# **TSP implementation - How-To**

### Requirements and Dependencies

- Python 3.5
- Numpy
- matplotlib
- scipy (for the plot generation and statistical analysis scripts)

## **Usage**

#### Configuration

The EA configuration parameters are defined in JSON files (see default\_args.json for an example).

Here are the key/value pairs that can be specified in the JSON config file:

- datafile The path to the data file, e.g. data/TSP\_Uruguay\_734.txt. Type: string
- popsize Population size. Type: integer
- initialize\_method Use random or k-means clustering for initialisation. Can be set to random or k-means. Type: string
- mp\_size Mating pool size. Must be a multiple of 2. Type: integer.
- tournament size Tournament size for tournament parent selection. Type: integer
- recombination The recombination/crossover operator used. Can be set to best\_order or cut\_crossfill. Type string
- mutation The mutation operator used. Can be set to cyclic, inversion, insertion, scramble, two\_opt, or permutation\_swap. Type: string
- crossover\_rate Crossover rate, a value between 0.0 and 1.0. Type: float
- mutation\_rate Mutation rate, a value between 0.0 and 1.0. Type: float
- generations The number of generations to run the EA for. Type: integer
- box\_cutting\_points\_n The number of crossover points for best order crossover. This parameter must always be larger than or equal to 5 and must always be less than the chromosome length minus 1.
   Type: integer
- kca\_k A value between 0.0 and 1.0, which acts as a proportion of the chromosome length to define
  the number of clusters for the k-means algorithm. Type: float
- kca\_iterations The number of iterations to refine the k-means clusters. Type: integer

**Note:** Make sure you use double quotes (") when editing the JSON file's key/value pairs. Also, make sure there is no trailing comma at the end of the JSON file. To check whether your config is valid JSON, you may run it through <a href="https://jsonformatter.curiousconcept.com/">https://jsonformatter.curiousconcept.com/</a>, which will report any JSON syntax errors (if present) in your config file.

### Running the algorithm

Here are the arguments that the main.py script takes:

- --args-file or -a: Used to specify a custom JSON config file. Default is default\_args.json. Example
   python3 main.py --args-file custom\_args.json
- --test-runs or -t: Number of times to run the algorithm, usually used in conjunction with the ---export argument. Default is 1. Example python3 main.py --test-runs 10

- --export or -e: Export the resulting best fitness values to a CSV file. Usually used in conjunction with the --test-runs argument to export the data for multiple runs of the algorithm to a CSV file. This option is how we generated the data for statistical comparisons and visualisations used in our report.
   Example python3 main.py --export
- --visualize or -v: Toggle the real-time visualisation. **Note** that you cannot use this option when you're using the --test-runs option. **Example** python3 main.py --visualize
- --debug or -d: Print performance and debugging output. Example python3 main.py --debug
- --help or -h: Print the help text. Example python3 main.py --help

Here are some examples of a few different ways you can run the script:

The following command uses a custom configuration file, runs the algorithm for 10 runs, and exports the data to CSV files:

```
python3 main.py --args-file custom_args.json --test-runs 10 --export
```

The following command uses a custom configuration file, turns on the real-time visualisation, and also prints the performance and debugging output:

```
python3 main.py --args-file custom_args.json --visualize --debug
```

### **Data Analysis**

We have also provided all the data used to conduct the statistical tests and generate the visualisations that were used in the report. All the relevant files are found under testing/:

- **testing/mutation/** contains all the data files, the plots and the script related to the different mutation parameters.
- **testing/recombination/** contains all the data files, the plots and the script related to the different recombination/crossover operators.
- **testing/parameters/** contains all the data file, the plots and the script related to the different tuning parameters for the evolutionary algorithm.

For example, in order to conduct the statistical tests and generate the visualisations for all the mutation parameters:

- 1. Go to the **testing/mutation/** directory.
- 2. Run: python3 analyze.py

Running the above command should print the results of the statistical tests for the various mutation parameters and should generate image files for the visualisations in the same directory (testing/mutation/).

You can perform the same steps for the **testing/recombination/** and **testing/parameters/** directories too.

