ADJACENCY CONSTRAINTS IN FORESTRY – A SIMULATED ANNEALING APPROACH COMPARING DIFFERENT CANDIDATE SOLUTION GENERATORS

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Abstract. Adjacency constraints along with harvest volume constraints are important in long term forest management planning. Simulated annealing (SA) has previously been successfully applied when addressing such constraints. The objective of this paper is to assess the performance of SA using three methods for generating candidate solutions. Biased probabilities in the management unit (MU) selection were introduced, one static and one dynamic. The first one (Method 1) is the conventional (static) method. The two other methods were implemented through a search vector used in the candidate solution generator. These methods are based on (Method 2) the number of treatment schedules and standard deviation of NPV within MUs and (Method 3) the MU’s potential improvement in the objective function value, the number of URM adjacency violations an MU is involved in, the period specific volume harvested in an MU and the number of times an MU is selected. The methods were tested on a large number of datasets including 300 hypothetical forest landscapes characterized by three different initial age class distributions, respectively young, normal and old. Evaluation of the methods was accomplished by means of objective function values and first feasible iteration. Solutions improved when introducing bias in the probabilities for MU selection (Methods 2 and 3) compared to the conventional method (Method 1) and when the probability bias for selecting MUs is dynamic (Method 3) rather than static (Methods 1 and 2). The mean improvement for the average GAP obtained by Method 3 for young, normal and old forest landscapes was 20.88%, 12.84% and 5.20%, respectively. Whereas for the minimum GAP the mean improvement was 21.96%, 14.30% and 6.05% for young, normal and old forest landscapes, respectively.

Keywords: OR in Natural Resources; static and dynamic search vectors; grid landscapes; heuristics; Unit restriction model.

1 Introduction

Since simulated annealing (SA) was introduced by Kirkpatrick et al. (1983) as a technique for solving optimization problems, many disciplines such as biology, telecommunications, geology, electronics and medicine have been using SA as a tool to provide good (eventually optimal) solutions (see e.g. Chibante, 2010). Also in the forestry sector there are numerous optimization problems that can benefit from the use of heuristics like SA. Forestry planning problems include optimization of long term forest plans with detailed information about where, when and how silvicultural treatments should occur (i.e. treatment schedule), transportation of various forest products (e.g. roundwood, forest residuals) to and from the industry, and optimization of complicated industrial processes.

Over time, long term forestry optimization problems have become more complex since not only the economic aspects are important but also because environmental and social aspects of forestry have received increased attention. This development inevitably has an effect on the development of long term forest planning and its complexity. Accordingly, the dependency on mathematical programming and information technology has also increased considerably. A typical example would be maximization of economic income over time, under temporal and spatial (adjacency) restrictions. One important aspect of long term forest planning is spatial considerations, typically imposed in order to preserve wildlife habitats or enhance scenic beauty. Such consid-
erations restrict harvesting of neighboring management units (MU, i.e. forest stands) in the same or consecutive time periods.

A lot of work has been done related to adjacency constraints in long term forest planning (see e.g. reviews of Baskent and Keles, 2005; Weintraub and Murray, 2006; Shan et al., 2009). Adjacency constraints, as defined by Murray (1999), are divided into the unit restriction model (URM), where clear cut is not allowed in neighboring MUs in the same time period, and the area restriction model (ARM), where the total area of neighboring MUs harvested in the same time period should not exceed a defined maximum. In addition, the concept of “green-up constraints” is introduced in order to guarantee a time buffer between two consecutive clear cuts. This concept can be used along with the URM and ARM approaches (e.g. Brunelle et al., 1998; Boston and Bettinger, 1999, 2006; McDill et al., 2002; Goycoolea et al., 2009; Strimbu et al., 2010). Also the “core area” concept has become important in long term forest planning for example due to conservation of wildlife habitats, where the formation of contiguous areas of old growth forest over time in a forest landscape is promoted (e.g. Ohman and Eriksson, 1998; Ohman, 2000; Rebain and McDill, 2003).

SA has been successfully applied in many forestry problems addressing adjacency constraints. Like other heuristic methods, SA is based on a neighborhood solution search approach. A broad definition of a neighborhood of a solution is “the set of solutions which differ slightly from the original one”. In forestry planning applications with adjacency constraints, the neighborhood for S-metaheristics such as simulated annealing and tabu search is usually defined by the set of new solutions that can be obtained or reached through a change in timing of clearcut harvests or a change in the application of management regimes (e.g. Lockwood and Moore, 1993; Boston and Bettinger, 1999; Ohman and Eriksson, 2002; Bettinger et al., 2002; Ohman and Lämäis, 2005; Liu et al., 2006; Bettinger and Kim, 2008; Borges et al., 2014).

Another important aspect, along with the neighborhood definition, is the candidate solution generator. The solution generator, among the set of neighboring solutions selects the one(s) that should be evaluated. In forestry applications, the common procedure when generating a candidate solution is first to assume a uniform probability distribution when selecting an MU, and then within that MU, another uniform probability distribution is applied to select either the period where the clear cut should occur or which treatment schedule to apply. The candidate solution generator, however, can be manipulated in order to improve the quality of solutions (e.g. better objective function values) of the adopted heuristic (e.g. Borges et al., 2014). This may be done by introducing a bias in the probability distribution for selection of MUs and even treatment schedules. Biased selection criteria in this context means that we are using a probability distribution which is not uniform.

Some previous studies have implemented biased approaches in the MU selection. O’Hara et al. (1989) used a heuristic that moved only through the feasible space, and three approaches with biased MU selection were compared with an unbiased (uniform) selection approach. The biased approaches were based on the improvement of the objective function value (harvested volume) and on the fewest effective adjacent units, i.e. the number of new units that cannot be harvested. Barrett and Gilless (2000) implemented bias in the MU selection by sorting the MUs in descending order according the MUs total net present value (NPV) and NPV per ha.

Biased criteria have been applied also in SA forestry applications. For example, Ohman and Eriksson (1998) applied SA maintaining the selection of MUs by first assuming a uniform probability distribution. However, after the MU selection only three treatment schedules among 14 were possible to select, i.e. two treatment schedules moving the period for clear cut one period forward or backward, respectively, and one treatment schedule maintaining the selected MU un-harvested over the entire time horizon. Baskent and Jordan (2002) maintained the selection of MUs by first assuming a uniform probability distribution, but the final harvest was assigned to the period that returned the lowest harvested volume in order to favor non-violation of the minimum harvest volume constraints. Borges et al. (2014) developed three biased approaches in the MU selection and compared them with the conventional approach where a uniform probability distribution in the MU and treatment schedule selection was assumed. The three biased criteria introduced in the MU selection took into account the number of treatment schedules and/or the standard deviation of the NPV within an MU. The biased criteria in that study, however, were defined beforehand, i.e. the probabilities of selecting an MU were calculated before the candidate solution generator started and remained constant throughout all SA iterations. This means that the candidate solution generator worked with static search vectors.

It is also possible to introduce bias in MU selection in a dynamic way where the probabilities are updated according to the current or best solution. For example, if penalty functions are adopted (see e.g. Lockwood and Moore, 1993), it is possible to consider the number of constraints that each MU is involved in. In a case where only feasible solutions are allowed (no use of penalty functions), one may in the MU selection take into ac-
count the number of feasible treatment schedules for the MUs.

SA is in general suitable for working in large neighborhood frameworks. This is particularly important in forestry problems with adjacency constraints. SA is fairly simple to implement and since it works with two solutions (current and candidate), it is relatively fast when evaluating many solutions. We are not aware of any previous work in the long term forestry planning literature that uses SA where the candidate solution generator applies dynamic probabilities for the MUs selection. The objective of this paper is therefore to assess the performance of SA under three different methods for generating candidate solutions.

The remainder of this paper is as follows: section 2 describes the general planning model, the case study based on a forestry problem, the simulated annealing approach including the methods for applying the candidate solution generator and finally the measures for how the three methods are compared. Section 3 presents results, section 4 contains a discussion on how the methods perform, whereas section 5 presents the conclusions.

2 Material and Methods

2.1 General model We use a standard formulation of a forestry planning problem where one treatment schedule is selected for each MU so that the NPV is maximized over an infinite time horizon. This objective schedule is selected for each MU so that the NPV is of a forestry planning problem where one treatment applied per MU, i.e. all the area of an MU should be manage by only one treatment schedule, equations (4) and (5) secure that only one MU within a clique can be harvested in each time period, i.e. they define the clique URM adjacency constraints, equation (6) relates the decision variables $y_{ik}$ and $x_{ik}$, equation (7) defines the total volume harvested in each time period, $v_{ikt}$ is the volume harvested per ha in period $t$ in MU $i$ in treatment schedule $k$.

The decision variables $y_{ik}$ takes the value 1 if treatment schedule $k$ is applied to MU $i$, and the value 0 otherwise. The decision variable $x_{ik}$ takes the value 1 if MU $i$ is clear cut in period $t$, and the value 0 otherwise. Thus, equation (1) defines the objective function which maximizes NPV, equation (2) defines NPV, equations (3) and (4) secure that only one treatment schedule is applied per MU, i.e. all the area of an MU should be harvested in each time period, $X_{it}$ is the area of MU $i$, $p_{ik}$ is the NPV per ha associated with MU $i$ when treated with treatment schedule $k$, $V H_k$ is the total volume harvested in period $t$, $v_{ikt}$ is the volume harvested per ha in period $t$ in MU $i$ in treatment schedule $k$.

The problem presented is the same as in Borges et al. (2014) but the formulation of the model is different. Rather than defining the URM adjacency constraints in a pairwise approach, we use the clique approach (e.g. Murray and Church, 1996b). For that, the additional decision variables $x_{it}$ need to be defined and related to the decision variables $y_{ik}$ (equation 5). This formulation is more compact than the formulation made by Borges et al. (2014), i.e. the total number of constraints (4) and (5) are less than the number of constraints needed to define the adjacency constraints in a pairwise approach.

2.2 Case study To compare the three methods for selection of a neighborhood solution, 300 artificial forests (from now on called datasets) were used. These datasets
represent three different forest landscapes with different initial age class distributions, namely young, normal and old (Table 1). Each dataset within a forest landscape encompasses 1,600 MUs distributed over a grid of 40 x 40 cells. An area of 1 ha was assigned to all cells. The grid configuration and the equal area assigned to each cell avoid effects of MU size and number of neighbors per MU when assessing the methods. Moreover, the datasets used in this work, are based on 8990 sample plots from the Norwegian national forest inventory (NFI).

Table 1: Age class distribution (%) for the different forest landscapes.

<table>
<thead>
<tr>
<th>Age class</th>
<th>Forest landscape</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Young</td>
</tr>
<tr>
<td>0–10</td>
<td>15</td>
</tr>
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<td>4</td>
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<tr>
<td>101–110</td>
<td>3</td>
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<tr>
<td>111–120</td>
<td>2</td>
</tr>
<tr>
<td>≥121</td>
<td>1</td>
</tr>
</tbody>
</table>

The growth simulator GAYA (Hoen and Eid, 1990; Gobakken, 2003) was applied to generate treatment schedules for the MUs representing the landscapes. The simulator takes as input a set of MUs (plots) and a set of rules which define how and when forest treatments may be applied. The output provides detailed information on common forest state variables (e.g. standing volume, harvested volume) as well as treatments (e.g. final harvest, thinning) and corresponding economic values. The total number of treatment schedules among the 300 datasets varied between 131,102 and 156,963 corresponding to an average of 82 and 98 treatment schedules per MU, respectively. Moreover, within a dataset the minimum number of treatment schedules for MUs was 1 and the maximum varied between 594 and 833.

2.3 Simulated Annealing SA is a meta-heuristic that establishes a relationship between the annealing of solids and optimization problems (Dowsland, 1995). The usual parameters in SA applications are the initial temperature, the number of iterations allowed at each temperature, the cooling rate, and the final temperature at which the search is finished (e.g. Bettinger and Kim, 2008). The solutions producing improvements in the objective function value are always accepted. In order to prevent the search to be trapped in local optima, solutions with poorer objective function values can also be accepted depending on a probabilistic threshold which is defined by the current temperature and the difference between the current and candidate solution values. The temperature is kept constant for a certain number of iterations and then gradually reduced, lowering the probability of accepting solutions with poorer objective function values. The candidate solutions are generated based on the current solution, and the search process usually finishes when certain criteria are fulfilled, typically when reaching the final temperature or after performing a certain number of iterations.

When applying heuristics, procedures for local search are typically used and a definition of the neighborhood is thus required. Two main approaches can be used. The first one is accepting only feasible solutions within the neighborhood (e.g. Murray and Church, 1995; Liu et al., 2006; O’Hara et al., 1989). However, this approach is costly in terms of computational time, since each new solution needs to be assessed in terms of feasibility (Liu et al., 2006). The second approach is considering infeasible solutions within the neighborhood (e.g. Öhman and Eriksson, 1998; Falcão and Borges, 2001, 2002; Borges et al., 2014). The use of this approach implies some relaxation of the problem, i.e. some constraints are included in the objective function as penalties in order to worsen the objective function value of solutions that do not satisfy the constraints. In this way, more movements will be considered feasible and less check of feasibility is needed (Liu et al., 2006). This approach is usually faster in terms of computational time.

Quadratic penalty functions are frequently applied because they ensure that larger deviations from the targets
are penalized more than small ones. However, to construct and interpret penalty functions is a difficult task (Falcão and Borges, 2001). Penalty functions should be calibrated properly to ensure convergence towards feasible solutions (Lockwood and Moore, 1993). Moreover, according to Lockwood and Moore (1993), the magnitude of the values of the penalty functions should also be close to the objective function values to secure that none of the components of the new objective function have influence over the others. The resulting evaluation function is usually named “fitness function” and the respective value in SA is named “energy”.

In addition, the quality of the solutions obtained depends on the SA parameter settings and on the penalty function(s) (if adopted). Therefore, several test runs are required to find suitable values for the parameters. Usually some values for each parameter are tested and the combination producing the best results is adopted. In the present study SA was implemented and each forest landscape was parameterized as described by Borges et al. (2014). This means that the neighborhood definition and parameters setting were the same for all datasets within each forest landscape. In detail, the starting temperature was set according to the following formula:

$$st = -\frac{p_1 \times (NPV_0)}{\ln(p_2)}$$

where, $NPV_0$ is the NPV from the initial solution produced by SA, $p_1$ is a percentage of the initial NPV value and $p_2$ is the probability to accept a solution that varies (worse value) the previous amount. For our datasets we set $p_1=1\%$ for all forest landscapes and $p_2=10\%$, $20\%$ and $1\%$ for young, normal and old forest landscapes, respectively. For all datasets, the cooling rate was set to 0.995, the number of iterations at each temperature was set to 5,000 and the neighborhood structure was defined by changing one treatment schedule of a selected MU at each iteration (typically called 1-opt moves). Thus, the assessment of solution feasibility is only dependent on the evaluation of URM and sequential flow constraints. As mention, when using the first approach (above) usually, as an SA search matures, a large number of unsuccessful moves are attempted and rejected (due to violations of constraints or due to severe changes in the objective function value). If the first approach is employed, a process for reducing the temperature every $x$ number of unsuccessful iterations is needed in order for the process to properly terminate. Thus, we decide for the second approach, i.e. accepting infeasible solutions regarding these two types of constraints but penalizing them in the objective function.

Rather than using the penalty function formula applied by Borges et al. (2014), we use a slightly different one where the component penalizing the violation of the adjacency constraints have been changed. The main reason for the modification has to do with the fact that for some runs of SA the last feasible solution reported was found relatively far from the end of the SA run. This will “deteriorate” the values obtained not only for the average but also for the standard deviation of the solution values. In Borges et al. (2014), the adjacency constraints were penalized by considering the product between the average NPV per MU and the total number of conflicts among the MUs at the power of two. However, in a situation where only one conflict is observed the total NPV associated with the two MUs that are in conflict can be larger than the respective penalty value if the total NPV associated with the MUs is larger than the average NPV per MU. Therefore, we introduce as penalty value for adjacency constraints, the sum product between the number of conflicts in each MU and the current NPV value associated with each MU. The formula used was the following:

$$\Phi(l) = \sum_{i=1}^{N} \text{NumConflicts}_{il} \times NPV_{it} + A_{SF} \left( \beta \sum_{p=1}^{T} \text{DevVH}_{ip} \right)^2, \quad A_{SF} > 0$$

where, $\text{NumConflicts}_{il}$ is the total number of clear cuts in neighboring MUs of MU$_i$ when MU$_i$ is also a clear cut at iteration $l$, the $NPV_{it}$ is the NPV of MU$_i$ at iteration $l$, and $\text{DevVH}_{ip}$ is the deviation of volume harvested from the allowed interval in period $p$ at iteration $l$. The scaling factor $\beta$ was introduced because URM and harvested volume flow constraints are measured in different units and the scaling factor can be seen as the value of one cubic meter deviation in URM adjacencies violations. This scaling factor is computed from the quotient between the total $\text{NumConflicts}$ and the total $\text{DevVH}$ in the initial SA solution and asserts that both components of the penalty function are in the same unit. Thus, the value $A_{SF}$ represents the monetary loss of one conflict and was set to represent the average NPV per MU. This average NPV per MU is computed by first calculating the average NPV for each MU according to the treatment schedule list and then the average for all MUs. The search stops when the final temperature (1% of the starting temperature) is achieved.

Our implementation was developed in VB.NET, framework 4.5. The SA runs were performed in an Intel(R) Xeon(R) X5650 with 2.67GHz CPU and optimal solutions were obtained using CPLEX 12.5 with the default settings.

2.4 Methods applied in the candidate solution generator In this section, we give a detailed description
of the conventional method and the methods that introduce bias (different probabilities) in the MU selection. For all methods, the process of generating candidate solutions is maintained by first selecting an MU and then selecting a treatment schedule within the MU. In this way we introduce different methods of bias in the MUs selection and force the candidate solution generator to work differently and thus, different candidate solutions will be generated. The generator works with a search vector where each entry $i$ corresponds to the cumulative probability of selecting an MU. The main advantage of building the search vector in this way is that it is sorted (ascending) and allows the use of binary search to find which MU to be selected.

### 2.4.1 Method 1 – Conventional method

Method 1 (i.e., the conventional method) assumes a uniform probability distribution for selection of both MUs and treatment schedules where we first randomly select MUs and then randomly select treatment schedules within MUs. This method does not need an associated search vector since it is possible to mimic the search vector approach by using the random number generator provided by the software used. The procedure to build the search vector is as follows:

1. Start by computing the probability of selection for each MU ($P(MU_i)$) using the following:

$$P(MU_i) = \frac{1}{\#N}$$

2. Then a search vector is created so that each entry $i$ corresponds to the cumulative probability of selecting $MU_i$. The procedure is as follows:

1. Create a vector with dimension equal to the number of MUs plus one and set the first entry to zero.

2. Then for each entry $i$, compute the summation between the previous entry and the $P(MU_i)$.

### 2.4.2 Method 2 – Combining number of treatment schedules and standard deviation of NPV

In Method 2, the bias is introduced in each MU in order to consider both the number of treatment schedules and the standard deviation of NPV within MUs (see Borges et al., 2014). This method was the best of the two biased and static methods tested in Borges et al. (2014). The main idea of this method is to give a high probability of selecting MUs that have a high number of treatment schedules and a high standard deviation for NPV (SD-NPV).

The search vector for this method is static since the values assigned to each entry of the vector do not change throughout an entire SA run. The resulting search vector is obtained by applying the procedure as described for Method 1 with modification for step 1, i.e., for each $MU_i$, the probability of selection ($P(MU_i)$) is computed by weighting two quotients; [1] the quotient of the number of treatment schedules within each MU ($\#TS_i$) and the total number of treatment schedules among all MUs, and [2] the quotient of SDNPV within each MU (SDNPV$_i$) and the sum of SDNPV for all MUs:

$$P(MU_i) = \alpha \times \frac{\#TS_i}{\sum_{i=1}^{N} TS_i} + (1 - \alpha) \times \frac{SDNPV_i}{\sum_{i=1}^{N} SDNPV_i}$$

The weight $\alpha \in [0,1]$ was set in order to minimize the absolute difference between the Pearson correlation coefficient obtained between the number of treatment schedules and the probability assigned and between the NPV standard deviation and the probability assigned.

### 2.4.3 Method 3 – Combining potential for objective function value improvement and constraints mitigation

In Method 3, the bias for each MU is defined according to four criteria: 1) the potential of an MU to improve the objective function value, 2) the number of conflicts that an MU is involved in, 3) the harvested volume in an MU in a specific period and 4) the number of times the MU is selected. The potential improvement in the objective function value from an MU ($POI_i$) is computed as the difference in NPV between the treatment schedule with highest NPV and the NPV associated to the treatment schedule assigned to the MU. The number of conflicts that an MU is involved in ($NoConflict_i$) is computed by summing all common clear cut periods among the MUs adjacent to MU. This value is afterwards multiplied by the NPV associated with the treatment schedule applied to MU. To introduce a bias in the MU selection to consider mitigation of sequential flow of harvest volume is not simple because a treatment schedule can include both thinning and final harvests. Thinning and final harvest obviously appear in different time periods, but they both contribute to total volume harvested. Therefore, to simplify this issue we focus our attention on the periods where the largest deviation from the targets in volume harvested is observed (|Dev$_{t1,t2}$|). One way to introduce a bias in the MUs selection is to consider the volume harvested in each MU in a specific period $t$ ($vh_t$) and prioritize the selection according to the difference between $vh_t$ and $|Dev_{t1,t2}|$. Thus, within the two periods ($t_1$ and $t_2$), if $Dev_{t1,t2} < 0$ we focus on period $t_1$, and the main idea is to remove excess harvested volume in period $t$. 

$$P(MU_i) = \alpha \times \frac{#TS_i}{\sum_{i=1}^{N} TS_i} + (1 - \alpha) \times \frac{SDNPV_i}{\sum_{i=1}^{N} SDNPV_i}$$
The number of times an MU is selected (PKi) is a simple counting system.

The main aim of Method 3 is to force the generation of solutions in a direction that might lead to higher objective function values without adjacency violations. The number of times an MU is selected is introduced to mitigate possible situations where the other criteria end up focusing the search in just a few MUs. Thus, the more an MU has already been selected, the lower the probability should be for that MU to be selected again. This can be accounted for by considering the inverse of the number of times an MU is selected. The procedure to calculate the probability of each MU to be selected is therefore computed by weighting each criterion as follows:

\[
P(MU_i) = \frac{\alpha \times POI_i}{\sum_{i=1}^{N} POI_i} + \frac{\beta \times NoConflict_i \times NPV_i}{\sum_{i=1}^{N} NoConflict_i \times NPV_i} + \frac{\lambda \times |\|v_{ihp} - Dev\|_p|}{\sum_{i=1}^{N} |\|v_{ihp} - Dev\|_p|} + \mu \frac{1}{\sum_{i=1}^{N} PK_i}
\]

Since at least one of these four criteria is changing for each new iteration, the computation of the MUs probabilities will be “expensive” in terms of computational time if it is done at each iteration. After some experimental runs, we decided to update the MUs probabilities only when the best solution found is updated in young and normal forest landscapes while in old forest landscapes we decided to update for each iteration of the temperature. For this method, the search vector is dynamic because the bias introduced in each MU is changing during an SA run.

We worked with four different weightings depending on whether the constraints are violated or not. Thus, weights are updated automatically: if both types of constraints are violated we used the following weights of each criterion (\(\alpha = 0.2, \beta = 0.6, \lambda = 0.2, \mu = 0\)), if only one type of constraint is violated we used (\(\alpha = 0.2, \beta = 0.6, \lambda = 0, \mu = 0.2\)) and (\(\alpha = 0.6, \beta = 0, \lambda = 0.2, \mu = 0.2\)), for the case of URM and sequential volume violation, respectively. If no constraints are violated we used (\(\alpha = 0.8, \beta = 0, \lambda = 0, \mu = 0.2\)).

### 2.5 Measures for comparison of the methods

Because SA has stochastic properties, we use penalty functions and therefore have no guarantee that the final solutions reported are feasible or optimal. We performed 10 runs with different seeds (same seeds were applied in all methods) for each dataset and all runs within a dataset started with the same initial random solution. When comparing the methods, we only considered runs that report solutions without violation of the constraints, i.e. the last feasible solution found during a SA run.

The evaluation of the three methods follows mainly two procedures proposed by Bettinger et al. (2009). We started by comparing the solutions obtained from each dataset and each SA run against the respective optimal solution of the dataset found by CPLEX. This difference (GAP) is a good measure to use since its value is a relative difference independent of the magnitude of the objective function values. We also recorded the first feasible iteration in each SA run for the three methods. Within a dataset, we computed the average, minimum and standard deviation of the GAPs. These calculations represent the average performance, the best case performance and the variation in solution values of the methods (Bettinger et al., 2009). We also computed the average of the first feasible iteration for comparison purposes. Additionally, we compared the methods statistically applying the Wilcoxon pairwise signed-rank test (Wilcoxon, 1945) for the average, minimum and standard deviation GAPs and for the average first feasible iteration. Here we test if the differences are statistically different zero. A relative difference approach (Borges et al., 2014) is also applied for the average and minimum GAP between the methods in order to assess the improvement in percentage. Here the conventional method is the reference method.

### 3 Results

Pairwise comparisons between the methods for the average and minimum GAP, and for the average first feasible iteration for young, normal and old forest landscapes are shown in Figures 3, 4 and 5, respectively. When dots (datasets) are far from the reference line, this means that the difference in performance between the methods for that specific dataset is big.

The results obtained for young and normal forest landscapes are in general similar (Figures 3). For these forests landscapes, the average and minimum GAPs were larger for Method 1 than for the other two methods since large parts of the datasets are below the reference line. When comparing Methods 2 and 3, the latter has considerably more datasets with smaller average and minimum GAPs. Moreover, when the opposite results occur, i.e. when Method 1 has smaller average and minimum GAPs than Method 2 and 3, and when Method 2 has smaller average and minimum GAPs than Method 3, the relative difference between the methods is smaller since we in general for these datasets observe that they are closer to the reference line. All these observations from Figures 3 were confirmed by statistical tests (Table 2). In Figure 3 displaying the results from the old forest landscapes, also reveal the same trends as
Table 2: p-values from pairwise Wilcoxon signed-rank statistical tests for each criterion (columns). Within a forest landscape each criterion and method is represented by one hundred values (number of datasets). *** is significance level =0.01, ** is significance level =0.05, * is significance level =0.10

<table>
<thead>
<tr>
<th>Forest landscape</th>
<th>Methods compared</th>
<th>Average GAP</th>
<th>Minimum GAP</th>
<th>StdDev GAP</th>
<th>Average first feasible iteration</th>
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<tr>
<td>Young</td>
<td>1 vs 2</td>
<td>1.91E-18 ***</td>
<td>1.09E-16 ***</td>
<td>1.47E-08 ***</td>
<td>1.98E-18 ***</td>
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<td>0.001225 ***</td>
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<td>6.94E-08 ***</td>
<td>0.182724</td>
<td>1.98E-18 ***</td>
</tr>
<tr>
<td>Old</td>
<td>1 vs 2</td>
<td>7.76E-17 ***</td>
<td>4.47E-16 ***</td>
<td>0.038134 **</td>
<td>1.06E-17 ***</td>
</tr>
<tr>
<td></td>
<td>1 vs 3</td>
<td>3.44E-18 ***</td>
<td>9.89E-16 ***</td>
<td>0.979267</td>
<td>1.98E-18 ***</td>
</tr>
<tr>
<td></td>
<td>2 vs 3</td>
<td>0.701870</td>
<td>0.194507</td>
<td>0.054423 *</td>
<td>1.98E-18 ***</td>
</tr>
</tbody>
</table>

Table 3: Relative difference between Method 1 against Method 2 and 3 for the criteria Average and Minimum GAP.

<table>
<thead>
<tr>
<th>Forest landscape</th>
<th>Methods compared</th>
<th>Average GAP</th>
<th>Minimum GAP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Min Mean Max Median</td>
<td>Min Mean Max Median</td>
</tr>
<tr>
<td>Young</td>
<td>1 vs 2</td>
<td>-29.38 % -16.23 % -2.51 %</td>
<td>-16.32 % -31.14 % -13.09 %</td>
</tr>
<tr>
<td></td>
<td>1 vs 3</td>
<td>-35.07 % -20.88 % -7.16 %</td>
<td>-21.51 % -45.54 % -21.96 %</td>
</tr>
<tr>
<td>Normal</td>
<td>1 vs 2</td>
<td>-22.33 % -9.88 % -1.40 %</td>
<td>-9.39 % -26.38 % -9.81 %</td>
</tr>
<tr>
<td></td>
<td>1 vs 3</td>
<td>-25.68 % -12.84 % -2.76 %</td>
<td>-12.32 % -33.74 % -14.30 %</td>
</tr>
<tr>
<td>Old</td>
<td>1 vs 2</td>
<td>-13.01 % -5.28 % 14.37 %</td>
<td>-5.18 % -20.21 % -5.62 %</td>
</tr>
<tr>
<td></td>
<td>1 vs 3</td>
<td>-11.71 % -5.20 % 1.63 %</td>
<td>-4.94 % -16.51 % -6.05 %</td>
</tr>
</tbody>
</table>

young and normal forest landscapes when comparing Method 1 against Methods 2 and 3. However, when comparing Method 2 and 3, no special trends regarding the average and minimum GAP were observed. The statistical test performed (Table 2) confirmed these observations. Moreover, in all datasets the minimum GAP never reached the value zero, i.e. an optimal solution was never found.

The standard deviation of the GAPs obtained for a dataset may be used as a measure to describe the consistency of the results obtained by each method. Having as reference the values in the axis (Figures 1, 2 and 3), all the methods become less consistent as the forests get older, i.e. standard deviations of the GAPs obtained in young forest landscapes are smaller than in normal forest landscapes which again are smaller than in old forest landscapes. In general, the standard deviations of the GAPs obtained by Method 2 were slightly smaller than for the other two methods, but exceptions occur in old forest landscapes, since for some datasets the standard deviation of GAP observed was considerably higher compared to the other two methods (Figure 3). The standard deviations are also in general lower for Method 3 than for Method 1 in all forests landscapes. The statistical test performed (Table 2) confirmed these observations.

Regarding the average first feasible iteration, Method 2 tends to use more iterations than Methods 1 and 3, while for Method 3 the average iteration when the first feasible solution is found is considerably smaller than for the other two methods. This occurs in all forest landscapes (Figure 4). In addition, having as reference the values in the axis (Figure 4), we observed that the average first feasible iteration tends to appear earlier in old forest landscapes than in young and normal forest landscapes. The statistical test performed (Table 2) confirmed these observations.

The improvements in percentage are shown in Table 3, the negative values represent improvements. In young forest landscapes, for methods 2 and 3 the mean improvement was 16.23%, 20.88% for the average GAP and 13.09% and 21.96% for the minimum GAP, respectively. The same trends were observed in normal forest landscapes for both the average and minimum GAP, i.e. Method 3 shows bigger improvements than Method 2. However, these improvements were smaller than the ones obtained in young forest landscapes. For old forest landscapes the mean improvement for the average
GAP obtained by Method 2 was slightly better than the improvement obtained by Method 3, respectively 5.28% and 5.20%. However, for the minimum GAP the improvement obtained by Method 3 was again higher than the mean improvement obtained by Method 2. Moreover, all the medians show that in a considerable number of datasets improvements were obtained.

In all the methods, the number of SA runs finding feasible solutions was high. Only in five and six datasets of old forest landscapes, one run was not capable to find a feasible solution in Methods 1 and 2, respectively.

4 Discussion

Long term forest management planning and optimization of forest management are complex tasks, and are highly dependent on mathematical programming and information technology. An important aspect of long term forest planning is spatial considerations and constraints on management in adjacent MUs, which are often imposed in order to preserve wildlife habitats or enhance scenic beauty. SA has been used widely to address this type of problems (e.g. Shan et al., 2009) and improving its performance is therefore always an asset. Thus, in addition to the conventional method (Method 1) to select MUs, we applied two methods for introducing bias in the MUs selection, one using a static search vector developed by Borges et al. (2014) (Method 2) and one using a dynamic search vector developed for the present study (Method 3). These vectors were used in the candidate solution generator of an SA application.
The methods were tested through a comprehensive experimental design where we, from NFI sample plots comprising a multitude of forest conditions, constructed 300 hypothetical datasets characterized by three different initial age class distributions (young, normal and old). Our experimental design with a large number of datasets provides a more solid base for evaluation of the developed methods as compared to many previous works with relatively few datasets (e.g. O’Hara et al., 1989; Boston and Bettinger, 1999; Falcão and Borges, 2001, 2002; Crowe and Nelson, 2003). A large number of datasets is important because of the stochastic nature of the method applied (SA). Consequently, this enables better assessments of the impacts of using different ways to introduce bias in the selection of MUs. The grid configuration of the forest landscapes with equal cell (stand) size (1 ha) was also important to exclude factors that may influence the evaluation of the methods, for instance the number of neighbors (Li et al., 2010) and the area of each MU. Even controlling such conditions in the experimental design it could be interesting to assess the performance of the methods in different forest conditions such as MUs with different size and shape.

In general, the average objective function values obtained from all datasets indicated that the biased methods perform better than the conventional method. Moreover, the dynamic approach (Method 3) produced lower GAPs than the static approaches (Methods 1 and 2) in the majority of the datasets (Figures 1, 2 and 3). Within forest landscapes, the results obtained in young and normal forest landscapes by the new method presented in this work (Method 3) was the best, whereas
Figure 3: Scatter plots for old forest landscapes comparing Methods 1 and 2 (upper panels), Methods 1 and 3 (middle panels) and Methods 2 and 3 (lower panels) for average, minimum and standard deviation of GAPs. Dashed lines are reference lines denoting the equality of the methods. Dots (datasets) above the reference line means that the method in Y axis has bigger values than the method in X axis.

for old forest landscapes none of the methods (Method 2 and 3) outperformed the other in regards to the objective function values. These results agree with our expectations; first because the results of Borges et al. (2014) in general showed that introducing bias through static search vectors in the MU selection performed better than the conventional method, and second because by selecting some MUs according to relevant criteria (management problem dependent) and by changing these selection probabilities dynamically, achieving better solutions is more likely.

When comparing the standard deviations of the GAPs, in general within a forest landscape, the solutions obtained from each method were similar (Table 2). Furthermore, as found in McDill and Braze (2000) old forest problems are more difficult to solve, and the results from our problems indicate the same, since not only the GAPs are bigger but also the increase of the standard deviations observed from young towards old forest landscapes may indicate that datasets of old forests landscapes are more difficult to solve. Although, the numbers of treatment schedules among all forest landscapes is the relative the same, more variables indicating clear cuts and consequently also more clique constraints were generated in old forest landscapes. This might explain why this type of forest landscape is more difficult to solve. Thus, in future research more focus should be put on this type of forest landscapes. Furthermore, the results obtained by Borges et al. (2014) for Methods 1 and 2 were improved, i.e. average GAPs became smaller and the difference between these two methods increased in all forest landscapes. This shows that the new penalty
function is more appropriate than the penalty function applied in the work of Borges et al. (2014).

Regarding the average first iteration when a feasible solution is found, there was undoubtedly a difference between the methods. Method 2 needs more iterations than Method 1 to find the first feasible solution while Method 3 uses considerably less iterations, showing that the bias introduced in MU selection in order to mitigate the two types of constraints was successful (Figures 3, 2 and 3). The value of that solution might not be very good. However, it can be used as a starting point for any other technique or the subset of following feasible solutions found can be used as initial population in evolution programs. When comparing Methods 1 and 2 these results are not entirely in agreement with Borges et al. (2014), where only in old forest landscapes it was evident that Method 2 in average needed more iterations to find the first feasible solution. This difference in results is due to the new penalty function applied.

In general Method 3 seems most robust as this method produces better objective function values and performs as consistent (standard deviations) as the other two methods. However, one disadvantage of using Methods 2 and 3, compared to Method 1, is the weighting criteria that have to be considered for the search vector. In Method 2, this disadvantage was reduced by an automatic procedure adapting to the problem at hand, i.e. to the specific dataset (see Borges et al., 2014). For Method 3, although the weights given to the criteria implemented on the search vector performed well, it would be of interest to perform a sensitivity analysis in order
to assess the impacts of using different weighting alternatives.

Heinonen and Pukkala (2004) showed that performing changes in two MUs are better than in only one MU and that this may improve the quality of the solution. Thus, an interesting challenge for future research could be not only applying dynamic search vectors but also to extend dynamically the size of the solution neighborhood. This could for instance be useful when solving adjacency violations, since they can occur in different disjoint cliques of MUs. Another interesting extension for the dynamic approach could be the introduction of bias also in the selection of the treatment schedules after the MUs are selected (e.g. Baskent and Jordan, 2002).

Spatial considerations and adjacency constraints are important aspects of long term forest planning. It is therefore crucial to develop efficient tools for decision support in forestry that are able to handle these aspects. SA is one of the most used metaheuristics to address this type of forest planning problems. Our work demonstrates that not only the parameterization of SA is important for the quality of the solutions but also the way the search in the solution space is performed. Moreover, guiding the search dynamically towards more promising areas of the solution space including mechanisms to mitigate constraints violations (Method 3) seems to be a better strategy than using static methods (Methods 1 and 2), and will in general produce higher objective function values, i.e. lower average and minimum GAPs.

5 Conclusions

Introducing bias in the candidate solution generator selecting MUs in SA (Methods 2 and 3) improved the performance compared to the conventional method (Method 1) when forestry planning problems with adjacency and sequential flow constraints were addressed. Considerable improvement and consistency in the objective function value of the solutions was also achieved by means of a new penalty function and by introducing the bias for selecting MUs dynamically (Method 3). The mean improvement for the average GAP obtained by Method 3 for young, normal and old forest landscapes was 20.88%, 12.84% and 5.20%, respectively. Whereas for the minimum GAP the mean improvement was 21.96%, 14.30% and 6.05% for young, normal and old forest landscapes, respectively. Introducing a dynamic bias in MUs selection considering the mitigation of constraints also shortened the appearance of the first feasible solution compared to the static methods. Moreover, not only the parameterization of the SA but also the way the search of the solution space is performed is important for the quality of the solutions.

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REFERENCES


