Programming Exercise 2


Algorithm description: Let \((K_n, c)\) be given, and consider a vector \(\lambda \in \mathbb{R}^n\). Define edge costs \(c_\lambda\) by
\[
c_\lambda(\{i, j\}) := c(\{i, j\}) + \lambda(i) + \lambda(j).
\]
Given a 1-tree \(T\), set
\[
c(T, \lambda) := c(T) + \sum_{v \in V(T)} (|\delta_T(v)| - 2) \cdot \lambda(v).
\]
Now, let \(T_\lambda\) be a min-\(c_\lambda\)-cost 1-tree (which also minimizes \(c(T_\lambda, \lambda)\)), and set
\[
HK(K_n, c, \lambda) := c(T_\lambda, \lambda).
\]
Then \(HK(K_n, c, \lambda)\) is a lower bound on the minimum weight of any tour in \((K_n, c)\), so \(HK(K_n, c) := \max_{\lambda \in \mathbb{R}^n} HK(K_n, c, \lambda)\) is also a lower bound on the minimum weight of a tour. In particular, if \(T_\lambda\) is 2-regular, it is an optimum tour. The idea of the algorithm is to use Branch-and-Bound to compute a 2-regular 1-tree of minimum cost.

It has been shown in the lecture that we can compute a vector \(\lambda\) that approximately maximizes \(HK(K_n, c, \lambda)\) using an iterative combinatorial algorithm, c.f. below.

In the Branch-and-Bound algorithm, we maintain an upper bound \(U\) on the cost of a shortest tour, and a set \(Q\) of branching nodes to be processed. Each node in the Branch-and-Bound tree is represented by two disjoint sets of edges \((R, F)\) which are initially empty, i.e., start with \(Q = \{()\}\). The node \((R, F)\) represents all tours in \(K_n\) where all edges in \(R\) are required, and all edges in \(F\) are forbidden. Clearly the node \((\emptyset, \emptyset)\) represents all tours in \(K_n\).

Now, while \(Q\) is not empty, select a node \((R, F) \in Q\) and set \(Q := Q \setminus \{(R, F)\}\). Compute \(\lambda\) s.t. the weight of a min-\(c_\lambda\)-cost 1-tree \(T\) with \(R \subseteq E(T) \subseteq E(K_n) \setminus F\) is approximately maximum and let \(T\) be such a 1-tree.

If \(HK(K_n, c, R, F) := c(T, \lambda) \geq U\), we know that the node \((R, F)\) does not represent a solution leading to better cost than \(U\), so we can discard \((R, F)\).

So assume \(c(T, \lambda) < U\). If \(T\) is 2-regular, it is an optimum tour represented by \((R, F)\), so we can update \(U\). Otherwise, there needs to be a vertex \(2 \leq i \leq n\) with \(|\delta_T(i)| > 2\). We can assume that \(|\delta_T(i) \setminus (R \cup F)| \geq 2\), because whenever two edges incident to a vertex \(v\) are required, we can forbid all other edges incident to \(v\). Let
$e_1, e_2 \in \delta_T(i) \setminus (R \cup F)$ be two distinct edges in $T$ incident to $i$ that we have not yet branched on. Partition $(R, F)$ into the three branching nodes

$$(R, F \cup \{e_1\}), (R \cup \{e_1\}, F \cup \{e_2\}), (R \cup \{e_1, e_2\}, F),$$

where the last node is omitted if there is already a required edge incident to $i$, and add these nodes to $Q$.

When $Q$ is empty, we know that $U$ equals the cost of a shortest tour in $(K_n, c)$, and we can return the tour that led to the current value of $U$.

**Usage:** Your program should be called as follows:

```
program_name --instance file.tsp [--solution file.opt.tour]
```

**Input:** The first argument, `file.tsp`, is mandatory (i.e., your program should exit with an error message if it is not present), and it encodes the graph $(K_n, c)$ for which your program should find a shortest tour. It is expected to be in TSPLIB format. The second argument is optional, and if present, an optimum solution should be written to the specified file. The TSPLIB format allows to encode arbitrary instances, however, we restrict to instances with rounded Euclidean costs. In the following, one possible way to parse Euclidean TSPLIB files is sketched.

First, normalize the input by removing all colons (`:`) and replacing consecutive spaces by a single one. Then, ignore all lines until a line of the form `DIMENSION n` is found, where $n$ is an integer specifying the number of vertices in the instance. Then, ignore all lines until a line of the form `NODE_COORD_SECTION` is found. The following $n$ lines specify the coordinates of the vertices. Each line has the form

```
i  x  y
```

where $1 \leq i \leq n$ is the vertex whose coordinates are specified, and $x$ and $y$ are floating point representations of the coordinates of $i$. Finally, the file ends with `EOF`.
Cost function: The cost of an edge between two points is its rounded Euclidean length, where we round to the nearest integer. The following C++ function computes the distance of two points \((x_1, y_1)\) and \((x_2, y_2)\):

```cpp
int distance(double x1, double y1, double x2, double y2) {
    return std::lround(std::sqrt((x1-x2)*(x1-x2) + (y1-y2)*(y1-y2)));
}
```

Output: Your program should write the cost of a shortest tour in \((K_n, c)\) to stdout. No other output may be written to stdout / std::cout. If you wish to output anything in addition (e.g. the current lower bound, upper bound, the number of branching nodes, runtime, etc.), use stderr / std::cerr. Furthermore, if the optional argument --solution file.opt.tour is specified, your program should write an encoding of the found optimum solution to the specified file in TSPLIB format. A tour is encoded as follows in TSPLIB format. The encoding starts with a header of the form:

```
TYPE : TOUR
DIMENSION : n
TOUR_SECTION
```

where \(n\) is the number of vertices in the instance. Now, \(n\) lines follow, each consisting of a single integer \(i\) with \(1 \leq i \leq n\). These lines specify the order in which the vertices are visited in an optimum tour. Finally, the encoding ends with these two lines:

```
-1
EOF
```
Implementation details: In [1], an effective implementation of the algorithm is described, which may be helpful. It is recommended to follow [1] as close as possible for at least the lower bound computation. They also describe heuristics to find better values of $U$ during the algorithm, which you do not need to implement. Also, you may start with $U = \infty$, although using a heuristic to determine a better value of $U$ before the algorithm may lead to better results.

Branching implementation: During branching, there are several choices to be made that will influence the runtime of the algorithm. Most importantly, it is not specified how to select the next element $(R, F) \in Q$ to be processed. The two most prominent choices are best-bound and depth-first. Best-bound always selects a node $(R, F)$ minimizing $HK(K_n, c, R, F)$, i.e., $Q$ is a priority queue. This approach processes a minimum amount of nodes, but may take a long time to find good values of $U$.

Depth-first always processes an element of $Q$ of maximum depth in the branching tree, i.e., $Q$ is a stack. This approach tends to quickly find feasible solutions and allows to delay the evaluation of $HK(K_n, c, R, F)$, but may enumerate more branching nodes. We recommend to use best-bound. Other choices to be made are the selection of the vertex $i$ to branch on, and the selection of the edges $e_1$ and $e_2$.

Lower bound implementation: In the lecture, it has been shown that we can find a vector $\lambda$ that approximately maximizes $HK(K_n, c, \lambda)$ by starting with $\lambda_0 \equiv 0$ and repeating the following step. In iteration $i \geq 0$, compute a min-$c_{\lambda_i}$-weight 1-tree $T_i$ and set

$$\lambda_{i+1}(x) := \lambda_i(x) + t_i \cdot (|\delta_{T_i}(x)| - 2),$$

where the sequence $t_i$ converges to 0 and satisfies $\sum_{i=0}^{\infty} t_i = \infty$, e.g., $t_i := \frac{1}{i+1}$.

In practice, a good choice of the step lengths $t_i$ is crucial for the algorithm to compute a good approximation within a bounded number of iterations. In particular, the step lengths of course should depend on the costs $c$. In [1], it is proposed to fix the number of iterations $N$ and the initial step length $t_0$ in advance, and then letting the $t_i$ converge to 0 (e.g., $t_N = 0$) with a constant second order difference

$$(t_i - t_{i+1}) - (t_{i+1} - t_{i+2}) \equiv \text{const}$$

under the (arbitrary) condition

$$(t_0 - t_1) = 3(t_{N-1} - t_N).$$
This can be achieved by defining $\Delta_0 = \frac{3n_0}{2N}$ and $\Delta \Delta = \frac{ln_0}{N^2 - N}$, and, for $0 \leq i \leq N - 1$,
\[
t_{i+1} = t_i - \Delta_i, \\
\Delta_{i+1} = \Delta_i - \Delta \Delta.
\]

It remains to choose $N$ and $t_0$. Since the computations for branching nodes $(R, F)$ that are not the root of the branching tree (i.e., $\emptyset, \emptyset$) can (and should!) start with the vector $\lambda$ of their parent, it is reasonable to use more iterations at the root node. Hence, [1] propose to use $N = \lceil \frac{n^2}{50} \rceil + n + 15$ at the root node and $N = \lceil \frac{n}{2} \rceil + 5$ at other nodes. To let $\lambda$ scale with $c$, at the root node set $t_0 = \frac{c(T_0)}{2n}$, where $T_0$ is the 1-tree computed in the first iteration, i.e., a minimum cost 1-tree of $(K_n, c)$. At other nodes use $t_0 = \frac{1}{2n} \sum_{i=1}^{n} |\lambda_{\text{root}}(i)|$, where $\lambda_{\text{root}}$ is the vector $\lambda$ computed at the root node.

Furthermore, Volgenant and Jonker [1] observed oscillating degrees during the iterations. To dampen these oscillations, for $i \geq 1$, they replace (1) by
\[
\lambda_{i+1}(x) = \lambda_i(x) + t_i \cdot \left( d \cdot (|\delta_{T_i}(x)| - 2) + (1 - d) \cdot (|\delta_{T_{i-1}}(x)| - 2) \right)
\]
with $d = 0.6$.

A minimum weight 1-tree can be obtained by starting with a MST on $\{2, \ldots, n\}$ and adding the two cheapest edges incident to vertex 1. This approach can be extended to computing a minimum weight $(R, F)$-1-tree, i.e., a 1-tree $T$ of minimum weight s.t. $R \subseteq E(T) \subseteq E(K_n) \setminus F$: Artificially set the costs of all edges in $F$ to $\infty$ and add all edges in $R$ first.

Finally, note that since these computations are done using floating point arithmetic, one must take care of not computing false lower bounds $L > OPT(K_n, c)$, which can be achieved by multiplying with $1 - \epsilon$ for a small value $\epsilon > 0$. Moreover, since all costs are integral, we can round up lower bounds.

Programming language: Your program should be written in C or C++, although the use of C++ is strongly encouraged. By default, your program will be compiled using clang-4.0.0 using C++14. Different compilers or compiler versions are available upon request. Your program will be compiled using \texttt{-pedantic -Wall -Wextra -Werror}, i.e., all warnings are enabled and each remaining warning will lead to compilation failure. Program evaluation will be performed on Linux. The standard library can be used as you wish, but no other libraries.

Algorithm evaluation: Make sure to implement core steps of the algorithm as fast as possible. Compute min-weight 1-trees in $O(n^2)$ time. Your algorithm should be able to solve each of the instances \texttt{berlin52.tsp, st70.tsp, eil176.tsp, rat99.tsp, rd100.tsp, eil101.tsp} and \texttt{lin105.tsp} in at most a few seconds.
**Code evaluation:** The elegance, cleanness and organization of your code will be evaluated. Make sure to add good documentation and give the variables, functions and types meaningful names that make their role clear. Break your complicated functions into small simple ones, break your program into a few units etc. Of course, your program may not trigger undefined behavior. In particular, your program must be **valgrind-clean**, i.e., must not leak memory and must not perform invalid operations on memory. Follow the guidelines presented in the first exercise class, which are still available on the website.

**Help:** On the website for the exercise classes, which you should visit regularly, you will find a precise definition of the TSPLIB format and a few instances with known optima that you can use to test your program.

**References**


Please submit your programs in groups of 2 students. (64 points)

**Deadline:** January 18th 2018, 14:15, via email to silvanus@or.uni-bonn.de. The websites for the lecture with all exercises and test instances can be found at:

[http://www.or.uni-bonn.de/lectures/ws17/co_exercises/exercises.html](http://www.or.uni-bonn.de/lectures/ws17/co_exercises/exercises.html)