



Improved heuristics for the regenerator location problem

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Abstract

Telecommunication systems use optical signals to transmit information. The strength of a signal in an optical network deteriorates and loses power as it goes farther from the source, mainly due to attenuation. Therefore, to enable the signal to arrive its intended destination with good quality, it is necessary to regenerate the signal periodically using regenerators. These components are relatively expensive and therefore it is desirable to deploy as few of them as possible in the network. In the regenerator location problem (RLP), we are given an undirected graph, positive edge lengths, and a parameter specifying the maximum length that a signal can travel before its quality deteriorates and regeneration is required. The problem consists in determining paths that connect all pairs of nodes in the graph and, if necessary, locating single regenerators in some of those nodes such that the signal never travels more than the maximum allowed distance without traversing a regenerator node. In this paper, we present new implementations of previous heuristics and two new heuristics—a GRASP and a biased random-key genetic algorithm—for the RLP. Computational experiments comparing the proposed solution procedures with previous heuristics described in the literature illustrate the efficiency and effectiveness of our methods.

Keywords: metaheuristics; network design; regenerator; location; GRASP; genetic algorithms

1. Introduction

Telecommunication systems use optical signals to transmit information. The strength of an optical signal deteriorates and loses power as it goes farther from the source, mainly due to attenuation. Therefore, to enable the signal to arrive its intended destination with good quality, it is necessary

to regenerate the signal periodically using regenerators. These components are relatively expensive and therefore it is desirable to deploy as few of them as possible in the network.

In the “regenerator location problem” (RLP), we are given an undirected graph $G = (V, E)$, where V is the node set, E is the set of edges, with each edge $(i, j) \in E$ having a real-valued length $d_{i,j} \in \mathbb{R}^+$. Furthermore, a parameter $D \in \mathbb{R}^+$ is given to specify the maximum length that a signal can travel before its quality deteriorates and regeneration is required. The problem consists in determining paths that connect all pairs of nodes in the graph and, if necessary, locating single regenerators in some of those nodes. Between each pair of nodes $\{s, t\} \in V \times V$, a “path” $\{(s, v_1), (v_1, v_2), \dots, (v_k, t)\}$ connecting these nodes is formed by one or more path segments. In this context, a “path segment” consists of a sequence of consecutive edges $\{(v_i, v_{i+1}), (v_{i+1}, v_{i+2}), \dots, (v_{q-1}, v_q)\}$ in the path, satisfying the condition that the “total length” of the segment

$$d_{v_i, v_{i+1}} + d_{v_{i+1}, v_{i+2}} + \dots + d_{v_{q-1}, v_q} \leq D. \quad (1)$$

If the total length of the path is not more than D , then the path consists of a single path segment. Otherwise, it will consist of two or more path segments. In this case, one or more regenerators will be located in the internal nodes of the path, separating consecutive path segments. The objective of the RLP is to determine paths that connect all pairs of nodes in the graph utilizing a minimum number of regenerators.

Figure 1a shows an example of a network with seven nodes and a maximum distance parameter $D = 100$. The numbers near the edges represent their lengths. Note that the length $d_{1,5} = 150$ of edge $(1, 5)$ is greater than D and therefore it cannot be part of any path. In this figure, we can see that the shortest path from node 1 to node 3 is $\{(1, 2), (2, 3)\}$ with a total length of $60 + 70 = 130 > 100$. Therefore, it must be divided into two path segments $\{(1, 2)\}$ and $\{(2, 3)\}$ and a regenerator must be placed in node 2 to connect node 1 with node 3 using this path. The shortest path connecting nodes 1 and 5 is the single edge path $\{(1, 5)\}$ with length 150. Since edge $(1, 5)$ is longer than D , then it cannot be used in any path. Therefore, to connect node 1 with node 5 we identify the shortest feasible path to be $\{(1, 2), (2, 3), (3, 5)\}$ with total length $60 + 70 + 90 = 220$. This path must be divided into three path segments, which requires two regenerators—one at node 2 and one at node 3. On the other hand, we can connect node 5 with node 7 using the shortest path $\{(5, 6), (6, 7)\}$ with a total length of $40 + 40 = 80$ and therefore we do not need to allocate any regenerator in this case. Finally, note that placing regenerators in nodes 2 and 7 allows communication between all pairs of nodes in the graph.

Regenerator placement in the context of traffic engineering with restoration was first discussed in Yetginer and Karasan (2003). Gouveia et al. (2003) address a multiprotocol label switching (MPLS) over wave division multiplexing (WDM) network design problem in which one type of constraint (called WDM path constraint) forbids path segments between two components that are longer than a given maximum length.

Chen and Raghavan (2007) and Chen et al. (2010) introduce the RLP and present a branch-and-cut procedure for the Steiner arborescence problem with a unit degree constraint on the root node, which is shown to be equivalent to the RLP. Furthermore, they introduce three heuristic procedures for this problem. Computational experiments using 740 test problems show that the heuristics obtain the optimal solutions in 454 instances while the branch-and-cut method finds the optimal solution in 536 instances. The same authors generalize the problem by defining a set of

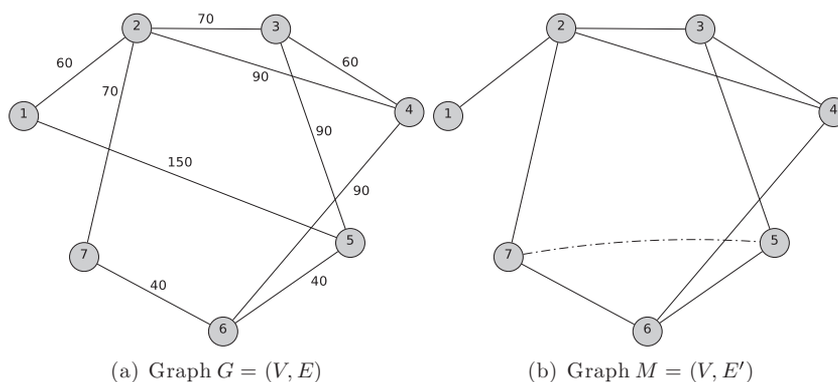


Fig. 1. Example of a seven-node graph G with edge lengths and corresponding communication graph M .

nodes where regenerators can be placed and a set of nodes that need to be connected (Chen et al., 2009). The RLP was shown to be NP-hard by Flammini et al. (2009) and Chen et al. (2010).

Chen et al. (2010) describe the “communication graph” that they use in their algorithms for the RLP. Given the weighted graph $G = (V, E)$, we first delete all the edges that have length greater than D . We add an edge between all nonadjacent pairs of nodes having length equal to the length of the corresponding shortest path if the edge length in G is smaller than D . Finally, we disregard all length information. The resulting unweighted graph is referred as $M = (V, E')$. If the resulting graph is complete, then there is no need for any regenerator. If the resulting graph is not connected, then the problem is infeasible. Alternatively, if the resulting graph is connected but not complete, then one or more regenerators will be required.

Figure 1b represents the communication graph M that results from the graph in the example of Fig. 1a. The dashed link in the graph between nodes 5 and 7 is the only link added during the procedure to create M from G .

The remainder of this paper is organized as follows. In Section 2, we propose new and efficient implementations of the three constructive algorithms originally proposed in Chen et al. (2010). In Section 3, we point out an error in the description of a local search proposed in Chen et al. (2010). We then combine randomized variants of three greedy algorithms proposed in Chen et al. (2010) with a correct version of their local search, resulting in three multistart greedy randomized adaptive search procedures (GRASP). A biased random-key genetic algorithm (BRKGA) for the RLP is proposed in Section 4. Computational experiments comparing the proposed GRASP and BRKGA with the heuristics of Chen et al. (2010) are described in Section 5. Concluding remarks are provided in Section 6.

2. Constructive heuristics

In this section, we use several constructive heuristics proposed in Chen et al. (2010) for the RLP: a greedy algorithm, and two heuristics called H1 and H2. We propose efficient implementations of these procedures that, as will be shown in our computational experience, outperform their original implementations by up to three orders of magnitude.

Table 1

$\mathcal{X}(u)$ and $g(u)$ for all candidate nodes u in first iteration of greedy algorithm on example of Fig. 1b

u	$\mathcal{X}(u)$	$g(u)$
1	\emptyset	0
2	$\{(1, 3), (1, 4), (1, 7), (3, 7), (4, 7)\}$	5
3	$\{(4, 5), (2, 5)\}$	2
4	$\{(2, 6), (3, 6)\}$	2
5	$\{(3, 7), (3, 6)\}$	2
6	$\{(4, 7), (4, 5)\}$	2
7	$\{(2, 6), (2, 5)\}$	2

2.1. Greedy algorithm

The greedy algorithm takes as input the set of pairs of nodes in the communication graph that are not directly connected (NDC) and builds a set R of regenerator nodes, one node at a time. At each iteration, the procedure computes, for each node u , $g(u)$, the number of yet-unconnected pairs in the communication graph that get connected with their inclusion in R . The method then determines a node u^* with maximum g -value. Node u^* is added to R and the communication graph is updated by adding to it the edges connecting those yet-unconnected pairs that get connected by placing a regenerator in u^* .

In our implementation of this method, we store for each vertex u , the set of its neighbors, $\mathcal{N}(u)$, in a hash table. Therefore, basic operations, such as adding, removing, or checking the membership of an element in $\mathcal{N}(u)$ can be practically performed in constant time. For example, to calculate $g(u)$ we need to compute the Cartesian product of $\mathcal{N}(u)$ in the communication graph, which is extremely fast with the hash table. Moreover, while computing $g(u)$ for each vertex u , we store the best so far, so that we do not have to go through them again to determine u^* , as apparently done in the original implementation of Chen et al. (2010).

Algorithm 1 shows a pseudocode of our adaptation of the greedy algorithm. It takes as input the communication graph $M = (V, E')$ and gives the set of regenerator nodes R as output. In line 1, the set R of regenerator nodes is initialized empty and the set C of candidate regenerator nodes is initialized with all of the nodes in the graph. In line 2, the set \bar{E}' of NDC pairs is initialized with the complement of the set E' of edges in M . The main loop of the greedy algorithm goes from lines 3 to 20. It is repeated until M is complete, that is, while $\bar{E}' \neq \emptyset$. For each candidate node $u \in C$, lines 4–14 compute the set $\mathcal{X}(u)$ of yet-unconnected pairs in M that would become connected if node u were to house a regenerator. Let $g(u)$ denote the cardinality of set $\mathcal{X}(u)$. In line 5, $\mathcal{X}(u)$ is initialized empty and in line 6, $g(u)$ is initialized to zero. Line 7 computes the set $\mathcal{N}(u)$ of nodes adjacent to u in M . For all pairs of nodes, both of which are in $\mathcal{N}(u)$, lines 8–13 verify if they are connected in M and if they are not connected we add them to $\mathcal{X}(u)$. In line 15, the node u^* that enables the connection of the largest number of yet-unconnected pairs in M is selected. Ties are broken at random. In lines 15–18, sets R , C , E' , and \bar{E}' are updated to reflect the inclusion of node u^* in set R . In line 24, the set R of regenerator nodes is returned.

Algorithm 1. Pseudocode for GREEDY: Greedy algorithm for the RLP

```

procedure Greedy
  Data: Communication graph:  $M = (V, E')$ 
  Result: Set of regenerator nodes:  $R \subseteq V$ 
  1  $R \leftarrow \emptyset; C \leftarrow V;$ 
  2  $\bar{E}' = \{(i, j) \in V \times V : (i, j) \notin E'\};$ 
  3 while  $\bar{E}' \neq \emptyset$  do
  4   for  $u \in C$  do
  5      $\mathcal{X}(u) \leftarrow \emptyset;$ 
  6      $g(u) \leftarrow 0;$ 
  7      $\mathcal{N}(u) = \{v \in V : (u, v) \in E'\};$ 
  8     for  $(i, j) \in \mathcal{N}(u) \times \mathcal{N}(u)$  do
  9       if  $(i, j) \notin E'$  then
 10          $\mathcal{X}(u) \leftarrow \mathcal{X}(u) \cup \{(i, j)\};$ 
 11          $g(u) \leftarrow g(u) + 1;$ 
 12       end
 13     end
 14   end
 15    $u^* \leftarrow \arg \max\{g(u) : u \in C\};$ 
 16    $R \leftarrow R \cup \{u^*\};$ 
 17    $C \leftarrow C \setminus \{u^*\};$ 
 18    $E' \leftarrow E' \cup \mathcal{X}(u^*);$ 
 19    $\bar{E}' \leftarrow \bar{E}' \setminus \mathcal{X}(u^*);$ 
 20 end
 21 return  $R;$ 

```

Consider as an example the communication graph depicted in Fig. 1. For each candidate u , Table 1 shows $\mathcal{X}(u)$, the set of pairs of nodes not connected that get connected if a regenerator is placed in node u and $g(u)$, which is the cardinality of $\mathcal{X}(u)$.

In iteration 1 of the greedy algorithm, node $u^* = 2$ is selected, since $g(2) = 5$ is the maximum $g(u)$ for all $u \in C$. Edges (1, 3), (1, 4), (1, 7), (3, 7), and (4, 7) are added to graph M resulting in the graph shown in Fig. 2 a. Similarly, in iteration 2, node $u^* = 7$ is selected and edges (1, 5), (1, 6), (2, 5), (3, 6), (4, 5), and (2, 6) are added to graph M making it complete, as can be seen in Fig. 2b. The set R of regeneration nodes obtained with this greedy algorithm is therefore $\{2, 7\}$.

2.2. H1 heuristic

The aim of heuristic H1 is to find a spanning tree on the communication graph M with the maximum number of leaf nodes. This way, the heuristic aims to minimize the number of internal nodes in the spanning tree. Regenerators are assigned to these internal nodes. The algorithm starts by computing $u^* \leftarrow \arg \min\{\deg(u) : u \in V\}$, a node of minimum degree (ties are broken by favoring the node with smallest index) and initializes the set of nodes in the spanning tree, $S \leftarrow \{u^*\}$, and its complement, $\bar{S} = V \setminus \{u^*\}$. The set R of regenerators is initialized empty, that is, $R \leftarrow \emptyset$. H1 then calls the recursive function, $\text{Tree}(u^*, S, \bar{S}, R)$, described in Algorithm 2.

Algorithm 2. Pseudocode for Tree used in H1

```

procedure Tree
  Data:  $u, S, \bar{S}, R$ 
  Result:  $R$ 
  1  $\mathcal{U}(u) \leftarrow \mathcal{N}(u) \cap \bar{S}$ ;
  2 if  $|S| > 1$  then
  3    $S \leftarrow S \cup \mathcal{U}(u)$ ;
  4    $\bar{S} \leftarrow V \setminus S$ ;
  5 end
  6 while  $\mathcal{U}(u) \neq \emptyset$  do
  7   Compute  $\deg_{\bar{S}}(v)$  for all  $v \in \mathcal{U}(u)$ ;
  8    $u^* \leftarrow \arg \max\{\deg_{\bar{S}}(v) : v \in \mathcal{U}(u)\}$ ;
  9   if  $\deg_{\bar{S}}(u^*) > 0$  then
 10     $R \leftarrow R \cup \{u^*\}$ ;
 11    Tree( $u^*, S, \bar{S}, R$ );
 12  end
 13   $\mathcal{U}(u) \leftarrow \mathcal{U}(u) \setminus \{u^*\}$ ;
 14 end
 15 return

```

Tree takes as input the vertex u with the lowest degree of the graph and recursively constructs a spanning tree. In step 3 of Algorithm 2, the spanning tree is augmented with the unspanned nodes that are neighbors of node u . Note that in the previous call to the procedure, a regenerator is placed in node u^* in step 10.

Let us illustrate the logic behind this procedure by using an example. Considering again the communication graph shown in Fig. 1, H1 selects vertex 1 as u^* since this vertex is the one with minimum degree. Therefore $S = \{1\}$, $\bar{S} = \{2, 3, 4, 5, 6, 7\}$, and $R = \emptyset$. With these four input arguments H1 calls the recursive procedure Tree. In the first iteration, it computes the unexplored neighbors of vertex 1 (i.e., $\mathcal{U}(1) = \mathcal{N}(1) \cap \bar{S} = \{2\} \cap \{2, 3, 4, 5, 6, 7\} = \{2\}$). It is important to remark that since Tree is

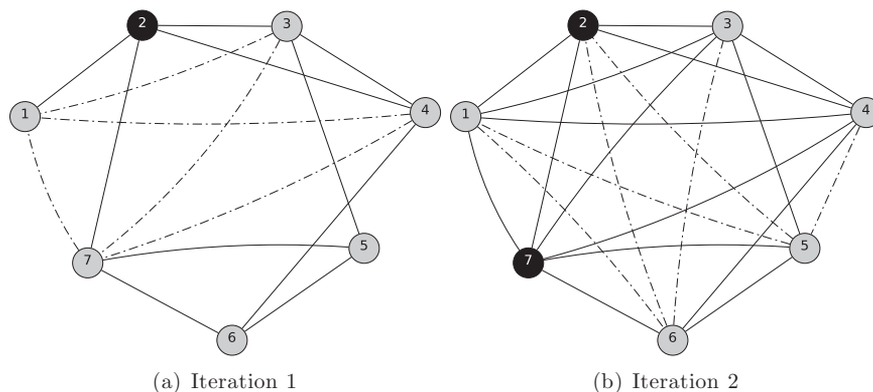


Fig. 2. Communication graph on two iterations of the greedy algorithm.

recursive, then in the first iteration, steps 2 to 5 are not executed. The procedure goes directly to step 6. There is only one vertex in $\mathcal{U}(1) = 2$ with $\deg_{\bar{S}}(2) = 3$. Notice that $\deg_{\bar{S}}$ is computed over the set \bar{S} . The degree of vertex 2 is then 3 instead of 4. Before procedure `Tree` is called again, the set of regenerators is updated (i.e., $R = \{2\}$).

In the second iteration, the procedure again computes the set of unexplored neighbors (i.e., $\mathcal{U}(2) = \mathcal{N}(2) \cap \bar{S} = \{1, 3, 4, 7\} \cap \{2, 3, 4, 5, 6, 7\} = \{3, 4, 7\}$). As opposed to the first iteration, in this iteration (and in all subsequent iterations) steps 2 to 5 are executed. Therefore, $S = S \cup \mathcal{U}(2) = \{1\} \cup \{3, 4, 7\} = \{1, 3, 4, 7\}$ and $\bar{S} = V \setminus S = \{2, 5, 6\}$. The procedure then computes the degree in \bar{S} of all the vertices in $\mathcal{U}(2)$. Thus, $\deg_{\bar{S}}(3) = 2$, $\deg_{\bar{S}}(4) = 2$, and $\deg_{\bar{S}}(7) = 3$, selecting vertex 7 as u^* since it presents the maximum value for $\deg_{\bar{S}}$. The procedure updates the set of regenerators ($R = \{2, 7\}$) and calls again procedure `Tree`.

In the third iteration, $\mathcal{U}(7) = \mathcal{N}(7) \cap \bar{S} = \{2, 5, 6\} \cap \{2, 5, 6\} = \{2, 5, 6\}$, $S = S \cup \mathcal{U}(7) = \{1, 2, 3, 7\} \cup \{2, 5, 6\} = \{1, 2, 3, 4, 5, 6, 7\}$, and $\bar{S} = \emptyset$. In this situation, the degree with respect to \bar{S} is always zero (since $\bar{S} = \emptyset$), then neither in this iteration nor in the subsequent iterations is the set of regenerators updated. At the end, the recursive procedure returns $R = \{2, 7\}$.

In our implementation of H1 we use a hash set to store the set of unvisited vertices instead of using the stack of the system. Additionally, our algorithm is iterative instead of recursive, as seems to be the intent of the original implementation of Chen et al. (2010). Finally, as we did in `Greedy`, the $\mathcal{N}(v)$ set is indexed with a hash table and therefore its associated operations can be performed in $O(1)$.

2.3. H2 heuristic

Heuristic H2 in some sense resembles the two previous methods. In each step, it first identifies the nodes of lowest degree and then examines the neighbors of these nodes and selects the one with the highest degree to assign a regenerator. Once a regenerator is assigned to a node, the communication graph is updated by adding edges with endpoints in newly connected pairs of nodes. H2 resembles `Greedy` in the sense that it updates the communication graph at each step and it resembles H1 in the sense that it is based on the degree of the nodes.

Contrasting our implementation of H2 with the original implementation in Chen et al. (2010), it is worth mentioning that our Java code employs hash-sets for an efficient update of the communication graph and the computation of the lowest and highest degree nodes. In each iteration, we compute the set of newly connected nodes by calculating the Cartesian product of the set of nodes adjacent to the node just made to house a regenerator. As in the previous methods, the use of hash-sets significantly accelerates this computation. In our experiments (see Section 5), we empirically compare both implementations.

3. GRASP for the RLP

A GRASP, or greedy randomized adaptive search procedure, is a multistart or iterative process in which each iteration consists of two phases: construction and local search (Feo and Resende, 1989, 1995; Festa and Resende, 2009a, b; Ribeiro et al., 2013; Moran-Mirabal et al., 2014). The

construction phase builds a feasible solution, whose neighborhood is explored until a local optimum is found after the application of the local search phase. At each iteration of the construction phase, GRASP maintains a set of candidate elements CL (candidate list) that can be feasibly added to the partial solution under construction. Every candidate element is evaluated according to a greedy function in order to select the next element to be added to the partial solution. A restricted CL (RCL) is created with the best elements in CL . This is the greedy aspect of the method. The element to be added to the partial solution is randomly selected from those in the RCL . This is the probabilistic aspect of the heuristic. Once the selected element is added to the partial solution, the CL is updated and its elements are evaluated. This is the adaptive aspect of the heuristic. We refer the reader to Resende and Ribeiro (2003, 2010) for two recent reviews of GRASP.

3.1. Construction procedures

We implemented three constructive algorithms, CG , $C1$, and $C2$ based on, respectively, the heuristics Greedy, H1, and H2 of Chen et al. (2010), described in Section 2.

Given the communication graph M , the CG constructive method builds a set R of regenerator nodes, one node at a time. At each iteration, CG first determines the set CL of candidate nodes to house a regenerator, which is simply the set of nodes in M in which no regenerator has been previously placed. Then, for each candidate node $u \in CL$, it computes the set $\mathcal{X}(u)$ of yet-unconnected pairs in M that would become connected if node u were to house a regenerator. Let $g(u)$ denote the cardinality of set $\mathcal{X}(u)$.

The RCL is formed with those nodes having a relatively good g -value. To this end, the method computes g_{min} and g_{max} as, respectively, the minimum and maximum values of g in CL . The algorithm next constructs an RCL with all the nodes in CL having a g -value greater than or equal to a specified cutoff value

$$g_{min} + \alpha \cdot (g_{max} - g_{min}),$$

where

$$g_{min} = \min_{u \in CL} g(u) \quad \text{and} \quad g_{max} = \max_{u \in CL} g(u).$$

CG randomly selects an element u^* in RCL , adds it to the set R , and updates the communication graph (by adding to it the edges connecting those yet-unconnected pairs that get connected by placing a regenerator in u^*). The construction ends when the graph M is complete. We test different values of α in the computational experiments in Section 5.

Heuristic H1 tries to find a spanning tree S on the communication graph M having the maximum number of leaf nodes, thus minimizing the number of internal nodes in S to which regenerators are assigned. At each iteration, the set CL of candidates nodes on which to place a regenerator, referred to as $\mathcal{U}(u)$ in Algorithm 2, is computed as $\mathcal{N}(u) \cap \bar{S}$, where \bar{S} is the complement of the spanning tree S under construction, and u is the node in which a regenerator was placed in the previous iteration. H1 selects the node $u^* \in CL$ with maximum degree $\deg_{\bar{S}}(u^*)$ with respect to \bar{S} and includes it in S . In the randomized construction $C1$, instead of selecting the node $u^* \in CL$ with

maximum degree, we compute the RCL with all the nodes in CL having a degree in \bar{S} greater than or equal to a specified threshold value

$$deg_{min} + \alpha \cdot (deg_{max} - deg_{min}), \quad (2)$$

where

$$deg_{min} = \min_{u \in CL} deg_{\bar{S}}(u) \quad \text{and} \quad deg_{max} = \max_{u \in CL} deg_{\bar{S}}(u).$$

Then, $C1$ randomly selects a node in RCL and adds it, together with its adjacent vertices, to the spanning tree S under construction.

Finally, constructive method $C2$ determines at each iteration the set CL of candidate nodes to house a regenerator as the set of nodes in M in which no regenerator has been previously placed. Then, for each candidate node $u \in CL$, it computes its degree $d(u)$ in M . Instead of selecting the node with the largest degree (as H2 does), $C2$ computes a RCL with those nodes having degree greater than or equal to a specified threshold value, as in (2), and randomly selects a node $u^* \in RCL$. The method adds u^* to the set R and updates the communication graph by adding to it the edges connecting those yet-unconnected pairs that get connected by placing a regenerator in u^* .

Algorithm 3. Pseudocode for MOVE: a 2-1 or 1-0 move.

procedure Move

Data: Communication graph $M = (V, E')$, regenerator set R ,
nodes i and j

Result: New regenerator set R'

```

1  $R' \leftarrow R \setminus \{i, j\}$ ;
2  $C \leftarrow V \setminus R'$ ;
3 for  $u \in (\mathcal{N}(i) \cup \mathcal{N}(j)) \setminus R'$  do
4   if  $\mathcal{N}(u) \cap R' = \emptyset$  then
5      $C \leftarrow C \cup \mathcal{N}(u)$ ;
6   end
7   if  $C = \emptyset$  then
8     return nil;
9   end
10 end
11 Select  $v \in C$  producing a feasible solution;
12  $R' \leftarrow R' \cup \{v\}$ ;
13 return  $R'$ ;

```

3.2. Local search

Given two nodes i and j in the set R of regenerator nodes, procedure $\text{Move}(i, j)$, shown in Algorithm 3, attempts to remove both regenerators from nodes i and j and replace them with a single regenerator (that eventually could be placed in i or j). A set C of potential candidates to host a regenerator is initialized in step 2. In steps 3–10, each non-regenerator node u adjacent to either i or

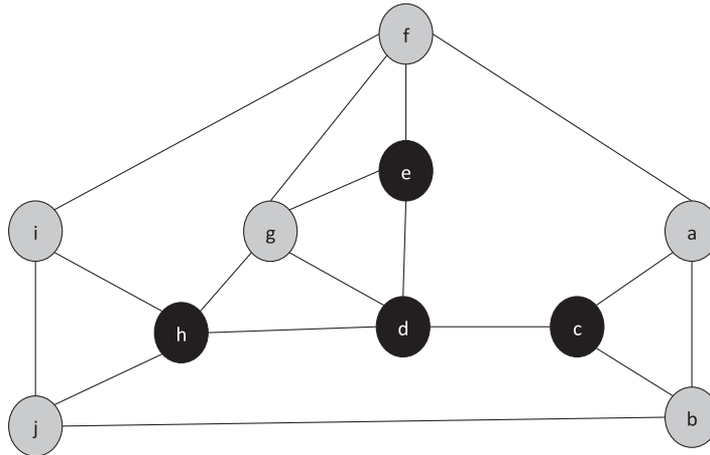


Fig. 3. Example where original MOVE fails.

j is analyzed. If u is not adjacent to an existing regenerator node (distinct from i and j), then in step 5 the set of potential candidate nodes is updated, eliminating from C the nodes not adjacent to u . If C is empty, then nodes i and j cannot be replaced in R with fewer than two nodes and the procedure returns a `nil` indicator. Otherwise, any element in the final set C can be selected to replace i and j . Note that i , j , or both i and j can be in C . In step 11, a node v is selected from C and is added to R' in step 12. A 1-0 move results if either i or j is selected. Otherwise, a 2-1 move is applied. Ties are broken by a node index. The node in C with the largest degree in the communication graph is selected.

Consider the example in Fig. 3 with the solution $R = \{c, d, e, h\}$ (depicted in black in the figure) and apply the MOVE procedure in Algorithm 3 to nodes d and e . In steps 3–9 we take each vertex $u \in (\mathcal{N}(d) \cup \mathcal{N}(e)) \setminus R = \{f, g\}$ and compute C , obtaining $C = \{a, g, i\}$. According to the original design of the MOVE procedure in Chen et al. (2010), we could replace the regenerators in d and e with a single regenerator allocated in any node in C . However, we can see that, in this case, this would produce an infeasible solution. For example, if we select a , we would obtain $R' = \{c, h, a\}$ leaving node e disconnected. To solve this problem, we have modified this method to include a condition on the feasibility of the resulting solutions (step 11 in Algorithm 3). Specifically, we check if the subgraph induced by the set of regenerators is connected. If it is, then the solution is feasible; otherwise, we discard this selection and resort to the next node in C . In the example in Fig. 3, none of the candidates in C produces a feasible solution.

Our local search procedure scans the regenerator nodes in the order given by their degrees in the original graph (where the node with the lowest degree is examined first). Nodes with low degree are expected to be good candidates to move their regenerator to an alternative node. The procedure then applies a first improvement strategy, performing the first associated move in its exploration. The local search finishes when there is no further improvement and returns the local optimum found.

4. BRKGA

Genetic algorithms with random keys, or “random-key genetic algorithms” (RKGAs), were first introduced by Bean (1994) for solving combinatorial optimization problems involving sequencing. In an RKGA, chromosomes are represented as vectors of randomly generated real numbers in the interval $[0, 1]$. A deterministic algorithm, called a “decoder,” takes as input a solution vector and associates with it a solution of the combinatorial optimization problem for which an objective value or fitness can be computed.

An RKGA evolves a population of random-key vectors over a number of iterations, called “generations”. The initial population is made up of p vectors of random keys. Each component of the solution vector is generated independently at random in the real interval $[0, 1]$. After the fitness of each individual is computed by the decoder in generation k , the population is partitioned into two groups of individuals: a small group of p_e “elite” individuals, that is, those with the best fitness values, and the remaining set of $p - p_e$ “non-elite” individuals. To evolve the population, a new generation of individuals must be produced. All elite individual of the population of generation k are copied without modification to the population of generation $k + 1$. RKGAs implement mutation by introducing “mutants” into the population. A mutant is simply a vector of random keys generated in the same way that an element of the initial population is generated. At each generation, a small number (p_m) of mutants is introduced into the population. With the p_e elite individuals and the p_m mutants accounted for in population $k + 1$, $p - p_e - p_m$ additional individuals need to be produced to complete the p individuals that make up the new population. This is done by producing $p - p_e - p_m$ offspring through the process of mating or crossover.

Bean (1994) selects two parents at random from the entire population to implement mating in an RKGA. A BRKGA (Gonçalves and Resende, 2011, 2014), differs from a RKGA in the way parents are selected for mating. In a BRKGA, each element is generated combining one element selected at random from the elite partition in the current population and one from the non-elite partition. Repetition in the selection of a mate is allowed and therefore an individual can produce more than one offspring in the same generation. “Parameterized uniform crossover” (Spears and DeJong, 1991) is used to implement mating in BRKGAs. Let $\rho_e > 0.5$ be the probability that an offspring inherits the vector component of its elite parent. Let n denote the number of components in the solution vector of an individual. For $i = 1, \dots, n$, the i th component $c(i)$ of the offspring vector c takes on the value of the i th component $e(i)$ of the elite parent e with probability ρ_e and the value of the i th component $\bar{e}(i)$ of the non-elite parent \bar{e} with probability $1 - \rho_e$.

When the next population is complete, that is, when it has p individuals, fitness values are computed for all of the newly created random-key vectors and the population is partitioned into elite and non-elite individuals to start a new generation.

A BRKGA searches the solution space of the combinatorial optimization problem indirectly by searching the continuous n -dimensional hypercube, using the decoder to map solutions in the hypercube to solutions in the solution space of the combinatorial optimization problem where the fitness is evaluated.

We propose a BRKGA heuristic for the RLP. As in the previous algorithms, we denote vertex and edge sets of the communication graph M as V and E' , respectively. The heuristic encodes the solution as a $|V|$ -vector x of random keys. The i th element of x corresponds to node $i \in V$. To decode x , we apply the greedy algorithm `Decoder` shown in Algorithm 4.

Algorithm 4. Pseudocode for DECODER.

```

procedure Decoder
  Data: Communication graph  $M = (V, E')$  and vector  $x$  of
           random keys
  Result: Set of regenerator nodes:  $R \subseteq V$ 
  1  $R \leftarrow \emptyset; C \leftarrow V;$ 
  2  $\bar{E}' = \{(i, j) \in V \times V : (i, j) \notin E'\};$ 
  3 Order the nodes in  $C$  w.r.t. their associated  $x$  value;
  4 while  $\bar{E}' \neq \emptyset$  do
  5   Select the next node  $k \in C$  following the  $x$ -order;
  6    $\mathcal{X}(k) \leftarrow \emptyset;$ 
  7    $A \leftarrow \mathcal{N}(k) = \{v \in V : (k, v) \in E'\};$ 
  8   for  $v \in A \cap R$  do
  9      $A \leftarrow A \cup \mathcal{N}(v);$ 
 10  end
 11  for  $(i, j) \in A \times A$  do
 12    if  $(i, j) \notin E'$  then
 13       $\mathcal{X}(k) \leftarrow \mathcal{X}(k) \cup \{(i, j)\};$ 
 14    end
 15  end
 16   $R \leftarrow R \cup \{k\};$ 
 17   $C \leftarrow C \setminus \{k\};$ 
 18   $E' \leftarrow E' \cup \mathcal{X}(k);$ 
 19   $\bar{E}' \leftarrow \bar{E}' \setminus \mathcal{X}(k);$ 
 20 end
 21 return  $R;$ 

```

In the initialization of the Decoder algorithm (see Algorithm 4), we order the nodes in C , initially containing all nodes in V , according to the random key vector x . In this way, nodes associated with components in x with a relative large value (close to 1) come first. The main loop of the Decoder algorithm goes from lines 4 to 20. It is repeated until M is complete, that is, while $\bar{E}' = \emptyset$. In this loop nodes are selected following the order induced by x . For each candidate node $k \in C$, lines 6–15 compute the set $\mathcal{X}(k)$ of yet unconnected pairs in M that would get connected if node k were to house a regenerator. In lines 16–19, sets R , C , E' , and \bar{E}' are updated to reflect the inclusion of node k in set R . In line 21, the set R of regenerator nodes is returned.

5. Experimental results

This section describes the computational experiments that we have performed to first test the efficiency of our different procedures and then compared them with the method in Chen et al. (2010). We have implemented the methods in Java SE 6 and all the experiments were conducted on a Pentium 4 computer at 3 GHz with 2 GB of RAM.

In our computational experiments we used the instances, which the authors kindly made available to us, from Chen et al. (2010). Specifically, these graphs come from two different sources—the

Table 2
Comparison with Chen et al. (2010) implementation

n	m	Orig. Greedy		Greedy		Orig. H1		H1		Orig. H2		H2	
		NR	RT	NR	RT	NR	RT	NR	RT	NR	RT	NR	RT
40	10	1.40	0.00	1.40	0.00	1.40	0.00	1.40	0.00	1.40	0.10	1.40	0.00
	30	2.00	0.20	2.00	0.01	2.00	0.10	2.00	0.00	2.00	0.00	2.00	0.00
	50	3.00	0.10	3.00	0.01	3.10	0.30	3.00	0.00	3.30	0.10	3.00	0.00
	70	4.60	0.70	4.50	0.01	4.70	0.50	4.60	0.00	4.60	0.20	4.40	0.00
	90	9.50	1.70	10.10	0.02	9.70	1.30	9.90	0.01	9.80	0.50	10.20	0.00
60	10	1.90	0.30	1.90	0.04	1.90	0.30	1.90	0.01	1.90	0.00	1.90	0.00
	30	2.10	0.80	2.00	0.03	2.10	0.30	2.00	0.01	2.20	0.20	2.00	0.00
	50	3.20	1.90	3.10	0.05	3.30	1.10	3.10	0.00	3.40	0.40	3.00	0.00
	70	5.30	4.30	5.10	0.06	5.40	3.90	5.10	0.00	5.30	0.70	5.00	0.01
	90	10.80	15.80	11.00	0.10	11.20	18.00	11.40	0.01	11.20	2.50	11.30	0.01
80	10	1.90	1.30	1.90	0.06	1.90	0.70	1.90	0.03	1.90	0.10	1.90	0.00
	30	2.90	3.90	2.60	0.09	2.80	2.00	2.70	0.02	2.80	0.30	2.70	0.01
	50	3.80	9.10	3.50	0.08	4.00	5.60	3.60	0.01	3.70	0.90	3.50	0.00
	70	6.00	19.40	5.80	0.13	5.90	17.30	5.70	0.00	5.80	3.00	5.70	0.01
	90	11.90	88.90	12.20	0.22	12.50	70.70	12.40	0.03	12.80	13.00	12.10	0.03
100	10	2.00	3.60	2.00	0.14	2.00	1.80	2.00	0.07	2.00	0.50	2.00	0.01
	30	2.90	9.60	2.70	0.17	2.90	7.10	2.80	0.03	2.80	1.40	2.80	0.01
	50	4.00	21.10	4.00	0.20	4.00	13.60	4.00	0.02	4.00	2.60	4.00	0.01
	70	6.00	49.00	6.00	0.25	6.40	56.00	5.80	0.01	6.10	8.20	6.00	0.01
	90	13.40	286.40	12.70	0.38	13.70	295.00	13.50	0.04	13.30	42.90	13.00	0.02
	Avg.	4.93	25.91	4.88	0.10	5.05	24.78	4.94	0.02	5.02	3.88	4.90	0.01

M -graph instances, in which they directly generated the communication graph; and random instances, in which they first generated a graph with random edge lengths and then computed the communication graph. The authors sent us a total of 280 instances, 200 from the first set and 80 from the second one, with $n = 40, 60, 80,$ and 100 .

In the first experiment, we compare the original implementation by Chen et al. (2010) of the heuristics Greedy, H1, and H2 coupled with the local search (labeled as Orig. Greedy, Orig. H1, and Orig. H2, respectively, in Table 2) with our new implementations of these methods (simply labeled as Greedy, H1, and H2). For each method and each instance size, we consider, in Table 2, the two columns reported in Table 1 of Chen et al. (2010): NR (number of regenerators) and RT (running time). Results in this table clearly indicate that our implementations outperform those in Chen et al. (2010) since they obtain slightly better solutions (lower number of regenerators) in significant lower running times (more than 100 times faster). The methods in Chen et al. (2010) were run on a Pentium D (3.0 GHz processor with 2 GB RAM) while, as described above, our implementations have been run on a similar computer (Pentium 4, 3 GHz with 2 GB of RAM). Note also that our implementations are in Java while those in Chen et al. (2010) were in C++.

In each of the following experiments, we compute for each instance the overall best solution value, *Best Value*, obtained by all executions of the methods considered. Then, for each method, we compute the relative percentage deviation between the best solution value obtained with that method and *Best Value* for that instance. We report the average of this relative percentage deviation

Table 3
GRASP constructive methods

	Value	#Best	Dev (%)	CPU (seconds)
CG				
$\alpha = 0.3$	7.00	7	25.77	12.93
$\alpha = 0.6$	5.65	13	5.92	10.00
$\alpha = 0.9$	5.35	19	0.45	7.96
C1				
$\alpha = 0.3$	6.80	8	20.42	0.77
$\alpha = 0.6$	6.10	10	10.79	0.74
$\alpha = 0.9$	6.25	11	12.10	0.76
C2				
$\alpha = 0.3$	6.30	7	16.59	0.57
$\alpha = 0.6$	6.10	11	10.64	0.56
$\alpha = 0.9$	5.95	12	8.51	0.54

(Dev) across all the instances considered in each particular experiment. We finally report, for each method, the number of instances (#Best) in which the value of the best solution obtained with this method matches *Best Value*.

In our preliminary experimentation we consider a set of 20 instances randomly selected from those instances in our set with $n = 80, 100$. In the first preliminary experiment, we study the parameter α in constructive methods CG, C1, and C2. Specifically, we test three values of this parameter on each method: 0.3, 0.6, and 0.9. We run CG, C1, and C2 100 times, thus obtaining 100 solutions for each method and instance pair. Table 3 reports, for this set of 20 instances and each value of α tested, the values of #Best and Dev. We also report the average objective function value (number of regenerators), Value, and the CPU time in seconds.

The results in Table 3 show that the best outcomes are obtained when the constructive method CG is run with a value of $\alpha = 0.9$. However, it consumes longer running times than its competing GRASP constructions C1 and C2, although the three of them are very fast.

In the second preliminary experiment, we compare the constructive with the local search methods for the RLP. Specifically we target the previous constructive and improvement methods Greedy + LS, H1 + LS, and H2 + LS of Chen et al. (2010), and our corresponding GRASP methods CG + LS, C1 + LS, and C2 + LS. We use the best value of α for each method according to the results of the first experiment. In particular, we set α to 0.9, 0.6, and 0.9 in CG, C1, and C2, respectively. As in the previous experiment, we consider the set of 20 instances and report, in Table 4, Value, #Best, Dev, and CPU time.

Table 4 shows that our three new GRASP heuristics are able to improve upon previous methods also based on construction plus local search. Specifically, CG, C1, and C2 present an average percent deviation from the best solutions of 1.45, 0.00, and 1.70, respectively, while Greedy + LS, H1 + LS, and H2 + LS obtain average percent deviations of 8.96, 9.96, and 7.50, respectively. It must be noted that these three previous methods are deterministic procedures that can only be run once. The randomized nature of the GRASP construction allows multiple runs, which provide a high-quality best solution over these runs. As expected, the GRASP-based constructions present

Table 4

Constructive with local search methods

	Value	#Best	Dev (%)	CPU (seconds)
Greedy + LS	5.65	11	8.96	0.17
H1 + LS	5.70	11	9.96	0.05
H2 + LS	5.55	14	7.50	0.01
GC(0.9) + LS	5.25	18	1.45	8.19
C1(0.6) + LS	5.15	20	0.00	2.24
C2(0.9) + LS	5.25	18	1.70	0.11

Table 5

Comparison of best methods

	Value	#Best	Dev (%)	CPU (seconds)
<i>n</i> = 40				
H2 + LS	3.96	59	4.46	0.003
C1(0.6) + LS	3.77	70	0.00	0.25
BRKGA	3.83	66	1.71	1.44
<i>n</i> = 60				
H2 + LS	4.37	62	1.74	0.006
C1(0.6) + LS	4.26	70	0.00	0.65
BRKGA	4.34	64	1.48	4.46
<i>n</i> = 80				
H2 + LS	4.90	46	8.25	0.01
C1(0.6) + LS	4.50	70	0.00	1.36
BRKGA	4.67	58	4.39	9.74
<i>n</i> = 100				
H2 + LS	5.27	50	5.69	0.02
C1(0.6) + LS	4.91	70	0.00	2.39
BRKGA	5.00	62	2.13	20.17

much longer running times than the deterministic methods, although they are still very moderate (less than 10 seconds).

In our final experiment, we compare our best method, C1 + LS, with the parameter α set to 0.6 with the previous best method, H2 + LS. We also include in this experiment the BRKGA, described in the previous section with $p = 100$, $p_e = 0.15$, $p_m = 0.10$, and $\rho_e = 0.7$, run for 20 generations. Table 5 reports the Value, #Best, Dev, and CPU time over the entire set of 280 instances divided according to their size.

Table 5 shows the merit of the proposed GRASP procedure. Our C1 + LS implementation consistently produces the best solutions with percent deviations smaller than those of the competing methods (and with number of best solutions found larger than the others). On the other hand, the H2 + LS algorithm is able to obtain relatively good solutions in short computational time. The BRKGA method performs well, better than H2 + LS, although it consumes significantly longer running times than the other methods.

We ran C1 + LS 100 times on a representative instance, stopping when a solution with objective value equal to the best known for this instance was found. For each run we recorded the running

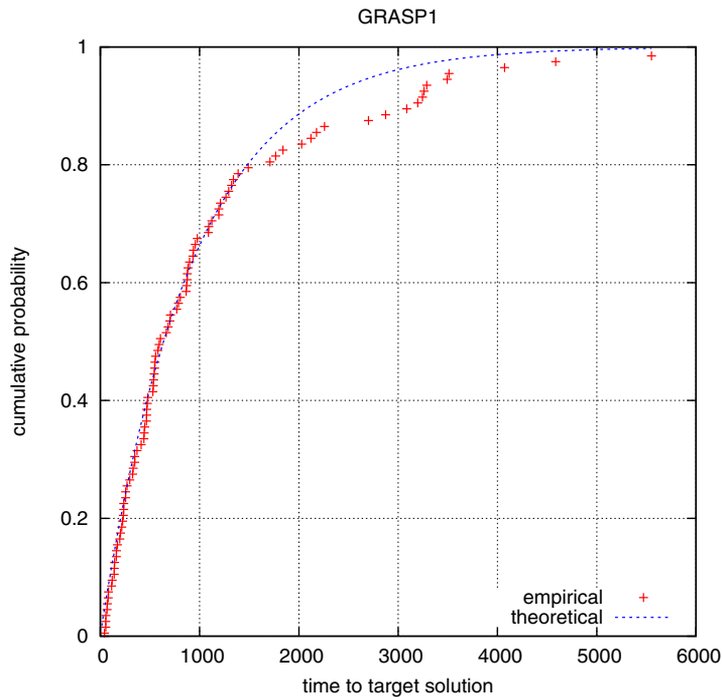


Fig. 4. Time-to-target plots (runtime distributions) for C1 + LS.

time. Each run was independent of the other, using a different initial seed for the random number generator. With these 100 running times, we plot the time-to-target plots (runtime distributions), shown in Fig. 4. This experiment confirms the expected exponential runtime distribution for our GRASP. Therefore, linear speed is expected if the algorithm is implemented in parallel.

6. Concluding remarks

This paper deals with the regenerator location problem (RLP) in optical telecommunication networks. In this problem, we are given a network represented by a weighted graph, where edge weights represent distances, and the maximum distance an optical signal can travel before it starts to degrade. Regenerators strengthen the signal to its original state. They are, however, expensive and one wants to deploy as few of them as possible in the network. We seek the locations of a minimum number of regenerators such that all pairs of nodes in the network are able to communicate using optical signals.

Three constructive procedures and a local improvement procedure were previously proposed by Chen et al. (2010). In this paper, we propose efficient implementations of the three constructive procedure as well as point out an error in the local search procedure for which we propose a correction. Not only are our implementations of the constructive procedures much faster (up to

7375 times faster in one case) than those of Chen et al. (2010), but, on average, they also find better solutions.

This paper also proposes three GRASP heuristics, each using one of the three constructive procedures of Chen et al. (2010) as well as the corrected local search procedure. The three (multistart) GRASP heuristics find much better solutions than the corresponding (single-start) constructive procedures with local search.

Finally, this paper proposes a bias random key genetic algorithm (BRKGA) for this RLP. The BRKGA uses a decoder based on the greedy constructive procedure of Chen et al. (2010). To enable the genetic algorithm to have running times similar to those of the other algorithms tested, a small population was used as well as a small number of generators were computed. Solutions found by the BRKGA were better than those found by the heuristics of Chen et al. (2010) but worse than those found by the GRASP heuristics. To find solutions of similar quality, we conjecture that the BRKGA would require larger populations, and more importantly, a greater number of generations.

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References

- Bean, J.C., 1994. Genetic algorithms and random keys for sequencing and optimization. *ORSA Journal on Computing* 6, 154–160.
- Chen, S., Ljubić, I., Raghavan, S., 2009. The generalized regenerator location problem. In Scutellà, M.G. (ed) Proceedings of the 2009 International Network Optimization Conference (INOC 2009), Copenhagen, Denmark.
- Chen, S., Ljubić, I., Raghavan, S., 2010. The regenerator location problem. *Networks* 55, 205–220.
- Chen, S., Raghavan, S., 2007. The regenerator location problem. Proceedings of the 2007 International Network Optimization Conference (INOC 2007), Spa, Belgium.
- Feo, T.A., Resende, M.G.C., 1989. A probabilistic heuristic for a computationally difficult set covering problem. *Operations Research Letters* 8, 67–71.
- Feo, T.A., Resende, M.G.C., 1995. Greedy randomized adaptive search procedures. *Journal of Global Optimization* 6, 2, 109–133.
- Festa, P., Resende, M.G.C., 2009a. An annotated bibliography of GRASP – Part I: Algorithms. *International Transactions on Operational Research* 16, 1–24.
- Festa, P., Resende, M.G.C., 2009b. An annotated bibliography of GRASP – Part II: Applications. *International Transactions on Operational Research* 16, 131–172.
- Flammini, M., Machetti, A., Monaco, G., Moscardelli, L., Zaks, S., 2009. On the complexity of the regenerator placement problem in optical networks. Proceedings of SPAA 2009, Calgary, pp. 154–162.
- Gonçalves, J.F., Resende, M.G.C., 2011. Biased random-key genetic algorithms for combinatorial optimization. *Journal of Heuristics* 17, 487–525.
- Gonçalves, J.F., Resende, M.G.C., 2014. An extended Akers graphical method with a biased random-key genetic algorithm for job-shop scheduling. *International Transactions in Operational Research* 21, 215–246.

- Gouveia, L., Patrício, P., Sousa, A., Valadas, R., 2003. MPLS over WDM network design with packet level QoS constraints based on ILP models. Proceedings of the IEEE Infocom, Vol. 1, San Francisco, CA, pp. 576–586.
- Moran-Mirabal, L.F., González-Velarde, J.L., Resende, M.G.C., 2014. Randomized heuristics for the family traveling salesperson problem. *International Transactions in Operational Research* 21, 41–57.
- Resende, M.G.C., Ribeiro, C.C., 2003. Greedy randomized adaptive search procedures. In Glover, F., Kochenberger, G. (eds) *Handbook of Metaheuristics*. Kluwer Academic Publishers, Norwell, MA, pp. 219–250.
- Resende, M.G.C., Ribeiro, C.C., 2010. Greedy randomized adaptive search procedures: advances and applications. In Gendreau, M., Potvin, J.-Y., (eds) *Handbook of Metaheuristics* (2nd edn). Springer Science and Business Media, New York, pp. 281–317.
- Ribeiro, C.C., Rosseti, I., Souza, R.C., 2013. Probabilistic stopping rules for GRASP heuristics and extensions. *International Transactions in Operational Research* 20, 301–323.
- Spears, W.M., DeJong, K.A., 1991. On the virtues of parameterized uniform crossover. Proceedings of the Fourth International Conference on Genetic Algorithms, San Diego, CA, pp. 230–236.
- Yetginer, E., Karasan, E., 2003. Regenerator placement and traffic engineering with restoration in GMPLS networks. *Photonic Network Communications* 6, 139–149.