

OPTIMIZATION AND PARALLELIZATION  
METHODS FOR THE DESIGN  
OF NEXT-GENERATION  
RADIO NETWORKS

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**Doctoral Dissertation**

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Doctoral Dissertation

**OPTIMIZACIJSKE IN VZPOREDNE METODE ZA  
NAČRTOVANJE RADIJSKIH OMREŽIJ  
NASLEDNJE GENERACIJE**

Doktorska disertacija

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## Abstract



# Povzetek

Prvi odstavek. Prvi odstavek. Prvi odstavek. Prvi odstavek. Prvi odstavek. Prvi odstavek.  
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# Abbreviations

## Abbreviations

2G	= Second Generation of mobile networks.
3G	= Third Generation of mobile networks.
4G	= Fourth Generation of mobile networks.
CPICH	= Common Pilot power Channel.
DB	= Database system.
GRASS	= Geographic Resources Analysis Support System.
GSM	= Global System for Mobile communications.
HPC	= High-perfomance computing.
HSDPA	= High Speed Downlink Packet Access.
HSPA	= High Speed Packet Access.
HSUPA	= High Speed Uplink Packet Access.
I/O	= Input/output.
LTE	= Long Term Evolution.
MW	= Master-worker parallel paradigm.
MWD	= Master-worker-database parallel paradigm.
PRATO	= Parallel radio-prediction tool.
RSCP	= Received signal code power.
RSG	= Regular square grid.
SHO	= Soft handover.
UMTS	= Universal Mobitel Telecommunications System.
WCDMA	= Wideband Code Division Multiple Access.

## Symbols

$\gamma^{\text{cov}}$  = Signal-to-interference ratio (SIR) coverage threshold.

$\gamma^{\text{cov}}$	= Signal-to-interference ratio (SIR) coverage threshold.
$\gamma^{\text{sho}}$	= SHO window.
$A_{\text{covered}}$	= Area under service coverage of the mobile network.
$A_{\text{total}}$	= Complete geographical area under optimization.
$as^{\max}$	= Maximum number of neighbouring cells in the active set.
$C$	= Set of antenna installations (cells) in a mobile network.
$C$	= Set of antenna installations (cells) in a mobile network.
$C_m$	= Subset of cells, $C_m \subset C$ , that cover a mobile $m \in M$ .
$c_m^*$	= Best-serving cell of mobile $m \in M$ .
$c_m^*$	= Best-serving cell of mobile $m \in M$ .
$cov(c, m)$	= Binary function to assert the coverage of a mobile $m \in M$ by from a cell $c \in C$ .
$cov(x, y)$	= Returns 1 if the coordinate $(x, y)$ is under mobile-network coverage.
$cov_{cm}$	= Returns 1 if the mobile $m \in M$ is under radio coverage of cell $c \in C$ .
$f_{\text{cov}}$	= Objective function for the service-coverage optimization problem.
$f_{\text{sho}}$	= Objective function of the SHO-balancing problem.
$L_{cm}^\downarrow$	= Downlink attenuation factor between cell $c \in C$ and mobile $m \in M$ .
$L_{cm}^\downarrow$	= Downlink attenuation factor between cell $c \in C$ and mobile $m \in M$ .
$L_{mc}^\uparrow$	= Uplink attenuation factor between mobile $m \in M$ and cell $c \in C$ .
$L_{mc}^\uparrow$	= Uplink attenuation factor between mobile $m \in M$ and cell $c \in C$ .
$M$	= Set of mobile devices or users of a mobile network.
$M$	= Set of mobile devices or users of a mobile network.
$P_c$	= Set of candidate CPICH power settings for cell $c \in C$ .
$P_c$	= Set of candidate CPICH power settings for cell $c \in C$ .
$p_c$	= Pilot-power setting of cell $c \in C$ .
$p_c$	= Pilot-power setting of cell $c \in C$ .
$p_m^\uparrow$	= Uplink transmit power of mobile $m \in M$ .
$SHO_m^\downarrow$	= Cell set to which a mobile may maintain concurrent downlink connections, i.e. downlink SHO.
$SHO_m^\uparrow$	= Cell set to which a mobile may maintain concurrent uplink connections, i.e. uplink SHO.
$sir(c, m)$	= Signal-to-interference ratio (SIR) from cell $c \in C$ to mobile $m \in M$ .
$sir(c, m)$	= Signal-to-interference ratio (SIR) from cell $c \in C$ to mobile $m \in M$ .

# 1 Introduction

Many researchers believe the computer has become the third method to do research, behind theory and experimentation, for both science and engineering. Although there is no complete agreement on the position intended for scientific computing with respect to the other two methods, it is undeniable that computational methods are an essential tool in most disciplines, particularly in those related to decision making.

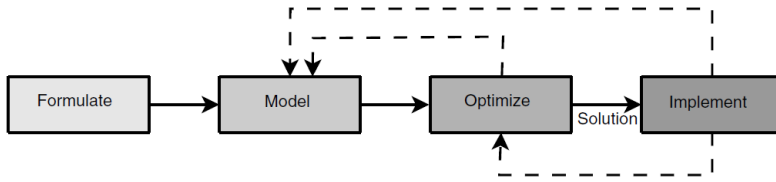
Nowadays, decision making is present practically everywhere. As scientists, engineers and managers have to make decisions in more complex and competitive circumstances every day, decision making involves dealing with rational and optimal approaches. According to Talbi [134], decision making consists in the following steps:

- formulate the problem,
- model the problem,
- optimize the problem, and
- implement a solution.

Formulating a decision problem means making an initial statement about it. Despite this first formulation may be imprecise, the objectives of the problem are outlined, together with internal and external factors that have some degree of influence over it. During the modelling of the problem, an abstract mathematical model is built for it. Sometimes this model is inspired by similar models in the literature, making it possible to tackle the problem with well-studied methods. After a model of the problem is available, the optimization step, i.e. generating “good” solutions for the problem, may begin. It is worth pointing out that the resulting solutions are given for the abstract model, and not for the original problem itself. Therefore, the performance of the obtained solution is indicative when the model is an accurate one [134]. In the last step, the obtained solution is practically tested by the decision maker and implemented if it is an “acceptable” or “good” one. In case of “bad” or “unacceptable” solutions, the decision-making process is repeated, possibly improving the model and/or the optimization algorithm. The process, as described here, is depicted in Figure 1.1.

Scientific computing, by means of computer-science methodology, makes possible the study of problems that are too complex to be treated analytically, or those that are very expensive or dangerous to be studied by direct experimentation. Real-world problems are typically very complex systems to be directly assessed by analytical models, and require a numerical simulation for their study. Computer simulations provide a resource to mimic the behavior of complex systems, by numerically evaluating a model and gathering its data to estimate their true characteristics [82].

A model is a simplified representation of a studied problem, and one of its purposes is to predict the effects of variations within the system. A good model is a balance between realism and simplicity. The system simulation, on the other hand, is the operation of the



**FIGURE 1.1** The classical process in decision making: formulate, model, solve, and implement. In practice, this process may be iterated to improve the optimization model or algorithm until an acceptable solution is found. Like life cycles in software engineering, the life cycle of optimization models and algorithms may be linear, spiral, or cascade.

Figure 1.1: The classical decision-making process, as presented in [134]. Multiple iterations of this process improve the optimization algorithm and/or model until an acceptable solution is found.

model. Its configuration can also be changed, allowing multiple experimental executions, something that might not be possible with the real system it represents [89]. However, it is important to understand that the models used in scientific simulations and engineering never offer a perfect model of the system they represent, but only a subset of its composition and dynamics. For this reason, experimentation and expert observation will always be essential as reference points for understanding the studied phenomena. Consequently, problems categorized as of large size and of considerable complexity represent a challenge, because of the different involved disciplines and the degree of difficulty of their modeling. Radio networks, in particular those of the third and fourth generations, fall under this characterization.

Radio networks represent one of the most fast-growing technology markets since the introduction of the Global System for Mobile communications (GSM) [3] as the second generation (2G) mobile networks more than twenty years ago. Its successor, the Universal Mobile Telecommunications System (UMTS) [4] marks an evolution from 2G, representing a milestone for the third generation of mobile radio networks (3G). In recent years, the first commercial networks implementing the Long Term Evolution (LTE), also known as fourth generation (4G) LTE, have also appeared [ref??]. The always increasing demand for more bandwidth has been one of the main forces behind the standardization and later implementation of systems delivering higher speed data services in order to improve the user's experience.

This evolution, first from 2G to 3G and later from 3G to 4G, has introduced not only the technology needed to increase both data-transfer capacity and voice quality, but also a greater complexity in terms of radio-network planning, deployment, and configuration. This fact has attracted the attention of the research community into areas such as the design and optimization of radio-networks.

In a traditional or manual approach, during the design phase of a radio network, a software tool would execute the analysis, while the human would make the change decisions. Therefore, a radio engineer configures the network parameters manually and the software tool analyzes the given configuration. If the obtained results are not acceptable, the analysis process has to be repeated several times, until the goal is achieved. We will refer to this process as manual radio-network optimization.

Advances in the last few years have improved the manual optimization process by introducing different problem-solving approaches that increase the role the computer has during the optimization of radio networks, consequently enlarging the scope of problems and instance sizes that may be subject to automated optimization. Still, there are some important aspects that restrict the utilization of these methods not only in real-world environments, but also when doing research in the area of optimization of radio networks:

- It is usually the case that the selected method is a compromise between solution quality and computational-time complexity. The proposed state-of-the-art approach for the evaluation of radio networks is the Monte-Carlo snapshot analysis. However, real-world environments, where radio-network design is carried out, require the evaluation of networks comprising up to thousands of base stations in a reasonable amount of time. Moreover, for applications involving radio-network optimization, multiple thousand evaluations are required to find a good solution, in which case also snapshot simulations are too time-consuming to be employed. Therefore, for such applications and environments, methods with improved time efficiency are required.
- A considerable number of publications in the field of radio-network optimization, of which we cite just a few for reference purposes [8, 122, 23, 26, 54, 117], base their simulations on platforms for which it is not possible to reproduce the experiments, either because they have used proprietary software or because the data is not available. This fact reduces the possibilities for comparing different approaches among each other, and significantly contrasts with other research areas, such as evolutionary computing or artificial intelligence, which have a set of open and well-known benchmarks for the community to use. Thus, an open and standardized framework would allow researchers to compare different methods and results in a simple and objective manner.
- Available commercial tools for radio-network evaluation have major drawbacks with respect to the size of networks that can be considered, simulation time, and especially the accuracy of the modeled system. Yet, if precise and fast methods were commercially available, they would lack the level of flexibility required by the scientific community. Consequently, an essential attribute of the framework is to be open source, so that anyone could extend it to meet some specific requirements. In the long term, this process should also extend the set of built-in functionality.
- Particularly for path-loss predictions, created with empirical mathematical models, the inaccuracy of the input data directly deteriorates the precision of the calculated results. Moreover, since the physical properties that influence the propagation of radio signals are not constant and every environment introduces its own deviations, the calculated coverage may be considered as not more than an approximation. Therefore, there is a need for a method to adapt state-of-the-art mathematical models for radio propagations so that the calculated path-loss predictions are as accurate as possible, despite the various sources of error noted before.

This thesis focuses on providing methods and tools to ??? the pointed drawbacks. Short summary of the introduced novelties??? It is important to note that some methods proposed in this thesis have been particularly designed for problems emerging in radio planning of 3G networks. Despite this, they may be adapted to other standards, e.g. GSM or LTE, without lose of generality.

## 1.1 Scope

In this thesis, we intend to contribute methods and tools that will provide solutions to the disadvantages pointed out in the previous section. The introduced methods and tools have a close correlation with the simulation and optimization of radio networks, especially those of the 3G and 4G. We will introduce the steps necessary to mitigate the problem of experimental reproducibility found in most published works, by simplifying setup, execution, and sharing of experimental results. With the development of the framework, we will evaluate and assess

the possibilities it offers as a support system for simulation and optimization problems of radio networks. By including parallel programming techniques for computer clusters and GPUs, we intend to go beyond the classical methodology provided by previous works, taking advantage of the inherent parallelism of some optimization techniques. We will also be looking into the application of many advances in HPC that should provide the framework with the computing power needed to improve the simulation process, thus enhancing its scalability to support real-world 3G radio networks. Finally, since we believe that this work will only become truly productive through the cooperation and long-term development of the scientific and engineering community, we will be releasing the source code, algorithms, documentation, and data to the public domain. This way, anyone will be able to use and to extend the framework for their own needs. Encouraging cooperation and sharing of experimentation-related tools and data should be a common goal from which everyone will benefit.

## 1.2 Hypothesis and aims

- The great proportion of published works about optimization problems for 3G radio networks is very difficult to reproduce, if possible at all.
- A common and open framework, designed for simulation of 3G radio networks, should mitigate the experimental reproducibility problem by simplifying sharing of experimental results.
- Parallelization techniques improve the scalability and time requirement of the framework, thus making it possible to process real-world data sets.
- Using hardware, specialized for parallel execution (e.g. GPU), improves the time requirement of parallel algorithms using threads or message-passing mechanisms.

### Aims

- Analyze the state-of-the-art of optimization problems for 3G radio networks in general, and UMTS in particular.
- Identify common obstacles in optimization problems for 3G radio networks that prevent them from being reproducible by other members of the research community.
- Design and implement framework tools to provide a common open environment that will enable the scientific community to easily share and reproduce different experimental conditions for maintenance and optimization of 3G radio networks.
- Integrate parallelization techniques to provide framework scalability, so that it will be able to deal with large real-world data sets.
- Evaluate the framework design by reproducing optimization environments and introducing new algorithms for tackling previously unsolved instances of optimization problems in 3G radio networks.
- Evaluate and analyze all experimental results and simulations in the context of real networks, using real-world data.

### 1.3 Methodology

The dissertation will use the following methodology to prove the hypothesis stated in the previous sections.

First, we will survey existing optimization problems and techniques for 3G radio networks in general, and UMTS in particular. The focus of the survey will be on optimization problems that do not provide the means required by experimental reproducibility and require the use of a snapshot-based model of the radio network.

Second, we will design and develop a parallel framework for radio network simulations. The development of the framework will focus on joining optimization and simulation, so that it will cover a wider range of use cases, namely as a maintenance and optimization tool for 3G radio networks. Additionally, we will make an intensive study of the mathematical models currently used, seeking to assess their reliability when used in simulations of 3G radio networks. Since the algorithms and simulations used in the optimization of 3G radio networks consume large amounts of data and require high computing power (due to the large number of simulations they need to run), we will improve the performance of the framework even more by adapting its most time-consuming parts for execution on GPUs.

Finally, we will evaluate the benefits of the above-mentioned framework by conducting reproducible experiments that will support the hypothesis outlined under ???, by tackling similar and bigger problem instances when solving both well-known and new optimization problems for 3G radio networks.

### 1.4 Contributions

The expected contributions of this dissertation to the fields of telecommunications and computer sciences include the following:

- State-of-the-art overview of optimization methods for 3G radio networks.
- Design and development of a framework that provides an open environment for radio network simulations, implemented for execution on computer clusters and GPUs. The framework will allow the scientific community to share a common domain to run the simulations needed by modern optimization methods, since most currently available simulation tools are proprietary and therefore unsuitable for experimental reproducibility.
- Improvement of quality and speed of renowned mathematical models, used for radio propagation predictions, by applying parameter optimization and parallelization techniques. The expected speed improvement should be of at least one order of magnitude.
- Proposal of a new algorithm, based on autonomous agents, to solve the service coverage problem. The solved problem instances should be of bigger size than ever solved in the literature and reach equal or better quality thereof. This should make our approach applicable for large real-world problem instances and data sets.
- Identification of a new optimization problem in 3G radio networks that deals with soft-handover alignment of downlink and uplink areas. By solving this problem, we should avoid abnormal network functioning in areas where there is soft-handover capability in the uplink, but none in the downlink. So far this problem has been solved manually by radio experts.
- Empirical comparison of the proposed metaheuristic algorithm against the existing state-of-the-art optimization algorithms on the soft-handover alignment problem.

## 1.5 Organization

The introduction provided in this chapter pretends to delimiter the context within which the dissertation will address.

The rest if this dissertation is organized as follows.

Chapter 2 present an overview of some well-known optimization problems that occur during deployment and configuration of mobile networks. A description of each optimization problem is given, followed by a short survey of recently proposed optimization methods. It closes by giving a conclusion regarding mobile networks and why they are a rich source of optimization problems.

Chapter 6 deals with the automatic tuning of parameters of the mathematical models, used for radio propagation predictions. Namely, based on field measurements, the configurable parameters of the mathematical models used by the framework may be automatically tuned to minimize the deviation from the prediction to the actual state of the network.

In Chapter 5, a static network simulator is used to find downlink and uplink SHO areas. By introducing a penalty-based objective function and some hard constraints, we formally define the problem of balancing SHO areas in UMTS networks. The state-of-the-art mathematical model used and the penalty scores of the objective function are set according to the configuration and layout of a real mobile network, deployed in Slovenia by Telekom Slovenije, d.d.. The balancing problem is then tackled by three optimization algorithms, each of them belonging to a different category of metaheuristics. We report and analyze the optimization results, as well as the performance of each of the optimization algorithms used.

Chapter ?? considers the problem of minimizing the total amount of pilot power subject to a full coverage constraint. Our optimization approach, based on parallel autonomous agents, gives very good solutions to the problem within an acceptable amount of time. The parallel implementation takes full advantage of GPU hardware in order to achieve impressive speed-up. We report the results of our experiments for three UMTS networks of different sizes based on a real network currently deployed in Slovenia.

# Part I

## Background and motivation



## 2 Background and motivation

Short introduction to the content of this chapter.

### 2.1 Optimization models

Optimization may be informally defined as the procedure of finding better solutions to a given problem that usually models some physical phenomenon. In our every day life, we are constantly solving small optimization problems, like choosing the shortest route to a friend's house, or organizing the appointments in our agenda. In general, these problems are small enough for us to find a good solution without extra help, but as they become larger and more complex, the aid of computers for their resolution is unavoidable.

Complex multidimensional optimization problems are popular in engineering, economics, physics and other scientific fields. When solving an optimization problem, the objective is to find a “good” solution in a “reasonable” computational time. In this respect, the field of mathematical optimization has received a lot of attention by the scientific community during the last decades. However, both “good” and “reasonable” are problem, application and context-specific concepts, in which the biggest challenge of selecting an appropriate optimization approach usually lays.

Mathematical optimization involves the process of finding solutions from a group of possible decisions, which may be defined as:

$$\min f(\vec{x}) \quad \vec{x} \in \Omega \subseteq \mathbb{R}^n, \quad (2.1)$$

where  $\vec{x} = (x_1, \dots, x_n)$  is a vector representing the decision variables,  $f(\vec{x})$  is the objective function measuring the quality of the decisions and  $\Omega$  is the set of feasible solutions of the problem, also known as search space. Note that the objective function  $f$  makes it possible to define a total order relation between any pair of solutions in the search space  $\Omega$ .

The search space  $\Omega$  may also be expressed as a solution to a system of equalities or inequalities, e.g.:

$$\begin{aligned} g(x_1, \dots, x_n) &\leq 0 \\ h(x_1, \dots, x_n) &= 0 \end{aligned} \quad (2.2)$$

Optimization problems involving the maximization of the objective function also fall into this category, since:

$$\max f(\vec{x}) = -\min(-f(\vec{x})) \quad (2.3)$$

A point  $\vec{x}^*$  is considered to be an unrestricted local minimum of a function  $f$  if it holds a better value than all its neighbours, i.e. there exists  $\epsilon > 0$  so that:

$$f(\vec{x}^*) \leq f(\vec{x}) \quad \forall \vec{x} \in \mathbb{R}^n \quad |\vec{x} - \vec{x}^*| < \epsilon \quad (2.4)$$

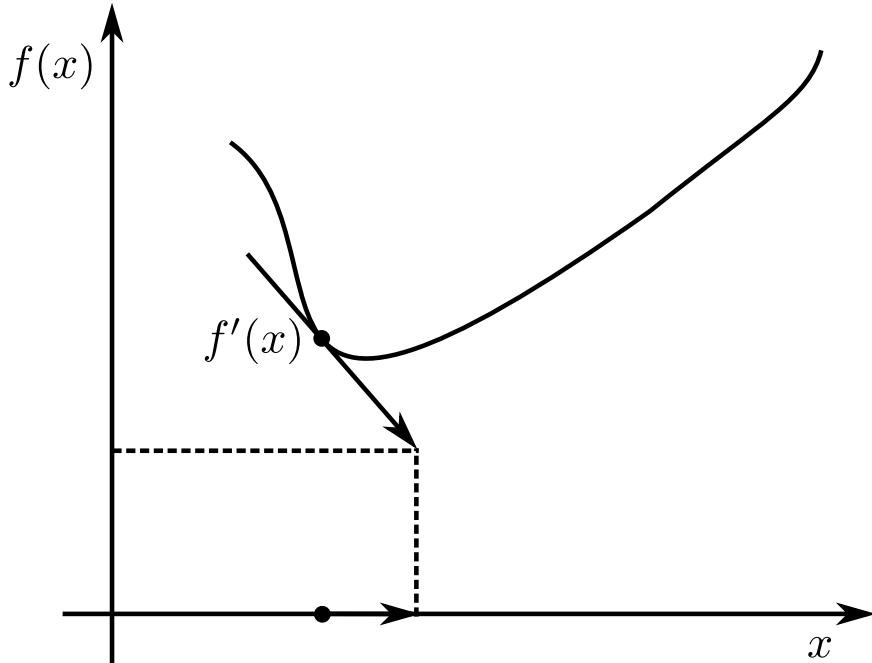


Figure 2.1: Gradient descent with a negative slope, i.e.  $x$  is increasing.

Similarly, a point  $\vec{x}^*$  is considered to be an unrestricted global minimum of a function  $f$  if it holds a better value than all others, i.e.:

$$f(\vec{x}^*) \leq f(\vec{x}) \quad \forall \vec{x} \in \mathbb{R}^n \quad (2.5)$$

The concepts of local and global minimum are considered strict if the inequalities of 2.4 and 2.5 are strict. Likewise, the definition of local and global maximum is given by the existing relation between a minimization and a maximization problem, as specified in 2.3, i.e. a point  $\vec{x}^*$  is a local or global maximum of a function  $f$  if and only if  $\vec{x}^*$  is a local or global minimum of function  $-f$ , respectively.

### 2.1.1 Gradient-based methods

Gradient-based methods are among the oldest and most studied optimization approaches. They are based on the derivative of the optimized function, using the first and even the second derivate of a function  $f$ . The name gradient follows from the derivative of multidimensional functions,  $\nabla f(\vec{x})$ , which is simply a vector where each element is the slope of  $\vec{x}$  in that dimension, i.e.  $\langle \frac{\partial f}{\partial x_1}, \dots, \frac{\partial f}{\partial x_n} \rangle$  [87].

The principle behind gradient-based methods is rather simple. Starting from an arbitrary value for  $x$ , we iteratively subtract (or add) a small positive value to it, e.g. for gradient descent:

$$x \leftarrow x - \alpha f'(x), \quad (2.6)$$

where  $\alpha$  is a small positive value. Consequently, a positive slope will make  $x$  decrease, whereas a negative slope will make it increase. Figure 2.1 shows an example of this behaviour. Therefore,  $x$  will gradually move down the function until it finds its minimum, where  $f'(x)$  is zero, causing it to stop.

However, gradient methods have certain drawbacks that make them unsuitable for tackling a wide range of optimization problems. Take, for example, the time they take to con-

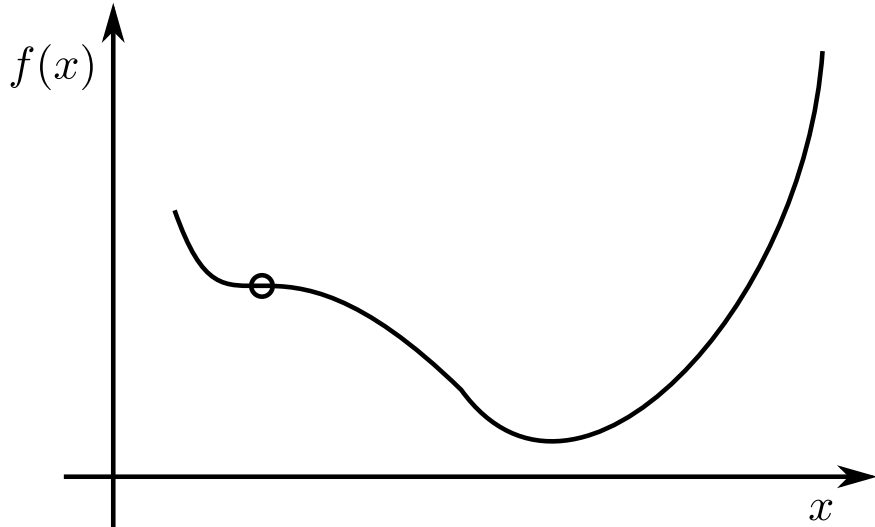


Figure 2.2: A saddle point or point of inflection, where the derivative is zero.

verge. As gradient descent approaches a function minimum, it will skip this point and land on the other side. In the next step, something similar will happen, but this time from the other side of the minimum point, thus slowly approaching to the target in a “zig-zag” way. This behavior is directly related to the slope of the function at the given point, i.e. a steepest slope translates into a larger jump, and may be alleviated by adjusting the value of  $\alpha$ . However, some functions (or regions of functions) may require smaller values, while for others a bigger value would be more appropriate. Newton’s method improves this by taking the second derivative of the function into account, i.e.:

$$x \leftarrow x - \alpha \frac{f'(x)}{f''(x)}, \quad (2.7)$$

thus adjusting the value of  $\alpha$  as it converges towards a point with zero slope [87].

Another issue is how other points are handled. Beside maxima and minima points, some functions also contain saddle points (known as inflection points in one-dimensional functions). Clearly, the first derivative of a saddle point is zero, meaning gradient descent will stop looking for the minimum, even though it hasn’t found it (see Figure 2.2). Newton’s method, on the other hand, does not help either. Moreover, in this case, we would be even dividing by zero! These observations clearly show how gradient methods get caught in local optima. We define local optima of a function as the optima (or minima in our case) of a local region. Similarly, global optima are defined as the optima of the whole domain of a function. It follows that gradient methods, as gradient descent or Newton’s method, are local optimization algorithms [87].

But maybe the biggest concern with gradient-based methods is they assume the function under optimization is derivable. This assumption holds only when optimizing a well-formed mathematical function. Unfortunately, this is generally not the case, since in most cases the gradient is not computable because the function is not known. The only available approach in such situations is creating inputs to the function in order to assess their quality. Metaheuristics are good candidates for this class of problems to solve moderate and large instances.

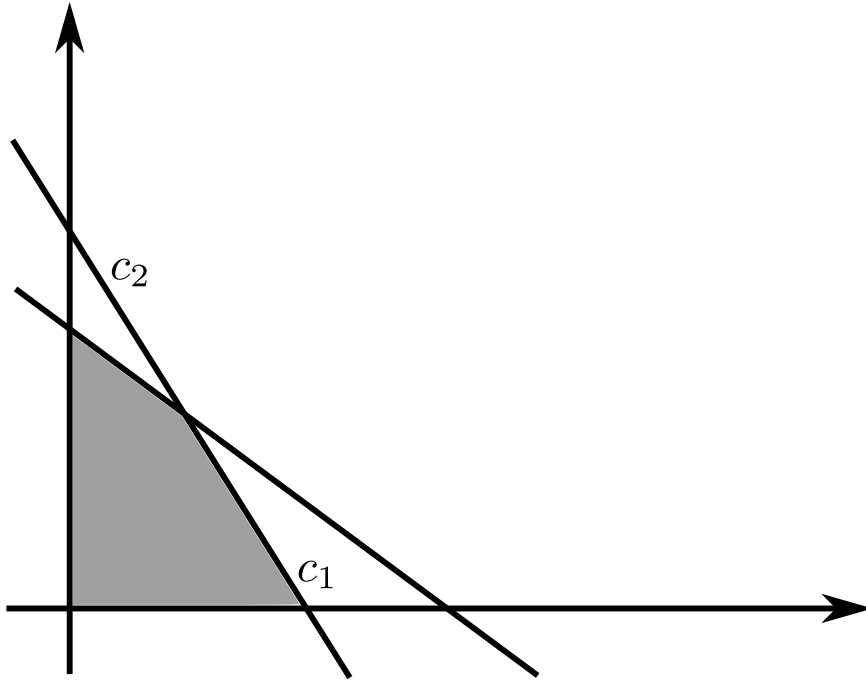


Figure 2.3: Graphical representation of a linear-programming example with two constraints,  $c_1$  and  $c_2$ . The greyed area is the polytope representing the region of feasible solutions.

### 2.1.2 Linear and non-linear programming

It was in the early 40s of the twentieth century, through the work of teams formed by mathematicians, economists and physicists, that the basis were established for the resolution of problems with a set of techniques known as linear and non-linear programming. Their initial goal was to solve different kinds of logistic problems during the second world war.

In a linear programming optimization problem, both the objective function  $f$  and a given set of constraints are linear functions. The constraints impose restrictions over  $\vec{x}$ , i.e. they must meet certain requirements as, for example, fullfil a limited availability of resources. An problem may be formulated as follows, e.g.:

$$\min f(\vec{x}) = c \cdot \vec{x} \quad (2.8)$$

subject to

$$\begin{aligned} A \cdot \vec{x} &\leq b \\ \vec{x} &\geq \vec{0} \end{aligned} \quad (2.9)$$

In the example above, the inequalities defined in 2.9 are the constraints to the linear program defined in 2.8.

For solving continuous linear optimization problems, efficient exact algorithms exist, such as the simplex method [36] or the interior-points method [75]. Indeed, linear programming is one of the most satisfactory models of solving optimization problems, since the feasible region of the problem is a convex set and the objective function is a convex function. It follows that the global optimum is a node of the polytope representing the feasible region [134]. See Figure 2.3 for a linear-programming example with several constraints.

Non-linear programming models, on the other hand, consider problems where the objective function  $f$  and/or the constraints are non-linear [13]. However, non-linear continuous

problems are more difficult to solve. Despite several existing techniques to linearize such models, they often not only introduce extra variables and constraints, but also some degree of approximation [51]. Moreover, some problem properties such as high dimensionality, parameter interaction, and multi-modality make these approaches ineffective.

Generally, when dealing with real-world problems, the availability of analytical optimization models, such as required by gradient methods or (non-)linear programming, is not guaranteed. Indeed, for some applications, only simulations or physical models are the available means for objective-function evaluation [43]. Once again, metaheuristics appear as good candidates to solve different instance sizes of this class of problems.

### 2.1.3 Metaheuristics

Metaheuristics, a term proposed by Glover in [52], represent a group of approximation algorithms designed to combine basic heuristic principles with advanced high-level guidance methods, targeted at improving the efficiency of a search process. These techniques are meant to find good solutions to a given problem, for which the mathematical function is not available or its search space is big enough for an exhaustive search to be unfeasible [79].

From the theoretical point of view, metaheuristics represent a subset of stochastic optimization, since they use some degree of randomness to find optimal (or as optimal as possible) solutions to hard problems. They are the most general of these kinds of algorithms, and are applied to a wide range of problems [87]. Unlike the exact optimization methods introduced in the previous sections, metaheuristics do not guarantee the optimality of the obtained solutions [134]. Moreover, they do not define how close the obtained solutions are from the optimal ones, as approximation algorithms do.

The characterization given by Blum and Roli [19] provides a clear overview of the fundamental properties associated with metaheuristics:

- metaheuristics are strategies that “guide” the search process;
- their goal is to efficiently explore the search space in order to find optimal or near-optimal solutions;
- they build upon techniques which range from simple local search procedures to complex learning processes;
- they are approximate and usually non-deterministic;
- they may incorporate mechanisms to avoid getting trapped in confined areas of the search space;
- their basic concepts permit an abstract-level description, which is not problem-specific;
- they may make use of domain-specific knowledge in the form of heuristics that are controlled by the upper level strategy;
- advanced metaheuristics use search experience (implemented as some form of memory) to guide the search process.

The strategies used by metaheuristics should provide a dynamic balance between the exploitation of the accumulated search experience (commonly called intensification) and the exploration of the search space (commonly called diversification) [19]. This balance provides the necessary means to quickly identify promising regions, and early discarding those which have already been explored or don't provide solutions of better quality. Promising regions

within the search space, which are identified by the obtained “good” solutions, are thoroughly explored during the intensification phase, hoping to find better solutions. On the other hand, during the diversification phase, not yet visited regions are explored, making sure the search space as a whole is evenly explored, thus avoid confining the search to a reduced number of regions. In this context, the ultimate search algorithm in terms of diversification is random search. Random search generates a random solution in the search space at each iteration, without using memory [134]. In terms of intensification, iterative local search is a representative algorithm. The steepest local search algorithm selects, at each iteration, the best neighboring solution that improves the current one [134].

Metaheuristics are applicable where state-of-the-art exact algorithms cannot tackle the given instances within the required time, either because of the size or the structure of the given problem instances. The meaning of “required time” within this context directly depends on the target optimization problem itself. A feasible or acceptable time may vary from some seconds to several months, again, depending on the target optimization problem, e.g. real-time decisions against structural-design problems.

Based on the characterization given by Talbi [135], let us summarize the essential properties of optimization problems that justify the use of metaheuristics:

- Very large problem instances. Even though exact polynomial-time algorithms might be known for solving the target problem, they are too expensive due to the size of instances.
- Problems with hard real-time constraints, where a “good solution” has to be found online. Metaheuristics appear as an alternative to exact algorithms in order to reduce the search time.
- A difficult problem of moderate size, which input instances have an intricate structure.
- Optimization problems with time-consuming objective function(s) and/or constraints. Indeed, various real-world optimization problems are characterized by the huge computational cost of the objective functions. Several radio-network design problems fall into this category.
- Problems that cannot be solved with exhaustive search due to the non-analytical models on which they are based. These problems are defined by a black-box evaluation of the objective function (see next section).

The influence of these conditions may increase in the presence of non-deterministic optimization models, e.g. problems with complex Monte Carlo simulations [33].

Undoubtedly, metaheuristics are rapidly gaining popularity as optimization problems are increasing in both size and complexity. Indeed, as the computing power of commodity hardware increases, the possibility of building models of greater complexity is available for developing more accurate models of real-world problems in engineering and science.

#### 2.1.4 Black-box optimization

A problem complexity is equivalent to the complexity of the best algorithm solving that problem [135]. If there exists a polynomial-time algorithm to solve a problem, we say the problem is easy or tractable. Similarly, if a problem is difficult or intractable, there is no known polynomial-time algorithm to solve it.

Many optimization problems cannot be formulated with a clear analytical mathematical notation. In such cases, the objective function may become a black box [74]. This is one of the main advantages when using metaheuristics, i.e. there is no need of a complete knowledge

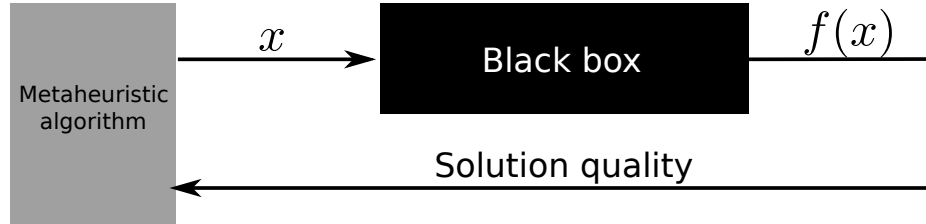


Figure 2.4: A metaheuristic algorithm process using a black box for the objective-function evaluation,  $f(x)$ , of a solution,  $x$ .

of the targeted model. Indeed, in a black box optimization, no analytical formulation of the objective exists [135], as Figure 2.4 shows.

More specifically, we say a function  $f(\vec{x})$ ,  $\vec{x} \in \mathbb{R}^n$ , is a black-box function if and only if [135]:

- the domain  $\vec{x}$  is known,
- it is possible to get the value of  $f$  for each  $\vec{x}$  based on simulation, and
- there is no other information available for function  $f$ .

Typically, the experiments associated with these kind of problems are very expensive in terms of time and cost, since a simulation must be forced to evaluate the solution. Generally speaking, the most time-consuming part of a metaheuristic optimization process is the evaluation of the objective function [134]. This is especially true when dealing with real-world problems of areas such as structural design [12], molecular docking [136] and, the field on which this thesis focuses, radio-network design [15]. A possible substitution for lengthy evaluations is to reduce their complexity by approximating the objective function, thus replacing the it with an approximation during the optimization process. This approach is known as meta-modeling [134]. However, when dealing with approximations, some degree of solution quality is inevitably sacrificed. As we will show in the following chapters, there is a very fine balance between the number of evaluations and the quality of the achieved solutions. Consequently, reducing the time spent in objective-function evaluation should favorably influence the solution quality achieved by a preferred metaheuristic algorithm. A major portion of this thesis is dedicated to improve this specific aspect on the area of radio-network optimization.

In practice, black-box evaluation of the objective function presents an inherent problem. In the context of research works related to radio-network optimization, there is an increasingly habit of not providing the black-box used to evaluate a given approach, and that leads to the results provided. A quick review of the state-of-the-art in radio-network optimization indicates that this fact has become increasingly regular in several published works [refs??]. This fact creates a barrier to one of the most important phases of scientific methodology: experimental reproducibility [46].

There are several reasons why the situation has reached this point. It is a known fact that proprietary software, providing good computational models for radio-network simulation, is a very expensive tool for science. Even neglecting the economical aspect, but considering the great variety of software packages and license combinations, it is practically impossible for a research laboratory to have at its disposal the whole palette of commercially-available solutions. Moreover, genuine users of these applications are generally not allowed to mention the formats, protocols, or algorithms used by the proprietary software, since their disclosure is explicitly forbidden by the commercial licenses.

If we turn our attention to non-commercial and open-source packages for radio-network simulation [refs!???], we would quickly run into mainly two difficulties: poor documentation and low scalability. The scalability issues shown by some projects [refs!???] retrain the packages to be used in real-world environments, where big problem instances are the rule. Despite this, it is a huge merit and acknowledgement to the authors of this packages, not only for providing it to the scientific community, including their source code, but fundamentally because providing an environment in which different kinds of simulations are completely reproducible. Regarding the lack of documentation, it represents a big hurdle when extending the base code, which becomes a difficult task without the help of the original authors of the package, who have a deep knowledge of the code-base. This knowledge is required in order to effectively expand the functionality of the open-source tool in question. This is especially true when dealing with complex simulation frameworks, as the ones used for radio networks.

### 2.1.5 Metaheuristic algorithms

Related literature groups metaheuristic algorithms due to their behaviour. For example, the following is a list of fundamentally different optimization algorithms, namely:

- differential evolution, from the family of evolutionary algorithms;
- differential ant-stigmergy algorithm, from the family of swarm-intelligence algorithms; and
- simulated annealing, from the group of classic metaheuristic algorithms, targeted at combinatorial optimization problems.

Each of these algorithms shall minimize an objective-function value by adopting essentially disparate approaches, hence the diversity of applying algorithms belonging to different families to solve the same optimization problem. In this way we want to find out whether any of the presented approaches is better suited for solving our problem.

In the following sections we give a short introduction about their functioning and controlling parameters.

### 2.1.6 Differential evolution

Differential evolution (DE) [129] is a simple and powerful evolutionary algorithm proposed for global optimization. A wide range of optimization problems have been solved by applying DE [37]. The algorithm exhibits a parallel direct search method, which utilizes  $D$ -dimensional parameter vectors. The balancing problem is expressed in each component of a vector  $X$  of the population, which maps to the CPICH power of one cell under optimization:

$$X_{aG} = \{x_1, x_2, \dots, x_i, \dots, x_D\}, \quad (2.10)$$

where  $x_i \in P_i$  represents a candidate CPICH power setting of cell  $i$ , and  $G$  indicates the generation of an individual  $a$  in the population. Since there are  $|N|$  cells in the mobile network, it follows that  $D = |N|$ .

In each generation, DE produces new parameter vectors by adding the weighted difference between two population vectors to a third one [129]. The resulting vector is retained if it yields a lower objective function value than a predetermined population member; otherwise, the old vector is kept.

There are different variants of DE. We have chosen the most popular one to solve our optimization problem, called *DE/rand/1/bin*. The nomenclature used to name this variant indicates the way the algorithm works:

- *DE* denotes the differential evolution algorithm,
- *rand* indicates that the individuals selected to compute the mutation values are randomly chosen,
- *1* specifies the number of pairs of selected solutions used to calculate the weighted difference vector, and
- *bin* means that a binomial recombination operator is used.

We considered four parameters to control the search process of DE: the population size, the maximum number of generations for the algorithm to run, the crossover constant, and the mutation scaling factor.

An extensive description of DE and its variants may be found in [105].

### 2.1.7 Differential ant-stigmergy algorithm

Based on the metaheuristic Ant-Colony Optimization (ACO) [39], the differential ant-stigmergy algorithm (DASA) [80] provides a framework to successfully cope with high-dimensional numerical optimization problems. It creates a fine-grained discrete form of the search space, representing it as a graph. This graph is then used as the walking paths for the ants, which iteratively improve the temporary best solution.

The mapping between the balancing problem and DASA is similar to the one depicted in Equation (2.10):

$$X_a = \{x_1, x_2, \dots, x_i, \dots, x_D\} \quad (2.11)$$

In this case, each ant,  $a$ , creates its own solution vector,  $X_a$ , during the minimization process. At the end of every iteration, and after all the ants have created solutions, they are evaluated to establish if any of them is better than the best solution found so far.

There are six parameters that control the way DASA explores the search space: the number of ants, the discrete base, the pheromone dispersion factor, the global scale-increasing factor, the global scale-decreasing factor, and the maximum parameter precision.

For a more in-depth explanation about these parameters and the DASA algorithm itself, we refer the reader to [80].

### 2.1.8 Simulated annealing

As the third optimization algorithm to tackle the balancing problem we have chosen simulated annealing (SA) [77], a classic metaheuristic algorithm often used when the search space is discrete. SA has proved to be a solid optimization algorithm, capable of giving high-quality solutions to a wide scope of optimization problems [130].

At each time step during the process, the system under optimization is in a given *state*. The objective function maps a system state to a value known as the *energy* of the system in that state. A *move* in the search space represents a change in the state of the system. After making a move, the system may exhibit lower or higher energy, depending on the results of the objective function. When dealing with minimization problems, a better state always describes lower energy than the previous one.

SA incorporates the notion of *temperature*, by which the probability of moving the current state of the system into a worst one is lowered as the temperature decreases. Exploration of the search space is thus induced at higher temperature, whereas exploitation appears at lower temperature, when only improving moves are accepted.

Table 2.1: Pseudo-code: a move in the search space of SA.

Step	
1	$i' = \text{random cell}(N)$
	<b>do</b>
2	<i>if rand() &lt; 0.5 then</i> $p_{i'}^{\text{NEW}} = p_{i'} + 0.01$
	<i>else</i> $p_{i'}^{\text{NEW}} = p_{i'} - 0.01$
3	<b>while</b> $p_{i'}^{\text{NEW}} \notin P_{i'}$
4	$p_{i'} = p_{i'}^{\text{NEW}}$

Table 2.1 shows the pseudo-code of a move in the search space of possible CPICH power settings, resulting in a new state of the system.

At the first step, a cell,  $i'$ , is randomly selected from the set of all cells in the network,  $N$ . In step 2, a change of +0.01 dB or -0.01 dB is applied with 50% probability to  $p_{i'}$ . The current CPICH power of cell  $i'$  is expressed in dBm. The randomly generated CPICH power setting,  $p_{i'}^{\text{NEW}}$ , is checked for validity in step 3, i.e. it must be an element of the set  $P_{i'}$ . If  $p_{i'}^{\text{NEW}}$  is not a valid CPICH power, step 2 is executed again, generating another random CPICH power. Finally, in step 4, the CPICH power of cell  $i$  is replaced by  $p_{i'}^{\text{NEW}}$ .

It is important to note that, as long as  $|P_{i'}| > 1$ , the algorithm shown in Table 2.1 shall never be trapped in an endless loop. On the other hand, if  $|P_{i'}| < 2$ , there are no candidate CPICH powers for cell  $i'$  and thus no possibility of optimization by means of CPICH power adjustment.

Notice also that the acceptance of a move in the search space is left to SA and its stochastic components.

## 2.2 Optimization of radio networks

Once a radio network is launched, an important part of its operation and maintenance deals with monitoring its quality characteristics and making changes in the configuration of the deployed equipment and parameter values in order to improve its overall performance.

The evolution from 3G to 4G has introduced not only the technology needed to increase data capacity and voice quality, but also a greater complexity in terms of network planning, deployment, and configuration, which have rendered some of the traditionally used methods to be ineffective. In a traditional approach (i.e. manual), during the network planning and maintenance processes, a radio-network simulation software executes the analysis, while an engineer makes the change decisions. Therefore, a radio-planning engineer adapts the network parameters manually and the network-planning tool analyzes the given configuration. If the obtained results are not acceptable, the analysis process has to be repeated several times, until a given goal is achieved.

The complexity of radio-network has grown even faster than their throughput capacity, thus making it impossible to plan 4G radio networks with traditional methods. In this sense, an examination of colored coverage maps in conjunction with some statistical analysis are no longer appropriate tools for troubleshooting a network. Moreover, since real-world 4G radio networks are large and many of their configuration parameters are interdependent, an engineer is not able to cope with the level of complexity present in such systems. For that reason, the computer, along with specialized software, guides the engineer to the most appropriate configuration for the network. In the context of this work, we will refer to this process as radio-network optimization.

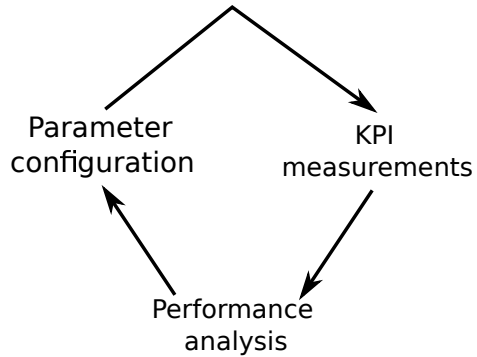


Figure 2.5: Typical optimization cycle for radio networks. This sequence is repeated until the achieved results are acceptable.

Radio-network optimization may be divided into two fundamental phases: analysis and decision [95]. The analysis phase consists on the examination of network performance, which mainly focuses on the definition and collection of several Key Performance Indicators (KPIs). KPIs are quantifiable measurements, agreed to beforehand, that reflect network quality factors. The second phase deals with the decision making, based on the analytical results collected in the previous phase, about the configuration of a particular configuration or parameter setting. This process, depicted in Figure 2.5, is repeated until the achieved results are acceptable.

A common limitation of several optimization methods appearing in the literature [ref!???] is their inability to meet the requirements needed by real-world radio networks, since their computational-time complexity make them unfeasible for practical use in industrial-sized scenarios. Particularly, this limitation is more common in implementations targeting traditional computer architectures of sequential execution [ref!???]. When analyzing real-world radio networks with thousands of transmitters it is necessary to reduce the execution time of the optimization processes, so that they are useful for practical use.

## 2.3 Survey of optimization problems for radio networks

Since radio networks are increasingly more sophisticated, the need for optimization methods, capable of coping with greater complexity, is far from declining. It has been established that most radio-network optimization problems are NP-hard, since the computational time grows non-polynomially as the problem size increases [7, 9, 56, 61, 83, 108, 123]. Moreover, there are other reasons directly related with the evolution of already deployed networks that greatly increase the need for optimization methods, as described in [95]:

**Network performance improvement** more users receive service coverage with the same physical infrastructure, making parameter optimization the less expensive and only viable short-term approach.

**Changes in users' profile** the introduction of new and faster services puts additional stress on the infrastructure, requiring additional optimization efforts.

**Changes in the propagation conditions** the allocation of a different frequency band for LTE compared to UMTS requires deployment of 4G sites, which radio propagation behaves differently than in 3G networks, mostly in urban areas.

In this context, network operators define different optimization targets, depending on the addressed optimization problem. These targets are formed by an objective function that

maps possible configurations into a real value. This number represents a quality rating of the proposed solution and it is used to compare different solutions among each other, and ultimately select the best one. Unfortunately, there is no definitive objective function in the field of radio-network optimization [95]. However, it is possible to optimize for different targets such as service coverage, base station locations, interference levels, etc.

In this work, we will address network optimization methods that are performed “off-line”, meaning that the optimization software is not an active functioning part of the radio network in operation. Statistical data about network operation is used as the input or feedback information for different optimization targets.

This paper gives an overview of well-known optimization problems in 3G mobile networks. At the beginning of each of the following sections, a description of an optimization problem is given, followed by a short survey of recently proposed optimization methods. Finally, a discussion about the introduced methods is given, before closing with some concluding remarks.

### 2.3.1 Optimizing base station locations

#### 2.3.1.1 Problem formulation

Some references [62, 140, 91] formulate the base station location problem in terms of the minimum set covering problem (shown in Figure 2.6). The coverage problem is defined by considering the signal level in every test point from all base stations and requiring that at least one level is above a fixed threshold.



Figure 2.6: The minimum set covering problem: (a) the problem input and (b) the solution.

A different formulation considers the site selection problem as a  $p$ -median problem, in which base station location is the only decision variable considered. To each of the candidates solutions, an installation cost is also associated. The  $p$ -median problem constitutes seeking  $p$  different locations each time, regardless of how distant the sites are. The problem is to select one candidate site from each region to install a base station such that the traffic capacity and the size of the covered area are maximized with the lowest installation cost.

#### 2.3.1.2 Proposed solutions

Aydin et al. [11] propose a solution to the  $p$ -median problem based on three meta-heuristic algorithms; a genetic algorithm, simulated annealing, and tabu search. Their experimental study focuses on performance comparison between the three algorithms.

A solution to the set covering problem is proposed by Hao et al. [62]. A simulated annealing implementation was developed to solve the formulated combinatorial problem. The results presented demonstrate the feasibility of the proposed approach. Tutschku [140]

presents a specialized greedy algorithm to solve the same problem. This work is part of the implementation of a planning tool prototype. Mathar and Niessen [91] propose a solution based on integer linear programming, which they claim finds optimal solutions in most cases. They also introduce simulated annealing as an approximate optimization technique. This approach substitutes linear programming whenever an exact solution is out of reach because of the complexity of the problem.

Amaldi et al. [?] offer a discussion about the computational results of two different heuristics: greedy search and tabu search. The problem formulation is based on a set of candidate sites where the base stations can be installed, an estimation of the traffic distribution and a propagation description of the area to be covered. Some years later, the same authors [?] extended the problem formulation by also considering base station configuration and hardware characteristics. In both works, they propose a mixed integer programming model with which they aim to maximize the trade-off between total traffic covered and total installation costs. The only difference between the models is the constraint definition of the linear program, where the constraints in [?] are a subset of the constraints in [?].

Finally, Whitaker et al. [107] focus on providing the required service coverage at the lowest possible financial cost. Their framework supports the use of any multiple objective optimization algorithm which seeks to approximate a Pareto front. The performance of four different algorithms is explored, namely SEAMO, SPEA2, NSGA-II and PESA.

### 2.3.2 Optimizing antenna parameters

There are many antenna parameters that control the coverage and interference in the network, since the antenna shapes the emitted energy. Two important parameters are the azimuth angle and the elevation angle (or tilt) of the antenna. The antenna azimuth (shown in Figure 2.7) is the direction in which the main beam of the horizontal pattern points [65]. The antenna tilt (shown in Figure 2.8) is defined as the angle of the main beam of the antenna relative to the horizontal plane [65]. Both of these parameters have a great influence on network quality, although antenna tilt requires less effort to implement, since most modern radio networks already support remote electrical tilt. The adjustment of these two parameters optimize some important aspects of the network, namely:

- path loss between the base station and the mobile phone, since less power is required for a connection, hence more power is available for traffic; and
- interference between neighboring cells, which leads to an overall capacity increase.

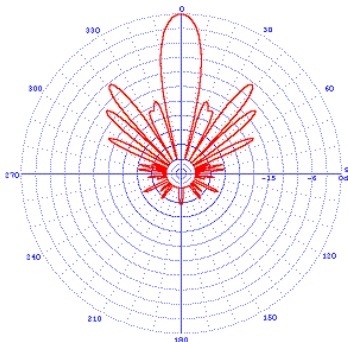


Figure 2.7: A typical antenna azimuth pattern.

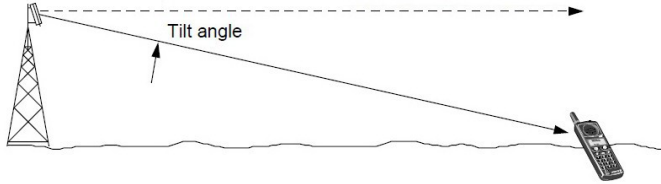


Figure 2.8: The antenna tilt angle with the horizontal plane.

### 2.3.2.1 Proposed solutions

Karner [76] proposes an “ad-hoc” strategy for adjusting antenna azimuth and downtilt by analyzing the structure of the network. The objective of this optimization is to improve the results presented in [48] by increasing the number of served users in the target area. In a similar line of work, Jakl [72] included in his doctoral thesis an antenna azimuth optimization algorithm, based on attempts of avoiding coverage holes by properly adjusting the azimuth settings.

Siomina and Yuan [118] propose a framework for automated optimization of antenna azimuth and tilt, including both mechanical and electrical tilt. The implementation introduces a simulated annealing algorithm that searches the solution space of possible antenna configurations. The goal of the optimization is targeted to address power sharing among cell channels and ultimately improve High-Speed Downlink Packet Access (HSDPA) [151] throughput.

Zhang et al. [55] present a method which is composed of two optimization loops: the inner one and the outer one. The inner loop concentrates on frequency planning while the outer loop focuses on finding the optimal setting of antenna azimuth and tilt for the current solution delivered by the inner loop. Although frequency planning is not directly related to 3G network optimization, we found this approach interesting enough to make a reference to it. The inner loop could be easily replaced with some other optimization target, e.g. common pilot channel power setting.

### 2.3.3 Optimizing common pilot channel power

The Common Pilot Channel (CPICH) is used as reference for handover [141], cell selection [1], and cell reselection [42]. Whenever a mobile phone is switched on, it tries to register with the cell providing the highest received CPICH level. It also defines the effective coverage area of the cell: according to the CPICH power level, the cell coverage area will enlarge or shrink. Consequently, by appropriately adjusting the CPICH power at the base stations, the number of served users per cell can be balanced among neighboring cells. This procedure is called load balancing and it reduces interference, stabilizes network operation, and facilitates radio resource management [65].

#### 2.3.3.1 Proposed solutions

Chen and Yuan [? ] optimize CPICH transmit power starting from a uniform allocation (i.e. all cells are set with the same transmit power level). Their objective is to enhance HSDPA performance at cell edges while preserving control of R99 soft handover [152]. The solution approach is based on a linear-integer mathematical model. A very similar problem and proposed solution is presented by Siomina and Yuan [121]. The problem definition

slightly differs from [? ] as there is no reference to soft handover control. The solution is also implemented as a linear program.

Olmos et al. [44] apply simulated annealing to the optimization of CPICH transmit powers in order to force mobile phones to transmit to the best available cell in a service area. Their results show decreased cell load with a consequently increased network capacity.

Chen and Yuan [? ] present a two-phase optimization algorithm for large-scale mobile networks. The algorithm uses the total CPICH power consumption as a minimization objective. The authors claim that the tabu-search-based algorithm can compute near optimal solutions within a few seconds, even for large networks.

### 2.3.4 Optimizing CPICH and antenna parameters

Both the antenna tilt and azimuth directly affect the direction and range in which the cell broadcasts its CPICH. Consequently, optimal CPICH power is highly dependent on how the antenna tilt and azimuth are configured at base stations. Ideal configuration of antenna tilt and azimuth network-wise, with the objective of optimizing the CPICH power consumption, is a challenging task. For this reason, many authors have considered optimization methods that address all three parameters.

#### 2.3.4.1 Proposed solutions

Varbrand et al. [115] approach the optimization of service coverage (see section 2.3.5) by considering all three configuration parameters, namely: CPICH transmit power, antenna tilt, and antenna azimuth. Their simulated annealing algorithm searches the solution space of possible configurations in order to find improvements in network performance and total transmitted power. Interestingly, the algorithm is efficient enough to optimize large networks without using excessive computing resources.

Siomina [120] combines optimization of base station antenna tilts and CPICH power to reduce the total interference level and to improve network capacity. The introduced algorithm optimizes the antenna downtilt setting so that the total CPICH power in the network is minimized.

Neubauer et al. [48] present two optimization algorithms for finding an optimal setting of antenna tilt and CPICH power of the base stations. The first algorithm builds on a rule-based approach, while the second one extends it by incorporating simulated annealing. An evaluation of both techniques shows that the second algorithm returns better results.

Jakl et al. [49] proposed a problem-specific genetic algorithm to tackle the optimization of antenna tilts and CPICH transmit powers. The goal of the optimization is to increase network capacity. The implementation involves a deterministic fitness selection scheme, a problem specific recombination operator and an improved mutation operator. After the initial identification of the best individuals, a local optimization technique is used to improve their fitness.

### 2.3.5 Optimizing coverage

Coverage is maybe the most common optimization objective considered in 3G network optimization. The objective function for coverage optimization may be defined as follows:

$$f_{\text{cov}} = \frac{A_{\text{covered}}}{A_{\text{total}}}$$

where  $A_{\text{covered}}$  represents the area covered by the network and  $A_{\text{total}}$  represents the total area under optimization. Thus, the expression  $f_{\text{cov}}$  represents the portion of the total area

that is actually under network coverage. This value ranges from 0 (no coverage) to 1 (total coverage).

The area being optimized is usually divided in squares (or pixels) of a certain size, creating a regular square grid (RSG) of a certain resolution. A pixel is considered covered if the signal to noise ratio is above a given threshold [2]. It is also common to use a test function  $cov(x, y)$ , which returns 1 if the pixel located at  $(x, y)$  is covered, and 0 otherwise.

### 2.3.5.1 Proposed solutions

Siomina and Yuan [124] consider the problem of minimizing pilot power subject to the coverage constraint. Their approach consists of mathematical programming models and methods, based on a linear-integer mathematical formulation of the problem. A special numerical analysis studies the trade-off between service coverage and pilot power consumption for different test networks.

Capone et al. [?] investigate mathematical programming models for supporting decisions on where to install new base stations and how to select their configuration (antenna height and tilt, sector orientations, maximum emission power, pilot signal, etc.) so to find a trade-off between maximizing coverage and minimizing costs. The overall model takes into account signal-quality constraints in both uplink and downlink directions, as well as the power control mechanism and the CPICH signal.

Valkealahti et al. [144] propose a method for automatic setting of CPICH power. The control algorithm applies total transmission power measurements from the base station, neighboring cells and mobile terminals to determine the pilot qualification. The CPICH power is then periodically updated based on a group of heuristic rules in order to improve coverage and load balance. A similar approach is described by Parkinnen et al. [143] where some network performance parameters are combined with service coverage in a cost function. The pilot power of a cell is periodically updated with a gradient descent method that minimizes the afore mentioned function.

### 2.3.6 Optimizing alignment of soft-handover areas

We have defined this new problem. About the connecting of this 3G-specific problem with LTE ...???

### 2.3.7 Discussion

In the field of 3G network optimization, comparing the efficiency of different optimization methods has proven to be a very difficult task. There are several reasons for this, among which we have identified the following:

- The networks on which the experiments are carried out are not standardized. Thus, it is problematic to set up an environment in which the presented results may be reproduced.
- There are no “open” implementations of 3G mobile network simulators. On the contrary, there is a vast variety of proprietary (or “closed”) network simulators that only increase the ambiguity regarding the experimental environment used for a certain network layout and configuration. Moreover, some of the “closed” network simulators do not even account on the built-in path loss estimation methods used.
- Only a small part of the referenced optimization methods perform their experiments on real-life networks, that have been already deployed and are currently running. This

fact creates a gap between the research field and the industry (i.e. the application area) and it is counterproductive for both the researchers and the network operators, since they don't benefit from mutual collaboration.

We believe that the creation of a standardized framework would be a great benefit in the field of 3G network optimization, as it would allow researchers to compare different methods, results and solutions in an easy, fast and objective manner. An effort in this direction is the MOMENTUM project [71]. It was created with the objective of setting up a standardized experimental environment that would facilitate research cooperation and would ultimately benefit from an improvement on the research field of 3G networks. The project includes complete data about the terrain, properties of the service area and hardware used in three different mobile networks. It offers the researcher detailed information about real-life networks based in Berlin, Lisbon and The Hague. As part of the package, there is also a Java application programming interface (API) that eases the implementation of some conventional tasks such as data parsing. The main focus of project is on data, thus there is no network simulator included. In any case, the researcher benefits from several included traffic snapshots that help assessing different network configuration settings. There have been no updates in the MOMENTUM project since 2005, when the last data corrections were introduced [71].

In the context of this survey and after reviewing many different papers on 3G network optimization, we can say that the MOMENTUM project has not been widely adopted by the scientific community. Moreover, several of the reviewed works do not offer detailed instructions on how to resemble the experimental environment. Consequently, it is very complicated (if not even impossible) to reproduce the presented results. Therefore, the selection of optimization methods, based on the results presented by their corresponding authors, is virtually meaningless, since the results are not comparable with each other. For this reason, we have decided to concentrate the following discussion on the optimization methods used, leaving the results aside.

### 2.3.8 Discussion about the presented methods

Regarding the optimizations methods presented in previous chapters, three distinctive groups emerge: genetic algorithms, linear programming and other search methods.

**Genetic algorithms** These algorithms work on a population of solutions that allows a more comprehensive search for optimal solutions. As a direct consequence, an increase in running time is commonly observed. The implementation effort of genetic algorithms is to some degree higher than for simpler search methods (e.g. local search), but their inherent structure greatly simplifies possible parallel implementations and execution.

**Linear programming** Linear optimization problems are widely used in different optimization areas and there are many good software packages to solve such problems. Consequently, if a problem can be modeled as a continuous linear problem, there is usually no difficulty in finding optimality. In the context of this survey, linear programming has proven useful for coverage optimization in early network planning stages.

**Other search methods** Other search methods<sup>1</sup> usually represent a compromise between running time and quality of results. They rely on evaluating a great number of alternative configurations. The number of parameters taken into account, as well as

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<sup>1</sup>Namely, local search, simulated annealing and tabu search in the context of this survey.

Table 2.2: A comparison among the presented optimization methods.

Algorithm	Running time	Typical application
Local search	Shorter	Solution quality improvement.
Tabu search	Shorter	Solution quality improvement.
Simulated annealing	Longer	Initial search of the solution space.
Genetic algorithm	Longer	Initial search of the solution space and solution quality improvement.
Linear programming	(formulation dependent)	Coverage network planning.

the evaluation precision, directly influence their running time. These methods don't excel in full simulation scenarios. On the other hand, some search methods (e.g. tabu search) have powerful mechanisms to escape local minima.

A short comparison of the optimization methods presented in this survey is shown in Table 2.2. The comparison variables arise from the context in which the methods were presented.

## Summary

The variety of optimization problems that have been introduced in the previous sections differ in many aspects like implementation, running time and solution quality. Picking the right method for a given situation depends on the optimization task and the desired results. Since computation time is usually an important restriction, simpler and faster methods may be preferable.

Beside the convenience of a survey such as the one presented in this work, it is very important to develop a feeling for the properties, advantages and drawbacks of the respective methods. Moreover, radio expert's recommendations regarding solution interpretation and feedback from everyday network operation, are an essential input for creating quality optimization methods. In this sense, and based on our own experience, the expert's advice is irreplaceable and a most valuable contribution to the research work.

Also, as it may be observed in some of the presented references, it is often advisable to combine different methods. Therefore, a simple optimization method may find a subset of reasonable parameter configuration, whereas a more complex method could be applied afterward, to refine the search. Sometimes it may also be useful to apply a simple search method at the end to find better solutions in the vicinity of a current promising one.

## 2.4 Principles of mobile radio networks

### 2.4.1 Quality of service

QoS...???

### 2.4.2 Handover and soft-handover

In mobile networks, handover is one of the main features that allows user's mobility [65]. The concept behind the handover operation is simple: when a user moves from the coverage area of a cell to the coverage area of a neighboring cell, the system creates a new connection with the latter cell and disconnects the user from the former one, while keeping the current connection active. Soft-handover (SHO), on the other hand, is a possibility available in mobile networks using the Wideband Code Division Multiple Access (WCDMA) technology, which the UMTS employs. SHO enhances handover functionality by allowing a user to

potentially operate on multiple radio links in parallel. Since different users are separated by unique spreading codes, the detection of single user's signal is implemented by despreading with the same code sequence used in the transmitter [65].

Every mobile terminal constantly monitors the common pilot power channel (CPICH) of the connected cell and its neighbors. The information about these measurements is sent to the network by the user terminal (i.e. mobile). The SHO condition depends on the relative received signal quality from different cells and the SHO window, which triggers the addition of a cell to the user's active set. Depending on radio propagation characteristics and different transceiver capabilities, the radio transmission can gain more than 3 dB out of a SHO situation [65]. From this point of view, SHO is a method to reduce interference and improve radio quality, particularly at the cell border where radio coverage is of inferior quality. In UMTS Release 99 [138], SHO is specified to work from the network towards the user (i.e. downlink), and from the user towards the network (i.e. uplink).

With the introduction of High Speed Packet Access (HSPA) as an improvement of the performance existing in WCDMA protocols, the role SHO plays in mobile network configuration and functioning slightly changed. The key difference is that High Speed Downlink Packet Access (HSDPA) does not support SHO, whereas the High Speed Uplink Packet Access (HSUPA) does. This particular distinction is discussed in Chapter 5, since it has some key implications in the balanced distribution of SHO areas, and thus in the quality of HSPA services [67].

#### 2.4.3 Pilot signal and power

The CPICH transmit power is typically between 5% to 10% of the total downlink transmit power of the base station [81], but there is no standardized method to find a CPICH power setting.

The CPICH transmits in the downlink of a UMTS cell system. The transmit power is usually between 5% and 10% of the total power available at the base station [66]. The capacity of a cell is limited by the amount of available power at the base station and the interference level at the mobile terminal. The coverage area of any cell is controlled by changing its pilot power, which consequently modifies the service area of the network.

The CPICH transmit power is common to many different planning and optimization problems in UMTS networks [96].

## 2.5 Principles of GPUs

### 2.5.1 OpenCL

We have chosen the Open Computing Language (OpenCL) [127] as the implementation platform of our optimization system on GPU.

OpenCL is an open parallel computing API designed to enable GPUs and other coprocessors to work together with the CPU, providing additional computing power. As a standard, OpenCL 1.0 was released in 2008, by The Khronos Group, an independent standards consortium [94]. For additional information about the OpenCL standard and API, we refer the reader to the numerous guides available online.

Our choice in using OpenCL was greatly influenced by the fact that its bitcode runs on a variety of hardware, including multicore CPUs and GPUs from different vendors. This provides a complete framework capable of comparing execution speed-up on different hardware without the need of changing the implementation.

Table 2.3: *Terminology translation between OpenCL and CUDA [78].*

<b>OpenCL</b>	<b>CUDA</b>
Grid	Grid
Work group	Block
Work item	Thread
__kernel	__global__
__global	__device__
__local	__shared__
__private	__local__
image2d_t	texture<type,n,...>
barrier(L M F)	__syncthreads( )
get_local_id(0 1 2)	threadIdx.x y z
get_group_id(0 1 2)	blockIdx.x y z
get_global_id(0 1 2)	(not implemented)

One unfortunate consequence of the vendor variety is that NVIDIA’s CUDA [100] and OpenCL documentation present disparate naming conventions for some key components. For the sake of consistency, in Table 2.3, we present a short “translation dictionary” between them. In the remaining of this work, we will stick to the naming convention used in the CUDA documentation.

Despite the use of OpenCL as the target platform for our implementation, the details described in the next sections may be equally applied on CUDA.

### 3 A parallel framework for radio-coverage planning and optimization

In this chapter, a parallel radio-coverage simulation framework is presented. The objective of the framework is to provide an environment for the radio-coverage prediction of large radio networks with several cells. Due to its high performance, the framework also enables the evaluation of larger and more complex optimization problems for radio networks.

The framework is implemented as a module of a geographical information system, since the prediction calculation employs digital elevation models and land-usage data in order to analyze the radio coverage of a geographical area. Following a classic master-worker parallel paradigm over a message-passing communication model proved to be a bottleneck for the performance of the parallel module. A new approach, that overcomes this performance constraint, is introduced in this chapter. The efficiency improvement is based on overlapping process execution and communication in order to minimize the idle time of the worker processes and thus improve the overall efficiency of the system. To this end, the intermediate calculation results are saved into an external database (DB) instead of sending them back to the master process. This approach is implemented as part of a parallel radio-prediction tool (PRATO) for the open-source Geographic Resources Analysis Support System (GRASS) [99]. An extended analysis of the experimental results is provided, which are based on real data from an LTE network currently deployed in Slovenia. Based on these results, which were performed on a computer cluster, the new technique exhibits better scalability than the traditional master-worker approach. Some real-world-sized data sets are presented, the radio-coverage predictions of which are calculated in a shorter time while saturating the hardware utilization.

The content of this chapter extends the research work published by the author in [14]. The rest of this chapter is organized as follows. Section 3.2 gives an overview of the relevant publications, describing how they relate to our work. Section 3.3 gives a description of the radio-coverage prediction problem, including the radio-propagation model. Section 3.4 concentrates on the design principles and implementation details of the radio-propagation tool, for the serial and parallel versions. Section 3.5 discusses the experimental results and their analysis.

There is a constant growing demand for hardware resources, longer-processing times and more memory to follow the evolution of 3G radio networks [88, 32, 125]. Fortunately, high-performance computer systems are increasingly accessible; something made possible because of the emergence of computer clusters and commodity hardware, capable of true parallel processing, e.g. multi-core CPUs [57] and GPUs [149]. Moreover, the highly parallel structure present on GPUs makes them more effective than CPUs for execution of algorithms

where large blocks of data need to be processed in parallel. Commodity GPUs have evolved from being a graphic accelerator into a general-purpose processor. They can achieve higher performance at lower power consumption and lower costs when compared to conventional CPUs. Additionally, the implementation of the framework will benefit from valuable advances in computer science and High Performance Computing (HPC), in order to perform faster and more reliable simulations [57, 149].

### 3.1 Motivation

Although Gordon Moore’s well-known and often cited prediction still holds [93], the fact is that for the past few years, CPU speeds have hardly been improving. Instead, the number of cores within a single CPU is increasing. This situation poses a challenge for software development in general and research in particular: a hardware upgrade will, most of the time, fail to double the serial execution speed of its predecessor. However, since this commodity hardware is present in practically all modern desktop computers, it creates an opportunity for the parallel exploitation of these computing resources to enhance the performance of complex algorithms over large data sets. The challenge is thus to deliver the computing power of multi-core systems in order to tackle a computationally time-consuming problem, the completion of which is unfeasible using traditional serial approaches. Indeed, improving the performance and data-set sizes a well-known approach may handle opens new possibilities for models and implementations of enhanced analytical methods.

A traditional approach when dealing with computationally expensive problem solving is to simplify the models in order to be able to execute their calculations within a feasible amount of time. Clearly, this method increases the introduced error level, which is not an option for a certain group of simulations, e.g., those dealing with disaster contingency planning and decision support [70, 154]. The conducted simulations during the planning phase of a radio network also belong to this group. Their results are the basis for the decision making prior to physically installing the base stations and antennas that will cover a certain geographical area. A larger deviation of these results increases the probability of making the wrong decisions at the time of the installation, which may considerably increase the costs or even cause mobile-network operators to incur losses.

Various groups have successfully deployed high-performance computing (HPC) systems and techniques to solve different problems dealing with spatial data [6, 10, 59, 70, 85, 154, 102, 131, 132, 133, 150]. This research has confirmed that a parallel paradigm such as master-worker, techniques like work pool (or task farming) and spatial-block partitioning are applicable when dealing with parallel implementations over large spatial data sets. However, it is well known that parallel programming and HPC often call for area experts in order to integrate these practices into a given environment [30]. Moreover, the wide range of options currently available creates even more barriers for general users wanting to benefit from HPC.

#### 3.1.1 GRASS

As the software environment for PRATO we have chosen GRASS [99], which is a free and open-source software project that implements a Geographical Information System (GIS). This GIS software was originally developed at the US Army Construction Engineering Research Laboratories and is a full-featured system with a wide range of analytical, data-management, and visualization capabilities. Currently, the development of GRASS GIS is supported by a growing community of volunteer developers.

The use of GRASS GIS as an environment for PRATO presents many advantages. First, the current development of GRASS is primarily Linux-based. Since the field of HPC is

dominated by Linux and UNIX systems, an environment with Linux support is critical for this work. Software licensing is another important consideration for choosing GRASS, since it is licensed under the GNU Public License [126] and imposes the availability of the source code. This allows us to make potential modifications to the system, thus adapting it for the parallel computation environment. Moreover, being an open system, GRASS provided us with a great deal of built-in functionality, capable of operating with raster and vector topological data that can be stored in an internal format or a DB.

## 3.2 Related work

The task-parallelization problem within the GRASS environment has been addressed by several authors in a variety of studies. For example, in [21], the authors present a collection of GRASS modules for a watershed analysis. Their work concentrates on different ways of slicing raster maps to take advantage of a Message Passing Interface (MPI) implementation.

In the field of HPC, the authors of [6] presented implementation examples of a GRASS raster module, used to process vegetation indexes for satellite images, for MPI and Ninf-G environments. The authors acknowledge a limitation in the performance of their MPI implementation for big processing jobs. The restriction appears due to the computing nodes being fixed to a specific spatial range, since the input data are equally distributed among worker processes, creating an obstacle for load balancing in heterogeneous environments.

Using a master-worker technique, the work [69] abstracts the GRASS data types into its own *struct* and MPI data types, thus not requiring the GRASS in the worker nodes. The data are evenly distributed by row among the workers, with each one receiving an exclusive column extent to work on. The test cluster contains heterogeneous hardware configurations. The authors note that data-set size is bounded by the amount of memory on each of the nodes, since they allocate the memory for the whole map as part of the set-up stage, before starting the calculation. Regarding the data sets during the simulations, the largest one contains 3,265,110 points. They conclude that the data-set size should be large enough for the communication overhead to be hidden by the calculation time, so that the parallelization pays off.

In [131], the authors employ a master-worker approach, using one worker process per worker node. The complete exploitation of the computing resources of a single computing node is achieved with OpenMP. The experimental environment features one host. The horizon-composition algorithm presents no calculation dependency among the spatial blocks. Consequently, the digital elevation model (DEM) may be divided into separate blocks to be independently calculated by each worker process. The authors present an improved algorithm that can also be used to accelerate other applications like visibility maps. The tasks are dynamically assigned to idle processes using a task-farming paradigm over the MPI.

Similarly, in [132] there is no calculation dependency among the spatial blocks. The experimental evaluation is made over multiple cores of one CPU and a GPU, and a master-worker setup is used for process communication.

In [154], the authors present a parallel framework for GIS integration. Based on the principle of spatial dependency, they lower the calculation-processing time using a knowledge database, delivering the heavy calculation load to the parallel back-end if a specific problem instance is not found in the database. There is an additional effort to achieve the presented goals, since the implementation of a fully functional GIS (or “thick GIS” as the authors call it) is required on both the desktop client and in the parallel environment.

An agent-based approach for simulating spatial interactions is presented in [53]. The presented technique decomposes the entire landscape into equally-sized regions, i.e., a spatial-

block division as in [131], which are in turn processed by a different core of a multi-core CPU. This work uses multi-core CPUs instead of a computing cluster.

Some years ago, grid computing received the attention of the research community as a way of accessing the extra computational power needed for the spatial analysis of large data sets [10, 147, 148]. However, several obstacles are still preventing this technology from being widely used. In particular, its adoption requires not only hardware and software compromises with respect to the involved parts, but also a behavioral change at the human level [10].

### 3.3 Radio-coverage prediction for mobile networks

#### 3.3.1 Background

The coverage planning of radio networks is a key problem that all mobile operators have to deal with. Moreover, it has proven to be a fundamental issue, not only in LTE networks, but also in other standards for mobile communications [109, 111, 124, 142]. One of the primary objectives of mobile-network planning is to efficiently use the allocated frequency band to ensure that some geographical area of interest can be satisfactorily reached with the base stations of the network. To this end, radio-coverage prediction tools are of great importance as they allow network engineers to test different network configurations before physically implementing the changes. Radio-coverage prediction is a complex task, mainly due to the several combinations of hardware and configuration parameters that have to be analyzed in the context of different environments. The complexity of the problem means that radio-coverage predictions are computationally-intensive and time-consuming, hence the importance of using fast and accurate tools (see Section 3.4.2 for a complexity analysis of the algorithm). Additionally, since the number of deployed transmitters keeps growing with the adoption of modern standards [109], there is a clear need for a radio-propagation tool that is able to cope with larger work loads in a feasible amount of time (see Section 3.4.2 for the running time of the serial version).

In the following, a high-performance radio-propagation prediction tool for GSM (2G), UMTS (3G) and LTE (4G) radio networks is presented. It can be used for planning the different phases of a new radio-network installation, as well as a support tool for maintenance activities related to network troubleshooting in general and optimization in particular. Specifically, automatic radio-coverage optimization requires the evaluation of millions of radio-propagation predictions in order to find a good solution set, which is unfeasible using other serial implementations of academic or commercial tools [92, 104, 68].

As a reference implementation, the publicly available radio-coverage prediction tool, presented in [68], was used. The authors developed a modular radio-coverage tool that performs separate calculations for radio-signal path loss and antenna-radiation patterns, also taking into account different configuration parameters, such as antenna tilting, azimuth and height. The output result, saved as a raster map, is the maximum signal level over the target area, in which each point represents the received signal from the best-serving transmitter (or cell). This work implements some well-known radio-propagation models, e.g., Okumura-Hata [63] and COST 231 [29]. The latter is explained in more detail in Section 3.3.2. Regarding the accuracy of the predicted values, the authors [68] report comparable results to those of an industrial tool. To ensure that the presented implementation is completely compliant with the previously mentioned reference, a comparison test that consists of running both tools with the same set of input parameters was designed. The test results from PRATO and the reference implementation were identical in all the tested cases.

### 3.3.2 Radio-propagation modeling

PRATO uses a modified version of the well-known Okumura-Hata empirical model for radio-propagation predictions [63]. This model has been largely studied and shown to be suitable for predicting the radio propagation in LTE networks [? 111]. In its primary form, the model distinguishes the distance from the receiver to the transmitter, the frequency used and the effective antenna height, i.e., the antenna height above the receiver's level. These variables are taken into account in order to calculate the path loss in line-of-sight (LOS) conditions. Additionally, for distinguishing non-line-of-sight (NLOS) conditions, the terrain profile and Earth shape are added to the original formula. In this context, a NLOS situation appears when the first Fresnel zone is obscured by at least one obstacle [153]. Equation (3.1) describes the path loss when there is LOS between the transmitter and the receiver.

$$\begin{aligned} pl_{\text{LOS}}(d_{(x,y)}, \beta) = & a_0 + a_1 \log(d_{(x,y)}) + a_2 \log(H_A) + \\ & a_3 \log(d_{(x,y)}) \log(H_A) - 3.2 [\log(11.75 \cdot H_R)]^2 + \\ & 44.49 \log(F) - 4.78 [\log(F)]^2, \quad (3.1) \end{aligned}$$

where  $\beta = (a_0, a_1, a_2, a_3)$  is the vector containing the tuning parameters of the model,  $d_{(x,y)}$  is the distance (in kilometers) from the transmitter to the topography point with coordinates  $(x, y)$ ,  $H_A$  is the effective antenna height (in meters) of the transmitter,  $H_R$  is the antenna height (in meters) of the receiver, and  $F$  is the frequency, expressed in MHz. On the other hand, in NLOS conditions, the path loss is calculated as in Equation (??).

$$pl_{\text{NLOS}}(d_{(x,y)}) = \sqrt{[\alpha K(d_{(x,y)})]^2 + E(d_{(x,y)})^2}, \quad (3.2)$$

where  $\alpha$  is the knife-edge diffraction control parameter, which value is calculated based on the level of obstruction of a Fresnel zone,  $K(d_{(x,y)})$  is the knife-edge diffraction loss (in dB), and  $E(d_{(x,y)})$  is the correction due to the Earth sphere (in dB). All three values depend on the characteristics of the topography point with coordinates  $(x, y)$ .

In this work, as well as in [? ], the terrain profile is used for LOS determination, i.e., obstacle obstruction in the first Fresnel zone of the transmitter. In order to adequately predict signal-loss effects due to foliage, buildings and other fabricated structures, additional loss factors based on the land usage (i.e., clutter data), are included. This technique is adopted by several propagation models for radio networks [? ? 98]. Consequently, an extra term is introduced for signal loss due to clutter, thus defining the model-predicted path loss as

$$pl(d_{(x,y)}, \beta) = pl_{\text{LOS}}(d_{(x,y)}, \beta) + pl_{\text{NLOS}}(d_{(x,y)}) + pl_{\text{CLUT}}(d_{(x,y)}), \quad (3.3)$$

where  $pl_{\text{CLUT}}(d_{(x,y)})$  represents the clutter loss at the topography point with coordinates  $(x, y)$ , expressed in dB.

## 3.4 Design and implementation

### 3.4.1 Design of the serial version

This section describes the different functions contained in the serial version of PRATO, which is implemented as a GRASS module. Their connections and data flow are depicted in Figure 3.1, where the parallelograms of the flow diagram represent the input/output (I/O) operations.

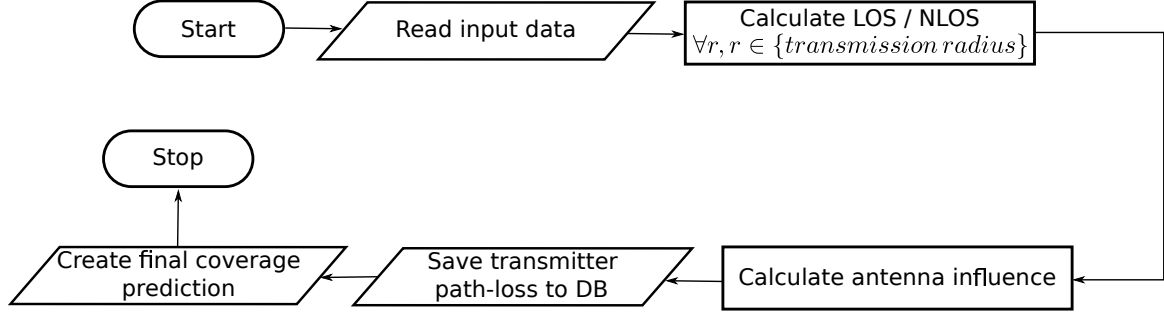


Figure 3.1: Flow diagram of the serial version.

The design follows a similar internal organization as the radio-planning tool presented in [68], but with some important differences. Specifically, the design presented here employs a direct connection to an external database server for intermediate result saving, instead of the slow, built-in GRASS database drivers. To explicitly avoid tight coupling with a specific database vendor, the generated output is formatted in plain text, which is then forwarded to the DB. Any further processing is achieved by issuing a query over the database tables that contain the path-loss results for each of the processed transmitters.

### 3.4.1.1 Input parameters

All input data are read in the first step (see “Read input data” in Figure 3.1). Their formats differ based on the data they contain, i.e.,

- GRASS raster files are used for the DEM and clutter data, whereas
- a text file is used for the transmitter configurations and other service-dependent options.

Since the module accepts a considerable amount of input parameters, they are read from a text-based initialization (INI) file. This is far more practical than passing them as command-line parameters, which would make them error-prone and difficult to read. Besides, the INI file may contain configuration parameters for many transmitters. The user selects which one(s) to use at run-time by passing a command-line option.

### 3.4.1.2 Isotropic path-loss calculation

This step starts by calculating which receiver points,  $r$ , are within the specified transmission radius (see “*transmission radius*” in Figure 3.1). The transmission radius is defined around each transmitter in order to limit the radio-propagation calculation to a reasonable distance. For these points, the LOS and NLOS conditions are calculated with respect to the transmitter (see “Calculate LOS/NLOS” in Figure 3.1). The following step consists of calculating the path loss for an isotropic source (or omni antenna). This calculation is performed by applying the radio-propagation model, which was previously defined in Equation (3.3), to each of the points within the transmission radius around the transmitter.

Figure 3.2 shows an example result of the isotropic path-loss calculation, only including the map area within the transmission radius. The color scale is given in dB, indicating the signal loss from the isotropic source of the transmitter, located at the center. Notice the hilly terrain is clearly distinguished due to LOS and NLOS conditions from the signal source.

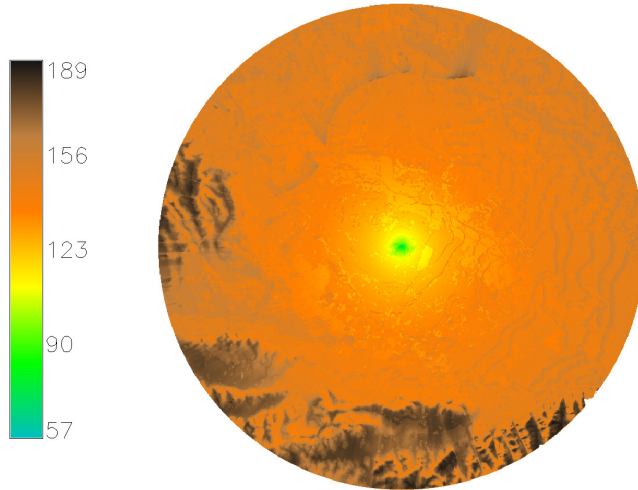


Figure 3.2: Example of raster map, showing the result of a path-loss calculation from an isotropic source.

#### 3.4.1.3 Antenna diagram influence

This step considers the antenna radiation diagram of the current transmitter and its influence over the isotropic path-loss calculation (see “Calculate antenna influence” in Figure 3.1). Working on the in-memory results generated by the previous step, the radiation diagram of the antenna is taken into account, including the beam direction, the electrical and the mechanical tilt. Figure 3.3 shows the map area within the transmission radius, where this calculation step was applied to the results from Figure 3.2. Notice the distortion of the signal propagation that the antenna has introduced.

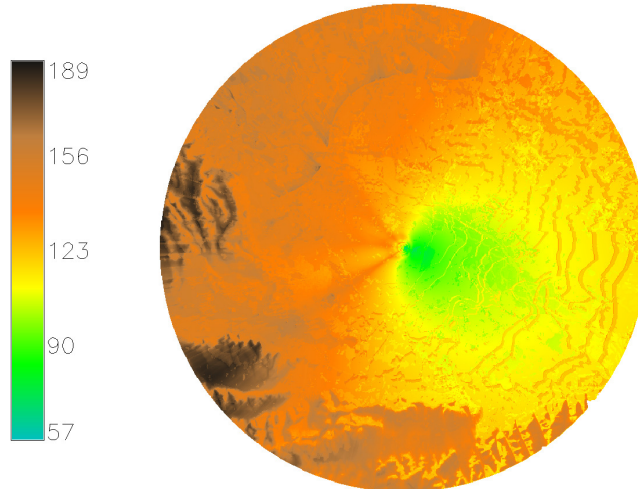


Figure 3.3: Example of raster map, showing the antenna influence over the isotropic path-loss result, as depicted in Figure 3.2.

#### 3.4.1.4 Transmitter path-loss prediction

In this step, the path-loss prediction of the transmitter is saved in its own database table (see “Save transmitter path-loss to DB” in Figure 3.1). This is accomplished by connecting

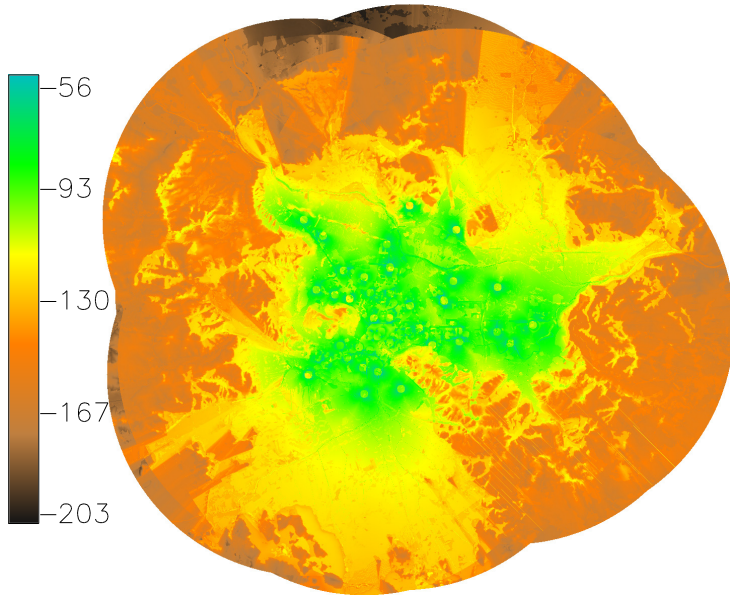


Figure 3.4: Example of a raster map, displaying the final coverage prediction of 136 transmitters over a geographical area. The color scale is given in dBm, indicating the received signal strength. Darker colors denote areas with a reduced signal due to the fading effect of the hilly terrain and clutter.

the standard output of the GRASS module with the standard input of a database client. Naturally, the generated plain text should be understood by the DB itself.

#### 3.4.1.5 Coverage prediction

The final radio-coverage prediction, containing the aggregation of the partial path-loss results of the involved transmitters, is created in this step (see “Create final coverage prediction” in Figure 3.1). The received signal strength from each of the transmitters is calculated as the difference between its transmit power and the path loss for the receiver’s corresponding position. This is done by executing an SQL query over the tables containing the path-loss predictions of each of the processed transmitters. Finally, the output is generated, using the GRASS built-in modules *v.in.ascii* and *v.to.rast*, which create a raster map using the query results as the input. The resulting raster map contains the maximum received signal strength for each individual point, as shown in Figure 3.4. In this case, the color scale is given in dBm, indicating the strongest received signal strength from the transmitters.

#### 3.4.2 Computational complexity of the radio-coverage algorithm

In this section, the time complexity of the radio-coverage prediction algorithm is presented, for which the pseudo code is listed in Algorithm 3.1.

The algorithm starts by loading the input, i.e., the DEM and the clutter data. Both regular square grids (RSGs) should account for the same area and resolution, consequently containing the same number of pixels,  $M$ . The transmitter data is then loaded into set  $T$ , the cardinality of which is denoted as  $n = |T|$ . For each transmitter  $t \in T$ , a smaller subarea of the DEM and clutter data (denoted  $DEM_t$  and  $Clut_t$ , respectively) is delimited around  $t$ , based on a given transmission radius. The number of pixels within this sub-area is denoted as  $m$ , and its value is the same for all  $t \in T$ . The visibility for an RSG pixel is computed using the *LineOfSight* function, by walking from the antenna of the transmitter to

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**Algorithm 3.1** Pseudo code of the radio-coverage prediction algorithm. The time complexity is given per line.

---

```

 $DEM \leftarrow$  Digital Elevation Model (DEM) of the whole area.  $\triangleright O(M)$ 
 $Clutter \leftarrow$  signal Losses due to land usage of the whole area.  $\triangleright O(M)$ 
 $T \leftarrow$  transmitter configuration data.  $\triangleright O(n)$ 
for all  $t \in T$  do  $\triangleright O(n \cdot m^2)$ 
     $DEM_t \leftarrow$  DEM area within transmission radius of  $t$   $\triangleright O(m)$ 
     $Clutt_t \leftarrow$  Clutter area within transmission radius  $t$   $\triangleright O(m)$ 
     $LoS_t \leftarrow$  LineOfSight ( $DEM_t$ )  $\triangleright O(m^2)$ 
     $PL_t \leftarrow$  PathLoss ( $DEM_t, Clutt_t, LoS_t$ )  $\triangleright O(m^2)$ 
     $Diag_t \leftarrow$  Antenna diagram of  $t$   $\triangleright O(1)$ 
     $PL_t \leftarrow$  AntennaInfluence ( $Diag_t, PL_t$ )  $\triangleright O(m)$ 
end for
for all  $t \in T$  do  $\triangleright O(n \cdot m)$ 
     $CoveragePrediction \leftarrow$  PathLossAggregation ( $t, PL_t$ )  $\triangleright O(m)$ 
end for
return  $CoveragePrediction$ 

```

---

the given pixel element, along the elements intersected by a LOS, until either the visibility is blocked, or the target is reached [38]. Regarding the *PathLoss* function, whenever a receiver point is in NLOS, the walking path from the transmitter has to be inspected for obstacles, calculating the diffraction losses for each of them, i.e.,  $\alpha$  and  $K(d_{(x,y)})$  from Equation (3.2). Hence, its quadratic complexity, which dominates the complexity of the algorithm, together with *LineOfSight*, resulting in an algorithmic complexity denoted by

$$O(M + n \cdot m^2). \quad (3.4)$$

Although  $n$  will generally be many orders of magnitude smaller than  $m^2$ , its computational-time complexity is relevant for practical use. For example, assuming the radio-coverage prediction for one transmitter completes in around 15 seconds using a serial implementation, the prediction for a mobile network comprising 10,240 transmitters would have an execution time of almost two days.

### 3.4.3 Multi-paradigm parallel programming

The implementation methodology adopted for PRATO follows a multi-paradigm, parallel programming approach in order to fully exploit the resources of each of the nodes in a computing cluster. This approach combines a master-worker paradigm with an external DB. To efficiently use a shared memory multi-processor on the worker side, and to effectively overlap the calculation and communication, PRATO uses POSIX threads [20].

To use the computing resources of a distributed memory system, such as a cluster of processors, PRATO uses the MPI [58]. The MPI is a message-passing standard that defines the syntax and semantics designed to function on a wide variety of parallel computers. The MPI enables multiple processes, running on different processors of a computer cluster, to communicate with each other. It was designed for high performance on both massively parallel machines and on workstation clusters.

PRATO also supports the execution of the most computationally-intensive parts of the radio-propagation algorithm on a GPU. Moreover, the GPU hardware is used if available on the computing nodes that host the worker processes (see Section 3.4.4.4 later in this chapter).

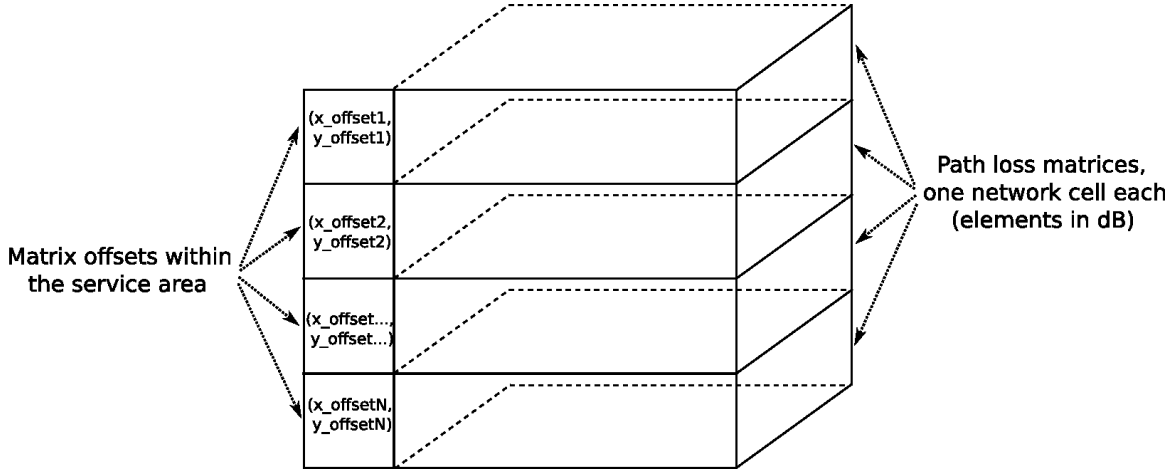


Figure 3.5: Memory organization of the input-spatial data.

In order to make the text clearer and to differentiate between the programming paradigms used from here on, a POSIX thread will be referred to simply as a ‘thread’ and a MPI process as a ‘process’.

### 3.4.4 Design of the parallel version

The focus here is on the practical usability and performance of PRATO. Therefore, to overcome the computational-time constraints that prevent a serial implementation of the radio-coverage prediction algorithm from tackling bigger problem instances in a feasible amount of time, a parallel implementation is presented.

A major drawback of the GRASS as a parallelization environment is that it is not thread-safe, meaning that concurrent changes to the same data set have an undefined behavior [18]. One technique to overcome this problem is to abstract the spatial data from the GRASS. For example, in [69], the authors achieve the GRASS abstraction by introducing a *Point* structure with four *double* attributes, where each pixel of the RSG is mapped to an instance of this structure. Another possibility is for one of the processes, e.g., the master, to read entire rows or columns of data before dispatching them for processing to the workers [6, 69]. In this case, an independence between row/column calculations is required, which is a problem-specific property. Here, abstraction from the GRASS is achieved by loading the spatial data into a 2D matrix (or matrices) of basic data-type elements, e.g., *float* or *double* depending on the desired accuracy. The geographical location of each element is calculated as the geographical location of the matrix plus the element offset within it (see Figure 3.5). To correctly locate the path-loss matrices within the service area, we supply the offset of the upper-left corner of each of them???. The advantage of this technique is having the geographical location of each pixel readily available with a minimum memory footprint. Moreover, a convenient consequence of this abstraction schema is that worker processes are completely independent of the GRASS, thus significantly simplifying the deployment of the parallel implementation over multiple computing hosts.

In the area of geographical information science, the master-worker paradigm has been successfully applied by several authors [5, 6, 21, 59, 69, 131, 132]. However, sometimes this technique presents certain issues that prevent the full exploitation of the available computing resources when deployed over several networked computers. Additionally, such issues are difficult to measure when the parallelization involves only one computing node [131, 132] (i.e., no network communication is required), or only a few processes deployed over a handful of

nodes [69]. Specifically, we are referring to network saturation and idle processes within the master-worker model. Generally speaking, a single communicating process, e.g., the master, is usually not able to saturate the network connection of a node. Using more than one MPI process per node might solve this problem, but possible rank-ordering problems may appear, thus restricting the full utilization of the network [106]. Another issue appears when the master process executes the MPI code, in which case other processes sleep, making a serial use of the communication component of the system. Consequently, the master process becomes the bottleneck of the parallel implementation as the number of worker processes it has to serve grows. This situation is also common when dealing with the metadata of a spatial region, which may relate to several elements of a RSG, making it a frequent cause of load imbalance [53, 64, 150]. In PRATO, the transmitter configuration and its antenna diagram represent metadata that are complementary to the sub-region that a transmitter covers.

Hybrid MPI-OpenMP implementations [131, 132], in which no MPI calls are issued inside the OpenMP-parallel regions, also fail to saturate the network [106]. A possible solution to this problem is to improve the communication overlap among the processes. To this end, PRATO features non-blocking point-to-point MPI operations, and an independent thread in the worker process to save the intermediate results to a DB. One such database system per computer cluster is used, which also serves the input data to the GRASS, in order to aggregate the partial results of the path-loss predictions or to visualize them. It is important to note that any kind of DB may be used. By this we mean relational, distributed [103] or even those of the NoSQL type [128]. Nevertheless, a central relational database system is used here, since they are the most popular and widely available ones. Additionally, the non-blocking message-passing technique used to distribute the work-load among the nodes provides support for heterogeneous environments. As a result, computing nodes featuring more capable hardware receive more work than those with weaker configurations, thus ensuring a better utilization of the available computing resources despite hardware diversity and improved load balancing.

#### 3.4.4.1 Master process

The master process, for which the flow diagram is given in Figure 3.6, is the only component that runs within the GRASS environment. As soon as the master process starts, the input parameters are read. This step corresponds to “Read input data” in Figure 3.6, and it is carried out in a similar way as in the serial version. The next step delivers the metadata that is common to all the transmitters and the whole region to all the processes (see “Metadata broadcasting” in Figure 3.6). Before distributing the work among the worker processes, the master process proceeds to decompose the loaded raster data into 2D matrices of basic-data-type elements, e.g., *float* or *double*, before dispatching them to the multiple worker processes. In this case, the decomposition applies to the DEM and the clutter data only, but it could be applied to any point-based data set. In the next step, the master process starts an asynchronous message-driven processing loop (see “Processing loop” in Figure 3.6), the main task of which is to assign and distribute the sub-region and configuration data of different transmitters among the idle worker processes.

The flow diagram shown in Figure 3.7 illustrates the “Processing loop” step of the master process. In the processing loop, the master process starts by checking the available worker processes, which will calculate the radio-coverage prediction for the next transmitter. It is worth pointing out that this step also serves as a stopping condition for the processing loop itself (see “Any worker still on?” in Figure 3.7). The active worker processes inform the master process that they are ready to compute by sending an idle message (see “Wait for

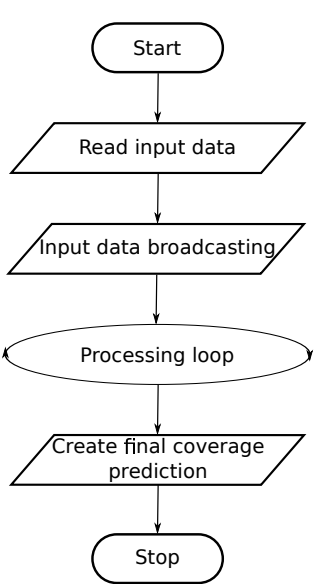


Figure 3.6: Flow diagram of the master process.

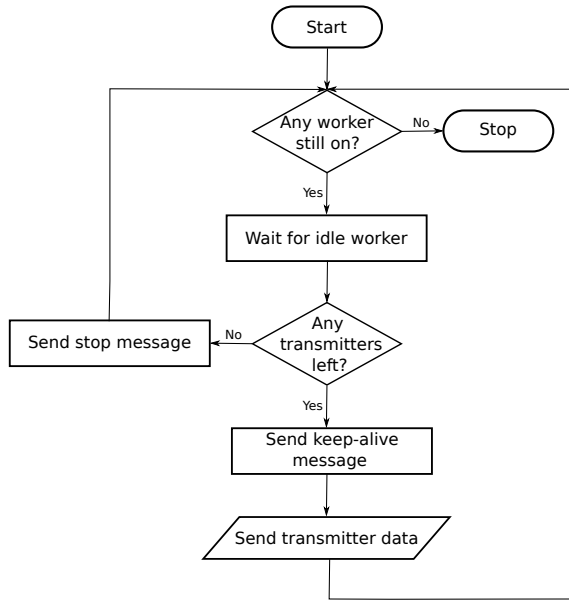


Figure 3.7: Flow diagram of the “Processing loop” step of the master process.

idle worker” in Figure 3.7). The master process then announces to the idle worker process that it is about to receive new data for the next calculation, and it dispatches the complete configuration of the transmitter to be processed (see “Send keep-alive message” and “Send transmitter data” steps, respectively, in Figure 3.7). This is only done in the case that there are transmitters for which the coverage prediction has yet to be calculated (see “Any transmitters left?” in Figure 3.7). The processing loop of the master process continues to distribute the transmitter data among the worker processes, which asynchronously become idle as they finish the radio-prediction calculations they have been assigned by the master process. When there are no more transmitters left, all the worker processes announcing they are idle will receive a shutdown message from the master process, indicating to them that they should stop running (see “Send stop message” in Figure 3.7). The master process will keep doing this until all the worker processes have finished (see “Any worker still on?” in Figure 3.7), thus fulfilling the stopping condition of the processing loop.

Finally, the last step of the master process is devoted to creating the final output of the calculation, e.g., a raster map (see “Create final coverage prediction” in Figure 3.6). The final coverage prediction of all the transmitters is an aggregation from the individual path-loss results created by each of the worker processes during the “Processing loop” phase in Figure 3.6, which provides the source data for the final raster map. The aggregation of the individual transmitter path-loss results is accomplished by issuing an SQL query over the database tables containing the partial results, in a similar way as in the serial version.

#### 3.4.4.2 Worker processes

An essential characteristic of the worker processes is that they are completely independent of the GRASS, i.e., they do not have to run within the GRASS environment nor use any of the GRASS libraries to work. This aspect significantly simplifies the deployment phase to run PRATO on a computer cluster, since no GRASS installation is needed on the computing nodes hosting the worker processes.

One possibility to overcome the thread-safety limitation of the GRASS is to save the transmitter path-loss predictions through the master process, thus avoiding concurrent ac-

cess. However, for the workers to send intermediate results back to the master process, e.g., as in [5, 69], is a major bottleneck for the scalability of a parallel implementation. The scalability is limited by the master process, because it must serially process the received results in order to avoid inconsistencies due to concurrent access. Instead, PRATO allows each of the worker processes to output its intermediate results into a DB, i.e., each path-loss prediction in its own table. Additionally, worker processes do this from an independent thread, which runs concurrently with the calculation of the next transmitter received from the master process. In this way, the overlap between the calculation and communication significantly hides the latency created by the result-dumping task, thus making better use of the available system resources.

The computations of the worker processes, for which the flow diagram is given in Figure 3.8, begin by receiving metadata about the transmitters and the geographical area from the master process during the initialization time (see “Receive broadcasted metadata” in Figure 3.8).

After the broadcasted metadata are received by all the worker processes, each one proceeds to inform the master process that it is ready (i.e., in an idle state) to receive the transmitter-configuration data that defines which transmitter path-loss prediction to perform (see “Send idle message” in Figure 3.8). If the master process does not give the instruction to stop processing (see “Has stop message arrived?” in Figure 3.8), the worker process collects the sub-region spatial data and the transmitter configuration (see “Receive transmitter data” in Figure 3.8). In the event that a stop message is received, the worker process will wait for any result-dumping thread to finish (see “Wait for result-dump thread” in Figure 3.8) before shutting down. The coverage calculation itself follows a similar design as the serial version (see “Coverage calculation” in Figure 3.8).

As mentioned before, the worker process launches an independent thread to save the path-loss prediction of the target transmitter to a database table (see “Threaded save path-loss to DB” in Figure 3.8). It is important to note that there is no possibility of data inconsistency due to the saving task being executed inside a thread, since path-loss data from different workers belong to different transmitters and are, at this point of the process, mutually exclusive.

#### 3.4.4.3 Master-worker communication

Similar to [131, 132], the message-passing technique used in this work enables a better use of the available computing resources, both in terms of scalability and load balancing, while introducing a negligible overhead. This last point is supported by the experimental results, introduced in Section 3.5.3.

The first reason to implement the message-passing technique is to support heterogeneous computing environments. In particular, our approach focuses on taking full advantage of the hardware of each computing node, thus explicitly avoiding the bottlenecks introduced by the slowest computing node in the cluster. This problem appears when evenly distributing the data among the worker processes on disparate hardware, e.g., as in [6, 69], being more noticeable with a larger number of computing nodes and processes. In other words, computing nodes that deliver better performance have more calculations assigned to them. Moreover, in real-world scenarios, it is often the case that a large number of dedicated computing nodes featuring exactly the same configuration is difficult to find, i.e., not every organization owns a computer cluster.

A second reason for selecting a message-passing technique is related to the flexibility it provides for load balancing, which is of greater importance when dealing with extra data or information besides spatial data [64]. This can be seen in Figure 3.7, where the master

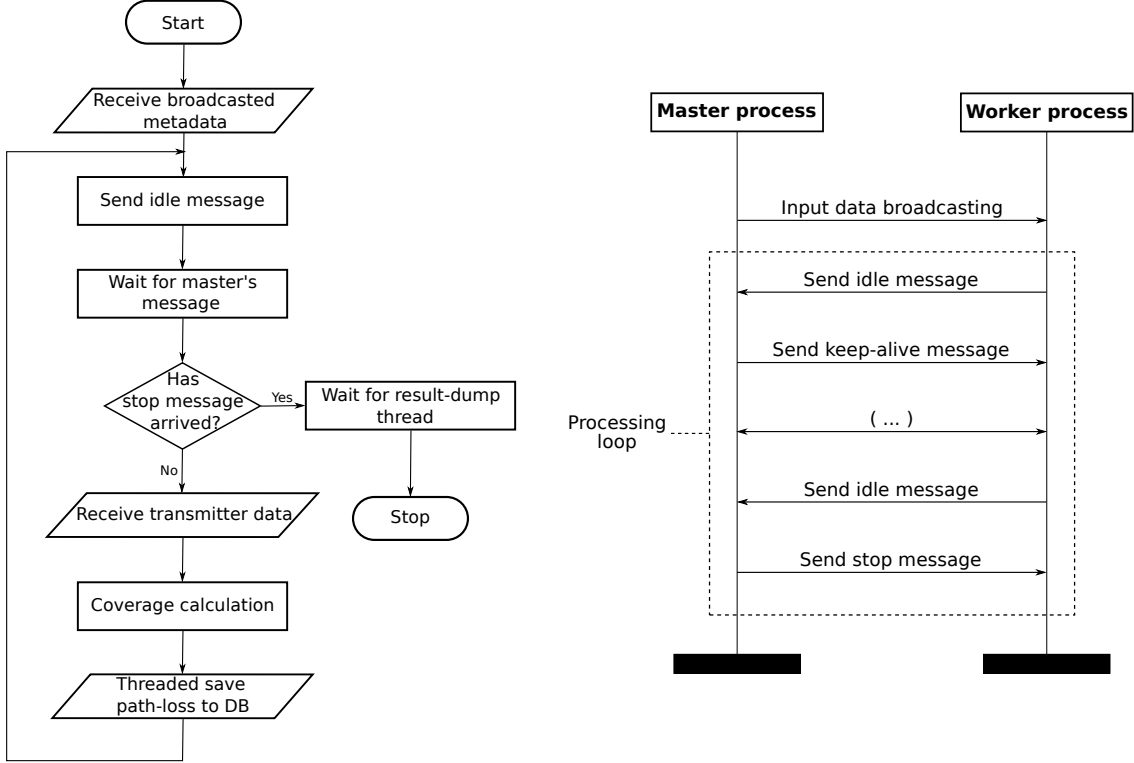


Figure 3.8: Flow diagram of a worker process.

Figure 3.9: Communication diagram, showing the message passing between the master and a worker process.

process, before delivering the spatial subset and transmitter-configuration data, sends a message to the worker process, indicating that it is about to receive more work. This a priori meaningless message plays a key role in correctly supporting the asynchronous process communication. Notice that the subset of spatial data that a worker process receives is directly related to the transmitter for which the prediction will be calculated. Similar to [131, 132], this problem-specific property enables the use of a data-decomposition technique based on a block partition of spatial data, e.g., the DEM and clutter data.

In general, there are many different ways a parallel program can be executed, because the steps from the different processes can be interleaved in various ways and a process can make non-deterministic choices [113], which may lead to situations such as race conditions [31] and deadlocks. A deadlock occurs whenever two or more running processes are waiting for each other to finish, and thus neither ever does. To prevent PRATO from deadlocking, message sending and receiving should be paired, i.e., an equal number of send and receive messages on the master and worker sides [113]. Figure 3.9 depicts the master-worker message passing, from which the transmitter-data transmission has been excluded for clarity. Notice how each idle message sent from the worker process is paired with an answer from the master process, whether it is a keep-alive or a stop message.

#### 3.4.4.4 GPU-accelerated worker processes

PRATO provides multi-GPU support for performance improvement on the computing nodes hosting the worker processes. The algorithmic adaptation from a CPU to a GPU is not a trivial task. This section focuses on the main modifications of the radio-propagation algorithm to work on GPU hardware.

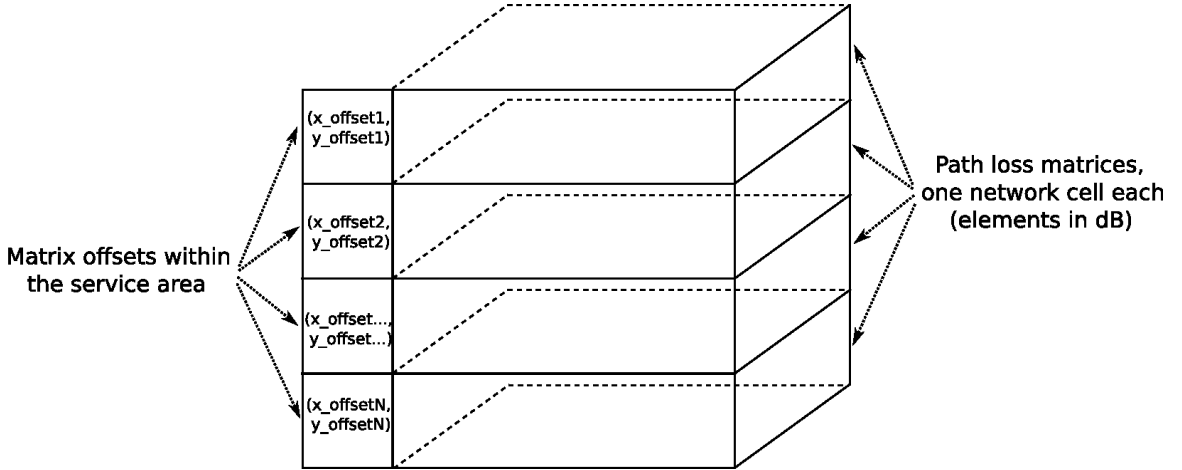


Figure 3.10: Memory system of a modern GPU, including random-access times in milliseconds.

GPUs have a totally different memory system (see Figure 3.10) and many times more processor cores than CPUs. Specifically, the actual throughput an application can achieve depends on issues related to the access to memory, in particular the high-latency accesses to global memory from the GPU chip, and the use of shared memory in the streaming multiprocessors to mitigate the high-latency effects [34]. For this reason, it is imperative to have as much data as allocated on the GPU itself.

In order to minimize the CPU-to-GPU memory transfers, the spatial data used by the radio-propagation algorithm was organized as explained before in Section 3.4.4, i.e., using geographically-located, offset-based 2D matrices. However, the internal representation of the matrix elements was changed to use less memory. To this end, the clutter-matrix elements were represented as *unsigned char*, since they express signal loss (in dB) due to land usage. It follows that for the radio-propagation prediction of one transmitter, the following matrices should be allocated on the GPU:

- one 2D matrix containing DEM data for the target subregion, the elements of which are *float* or *double*;
- one 2D matrix containing clutter data for the target subregion, the elements of which are *unsigned char*; and
- one 2D matrix containing the resulting path-loss prediction.

The dimension of all matrices is based on the transmission radius, within which the radio-propagation prediction should be calculated. The contents of the DEM and clutter matrices is constant throughout the calculation process. For this reason they were saved into read-only texture memory to take advantage of the faster access time. Regarding the resulting path-loss matrix, each step of the radio-prediction algorithm is applied over the results of the previous step (see Figure 3.1 for a flow diagram of the steps involved), thus avoiding the allocation of extra memory or a data-transfer from/to the CPU.

### 3.5 Simulations

Considering the large computational power needed for predicting the radio-coverage of a real mobile network, the use of a computer cluster is recommended. A computer cluster

is a group of interconnected computers that work together as a single system. Computer clusters typically consist of several commodity PCs connected through a high-speed local-area network (LAN) with a distributed file system, like NFS [112]. One such system is the DEGIMA cluster [60] at the Nagasaki Advanced Computing Center (NACC) of the Nagasaki University in Japan. This system ranked in the TOP 500 list of supercomputers until June 2012<sup>1</sup>, and in June 2011 it held third place in the Green 500 list<sup>2</sup> as one of the most energy-efficient supercomputers in the world.

This section presents the simulations, and an exhaustive analysis of the performance and scalability of the parallel implementation of PRATO. The most common usage case for PRATO is to perform a radio-coverage prediction for multiple transmitters. Therefore, a straight-forward parallel decomposition is to divide a given problem instance by transmitter, for which each coverage prediction is calculated by a separate worker process.

The following simulations were carried out on 34 computing nodes of the DEGIMA cluster. The computing nodes are connected by a LAN, over a Gigabit Ethernet interconnect. As mentioned before, the reason for using a high-end computer cluster such as DEGIMA is to explore by experimentation the advantages and drawbacks of the considered methods. However, this does not imply any loss of generality if applying these principles over a different group of networked computers that do not operate as a computer cluster.

Each computing node of DEGIMA features one of two possible configurations, namely:

- Intel Core i5-2500T quad-core processor CPU, clocked at 2.30 GHz, with 16 GB of RAM; and
- Intel Core i7-2600K quad-core processor CPU, clocked at 3.40 GHz, also with 16 GB of RAM.

During the simulation runs, the nodes equipped with the Intel i5 CPU host the worker processes, whereas the master process and the PostgreSQL database server (version 9.1.4) each run on a different computing node, featuring an Intel i7 CPU. The database server performed all its I/O operations on the local file system, which was mounted on an 8 GB RAM disk. During the simulations, the path-loss predictions of 5,120 transmitters occupied less than 4 GB of this partition. No GPU hardware was used for the following simulation sets.

A 64-bit Linux operating system (Fedora distribution) was the operating system used. The message-passing implementation OpenMPI, version 1.6.1, was manually compiled with the distribution-supplied gcc compiler, version 4.4.4.

### 3.5.1 Test networks

The parallel performance of PRATO was tested with real radio networks of different sizes. In order to create the synthetic test data sets with an arbitrary number of transmitters, a group of 2,000 transmitters of a real network was used. The configuration parameters of these transmitters resembled those of the LTE network deployed in Slovenia by Telekom Slovenije, d.d., which were, in turn, randomly replicated and distributed over the whole target area. The path-loss predictions were calculated using the radio-propagation model introduced in Section 3.3.2. The DEM area, as well as the clutter data, covered 20,270 km<sup>2</sup> with a pixel resolution of 25 m<sup>2</sup>. The clutter data contained different levels of signal loss due to land usage. For all the points within a radius of 20 km around each transmitter, we assume that the receiver is positioned 1.5 m above the ground, and the frequency is set to 1,843 MHz.

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<sup>1</sup><http://www.top500.org>

<sup>2</sup><http://www.green500.org>

Table 3.1: Running-time gain (in percent) of the simulations for the weak-scalability of the MWD setup relative to the classic MW approach.

TX/core	Number of cores						
	1	2	4	8	16	32	64
5	-11.39	-10.42	-11.14	-0.95	11.75	26.15	32.53
10	-5.84	-7.78	-7.67	0.91	12.81	33.28	33.55
20	-8.59	-10.88	-1.04	1.95	14.29	35.23	35.27
40	-5.26	-6.90	-3.68	-0.67	17.27	36.23	36.65
80	-5.29	-7.11	-3.20	-0.31	17.94	36.32	36.57

### 3.5.2 Weak scalability

The weak-scalability experiments are meant to analyze the scalability of the parallel implementation in cases where the workload assigned to each process (one MPI process per processor core) remains constant as the number of processor cores is increased. It follows that the number of transmitters deployed over the target area is directly proportional to the number of processor cores and worker processes. This is accomplished by assigning a constant number of transmitters per core, while increasing the number of cores hosting the worker processes. Here, the following numbers of transmitters per worker/core were tested  $\{5, 10, 20, 40, 80\}$ , by progressively doubling the number of worker processes from 1 to 64.

Problems that are particularly well-suited for parallel computing exhibit computational costs that are linearly dependent on the size of the problem. This property, also referred to as algorithmic scalability, means that proportionally increasing both the problem size and the number of cores results in a roughly constant time to solution.

The master-worker (MW) configuration performs result aggregation continuously, i.e., while receiving the intermediate results from the worker processes. In contrast, the master-worker-DB (MWD) setup performs the result aggregation as the final step. This set of experiments is meant to investigate how the proposed MWD technique compares with the classic MW approach in terms of scalability when dealing with a constant computational load per core.

An important fact about the presented simulations when using multi-threaded implementations is to avoid oversubscribing a computing node. For example, if deploying four worker processes over a quad-core CPU, the extra threads will have a counter effect on the parallel efficiency, since the CPU resources would be exhausted, which slows the whole process down. For this reason, we have deployed three worker processes per computing node, leaving one core free for executing the extra threads.

#### 3.5.2.1 Results

The results represent the best running time out of a set of 20 independent simulation runs, in which we randomly selected both the transmitters and the rank ordering of the worker processes. The collected running times for the weak-scalability experiments are shown in Figure 3.11. All the measurements express wall-clock times in seconds for each setup and problem instance, defined as the number of transmitters per process (Tx/process). The wall-clock time represents the real time that elapses from the start of the master process to its end, including the time that passes while waiting for the resources to become available. The running-time improvements of the MWD versus the MW setup are shown in Table 3.1.

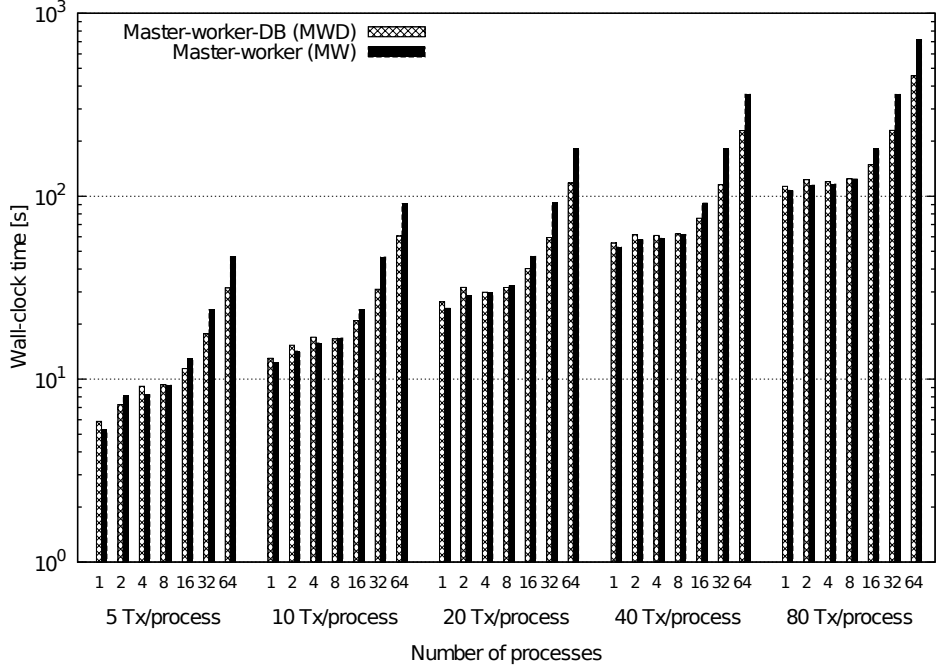


Figure 3.11: Measured wall-clock time for weak-scalability experiments, featuring MW and MWD setups. Experiments allocate one MPI worker process per core. The wall-clock time axis is expressed in a base-10 logarithmic scale, whereas the axis representing the number of cores is expressed in a base-2 logarithmic scale.

The time measurements observed from the weak-scalability results show that the classic MW approach performs well for up to four worker processes. When using eight worker processes, the MW setup is practically equivalent to the MWD approach, indicating that the master process is being fully exploited. When increasing the problem size and the number of worker processes to 16, the running-time gain is already clear, favoring the MWD configuration. This gain keeps growing, although slower, as we increase the number of worker processes to 32 and 64, confirming the hypothesis that in a classic MW approach, the parallel efficiency is bounded by the capacity of the master process to serve an increasing number of worker processes. Interestingly, the gain when using 32 and 64 worker processes is almost the same. After further investigation, the reason for this behavior was found: the new bottleneck was the LAN being completely saturated by the worker processes. Consequently, they have to wait for the network resources to become available before sending or receiving data, which is not the case when running the MW setup. Therefore, using the MWD approach a hardware constraint is hit, meaning that the bottleneck is no longer at the implementation level. Moreover, since the master process is far from overloaded when serving 64 worker processes, it can be expected that the MWD approach will keep scaling if a faster network infrastructure is used, e.g., 10-gigabit Ethernet or InfiniBand.

Certainly, the parallel version of PRATO, when using the MWD approach, scales better when challenged with a large number of transmitters (5,120 for the biggest instance) over 64 cores. This fact shows PRATO would be able to calculate the radio-coverage prediction for real networks in a feasible amount of time, since many operational radio networks have already deployed a comparable number of transmitters, e.g., the 3G network within the Greater London Authority area, in the UK [101]. For a more in-depth discussion and experimentation about real-world planning scenarios, see Chapter 7.

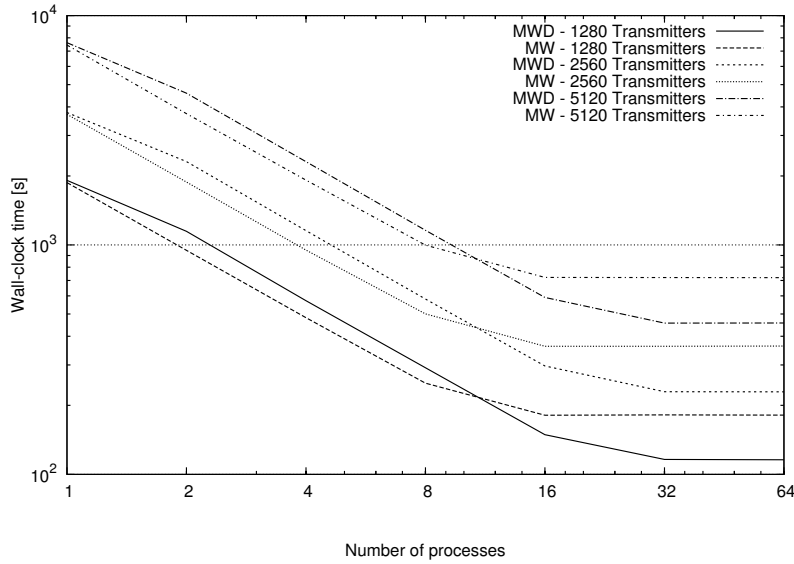


Figure 3.12: Measured wall-clock time for strong-scalability experiments, featuring MW and MWD setups. Experiments assigned one MPI worker process per core. The wall-clock time axis is expressed in a base-10 logarithmic scale, whereas the axis representing the number of cores is expressed in a base-2 logarithmic scale.

Not being able to achieve perfect weak scalability using the MWD setup is due to a number of factors. Specifically, the overhead time of the serial sections of the parallel process grows proportionally with the number of cores, e.g., aggregation of the intermediate results, although the total contribution of this overhead remains low for large problem sizes. Moreover, the communication overhead grows linearly with the number of cores used. Consequently, the findings of Huang et al. [69] can be confirmed, who concluded that the data-set size should be large enough for the communication overhead to be hidden by the calculation time. This ensures profitable parallelization in terms of running-time reduction.

### 3.5.3 Strong scalability

This set of simulations is meant to analyze the impact of increasing the number of computing cores for a given problem size, i.e., the number of transmitters deployed over the target area does not change, but only the number of worker processes used is increased. Here, the following number of transmitters are tested  $\{1,280, 2,560, 5,120\}$ , by gradually doubling the number of workers from 1 to 64 for each problem size.

#### 3.5.3.1 Results

Similar to the weak-scalability experiments, these time measurements show that when applying a classic MW approach the running-time reduction starts flattening when more than eight worker processes are used. Moreover, the running times for 16, 32 and 64 worker processes are the same, i.e., it does not improve due to the master process being saturated. In contrast, when using the proposed MWD technique, the running-time reduction improves for up to 32 worker processes, after which there is no further improvement since the network is being fully exploited. These results clearly show that when applying parallelization using a larger number of worker processes, the master process becomes the bottleneck of the MW approach. When using the MWD configuration, a steady running-time reduction is observed, until a hardware constraint is hit, e.g., the network infrastructure.

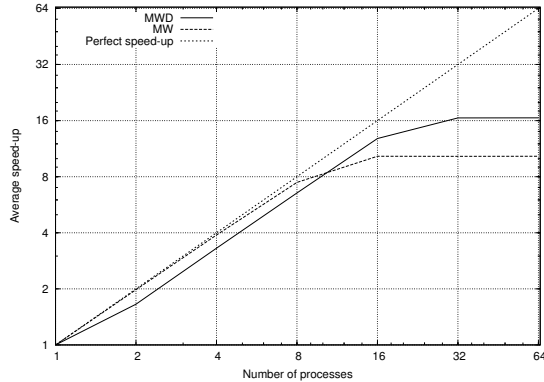


Figure 3.13: Average speedup for strong-scalability experiments. Both axes are expressed in a base-2 logarithmic scale.

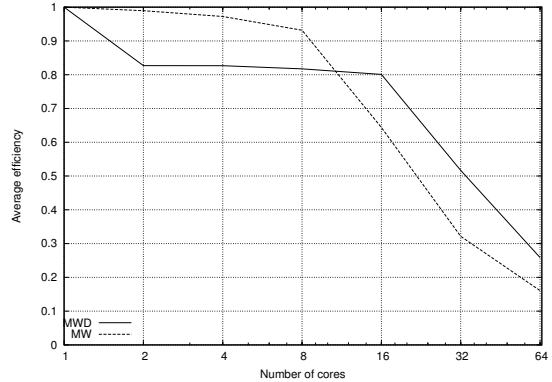


Figure 3.14: Average parallel efficiency for strong-scalability experiments. The parallel-efficiency axis is expressed in a linear scale, whereas the axis representing the number of cores is expressed in a base-2 logarithmic scale.

The overhead of sending/receiving asynchronous messages in order to support heterogeneous systems was also measured. It was found that this overhead never exceeds 0.02% of the total running time for the MW experiments, and 0.01% for the MWD experimental set.

### 3.5.3.2 Speedup

In order to further analyze how well the PRATO scales using the MW and MWD approaches, the performance of the parallel implementation in terms of its speedup was measured, which is defined as

$$S(NP) = \frac{\text{execution time for base case}}{\text{execution time for } NP \text{ cores}}, \quad (3.5)$$

where  $NP$  is the number of cores executing the worker processes. The parallel implementation running on only one core was the base case for comparisons. The serial implementation is not a good base comparison for the parallel results as it does not reuse the resources between each transmitter-coverage calculation and it does not overlap the I/O operations with the transmitter computations. In practice, this means that several concatenated runs of the serial version would be considerably slower than the single-worker configuration.

Using the speedup metric, linear scaling is achieved when the obtained speedup is equal to the total number of processors used. However, it should be noted that a perfect speedup is almost never achieved, due to the existence of serial stages within an algorithm and the communication overhead of the parallel implementation.

Figure 3.13 shows the average speedup of the parallel implementation for up to 64 worker processes, using the standard MW method and the proposed MWD approach. The average speedup was calculated for the three different problem instances, i.e., 1,280, 2,560, and 5,120 transmitters deployed over the target area. The number of transmitters used in these problem sizes is comparable to several real-world radio networks that were already deployed in England, e.g., Hampshire County with 227 base stations, West Midlands with 414 base stations, and Greater London Authority with 1,086 base stations [101]. Note that it is common for a single base station to host multiple transmitters.

The plotted average speedup clearly shows the minimal overhead of the MWD approach when using a small number of worker processes. This overhead accounts for the final aggre-

gation of the intermediate results at the DB, which in the MW configuration is performed along worker processing. Like before, the DB component allows the parallel implementation to fully exploit the available computing resources when deploying a larger number of worker processes, until the network-capacity limit is met. Of course, these results are directly correlated with the wall-clock times shown in Figure 3.12.

### 3.5.3.3 Efficiency

Another measure to study how well PRATO utilizes the available computing resources considers the parallel efficiency of the implementation. The definition of parallel efficiency is as follows

$$E(NP) = \frac{S(NP)}{NP}, \quad (3.6)$$

where  $S(NP)$  is the speedup as defined in Equation (3.5), and  $NP$  is the number of cores executing worker processes. Figure 3.14 shows the average parallel efficiency of the parallel implementation for different problem sizes as the number of processing cores was increased. Like for the speedup measure, the average parallel efficiency from the three problem instances was calculated.

The ideal case for a parallel application would be to utilize all the available computing resources, in which case the parallel efficiency would always be equal to one as the core count increases. From the plot in Figure 3.14, it can be observed that the efficiency of the MWD approach is better than in the MW case for larger number of processes and as long as there is still capacity at the LAN level. In accordance to the previous analysis, the under utilization of the computing resources is more significant when the master process is overloaded (in the MW case) than when the network infrastructure is saturated (in the MWD case). The lower efficiency is directly proportional to the number of idle worker processes that wait either for the master process (MW case) or for network access (MWD case).

Overall, the experimental results confirm that the objective of fully exploiting the available hardware resources is accomplished when applying the presented MWD approach, thus improving the scalability and efficiency of PRATO when compared to a traditional MW technique.

### 3.5.4 GPU performance

## 3.6 Summary

PRATO, a parallel radio-coverage prediction tool for radio networks, has been presented in this chapter. The tool is intended to be used for radio-network planning analysis and decision support. Its high-performance capabilities make it ideal for automatic-optimization tasks that require a large number of evaluations.

The parallel implementaion of PRATO includes a novel parallel technique for master-worker configurations. The introduced MWD technique, which combines the use of a database system with a work-pool approach, delivers improved performance when compared with a traditional MW setup. Moreover, the presented system provides parallel and asynchronous computation that is completely independent of the GIS used, in this case the GRASS environment. Consequently, a GIS installation is only needed on the master node, thus simplifying the required system setup and greatly enhancing the applicability of this methodology in different environments.

The extensive simulations, performed on the DEGIMA cluster of the Nagasaki Advanced Computing Center, were analyzed to determine the level of scalability of the implementation,

as well as the impact of the presented methods for parallel-algorithm design aimed at spatial-data processing. The conducted analyses show that when using the MWD approach, PRATO is able to calculate the radio-coverage prediction of real-world mobile networks in a reduced amount of time. Moreover, the experimental results show PRATO has a better scalability when using the MWD approach than the standard MW setup, since it is able to completely saturate the network infrastructure of the computer cluster. These promising results also show the great potential of the MWD approach for parallelizing different time-consuming problems dealing with spatial data, where databases form an intrinsic part of almost all GIS. Furthermore, the automatic optimization of radio networks, where millions of radio-propagation predictions take part in the evaluation step of the optimization process, is also an excellent candidate for PRATO. Indeed, this last point will be further discussed and validated in the following chapters.

The performance of the worker processes has been additionally improved by including the implementation of the radio-propagation algorithm on GPU. The use of GPU hardware is optional, i.e., it is exploited only if it is available on the computing nodes that host the worker processes. The experimental simulations showed a significant speedup due to the GPU implementation, when compared to the CPU-only version.

To the best of the author's knowledge, neither the MWD parallel technique nor the GPU implementation of the radio-prediction algorithm as presented in this chapter, have yet been described in the related literature.

## Part II

# Experimental evaluation



## 4 The service-coverage problem

The high-performance of PRATO, the radio-coverage simulation framework presented in Chapter 3, allows dealing with big problem instances in a reduced amount of time. Additionally, it enables tackling optimization problems that, because of their size, are out-of-reach of traditional approaches, mainly due to the computational-time complexity of their objective-function evaluation.

In this chapter, the challenge is to exploit PRATO for solving one of the classic optimization problems of radio networks: the service-coverage problem. Considering the minimization of the total amount of pilot power subject to a full coverage constraint, a novel optimization approach is introduced. The presented method, based on parallel autonomous agents, gives very good solutions to the problem in an acceptable amount of time. The parallel implementation takes full advantage of GPU hardware in order to achieve considerable speed-up. The interpretation of the experimental results, considering six real-world, radio networks of different sizes, analyzes solution-quality and performance aspects.

The content of this chapter extends the research work published by the author in [15] and [17]. The rest of this chapter is organized as follows. Section 4.1 gives a description of the coverage problem and its motivation from the mobile operator's perspective. In Section 4.2, a short overview of related research works is given, before formally introducing the key elements of the service-coverage problem in Section 4.3. The parallel-agent approach, as well as the strategies used for result comparison, are presented in Section 4.4. A detailed description of the tested implementations is given in Section 4.5, followed by the simulations and their analyses in Section 4.6.

### 4.1 Motivation

Solving the service-coverage problem for radio networks has received a great deal of attention in the past years. Its complexity demands the confluence of different skills in areas such as propagation of radio signals, telecommunications and information systems, among others.

Even several decades after the launch of the first commercial GSM network, service-coverage planning remains a key problem that all mobile operators have to deal with. Its intricacy arises from the wide range of different combinations of configuration parameters and their evaluation-time complexity. One crucial parameter, which is mainly subject of adjustment, is the transmit power of the pilot signal (see Section 2.4.3).

Regardless of the mobile technology used, e.g., GSM, UMTS or LTE, reducing pilot-power usage is related to issues regarding human exposure to the electromagnetic fields generated by base-station antennas [41]. During the past few years, public opinion has been extremely sensitive regarding this issue, and thus many countries have already imposed safety standards to limit the electromagnetic field levels produced by antennas in a given range.

From the UMTS perspective, minimizing pilot-power usage leaves more power available for increased network capacity. This is especially important if the traffic and other channels are configured relative to CPICH [66]. Moreover, as the users' demand for internet access

and data services increases [35], so does the pressure on existing network infrastructure, making parameter optimization the only viable solution in the short-term [96].

The idea of using autonomous agents for optimization is not new. It has proven to be a solid optimization approach for solving different types of problems, not only within the area of mobile networks [41, 28], but also in other fields [146, 142]. Nevertheless, we have not observed any similar optimization method for solving the service coverage problem in mobile networks [15]. Moreover, this is, to the best of our knowledge, the first work to experiment with optimization of a real UMTS network on a fully-enabled GPU environment.

Our optimization approach is based on a state-of-the-art mathematical model, that has been previously used to solve a comparable problem [24]. The large computational-time complexity when dealing with big problem instances is tackled using a parallel implementation of the agent-based algorithm for on GPU. This minimizes the overhead when deploying a larger number of agents working in parallel over the service area, only limited by the amount of memory available.

The algorithm on subsets of a real radio network deployed in Slovenia. All network-related data were provided by the engineers of the Radio Network department at Telekom Slovenije, d.d. The results show that the solution quality is greatly improved, compared to classic coverage-planning techniques.

## 4.2 Related work

There are several approaches in the literature that address solving the service-coverage problem in radio networks [96, 116]. Some of them even claim to achieve near-optimal solutions [124]. As a matter of fact, most formulations are only useful for small network instances and often fail when challenged with larger and real-world networks.

A genetic-algorithm approach for solving the service-coverage problem for GSM networks is presented in [86]. The proposed solution is based on the physical distribution of base-station antennas in order to maximize coverage. The simulations were performed on a test network with 40 candidate sites for base-station antennas.

In [124], Siomina and Yuan considered the problem of minimizing the total amount of pilot power for UMTS networks subject to a full coverage constraint. They tackled the problem with an iterative linear-programming approach, reporting very good results for some test networks, containing from 15 to 65 base stations. The authors noted that bigger problem instances could not be solved because of hardware constraints on the target platform.

As for LTE networks, the service-coverage problem is presented in [137]. The authors presented an algorithm, based on reinforcement learning, to tackle three aspects of the coverage problem, i.e., coverage holes, weak coverage and pilot pollution. The experimental simulations, performed on 3 base stations, used different antenna-tilt configurations as the proposed solutions. The service-coverage problem as presented in this chapter corresponds to achieving full coverage of the target area, i.e., without coverage holes.

## 4.3 Radio-network model

Extending the representation of a radio-network model from [97], this section addresses the definitions of all the elements included in the mathematical model used for the simulations.

Our goal here is to analyze the state of the network in a given situation, e.g. a ‘snapshot’ at an arbitrary instance. A snapshot consists of a set of mobiles (or users) having individual properties, such as location, and equipment type. The static approach inherently ignores dynamic effects that influence the system, like fast power control [97].

For additional information regarding mathematical models of comparable problems, see [96].

### 4.3.1 Basic elements

Consider a UMTS network with a set of antenna installations (cells),  $C$ . A RSG of a given resolution represents a geographical area,  $A_{\text{total}}$ , within which a set of mobiles,  $M$ , is spatially distributed over the pixels of  $A_{\text{total}}$ . Further,  $L_{cm}^{\downarrow}$  is defined as the downlink attenuation factor between cell  $c \in C$  and mobile  $m \in M$ . Similarly,  $L_{mc}^{\uparrow}$  represents the uplink attenuation factor between mobile  $m$  and cell  $c$ . The attenuation factor values are calculated by performing signal propagation predictions for every pair  $(c, m)$ ,  $c \in C$ ,  $m \in M$ , using the radio-propagation model introduced in ...???. These predictions already include losses and gains from cabling, hardware, and user equipment.

The amount of power allocated to the pilot signal of cell  $c$  is denoted as  $p_c$ , and it can adopt any value from the sorted set of available pilot power levels,  $P_c = \{p_c^1, p_c^2, \dots, p_c^k\}$ , where  $p_c^k$  is the maximum power.

Based on the introduced elements, the received pilot power from cell  $c$  to mobile  $m$  is  $L_{cm}^{\downarrow} p_c$ .

### 4.3.2 Coverage

A mobile  $m$  within the area  $A_{\text{covered}}$  is under service coverage, meaning that, at least, one cell  $c$  covers it. Cell coverage is provided to a mobile  $m$  from a cell  $c$  if its signal-to-interference ratio,  $sir(c, m)$ , at the RSG pixel where  $m$  is located, is not lower than a given threshold,  $\gamma^{\text{cov}}$ , i.e.,

$$sir(c, m) = \frac{L_{cm}^{\downarrow} p_c}{\tau_0 + \sum_{i \in C} L_{im}^{\downarrow} p_i} \geq \gamma^{\text{cov}} \quad (4.1)$$

where  $\tau_0$  is the thermal noise. For convenience, a binary function is defined to determine the coverage of a mobile  $m$  by a cell  $c$ . So, for any pair  $(c, m)$ ,  $c \in C$ ,  $m \in M$ , the coverage of mobile  $m$  by cell  $c$  is defined as

$$cov(c, m) = \begin{cases} 1 & \text{if } sir(c, m) \geq \gamma^{\text{cov}} \\ 0 & \text{otherwise} \end{cases}. \quad (4.2)$$

A set, denoted as  $C_m$ ,  $C_m \subset C$ , contains all the cells covering a mobile  $m$ . From this set, the cell with the highest  $sir(c, m)$  is referred to as the best server, and denoted as  $c_m^*$ . Notice that the described radio-network model is easily adaptable for different mobile technologies, e.g., GSM, UMTS and LTE. For example, if solving the service-coverage problem for UMTS, it would be reasonable to assume that all cells in the network operate at maximum power, and adapt Equation (4.1) accordingly. This is, from the interference point of view, the worst-case scenario [124, 24]. This assumption guarantees, that even under heavy user traffic, full coverage of the service area is maintained due to the cell-breathing principle [66].

### 4.3.3 Problem complexity

It has been proved that the problem of pilot power optimization for full coverage of the service area is  $NP$ -hard, since it can be reduced to the set-covering problem [145]. Consequently, as long as  $P \neq NP$ , it is unfeasible that a polynomial-time algorithm exists, which is able to find an exact solution to this problem.

### 4.3.4 Optimization objective and constraints

In the problem of optimization of pilot powers for service coverage, the objective is to find a set of pilot power settings for all cells in the network, such that the total pilot power used is minimized, and a given service coverage criteria is fulfilled. In other words, solving the service-coverage problem corresponds to finding the pilot power levels  $p_c$ , for all cells  $c \in C$ , such that coverage of at least  $b$  mobiles is guaranteed, while the total amount of pilot power used is minimized. Since we are considering full coverage, we denote  $b = |M|$ . Consequently, the optimization objective is defined as follows

$$P^* = \min \sum_{c \in C} p_c, \quad (4.3)$$

subject to

$$\frac{\sum_{m \in M} \text{cov}(c_m^*, m)}{b} = 1. \quad (4.4)$$

## 4.4 Optimization approaches

Since the problem instances we will be analyzing are part of a real mobile network deployed in Slovenia by Mobitel Telecommunication Services, Inc., there are no references in the literature of other optimization techniques dealing with exactly the same data set. For this reason, we will introduce two different strategies for setting the pilot power, that shall enable us to compare the experimentation results. The first strategy is the attenuation-based pilot power, presented in [116], in which a pixel of the service area is always covered by the cell with the maximum attenuation-factor value, i.e., the minimum path loss. The second strategy is our parallel-agent approach, based on ideas inspired by two-dimensional cellular automata [110] and metaheuristics [134]. A detailed description is given in Section 4.4.2.

Similar criteria for result comparison have also been used in [124].

### 4.4.1 Attenuation-based approach

The first heuristic for setting the pilot power of all cells in the network is known as attenuation-based, since it relies on the downlink-attenuation factor,  $L_{cm}^\downarrow$ . A mobile located on some pixel of the service area is always covered by the cell with the maximum  $L_{cm}^\downarrow$ . Whenever the maximum available power,  $p_c^k$ , is the same for all the cells in the network, this is equivalent to selecting the cell with the minimum required pilot power to cover a mobile  $m$ . Hence, under this assumption, we identify the cell  $c$  covering mobile  $m$  as

$$p_{cm}^{\text{att}} = \min p_c \forall c \in C \iff \text{cov}(c, m) = 1 \quad (4.5)$$

Picking the cells conforming to Equation (4.5) and setting the pilot powers accordingly, we achieve full coverage of the service area with a solution exhibiting a total pilot power of

$$P^{\text{att}} = \sum_{c \in C} \max p_{cm}^{\text{att}} \quad (4.6)$$

The procedure to find a cell  $c$  for every mobile  $m$  in the service area consists in sorting, in descending order, all mobiles by their attenuation-factor values,  $L_{cm}^\downarrow$ . The solution is thus established by the first  $b$  mobiles of the sorted sequence, taking the maximum pilot-power setting for a cell into account, i.e.,  $p_{cm}^{\text{att}}$ .

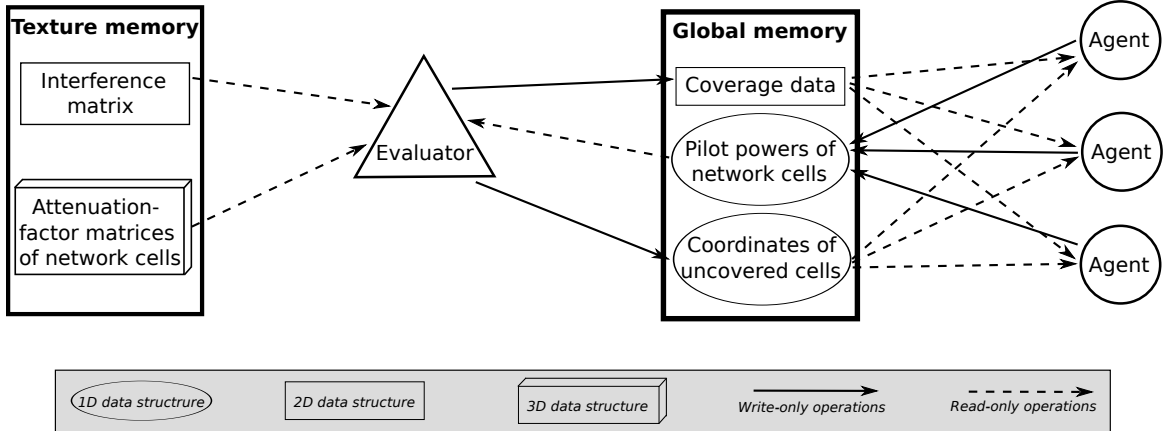


Figure 4.1: Architecture of the optimization system on GPU.

#### 4.4.2 Parallel-agent approach

In the parallel-agent approach, a set of autonomous worker agents explore the target geographical area,  $A_{\text{total}}$ , in order to optimize the pilot-power consumption. Each agent randomly moves over the  $A_{\text{total}}$  as it dictates different changes to the pilot power of the cells. PRATO, the radio-coverage framework presented in Chapter 3, performs the objective-function evaluations and radio-propagation predictions based on the proposed changes of each agent.

The moving process during the optimization is strictly random. However, several physical properties that are exclusive to the service-coverage problem are being exploited during the exploration of the search space. Additionally, whenever the current solution breaks any of the given constraints, the optimization process is guided back to the space of valid solutions, providing a mechanism for improving exploration and escaping from local optima.

Because the behaviour of the agents is independent between each other, a parallel implementation is fairly straight-forward to achieve. Figure 4.1 shows the architecture of the agent-optimization system. In this GPU-only architecture, agents work in a parallel and autonomous manner, while the evaluator reacts to their changes.

##### 4.4.2.1 The agents

The agents apply the pilot-power changes only considering local information. Each of them encapsulates a set of steps that is consistently applied as it randomly moves through the service area of the network. Whenever an agent arrives at a new location, the set of cells covering it is calculated, e.g.,  $C_m$ .

The step set an agent applies following this point is directly related to  $|C_m|$ , whereas its movement is determined by  $|\overline{A}_{\text{covered}}|$ , i.e., the cardinality of  $\overline{A}_{\text{covered}}$ , denoting the locations without service coverage.

The behavior of an agent is dictated by the pseudo-code shown in Algorithm 4.1. The first four steps are responsible for guiding its movements. The coordinates are randomly selected from two sets,  $A_{\text{total}}$  and  $\overline{A}_{\text{covered}}$ . Only “special” agents may move to a location without service coverage, and they apply the step set  $SS_0$  for as long as the solution is not valid. The portion of “special” agents used for correcting a solution is a parameter of the optimization process. During the following steps of Algorithm 4.1, the agent applies step sets  $SS_0$  and  $SS_1$  based on the number of cells in  $C_m$ .

If the current location of the agent is not covered by any cell, i.e.,  $|C_m| = 0$ , the step set  $SS_0$  is applied (see Algorithm 4.2). At the beginning, the cell with the highest attenuation

---

**Algorithm 4.1** Pseudo-code representing the behaviour of an agent.

---

```

repeat
  if is_special_agent() and  $\overline{A_{\text{covered}}} > 0$  then
     $l \leftarrow \text{pick\_random\_location}(\overline{A_{\text{covered}}})$ 
  else
     $l \leftarrow \text{pick\_random\_location}(A_{\text{total}})$ 
  end if
  move(l)
  if  $|C_m| = 0$  then
    apply(SS0)
  else
    if  $|C_m| \geq 1$  then
      apply(SS1)
    end if
  end if
until stopping_criterion()

```

---

**Algorithm 4.2** Pseudo-code representing step set  $SS_0$ , which is applied by the agents in areas with no service coverage.

---

```

repeat
   $c' \leftarrow \text{next\_cell\_with\_max\_att}(m)$ 
   $p_{c'} \leftarrow \text{adjust\_power}(c', \text{inc\_rate})$ 
until  $p_{c'} \in P_{c'}$ 

```

---

factor that may cover a mobile at this location,  $c'$ , is selected. If several cells have the same  $L_{cm}^{\downarrow}$  value, one of them is randomly chosen. Once  $c'$  is uniquely identified, the agent changes its pilot power by *incrate* dB.

The step set  $SS_1$  listed in Algorithm 4.3 is applied if the location of the agent is under the coverage of one or more cells, i.e.,  $|C_m| \geq 1$ . The first step randomly selects a cell from the set  $C_m$ , followed by a decrease of the pilot power of cell  $c'$ . This practice keeps the coverage constraint valid over  $A_{\text{total}}$ , although it might potentially break it on other areas. Ideally, every pixel of the geographical area has to be covered by exactly one network cell, although this is just a representation of a perfect solution that is unreachable because of irregularities in network topology and terrain.

In both step sets,  $SS_0$  and  $SS_1$ , the agent makes sure that the new pilot power setting,  $p_{c'}$ , is an element of  $P_{c'}$ . If this is not the case, cell  $c'$  is discarded and another cell is repeatedly selected at the beginning of both step sets, until this condition is satisfied.

The values *incrate* and *decrate* are configurable parameters that should be set before starting the optimization process. They indicate the relative adjustment (expressed in dB) of the pilot power of cell  $c'$ . On the one hand, lowering the pilot power of a cell decreases the interference it creates within its coverage area and those of their neighbours. Since

**Algorithm 4.3** Pseudo-code representing step set  $SS_1$ , which is applied by the agents in areas with service coverage.

---

```

repeat
   $c' \leftarrow \text{pick\_random\_cell}(C_m)$ 
   $p_{c'} \leftarrow \text{adjust\_power}(c', \text{dec\_rate})$ 
until  $p_{c'} \in P_{c'}$ 

```

---

the  $sir(c, m)$  value increases with lower interference, the coverage of  $m$  may be achieved by a neighbor cell with the same or lower pilot power. On the other hand, increasing the pilot power of the cell with the maximum attenuation factor improves coverage by evenly distributing the power among different network cells. This cell is, on average, the nearest one to the location of a mobile  $m$ .

#### 4.4.2.2 The evaluator

The evaluator represents a central component of the optimization system. It reacts to the pilot-power changes by recalculating the objective-function value. Recall that the objective-function evaluation involves the radio-coverage prediction of the service area and the calculation of the total pilot power used by the cells in the target network.

After a short initialization, during which the attenuation-factor matrices of all the cells and the interference matrix are calculated, the evaluator computes the coverage of the service area based on the pilot powers supplied as the initial solution. Initial solutions are randomly generated from valid pilot-power settings that conform to the full coverage constraint.

The evaluator also maintains a special part of the memory (see “Coordinates of uncovered cells” in Figure 4.1) that is intended for registering uncovered areas, i.e.,  $\overline{A_{\text{covered}}}$ . If Equation (4.4) does not hold, the “special” agents randomly select a location from this portion of memory so that a valid solution may be reached again.

It is worth mentioning that the evaluator itself has no influence in the optimization process from a quality point-of-view. Its task is to provide feedback and updated information to the agents that move through the service area. From a performance point-of-view, the importance of the evaluator is significant, as it will be shown in the following sections.

## 4.5 Implementation

The evaluation of the objective function was completely implemented on the GPU using OpenCL (see Section 2.5.1). The reason behind this decision is the impact objective-function evaluation has on the performance of the optimization system as a whole, as discussed in Section 2.1.4. Agents’ implementation is also based on the GPU, which drastically reduces the number of data transfers between CPU and GPU, since all problem elements are available on the GPU during the optimization process. In this sense, the first hardware restriction we have to overcome is the amount of memory on the GPU device. Consequently, careful memory utilization and organization are critical to successfully accommodate all involved problem elements on the GPU, the memory of which is significantly smaller than the RAM memory available in desktop computers.

### 4.5.1 Parallel agents on GPU

With the objective-function evaluation running on the GPU, a new performance bottleneck appeared. The limitation factor in this case was the CPU-to-GPU data transfers in each iteration of the optimization process.

The GPU kernel of the agents is launched as one thread block that contains one thread per deployed agent. The thread block is organized in a one-dimensional grid. The initial location of each agent is randomly generated using the current system time as a random seed. Since OpenCL provides no function for random-number generation, a simplified version of Marsaglia’s generator [90] was implemented.

The analysis each agent does about the received signals at the current location is saved into shared memory of the thread block. It contains the network cell and its pilot-power

setting. Since both numbers are of type *short*, there is enough space in a 16 KB shared-memory block to allocate 4,096 agents. The last step involves saving the new pilot powers into global memory. This step is performed by only one of the threads within the thread block in order to avoid memory-access conflicts. Updated pilot powers are saved in negative form to indicate that coverage re-calculation is needed for these cells. In case there are several updated pilot powers for one network cell, the median is calculated and applied as the new pilot power.

Even though coalesced access is not achieved by the GPU kernel of the agents, its sole implementation provided enhanced performance. This performance gain appears because of the lower number of data transfers between the CPU and the GPU, since most data is available in global memory. Moreover, the GPU kernel also produces the truly parallel behavior of the agents, as they all apply the pilot-power changes at the same time.

## 4.6 Simulations

### 4.6.1 Test networks

The test networks, Net<sub>1</sub>, Net<sub>2</sub> and Net<sub>3</sub> are subsets of the real UMTS network deployed by Telekom Slovenije, d.d. The path-loss predictions were calculated using the ...??? model presented in Section ...???. A DEM of 100 m<sup>2</sup> resolution was used as the terrain-profile data. The requirements for the coverage threshold,  $\gamma^{\text{cov}}$ , were provided by experts of the Radio Network department of Telekom Slovenije, d.d.

Net<sub>1</sub> is deployed over a densely populated urban area. For this reason, the value of  $\gamma^{\text{cov}}$  is lower here, since network capacity is the dominating factor, whereas coverage is flexible because of a larger cell density, i.e., more base stations per surface unit. Net<sub>2</sub> represents a network deployed over a rural area, meaning that network capacity can be reduced at the cost of better coverage, since user density is lower. The last network, Net<sub>3</sub>, represents a suburban area with a densely populated, but relatively small, downtown center, where a compromise between network capacity and coverage has to be achieved.

The second group of test networks, including Net<sub>4</sub>, Net<sub>5</sub> and Net<sub>6</sub>, is part of the publicly available MOMENTUM project [71]. Test network Net<sub>4</sub> represents the city of Berlin (Germany), Net<sub>5</sub> represents the city of The Hague (Netherlands), and Net<sub>6</sub> is one of the networks optimized in [124], representing a reduced version of Net<sub>4</sub>. All networks include information about site locations, path-loss predictions and realistic antennas, which are part of the scenarios provided by the MOMENTUM project.

Network configurations, that represent what could be an initial-network setup by common planning standards [66], were produced using the attenuation-based approach. Such configurations can be easily calculated by a network planner. Table 4.1 lists the number of base stations and cells per test network, as well as the size of the geographical area. Different network-parameter values used during the simulations are shown in Table 4.2.

### 4.6.2 Parameter settings of the parallel-agent approach

The parameter settings for the optimization algorithm were determined after some experimentation with the test networks. The parameter settings for each test networks are listed in Table 4.3.

Using a higher *inc\_rate* than *dec\_rate* reflects the behavior of the agents when full coverage of the service area is not guaranteed. In practice, areas without service coverage usually appear as irregular islands. The stopping criteria were set by limiting the total number of pilot-power changes an agent is allowed to make. The value was set to 10,000

Table 4.1: Sizes of the test networks used, in terms of equipment and geographical area.

	Number of base stations	Number of cells	Surface (km <sup>2</sup> )	Resolution (m <sup>2</sup> )
Net <sub>1</sub>	26	77	100.00	25
Net <sub>2</sub>	8	23	306.25	25
Net <sub>3</sub>	45	129	405.00	25
Net <sub>4</sub>	65	193	56.25	50
Net <sub>5</sub>	12	36	16.00	50
Net <sub>6</sub>	50	148	56.25	50

Table 4.2: Network parameters of the test networks used.

	$p_c^k$	$\tau_0$	$\gamma^{\text{cov}}$
Net <sub>1</sub>	15.00 W	$1.55 \cdot 10^{-14}$ W	0.010
Net <sub>2</sub>	19.95 W	$1.55 \cdot 10^{-14}$ W	0.020
Net <sub>3</sub>	15.00 W	$1.55 \cdot 10^{-14}$ W	0.015
Net <sub>4</sub>	19.95 W	$1.55 \cdot 10^{-14}$ W	0.010
Net <sub>5</sub>	19.95 W	$1.55 \cdot 10^{-14}$ W	0.010
Net <sub>6</sub>	19.95 W	$1.55 \cdot 10^{-14}$ W	0.010

Table 4.3: Parameter settings of the parallel-agent approach for each test network.

	Agents	<i>inc_rate</i> (dB)	<i>dec_rate</i> (dB)	Pilot-power changes
Net <sub>1</sub>	16	0.2	-0.1	10,000
Net <sub>2</sub>	16	0.2	-0.1	10,000
Net <sub>3</sub>	16	0.2	-0.1	10,000
Net <sub>4</sub>	6	1.0	-0.1	10,000
Net <sub>5</sub>	2	1.0	-0.1	10,000
Net <sub>6</sub>	6	1.0	-0.1	10,000

Table 4.4: Optimization results of all test networks after applying different approaches for solving the service-coverage problem. All values are expressed in Watts.

Attenuation-based			Parallel agents	
	Total power	Average pilot power	Total power	Average pilot power
Net <sub>1</sub>	419.292	5.445	137.064	1.780
Net <sub>2</sub>	78.297	3.404	33.344	1.450
Net <sub>3</sub>	1,014.113	7.861	582.954	4.519
Net <sub>4</sub>	179.876	0.932	145.715	0.755
Net <sub>5</sub>	73.872	2.052	34.884	0.969
Net <sub>6</sub>	147.014	0.993	112.332	0.759

even though for some of the test networks the best solutions were found in the first quarter of the experiment.

#### 4.6.3 Experimental environment

All experiments were performed on a multi-core Intel i7 2.67 GHz desktop computer with 6 GB of RAM running a 64-bit Linux operating system. The GPU hardware used was an ATI HD5570 with 1 GB of DDR3 RAM. The implementation language used was C, combined with OpenCL and OpenMPI extensions.

### 4.7 Results

The results achieved by the parallel-agent approach, listed in Table 4.4, improved the optimization objective significantly. They show that the pilot-power usage was reduced in all networks while the service area was kept under full coverage. Moreover, the parallel-agent solution for Net<sub>1</sub> improved the attenuation-based setting by more than 300%. As for Net<sub>2</sub>, the observed improvement is around 232%, while the improvement for Net<sub>3</sub> is more than 170%.

As mentioned in Section 4.1, the interpretation of these results depends on the mobile technology used. For GSM, ...???. As for UMTS, the capacity of the network has been significantly increased in all three problem instances. Therefore, a greater number of users should be able to access services provided by the mobile network, since coverage is assured. Moreover, an increased speed in data services should be observed [66]. Regarding LTE, ...???

The last test network, Net<sub>6</sub>, is the same as in [124]. When comparing these results to those of [124], an improvement of almost 3% can be observed in the solution provided by the parallel-agent approach.

#### 4.7.1 Convergence analysis

The graphs shown in Figures 4.2, 4.3 and 4.4 depict the convergence of the parallel-agent approach after ten independent runs for test networks Net<sub>1</sub>, Net<sub>2</sub> and Net<sub>3</sub>, respectively. Only feasible solutions were plotted, i.e., solutions that meet the full-coverage constraint. Unfeasible solutions were marked with a value of inferior quality than the worst solution found: 428 for Net<sub>1</sub>, 129 for Net<sub>2</sub>, and 1,435 for Net<sub>3</sub>.

From the graphs of Net<sub>1</sub> (see Figure 4.2) and Net<sub>2</sub> (see Figure 4.3) a good initial convergence can be observed. This is followed by a steady improvement of the intermediate

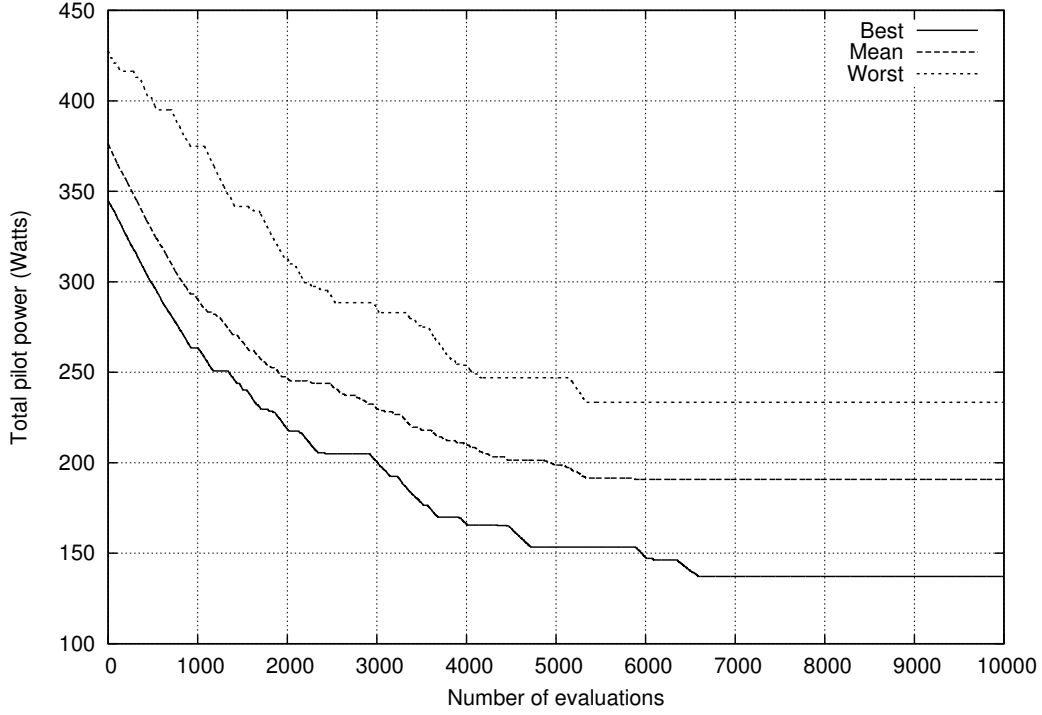


Figure 4.2: Convergence profile of the parallel-agent approach for the test network  $\text{Net}_1$ .

solutions. In  $\text{Net}_1$ , an additional solution improvement is noticed towards the end of the optimization process. This fact suggests that longer runs would potentially find better solutions in this case. In contrast, the solution improvement of  $\text{Net}_2$  shows a flat profile towards the end, suggesting that the stopping criterion is appropriate for this instance. From the graph of  $\text{Net}_3$  (see Figure 4.4), a slower initial convergence, followed by a steady improvement of intermediate solutions and no significant solution enhancement towards the end, can be observed. This convergence profile suggests that this problem instance presents a more difficult optimization case than for  $\text{Net}_1$  and  $\text{Net}_2$ . Indeed, this is the largest test network in terms of surface area. However, further investigation is needed to confirm this hypothesis. Nevertheless, the parallel-agent approach improved the pilot-power usage of this test network by almost 75%.

#### 4.7.2 Performance analysis

In this section, the performance analysis of the experimental results is presented. This analysis covers the running times during the optimization of the first three test networks, i.e.,  $\text{Net}_1$ ,  $\text{Net}_2$  and  $\text{Net}_3$ . The running times were measured for each implementation, and the average times, calculated after ten independent runs, are given. The number of pilot-power changes per agent was limited to 100, while all other algorithm parameters were kept at the same values as in Section 4.6.2.

Table 4.5 lists the average wall-clock times in seconds for the different implementations and test networks. The implementations include: the CPU-MPI implementation that consists of objective-function evaluation on CPU and parallel agents over MPI, the GPU-MPI implementation that consists of objective-function evaluation on GPU and parallel agents over MPI, and the GPU-GPU implementation that consists of objective-function evaluation and parallel agents on the same GPU. The CPU-MPI implementation is the basis for the speed-up calculation of the other implementations.

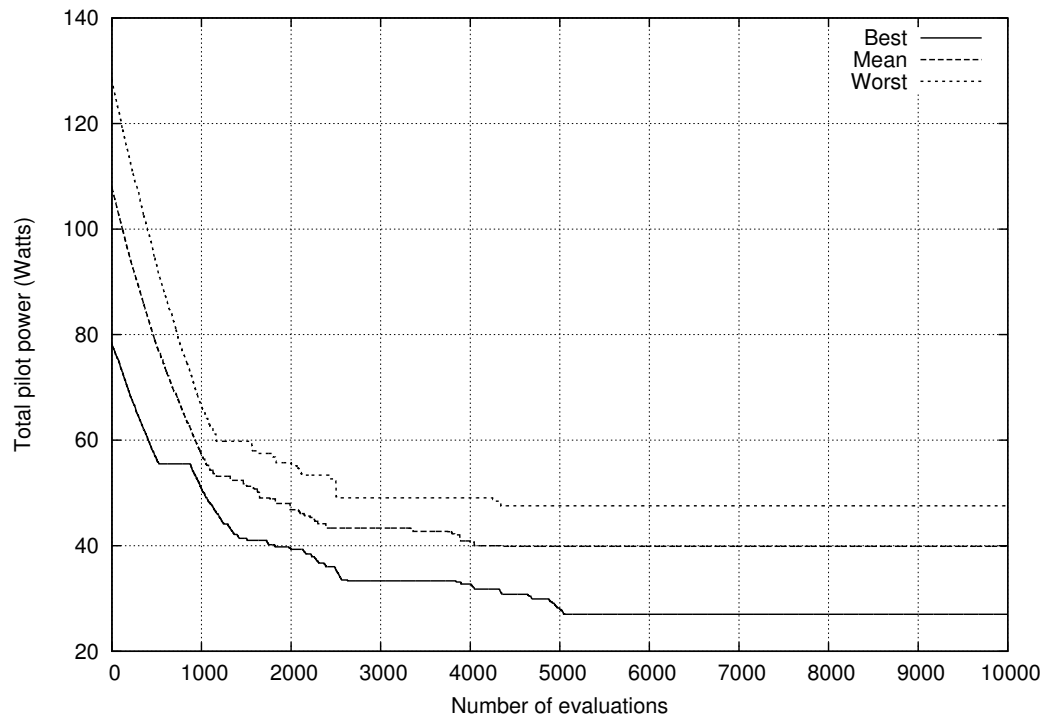


Figure 4.3: Convergence profile of the parallel-agent approach for the test network Net<sub>2</sub>.

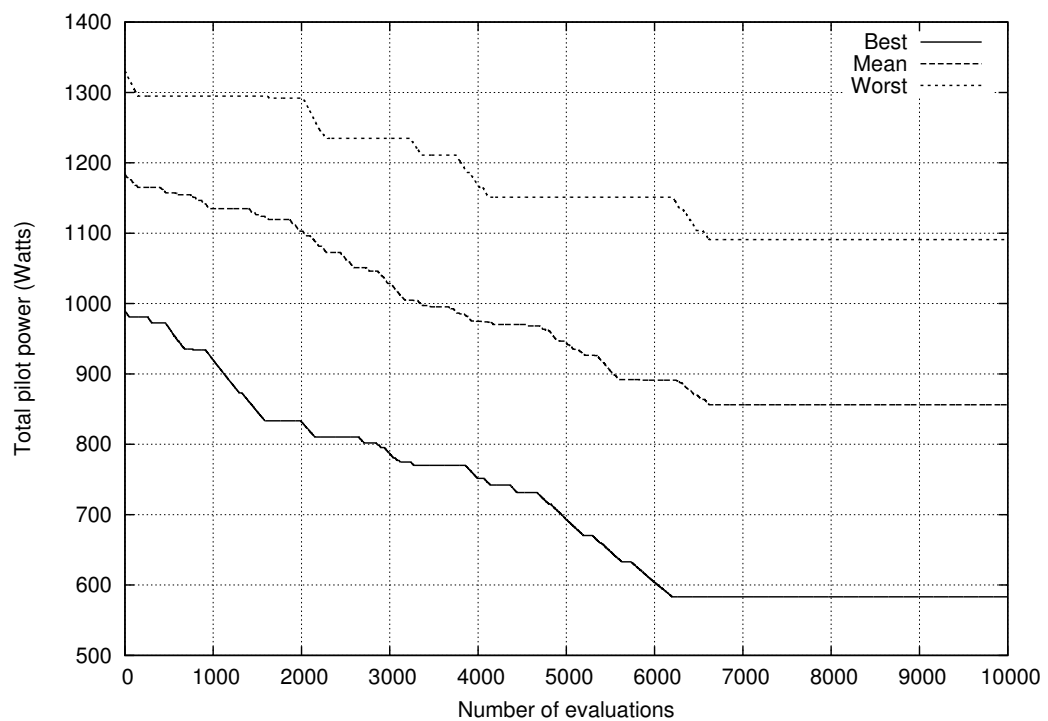


Figure 4.4: Convergence profile of the parallel-agent approach for the test network Net<sub>3</sub>.

Table 4.5: Wall-clock times (in seconds) and speed-up factors for different implementations of the objective-function evaluation and the parallel agents.

	CPU-MPI		GPU-MPI		GPU-GPU	
	Avg. time		Avg. time	Speed-up	Avg. time	Speed-up
<i>Net</i> <sub>1</sub>	105,455		346	305x	67	1574x
<i>Net</i> <sub>2</sub>	33,700		195	173x	46	733x
<i>Net</i> <sub>3</sub>	191,900		506	379x	117	927x

Function evaluation on the GPU communicating with agents over MPI provides the second measured setup. The evaluator implementation takes advantage of shared memory for thread collaboration within a thread block and texture memory for constant elements, as is has been explained in Section ???. Still, the speed-up is considerable but improvable, since numerous data transfers between CPU and GPU are needed for the agents to access optimization-related information.

The last result set presents measurements for complete GPU implementation, including objective-function evaluation and agents on the same device. The substantial speed-up delivered by this combination highlights the great impact that CPU-to-GPU memory transfers have on overall system performance. This fact is supported by the speed-up between the second and third measured setups, which exhibit, on average, an improvement of more than 400%.

## 4.8 Summary

This chapter presented a novel optimization approach for solving the well-known service-coverage problem in radio networks. The problem addressed the full coverage of a geographical area using a minimum amount of pilot power. The newly introduced parallel-agent approach was successfully tested in six networks that represent real-world scenarios of deployed radio networks. The experimental results show that the parallel-agent approach is able to find better solutions than other common radio-planning methods [66]. Moreover, the algorithm successfully tackled larger networks, thus overcoming the obstacles of other state-of-the-art optimization methods regarding problem-instance size [124].

Compared to a different optimization approach in the literature [123], the solution-quality of the parallel-agent approach showed an improvement. The proposed solutions, calculated for the same problem instance as in [124], were improved at the cost of longer running time. It worth mentioning that it is feasible for the optimization algorithm to take a longer time to reach the solution, since design problems, as the service-coverage one, are usually solved offline.

Different implementations of the parallel-agent approach, combining a serial version on CPU, parallel processes over MPI and GPU kernels, were presented. In particular, GPU architectures enable the implementation of parallel heuristics in a natural way while substantially improving the computational-time performance. To the best of the author's knowledge, the GPU implementation of the parallel-agent approach as presented in this chapter, has not yet been described in the related literature.



## 5 The SHO-balancing problem

In Chapter 4, an application exploiting the advantages of faster evaluation methods has been introduced. Solving the service coverage problem for real-world networks capitalizes on the ability to tackle bigger problem instances, i.e. problems that, because of their size, were previously unsolvable in a feasible amount of time. This improved performance also allows solving optimization problems with a higher degree of complexity, usually represented by the evaluation of multi-dimensional non-convex objective functions.

This chapter focuses on solving a new optimization problem for 3G networks, dealing with downlink and uplink SHO areas (see Section 2.4.2). By introducing a penalty-based objective function and some hard constraints, the formal definition of the SHO-balancing problem in UMTS networks is given. The state-of-the-art mathematical model used and the penalty scores of the objective function are set according to the configuration and layout of a real mobile network, deployed in Slovenia by Telekom Slovenije, d.d. The balancing problem is then tackled by three optimization algorithms, each of them belonging to a different category of metaheuristics.

To the best of the author's knowledge, there is no reference in the literature to a simulation-based approach to find active downlink and uplink SHO areas. Additionally, there are no formal optimization methods known to the author that tackle the SHO balancing problem as described here. The approach described in this chapter extends the approach published by the author in [16].

The remainder of this chapter is organized as follows. Section 5.1 describes the motivation behind the SHO-balancing problem, whereas Section 5.2 gives an overview of other works related to CPICH-power and SHO optimization in UMTS. The static network model is presented in Section 5.3, where all the elements of the mathematical model and the objective function are defined. In Section 5.4 a short description of the optimization algorithms used is given, including their mapping to the SHO-balancing problem. The simulations, including their environment and parameter setup, are introduced in Section 5.5, followed by their results in Section 5.6.

### 5.1 Motivation

Despite several built-in mechanisms that allow the radio network to overcome different problems due to the lack of SHO during a HSDPA connection, some abnormal cases do arise, especially in those areas where there is SHO capability in the uplink, but none in the downlink. An example of such a case is depicted in Figure 5.1, which shows interference behavior during a HSPA connection in normal SHO conditions (a), and in unbalanced SHO conditions (b). Graph data are actual radio network statistics, taken from the mobile network deployed in Slovenia by Telekom Slovenije, d.d.. The graph on the left (a) shows a normal HSUPA-enabled service situation, in which the measured interference is proportional to the traffic being served. Note how the noise rises with the increased traffic on cell 1, while its neighbor (cell 2) has almost no interference nor traffic. Moreover, the graph profile for both traffic and noise of cell 1 are almost identical. The graph on the right (b) depicts a problematic

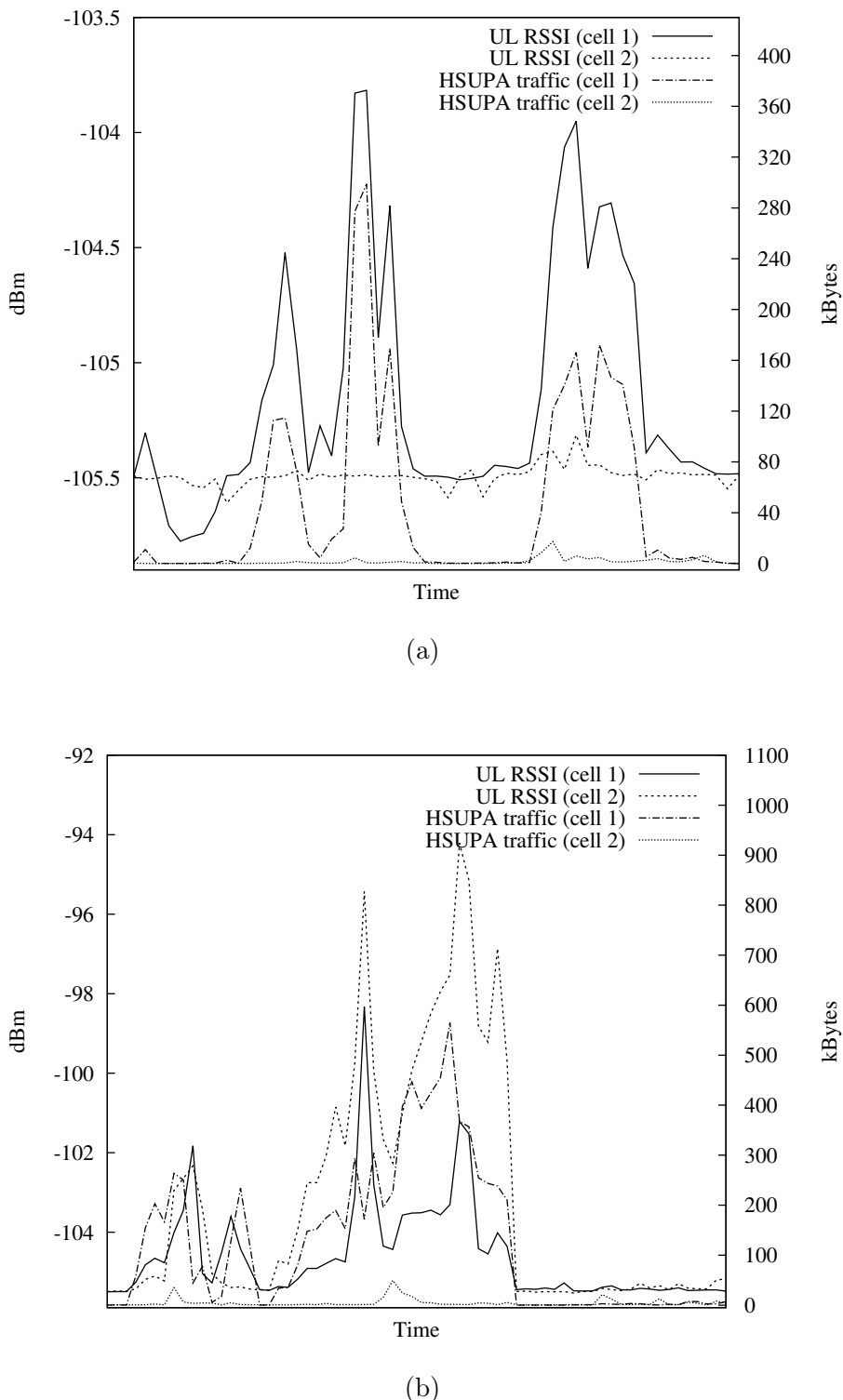


Figure 5.1: HSUPA traffic and uplink interference with: (a) balanced downlink and uplink SHO conditions; (b) unbalanced downlink and uplink SHO conditions.

situation, where the noise level does not only rise on the cell serving the HSUPA services (cell 1), but also on the neighboring one. Notice how the interference level rises on the cell that has almost no traffic (cell 2). It is clear that the source of this noise rise is generated by the active connection on cell 1, which shows an increase in HSUPA traffic. However, the noise level profile on cell 2 does not follow its traffic, as it did in the normal situation (a). This is due to cell 2 not being part of the active set. Such situations appear when the UL coverage is larger than the DL coverage. Interestingly enough, this seems to be an exceptional case, as Holma and Toskala write in [67] when describing soft handover in chapter 5:

"... There is no obvious reason why the serving E-DCH cell would not be the same as the serving HSDPA cell, and this is also required to be the case in the specifications."

Given the described context, the challenge is to achieve the correct balance or distribution of downlink and uplink SHO areas within a working UMTS network. Therefore, the network has to be fine-tuned to achieve a better SHO-area balancing, and thus avoiding the exceptional appearance of problematic situations as shown in Figure 5.1. This clearly implies that the mobile network configuration should not be excessively altered, since other aspects of the network are working well before starting the optimization process. Hence, the objective of the optimization problem is to find a CPICH power level configuration for all the cells in the target network, such that the balance of downlink and uplink SHO areas is improved and other network aspects are preserved. The optimization process takes into account different kinds of hardware (e.g. amplifiers, cables, and antennas), changing only the CPICH powers of the cells.

PRATO is used as the evaluation framework, as defined in Chapter 3. A state-of-the-art mathematical model [97] describes downlink and uplink SHO areas, considering the static nature of the evaluation. By introducing a penalty-based objective function and some hard constraints, a formal definition of the SHO-balancing problem in UMTS networks is given. The mathematical model and the penalty scores of the objective function are set according to the configuration and layout of a real mobile network, deployed in Slovenia by Telekom Slovenije, d.d. The SHO settings are also taken from actual network configuration, still they were adapted to closely model interference and other dynamics present in the network. The balancing problem is then tackled by three optimization algorithms, each of them belonging to a different category of metaheuristics. The optimization results, as well as the performance of each of the optimization algorithms used, are afterwards analyzed.

## 5.2 Related work

SHO optimization has received quite some attention from the scientific community in the last years. This mainly relates to the importance it has within deployed networks that provide high speed services such as video telephony [22] and Internet access by means of HSPA [27].

Some authors tackle optimization problems at the planning stage of the network [40, 50], considering, among other variables, base-station locations and hardware. However, most mobile operators are unable to apply these contributions to a live network since the planning phase has long been concluded. Moreover, the great majority of the base stations have already been deployed and their hardware also installed. Therefore, from the mobile operator's point of view, mainly parameter and software optimization are the only tools available when it comes to QoS improvement (see Section 2.4.2) and network troubleshooting in the short term.

Optimizing SHO by means of CPICH is an established way of enhancing network capacity when high speed services like HSDPA and HSUPA coexist with legacy technologies [25]. The CPICH transmit power is typically between 5% to 10% of the total downlink transmit power of the base station [81], but there is no standardized method to find a CPICH power

setting. A number of existing approaches to resolve this issue exist in the related literature (see [65, 114, 155]). The most effective ones are those based on optimization methods [40, 45, 81, 84, 119]. Such a wide spectrum of available procedures is directly related to the diverse criteria taken into account when assigning the CPICH power of a cell. The fundamental reason behind this fact is that the CPICH power is a common factor of various optimization problems in UMTS networks.

### 5.3 Radio-network model

Extending the representation of a radio-network model from [97], this section addresses the definitions of all the elements included in the mathematical model used for the simulations.

Our goal here is to analyze the state of the network in a given situation, e.g. a ‘snapshot’ at an arbitrary instance. A snapshot consists of a set of users (or mobiles) having individual properties, such as location, and equipment type. The static approach inherently ignores dynamic effects that influence the system, like fast power control [97].

The mathematical model links the SHO settings with CPICH power settings for each cell, the best-server pattern, and the network coverage.

For additional information regarding mathematical models of comparable problems, see [96].

#### 5.3.1 Basic elements

Consider a UMTS network with a set of antenna installations (cells),  $C$ . A RSG of a given resolution represents the service area,  $A_{\text{total}}$ , within which a set of mobiles,  $M$ , is spatially distributed over the pixels of  $A_{\text{total}}$ . Further,  $L_{cm}^{\downarrow}$  is defined as the downlink attenuation factor between cell  $c \in C$  and mobile  $m \in M$ . Similarly,  $L_{mc}^{\uparrow}$  represents the uplink attenuation factor between mobile  $m$  and cell  $c$ . The attenuation factor values are calculated by performing signal propagation predictions for every pair  $(c, m)$ ,  $c \in C$ ,  $m \in M$ , using the radio-propagation model introduced in ...???. These predictions already include losses and gains from cabling, hardware, and user equipment.

The amount of power allocated to the pilot signal of cell  $c$  is denoted as  $p_c$ , where  $p_c$  may adopt any value from a finite set of possible pilot power levels,  $P_c = \{p_c^1, p_c^2, \dots, p_c^K\}$ . Consequently, the received pilot power from cell  $c$  to mobile  $m$  is  $L_{cm}^{\downarrow} p_c$ .

Service coverage is guaranteed when a mobile has, at least, one cell covering it. Cell coverage is provided to a mobile  $m$  from a cell  $c$  if its signal-to-interference ratio,  $sir(c, m)$ , at the RSG pixel where  $m$  is located, is not lower than a given threshold,  $\gamma^{\text{cov}}$ , i.e.,

$$sir(c, m) = \frac{L_{cm}^{\downarrow} p_c}{\tau_0 + \sum_{i \in C} L_{im}^{\downarrow} p_i} \geq \gamma^{\text{cov}} \quad (5.1)$$

where  $\tau_0$  is the thermal noise.

The mathematical model links the SHO settings with CPICH power settings for each cell, the best-server pattern, and the network coverage.

By introducing a change step of 0.01 dB and bounding the CPICH power of a cell  $c$  to  $\pm 2$  dB, relative to the CPICH power setting the cell had before optimization, the number of elements in the set  $P_c$  is reduced. The purpose of this reduction is twofold. First, since the optimization targets a live network, there is no need for the algorithms to create complete new configurations, but just to fine-tune existing ones. Second, the problem complexity is lowered, because the size of the search space is smaller and discrete.

### 5.3.2 Coverage

A mobile  $m$  within the area  $A_{\text{covered}}$  is under network coverage if at least one cell  $c$  covers it. We define the downlink coverage by means of the received signal code power (RSCP) [65]. Following real network settings, and including a margin for interference derived from HSDPA [67], the RSCP threshold is set to  $-115 \text{ dBm}$  ( $3.16227766 \cdot 10^{-12} \text{ mW}$ ). So, for any pair  $(c, m)$ ,  $c \in C$ ,  $m \in M$ , the coverage of mobile  $m$  by cell  $c$  is defined as

$$\text{cov}_{cm} = \begin{cases} 1 & \text{if } \text{RSCP} \geq -115 \text{ dBm} \\ 0 & \text{otherwise} \end{cases}. \quad (5.2)$$

In case  $m$  is covered by several cells, the one with the highest RSCP is referred to as the best server, and denoted as  $c_m^*$ .

### 5.3.3 SHO areas

To obtain a realistic outline of the areas where a mobile may potentially maintain connections to more than one cell, a static version of the active set, as defined in [97], is used. To this end, a SHO window,  $\gamma^{\text{sho}}$ , and a maximum active-set size,  $as^{\max}$ , are introduced. Both parameters are taken from the working configuration of the real network. It follows that the cells to which a mobile  $m \in M$  may maintain concurrent downlink connections are part of the set

$$\begin{aligned} SHO_m^\downarrow &= \left\{ c \mid L_{c^*m}^\downarrow p_{c^*} - L_{cm}^\downarrow p_c \leq \gamma^{\text{sho}} \right\}, \\ |SHO_m^\downarrow| &\leq as^{\max}, \end{aligned} \quad (5.3)$$

where  $c \in C$ ,  $L_{c^*m}^\downarrow$  is the downlink attenuation factor of the best-serving cell, and  $p_{c^*}$  is its CPICH power. Since the number of elements in  $SHO_m^\downarrow$  is at most  $as^{\max}$ , the weakest links are removed if there are several present. This method is well suited for configurations with no hysteresis, since dynamic effects are ignored in static models [97].

Additionally, in the uplink, the set of cells to which a mobile can potentially be in SHO is defined as

$$SHO_m^\uparrow = \left\{ c \mid L_{mc}^\uparrow p_m^\uparrow \geq 3.16227766 \cdot 10^{-12} \text{ mW} \right\}, \quad (5.4)$$

where  $L_{mc}^\uparrow$  is the uplink attenuation factor from mobile  $m$  to cell  $c$ , and  $p_m^\uparrow$  is the uplink transmit power of mobile  $m$ .

The static nature of the model intentionally neglects mobility and interference by narrowing  $\gamma^{\text{sho}}$  down to 2 dB [97].

### 5.3.4 Optimization objective

Using the elements defined in Section 5.3, we have constructed an objective function in cooperation with a team of radio engineers of the Radio Network Department at Telekom Slovenije, d.d. The objective function is constructed as a weighted sum, containing different costs that penalize the occurrence of specific SHO conditions in downlink and uplink, which may potentially cause the aforementioned malfunctioning, introduced in Section 5.1.

A cost-based objective function is the most natural and straight-forward way of defining the optimization objective. Besides it is easily extendable to include other future situations,

also defining the mutual importance of the different phenomena taken into account at the optimization phase.

Hence, the definition of the objective function for the SHO-balancing problem is the minimization of the sum of penalty scores given as

$$\min f_{\text{sho}} = \sum_{c \in C} \sum_{m \in M} p f_{\text{cov}} (1 - cov_{cm}) + p f_{\text{sho}}^{\uparrow} sho_{cm}^{\uparrow} (1 - sho_{cm}^{\downarrow}) + p f_{\text{sho}}^{\downarrow} sho_{cm}^{\downarrow} (1 - sho_{cm}^{\uparrow}), \quad (5.5)$$

where

$$sho_{cm}^{\downarrow} = \begin{cases} 1 & c \in SHO_m^{\downarrow}, \\ 0 & \text{otherwise} \end{cases}, \quad (5.6)$$

$$sho_{cm}^{\uparrow} = \begin{cases} 1 & c \in SHO_m^{\uparrow}, \\ 0 & \text{otherwise} \end{cases}, \quad (5.7)$$

and

- $p f_{\text{cov}}$  represents the penalty factor for uncovered areas,
- $p f_{\text{sho}}^{\uparrow}$  represents the penalty factor for uplink SHO areas where SHO is not possible in the downlink, and
- $p f_{\text{sho}}^{\downarrow}$  represents the penalty factor for downlink SHO areas where SHO is not possible in the uplink.

## 5.4 Optimization algorithms

The SHO-balancing problem has been tackled using three fundamentally different optimization algorithms, namely:

- differential evolution (DE, see Section 2.1.6), from the family of evolutionary algorithms;
- differential ant-stigmergy algorithm (DASA, see Section 2.1.7), from the family of swarm-intelligence algorithms; and
- simulated annealing (SA, see Section 2.1.8), from the group of classic metaheuristic algorithms, targeted at combinatorial optimization problems.

Each of these algorithms shall minimize the objective function value by adopting essentially disparate approaches, hence the diversity of applying algorithms belonging to different families to solve the same optimization problem. Therefore, the result analysis shall establish which of the presented approaches is better suited for solving the SHO-balancing problem.

The following sections describe how the SHO-balancing problem is represented (or mapped) by the internal structure of each of the selected algorithms and their controlling parameters.

### 5.4.1 DE mapping

The DE algorithm features a parallel direct search method, which utilizes a population of  $D$ -dimensional parameter vectors. The SHO-balancing problem is expressed in each component of a vector  $X$  of the population, which maps to the CPICH power of one cell under optimization, i.e.

$$X_{aG} = \{x_1, x_2, \dots, x_c, \dots, x_D\}, \quad (5.8)$$

where  $x_c \in P_c$  represents a candidate CPICH power setting of cell  $c$ , and  $G$  indicates the generation of an individual  $a$  in the population. Since there are  $|C|$  cells in the mobile network, it follows that  $D = |C|$ .

From the different variants of DE, the most popular one is used here, called *DE/rand/1/bin*. The nomenclature used to name this variant indicates the way the algorithm works:

- *DE* denotes the differential evolution algorithm,
- *rand* indicates that the individuals selected to compute the mutation values are randomly chosen,
- 1 specifies the number of pairs of selected solutions used to calculate the weighted difference vector, and
- *bin* means that a binomial recombination operator is used.

Four control parameters of the search process for DE were considered: the population size ( $NP$ ), the number of generations for the algorithm to run ( $G_{max}$ ), the crossover constant ( $CR$ ), and the mutation scaling factor ( $F$ ).

### 5.4.2 DASA mapping

The mapping between the balancing problem and DASA is similar to the one for DE, depicted in Equation (5.8):

$$X_a = \{x_1, x_2, \dots, x_i, \dots, x_D\} \quad (5.9)$$

In this case, each ant,  $a$ , creates its own solution vector,  $X_a$ , during the minimization process. At the end of every iteration, and after all the ants have created solutions, they are evaluated to establish if any of them is better than the best solution found so far.

There are six parameters that control the way DASA explores the search space: the number of ants ( $m$ ), the discrete base ( $b$ ), the pheromone dispersion factor ( $q$ ), the global scale-increasing factor ( $s_+$ ), the global scale-decreasing factor ( $s_-$ ), and the maximum parameter precision ( $e$ ).

### 5.4.3 SA mapping

From the SA perspective, the system under optimization is in a given *state* at each time step during the process. The objective function maps a system state to a value known as the *energy* of the system in that state. A *move* in the search space represents a change in the state of the system. After making a move, the system may exhibit lower or higher energy, depending on the results of the objective function.

Algorithm 5.1 shows the pseudo-code of a move in the search space of possible CPICH power settings, resulting in a new state of the system.

---

**Algorithm 5.1** A move in the search space of SA.

---

```

 $c' \leftarrow \text{pick\_random\_cell}(C)$ 
repeat
    if  $\text{rand}() < 0.5$  then
         $p_{c'}^{\text{new}} \leftarrow p_{c'} + 0.01$ 
    else
         $p_{c'}^{\text{new}} \leftarrow p_{c'} - 0.01$ 
    end if
until  $p_{c'}^{\text{new}} \in P_{c'}$ 
 $p_{c'} \leftarrow p_{c'}^{\text{new}}$ 

```

---

Table 5.1: Technical characteristics of the test network used.

Number of cells	25
Coverage threshold (RSCP)	-115 dBm
SHO window ( $\gamma^{\text{sho}}$ )	2 dB
User equipment ( $p_m^{\uparrow}$ )	21 dBm, power class 4
Pixel resolution	25 m <sup>2</sup>
Population density	398/km <sup>2</sup>

At the first step, a cell,  $c'$ , is randomly selected from the set of all cells in the network,  $C$ . In step 2, a change of +0.01 dB or -0.01 dB is applied with 50% probability to  $p_{c'}$ . The CPICH power of cell  $c'$  is expressed in dBm. The randomly generated CPICH power setting,  $p_{c'}^{\text{new}}$ , is checked for validity in step 3, i.e. it must be an element of the set  $P_{c'}$ . If  $p_{c'}^{\text{new}}$  is not a valid CPICH power, step 2 is executed again, generating another random CPICH power. Finally, in step 4, the CPICH power of cell  $c$  is replaced by  $p_{c'}^{\text{new}}$ .

It is important to note that, as long as  $|P_{c'}| > 1$ , the pseudo-code shown in Algorithm 5.1 shall never be trapped in an endless loop. On the other hand, if  $|P_{c'}| < 2$ , there are no candidate CPICH powers for cell  $c'$  and thus no possibility of optimization by means of CPICH power adjustment. Notice also that the acceptance of a move in the search space is left to SA and its stochastic components.

SA has two parameters to control the search process: the initial temperature ( $t_{\text{initial}}$ ) and the total number of iterations or evaluations ( $it$ ).

## 5.5 Simulations

The simulations are performed using a standard Monte-Carlo method, assuming the mobile users are uniformly distributed. The SHO conditions of different users depend on the relative received signal quality from different cells and the SHO window, which triggers the addition of a cell to the user's active set [65].

### 5.5.1 Test network

The test network used for the simulations is a subset of the real UMTS network deployed in Slovenia by Telekom Slovenije, d.d. It represents a network extending over a hilly terrain, combining both rural and middle-dense suburban areas, which contains 25 cells within an area of more than 150 km<sup>2</sup>. Table 5.1 shows some characteristics of the test network used, and Figure 5.2 shows the area under radio coverage,  $A_{\text{covered}}$ , within  $A_{\text{total}}$ .

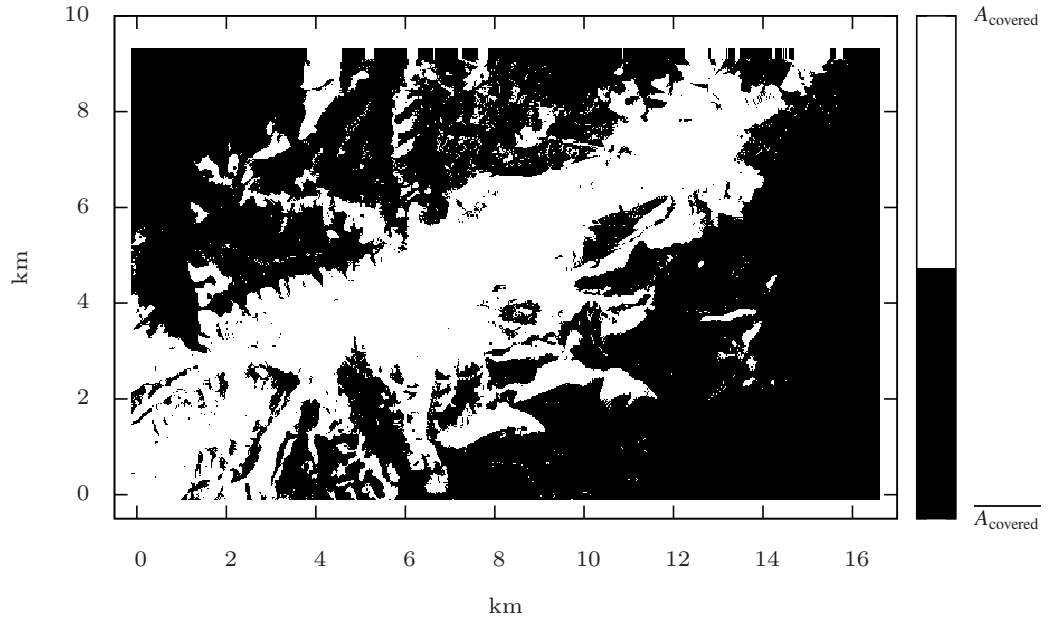


Figure 5.2: Area under radio coverage,  $A_{\text{covered}}$ , and without radio coverage,  $\overline{A}_{\text{covered}}$ , within the complete geographical area,  $A_{\text{total}}$ .

### 5.5.2 Penalty factors

After extensive experimentation, and working in cooperation with the radio engineers from the Radio Network Department at Telekom Slovenije, d.d., the penalty factors from Equation (5.5) are set to the following values:

- $pf_{\text{cov}} = 15$ ,
- $pf_{\text{sho}}^{\uparrow} = 13$ , and
- $pf_{\text{sho}}^{\downarrow} = 3$ .

It is clear that coverage is the most important quality aspect from the network point of view (penalty factor  $pf_{\text{cov}}$ ). Moreover, it imposes the biggest constraint to the optimization process, since the balance between SHO areas should not sacrifice network coverage. Another important characteristic that emerges from these values is the preference for minimizing areas where SHO capability is available in the uplink, but not in the downlink (penalty factor  $pf_{\text{sho}}^{\uparrow}$ ). As it has been described in Section 5.1, consequences of such SHO arrangement produce serious interference rise in neighboring cells (Figure 5.1), which may also result in service inaccessibility. The last factor  $pf_{\text{sho}}^{\downarrow}$  imposes a penalty value over areas where SHO capability is available in the downlink, but not in the uplink. Recall that when accessing HSPA services, SHO is available only in the uplink. For this reason, the link throughput may benefit from SHO in the uplink if it is available. The relative lower importance of the last penalty factor compared with the other ones is directly related to the consequences of such unbalancing of SHO areas may have on the network. In this case only HSPA throughput is

affected, while the service accessibility should not be an issue, given there is enough uplink coverage [67].

### 5.5.3 Algorithm parameters

In this section the algorithm parameter setup, as used during the simulations, is given. In all three cases we have followed the naming conventions as they have been previously introduced in Section 5.4.

#### 5.5.3.1 DE

The parameters controlling the behavior of the DE algorithm have been set as follows:

- $NP = 100$ , the population size;
- $G_{max} = 1000$ , the maximum number of generations for the algorithm to run;
- $CR = 0.8$ , the crossover constant; and
- $F = 0.5$ , the mutation scaling factor.

#### 5.5.3.2 DASA

As for DASA, we have set the parameters to the following values:

- $m = 10$ , the number of ants;
- $b = 10$ , the discrete base;
- $q = 0.2$ , the pheromone dispersion factor;
- $s_+ = 0.01$ , the global scale-increasing factor;
- $s_- = 0.01$ , the global scale-decreasing factor; and
- $e = 1.0^{-2}$ , the maximum parameter precision.

#### 5.5.3.3 SA

There are only two parameters controlling SA:

- $t_{initial} = 125$ , the initial temperature; and
- $it = 100,000$ , the total number of iterations.

SA also allows to define the way the temperature is lowered during the annealing process. In this case, the exponential-lowering schema has been used.

### 5.5.4 Experimental environment

All experiments were carried out on a 4-core Intel i7 2.67 GHz desktop computer with 6 GB of RAM running a 64-bit Linux operating system. The implementation languages used were C and Python, with the latter mostly used as ‘glue’ to hold the different implementation parts together, as well as for I/O operations. To lower the time needed to run one optimization round, we have implemented the entire objective-function evaluation using OpenCL and executed it on a nVidia GeForce GTX 260. This individual improvement exhibited more than 15x execution time speed-up when compared to the original CPU-only version.

Table 5.2: Solution-quality performance of the three algorithms, after 30 independent runs.

	Best	Worst	Mean	Std. deviation
DE	2,286,292.00	2,286,541.00	2,286,517.09	62.06
DASA	2,286,446.00	2,286,633.00	2,286,592.00	26.19
SA	2,293,350.00	2,295,570.00	2,294,626.50	663.75

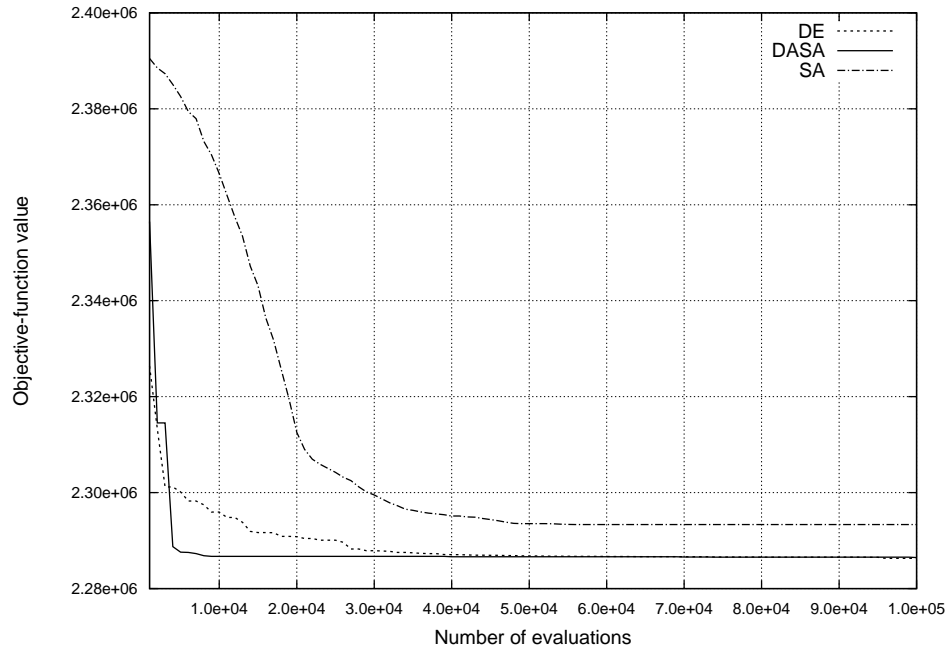


Figure 5.3: Convergence analysis for each algorithm, showing the best results obtained.

## 5.6 Results

### 5.6.1 Algorithm performance

In this section the performance of the selected algorithms is presented. The analysis includes aspects related to solution quality and convergence speed. All experimental results were obtained after 30 independent runs, each of them limited to a maximum of 100,000 evaluations. The gathered results are shown in Table 5.2.

It may be observed that DE reaches the lowest objective function value, closely followed by DASA. Likewise, both algorithms reach very similar results for the worst, mean and standard deviation values. SA, on the other hand, did not achieve comparable values, since its results are behind those of DE and DASA. Notice that even the best SA solution is no better than the worst solution of DASA. Moreover, the standard deviation exhibited by SA is many times bigger to those of DASA and DE, induced by the greater level of variance of its results.

The convergence of the best-recorded run of each of the three algorithms is shown in Figure 5.3. It is worth mentioning that every optimization run starts from a different solution,

Table 5.3: Improvement analysis for each of the achieved best solutions.

	Uncovered	Covered, no SHO	SHO	no SHO $\downarrow$ , SHO $\uparrow$	SHO $\downarrow$ , no SHO $\uparrow$	Total
Before opt.	63.00 %	15.11 %	15.73 %	1.80 %	4.36 %	100.00 %
DE sol.	60.23 %	16.13 %	16.09 %	1.47 %	6.08 %	100.00 %
DASA sol.	60.24 %	16.16 %	16.90 %	1.46 %	5.24 %	100.00 %
SA sol.	60.42 %	16.55 %	15.97 %	1.56 %	5.50 %	100.00 %
DE impr.	+4.40 %	+6.75 %	+2.29 %	+18.33 %	-39.45 %	—
DASA impr.	+4.38 %	+6.95 %	+7.44 %	+18.88 %	-20.18 %	—
SA impr.	+4.09 %	+9.53 %	+1.52 %	+13.33 %	-26.15 %	—
Avg. impr.	+4.29 %	+7.74 %	+3.75 %	+16.85 %	-28.59 %	—

randomly constructed by picking a CPICH power setting,  $p_c^k$ , from every  $P_c$ ,  $1 \leq k \leq |P_c|$ ,  $\forall c \in C$ . Notice how fast DASA converges to a good solution. After a number of evaluations without improvement, DASA resets itself and continues searching from a new random point within the search space [80]. Similarly, DE converges considerably fast, although not as fast as DASA does. In this case, DE does not reset itself if the current solution cannot be improved. Despite this, and based on the flat profile the graph exhibits towards the end of the optimization run, we are confident that 100,000 evaluations is an adequate stopping criterion for this algorithm. The third algorithm, SA, slowly converges towards the best solution found, even though it is not as good as the solutions found by DE and DASA.

The three convergence profiles shown in Figure 5.3 give a clearer notion about the way these algorithms explore the search space of the SHO-balancing problem.

The simulation running times have been intentionally omitted, since the implementations used are fundamentally different and therefore not comparable.

### 5.6.2 Interpretation

Table 5.3 presents the analysis of the obtained results from the network point of view. After 30 independent runs of each of the three algorithms, the best results obtained were evaluated for improvement and decline of each of the measured network aspects. The results are shown in Table 5.3, where '+' indicates improvement and '-' indicates decline of a given criteria. Overall, it may be observed that the measured criteria have been significantly improved. The only exception is the measure labeled as 'SHO $\downarrow$ , no SHO $\uparrow$ ', which shows an expected decline, since it is the optimization aspect with the lowest penalty factor value.

Coverage has been improved with an average of 4.29%, whereas the coverage area where there is no SHO capability has been increased 7.74% in average. Areas where SHO is available in both downlink and uplink have also been improved, 3.75% in average. This particular improvement is interesting from the optimization point of view, because it had no explicit penalty factor set. Therefore, it may be understood as a consequence of the criteria completeness the objective function has taken into account.

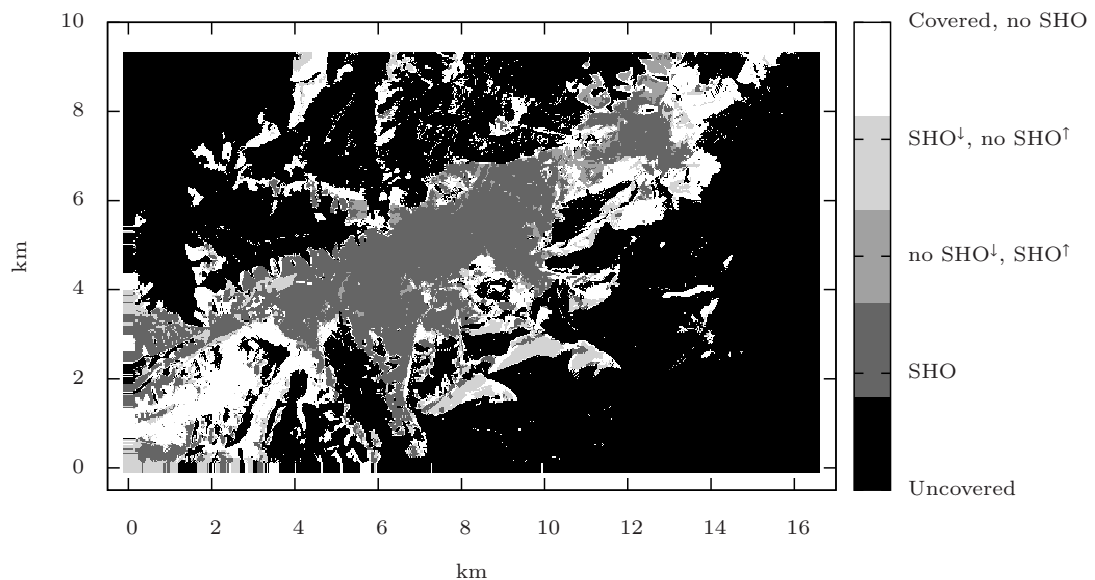


Figure 5.4: Spatial distribution of SHO areas before the optimization.

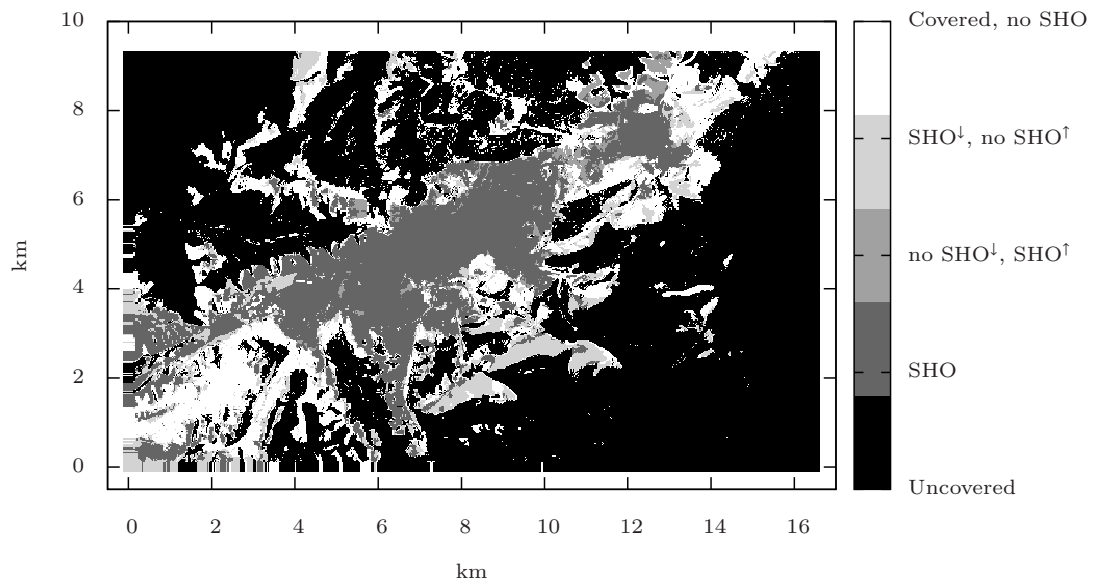


Figure 5.5: Spatial distribution of SHO areas after the optimization.

The second most important optimized aspect in the SHO-balancing problem is the proportion of areas with uplink SHO and no SHO in the downlink (labeled as ‘ $\text{no SHO}^{\downarrow}, \text{SHO}^{\uparrow}$ ’)

in Table 5.3). This particular condition has been improved by almost 17% in average, greatly reducing the possibility of interference in neighboring cells when serving HSPA traffic. The last measured aspect takes into account areas with downlink SHO and no SHO in the uplink (labeled as ‘SHO $\downarrow$ , no SHO $\uparrow$ ’ in Table 5.3). This condition, although it hasn’t improved, does not expose the mobile network to malfunctioning, only to reduced throughput within these specific areas. However, the reduced throughput is relative, since there are many cells capable of serving HSDPA data access, as the downlink SHO condition confirms. For this reason, the serving cell should not only deliver HSDPA, but also take care of the user signaling and power control, received in the uplink. Obviously, this is only feasible in areas where uplink coverage is guaranteed.

It is worth mentioning that the simulation results were obtained for a real mobile network with actual configuration data. Moreover, the hard constraints imposed to the optimization process (CPICH power limited within the  $\pm 2$  dB interval) ensure that the resulting configuration may be immediately applied to the mobile network. This fact can be contrasted with the spatial distribution of each of the optimized aspects, before and after applying the optimization results, as it is shown in Figures 5.4 and 5.5. The lack of any prominent visual change in Figures 5.4 and 5.5 is a desired consequence of the fine-tuning procedure the network has been exposed to. Still, the improvements are present precisely over the areas that are most exposed to malfunctioning due to unbalanced SHO, e.g. their borders.

## 5.7 Summary

This chapter formally introduced a new optimization problem for 3G networks: the SHO-balancing problem. A characterization of the consequences that unbalanced SHO areas have on the quality of HSPA services was also given. Particularly, tackling the SHO-balancing problem was possible due to the improved performance delivered by the evaluation framework PRATO (see Chapter 3).

Using a extension of the state-of-the-art radio-network model presented in ...???, the penalty scores of the objective function were set according to the configuration and layout of a real mobile network, deployed in Slovenia by Telekom Slovenije, d.d.

The balancing problem has been tackled by three optimization algorithms, namely DE, DASA and SA. All three algorithms were able to improve the given network configuration, being DE the most successful one. The presented results confirm that a great proportion of the SHO areas, that were not balanced before the optimization, were correctly balanced, therefore significantly reducing the possibility of HSPA-service failures. Additionally, radio coverage was improved, while all other essential network services were not altered.

One of the key advantages of the presented method is that it targets the optimization of a deployed network, for which the focus is to fine-tune the existing configuration instead of creating complete new solutions. Furthermore, a deployed network has a great number of hard-constraints that should be taken into account at the optimization stage. Yet, the presented approach is simple and versatile enough to be used in practically any working UMTS network. Moreover, the introduced model is applicable for mobile networks in heterogeneous environments, because it imposes no restrictions regarding cell layout or radio-propagation characteristics, which are completely adaptable through PRATO.

# Part III

## Radio-network planning



## 6 Framework automatic tuning

The assessment of the radio coverage is essential for network planning and optimization. Consequently, the planning tools used to gather this information play a key role in the decisions made during the planning phase of a radio network. But acquiring the necessary information to support the decision making in this context is a challenging task. Particularly, planning tools have to be adapted for a specific environment and technology in order to improve their accuracy. The complexity of this problem means that radio-coverage prediction is generally a computationally-intensive and time-consuming task, hence the importance of fast and accurate prediction tools.

Within this context, we present an open-source simulation framework for planning and optimization of radio networks. We provide two use cases for the newly deployed LTE network in Slovenia. The first one involves the parameter tuning of an empirical radio-propagation model using a snapshot of field measurements. The other one involves the optimization of clutter losses over different regions of the country, therefore adapting the losses due to land usage to the local conditions of each region.

We report the results of our experimental simulations over three regions of the real LTE network, deployed by Telekom Slovenije, d.d., thus showing the suitability of the parallel framework for tackling real-world planning and optimization problems.

### 6.1 Introduction

With the advent of long-term evolution (LTE) as the fourth generation (4G) in cellular technology, mobile operators are facing the challenges of deploying a new network. LTE follows the well established universal mobile telecommunication system/high-speed packet access (UMTS/HSPA) combo, targeting higher peak data rates, higher spectral efficiency and lower latency [?].

The deployment of a new mobile network is always a challenge for mobile operators, who constantly struggle to find the optimal investment in order to provide a competitive network in terms of coverage and quality of service. Indeed, coverage planning remains a key problem that all operators have to deal with. It has proven to be a fundamental issue since the deployment of the first GSM networks, more than 20 year ago.

One of the primary objectives of radio-coverage planning is to efficiently use the allocated frequency band for a geographic area to be satisfactorily reached with the radio stations of the network. To this end, radio-coverage prediction tools are of great importance as they allow network engineers to test different configurations before physically implementing the changes. However, to accurately predict the radio coverage of a mobile network is a very complex task, mainly due to the wide range of various combinations of hardware and configuration parameters which have to be analyzed in the context of different environments. The complexity of the problem means that radio-coverage prediction is generally a computationally-intensive and time-consuming task, hence the importance of fast and accurate prediction tools.

Although different mathematical models have been proposed for radio-propagation modeling, none of them excels in a network-wide scenario [111]. Empirical propagation models

usually give good results with a limited computational effort. However, for improved accuracy, the model parameters have to be adapted to better fit a specific network or region within it, mainly because of inaccuracies in input data and environmental changes in the network region, e.g. foliage of trees or snow. Consequently, a combination of different parameters is generally needed in order to reliably calculate radio-propagation predictions for particular environments. Moreover, since the number of deployed cells (transmitters) keeps growing with the adoption of modern standards [109], there is a clear need for a radio propagation tool that is able to cope with larger work loads in a feasible amount of time.

To address the afore-mentioned issues, we adapt the parameters of an empirical propagation model to a set of field measurements. The parameter tuning is analytically calculated per cell, in order to increase the accuracy of the calculated predictions. Moreover, we fine tune the signal losses due to land usage (clutter) in a regional basis, using an optimization approach. As a working framework to tackle the presented problems, we use a parallel radio-prediction tool [?], thus showing the suitability of the presented framework for real-world planning and optimization of LTE radio networks. Particularly, we show the tool capabilities to handle several parallel radio-prediction runs using a metaheuristic algorithm and distributed objective-function evaluation, while resolving the clutter optimization problem.

This paper is organized as follows. Section 6.2 gives a description of the parallel simulation framework, including some implementation details and performance figures. Section 6.3 introduces principles of radio-propagation prediction, and the mathematical model used. The parameter-tuning problem and the analytical approach for tackling it are presented in Section 6.4, including the simulations performed on the framework and their results. Section 6.5 concentrates on describing the optimization problem involving the regional adaptation of signal losses due to clutter, including the performed simulations the achieved results. Finally, Section 6.6 gives an overview of relevant publications, describing how they relate to our work, before drawing some conclusions.

## 6.2 Simulation framework

As if was mentioned before, we use the parallel radio-prediction tool PRATO [?] as our simulation and optimization framework. PRATO deals with complex calculations over large data sets by applying a work-pool parallel paradigm over a message-passing communication model. As such, PRATO is ready to be deployed over small groups of networked computers as well as over a computer cluster with hundreds of nodes. Since the prediction calculation employs digital elevation models and land-usage data in order to analyze the radio coverage of a geographic area, the framework is implemented as a module of the open source Geographic Resources Analysis Support System (GRASS) [99].

### 6.2.1 Parallel computation on computer clusters

Considering the high computational power needed for evaluating the radio coverage of real mobile networks during optimization, the use of a computer cluster is preferred. A computer cluster is a group of interconnected computers that work together as a single system. To reach high levels of parallel performance and scalability, PRATO performs the parallel decomposition big data sets, distributing the computational load among the computing nodes that belong to the cluster.

Computer clusters typically consist of several commodity PCs connected through a high-speed local network with a distributed file system, like NFS [112]. One such system is the DEGIMA cluster [60] at the Nagasaki Advanced Computing Center (NACC) of the Nagasaki University in Japan. This system ranked in the TOP 500 list of supercomputers until June

2012<sup>1</sup>, and in June 2011 held the third place of the Green 500 list<sup>2</sup> as one of the most energy-efficient supercomputers in the world.

### 6.2.2 Multi-paradigm parallel programming

The implementation methodology adopted for PRATO follows a multi-paradigm parallel programming approach in order to fully exploit the resources of each of the nodes in a computing cluster. To effectively use a shared memory multi-processor, PRATO uses POSIX threads to implement parallelism [20]. By using POSIX threads, multiple threads can exist within the same process while sharing its resources. For instance, an application using POSIX threads can execute multiple threads in parallel by using the cores of a multi-core processor, or use the system resources more effectively, thus avoiding process execution-halt due to I/O latency by using one thread for computing, while a second thread waits for an I/O operation to complete.

To use the computing resources of a distributed memory system, such as a cluster of processors, PRATO uses the Message Passing Interface (MPI) [58]. MPI is a message-passing standard, which defines syntax and semantics designed to function on a wide variety of parallel computers. MPI enables multiple processes running on different processors of a computer cluster to communicate with each other. It was designed for high performance on both massively parallel machines and on workstation clusters. Its development is supported by a broadly-based committee of vendors, developers, and users.

### 6.2.3 Master-worker model

PRATO follows a master-worker paradigm [? ], where the main process, i.e. the master, produces many sub-problems, which are delivered to be executed by the worker processes. These sub problems are, in this case, the radio-coverage prediction for individual cells of the radio network under optimization.

The master process is the only component that should be run from within the GRASS environment. It is responsible for dynamically starting the worker processes using the available computing nodes, based on the amount of network cells for which the coverage prediction should be calculated. For distributing the work among the worker processes, the master process dispatches the loaded geographic data, before dispatching them to the multiple worker processes. In this case, the decomposition of the data applies to the digital-elevation and the clutter data only, but it could be applied to any point-based data set, vector or raster. In the next step, the master process starts a message-driven processing loop, which main task is to evaluate the radio-coverage prediction of different transmitters among idle worker processes.

The worker processes, on the other hand, are completely independent from GRASS, i.e. they do not have to run within the GRASS environment nor use any of the GRASS libraries to work. This aspect significantly simplifies the deployment phase to run PRATO on a computer cluster, since no GRASS installation is needed on the computing nodes hosting the worker processes. During the result-saving phase, each of the worker processes sends its results independently from each other, following an asynchronous and decoupled design. Moreover, worker processes do this from an independent thread, which runs concurrently with the radio-prediction evaluation of the next transmitter received from the master process. The overlap between calculation and communication achieved by the use of an auxiliary thread

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<sup>1</sup><http://www.top500.org>

<sup>2</sup><http://www.green500.org>

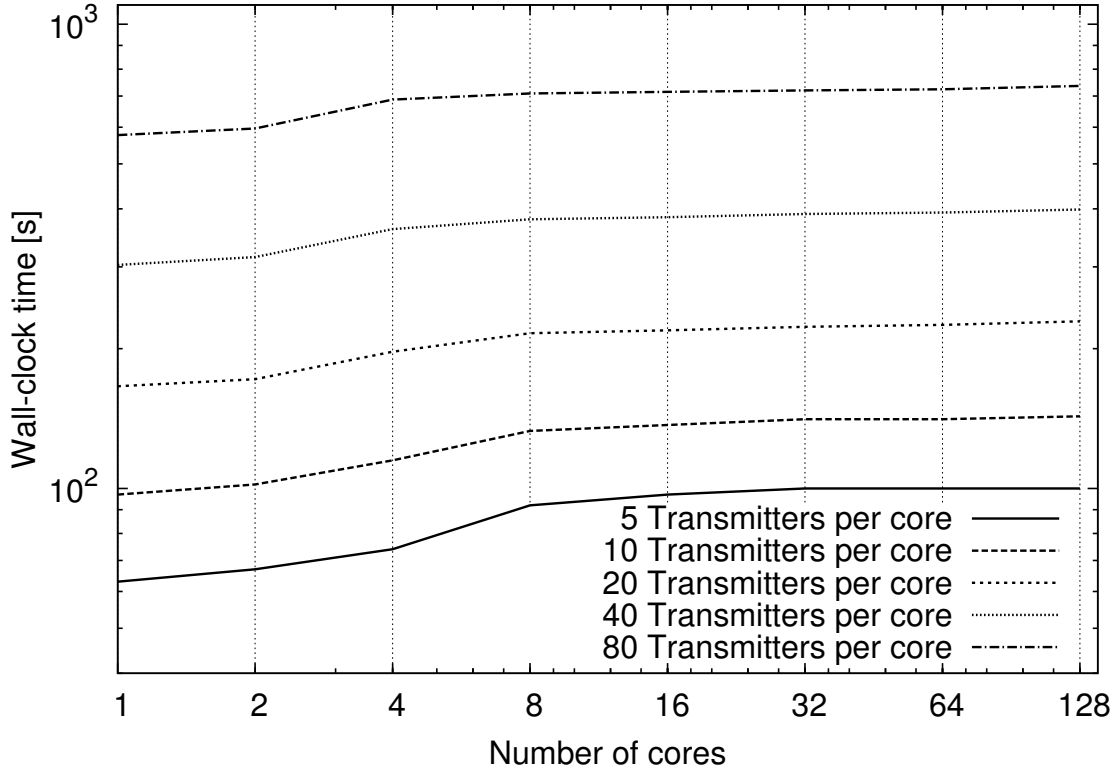


Figure 6.1: Measured wall-clock time for weak-scalability experiments. Experiments performed assigned one MPI worker process per available core. The wall-clock time axis is expressed in base-10 logarithmic scale, whereas the axis representing the number of cores is expressed in base-2 logarithmic scale.

completely hides the latency created by the result-dumping task, and makes better use of the system resources.

#### 6.2.4 Performance

From the performance point of view, PRATO is capable of reaching levels of high efficiency, as the following measurements show.

Figure 6.1 shows the time measurements of a weak-scalability experiment. To measure the weak-scalability properties of PRATO means analyzing the scalability of the parallel implementation in cases where the workload assigned to each MPI process (one process per processor core) remains constant as the number of processor cores, and thus the total size of the problem, is increased.

The time measurements observed from the weak-scalability results show that the wall-clock times are almost constant for bigger problem instances, revealing that the achieved level of scalability gets close-to-linear as the amount of transmitters-per-core increases. Specifically, PRATO scales especially well when challenged with a big number of cells or transmitters (10,240 for the biggest instance) over 128 cores.

Another aspect is the strong-scalability capabilities of PRATO, which represents the performance when increasing the number of computing cores for a given problem size, i.e. the number of cells deployed over the target area does not change, while only the number of cores used is increased. Figure 6.2 shows the speedup factors achieved for a set of strong-scaling experiments. We may see that a close-to-linear scaling is achieved, especially for

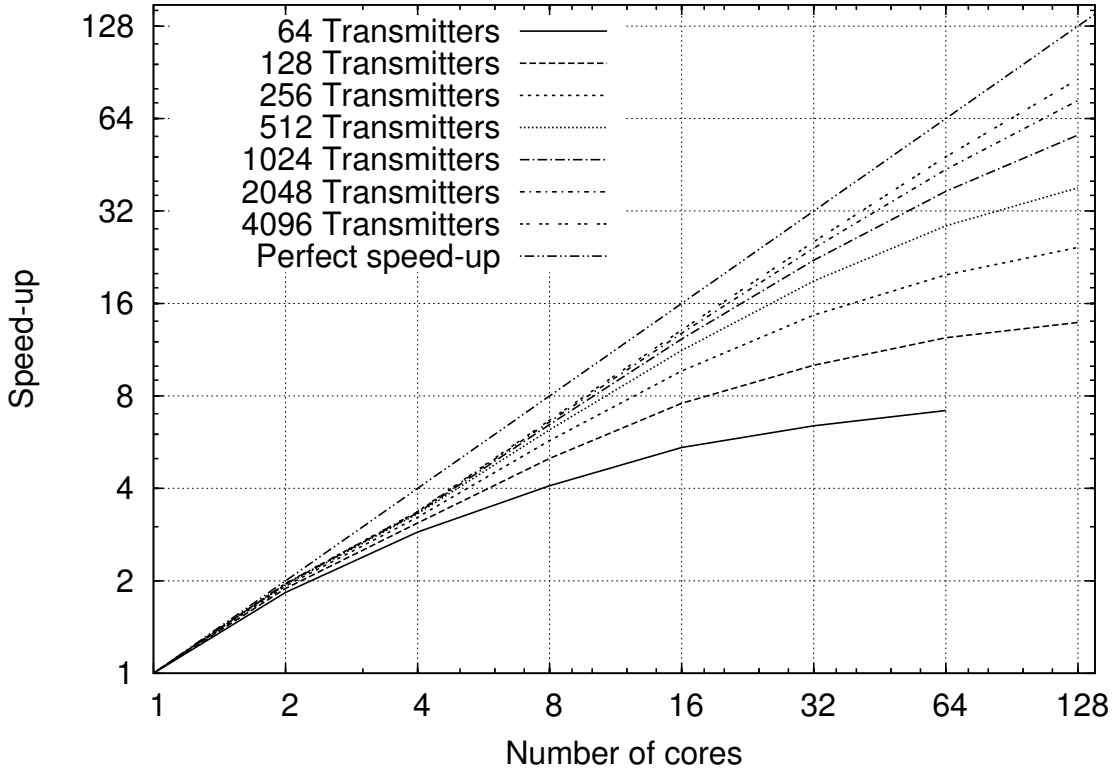


Figure 6.2: Measured speedup for strong-scalability experiments. The speedup axis is expressed in base-2 logarithmic scale, whereas the axis representing the number of cores is expressed in base-2 logarithmic scale.

the biggest problem sizes. Linear scaling occurs when the obtained speedup is equal to the total number of processors used. However, it should be noted that perfect speedup is almost never achieved, due to the existence of serial stages within an algorithm and communication overheads of the parallel implementation.

For more detailed information about PRATO, its design and capabilities, we refer the reader to [? ].

### 6.3 Radio-propagation prediction

The objective here is to improve the quality performance of a given mathematical model, used for radio-propagation calculations, by fine-tuning its configurable parameters as per-cell basis. In order to do this, we apply a state-of-the-art analytical approach, based on least squares method [? ], thus combining field measurements to fit the parameters of the mathematical model.

The idea is to automatically adapt the parameters of the mathematical model for each of the cells targeted by the optimization. That is, starting from an a-priori best-known set of parameters, empirically calculated by the radio engineers at Telekom Slovenije, d.d., this approach adapts the model parameters so that the deviation of the radio-propagation prediction to a given set of field measurements is minimized.

To calculate the radio-coverage predictions we use a mathematical model based on the well-known empirical Okumura-Hata formula [? ]. Other more accurate methods exist, like the ones based on ray tracing [? ? ]. However, these methods are more sensible to deviations

in input data, like digital elevation models and buildings, and are still inefficient in terms of the computational effort required to achieve satisfying results.

Empirical methods for radio-propagation predictions, on the other hand, give acceptable results within a feasible amount of time. For this reason, they have become the industry standard for non-deterministic propagation-loss calculations [? ? 29? ?, 68].

### 6.3.1 Radio-prediction model

The Okumura-Hata model has been largely studied and shown to be suitable for predicting radio propagation of LTE networks [? ]. Moreover, this mathematical model is especially appropriate for tuning, as it contains a number of adjustable parameters, which adapt the model according to a given scenario and its local conditions. In its primary form, the model distinguishes the distance of receiver from the transmitter, the frequency used and the effective antenna height, i.e. the antenna height above the receiver's level, in order to calculate the path loss in line-of-sight (LOS) conditions. However, for distinguishing non-line-of-sight (NLOS) conditions, the terrain profile and Earth shape are added to the original formula. Moreover, empirical corrections due to different land usage are also included to improve the quality of the calculated predictions. Equation (6.1) describes the path loss when there is LOS between the transmitter and the receiver.

$$\begin{aligned} pl_{\text{LOS}}(d_{(x,y)}, \beta) = & a_0 + a_1 \log(d_{(x,y)}) + a_2 \log(H_A) + \\ & a_3 \log(d_{(x,y)}) \log(H_A) - 3.2 [\log(11.75 \cdot H_R)]^2 + \\ & 44.49 \log(F) - 4.78 [\log(F)]^2, \quad (6.1) \end{aligned}$$

where  $\beta = (a_0, a_1, a_2, a_3)$  is the vector containing the tuning parameters of the model,  $d_{(x,y)}$  is the distance (in kilometers) from the transmitter to the topography point with coordinates  $(x, y)$ ,  $H_A$  is the effective antenna height (in meters) of the transmitter,  $H_R$  is the antenna height (in meters) of the receiver, and  $F$  is the frequency, expressed in MHz. On the other hand, in NLOS conditions, the path loss is calculated as in Equation (6.2).

$$pl_{\text{NLOS}}(d_{(x,y)}) = \sqrt{[\alpha K(d_{(x,y)})]^2 + E(d_{(x,y)})^2}, \quad (6.2)$$

where  $\alpha$  is the knife-edge diffraction control parameter,  $K(d_{(x,y)})$  is the knife-edge diffraction loss (in dB) and  $E(d_{(x,y)})$  is the correction due to the Earth sphere (in dB). The latter two values are calculated on the topography point with coordinates  $(x, y)$ .

In this work, the terrain profile is used for LOS determination. In order to adequately predict propagation effects due to foliage, buildings and other fabricated structures, additional loss factors based on the land usage (clutter data), are included. This technique is adopted by several propagation models for radio networks [? ? 98]. Consequently, we introduce an extra term for signal loss due to clutter, thus defining the model-predicted path loss as

$$pl(d_{(x,y)}, \beta) = pl_{\text{LOS}}(d_{(x,y)}, \beta) + pl_{\text{NLOS}}(d_{(x,y)}) + pl_{\text{CLUT}}(d_{(x,y)}), \quad (6.3)$$

where  $pl_{\text{CLUT}}(d_{(x,y)})$  is clutter loss at the topography point with coordinates  $(x, y)$ , expressed in dB.

Table 6.1: Clutter-category label numbers and their land-usage meanings for the radio-prediction model.

Clutter category	Description
0	Urban area without buildings, mostly roads
1	Suburban area
2	Urban area
3	Dense urban area
4	Agricultural area
5	Forestall area
6	Swamp area
7	Dry open land area with special vegetation
8	Dry open land area without special vegetation
9	Water area
10	Industrial area
11	Park area

In our case, we recognize twelve different clutter categories representing loss values due to different land usage. These categories, including their label numbers and descriptions, are depicted in Table 6.1.

In any case, the effectiveness of decision-making during radio-network planning is tightly coupled with the precision achieved by the propagation model used. In order to obtain a radio-propagation model that most accurately reflects the propagation characteristics of the area covered by each radio cell in the network, the parameters of the mathematical model are tuned using field-measurement campaigns. Parameter tuning using this method depends on existing field-measurement data, which are taken before hand over the area covered by the target network cells.

## 6.4 Parameter tuning of the radio-prediction model

The objective here is thus improve the quality performance of a given mathematical model, used for radio-propagation calculations, by fine-tuning its configurable parameters as per-cell basis. In order to do this, we apply a state-of-the-art analytical approach, based on least squares method [?], thus combining field measurements to fit the parameters of the mathematical model.

The idea is to automatically adapt the parameters of the mathematical model for each of the cells targeted by the optimization. That is, starting from an a-priori best-known set of parameters, empirically calculated by the radio engineers at Telekom Slovenije, d.d., this approach adapts the model parameters so that the deviation of the radio-propagation prediction to a given set of field measurements is minimized.

In the context of our work, this translates to fine tuning the parameters of the LOS part of the path-loss prediction model, i.e.  $pl_{\text{LOS}}(d_{(x,y)}, \beta)$ . The adjustable parameters are the elements of vector  $\beta = (a_0, a_1, a_2, a_3)$ , namely

- $a_0$  the reference loss or offset;
- $a_1$  the loss slope due to distance of the receiver from the transmitter;
- $a_2$  the loss slope due to height of the transmitter antenna;
- $a_3$  the loss slope due to the combined effect of the distance and height of the antenna.

The parameter tuning is performed per cell to improve local fitting of the radio predictions. The steps to be completed are: collect and process field-measurement data, and solve the particular system of linear equations. The resulting solution is the vector  $\beta$  of the target cell, with its values locally adapted.

In the following, we denote the default parameters of the radio-propagation model with these values

$$a_0 \ 38.0,$$

$$a_1 \ 32.0,$$

$$a_2 \ -12.0, \text{ and}$$

$$a_3 \ 0.1.$$

#### 6.4.1 Field measurements

In mobile networks, a moving mobile constantly performs cell selection/reselection and handover in order to keep the best possible connection to the network. Within this context, the best connection is selected by measuring the signal strength or quality of the neighboring cells. In LTE networks, a mobile measures two parameters on reference signal, namely the Reference Signal Received Power (RSRP) and the Reference Signal Received Quality (RSRQ).

For a certain frequency bandwidth, RSRP measures the average received power over the resource elements that carry cell-specific reference signals. RSRP is applicable in idle (e.g. waiting for a call) and connected (e.g. during a call) modes. During the procedure of cell selection/reselection in idle mode, RSRP is used. On the other hand, RSRQ is only applicable when the mobile is in connected mode.

The radio-coverage calculation involves predicting the network coverage over a certain region, and thus to the users' mobiles within it. Hence, in the first place, we are interested on accurately predicting the best connection the mobile would select in idle mode and the RSRP field measurements it uses.

In our case, the field measurements, representing the RSRP at a given location, are collected using a small truck equipped with the spectrum analyzer Rohde & Schwarz. The spectrum analyzer is connected to an external omni antenna mounted on the roof of the truck, at roughly 2 meters above the ground, taking measurements at a rate of 50 Hz???, with the symbol rate set to ??? Mhz. To accurately establish the measurement location points, a GPS unit was used. The measurement locations covered most of the streets within the target area, with over 300,000 individual field-measurement points taken at more than 150 network cells.

To minimize the error impact in measured RSRP and estimated location, all field measurements are processed so that a single value, the median<sup>1</sup>, is calculated for each of the measured locations. The resulting RSRP is then used to estimate the path-loss prediction at the location corresponding to each of the measurements, which resolution matches those of the digital elevation model and clutter data.

#### 6.4.2 Linear least squares

It is important to note the linear relationship between the predicted loss  $pl(d_{(x,y)}, \beta)$  and the components of vector  $\beta$ , for this is a necessary condition for the least-squares method to find a solution. Following this method, the parameter fitting of the propagation model is based

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<sup>1</sup>The median is the numerical value separating the higher half of a sample set from the lower half.

Table 6.2: Percentage of clutter-category proportions for each of the test networks used. The category legend is given in Table 6.1.

	Cat. 0	Cat. 1	Cat. 2	Cat. 3	Cat. 4	Cat. 5	Cat. 6	Cat. 7	Cat. 8	Cat. 9	Cat. 10	Cat. 11
Net <sub>1</sub>	0.53	4.53	1.68	0.45	71.89	17.94	0.07	0.00	0.03	2.21	0.67	0.00
Net <sub>2</sub>	0.91	5.53	9.48	3.84	29.73	48.57	0.14	0.03	0.03	0.76	0.86	0.12
Net <sub>3</sub>	0.15	3.99	1.14	0.11	26.50	67.13	0.26	0.00	0.00	0.36	0.36	0.00

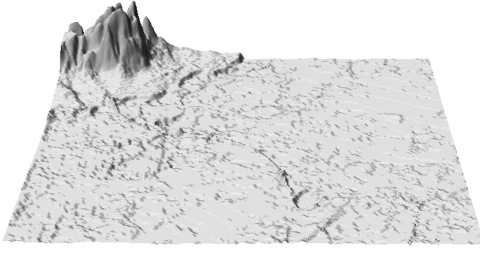


Figure 6.3: Terrain profile for network Net<sub>1</sub>, dominated by an agricultural area.

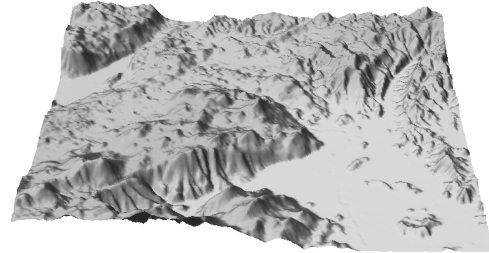


Figure 6.4: Terrain profile for network Net<sub>3</sub>, dominated by hills.

on the minimization of an observed error, i.e. the sum of the squared difference between the predicted and measured RSRP,

$$E(c) = \sum_{i=1}^{m_c} (p_c - pl_c(d_{(x,y)}, \beta_c) - fm_i)^2, \quad (6.4)$$

where  $E(c)$  is the observed error for cell  $c$ ,  $p_c$  is the transmit power of cell  $c$ , and  $fm_i$  is the  $i$ -th field measurement out of a set of measurements of cell  $c$  with cardinality  $m_c$ .

#### 6.4.3 Simulations

The simulations carried out for this part of our work comprise building the matrices of observed-error values,  $E(c)$ , for each cell  $c$  in the target network. The linear systems of equations are then individually solved by applying the linear least squares method [? ], which involves the evaluation of one radio-coverage prediction per cell. Each system holds a unique solution for each cell  $c$ , denoted by the vector  $\beta_c$ .

##### 6.4.3.1 Test networks

All the test networks, Net<sub>1</sub>, Net<sub>2</sub>, and Net<sub>3</sub> are subsets of a real LTE network deployed in Slovenia by Telekom Slovenije, d.d. The path-loss predictions are calculated using a digital elevation model and clutter map of 25 m<sup>2</sup> resolution and a receiver height of 1.5 m above ground level. A radius of 16 km is taken around each network cell, thus limiting the path-loss prediction to a distance where it is still feasible for the mobile to connect to a cell, with a RSRP greater or equal to -124 dBm [? ]. At the same time, the selected calculation radius provides enough overlap among neighboring cells to also calculate the network coverage over the whole region. Table 6.3 provides more information about the test networks used.

Net<sub>1</sub> represents a network deployed over a dominant agricultural area with almost flat terrain, some forests and waters streams. Net<sub>2</sub>, on the other hand, is deployed over a densely

Table 6.3: Several properties of the test networks used for the experimental simulations.

	Number of cells	Area (km <sup>2</sup> )	Field-measurement proportion (%)
Net <sub>1</sub>	12	103.74	5.41
Net <sub>2</sub>	130	1298.02	12.02
Net <sub>3</sub>	6	386.38	2.30

populated urban area, containing high buildings, parks and avenues. The last network, Net<sub>3</sub>, represents a network deployed over hilly terrain, including some smaller villages and vast forests. It is important to note that the number of deployed cells is directly related to the population density within the region of each test network, as the number of cells from Table 6.3 show. For a clearer representation of the test networks used, we present the proportion of each land-use (clutter) category in Table 6.2.

The terrain profiles are most relevant for Net<sub>1</sub> and Net<sub>3</sub>, since none of them comprehends an urban environment. Note that the terrain shown in Figure 6.3 is mostly flat, since the agricultural area is predominant there. In contrast, the terrain for Net<sub>3</sub> is dominated by hills, which are mostly covered by dense forests, with some small villages in the valleys (see Figure 6.4).

#### 6.4.3.2 Experimental environment

The simulations were carried out on 40 computing nodes of the DEGIMA cluster [60] at the Nagasaki Advanced Computing Center (NACC) of the Nagasaki University in Japan. This system ranked in the TOP 500 list of supercomputers until June 2012<sup>1</sup>, and in June 2011 held the third place of the Green 500 list<sup>2</sup> as one of the most energy-efficient supercomputers in the world.

The computing nodes are connected by a LAN, over a Gigabit Ethernet interconnect. The reason for using a high-end computer cluster as DEGIMA is to exploit the parallel nature of PRATO. However, this does not mean that the presented simulation cannot be carried out serially in a commodity desktop computer. PRATO does support serial calculation of radio-propagation predictions for several cells.

Each computing node of DEGIMA features one of two possible configurations, namely:

- Intel Core i5-2500T quad-core processor CPU, clocked at 2.30 GHz, with 16 GB of RAM; and
- Intel Core i7-2600K quad-core processor CPU, clocked at 3.40 GHz, also with 16 GB of RAM.

During the simulation runs, the nodes equipped with the Intel i5 CPU host the worker processes, whereas the master process runs on a computing node featuring an Intel i7 CPU, and performing all its I/O operations on the local file system.

All the nodes are equipped with a Linux 64-bit operating system (Fedora distribution). As the message passing implementation we use OpenMPI, version 1.6.1, which has been manually compiled with the distribution-supplied gcc compiler, version 4.4.4.

<sup>1</sup><http://www.top500.org>

<sup>2</sup><http://www.green500.org>

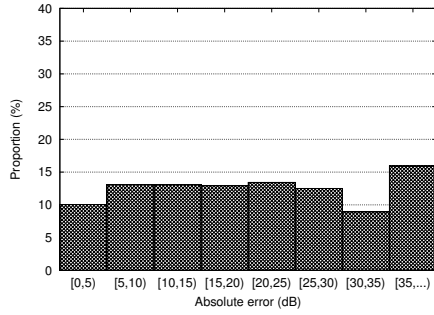


Figure 6.5: Error distribution of the radio prediction for network Net<sub>1</sub> with default parameter values.

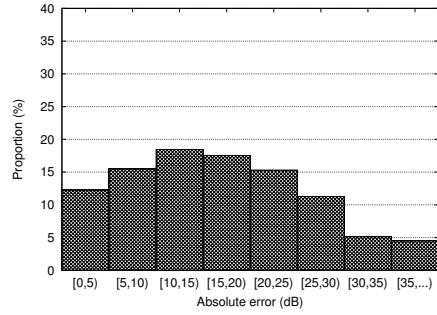


Figure 6.6: Error distribution of the radio prediction for network Net<sub>1</sub> with fitted parameter values.

#### 6.4.3.3 Results

The results of applying the linear least squares method to fit the parameters of the radio-prediction model to a set of field measurements presented in this section. We have prepared bar charts showing the cumulative distribution of the absolute error between the prediction points and the field measurements. Each bar represents an open interval, expressed in dB, denoting the proportion of points that deviate from the prediction for the expressed number of dB. For example, in Figure 6.5, we may see that the proportion of predicted points differing from the field measurements in 35 dB or more is around 16%, whereas the proportion differing in less than 5 dB is 10%. These numbers represent the test network Net<sub>1</sub> before applying the model-parameter fitting. For comparison, in Figure 6.6, the absolute-error distribution for the same test network is given, but with the model parameters fitted to the available field measurements. Notice how the proportions describing the biggest deviation have dropped to under 5% (35 dB and more) and less than 6% (30 dB to 35 dB). Moreover, it is clear how all proportions improved, raising the bars towards the left-hand side.

The error distributions of the radio-propagation prediction for test network Net<sub>2</sub> using default parameters and fitted ones are given in Figures 6.7 and 6.8, respectively. In this case, the improvement is even more significant than for the first test network, clearly showing that the tuned propagation model represents the local radio propagation conditions within this environment more accurately.

For the third test network, Net<sub>3</sub>, the error distributions are depicted in Figure 6.9 using the default parameters, and Figure 6.10 for the tinned ones. Similarly as for the first test network, Net<sub>1</sub>, we may see a how the proportion of highest error deviation has been lowered over those with lower deviation values.

The overall results confirm that fitting the parameters of the radio-propagation model to the field measurements within a certain region significantly improve the quality of the calculated propagation prediction.

Despite the presented results are encouraging, there are some specific reasons behind the considerable better results for Net<sub>2</sub>, when compared to those of Net<sub>1</sub> and Net<sub>2</sub>. Clearly, the number of available field measurements directly affects the quality of the calculated results, making the least squares approximation rougher and thus less precise (see Table 6.3 in Section 6.4.3.1). However, it is out of the scope of this work, in which we focus is on the applicability of PRATO as a tool for planning and optimization of LTE networks, to assess the quality of the achieved results. Nevertheless, for the sake of completeness, we would like to point out that some researchers have already started working on different ways on how to improve this aspect [? ? ].

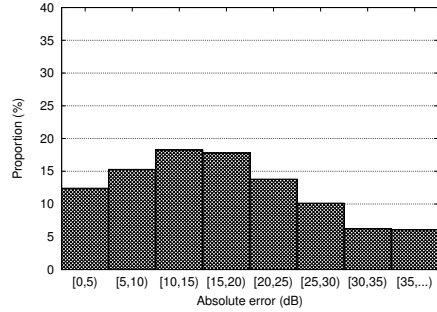


Figure 6.7: Error distribution of the radio prediction for network  $\text{Net}_2$  with default parameter values.

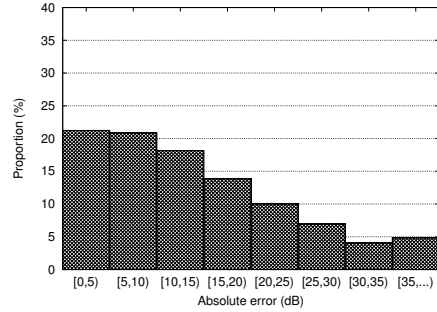


Figure 6.8: Error distribution of the radio prediction for network  $\text{Net}_2$  with fitted parameter values.

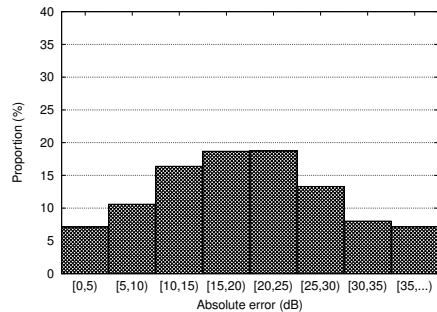


Figure 6.9: Error distribution of the radio prediction for network  $\text{Net}_3$  with default parameter values.

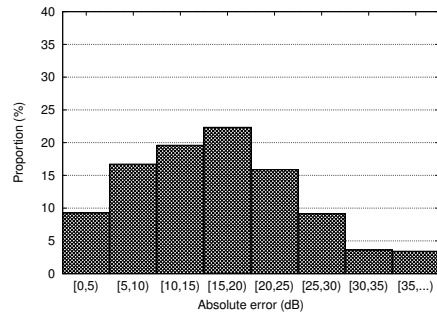


Figure 6.10: Error distribution of the radio prediction for network  $\text{Net}_3$  with fitted parameter values.

## 6.5 Clutter optimization

In order to better adapt the radio-prediction calculation to a given regional environment, the signal losses due to clutter have to be optimized. As it was mentioned before, there are several reasons for the signal loss values to be inaccurate. Among them, we can mention seasonal changes, like tree foliage or snow, construction or demolition of buildings and parks, different kinds of forests or agricultural areas. These changes are only noticeable through regular field-measurement campaigns and land-usage data updates. The challenge is thus to combine both in order to improve the prediction results even further.

In the following, we use a metaheuristic algorithm in order to optimize the clutter losses, i.e. the signal loss due to terrain influence, of several regions within the mobile network under optimization. This is done over a group of network cells within different regions of the target network, e.g. agricultural, urban or hilly. In terms of radio-network planning, a regional classification of signal loss due to clutter further improve the accuracy of the radio-coverage prediction, thus enhancing the model by distinguishing, for example, different types of agricultural, park and vegetation areas.

Compared to the parameter tuning of the mathematical model presented in Section 6.3.1, we are not using an analytical approach for tackling this problem, since we wish to potentially include corrections for possible errors in the NLOS part Equation 6.3. For this reason, we turn our attention to metaheuristic algorithms [134] in general and swarm intelligence in particular [? ]. From this last family of metaheuristic algorithms, we choose the differential ant-stigmergy algorithm (DASA) [? ].

Several authors have shown the benefits of using metaheuristic algorithms for tackling radio-related problems [16? ? ? ]. In our case, an analytical approach may also be possible, but our objective is to show the capabilities of PRATO when combining an optimization algorithm with parallel and distributed objective-function evaluation. For the following simulations, the DASA runs as the master process, whereas the workers evaluate the current solution in parallel for all the involved cells (see Figure 6.11).

### 6.5.1 Optimization objective

The optimization objective consists of adjusting the loss values of the different clutter categories, i.e.  $pl_{CLUT}(d_{(x,y)})$  of Equation (6.3), to best fit a set of field measurements in a given region containing multiple cells. We have three data sets at our disposal, one for each of the three regions: the first for Net<sub>1</sub>, the second for Net<sub>2</sub>, and the third for Net<sub>3</sub>. Each region is independently optimized, so that the radio-propagation predictions of its network cells, which already have their model parameters fitted, minimize the mean-squared error against the field measurements, namely

$$F^*(R) = \sum_{i=1}^{m_c} \frac{(p_c - pl_c(d_{(x,y)}, \beta_c) - fm_i)^2}{m_R} \quad \forall c \in R, \quad (6.5)$$

where  $F^*(R)$  is the optimization objective to be minimized,  $R$  is one of the three regions,  $p_c$  is the transmit power of cell  $c$ ,  $fm_i$  is the  $i$ -th field measurement out of a set of measurements of cell  $c$  with cardinality  $m_c$ ,  $m_R$  is the number of field measurements of all the cells within the region  $R$ , and  $pl_c(d_{(x,y)}, \beta)$  is the path loss of cell  $c$  at coordinate  $(x, y)$ , where  $\beta$  contains per-cell fitted parameter values of the prediction model.

For a reference of the different clutter categories our prediction model recognizes, see Table 6.1.

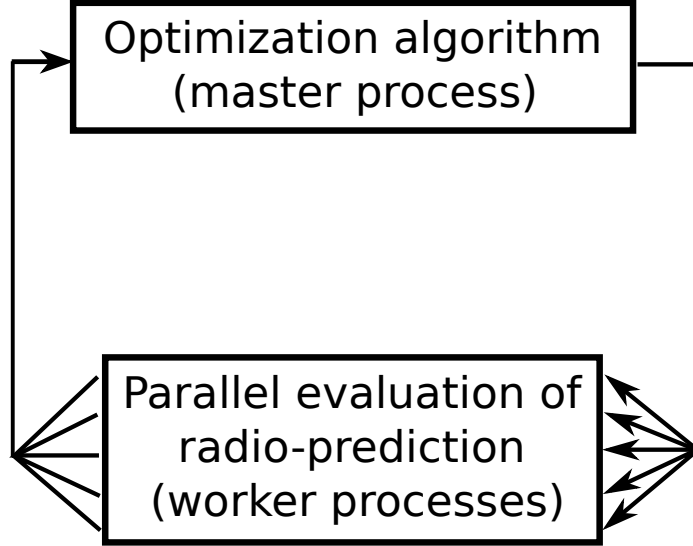


Figure 6.11: *PRATO architecture and data flow during the clutter-optimization phase.*

### 6.5.2 Differential ant-stigmergy algorithm

As it has been mentioned before, the optimization algorithm we have chosen for the clutter-optimization problem is the (DASA). Based on the metaheuristic Ant-Colony Optimization (ACO) [?], the DASA [?] provides a framework to successfully cope with high-dimensional numerical optimization problems. It creates a fine-grained discrete form of the search space, representing it as a graph. This graph is then used as the walking paths for the ants, which iteratively improve the temporary best solution.

The mapping between the clutter-optimization problem and DASA is as follows:

$$X_a = \{x_1, x_2, \dots, x_i, \dots, x_{12}\} \quad (6.6)$$

where  $X_a$  is the solution vector of ant  $a$  during the minimization process, and  $x_i$  represents the  $i$ -th clutter category within a given region. At the end of every iteration, and after all the ants have created solutions, they are evaluated to establish if any of them is better than the best solution found so far.

There are six parameters that control the way DASA explores the search space: the number of ants, the discrete base, the pheromone dispersion factor, the global scale-increasing factor, the global scale-decreasing factor, and the maximum parameter precision.

For a more in-depth explanation about these parameters and the DASA algorithm itself, we refer the reader to [?].

### 6.5.3 Simulations

In this case, the simulations for each test network comprise multiple iterations of several steps. An iteration begins by generating a solution vector for each of the ants in the DASA population. The following step involves the parallel evaluation of each solution, i.e. one radio-coverage prediction per cell. The final step is calculating the objective-function value, as defined in Equation (6.5), before sending it to the DASA for it to generate the next set of solutions. Figure 6.11 depicts the way PRATO performs the objective-function evaluation in parallel over the worker processes, while the optimization algorithm runs on the master one.

Table 6.4: Clutter-category losses after the optimization. The default losses for each clutter category are given along the solutions for each of the test networks. All values are expressed in dB.

Clutter category	Default	Net <sub>1</sub>	Net <sub>2</sub>	Net <sub>3</sub>
0	5.0	13.71	11.30	17.90
1	15.0	12.39	16.67	-
2	13.0	16.04	17.04	15.69
3	28.0	19.59	18.01	23.00
4	12.0	11.48	9.71	10.80
5	20.0	16.26	11.62	16.26
6	15.0	-	-	-
7	8.0	-	13.49	-
8	5.0	-	13.50	-
9	1.0	17.50	5.60	-
10	20.0	8.26	16.75	16.63
11	8.0	-	18.93	-

Compared to the experiments presented in Section 6.4.3, a much higher number of evaluations is needed for this kind of optimization. In this context, it is of key importance to perform the radio-coverage prediction for multiple cells in a parallel manner. Otherwise, such an approach would not be feasible, since the time required to reach a reasonable solution would be excessive.

The loss value (in dB) each clutter category may take has been limited to the interval [0,40]. This information has been provided by the radio experts of the Radio Network department at Telekom Slovenije, d.d.

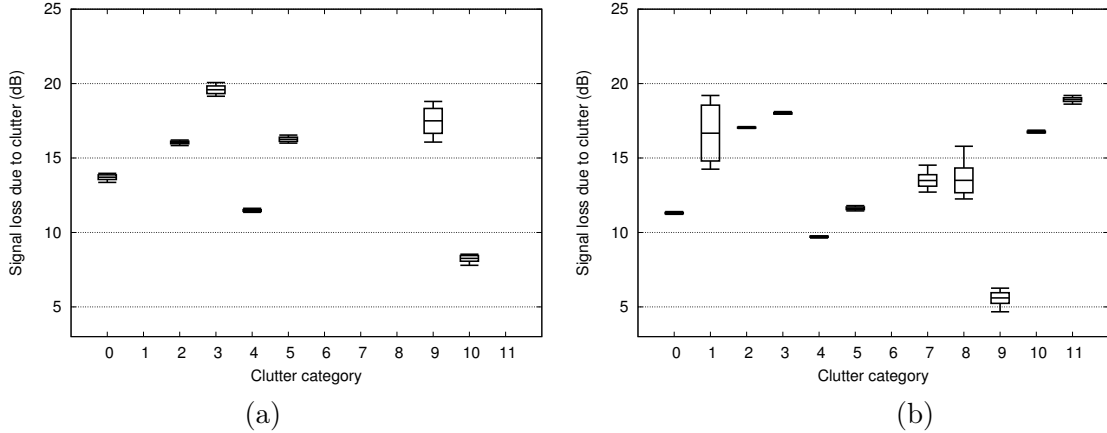
As stopping criteria of the optimization runs for Net<sub>1</sub> and Net<sub>3</sub>, we have fixed the maximum number of iterations to 200, whereas for Net<sub>2</sub> this values has been set to 500 iterations, since this networks comprises the largest number of network cells. Overall, the framework completed 48,000 objective-function evaluations in the former case, and 120,000 in the latter one.

Regarding the parameters that control algorithm behavior, we have set them to the following values

- $m = 240$ , the number of ants;
- $b = 10$ , the discrete base;
- $q = 0.2$ , the pheromone dispersion factor;
- $s_+ = 0.01$ , the global scale-increasing factor;
- $s_- = 0.01$ , the global scale-decreasing factor; and
- $e = 1.0^{-2}$ , the maximum parameter precision.

#### 6.5.4 Results

The results calculated by the optimization process are shown in Table 6.4. The solutions are given for each of the test networks, along with typical default loss values due to clutter. Hyphens represent clutter categories for which there are no field measurements available. Consequently, it is not possible to calculate an objective-function value for them.



The optimized loss for the first clutter category, 0, representing urban area without buildings, is larger than the default value in all three networks. This may be attributed to the fact that ...??? As for the category 1, representing suburban area, the value for Net<sub>1</sub> is lower than the default one, mainly because this network is deployed over a predominant agricultural area, i.e. suburban areas are less dense here. On the other hand, the value for Net<sub>2</sub> is larger, indicating a building density above the average, whereas for Net<sub>3</sub>, the value could not be calculated due to lack of measurements. A similar behavior may be observed for category 2, representing urban area. However, for category 3, representing dense urban area, the optimized losses of all three networks are lower than the default value, indicating dense urban areas here are not as dense as the average case. Representing agricultural area, category 4 gets a value very close to the default one for Net<sub>1</sub> and Net<sub>3</sub>, whereas for Net<sub>2</sub> the value is lower, indicating the most of this kind of land is not being exploited near the city. As for category 5, representing forests, the results are well corresponded with the kind of forest dominating each of the test networks, being those of Net<sub>1</sub> and Net<sub>3</sub> more dense due to leave foliage, whereas as for Net<sub>2</sub> the forests are mostly coniferous and more sparse. Keeping the default loss values for categories 6, 7 and 8, we move on to category 9, representing water, for which the results of all three networks indicate creeks and rivers in these areas are mostly surrounded by forests (Net<sub>1</sub>) or buildings (Net<sub>2</sub>), since none of the regions lays by the sea. As for the industrial areas, denoted by clutter category 10, show lower loss values than the typical default, indicating very sparse industrial buildings (Net<sub>1</sub>) and a higher density of mostly commercial buildings for Net<sub>2</sub> and Net<sub>3</sub>).

These results clearly show the benefit of the optimization of losses due to land usage or clutter, by finely adapting these values to local conditions within the environment where these networks are deployed.

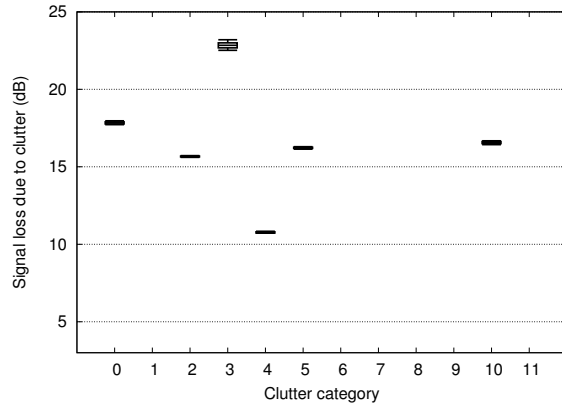
#### 6.5.4.1 Statistical analysis

Because of the stochastic nature of the optimization algorithm, we have collected the results from 30 independent runs, in order to have enough data for the results to be statistically relevant. In other words, the robustness of the solutions presented in the previous section is analyzed here.

To this end, Table 6.5 shows the solutions reached by the DASA for each of three test networks. The calculated signal losses are thus depicted with the minimum, maximum and average values for every clutter category, along with their standard deviation. Similarly as before, hyphens represent clutter categories for which there are no field measurements available, and thus they cannot be optimized.

Table 6.5: Statistical analysis of the optimization solutions for each test network. All values are expressed in dB.

Category	Net <sub>1</sub>				Net <sub>2</sub>				Net <sub>3</sub>			
	Min	Max	Avg	St.dev.	Min	Max	Avg	St.dev.	Min	Max	Avg	St.dev.
0	13.36	13.97	13.71	0.15	11.22	11.40	11.30	0.04	17.72	17.85	17.90	0.07
1	-	-	-	-	14.25	19.20	16.67	1.87	-	-	-	-
2	15.84	16.21	16.04	0.08	16.99	17.11	17.04	0.03	15.64	15.72	15.69	0.03
3	19.15	20.07	19.59	0.25	17.95	18.12	18.01	0.04	22.68	23.20	23.00	0.16
4	11.35	11.62	11.48	0.05	9.63	9.77	9.71	0.03	10.73	10.84	10.80	0.03
5	16.00	16.54	16.26	0.14	11.45	11.80	11.62	0.08	16.19	16.30	16.26	0.04
6	-	-	-	-	-	-	-	-	-	-	-	-
7	-	-	-	-	12.71	14.52	13.49	0.39	-	-	-	-
8	-	-	-	-	12.25	15.79	13.50	0.83	-	-	-	-
9	16.07	18.80	17.50	0.83	4.67	6.26	5.60	0.35	-	-	-	-
10	7.79	8.54	8.26	0.19	16.68	16.87	16.75	0.05	16.50	16.68	16.63	0.07
11	-	-	-	-	18.62	19.20	17.04	0.13	-	-	-	-



(c)

Figure 6.12: Box plots showing the statistical analysis values of Table 6.5, showing the solutions of the clutter-optimization process for each test network: (a) Net<sub>1</sub>; (b) Net<sub>2</sub>; (c) Net<sub>3</sub>.

## 6.6 Related work

There are a few examples of radio-network simulators available for LTE [92, 104? ], mostly developed for academic environments, they are not targeted at real-world environments. A Matlab-based LTE simulator has been proposed in [92]. It implements a standard LTE downlink physical layer, including Adaptive Modulation and Coding (AMC), multiple users, MIMO transmission and scheduler. Despite being open source and freely available, the fact of being implemented in Matlab make it very restrictive in terms of tackling bigger problem instances of real networks. A promising tool in this sense is presented in [104], where the authors implement a full-stack LTE system in C++. Although the tool has no capability for graphically displaying the simulations, it could be implemented, since the source code is available. In our opinion, the main drawback of this tool is the lack of documentation, which makes it very time-consuming to continue extending this work without the direct help of one of the original authors. In [68], the authors present a tool for radio-coverage calculations of wireless networks. Implemented as a module of the open-source GRASS system, it made an ideal basis for expanding it with parallel-computation capabilities.

As an extension to the well-known NS-2 network simulator, Filiposka and Trajanov [?] introduced a module for radio-propagation predictions, which takes the terrain profile into account. In this case, the authors focus on the relief and leave out signal loss due to land-usage, which is a key factor for acquiring more realistic radio-propagation predictions.

Following an optimization-oriented approach [? ], the authors study the effects of the location accuracy while performing a semi-automatic optimization of the parameters of a radio-propagation model. When compared to our work, where we take a fully automatic optimization approach, the advantage is clear, since no human intervention is needed during the optimization process. Besides, they conclude that locations with a median accuracy of around 60 mts, may be used for parameter tuning. In this sense, the GPS-informed location of the field measurements we had available, has been tested to be within this limit.

## 6.7 Sumary

We have presented an open-source simulation framework for planning and optimization of radio networks (PRATO). Based on extensive experimental simulations, we have shown the suitability of PRATO by tackling two planning and optimization cases, tested over the newly deployed LTE network in Slovenia. The first one involved the parameter tuning of an empirical radio-propagation model using a snapshot of field measurements. The other one consisted in optimizing the clutter losses over different regions of the country, therefore adapting the losses due to land usage to the local conditions of each region.

The encouraging results indicate that PRATO is applicable in planning and optimization of real-world radio networks in general, and LTE in particular. Moreover, even computational intensive tasks such as stochastic optimization, involving thousands of radio-prediction evaluations, are feasible thanks to the parallelization capabilities provided by the framework.

## Acknowledgments

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## 7 Performance assessment within real network-planning scenarios

In this chapter, real-world, network-planning activities on different radio networks are presented. The objective is to compare PRATO with a radio-prediction enterprise tool in terms of quality and performance. To this end, the engineers of the Radio Network department at Telekom Slovenije, d.d., provided us with their radio-prediction industrial software and guidelines to perform some typical radio-planning activities.

### 7.1 Measurements and simulation comparison

We have also developed two modules for the evaluation of the accuracy of the designed radio coverage prediction tool with respect to the actually measured results. The first module, db.CompareResults, compares the calculated values with the values obtained during a field measurement campaign. The module reads each row of the text file with field measurement data and saves the x and y coordinates, the measured signal level and the name of the cell. The module first maps the coordinates to the nearest coordinates in the database (as they do not necessarily coincide), finds the corresponding row in the database and then extracts the received signal level for the required cell. The data for the measured and calculated signal level, their difference in decibels along with the location, the used path loss prediction model and the cell name are finally written to the output text file (Table 3).

The second module, r.EvaluateSimulations, compares the field measurements and the strongest simulation signal levels, neglecting the information about the serving cell. The module output is a textual file including the same fields as the output of the db.CompareResults module. The module enables comparison of field measurements with simulation results calculated with the designed radio coverage prediction tool or the commercial tool TEMS. Because of different input raster files (a GRASS raster file includes values for maximal received power while a TEMS raster file contains path loss values) additional selection is done with the “GRASS MaxPower raster”. In the case of the TEMS input raster file, an average transmitted power value must be entered for the receive power calculation.

### 7.2 Coverage-prediction performance analysis

The performance and accuracy of the developed modules for radio signal coverage prediction was investigated by comparing simulation results and field measurements. The reference values were obtained by comparing field measurements with simulation results acquired from the professional radio signal coverage prediction program TEMS. In both simulation tools we used the modification of the Okumura-Hata propagation model [23].

The performance of the new software package was investigated for different types of networks (GSM, UMTS) and terrains (hilly and almost flat rural, urban, and suburban). The evaluation of the developed software for different frequency bands is presented first, followed by an analysis for a different terrain type.

The accuracy of the GRASS prediction software can be verified from the charts on Fig. 6, 7 and 8. On the left side, the charts comparing the measurements and calculations with the GRASS radio coverage prediction software are depicted, while graphs showing the comparison between the measurements and calculations with the TEMS software package are on the right.

Fig. 6 and 7 are presenting the simulation results from both tools and the field measurements in suburban environment for 900MHz and 2040MHz. Received power charts clearly shows that the simulation results match the measured values rather well. Slightly better agreement can be perceived in the 900MHz frequency band (Fig. 7). The deviation among the measurements and simulations for both software applications is depicted in the second raw of diagrams in Fig. 6 and 7. It is evident that the difference between the diagrams on the left- and on the right-hand side for both frequencies is minor. Thus, it can be concluded that the results from the developed radio coverage tool are comparable with the results from the TEMS application and are independent from the used frequency band.

Additional analyses were done on different terrain types. The analyses for the flat rural environment are depicted in Fig. 8. The curves on the charts showing the difference between the measurements and simulations for both software applications have similar course. This confirms applicability of the developed software also for arbitrary terrain types.

The developed radio coverage software gives similar results as the professional TEMS software irrespective of the operational frequency or chosen terrain type. The computed values are comparable also for different distances between the base station and the receiver. Some negligible differences between the results originate from the fact that the implemented path loss model in the TEMS software used in the simulation is not entirely available and thus cannot be realized in the GRASS software in a completely identical way.

Additionally, execution performance of the developed modules was evaluated in terms of the required processing times on our system (processor Intel Core2 Quad CPU 2,66GHz, disk WD2500KS, OS Linux RHEL5). The simulated configuration included eight transmission antennas on four locations (base stations), therefore requiring four model and eight sector computations. Two different geographic regions were used: a small one, "Ljutomer", encompassing all transmission locations (15 x 13km, resolution 25m) and a large one "Slovenia" (whole Slovenia, 285 x 185km, resolution 100m). The effective transmission radius was limited to 10km. The results for single core execution are given in Table 4. The hataDEM model was not simulated for the whole Slovenia region since its clutter map was not available to us. The r.MaxPower module was not run with DBF database on the whole Slovenia region since the internal GRASS DBF processing is very memory inefficient, keeping the whole database in the main memory and hence running out of memory for large regions.

### 7.3 Summary

Precise and efficient planning of the wireless telecommunication systems requires efficient and exact radio signal coverage calculations. The high price and limited functionalities of the existing professional network planning tools compels to look for alternative solutions. The needs can be fulfilled using an open-source system which gives possibilities to improve the existing models based on measurements, or to develop entirely new path loss prediction models. This paper presented a radio signal coverage prediction software tool developed for the open-source GRASS system. After a short introduction of the GRASS GIS system, a detailed description of the coverage prediction software was given. The tool enables a high level of flexibility and adaptability. It is composed of several GRASS modules for path loss calculation, a sectorization module, a module for radio signal coverage calculation, and additional modules for preparing input data and analyzing simulation results. Modules can be

used individually or through the r.radcov module, written in Python, which interconnects individual modules into a complete radio signal propagation software package. At the end, the developed software was evaluated by comparing with the field measurements and simulation results obtained from a professional software application. The radio signal coverage prediction software implementation was quite straightforward, as API is well developed and documented. The set of built-in C functions is adequate. The possibility to study parts of the already implemented code is also very helpful. Extensive performance analyses showed satisfactory results. Compared to a professional network planning tool, the computation speed is slightly lower while the result accuracy is completely comparable irrespective of the terrain type or operational frequency. For better agreement between simulations and measurements, additional model tuning will be performed. In our future work, we also plan to expand the functionalities of the developed software package and build additional path loss modules for the urban and hilly rural environments that will also include the elements of ray tracing techniques and additional environment data [29]. The achievement made so far represents a strong base for future work and is interesting both from the point of view of researchers as well as network developers. The whole source code of the radio signal coverage prediction tool together with detailed instructions will be publicly available at <http://commsys.ijs.si/en/software/grassradiocoverage> tool. The tool can be freely used, modified and upgraded with new path loss modules.



## 8 Conclusion and further work

Encouraged by the favorable results, further work will include abstracting the introduced MWD principle into a multi-purpose parallel framework such as Charm++ [73], which provides a functionality for overlapping execution and communication, as well as fault tolerance.

In addition, as PRATO is also a free and open-source software project<sup>1</sup>, it can be readily modified and extended to support, for example, other propagation models and post-processing algorithms. This characteristic provides it with a clear advantage when compared to commercial and closed-source tools.

Comparison of our experimental results with other algorithms dealing with the same and similar problems would be useful. However, this task is not straightforward, since the results of several works (e.g. [47, 139]) depend on black-box evaluations, making experimental association very difficult, if possible at all.

All in all, we consider that the present work provides a robust foundation for future work on grid-based metaheuristics with expensive objective-function evaluation.

In future work, we will consider further analysis of our parallel-agent approach, including experimentation with different parameters, in order to gain better understanding of the dynamics leading the metaheuristic during the search process. Multi-GPU environments present an interesting possibility, where evaluator(s) and worker agents are run on separate GPU devices.

To further improve the presented results, dynamic effects, such as fast power control, should be included in the simulations, particularly for recognizing dynamic functionality like SHO in a WCDMA mobile system. Another extension of the current work is to incorporate antenna tilt as an additional objective of the optimization process. This should certainly include experimentation with models and algorithms that support multiobjective optimization.

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<sup>1</sup>The source code is available for download from <http://cs.ijs.si/benedicic/>



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