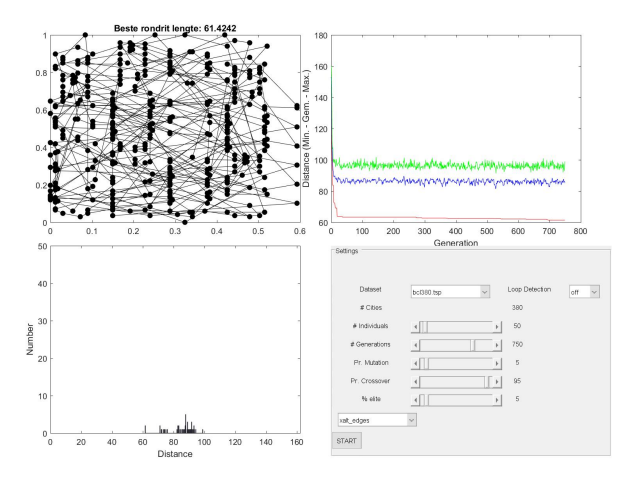
We can see that the minimum distance has decreased, from 64 to 61, not a big decrease, but significant enough. Increasing the number of individuals to 200 is then to be considered a good measure. Timewise, the test was executed in 31.61 seconds. More than twice the time for the first, which, given the increase in the number of individuals, is not out of the expected.

For the next test, this time the number of generations has been increased. As previously stated, the best number was 200, but a higher number was also suitable for some cases. We tried, respectively, with 200, 750, and 1000 generations, thus having

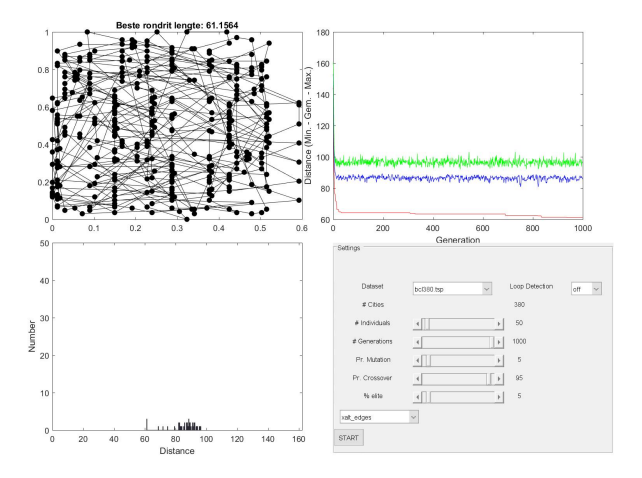
[Nind, gens, elitism, crossover, mutation] =[50,200,5%,95%,5%]



 $[\mathrm{Nind,\,gens,\,elitism,\,crossover,\,mutation}] = [50,750,50,5\%,95\%,5\%]$ 

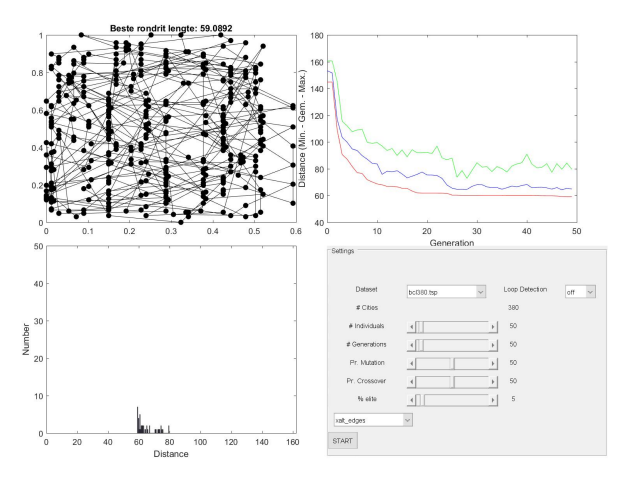


[Nind, gens, elitism, crossover, mutation] =[50,1000,5%,95%,5%]

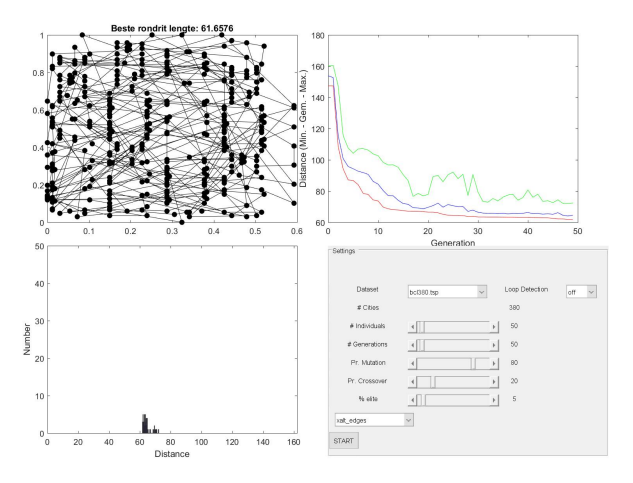


We can observe that the result is slightly better, with each increment of generations, but the cost is excesive, when it comes to time. The times for the tests are 46.01, 171.67, and 233.85 seconds, which means, 4, 13.5 and 18.5 times more than the base case. As the elitism parameter was abundantly clear that was to be kept at 0.05 (or up to 0.1 at most), and it is already the value for the base case, there is no specific test for the elitism. There is, however, for the percentage of crossover and mutation. Since the result was a bit unexpected, we did not only try with the apparent best result (0.5 percentage of each), but with cases of 0.2 crossover — 0.8 mutation, and viceversa. The results for the tests are

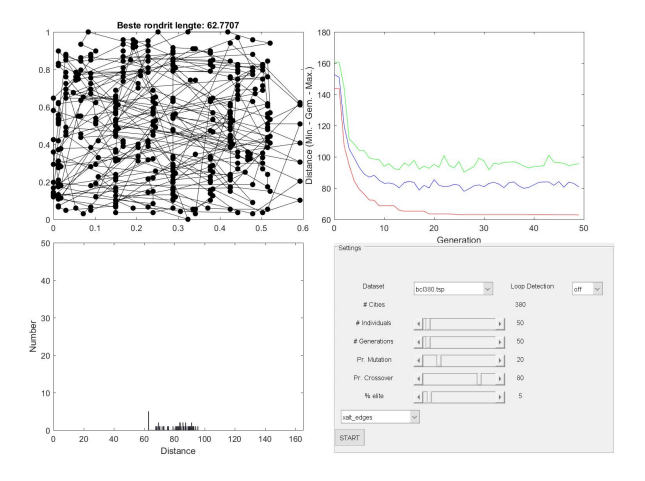
[Nind, gens, elitism, crossover, mutation] = [50,50,5%,50%,50%]



[Nind, gens, elitism, crossover, mutation] =[50,50,5%,20%,80%]



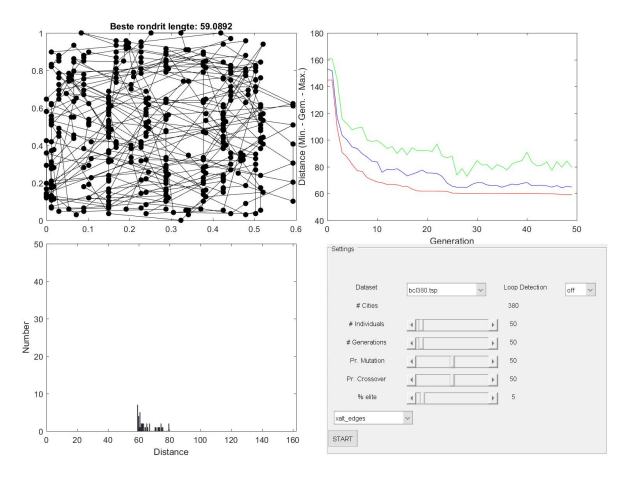
[Nind, gens, elitism, crossover, mutation] = [50,50,5%,80%,20%]



The test confirms the results obtained in the general test. The best ratio of crossover—mutation is 50%—50%, since, out of the 3 configurations, it has the best result, while all of them are over the base case. Although not as relevant in this case, the times of the tests are equal or even lower than the original, 11.3, 9.71, 12.92 seconds respectively.

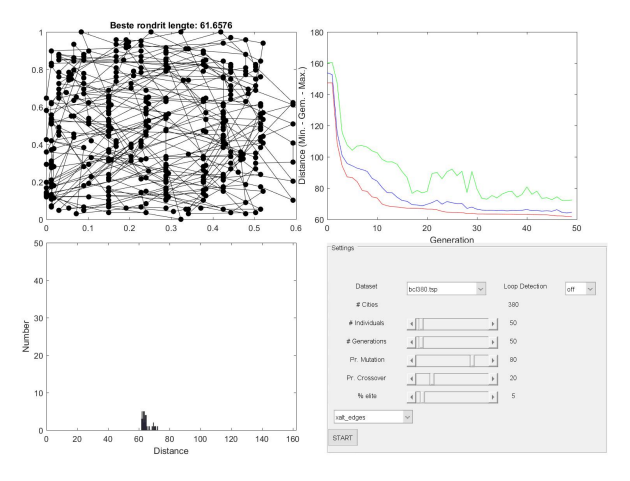
Finally, the combination of the good results is what made them get outstanding results. In order to avoid bias, the first combination is just modifying the number of individuals, and the number of generations

[Nind, gens, elitism, crossover, mutation] = [200,200,5%,95%,5%]



We can see an improvement, even slightly better than those obtained with the increase in number of individuals or generations alone. The number of generations was choosen to be 200 because we deemed it good enough, and while costly, not as costly as higher quantities.

[Nind, gens, elitism, crossover, mutation] = [200,200,5%,50%,50%]

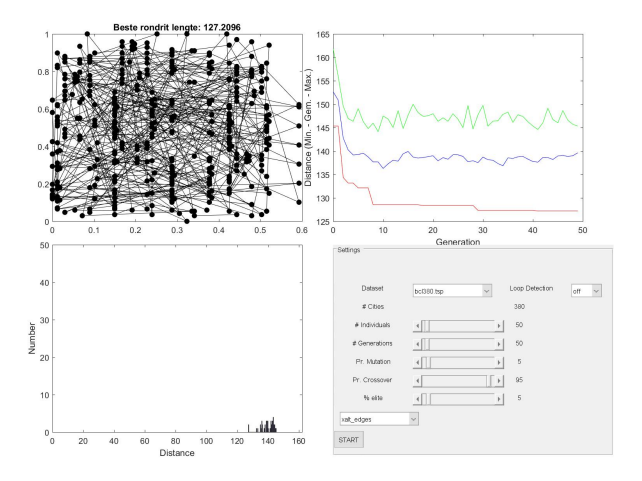


And the final result, when combining every single improvement, is the best of all. The decrease of distance is 30.8%, which could be said to be a great decrease. The time spent on this test was 76.74 seconds, around 6.1 times more than the original, although high, we consider that it is not sufficiently high as to consider it bad.

### 2.2.2 Path representation

The same structure as the previous representation will be followed. One parameter at a time will be tested, stating the change, after performing a base test

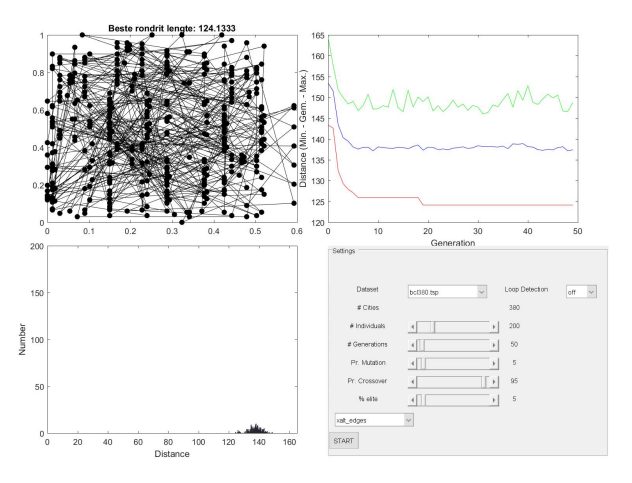
[Nind, gens, elitism, crossover, mutation] = [50,50,5%,95%,5%]



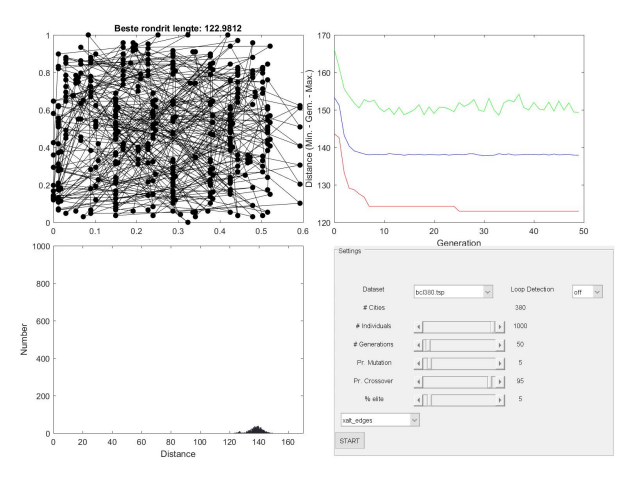
The first noticeable thing that we can observe here is that the distance is far higher than the adjacency representation, although looking at the time, it is better, since it takes only 6.88 seconds.

We have the same situation that we had with the number of generations in the previous representation, but this time with the number of individuals. While 200 was the amount with the generally lower distances, higher numbers also performed somewhat ok. Thus we tested for 200, 1000, and 1250.

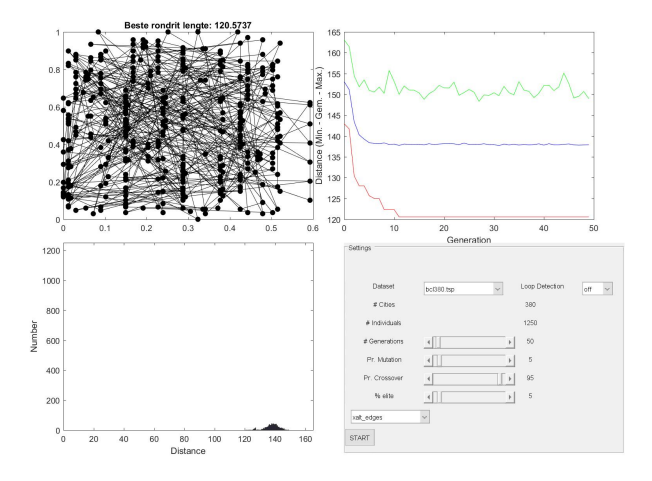
[Nind, gens, elitism, crossover, mutation] =[200,50,5%,95%,5%]



[Nind, gens, elitism, crossover, mutation] = [1000,50,50,5%,95%,5%]



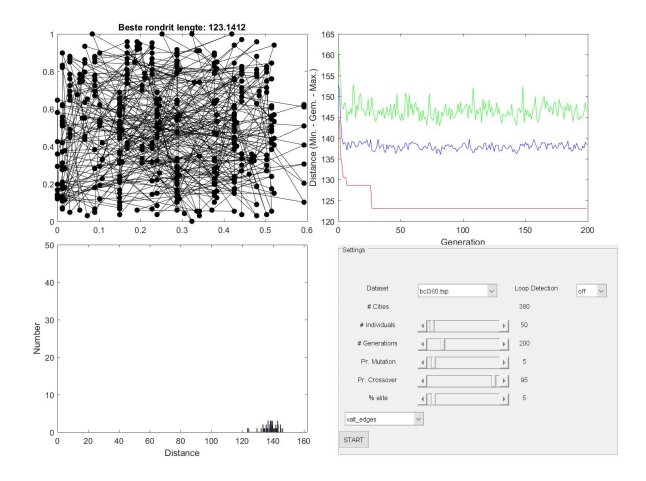
[Nind, gens, elitism, crossover, mutation] = [1250,50,5%,95%,5%]



Although the change is not big, there is improvement, with each increase of individuals. And, what's more, as it is individuals, and not generations that we are increasing now, there is higher cost, but not as high. The times for the tests are 9.08, 29.44, and 37.99 seconds, being the last not even 6 times higher. As there is the increase is the highest in the last test (1250 individuals), and the time is not humongously high, that value will be the one to be tested in the combined test.

The general test for the number of generations revealed that the best quantity was 200.

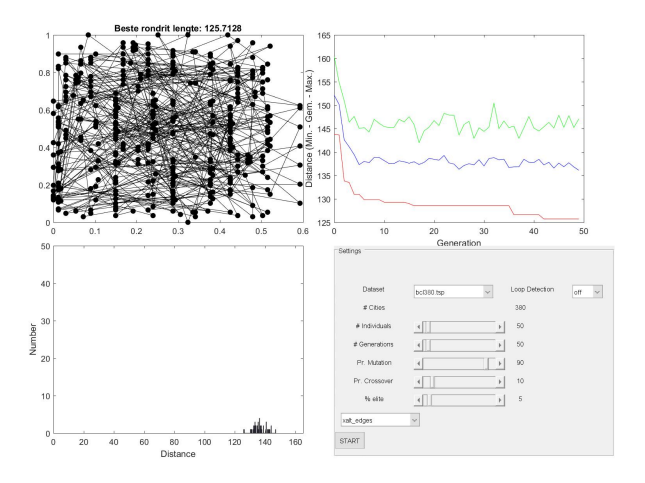
[Nind, gens, elitism, crossover, mutation] =[50,200,5%,95%,5%]



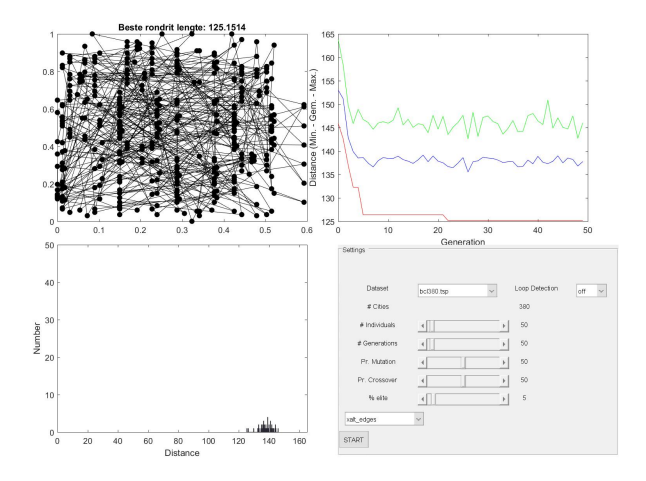
We can observe a slight improvement against the base case, 4 units of distance. Timewise, it is much more demanding that increasing individuals, since the test took 23.73 seconds to end, 3.5 times more than the base case. Any higher number of generations will only hinder the process by making it tediously slow.

As happened in the previous representation tests, the general test for path rep. showed that the value 5% for elitism was without a doubt the best that could be set. So, the last to test is crossover and mutation, and here comes the tricky part for this representation. We found in our general test that no matter what the percentage for mutation or crossover was being tested, the results were almost independant from them. So, given that the base case already tested for a high crossover, and low mutation, the designed tests are, for low crossover and high mutation, and for a 50—50.

[Nind, gens, elitism, crossover, mutation] = [50,200,5%,50%,50%]



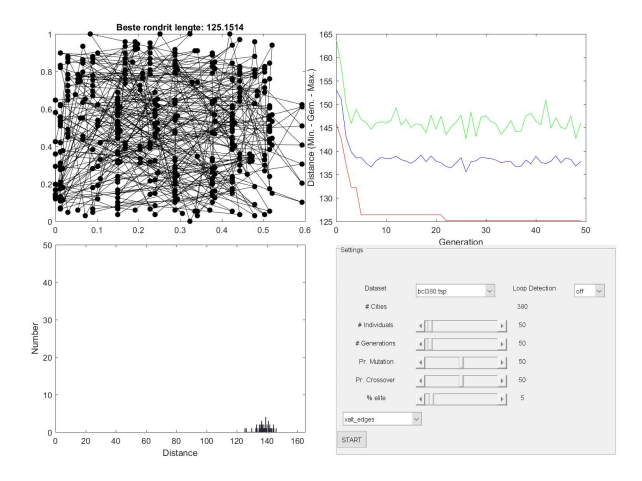
[Nind, gens, elitism, crossover, mutation] =[50,200,5%,50%,50%]



Although somewhat better than the base case, the result from one test to the other differ so little that it confirms the result of the general test, stating that, for this represention, given our implementation with our operators, the percentage is not really relevant. But, as it's still better than the base, and the best of the two, the percentages chosen for the combined test are 50%—50%. Obviosly, as the number of indviduals or generations did not change, the time is almost identical to the base case

For the combined test, in hope to get the best result, this is the test made

[Nind, gens, elitism, crossover, mutation] = [1250,200,5%,50%,50%]



Unlike the previous combined test, the big expected improvement is nowhere to be seen. Going from a length of 127 to 120 is just a mere 5.2% of decrease. Furthermore, the combined increase of individuals and generations, given that the former is very high, makes the test to take extremly long compared to the base, going from 6.88 seconds to 141.02 seconds, 20.5 times more. TODO: Añadir posibles causas de que no mejore, y mencionar cuál puede ser la razón de que no cambie prácticamente nada cuando se modifica mutacion y crossover

### References

- [1] A.E. Eiben, James E Smith. Introduction to Evolutionary Computing. Springer
- [2] P. Larrañaga, C.M.H. Kuijpers, R.H. Murga, I. Inza, S. Dizdarevic Genetic algorithms for the travelling salesman problem: A review of representations and operators. University of the Basque Country
- [3] Dirk Roose Genetic algorithms lectures and slides KU Leuven
- [4] mnemstudio.org Genetic algorithms mutations http://mnemstudio.org/genetic-algorithmsmutation.htm

### 3 Apendix

### 3.1 tsp\_ImprovePopulation.m

```
1 % tsp_ImprovePopulation.m
2 % Author: Mike Matton
4 % This function improves a tsp population by removing local loops
     from
5 % each individual.
6 %
7 | Syntax: improvedPopulation = tsp_ImprovePopulation(popsize,
     ncities, pop, improve, dists)
 1%
9 | % Input parameters:
10 %
      popsize
                          - The population size
      ncities
                          - the number of cities
_{12} \%
      pop
                          - the current population (adjacency
     representation)
_{13} \%
                          - Improve the population (0 = no improvement)
      improve
     <>0 = improvement)
 1%
      dists
                          - distance matrix with distances between the
      cities
_{15} \%
16 \% Output parameter:
      improvedPopulation - the new population after loop removal (
     if improve
18 %
                               \Leftrightarrow 0, else the unchanged population).
 function newpop = tsp_ImprovePopulation(popsize, ncities, pop,
     improve, dists)
21
  if (improve)
22
     for i=1:popsize
23
24
       result = improve_path(ncities, pop(i,:), dists);
25
26
       pop(i,:) = path2adj(result);
27
     end
29
 end
30
_{32} newpop = pop;
```

### $3.2 \quad run\_ga.m$

```
function run_ga(x, y, NIND, MAXGEN, NVAR, ELITIST, STOP_PERCENTAGE
     , PR_CROSS, PR_MUT, CROSSOVER, LOCALLOOP, ah1, ah2, ah3)
2 % usage: run_ga(x, y,
3 %
                   NIND, MAXGEN, NVAR,
                   ELITIST, STOP_PERCENTAGE,
5 %
                   PR_CROSS, PR_MUT, CROSSOVER,
                   ah1, ah2, ah3)
9 \% x, y: coordinates of the cities
10 % NIND: number of individuals
11 % MAXGEN: maximal number of generations
12 % ELITIST: percentage of elite population
13 % STOP PERCENTAGE: percentage of equal fitness (stop criterium)
14 % PR_CROSS: probability for crossover
15 \% PRMUT: probability for mutation
16 % CROSSOVER: the crossover operator
17/% calculate distance matrix between each pair of cities
18 % ah1, ah2, ah3: axes handles to visualise tsp
 {NIND MAXGEN NVAR ELITIST STOP_PERCENTAGE PR_CROSS PR_MUT
    CROSSOVER LOCALLOOP \;
20
          tic;
21
          GGAP = 1 - ELITIST;
          mean_fits = zeros(1, MAXGEN+1);
          worst=zeros(1,MAXGEN+1);
2.4
          Dist=zeros (NVAR, NVAR);
25
          for i=1:size(x,1)
26
               for j=1:size(y,1)
                   Dist(i,j)=\mathbf{sqrt}((x(i)-x(j))^2+(y(i)-y(j))^2);
28
              end
          end
30
          % initialize population
          Chrom=zeros (NIND, NVAR);
32
          for row=1:NIND
            %Chrom(row,:)=path2adj(randperm(NVAR));
34
              Chrom(row,:) = randperm(NVAR);
35
          end
36
          gen=0;
          % number of individuals of equal fitness needed to stop
          stopN=ceil(STOP_PERCENTAGE*NIND);
39
          % evaluate initial population
          ObjV = tspfun (Chrom, Dist);
41
```

```
best=zeros(1,MAXGEN);
42
          % generational loop
43
          while gen<MAXGEN
44
              sObjV=sort (ObjV);
45
               best (gen+1)=min(ObjV);
            minimum = best (gen + 1);
47
               mean_fits(gen+1)=mean(ObjV);
               worst(gen+1)=max(ObjV);
49
               for t=1:size(ObjV,1)
50
                   if (ObjV(t)=minimum)
                       break;
                   end
              end
              %visualizeTSP(x,y,adj2path(Chrom(t,:)), minimum, ah1,
                  gen, best, mean_fits, worst, ah2, ObjV, NIND, ah3);
               visualizeTSP(x,y,Chrom(t,:), minimum, ah1, gen, best,
                  mean_fits, worst, ah2, ObjV, NIND, ah3);
58
               if (sObjV(stopN)-sObjV(1) \le 1e-15)
                     break;
              end
61
            %assign fitness values to entire population
62
            FitnV=ranking (ObjV);
            %select individuals for breeding
64
            SelCh=select ('sus', Chrom, FitnV, GGAP);
            %recombine individuals (crossover)
66
               SelCh = recombin (CROSSOVER, SelCh, PR_CROSS);
67
              %SelCh=mutateTSP('inversion', SelCh, PR_MUT);
68
               SelCh=mutateTSP('insertion', SelCh, PR_MUT); % <-- line
69
                  changed, now insertion mutation is used
              %evaluate offspring, call objective function
70
            ObjVSel = tspfun(SelCh, Dist);
71
              %reinsert offspring into population
             [Chrom, ObjV] = reins (Chrom, SelCh, 1, 1, ObjV, ObjVSel);
73
74
              Chrom = tsp_ImprovePopulation(NIND, NVAR, Chrom,
                 LOCALLOOP, Dist);
            %increment generation counter
            gen=gen+1;
          end
          toc;
79
          minimum
 end
```

### 3.3 insertion.m

```
1 % low level function for TSP mutation
2 \% Representation is an integer specifying which encoding is used
_{3}|\% 1 : adjacency representation
_{4} % 2 : path representation
  function NewChrom = insertion (OldChrom)
       NewChrom = OldChrom;
       % select two positions in the tour
       rndi = zeros(1,2);
       while \operatorname{rndi}(1) = \operatorname{rndi}(2)
            rndi=rand_int(1,2,[1 \text{ size}(NewChrom},2)]);
       end
       rndi = sort(rndi);
      % get the value of the first random position
17
       temp = NewChrom(rndi(1));
1.8
       % insert this value in the second random position
       NewChrom = insertAt (NewChrom, temp, rndi(2));
20
      % remove the first random position
21
       NewChrom(rndi(1)) = [];
2.5
      % End of function
23
  end
24
  function arrOut = insertAt(arr, val, index)
26
       if index = numel(arr)+1
27
            arrOut = [arr val];
28
       else
            \operatorname{arrOut} = [\operatorname{arr}(1:\operatorname{index}-1) \ \operatorname{val} \ \operatorname{arr}(\operatorname{index}:\operatorname{end})];
30
       end
32 end
```

#### 3.4 order crossover.m

```
% Syntax: NewChrom = order_crossover(OldChrom, XOVR)
% Input parameters:
% OldChrom - Matrix containing the chromosomes of the old population. Each line corresponds to one individual
% (in any form, not necessarily real values).
```

```
7 %
       XOVR
                  - Probability of recombination occurring between
     pairs
8 %
                     of individuals.
% Output parameter:
 %
       NewChrom
                  - Matrix containing the chromosomes of the
     population
_{12} %
                     after mating, ready to be mutated and/or
     evaluated,
 %
                     in the same format as OldChrom.
_{14} %
  function NewChrom = order_crossover(OldChrom, XOVR)
18 if nargin < 2, XOVR = NaN; end
  [rows, \tilde{z}] = size(OldChrom);
2.0
     maxrows=rows;
21
     if rem(rows, 2)^{\sim}=0
       maxrows = maxrows - 1;
     end
24
     for row = 1:2: maxrows
26
      % crossover of the two chromosomes
28
      % results in 2 offsprings
    if rand<XOVR
                        % recombine with a given probability
30
           MatrixChrom = order_low_level([OldChrom(row,:);OldChrom(
31
              row + 1 ...);
      NewChrom(row,:) = MatrixChrom(1,:);
32
      NewChrom(row+1,:) = MatrixChrom(2,:);
33
    else
34
      NewChrom(row,:) = OldChrom(row,:);
35
      NewChrom (row + 1,:) = OldChrom (row + 1,:);
36
    end
37
     end
38
39
     if rem(rows, 2) = 0
40
       NewChrom(rows,:)=OldChrom(rows,:);
     end
42
44 % End of function
```

### 3.5 order\_low\_level.m

```
1 % low level function for calculating an offspring
2 % given 2 parent in the Parents - agrument
3 Parents is a matrix with 2 rows, each row
4 % represent the genocode of the parent
6 % Returns a matrix containing the offspring
 function Offspring=order_low_level(Parents)
10
      cols = size(Parents, 2);
11
      Offspring=zeros(2, cols);
13
      start_index = rand_int(1, 1, [1, cols - 1]);
15
      end\_index = rand\_int(1, 1, [start\_index + 1, cols]);
16
17
      Offspring(1, start_index:end_index) = Parents(2, start_index:
18
         end_index);
      Offspring(2, start_index:end_index) = Parents(1, start_index:
         end_index);
20
21
      for off = 1:2
          Buff = Parents (off,:);
23
          Buff = [Buff(end\_index + 1:end), Buff(1:end\_index)];
2.4
25
          members = ismember(Buff, Offspring(off, :));
26
          Buff (members == 1) = 0;
28
          ii = 1;
          X = find(Buff);
30
          for jj=1:start\_index - 1
              if Buff (X(ii)) = 0
32
                 Offspring (off, jj) = Buff (X(ii));
                 Buff(X(ii)) = 0;
34
                 ii = mod(ii, cols) + 1;
35
             end
36
          end
37
          ii = 1;
39
          X = find(Buff);
          for jj=end_index + 1:cols
```

```
if Buff(X(ii)) \tilde{}=0
42
                  Offspring (off, jj) = Buff(X(ii));
43
                  Buff(X(ii)) = 0;
44
                  ii = mod(ii, cols) + 1;
45
              end
           end
47
          %Offspring(off, end_index+1:end) = Buff(start_index:end);
          \%Offspring(off, 1:start\_index - 1) = Buff(1:start\_index - 1)
49
              1);
      end
 |\%| end function
```

### 3.6 tspgui.m

```
CROSSOVER = 'order_crossover';

crossover = uicontrol(ph, 'Style', 'popupmenu', 'String', {'
order_crossover'}, 'Value', 1, 'Position', [10 50 130 20], '
Callback', @crossover_Callback);
```

### 3.7 tspfun.m

```
1 %
2 % ObjVal = tspfun (Phen, Dist)
3 % Implementation of the TSP fitness function
4 % Phen contains the phenocode of the matrix coded in path
5 % representation
6 % Dist is the matrix with precalculated distances between each
    pair of cities
7 % ObjVal is a vector with the fitness values for each candidate
    tour
8 %
     (=each row of Phen)
9 %
 function ObjVal = tspfun(Phen, Dist)
11
     % the objective function works with adjacency representation.
         In this
     % version, path representation is used, so the fitness
13
         function should
     \% be adapted. Now, the phenotype is converted to adjacency
     % representation first, and then, the Objective Value is
        computed as it
     % was computed in the original version.
16
      adj = zeros(size(Phen));
```

```
for row=1:size(Phen)
    adj(row,:) = path2adj(Phen(row,:));
end

ObjVal = Dist(adj(:,1), 1);
for t=2:size(adj,2)
    ObjVal=ObjVal + Dist(adj(:,t), t);
end

End

End

End

End

For row=1:size(Phen)
    adj(row,:) = path2adj(Phen(row,:));
end
```

### 3.8 mutateTSP.m

```
1 MUTATETSP.M
                       (MUTATion for TSP high-level function)
2 %
3 % This function takes a matrix OldChrom containing the
4 % representation of the individuals in the current population,
_{5} mutates the individuals and returns the resulting population.
7 Syntax: NewChrom = mutate(MUT.F, OldChrom, MutOpt)
8 %
9 % Input parameter:
       MUT<sub>F</sub>
                  - String containing the name of the mutation
     function
11 %
       OldChrom - Matrix containing the chromosomes of the old
12 %
                    population. Each line corresponds to one
     individual.
13 %
       MutOpt
                  - mutation rate
14 % Output parameter:
_{15} \%
       NewChrom - Matrix containing the chromosomes of the
     population
 %
                    after mutation in the same format as OldChrom.
16
17
 function NewChrom = mutateTSP(MUT_F, OldChrom, MutOpt)
19
20
 % Check parameter consistency
21
     if nargin < 2, error ('Not_enough_input_parameters'); end
22
  [rows, \tilde{z}] = size(OldChrom);
 NewChrom=OldChrom;
25
 for r=1:rows
27
    if rand<MutOpt
```

```
NewChrom(r,:) = feval(MUT.F, OldChrom(r,:));

end

end

% End of function
```