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Parallel implementation of simulated annealing to reproduce multiple-point statistics

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Abstract

This paper shows an innovative implementation of simulated annealing in the context of parallel computing. Details regarding the use of parallel computing through a cluster of processors, as well as the implementation decisions, are provided. Simulated annealing is presented aiming at the generation of stochastic realizations of categorical variables reproducing multiple-point statistics.

The procedure starts with the use of a training image to determine the frequencies of occurrence of particular configurations of nodes and values. These frequencies are used as target statistics that must be matched by the stochastic images generated with the algorithm. The simulation process considers an initial random image of the spatial distribution of the categories.

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Nodes are perturbed randomly and after each perturbation the mismatch between the target statistics and the current statistics of the image is calculated. The perturbation is accepted if the statistics are closer to the target, or conditionally rejected if not, based on the annealing schedule.

The simulation was implemented using parallel processes with C++ and MPI. The message passing scheme was implemented using a speculative computation framework, by which prior to making the decision of acceptance or rejection of a proposed perturbation, processes already start calculating the next possible perturbation at a second level; one as if the perturbation on level one is accepted, and another process as if the proposed perturbation is rejected. Additional levels can start their calculation as well, conditional to the second level processes. Once a process reaches a decision as to whether accept or reject the suggested perturbation, all processes within the branch incompatible with that decision are dropped. This allows a speed up of up to $\log_n(p+1)$, where $n$ is the number of categories and $p$ the number of processes simultaneously active.

Examples are provided to demonstrate improvements and speed ups that can be achieved.

Key words: Geostatistics, Stochastic Simulation, Multipoint, Speculative
1. Introduction

Numerical modeling with geostatistical techniques aims at characterizing natural phenomena by summarizing and using the spatial correlation for the statistical inference of parameters of the distribution of uncertainty at unsampled locations in space. In simulation techniques, this spatial correlation is imposed into a model commonly constructed on a regular lattice. The models must reproduce the statistical (histogram) and spatial distribution (variogram or other spatial statistics) and their quality is often judged in terms of the reproduction of geological features (Ortiz and Peredo, 2009).

Conventional techniques in geostatistics address the modeling using statistical measures of spatial correlation that quantify the expected dissimilarity (transition to a different category) between locations separated by a given vector distance, in reference to a given attribute, such as the facies, rock type, porosity, grade of an element of interest, etc. This is done using the variogram. Limitations of these techniques have been pointed out in that they only account for two locations at a time when defining the spatial structure. Much richer features can be captured by considering multiple-point statistics that consider the simultaneous arrangement of the attribute of interest.
at several locations, providing the possibility to account for complex features, such as hierarchy between facies, delay effects, superposition, curvilinearity, etc. There are several approaches to simulate accounting for multiple-point statistics. Modifications of conventional methods to impose local directions of continuity using the variogram is a simple approach to impose some of the complex geological features (Xu, 1996; Zanon, 2004). Object based methods and methods inspired in the genetic rules and physics of the deposition of sediments in different environments also seek to overcome the limitations of conventional categorical simulation techniques, with significant progress (Deutsch and Wang, 1996; Tjelmeland, 1996; Pyrcz and Strebelle, 2008).

Presently, the most popular method is a sequential approach based on Bayes' postulate to infer the conditional distribution from the frequencies of multiple-point arrangements obtained from a training image. This method, originally proposed by Guardiano and Srivastava (1993), and later efficiently implemented by Strebelle and Journel (2000), is called single normal equation simulation (snesim) (see also Strebelle, 2002). This method has been the foundation for many variants such as simulating directly full patterns (Arpat and Caers, 2007; Eskandari and Srinivasan, 2007) and using filters to approximate the patterns (Zhang, Switzer and Journel, 2006). The use of
a Gibbs Sampling algorithm to account directly for patterns has also been proposed (Boisvert, Lyster and Deutsch, 2007; Lyster and Deutsch, 2008). A sequential method using a fixed search pattern and a ‘unilateral path’ also provides good results (Daly, 2005; Daly and Knudby, 2007; Parra and Ortiz, 2009). Other approaches available consider the use of neural networks (Caers and Journel, 1998; Caers and Ma, 2002), updating conditional distributions with multiple-point statistics as auxiliary information (Ortiz, 2003; Ortiz and Deutsch, 2004; Ortiz and Emery, 2005) or secondary variable (Hong, Ortiz and Deutsch, 2008), and simulated annealing (Deutsch, 1992).

Simulated annealing provides a very powerful framework to integrate different types of statistics, and potentially, generate models subject to constraints that cannot be handled by other methods. This motivates studying approaches to speed up the iterative process involved in annealing simulation. The recent increase in availability of powerful multiple-processor computers and multiple-core central processing units (CPU), as well as the use of graphics processing units (GPU) for parallelizing the calculations required for some heavy computing tasks, motivates researching new applications in this framework and lets us revisit algorithms that were too demanding for the technology existing a few years ago. In this paper, we explore a known paradigm in
Computer Science and implement simulated annealing to impose multiple-point statistics. The approach is known as speculative computing, and works in a parallel computing setting, aiming at reducing the time of computation for complex problems. Performance is assessed regarding speedup in computation time and statistical reproduction of multiple-point frequencies in the simulated realizations.

2. Simulated Annealing

Simulated annealing is a general optimization algorithm that can generate numerical models by reducing an objective function -usually minimizing a weighted sum of mismatch terms with respect to reference values-, reproducing different spatial statistics and respecting constraints imposed in that objective function (Besag, 1986; Farmer, 1992; Geman and Geman, 1984; Kirkpatrick et al., 1983; Rothman, 1985).

In a spatial context, the algorithm can be used to generate models in a lattice where each node of the grid has a value for a particular property being simulated and the objective function considers some statistical parameter that relates those nodes spatially. Typical applications have been the simulation of categorical variables such as facies or rock types, or the repro-
duction of continuous variables, such as petrophysical properties (porosity, permeability), conductivity or concentrations of elements (Goovaerts, 1996; Fang and Wang, 1997).

In essence, the algorithm works by perturbing one or more nodes at a time of an initial model, which usually is a model with a random spatial distribution of values. At every step of the process, the mismatch between the current statistics of the model and those required (target statistics) is quantified. If a perturbation reduces the mismatch, this means the simulated model has statistics closer to the target ones, hence the perturbation is kept. If a perturbation increases the mismatch, this means the model has statistics that are more different from the target ones, hence the perturbation should be rejected. However, in simulated annealing some of the unfavourable perturbations are kept, in order to allow the model moving away from local minima and reaching a lower mismatch later on. If a local minimum is reached with a given perturbation and all unfavourable changes are rejected, then it will be impossible to move out of that local minimum to get closer to the global minimum. Depending on the topology of the solution space, unfavourable changes should be rejected with a higher probability. The rate of acceptance of unfavourable perturbations is controlled by the “annealing
schedule”, which defines the probability of acceptance of bad changes and also controls the way this probability changes as the simulation progresses.

The annealing schedule is defined by an initial temperature and a procedure to lower that temperature as simulation progresses. In addition to these parameters, a perturbation mechanism is required, e.g. swapping nodes, randomly perturbing one or more nodes, etc, and one or more stopping criteria.

The general formulation of the algorithm considers an objective function of the following type:

\[ O = \sum_{i=1}^{N_c} w_i O_i \]  

(1)

where \( N_c \) is the number of components in the objective function, \( w_i \) are the weights assigned to each one of the components, and \( O_i \) is the mismatch value for component \( i \). For example, this function could be composed by the mismatch in histogram reproduction, defined as the squared difference in the cumulative frequencies measured at some quantiles for the model simulated versus the target histogram, a mismatch in variogram reproduction, composed by squared differences between the target variogram model and the variogram calculated from the realization being perturbed, for a number of lag distances, and a mismatch in the reproduction of multiple-point statistics.
In general any constraint can be ensured in a similar fashion. Conditioning data can be imposed simply by not allowing the nodes to be perturbed at sample locations. An important requirement to achieve a low mismatch between model and target statistics is that all the statistics and constraints must be consistent. Since imposing the multiple-point statistics implicitly defines lower-order statistics, if statistics are inferred from multiple sources, some inconsistencies are expected. Most often, we do not have control over the consistency of the statistics we try to impose, as these may come from different sources. The use of training images as the basis for inferring the multiple-point statistics can generate problems when its lower order statistics (histogram and variogram) are different than those on the sample data (Ortiz, Lyster and Deutsch, 2007).

In general, the reproduction of a variogram map, indicator variograms, a histogram of multiple-point statistics for some pattern sizes and the requirement of honouring conditional information can be imposed through elements of the objective function.

The typical procedure for implementing the algorithm is:

1. Start with a spatially random distribution of values over all the nodes on the lattice to be simulated, but ensuring that conditioning values at
2. Compute the global objective function, considering all target statistics and set the initial parameters of the annealing schedule.

3. Select a node at random and propose a different value for that location.

4. Evaluate the change in the objective function. This updating can be done by only computing statistics affected by the change. This avoids the lengthy calculation of a usually complex global objective function.

5. Calculate the probability of acceptance of the proposed perturbation. If the change is favourable, the perturbation is accepted; otherwise, the perturbation may be conditionally accepted as dictated by a probability distribution defined by the annealing schedule. The probability of acceptance is given by the Boltzmann distribution:

\[
P(\text{accept}) = \begin{cases} 
1 & \quad O_{\text{new}} \leq O_{\text{old}} \\
\exp\left(-\frac{O_{\text{old}} - O_{\text{new}}}{t}\right) & \quad \text{otherwise} 
\end{cases} \tag{2}
\]

where \( t \) is a parameter equivalent to the product of the Boltzmann constant \( k_b \) and the temperature \( T \) in the application to the physical process. By analogy, \( t \) is called the temperature in simulated annealing; \( O_{\text{old}} \) and \( O_{\text{new}} \) are the values of the objective function before and after the perturbation, equivalent to the difference in Gibbs free en-
ergy $\Delta E$ in the physical process of annealing. In simulated annealing, the temperature must be lowered as the algorithm runs, emulating the cooling that occurs during the physical process that lets the molecules reorganize to lower energy states.

6. Update the objective function if the change was accepted.

7. Decide if the temperature should be lowered.

8. Check the stopping criteria and stop or go back to 3.

A variety of options regarding the perturbation mechanism (Deutsch and Wen, 2001; Deutsch, 2002), updating the objective function (Deutsch, 2002), and defining the initial temperature (Norrena and Deutsch, 2000) have been proposed. Usually, these parameters strongly depend upon the problem at hand and are set empirically after some trial and error runs.

3. Implementation

A 2D implementation of simulated annealing to impose multiple-point statistics in a parallel computation framework is presented. The tools used are C++ and MPI (Pacheco, 1996). This implementation could be extended to 3D and multiple classes, although dimensionality may be an issue as the pattern size and number of categories increase.
3.1. 2D Patterns

The 2D patterns used to calculate the multiple-point statistics are stored using the class `map<string,int>` from the C++ Standard Template Library. The implementation is made to calculate rectangular patterns and supports two categories of data (binary case), represented by black and white pixels in Figure 1. Each pattern is represented by a string of 1’s, 0’s and X’s. The X indicates that there is no information available at that node.

![Figure 1: Pixels and patterns](image)

For example, in Figure 1, the pattern A is represented by the string

```
110111101
```

and the pattern B by

```
001X001X101XXXXX.
```
The class `map<string,int>` is used to store the occurrence of the patterns represented by these strings in a training image. That class allow assignment of an integer to each string which acts as a counter. The counter is increased every time an occurrence of the string is detected in the training image.

If the class `map<string,int>` is instantiated by the variable `hash` and `code="110111101"` then, if it is the first occurrence of the pattern represented by that string code, an insertion is performed with frequency equal to 1: `hash.insert(make_pair(code,1))`, and in case it has already been detected, an increment of 1 in the frequency is performed: `hash[code]++`.

### 3.2. Parallel Simulated Annealing

Simulated annealing is an inherently sequential algorithm because a decision about accepting or rejecting the current state, defined by the proposed perturbation, must be made and this decision conditions all subsequent states of the model. Implementing simulated annealing in a framework called Speculative Computation (Burton, 1985; Witte, Chamberlain and Franklin, 1991) allows reducing the computation time to achieve models that converge to the required statistics. In that framework, a binary tree of communications is designed, where each node of the tree represents a process launched by an independent CPU. Every time a process finishes the calculation step, it sends
the information to the root process. This results in the evaluation of several
states in a time close to the time required for computing a single state in a
single process, but requires additional processing capacity to be available.

Suppose that we are in the state $i$ and the objective state is the state $i+3$.
The path to be followed before reaching that state is $i \rightarrow i+1 \rightarrow i+2 \rightarrow i+3$
and for each step the decision of perturbing the state must be calculated,
taking a considerable amount of time (it involves the calculation of the ob-
jective function). In this case, 3 decision evaluations must be calculated, so
the decision tree has $2^3 - 1$ nodes (balanced binary tree). In the sequen-
tial simulated annealing, the only process involved must evaluate 3 tasks
sequentially. In the parallel simulated annealing, where each node of the
decision tree if evaluated by a different process, the tasks are the following
(one task per process):

- The decision for state $i$.
- The decision for state $i + 1$, assuming that the decision for state $i$ was
  accept.
- The decision for state $i + 1$, assuming that the decision for state $i$ was
  reject.
• The decision for state $i + 2$, assuming that the decision for state $i$ was accept and assuming that the decision for state $i + 1$ was accept.

• The decision for state $i + 2$, assuming that the decision for state $i$ was accept and assuming that the decision for state $i + 1$ was reject.

• The decision for state $i + 2$, assuming that the decision for state $i$ was reject and assuming that the decision for state $i + 1$ was accept.

• The decision for state $i + 2$, assuming that the decision for state $i$ was reject and assuming that the decision for state $i + 1$ was reject.

With this scheme it is expected that the entire decision tree will be calculated in an amount of time similar to the time of evaluation of the first task (decision for state $i$), so when the first task is completed, giving a result of accept or reject, the path to the corresponding leaf of the tree will be already calculated and the new state will be $i + 3$ (see figure 2).
3.3. Message passing scheme

An example with 7 processes in a balanced binary tree topology illustrates the message passing scheme. Considering an initial state of the model, the first step consists in applying perturbations in each node of the tree assuming that the corresponding parent node accepted the proposed perturbation.

In Figure 3 the process 0 obtains random coordinates $C_0$ (perturbation of process 0) and sends the coordinates to process 1 (accepter). The process 1 must update its configuration as soon as it gets the coordinates from its parent node. Then, the process 1 sends $C_0$ to the processes 3 and 4, and sends its random coordinates $C_1$ (perturbation of process 1) to process 3 (ac-
cepter). The process 3 must update its configuration in the coordinates $C_0$
and then $C_1$, in that order, and the process 4 must update its configuration
in the coordinate $C_0$. A similar situation occurs in the other branch with the
processes 2, 5 and 6 (rejectors).

In summary, each process performs the updates corresponding to the
random coordinates obtained by its parent, and the parent of the parent
and so on, depending on its condition of accepter or rejector, storing the
coordinate set from its parent and its own coordinate, which will be evaluated
in the next step.
In the second step, after updating the configurations for every node, the decision of accepting or rejecting the random coordinate chosen by each process must be computed. If the process 0 accepts a perturbation in $C_0$, the rejection branch (processes 2, 5 and 6) is discarded, sending a STOP signal to those processes (Figure 4). The procedure is performed in the next node, in this case, the node associated with process 1. In Figure 3 the final path is $C_0 \rightarrow \phi \rightarrow C_4$, where $\phi$ indicates that no perturbation is performed (a rejection branch).
The next step is to report every process about the path of perturbations, $C_0 \rightarrow \phi \rightarrow C_4$. This is done in two sub-steps: (1) the path must be sent back to the root node, associated with process 0. To send the path of perturbations to process 0, it has to travel backwards through all the nodes associated with the path to which a GO signal was sent. In this example, this occurs with processes 4 and 1. In Figure 5, process 4 sends $C_4$ to process 1 because it was decided to accept $C_4$, then process 1 sends $\phi$ and $C_4$ to process 0 ($\phi$ represents a rejection). Once those values are received by process 0, a decision is made.
whether to include or not $C_0$ in the path. In this example, $C_0$ is accepted and the final path to be broadcasted is $C_0 \rightarrow \phi \rightarrow C_4$. And, (2) the path must be sent from the root to every other node using a broadcast function from MPI (MPI\_Bcast) and then each process synchronizes its configuration with respect to the root node associated with process 0.

Figure 5: Third step in the message passing scheme
3.4. Speedup

3.4.1. Amdahl’s Law

To obtain a theoretical estimation for the speedup using the speculative framework, let us first recall Amdahl’s law (Amdahl, 1967) which gives a theoretical bound for the speedup of a general potentially parallel application.

The speedup is given by

$$S(P) = \frac{T_\sigma}{(1-r)T_\sigma + \frac{rT_\sigma}{P}}$$  \hspace{1cm} (3)
where $P$ is the number of processes, $T_\sigma$ represents the sequential processing time (with one process) and $r$ is the percent of possible parallel computations. $S(P)$ is the ratio between the sequential processing time and the parallel processing time. Cancelling $T_\sigma$ and calculating the derivative with respect to $P$ gives

$$\frac{dS(P)}{dP} = \frac{r}{((1 - r)P + r)^2} \quad (4)$$

which is positive for all $r \in [0, 1]$, then $S(P)$ is a monotone increasing function with limit $\frac{1}{1 - r}$ when $P \to \infty$. Therefore there is an upper bound for the speedup in a parallel application. This bound depends on the percentage of possible operations that can be calculated in parallel and the problem should be analyzed to characterize this bound.

3.4.2. Balanced Trees

Using the speculative framework to parallelize simulated annealing with a balanced binary tree of processes yields a theoretical speedup of $S(P) = \log_2(P + 1)$, where $P$ is the number of processes/nodes. The theoretical speedup coincides with the number of levels in the tree. For example, if the tree has 3 nodes (3 processes), the number of levels is 2 ($\log_2(3 + 1) = 2$) which coincides with the speedup. The approach could be generalized to be
implemented using a non balanced tree (Witte, Chamberlain and Franklin (1991)), although this is not covered in this work.

Tests conducted with 3 and 7 processes, confirmed the theoretic speedup in the balanced scenario, giving evidence of the power of the method by increasing the number of processes in the tree.

4. Examples

Some examples are presented to illustrate the implementation and results over three simple cases. Several tests were carried out to evaluate convergence and speedup of the algorithm, in particular considering multiple-point statistics obtained for squared patterns of adjacent nodes of size 2x2, 3x3 and 4x4 pixels. This is equivalent to trying to reproduce the multiple-point histogram (Deutsch, 1992).

Two objective functions were tested, one that assigns equal importance to all patterns:

\[ O = \sum_{i=1}^{N_{MPE}} (f_{target}^i - f_{model}^i)^2 \]  

where \( f_{target}^i \) are the frequencies of the multiple-point events from the training image, \( f_{model}^i \) are the statistics for the same multiple-point events
in the current state of the simulated model, and $N_{MPE}$ is the total number
of possible multiple-point events for the pattern size considered. A second
implementation is done assigning a weight inversely proportional to the fre-
quencies of multiple-point events, in order to increase the relative importance
of less frequent events in the objective function:

$$O = \sum_{i=1}^{N_{MPE}} \lambda_i (f_{i\text{target}} - f_{i\text{model}})^2$$

(6)

where $\lambda_i$ is a standardized weight inversely proportional to the target
frequency of the multiple-point event:

$$\lambda_i = \begin{cases} \frac{1}{f_{i\text{target}}} & f_{i\text{target}} \neq 0 \\ 0 & f_{i\text{target}} = 0 \end{cases}$$

(7)

All tests were run 10 times (10 realizations) with 1 process (sequential
implementation), 3 processes (2-level balanced tree) and 7 processes (3-level
balanced tree), so a total amount of 180 tests were performed for each ex-
ample (10 realizations × {1p, 3p, 7p} × {2x2, 3x3, 4x4} × {equal weight,
inversely proportional weight}). The tests performed with 1 processor consti-
tutes the sequential implementation (base case for measuring the speedup).
4.1. Example 1: channels

The first example consists of simulating a geological setting of sinuous channels in a background, using the training image presented in Figure 7 (Strebelle, 2002).

Figure 7: Training image: Channels

The average speedup of the parallelization for this example is presented in table 1, in which each test was performed 10 times and all tests are measured using the same annealing schedule. The maximum and minimum speedups are presented in the tables 2 and 3. An example of the simulated image, statistical validation and the evolution (in percentage) of the objective function for a particular test are presented in Figures 8 (case of equal weighting) and 9 (inversely proportional weighting).
Table 1: Average speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 7

<table>
<thead>
<tr>
<th></th>
<th>2 x 2</th>
<th>3 x 3</th>
<th>4 x 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 process</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>3 processes</td>
<td>1.75</td>
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<tr>
<td>7 processes</td>
<td>3.98</td>
<td>1.92</td>
<td>2.24</td>
</tr>
</tbody>
</table>

Table 2: Maximum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 7

<table>
<thead>
<tr>
<th></th>
<th>2 x 2</th>
<th>3 x 3</th>
<th>4 x 4</th>
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<tr>
<td>1 process</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>3 processes</td>
<td>2.47</td>
<td>2.59</td>
<td>1.88</td>
</tr>
<tr>
<td>7 processes</td>
<td>4.09</td>
<td>2.48</td>
<td>2.54</td>
</tr>
</tbody>
</table>
Table 3: Minimum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 7.

<table>
<thead>
<tr>
<th></th>
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<td>1.00</td>
<td>1.00</td>
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</tr>
<tr>
<td>3 processes</td>
<td>1.36</td>
<td>1.40</td>
<td>1.64</td>
<td>3 processes</td>
<td>1.42</td>
<td>1.69</td>
<td>1.84</td>
</tr>
<tr>
<td>7 processes</td>
<td>2.87</td>
<td>1.64</td>
<td>2.17</td>
<td>7 processes</td>
<td>1.49</td>
<td>1.55</td>
<td>2.45</td>
</tr>
</tbody>
</table>

Figure 8: Simulated images: Channels (equal weighting)
<table>
<thead>
<tr>
<th>Image</th>
<th>Patterns frequency</th>
<th>Convergence</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="2x2 Image" /></td>
<td><img src="image2.png" alt="2x2 Patterns Frequency" /></td>
<td><img src="image3.png" alt="2x2 Convergence" /></td>
</tr>
<tr>
<td><img src="image4.png" alt="3x3 Image" /></td>
<td><img src="image5.png" alt="3x3 Patterns Frequency" /></td>
<td><img src="image6.png" alt="3x3 Convergence" /></td>
</tr>
<tr>
<td><img src="image7.png" alt="4x4 Image" /></td>
<td><img src="image8.png" alt="4x4 Patterns Frequency" /></td>
<td><img src="image9.png" alt="4x4 Convergence" /></td>
</tr>
</tbody>
</table>

Figure 9: Simulated images: Channels (inversely proportional weighting)

The experimental speedup in the inversely proportional weighting case coincides more accurately with the theoretical speedup. The simulated images using this weighting more resemble the training image. It should be emphasized that the validation of the realizations is done from a statistical point of view, in terms of the reproduction of multiple-point frequencies. Visually, a better result would be obtained by increasing the pattern size or by implementing a multi-scale approach, such as a multiple grid (Strebelle,
A slower convergence of the objective function is the trade-off of this resemblance.

4.2. Example 2: ellipses 0

The second example consists in simulating an artificial setting of ellipses in a background, with their major semi-axis aligned with the vertical direction, using the training image presented in Figure 10.

![Figure 10: Training image: Ellipses 0](image_url)

The average speedup of the parallelization for this example is presented in table 4, in which each test was performed 10 times and all tests are measured using the same annealing schedule. The maximum and minimum speedups are presented in the tables 5 and 6. An example of the simulated image, statistical validation and the evolution (in percentage) of the objective function.
for a particular test are presented in Figures 11 (case of equal weighting) and 12 (inversely proportional weighting).

<table>
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</tr>
<tr>
<td>3 processes</td>
<td>2.19</td>
<td>2.38</td>
<td>1.71</td>
</tr>
<tr>
<td>7 processes</td>
<td>3.71</td>
<td>2.82</td>
<td>2.22</td>
</tr>
</tbody>
</table>

Table 4: Average speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for Figure 10

<table>
<thead>
<tr>
<th></th>
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<td>1.00</td>
</tr>
<tr>
<td>3 processes</td>
<td>3.45</td>
<td>2.99</td>
<td>1.85</td>
</tr>
<tr>
<td>7 processes</td>
<td>7.07</td>
<td>4.08</td>
<td>2.33</td>
</tr>
</tbody>
</table>

Table 5: Maximum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 10
Table 6: Minimum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 10

<table>
<thead>
<tr>
<th>Image Patterns frequency</th>
<th>Convergence</th>
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</tr>
<tr>
<td>1 process</td>
<td>1.00</td>
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<td>1.27</td>
</tr>
<tr>
<td>7 processes</td>
<td>1.31</td>
</tr>
</tbody>
</table>

Figure 11: Simulated images: Ellipses 0 (equal weighting)
There are no major differences in the experimental speedup for the equal and inversely weighted cases. The simulated images in both cases resemble the training image, except in the case of a 4x4 pattern using inversely proportional weighting in which there are convergence problems. Also, a slower convergence and a slight bias in the multiple-point statistics reproduction occur when the inversely proportional weights are used.
4.3. Example 3: ellipses 45

The third example consists in simulating a setting of ellipses in a background, each of them rotated in 45° with respect to the horizontal axis, using the training image presented in Figure 13.

![Training image: Ellipses 45](image)

Figure 13: Training image: Ellipses 45

The average speedup of the parallelization for this example is presented in table 7, in which each test was performed 10 times and all tests are measured using the same annealing schedule. The maximum and minimum speedups are presented in the tables 8 and 9. An example of the simulated image, statistical validation and the evolution (in percentage) of the objective function for a particular test are presented in Figures 8 (case of equal weighting) and 9 (inversely proportional weighting). Some problems related to edge effects...
are visible in these results. These can be explained by the statistical weight
that a perturbation node has on an edge which should be equivalent to the
one of a node at the center of the simulation domain. However, since a node
near the edge may affect a smaller number of patterns when perturbed, in
the current formulation of the objective function, it has a smaller influence
in the overall performance causing the edge effects. This can be solved by
modifying the objective function or by simulating a larger domain that is
later eroded to the size of interest.

<table>
<thead>
<tr>
<th></th>
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<th>3 x 3</th>
<th>4 x 4</th>
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<td>1.00</td>
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<td>3.04</td>
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</table>

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<tbody>
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<td>1.00</td>
<td>1.00</td>
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<tr>
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<td>4.15</td>
<td>3.03</td>
<td>2.53</td>
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</tbody>
</table>

Table 7: Average speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for Figure 13
Table 8: Maximum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 13

<table>
<thead>
<tr>
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</thead>
<tbody>
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<td>1.00</td>
<td>1.00</td>
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<table>
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</thead>
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<tr>
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<td>1.00</td>
<td>1.00</td>
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</tr>
<tr>
<td>7 processes</td>
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<td>6.92</td>
<td>2.63</td>
</tr>
</tbody>
</table>

Table 9: Minimum values for the speedup of the parallel implementation for the equal weighting (left) and inversely proportional weighting (right) for the examples using the training image in Figure 13

<table>
<thead>
<tr>
<th></th>
<th>2 x 2</th>
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<th>4 x 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 process</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>3 processes</td>
<td>1.12</td>
<td>1.27</td>
<td>1.80</td>
</tr>
<tr>
<td>7 processes</td>
<td>1.69</td>
<td>1.13</td>
<td>2.47</td>
</tr>
</tbody>
</table>
Figure 14: Simulated images: Ellipses 45 (equal weighting)
Figure 15: Simulated images: Ellipses 45 (inversely proportional weighting)
The experimental speedup in both cases coincides with the theoretical speedup, with small differences. As in example 2, the simulated images in both cases resemble the training image, with some convergence problems in the case of a 4×4 pattern considering the inversely proportional weighting case. Again, a slower convergence in the inversely proportional weighting case is observed.

5. Conclusions

We have shown a parallel implementation of simulated annealing under a speculative framework that allows speeding up sequential implementations. Several conclusions can be drawn from this work. The most relevant result is that a significant speedup can be achieved using multiple processors to run the simulation under the speculative scheme. The theoretical speedup log₂(\(P+1\)) on the balanced binary tree topology was approximately obtained in all of the test cases performed. The inclusion of a weighted objective function slightly improved the results in terms of reproduction of multiple-point statistics. However, the trade-off is an increase in the simulation time for larger templates (4×4 or more). Also, a slight bias in the statistical reproduction of multiple-point events could be seen. It should be pointed out
that this work aims at showing the potential of simulated annealing in terms of the statistical reproduction of multiple-point frequencies, rather than obtaining geologically appealing simulated images. This last point is relevant, as the final goal of the numerical algorithms is to assist in the study of the geological phenomenon. However, the geological quality of the resulting realizations depends on factors such as the pattern configurations and size used. Our focus was on the speedup that can be obtained with the speculative approach, considering that simulated annealing is powerful enough to include many more statistics and constraints in the objective function than presented in the examples provided here.

There are many avenues of further research. The optimization of data structures to manage the patterns and templates is important. The time of simulation grows exponentially with the size of the template so this topic is a main priority in the development of this technique. Also, continuing the research with non-balanced trees in the speculative scheme for simulated annealing is of interest.
6. Acknowledgments

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