Tools and Models for High Level Parallel and Grid Programming

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By

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2008
A tutta la mia famiglia,
con un particolare pensiero per lo zio Giancarlo.
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Producer-filter-consumer with parallel filter (farm skeleton).

Reconfiguration overhead: Stop.

Reconfiguration overhead: New.

Reconfiguration overhead: Restart.
Self-optimization experiment.
First of all I wish to thank my supervisor Marco Danelutto who always supported and encouraged me, leaving me the freedom of experimenting with many different topics, far beyond I have expected. If I wouldn’t dread being disrespectful I would say he is a real good friend. I am very grateful to Domenico Laforenza the head of the High Performance Computing Laboratory, where I spent most of time of the last three years. I give my thanks also to my co-workers: R. Perego, R. Baraglia, S. Orlando, F. Silvestri, N. Tonellotto, C. Lucchese, M. Coppola, D. Bocci, F. M. Nardini, G. Capannini and G. Tolomei. Many thanks go also to who co-authored my works: M. Aldinucci, S. Campa, L. Presti, A. Panciatici, M. Pasin, M. Vanneschi and P. Kilpatrick. I thank Marco Paquali, he was both a valuable co-worker and a precious co-author but, above all, he is a very good friend. I am very grateful both to my reviewers Vladimir Getov and Christian Perez, and to the member of my internal thesis committee Gianluigi Ferrari and Paolo Ciancarini for giving me many helpful suggestions. I express my gratitude to all my friends who shared with me the spare time. Among them Francesco, with whom I had many discussions about almost everything.

Now let me switch to Italian to... ringraziare tutta la mia preziosa famiglia, con particolare riferimento ai miei genitori che da anni mi supportano sia col loro affetto sia economicamente. Sono fermamente convito che possiate ritenervi i migliori genitori che un figlio possa avere. L’ultimo ringraziamento va alla persona che da anni ha il monopolio del mio cuore, Chiara, trovare le parole per ringraziarti è un compito impossibile, qualsiasi frase o parola non sarebbe mai abbastanza.
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Abstract

When algorithmic skeletons were first introduced by Cole in late 1980 (50) the idea had an almost immediate success. The skeletal approach has been proved to be effective when application algorithms can be expressed in terms of skeletons composition. However, despite both their effectiveness and the progress made in skeletal systems design and implementation, algorithmic skeletons remain absent from mainstream practice. Cole and other researchers, respectively in (51) and (19), focused the problem. They recognized the issues affecting skeletal systems and stated a set of principles that have to be tackled in order to make them more effective and to take skeletal programming into the parallel mainstream. In this thesis we propose tools and models for addressing some among the skeletal programming environments issues. We describe three novel approaches aimed at enhancing skeletons based systems from different angles. First, we present a model we conceived that allows algorithmic skeletons customization exploiting the macro data-flow abstraction. Then we present two results about the exploitation of metaprogramming techniques for the run-time generation and optimization of macro data-flow graphs. In particular, we show how to generate and how to optimize macro data-flow graphs accordingly both to programmers provided non-functional requirements and to execution platform features. The last result we present are the Behavioural Skeletons, an approach aimed at addressing the limitations of skeletal programming environments when used for the development of component-based Grid applications. We validated all the approaches conducting several test, performed exploiting a set of tools we developed.
Chapter 1

Introduction

Computers are becoming tools of vital importance. They are used almost everywhere, they are used for work, for study, for fun and actually for solve problem. Unfortunately, many problems require a huge amount of computational power to solve (as an example: genome mapping, portfolio risk-analysis, protein folding). Such a power cannot be obtained using a single processor. The only suitable solution is to distribute the application workload across many different computational resources. Resources those contemporaneously (“in parallel”) execute parts of the whole application. Programming applications that make use of several computational resources at the same time introduces some difficulties, as an example the communication and synchronization among the resources, or the application code and data decomposition and distribution. In order to ease this burden, since the early steps of computer science, researchers conceived and designed programming models and tools aiming at supporting the development of parallel applications. Throughout the ages, a lot of models and tools have been proposed, presented in several different (sometime exotic) forms. Nevertheless, the main goal is always the same: find a good trade-off between simplicity and efficiency. Indeed, a very abstract model simplifies the programming activity but can lead to a very inefficient exploitation of computing resources. Instead, a low-level model allows programmers to efficiently exploit the computational
resources but requires to programmers a tremendous effort when the number of resources grows. Since the nineties, several research groups have proposed the structured parallel programming environments (SPPE). Since the structured parallel programming model was conceived, several works have been done about it. Programming environments relying on this paradigm ask programmers to explicitly deal with the qualitative aspects of parallelism exploitation, namely the application structure and problem decomposition strategies. All the low-level parallelism exploitation related aspects like communication, synchronization, mapping and scheduling are managed by compiler tools and run-time support. In these environments parallelism is exploited by composing “skeletons”, i.e. parallelism exploitation patterns. From language viewpoint, a skeleton is a higher-order function that behaves as a pure function (no side-effects). Several real world, complex applications have been developed using these environments. The skeletal approach has been proved to be quite effective, when application algorithms can be somehow expressed in terms of skeleton composition. Notwithstanding, skeletal programming has still to make a substantial impact on mainstream practice in parallel applications programming.

1.1 Contribution of the thesis

This thesis originates from the wish to address the issues that have limited the diffusion of structured parallel programming environments. These issues are well-known by the structured parallel programming models scientific community. They have been organically reported in two key papers \cite{19,51} where the authors describe both the issues and the features that the next generation of structured parallel programming environments have to support in order to address them. The features “check-list” includes, as an example, the ease of use, the integration of structured and unstructured form of parallelization, the support for code reuse, the heterogeneity and dynamicity handling. Drawing a parallel with web programming model we can refer as “Skeletons 2.0” the next generation of structured parallel programming environments that address the
issues that prevent the skeleton environment to became part of the main-stream practice in parallel applications programming. Some groups of re-searchers involved in structured parallel programming developed skele-ton systems that have partially addressed the “Skeletons 2.0” principles to different degrees in different combinations. Nevertheless, the research for addressing the presented issues has just started. Indeed, up to now, tools and models that are generally recognized as the best solutions for addressing the issues still do not exist.

The main goal of this thesis is to present an organic set of tools and models conceived, designed and developed to address most of these issues, therefore form the base of a next generation skeleton system. The scientific contribution of the thesis is organized in three main parts. They reports four results we obtained in the last three years. These research results as has been already presented in published papers. Some results have been achieved with actual experiments conducted using software tools and packages designed and developed to the purpose. Some of them are simple, proof-of-concept tools, like JJPF \cite{59} or PAL \cite{61}. Some others are custom version of existing framework, like muskel with the support for developing unstructured form of parallelism \cite{21} or muskel with an aspect oriented programming support \cite{60}. Others are part of complex international research project focused on Grid computing, like the Behavioural Skeletons \cite{17}.

Our first contribution copes with the lack of models supporting the integration of unstructured form of parallelization in skeleton systems. In fact, if on the one hand structured parallel programming environments raise the level of abstraction perceived by programmers and guarantee good performance, on the other hand they restrict the freedom of pro-grammers to implement arbitrary parallelism exploitation patterns. In or-der to address this issue we propose a macro data-flow based approach that can be used to implement mixed parallel programming environments providing the programmer with both structured and unstructured ways of expressing parallelism. Structured parallel exploitation patterns are implemented translating them into data-flow graphs executed by a distributed macro data-flow interpreter. Unstructured parallelism exploita-
tion can be achieved by explicitly programming data-flow (sub)graphs. To validate the approach, we modified a skeleton system that in its original form does not deal with unstructured parallelism: muskel. We extended muskel, in collaboration with the research staff that developed it. Our customized muskel is implemented exploiting (macro) data-flow technology, rather than more usual skeleton technology relying on the usage of implementation templates. Using data-flow, the extended muskel supports the development of both classical, predefined skeletons, and programmer-defined parallelism exploitation patterns. Our extended version provides two mechanisms to the muskel programmers for unstructured parallelism exploitation. First, we provide primitives that allow to access the fundamental features of the data-flow graph generated out of the compilation of a skeleton program. Namely, methods to deliver data to and retrieve data from data-flow graph. We provide to programmers the ability to instantiate a new graph in the task pool by providing the input task token and to redirect the output token of the graph to an arbitrary data-flow instruction in the pool. Second, we provide the programmer with direct access to the definition of data-flow graphs, in such a way he can describe his particular parallelism exploitation patterns that cannot be efficiently implemented with the available skeletons. The two mechanisms can be jointly used to program all those parts of the application that cannot be easily and efficiently implementing using the traditional skeletons subsystem. Unfortunately, this approach is not free from shortcomings. In fact, exploiting unstructured parallelism interacting directly with data-flow graph requires to programmers to reason in terms of program-blocks instead of a monolithic program.

In order to ease the generation of macro data-flow blocks and in general to provide mechanism easing the use of structured parallel programming environment, we exploited some metaprogramming techniques. Exploiting these techniques the programmers are no longer requested to deal with complex application structuring but simply to give hints to the metaprogramming support using high-level directives. The directives drive the automatic application transformation. In this thesis we present two results we obtained regarding the exploitation of metaprogramming
techniques for parallel programming. The first result is “Parallel Abstraction Layer” (PAL). A java annotation based metaprogramming framework that restructures applications at bytecode-level at run-time in order to make them parallel. The parallelization is obtained asynchronously executing the annotated methods. Each method call is transformed in a macro data-flow block that can be dispatched and executed on the available computing resources. PAL transformations depend on both on the resources available at run-time and the hints provided by programmers.

The other result concerns the integration of the Aspect Oriented Programming mechanisms with our modified muskel skeleton framework. We make this integration in two distinct phases, in the first phase we integrated the AOP mechanisms in order to achieve very simple code transformation. In the second phase we implemented a more complex integration to obtain a support enabling the development of workflows which structure and processing are optimized at run-time depending on the available computational resources.

In this thesis we present also a model to address two other issues: the lack of support for code reuse, and the lack of support for handling of dynamicity. The muskel framework, addresses this last point through the definition of the Application Manager, namely an entity able to observe, at run-time, the behavior of the parallel application and in case of faults or application non-functional requirement violations it reacts aiming to fix the problem. The dynamicity handling is a very important feature for next generation parallel programming systems, especially for the ones designed for computational Grids. Indeed, Grid are often composed by heterogeneous computer and managed by different administration policies. To address these additional difficulties most of the models and tools conceived and developed for parallel programming have to be re-thought and adapted. Actually, the muskel framework, at least in its original form, is designed to be exploited in cluster and network of workstations rather than in Grids. Indeed, some of the implementation choices done when it was developed limit its exploitation on Grids, in particular the ones related with communication protocol and with the mechanisms for recruiting computational resource. On the other hand, several studies
recognized that component technology could be leveraged to ease the development of Grid Application (25; 72). Indeed, a few component based model have been proposed by parallel computing scientific community for programming Grids, as CCA (5), CCM (67) and GCM (52). The GCM represents one of the main European scientific community efforts for designing and developing (3) a grid component model. We contributed to the design of GCM and its reference implementation together with the research group that developed muskel and with several European research groups. In particular, we focused our contribution on the GCM autonomic features. We referred to the muskel Application Manager approach, generalizing it and extending the approach to make it suitable for components based models. Indeed, each GCM component with a complete support of autonomic features has an Autonomic Manager that observes the component behavior. In case the behavior turns out to be different from the expected one the manager trigger a component reconfiguration. In other words, GCM autonomic features provide programmers with a configurable and straightforward way to implement autonomic grid applications. Hence, they ease the development of application for the Grids. Nevertheless, they rely fully on the application programmer’s expertise for the setup of the management code, which can be quite difficult to write since it may involve the management of black-box components, and, notably, is tailored for the particular component or assembly of them. As a result, the introduction of dynamic adaptivity and self-management might enable the management of grid dynamism, and uncertainty aspects but, at the same time, decreases the component reuse potential since it further specializes components with application specific management code. In order to address this problem, we propose the Behavioural Skeletons as a novel way to describe autonomic components in the GCM framework. Behavioural Skeletons aim to describe recurring patterns of component assemblies that can be equipped with correct and effective management strategies with respect to a given management goal. Behavioural Skeletons help the application designer to i) design component assemblies that can be effectively reused, and ii) cope with management complexity. The Behavioural Skeletons model is an effective solution for handling dynam-
icity, supporting reuse both of functional and non-functional code. We want to point out that we have not the “sole rights” concerning the Behavioural Skeletons model. Indeed, it has been developed in conjunction with the other authors of the two papers about Behavioural Skeletons we published [16; 17].

This thesis is not our first attempt of design programming model for parallel programming. In a previous work we developed JJPF, a Java and Jini based Parallel Framework, and investigated the possibilities offered by structured parallel programming. In [59] we described the architecture of JJPF. JJPF was specifically designed to efficiently exploit affordable parallel architectures, such as a network of workstations. Its reactive fault-tolerance support and its dynamic support for task distribution as well as for resources recruiting were designed to enable an efficient exploitation of resources in highly dynamic environment. In particular, JJPF exploits the Jini technologies to dynamically find and recruit the available computational resources. JJPF provide to programmers an API enabling the development of task-parallel application following the master-slave paradigm. It also provides an high-level support for data sharing among slaves. JJPF ease the parallel programming task hiding most of low-level error prone issues to programmers. As we stated above, JJPF is implemented in Java. It simplifies the code portability among heterogeneous architectures. For the communications among master and slaves JJPF exploits the JERI. It is a variant of RMI allowing the protocol customization and as a consequence an optimization of its performance in several situations. For the performance purpose JJPF also provides an alternative to the java distributed class-loader that reduces the class-loading latency in some situations. Some problems encountered during the design of JJPF still remain open. Moreover, during the realization of JJPF we faced directly with the development complexity of this kind of software so we think that some kind of software engineering is needed to facilitate reuse and maintenance of source code.
1.2 Thesis Outline

As we already stated, in this thesis we report our contribution to address the issues that are typical of traditional structured parallel programming environments. The contribution is organized in three main parts. Each part is presented in a dedicated chapter. Moreover, there are three more chapters: an Introduction chapter (this one, actually), a Conclusion chapter and another one that introduces the problems we face in this thesis and outlines the state-of-the-art of existing solutions. In the remain of this section we describe the content of each chapter.

Chapter 2 In this chapter we take into account the problems related to programming parallel applications, the existing solutions and their main limitations. In particular, after a general introduction to the different parallel programming models, the topic is focused on the limitations that prevent the structured parallel programming models from spreading and to become part of the mainstream practice. Section 2.1 gives a bird’s-eye view both on the parallel architectures and on the fields in which parallelism has traditionally been employed. Section 2.2 reports a selection of the main parallel programming models distinguishing between the implicit (Section 2.2.1) and explicit (Section 2.2.2) approaches. The explicit approaches are further discussed subdividing them, with respect to the abstraction presented to programmers, in high-level (Section 2.2.3) and low-level (Section 2.2.4) ones. For each of them are presented the Pros and Cons. The chapter reports also some other notable approaches (Section 2.2.5). Then the Chapter present the structured approach, an approach conceived in order to overcome the limitations of traditional approaches (Section 2.2.6). Some tools based on the structured parallel programming models are presented (Section 2.2.6) and others are reported as well as references to the literature. The models are presented highlighting their features and main limitations. Section 2.3 reports the issues that next generation skeleton system should own to address the existing limitations. Finally, the chapter introduces (Section 2.4) our contributions to the field placing them in the proper context, showing how such contributions can
be exploited for addressing the issues related to structured parallel programming environments.

Chapter 3 In this Chapter we discuss a methodology that can be exploited in order to provide to programmers the possibility to mix structured and unstructured ways of expressing parallelism while preserving most of the benefits typical of structured parallel programming models. The methodology is based on the data-flow model. Unstructured parallelism exploitation is achieved by explicitly programming data-flow graphs. Section 3.1 briefly recalls the structured programming models outlining their main advantages and limitations. In particular, the section focuses on the skeleton customization issue. Namely the lack of flexibility of skeletal systems in expressing parallel form different from the ones that are “bundled” with the skeleton framework and their compositions. Then the section introduces the macro data-flow based approach we conceived in order to address of this limitation and reports the related work: alternative approaches addressing the structured parallel programming limitations. Section 3.2 introduces both the classical template-based implementation of skeleton systems and the more recent data-flow technologies based one used in muskel. Section 3.3.1 describes the details of our contribution, i.e. how we exploited the methodology presented to extend the muskel framework. Finally, Section 3.4 reports the experimental results we obtained conducting some test using our customized muskel framework.

Chapter 4 In this Chapter we introduce some novel metaprogramming techniques for the generation and optimization of macro data-flow blocks. This Chapter presents our efforts aimed at providing metaprogramming tools and models for optimizing at run-time the execution of structured parallel applications. The approaches are based on the run-time generation of macro data-flow blocks from the application code. The Chapter discusses how we exploited these techniques both in our modified muskel framework as well as in other frameworks we developed. Section 4.1 presents the motivations behind our contributions. Section 4.2
presents PAL, our first result in the field. The core of PAL framework is its metaprogramming engine that transforms at run-time an annotated sequential java code in a parallel program exploiting both programmer hints and information about executing platforms. Section 4.2.1 describes the details of our PAL prototype implementation. Section 4.2.2 reports the experimental results we obtained testing PAL framework. Section 4.2.3 discusses the motivations that convinced us to integrate the PAL approach to our modified muskel framework. Section 4.3 describes the preliminary attempts we made integrating metaprogramming techniques in muskel showing how Aspect Oriented Programming can be exploited to do some simple code transformations. Section 4.4 describes how we further enhanced muskel making it able to exploit metaprogramming for run-time code optimizations. In particular, how it can be exploited to optimize the parallel execution of computations expressed as workflows. Section 4.4.2 describes the implementation details of workflows transformations and Section 4.4.3 presents the results of some experiments we conducted. Finally Section 4.5 presents a comparison of the two approaches.

Chapter 5 In this Chapter we present some results about the customization of skeletons applied to the Grid Component Model. In this chapter we present the Behavioural Skeletons model, an approach, we contribute to conceive and validate, aimed at provide programmers with the ability to implement autonomic grid component-based applications completely taking care of the parallelism exploitation details by simply instantiating existing skeletons and by providing suitable, functional parameters. The model has been specifically conceived to enable code reuse and dynamicity handling. Section 5.1 describes how component-based applications can ease the task of developing grid applications. Section 5.2 outlines the grid component model focusing on its autonomic features. After, Section 5.4 presents the Behavioural Skeletons model, Section 5.5 reports a set of noteworthy Behavioural Skeletons and Section 5.6 describe their GCM implementation. Section 5.7 describes a set of experiment we conducted to validate the Behavioural Skeletons model.
Chapter 6 This Chapter summarizes the materials contained in the previous chapters and discusses the conclusions of the thesis. Finally, the future work related to the thesis is introduced.
Chapter 2

High-Level Parallel Programming

As we already stated in the Introduction, using several processors (or computational resources) at the same time (in parallel), however, introduces some difficulties. The conceptual barrier encountered by the programmers in efficiently coordinating many concurrent activities towards a single goal is an example of such barriers. To address these difficulties software developers need high-level programming models for sensibly raising the abstraction of computational resources. This is a fundamental requirement to avoid programmers having to deal with low-level coordination mechanisms. In fact, low-level parallel programming is an error prone approach that distracts programmers from qualitative aspects of parallelization. Throughout the ages, researchers conceived and developed several models for high-level parallel programming. However, most of current implementations of very high-level programming models often suffer from low performance. This because of the abstraction penalty, which actually has historically limited the usage of high-level programming techniques in high performance computing. For this reason, nowadays most of parallel programs are developed exploiting lower-level language, even if a higher-level language would make the coding easier. Structured parallel programming models were conceived to be an al-
ternative both to very high-level models and to low-level models. Structured parallel programming models ask programmers to explicitly deal with the qualitative aspects of parallelism exploitation, namely the application structure and problem decomposition strategies. Compilers and run-time supports manage all the low-level parallelism exploitation related aspects like communication, synchronization, scheduling and mapping. The Structured Way is driven by those two observations: there are some things programmers do better than compilers, and there are some things that compilers do better than programmers. Nevertheless, also the structured models are not perfect and free from limitations. In fact, for some years researchers very expert in structured parallel programming models have outlined the features that the next generation of structured models have to provide in order to address these limitations (19; 51). In next three chapters of this thesis we present some results we obtained as an attempt of address some of these limitations.

Chapter road-map The chapter starts with a bird’s-eye view both on the parallel architectures and on the fields in which parallelism has traditionally been employed (Section 2.1). Then, it reports the main parallel programming models (Section 2.2) distinguishing between the implicit (Section 2.2.1) and explicit (Section 2.2.2) approaches. The explicit approaches are further subdivided in high-level (Section 2.2.3), and low-level (Section 2.2.4) ones. The chapter reports also some other notable approaches (Section 2.2.5). Then the Chapter present the structured approach, an approach conceived in order to overcome the limitations of traditional approaches (Section 2.2.6). Some tools based on the structured parallel programming models are presented (Section 2.2.6) highlighting their features and main limitations. Then Section 2.3 reports the issues that next generation skeleton system should own to address the existing limitations. Finally, the chapter introduces (Section 2.4) our contributions to the field placing them in the proper context, showing how they can be exploited for addressing some of the issues related to structured parallel programming environments.
2.1 From sequential to parallel architectures

The Von Neumann architecture is a very common and well-known computer design model. It has a very simple formulation and can be described as a sequential process running in a linear address space. It consists in a processing unit and a single separate storage structure to hold both instructions and data. The Von Neumann model “implements” a universal Turing machine. It represents the common “referential model” of specifying sequential architectures, in contrast with parallel architectures. In a parallel architecture many instructions are carried out simultaneously. Parallel computers operate on the principle that large problems can almost always be divided into smaller ones, which may be carried out at the same time. Parallel architectures exist in several forms and levels. They range from superscalar processors to computational Grids. In this section we briefly mention some of the most common forms of parallelism, without claiming to be exhaustive but only to give an idea of the variety of the existing forms of parallelism.

Bit-level parallelism is a form of parallelization based on increasing processor word size. It leads to a reduction of the number of instructions the processor must execute in order to perform an operation on variables whose sizes are greater than the length of the word. (For instance, consider a case where a 16-bit processor must add two 32-bit numbers. The processor must first add the 16 lower-order bits from each number, and then add the 16 higher-order bits, and the carry from the previous add requiring two instructions to complete a single operation. A 32-bit processor would be able to complete the operation using a single instruction). Historically, 4-bit microprocessors were replaced with 8-bit, then 16-bit, then 32-bit microprocessors. This trend generally came to an end with the introduction of 32-bit processors, which has been a standard in general purpose computing for two decades. Only recently, with the proliferation of processors based both on the IBM PowerPC G5 processor and on the x86-64 architectures, the 64-bit processors have become commonplace.
Instruction-level parallelism is a form of parallelization based on the simultaneous execution of instructions part of a computer program. Even if ordinary programs are typically written according to a sequential execution model where instructions execute one after the other and in the order specified by the programmer, in some significant cases there is no need to follow this order. ILP allows the compiler and the processor to overlap the execution of multiple instructions or even to change the order in which instructions are executed. Due to its nature, ILP requires an hardware support; micro-architectural techniques that are used to exploit ILP include (for a better description see (104)):

- Instruction pipelining, where the execution of multiple instructions can be partially overlapped.

- Superscalar execution, in which multiple execution units are used to execute multiple instructions in parallel. In typical superscalar processors, the instructions executing simultaneously are adjacent in the original program order.

- Out-of-order execution, where instructions execute in any order that does not violate data dependencies. Note that this technique is orthogonal w.r.t. both pipelining and superscalar.

- Register renaming, which refers to a technique used to avoid unnecessary serialization of program operations imposed by the reuse of registers by those operations, used to enable out-of-order execution.

- Speculative execution, which allows the execution of complete instructions or parts of instructions before being certain whether this execution should take place or not. A commonly used form of speculative execution is control flow speculation where instructions following a control flow instruction (e.g., a branch) are executed before the target of the control flow instruction is determined. Several other forms of speculative execution have been proposed and are in use including speculative execution driven by value prediction, memory dependence prediction and cache latency prediction.
- Branch prediction, which is used to avoid stalling for control dependencies to be resolved. Branch prediction is used with speculative execution.

**Data parallelism** is a form of parallelization of computer code across multiple processors in parallel computing environments. This paradigm is useful for taking advantage of the large amounts of data parallelism that is available in many scientific/numeric applications. The data parallelism is exploited by performing the same operation on a large amount of data, distributed across the processors of the machine. From the programmer viewpoint, languages based on data-parallel paradigm (such as HPF, sketched in Section 2.2.5) are pretty similar to sequential languages. The main difference is that certain data types are defined to be parallel. Parallel data values consist of a collection of standard, scalar data values.

The data-parallel paradigm has some main virtues that have led to its success. Parallel data types are typically static in size (e.g. arrays); their distribution across the machine is usually done at compile-time. Any synchronization or communication that is needed to perform an operation on a parallel value is automatically added by the compiler/run-time system. The processors collectively compute operations on parallel data values; computation load usually distributed directly linking data values and computations through the owner computes rule. As data values, computation load is statically distributed across the processors of the system. The data parallelism approach typically offers very good scalability. Because operations may be applied identically to many data items in parallel, the amount of parallelism is dictated by the problem size. Higher amounts of parallelism may be exploited by simply solving larger problems with greater amounts of computation. Data parallelism is also simple and easy to exploit. Because data parallelism is highly uniform, it can usually be automatically detected by an advanced compiler, without forcing the programmer to manage explicitly processes, communication, or synchronization. Many scientific applications may be naturally specified in a data-parallel manner. In these settings, programs data layout is often fixed; the most used data structures are large arrays. Operations
on whole data structures, such as adding two arrays or taking the inner product of two vectors, are common, as are grid-based methods for solving partial differential equations (PDEs). In spite of this, data parallelism has a significant drawback: the limited range of applications for which data-parallel is well suited. Applications with data parallelism tend to be static in nature; the control flow of a data-parallel program is mostly data independent. Many applications are more dynamic in nature and do not have these characteristics. To run in parallel, these dynamic applications need to perform independent operations at the same time. These applications, which may be as simple as recursively computing Fibonacci numbers or as complex as computer chess and n-body simulations, are nearly impossible parallelize using data parallelism.

Task parallelism is a form of parallelization of computer code across multiple processors in parallel computing environments. Task parallelism focuses on distributing execution processes across different parallel computing nodes. In the task-parallel paradigm the program consists of a set of (potentially distinct) parallel tasks that interact through explicit communication and synchronization. Task parallelism may be both synchronous and asynchronous. A major advantage of task parallelism is its flexibility. Many scientific applications contain task parallelism. For example, in a climate model application the atmospheric and ocean circulation may be computed in parallel. A task-parallel language can express this relationship easily, even if different methods are used for the two circulation models. Another natural application of task-parallel languages is reactive systems in which tasks must produce output in response to changing inputs, in a time-dependent manner. Another common structured paradigm exploits parallelism on different data items through task replication. For example, the elaboration of a video stream may involve the filtering on each single frame. In a task-parallel language the filter may be farmed out by spreading different frames on different worker processes, each of them computing the same function. In the task parallelism approach the interactions between tasks are explicit, thus the programmer can write programs that exploit parallelism not detectable
automatically by compiler techniques. In general, task parallelism is less
dependent on advanced compiler technology than the data parallelism;
in many cases, all that is strictly necessary is the translation of task inter-
actions into appropriate low-level primitives on the target architecture. A
disadvantage of the task-parallel programming model is that it requires
extra effort from the programmer to create explicit parallel tasks and man-
age their communication and synchronization. Because communication
and synchronization are explicit, changing the manner a program is par-
allelized may require extensive modifications to the program text.

Due to their nature data and task parallelism (unlike the bit level and
instruction level parallelism) cannot be fruitfully exploited using a sin-
gle CPU system but they are well-tailored for multi-processors or cluster
computers, typically referred as parallel computers.

For many years parallel computers has been mainly used in high per-
formance computing, but they have spread in recent years as convenient
and effective way to increase the computational power of personal com-
puters and workstations due to physical constraints preventing frequency
scaling of CPUs. Hence, parallel architectures are becoming the dominant
paradigm in computer architecture, mainly in the form of multicore pro-
cessors (28). Indeed, if a problem requires a huge computational capacity
to be rapidly solved and such a power cannot be obtained using a single
processing element (PE) the only suitable solution is to use many proces-
sors simultaneously. Traditionally, parallel architectures have been mo-
tivated by numerical simulations of complex systems and “Grand Chal-
lenge Problems” such as:

- weather and climate forecasting
- chemical and nuclear reactions simulations
- biological, human genome analysis
- geological, seismic activity analysis
- mechanical devices and electronic circuits’ behavior simulations
Today, also commercial applications need the development of faster and faster computers. These applications require to process large amounts of data in sophisticated ways. Example applications include:

- parallel databases, data mining
- web search engines, web based business services
- computer-aided medical diagnosis
- management of national and multi-national corporations
- advanced graphics and virtual reality, particularly in the entertainment industry
- networked video and multi-media technologies
- collaborative working environments

Unfortunately, as we already stated before, using several PEs at the same time introduces some difficulties. Among the others: (i) code and data have to be decomposed and distributed among the computational resources, (ii) work and communications of resources have to be simultaneously coordinated and (iii) fault-tolerance has to be managed. Thus, the design and implementation of software systems that can ease this burden is very important. Indeed, since the early steps of computer science, researchers conceived and designed programming models, systems and tools aiming at supporting the development of parallel applications. Such systems must find a good balance between the simplicity of the interface presented to the programmers and their implementation efficiency. Finding a good trade-off is a grand challenge. Indeed, a very abstract model simplifies the programming activity but can lead to a very inefficient exploitation of computing resources. Instead, a low-level model allows programmers to use efficiently the computational resources but requires tremendous efforts from the programmers when the number of resources grows.
2.2 Parallel programming models: State-of-the-art

A good way to organize the state of art of parallel programming models for reporting purpose is to divide them with respect to their level of abstraction. Therefore, in this section we report a selection of the main parallel programming models, proposed by computer scientist over the years, classifying them with respect to the level of abstraction provided to programmers. With respect to this aspect, the parallel programming models can be roughly partitioned in two main classes: the implicit parallel models and the explicit ones. The former completely cover up parallelism to programmers. Typically, they are exploited by functional and logic languages. The latter ask programmers to deal directly with parallelism. These models can be further partitioned, w.r.t. the abstraction perspective, in three categories: high, medium and low-level programming models.

In the remaining of this section we describe for each category, by way of examples, some programming models and tools belonging to it showing the models Pros & Cons. In particular, Section 2.2.1 describes the functional and logic models as an example of implicit models for parallel programming, Section 2.2.3 shows the data-flow model as a representative of high-level explicit models. In Section 2.2.4 we outline the low-level approaches describing the OpenMP and MPI frameworks. Then, in Section 2.2.5 we report some other notable approaches. Finally, we describe the structured approach in Section 2.2.6, it is one of the main medium-level models. Here we describe also some our past contributions in the field (Section 2.2.6).

2.2.1 Implicit approaches

These systems present to programmers a programming model entirely devoid of parallelism and completely isolated from the underlying implementation mechanism. Such systems typically present functional or logical models of computation. They are often referred to as being “declara-
tive” systems, since the programmer makes a series of declarations defining the properties of a solution to some problem, rather than specifying a precise series of operations which will lead to the solution. Thus, languages of this type are neither parallel nor sequential, having no notion at all of a flow of control.

Functional languages are based on the lambda calculus. It is a very simple, but powerful language to define expressions and their transformation rules. The only objects present are identifiers, single argument function definitions (“abstractions”) and applications of functions to arguments. A “program” consists of a collection of such objects. The program execution is performed applying a top-level function to an argument. This type of function application is the only operation present and involves the replacement of a function-argument pair with a copy of the function body (from its definition) in which occurrences of the “free” variable have been replaced by copies of the actual argument. This simple system can be shown to provide as much computational power as any other fundamental computing mechanism (e.g. the Turing machine). A particularly powerful aspect of the model is the ability to define “higher order functions”, namely, functions taking functions as input parameter. Other convenient features such as multiple argument functions, localized definitions and data structures may all be defined as lambda expressions.

In the same way, a high-level functional program is simply a function definition that refers to other functions in its body. A “call” of the program involves supplying arguments to this function and “execution” consists of using the function definitions (conceptually using the application by substitution technique from the lambda calculus) to obtain an alternative, but equivalent representation of the function and arguments pair, namely a more useful representation of the original program and the “input”.

The key point of this approach is that execution may progress from the initial to the final representation in any fashion that preserves the equivalence. In particular, it will often be possible to execute many transformation steps concurrently since the conventional problems associated with changes of state have been discarded along with the notions of state and
store themselves. A quite common way to represent the program as it evolves is as a graph, in which nodes represent function applications and the children of a node are the ("input") arguments of the corresponding application. The process of expanding and contracting the graph is referred to as "graph reduction".

Exploiting this approach, the parallelization via decomposition is simple. The abstract execution model allows candidate nodes to be expanded at any time, while function applications may be evaluated as soon as arguments are available. Thus, a potentially parallel process is generated every time a node reaches one of these states.

It is important to realize that this approach does not imply that every functional program is a highly parallel one. As a trivial, well-known, example, consider defining a function to compute factorials.

The obvious definition will look something like this:

\[
\begin{align*}
\text{factorial } 0 &= 1 \\
\text{factorial } n &= n \times \text{factorial}(n - 1)
\end{align*}
\]

Such a function would execute in a sequential way on a typical graph reduction machine, irrespective of the number of available processors. A more complex definition notes that

\[
\begin{align*}
\text{factorial } 0 &= 1 \\
\text{factorial } n &= \text{product } 1 \ n \\
\text{product } a \ a &= a \\
\text{product } a \ b &= (\text{product } a \lfloor \frac{a+b}{2} \rfloor) \times (\text{product } \lfloor \frac{a+b}{2} \rfloor + 1 \ b)
\end{align*}
\]

This definition produces significant potential parallelism. Although declarative systems involve no explicit notion of execution sequence, it is unfortunately clear that, in order to optimize the parallel execution programmers must be aware of the execution mechanisms.

An alternative approach recognizes the difficulty of automating distribution process and introduces program annotations that programmers
exploit to drive the execution mechanism in order to improve its efficiency. Such additions may be argued to move the model out of this category, in that the programmer is now partially responsible (and aware) for the task of parallel decomposition. Similarly, (81) discusses a language which allows program partitioning and interconnection structure to be described in a declarative style.

Another category of implicit systems consists in parallel logic languages. They are based on Horn clauses, a restricted form of first order logic. The computational model focuses on the definition and investigation of relationships described as predicates, among data objects described as input arguments to these predicates. As in functional programming, the specification of a computation consists of a collection of predicates and clauses. In the logic model the role of the outermost function application, is played by the outermost predicate together with its arguments. The arguments interpretation is similar: “execution” consists of deciding whether the predicate is true given the arguments and the associated definitions. Furthermore, it is possible to specify the outermost predicate with unbound arguments to find bindings to the arguments that allow the predicate to be satisfied, or to determine that no such bindings exist.

At an abstract level, the process of evaluation may be seen as expanding and searching a tree of possibilities presented by consideration of the various dependencies between appropriate predicates and clauses. As with graph reduction, the semantics of pure logic languages often allow this process to proceed at many points in parallel. Four principal kinds of (implicitly exploitable) parallelism can be identified in logic programs:

- **Unification parallelism** arises when arguments of a goal are unified with those of a clause head with the same name and arity. The different argument terms can be unified in parallel as can the different subterms in a term (34). Unification parallelism is very fine-grained and has been exploited by building specialized processors with multiple unification units.

- **Or-parallelism** arises when more than one rule defines some relation
fib(0, 1).
fib(1, 1).
fib(M, N) :- [ M1 is M - 1, fib(M1, N1) ],
              [ M2 is M - 2, fib(M2, N2) ],
              N is N1 + N2.

Figure 1: Fibonacci program parallelizable with independent and-parallelism

and a procedure call unifies with more than one rule head; the corresponding bodies can then be executed in parallel with each other. Or-parallelism is a way of efficiently searching for solutions to the query, by exploring alternative solutions in parallel.

- **Independent and-parallelism** arises when more than one goal is present in the query or in the body of a procedure, and the run-time bindings for the variables in these goals are such that two or more goals are independent of one another, i.e., their resulting argument terms after applying the bindings of the variables are either variable-free or have non-intersecting sets of variables. Parallel execution of such goals result in and-parallelism.

- **Dependent and-parallelism** arises when two or more goals in the body of a procedure have a common variable and are executed in parallel. Dependent and-parallelism can be exploited in two ways: (i) the two goals can be executed independently until one of them accesses/binds the common variable. (ii) Once the common variable is accessed by one of the goals, if it is bound to a structure, or stream (the goal generating this binding is called the producer), and this structure is read as an input argument of the other goal (called the consumer) then parallelism can be further exploited by having the consumer goal compute with one element of the stream while the producer goal is computing the next element. Case (i) is very similar to independent and-parallelism. Case (ii) is sometimes also referred to as stream-parallelism and is useful for speeding up producer-consumer interactions.

Figure 1 shows a simple program for computing the Fibonacci number.
The two lists of goals, each enclosed within square brackets above, have no data-dependencies among themselves and hence can be executed independently in parallel with each other. However, the last subgoal \( N = N_1 + N_2 \) depends on the outcomes of the two and-parallel subgoals, and should start execution only after \( N_1 \) and \( N_2 \) get bound. Consider that, as in case of functional languages, the programmers in order to exploit the potential application parallelism should give a proper structure to the program.

It should be pointed out that exist some extensions for logic programming language with explicit constructs for concurrency. They can be largely put into three categories:

- those that add explicit message passing primitives to Prolog, e.g., Delta Prolog \(^{[95]}\) and CS-prolog \(^{[71]}\). Multiple Prolog processes are run in parallel that communicate with each other via messages.
- those that add blackboard primitives to Prolog, e.g., Shared Prolog \(^{[46]}\). These primitives are used by multiple Prolog processes running in parallel to communicate with each other via the blackboard.
- those based on guards, committed choice, and data-flow synchronization, e.g., Parlog \(^{[48]}\), GHC \(^{[112]}\), and Concurrent Prolog \(^{[102]}\).

As for the functional languages, the extensions of parallel logic languages move the approach outside the category of implicit parallel programming models.

Similarities between functional and logic styles are emphasized in \(^{[65]}\).

**Summarizing Pros and Cons** Implicit parallel programming models provide programmers a very expressive programming metaphor: programmers can implement parallel application without actually deal with parallelism. Unfortunately, this ease is paid in terms of efficiency. In order to address such performance issues researchers introduced some annotation mechanisms and communication primitives, through which programmers can drive the code parallelization. Nevertheless, such additions place the model out of highly abstract systems category because the programmer exploiting annotations is partly responsible and aware for the task of decomposition.
2.2.2 Explicit models

The inefficient exploitation of available parallelism caused by the absence of parallel structure in implicit parallel programs is the main reason why explicit parallel programming models exist. These models are based on the assumption that programmers are often the best judges of how parallelism can be exploited for a particular application. Actually, in nearly every case the use of explicit parallelism will obtain a better efficiency than implicit parallelism models.

2.2.3 High-level explicit models: data-flow

The models belonging to this category still not require programmers to deal with the several issues related with parallel programming. For instance communications, fault-tolerance, heterogeneity, data decomposition and task granularity. Programmers are only required to write their applications as a set of independent instructions that interact each other through well-known interfaces, so that automatic tools can execute it in parallel. The data-flow model of computation is the main representative of this class of models.

In the data-flow model (for a deep description see (26; 80; 103; 115)) the computations are represented by a graph of “operator” or “instruction” nodes connected by edges along which data items flow. Each node receives by its input edges the data “tokens”, it performs some simple, stateless, calculation and distributes resultant data tokens on its output edges. A node may only perform its operation once it has received all the data tokens required, from all of its inputs. Thus, each node may compute in parallel, subject only to the availability of data. The processes of associating output tokens with appropriate operator nodes and of deciding which are ready for execution is known as “matching” process.

Under this paradigm there is no current operation, and each operator is free to execute when all its input tokens are available. The model is naturally concurrent, and the concurrency grain depends on the operations grain.

The data-flow model has the single-assignment property. Values are
data tokens that are carried from their producing node to the node that consumes them; there is no concept of a variable with a state that can be arbitrarily updated later. In data-flow, identifiers may be used to name these data tokens. Such identifiers are thus either undefined (not yet produced) or carry a single unique value; they cannot be updated. A node with all input data available is called “fireable”. When a node is “fireable” is ready to be run on a data-flow interpreter. Each data-flow interpreter is called “actor”. The features of a data-flow model were listed by Ackerman in its 1982 milestone paper (10). They are:

- side effects free;
- locality of effect;
- equivalence of instruction scheduling with data dependencies;
- single-assignment semantics;
- an unusual notation for iterations;
- lack of history sensitivity in procedures.

Synchronization is automatically provided by the token transport mechanism. Parallelism is exploited in data-flow architectures by allowing any actor to execute on any processor and by allowing as many enabled actors to fire as there are processors to execute them. When there are a sufficiently large number of processors, only actors that do not have the input data available are not enabled.

A key feature of the model is that the order of actor execution does not affect the result. Thus, the data-flow model naturally achieves high degrees of parallelism. Nevertheless, traditional data-flow presents three major problems when considered for large distributed (grid) environments.

- The granularity of traditional data-flow is too small for many distributed architectures, for instance related to distributed memory access time (where latencies are measured in hundreds to thousands of microseconds). The overhead of token transport and actor

28
scheduling and instantiation requires that the granularity of computation be at least hundreds of thousands, and perhaps million of instructions.

- The programming abstraction provided to programmers is quite different with respect to the traditional sequential one.

The main difference between this approach and those discussed above is that whereas a graph reducer manipulates the graph by modifying both data and the “instruction code” itself, a data-flow graph is statically defined by the program and only data is manipulated.

Data-flow based languages may be dressed up to resemble sequential imperative languages (27), particularly in case of “scientific” applications. The compilation process from high-level language to the underlying data-flow graph is quite similar to the process of expansion in graph reduction. It is equivalent to the decomposition phase of parallel implementation.

All the problems of distribution, communication and synchronization are associated with the data-flow graph and the interactions between its node operators. Although the structure of the graph is static, it will only be apparent during (or even after) execution that some sections of the graph were more active than others. Thus, a good distribution scheme is difficult to obtain without any additional information, for instance in the form of programmer annotations.

**Macro-Dataflow approaches**

The macro data-flow model extends the traditional data-flow model addressing its main problems. There are two principal differences with traditional data-flow. First, the granularity of the actors is considerably larger (indeed in this case they are named “macro” actors). This allows to achieve a good scalability when the degree of parallelism, namely the number of recruited PEs, increases. Second, some actors (76) can maintain state information between firings, providing an effective way to model side-effects and non-determinism, these actors are called “persistent” actors. Some examples of existing and widely used macro-actors implement high-level functions such as: matrix multiplication, Gaussian elimination
or image convolution instead of individual machine instructions. Macro actors can be described as follows.

**Regular actors** are similar to actors in the data-flow model. Specifically, all regular actors of a given type are functionally equivalent. A regular actor is enabled and may execute when all of its input tokens are available. It performs some computation, generating output tokens that depend only on its input tokens. It may maintain internal state information during the course of a single execution, but no state information is preserved from one execution to another; regular actors, therefore, represent pure functions.

**Persistent actors** maintain state information that is preserved from one execution to the next. Output tokens generated by a persistent actor during different executions are not necessarily the same for the same input tokens. The state corresponds to member variables (instance variables) in the object-oriented paradigm. This correspondence implies that several different actors may share the same state, (as an example with the enqueue and dequeue operations on a queue). The model guarantees that the actors that share state will be executed in mutual exclusion, that is, no two actors that share the same state will ever be executing simultaneously. (This can be modeled in stateless data-flow using a single “state” token and a non-deterministic merge operator (9)). The introduction of state means that the arcs of the program graph no longer model all dependencies in the program; there are implicit dependencies via the shared state. For example, consider the program graph fragment in Figure 2. Suppose that actors A and B share state. If the execution of A occurs first, there is a hidden dependency, based on the state, between A and B. Because of this hidden dependency, the results of the A and B operations depend not only on their arguments and the object history, but also on the order of execution.

If on the one hand the persistent macro actors approach addresses the one limitation of the traditional data-flow model, on the other hand it
makes the programming model more complicated and requires to programmers to pay more attention when programming parallel applications. In particular, the introduction of state has one very important consequence: some programs will be deterministic, and others not. Non-determinism is not necessarily bad. There are in fact many “correct” non-deterministic applications. Thus, it is the responsibility of the programmer to guarantee higher-level notions of correctness. Due to the additional complexity they introduce, several existing macro data-flow systems do not support persistent actors.

**A notable MDF approach: the Mentat framework**

Mentat is one of the most known and used macro data-flow system (75). It is an object-oriented parallel processing system for MIMD architectures developed at the University of Virginia. The computation model used in Mentat is a data-driven macro data-flow computation model based on the object-oriented paradigm. There are two primary components of Mentat: the Mentat Programming Language (MPL) and the Mentat runtime system. MPL is an object-oriented programming language based on
C++. The computational grain of the macro data-flow block is the Mentat class instance, which consists of contained objects (local and member variables), their procedures, and a thread of control. Programmers are responsible for identifying those object classes that are of sufficient computational complexity to allow efficient parallel execution. Instances of Mentat classes are used just like ordinary C++ classes. The data and control dependencies between Mentat class instances involved in invocation, communication, and synchronization are automatically detected and managed by the compiler and run-time system without programmer intervention.

**MPL** is an extended C++ designed for developing parallel applications by providing parallelism encapsulation. Parallelism encapsulation takes two forms, intra-object encapsulation and inter-object encapsulation. In intra-object encapsulation of parallelism, callers of a Mentat object member function are unaware of whether the implementation of the member function is sequential or parallel, i.e., whether its program graph is a single node or a parallel graph. In inter-object encapsulation of parallelism, programmers of code fragments (e.g., a Mentat object member function) need not concern themselves with the parallel execution opportunities between the different Mentat object member functions they invoke. The basic idea in the MPL is to allow the programmer to specify those C++ classes that are of sufficient computational complexity to warrant parallel execution. Programmers can select which classes should be executed in parallel using a `mentat` keyword in the class definition. Instances of Mentat classes are called Mentat objects. Mentat classes are very similar to C++ class instance but with some minor differences (described below). The compiler generates code to construct and execute data dependency graphs in which the nodes are Mentat object member function invocations, and the arcs are the data dependencies found in the program. Thus, it transparently generates inter-object parallelism encapsulation. All the communications and synchronizations are managed by the compiler. MPL is built around four main extensions to the C++ language. The extensions are Mentat classes, Mentat object instantiation, the
A key feature of Mentat is the transparent encapsulation of parallelism within and between Mentat object member function invocations. The hiding of whether a member function implementation is sequential or parallel is called intra-object parallelism encapsulation. Similarly, the inter-object parallelism encapsulation consists in the exploitation of parallelism opportunities between Mentat object member function invocations in a transparent way to the programmer. Intra-object parallelism encapsulation and inter-object parallelism encapsulation can be combined. Indeed, inter-object parallelism encapsulation within a member function implementation is intra-object parallelism encapsulation as far as the caller of that member function is concerned. Thus, multiple levels of parallelism encapsulation are possible, each level hidden from the level above.

Not all class objects should be Mentat objects. In particular, objects that do not have a sufficiently high communication ratio, i.e., whose object operations are not sufficiently computationally complex, should not be Mentat objects. The programmer defines a Mentat class by using the keyword `mentat` in the class definition. The programmer may further specify whether the class is persistent, sequential, or regular. Persistent and sequential objects maintain state information between member function invocations, while regular objects do not. Thus, regular object member functions are pure functions. Because they are pure functions, the system is free to instantiate new instances of regular classes at will. Regular classes may have local variables much as procedures do, and may maintain state information for the duration of a function invocation. The programmer binds Mentat variables to persistent Mentat objects using two reserved member functions for all Mentat class objects: `create()` and `bind()`. The `create()` call tells the system to instantiate a new instance of the appropriate class whereas the `bind()` function binds Mentat variables to an already existing instance. The member function `destroy()` destroys the named persistent Mentat object. The return-to-future function (`rtf()`) is the Mentat analog to the return of C. Its purpose is to allow Mentat member functions to return a value to the successor nodes in the macro data-flow graph in which the member function appears. The select/accept state-
ments of Mentat is a guarded statement that derives directly from the ADA one. Guarded statements permit the programmer to specify a set of entry points to a monitor-like construct. The guards are boolean expressions based on local variables and constants. A guard is assigned to each possible entry point. If the guard evaluates to true, its corresponding entry point is a candidate for execution. The rules vary for determining which of the candidates is chosen to execute. It is common to specify in the language that it is chosen at random. This can result in some entry points never being chosen. There are two types of guard-actions supported by Mentat: accepts, tests, and non-entries. Accept is similar to the accept of ADA. Tests are used to test whether a particular member function has any outstanding calls that satisfy the guard. When a test guard-action is selected, no parameters are consumed. In Mentat there is no “else” clause as in ADA. However, using the priority options, the programmer can simulate one by specifying that the clause is a non-entry statement and giving the guard-statement a lower priority than all other guard-statements. Then, if none of the other guards evaluates to true, it will be chosen. The priority of the guard-statement determines the order of evaluation of the guards. It can be set either implicitly or explicitly. The token priority determines which call within a single guard-statement priority level will be accepted next. The token priority is the maximum of the priorities of the incoming tokens. Within a single token priority level, tokens are ordered by arrival time.

To give an idea of the programming model in Figure 3 we report a simple Mentat program. The program computes recursively the Fibonacci number. It is composed by two classes, the first one recursively computes the Fibonacci number exploiting the second one for computing the sum of partial results. Clearly, in this case the efficiency is low because the amount of computation done by the macro actors computing the mentat object adder class is very small.

Unfortunately, there are a number of issues and limitation that MPL programmers must be aware of that can lead to unpredictable program behavior, related both to Mentat implementation and model. Among the others:
Figure 3: Fibonacci computation with Mentat

- The use of static member variables for Mentat classes is not allowed. Since static members are global to all instances of a class, they would require some form of shared memory between the instances of the object.

- Mentat classes cannot have any member variables in their public definition. If data members were allowed in the public section, users of that object would need to be able to access that data as if it were local. If the programmer wants the effect of public member variables, appropriate member functions can be defined.

- Programmers cannot assume that pointers to instances of Mentat classes point to the member data for the instance.

- Mentat classes cannot have any friend classes or functions. This restriction is necessary because of the independent address space of Mentat classes.

- It must be possible to determine the length of all actual parame-
ters of Mentat member functions, either at compile-time or at run-time. This restriction follows from the need to know how many bytes of the argument to send. Furthermore, each actual parameter of a Mentat member function must occupy a contiguous region of memory in order to facilitate the marshaling of arguments.

- Mentat object member function parameter passing is call-by-value. All parameters are physically copied to the destination object. Similarly, return values are by-value.

- if a Mentat member function returns a pointer, the programmer must explicitly delete the reference when the function is finished using the value.

- semantic equivalence to the sequential program is not guaranteed when persistent objects are used. This is trivially true for programs that have select/accept statements; there are no serial equivalents.

Summarizing Pros and Cons  Data-flow model is inherently parallel, it represents each computation as a graph made by operators and instructions where each node can be potentially executed in parallel. This model permit to programmers to express parallel applications in a very abstract way, indeed programmers are not required to deal with low-level issues related to the running architecture. The main problem of Data-flow model is the fine-granularity of instruction that prevent its exploitation in most distributed architectures and in large grid environments. This limitation led to the development of the macro data-flow model (MDF). The MDF model allows programmers to define code fragment in place of instruction as nodes in DF graph. Unfortunately, such additions impair the high-level abstraction, like in case of the implicit models. Hence, programmers have both to deal with data/application decomposition and to assure semantic equivalence with respect to the sequential program, especially when exploiting persistent actors.
2.2.4 Low-level explicit models: MPI and OpenMP

The low-level approaches provide to the programmers a programming metaphor where parallelism is represented by means of primitives in the form of special-purpose directives or function calls. Most parallel primitives are related to process synchronization, communication or task partitioning. The total amount of computational cost for executing these primitive is considered as parallelization overhead. The advantage of explicit parallel programming is the absolute programmer control over the parallel execution. A very skilled parallel programmer takes advantage of explicit parallelism to produce very efficient code. However, programming with explicit parallelism is often difficult and error prone, because of the extra work involved in planning the task division and synchronization of concurrent processes. In this section we report two of the main approaches to low-level parallel computing: MPI and OpenMP. The former is suitable for distributed architectures whereas the latter is appropriate for multicore and multiprocessor architectures.

MPI

MPI is a message-passing library, proposed as a standard by a broadly based committee of vendors, implementors, and programmers. MPI was designed for high performance on both massively parallel machines and on workstation clusters. The Message Passing Interface is meant to provide essential synchronization and communication functionality between a set of processes, mapped into different computer instances, in a language independent way, plus a few features that are language specific. The programming metaphor of MPI is based on the “process” concept. An MPI program consists of autonomous processes, executing their own code, in a Multiple Instructions, Multiple Data stream (MIMD) style, i.e. Multiple autonomous processors simultaneously executing different instructions on different data. Distributed systems are generally recognized to be MIMD architectures. The processes communicate exploiting MPI communication primitives. Typically, each process executes in its own address space, although shared-memory implementations of MPI are possi-
MPI does not specify the execution model for each process. A process can be sequential, or can be multi-threaded, with threads possibly executing concurrently. The intended interaction of MPI with threads is that concurrent threads be all allowed to execute MPI calls, and calls be reentrant; a blocking MPI call blocks only the invoking thread, allowing the scheduling of another thread. MPI does not provide mechanisms to specify the initial allocation of processes to an MPI computation and their binding to physical processors. MPI mapping of processes on PEs happens at run-time, through the agent that starts the MPI program, normally called `mpirun` or `mpiexec`.

MPI primitives include, but are not limited to, point-to-point rendezvous type send/receive operations, combining partial results of computations (gathering and reduction operations), choosing between a Cartesian or graph-like logical process topology, exchanging data between process pairs (send and receive operations), synchronizing nodes (barrier operation) as well as obtaining network-related information such as the number of processes in the computing session, current processor identity that a process is mapped to, neighboring processes accessible in a logical topology, and so on. Point-to-point operations come in synchronous, asynchronous, buffered, and ready forms in order to allow both relatively stronger and weaker semantics for the synchronization aspects of a rendezvous-send. Many outstanding operations are possible in asynchronous mode, in most implementations. Figure 4 reports the main classes of MPI primitives. There are two versions of the MPI standard that are currently popular: version 1.2 (also called MPI-1), which empha-
sizes message passing and has a static run-time environment, and MPI-2.1 (MPI-2), which includes features such as parallel I/O, dynamic process management and remote memory operations. Figure 5 show a simple Hello World MPI program. It defines two roles: master and slave. The master ask slaves to process the “Hello word” string and then return it. The master eventually print on screen the string received by slaves. The roles are specified by means of the MPI process id. The process number 0 is the master whereas the others are slaves.

As shown in Figure 5 MPI Hello World programmer is in charge of:

- initialize MPI
- find the available resources and manage them
- implement by hands a way to differentiate the master and the slaves
- prepare the data the master sends
- send the data to slaves
- make the slaves receive the data
- implement the slave data processing
- prepare the data the slaves send
- make the master receive the data, collecting it and processing it
- finalize MPI

Furthermore, he must allocate memory buffers, manage fault(s) and distribute data by hands. It is easy to understand that implement a complex application with MPI is a very difficult and error prone task because MPI programmers must manage all the aspects of the application parallelization. On one hand, it guarantees maximum programming flexibility, but on the other hand such a freedom is paid in terms of programming complexity.
```c
#include <mpi.h>
#include <stdio.h>
#include <string.h>
#define BUFSIZE 128
#define TAG 0

int main(int argc, char *argv[]) {
    char idstr[32], buff[BUFSIZE];
    int numprocs, myid, i;
    MPI_Status stat;
    /* MPI programs start with MPI_Init; all ’N’ processes exist thereafter */
    MPI_Init(&argc,&argv);
    /* find out the number of available PEs */
    MPI_Comm_size(MPI_COMM_WORLD,&numprocs);
    /* and this processes’ rank is */
    MPI_Comm_rank(MPI_COMM_WORLD,&myid);
    /* At this point, all the programs are running equivalently, the rank is
    used to distinguish the roles of the programs in the SPMD model */
    if(myid == 0) {
        /* rank 0 process sent a string to all the other processes */
        for(i=1;i<numprocs;i++) {
            sprintf(buff, "Hello %d! ", i);
            MPI_Send(buff, BUFSIZE, MPI_CHAR, i, TAG, MPI_COMM_WORLD);
        }
    /* rank 0 process sent a string to all the other processes */
    for(i=1;i<numprocs;i++) {
        MPI_Recv(buff, BUFSIZE, MPI_CHAR, i, TAG, MPI_COMM_WORLD, &stat);
        printf("%d: %s\n", myid, buff);
    }
    } else {
        /* receive from rank 0: */
        MPI_Recv(buff, BUFSIZE, MPI_CHAR, 0, TAG, MPI_COMM_WORLD, &stat);
        sprintf(idstr, "Processor %d ", myid);
        strcat(buff, idstr);
        strcat(buff, "reporting for duty\n");
    /* send to rank 0: */
        MPI_Send(buff, BUFSIZE, MPI_CHAR, 0, TAG, MPI_COMM_WORLD);
    }
    /* MPI Programs end with MPI Finalize */
    MPI_Finalize();
    return 0;
}
```

**Figure 5:** Hello Word example implemented using MPI
Like MPI, OpenMP (Open Multi-Processing) is a specification defined by a group of major computer hardware and software vendors for multi-platform multiprocessing programming. It consists of a set of compiler directives, library routines, and environment variables that influence runtime behavior. Unlike MPI, it is mainly targeted to shared memory multiprocessing. Indeed, it is used in conjunction with MPI on distributed architectures made of multicore/multiprocessor machines. OpenMP uses multiple, parallel threads to accomplish parallelism. A thread is a single sequential flow of control within a program. OpenMP uses a directive-based method to tell explicitly to the compiler how to distribute programs across parallel threads.

The core elements of OpenMP are the constructs for thread creation, workload distribution (work sharing), data environment management, thread synchronization, user level run-time routines and environment variables. OpenMP programmers exploit such constructs to manage all the aspects of application parallelization. Figure 6 shows the classes of existing OpenMP language extensions.

Even if the OpenMP approach to parallel programming has to be considered as a low-level one, OpenMP code is more straightforward than

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**Figure 6: OpenMP language extensions**

<table>
<thead>
<tr>
<th>Parallel Control Structures</th>
<th>Work Sharing</th>
<th>Data Environment</th>
<th>Synchronization</th>
<th>Runtime Functions, Env. Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Parallel Directive</strong></td>
<td><strong>Distributes Work Among Threads</strong></td>
<td><strong>Scopes Variables</strong></td>
<td><strong>Coordinates Thread Execution</strong></td>
<td><strong>Runtime Environment</strong></td>
</tr>
<tr>
<td><code>parallel</code> directives</td>
<td><code>do/parallel do</code> and <code>section</code> directives</td>
<td><code>shared and private</code> clauses</td>
<td><code>critical and atomic</code> directives</td>
<td><code>omp_set_num_threads()</code></td>
</tr>
<tr>
<td><code>omp_get_thread_num()</code></td>
<td><code>omp_num_threads</code></td>
<td><code>omp_get_thread_num()</code></td>
<td><code>omp_set_schedule</code></td>
<td></td>
</tr>
</tbody>
</table>

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**OpenMP**

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The core elements of OpenMP are the constructs for thread creation, workload distribution (work sharing), data environment management, thread synchronization, user level run-time routines and environment variables. OpenMP programmers exploit such constructs to manage all the aspects of application parallelization. Figure 6 shows the classes of existing OpenMP language extensions.

Even if the OpenMP approach to parallel programming has to be considered as a low-level one, OpenMP code is more straightforward than
MPI code. This is mainly due to the memory model indeed, relying on a shared memory model. The OpenMP application does not need to deal with message passing hence data are not directly split and divided among PEs but handled through compiler directives.

An OpenMP program is a C++ or Fortran program with OpenMP pragma statements/directives placed at appropriate points. The pragma statement directs the compiler how to process the block of code that follows the pragma. An OpenMP-enabled compiler recognizes the pragma directives and produces a parallelized executable suitable for running on a shared-memory machine. In C/C++, an OpenMP directive has the general form:

```
#pragma omp directive – name [clause,...] newline
```

The #pragma omp directive tags a block for parallel or various types of work sharing execution, variable scoping and synchronization considerations. One or more clauses are optional and may be in any order. The clauses are used to explicitly define the scoping of enclosed variables. In OpenMP there are two main constructs:

- A parallel region is a block of code that will be executed by multiple threads. This is the fundamental parallel construct.

- A work-sharing construct divides the execution of the enclosed code region among the members of the team that encounter it. Work-sharing constructs do not launch new threads. These constructs are identified by DO/FOR, SECTIONS and WORKSHARE (Fortran only) directives.

Since OpenMP is a shared memory programming model, most variables in OpenMP code are visible to all threads by default. However, sometimes private variables are necessary to avoid a race condition and there is a need to pass values between the sequential part and the parallel region. Another important issue is the synchronization and scheduling of the threads. These are managed through clauses appended to the OpenMP directive. Thus, the different types of clauses are Data Scoping, Synchronization and Scheduling clauses.
int main (int argc, char *argv[]) {
    int nthreads, tid, i, chunk;
    float a[N], b[N], c[N];

    /* Some initializations */
    for (i=0; i < N; i++) a[i] = b[i] = i * 1.0;
    chunk = CHUNKSIZE;

    #pragma omp parallel shared(a,b,c,nthreads,chunk) private(i,tid)
    {
        tid = omp_get_thread_num();
        if (tid == 0) {
            nthreads = omp_get_num_threads();
            printf("Number of threads = %d\n", nthreads);
        }
        printf("Thread %d starting...\n",tid);
        #pragma omp for schedule(dynamic,chunk)
        for (i=0; i<N; i++) {
            c[i] = a[i] + b[i];
            printf("Thread %d: c[%d]= %f\n",tid,i,c[i]);
        }
    } /* end of parallel section */
}

Figure 7: Factorial example in OpenMP

In Figure 7 we report an OpenMP example program. The example uses two pragma directives. The outer #pragma omp parallel tags a block for parallel execution. The shared() clause specifies common variables, and private() specifies the variables restricted to exclusive use by a process. The inner #pragma omp for schedule directive specifies distribution across threads. The threads share the variables a, b, c and chunk; the iteration variable i is private in each thread. The expression tells the compiler to perform parallel execution of the for-loop and to split the iteration space into blocks of size chunk.

The current version of OpenMP presents some issues, some related to the implementation and others related to the model. For instance a reliable error handling, fine-grained mechanisms for controlling thread-processor mapping or synchronization among a subset of threads. The model related issues, clearly more difficult to overcome include inefficient parallelism exploitation in distributed-memory platforms and a limited scalability that actually depends by memory architecture.
Summarizing Pros and Cons  Low-level approaches allow programmers to control all the aspects of parallel applications and their execution. Exploiting low-level approaches skilled programmers can implement very efficient parallel applications. The freedom and efficiency allowed by the model are paid in terms of expressiveness and ease of use. Indeed, programmers have to manage “by hand” all the issues related to data and program decomposition, fault tolerance, load balancing and communications.

2.2.5 Other notable approaches

Other two noteworthy explicit parallel approaches are Cilk and High Performance Fortran.

The first one is quite similar to OpenMP, indeed it consists in an enriched version of C language, it requires that the computing resources share the main memory hence can be used for programming parallel applications running in multiprocessor machines but not in distributed architecture like clusters. It enriches GNU C with a few Cilk-specific keywords. Using them programmers expose the parallelism identifying elements that can safely be executed in parallel. Using such information the run-time environment, in particular the scheduler, decides during execution how to distribute the work among processors. The first Cilk keyword is \texttt{cilk}, which identifies a function written in Cilk. Since Cilk procedures can call C procedures directly, but C procedures cannot directly call or spawn Cilk procedures, this keyword is needed to distinguish Cilk code from C code. Other keywords are: \texttt{spawn}, \texttt{sync}, \texttt{inlet} and \texttt{abort}. The first two keywords are all Cilk programmers have to use to start using the parallel features of Cilk: \texttt{spawn} indicates that the procedure call it modifies can safely operate in parallel with other executing code. Note that from the point of view of the scheduler it is not mandatory to run this procedure in parallel; the keyword only inform the scheduler that it can run the procedure in parallel. \texttt{sync} indicates that execution of the current procedure cannot proceed until all previously spawned procedures have completed and returned their results to the parent frame. The two remaining Cilk keywords are slightly more advanced, and concern the use
of inlets. Typically, when a Cilk procedure is spawned, it can only return its results to the parent procedure by putting those results in a variable in the parent’s frame, as we assigned the results of our spawned procedure calls in the example to x and y. The alternative is to use an inlet. An inlet is a function internal to a Cilk procedure that handles the results of a spawned procedure call as they return. One major reason to use inlets is that all the inlets of a procedure are guaranteed to operate atomically with regards to each other and to the parent procedure, thus avoiding the bugs that could occur if the multiple returning procedures tried to update the same variables in the parent frame at the same time. The abort keyword can only be used inside an inlet; it tells the scheduler that any other procedures that have been spawned off by the parent procedure can safely be aborted.

High Performance Fortran is an extension of Fortran 90 defined by the high performance fortran forum with constructs that support data-parallel computations. It consists in a portable language for data-parallel computations. HPF uses a data parallel model of computation to support spreading the work of a single array computation over multiple processors. This allows efficient implementation on both SIMD and MIMD style architectures. It provides a number of basic data parallel functions as built-in array operators and intrinsic functions. It also provides constructs, such as the where and the forall, which assist in programming more complex data parallel functions. The simplest data parallel operations are the elementwise operations. For any base operation on a data type, programmers can extend that operation to an array operation. For binary (and higher degree) operations, the arrays must have the same shape. The result of the operation is another array of that shape, in which the elements are defined by the elementwise extension of the base operation. A more advanced set of operations operate on an entire array to produce a single answer, they implement a behavior generally known as reduction. Reduction can be defined for any associative, binary operation that produces a result of the same element type by successively accumulating the results of applying that operation to elements of the array. Commonly used operations include arithmetic operators like addition,
multiplication, maximum, and minimum and boolean operators. As an example, HPF programmers can define reduction with addition, usually called sum reduction, over any array whose element type can be added.

2.2.6 Structured approach

Highly abstract approaches and low-level approaches represent the two extremes in parallel programming models. The formers completely automate the aspects of parallelization, namely do not ask programmers (at least in their “pure” version) to give any information about application, like data distribution and synchronization, communication mechanisms, executing environment or code sequences to run in parallel. The latter, opposite, approaches do not automate anything and ask programmers to deal, almost entirely, with the application parallelization aspects.

As we outlined in previous sections, several researchers have tried to address the limitation of these approaches enriching them with additional features. Some other work was done trying to conceive alternative models. In particular, since the nineties, several research groups have proposed the structured parallel programming environments (SPPE). Since the structured parallel programming model was conceived, several works have been done about it, also from a foundational point of view (20), (11), (50). Programming environments relying on this paradigm (i.e. (101)) ask programmers to explicitly deal with the qualitative aspects of parallelism exploitation, namely the application structure and problem decomposition strategies. All the low-level parallelism exploitation related aspects like communication, synchronization, mapping and scheduling are managed by compiler tools and run-time support.

The structured way is driven by two observations: that there are some things people do better than compilers, and that there are some things that compilers do better than people. Rather than have either do the complete job, it exploits the comparative advantages of each. Indeed the management of tens to thousands of asynchronous tasks, where timing-dependent errors are quite common, is beyond the capacity of most programmers whereas compilers are very good at ensuring that events hap-
pen in the right order and can more readily and correctly manage com-
munication and synchronization than programmers. On the other hand, 
data decomposition strategies and computational grain can be successful 
managed by programmers but not efficiently by compilers.

The environments following this way are those based on the algorithmic 
skeleton concept. A skeleton, is a known and widely used pattern of 
parallelism exploitation originally conceived by Cole (50) and later on by 
different research groups to design high-performance structured parallel 
programming environments.

Basically, structured parallel programming systems allow a parallel 
application to be coded by properly composing a set of basic parallel 
skeletons. These basic skeletons usually include skeletons modeling em-
barrassingly parallel computations (farms), computations structured in 
stages (pipelines) as well as common data parallel computation patterns 
(map/forall, reduce, scan). Each skeleton is parametric; in particular, 
it accepts as a parameter the kind of computation to be performed ac-
cording to parallelism exploitation pattern it models. As an example, a 
farm skeleton takes as a parameter the worker, i.e. the computation to be 
performed on the single input task (data item). As a further example, a 
pipeline takes as parameters the pipeline stages. Such parameters may be 
either parameters modeling sequential portions of code (sequential skele-
tons) or even other skeletons, in turn. Therefore, a farm skeleton may take 
as a worker a two stage pipeline. The composition of the two expresses 
embarrassingly parallel computations where each input task (data item) 
is processed by two stages. Parallelism is exploited both by using differ-
et resources to compute independent input tasks and by using different 
resources to compute the first and the second stage onto a single input 
task.

A skeleton (in its original formulation) is formally an higher order 
function taking one or more other skeletons or portions of sequential code 
as parameters, and modeling a parallel computation out of them. Cole’s 
skeletons represent parallelism exploitation patterns that can be used (in-
stanced) to model common parallel applications. Later, different authors 
figure out that skeletons can be used as constructs of an explicitly paral-
lel programming language, actually as the only way to express parallel computations in these languages (30; 64). Recently, the skeleton concept evolved, and became the coordination layer of structured parallel programming environments ((29; 32; 113)). In any case, a skeleton can be considered as an abstraction modeling a common, reusable parallelism exploitation pattern. Skeletons can be provided to the programmer either as language constructs (29; 30; 32) or as libraries (12; 55; 62; 88). Usually, the set of skeletons includes both data-parallel and task parallel patterns.

**Traditional skeleton approaches**

From the nineties, several research groups proposed or currently propose programming environments supporting parallel computations based on the algorithmic skeleton concept. They are implemented as frameworks, languages or libraries. Among the others, we mention Kuchen’s C++ MPI skeleton library (88), Serot’s SKiPPER environment, $P^3L$, Lithium, a first version of muskel and JJPF. In particular, the last one, JJPF, represents our approach to traditional SPPE. In the rest of this section we present a more detailed description about the programming model of $P^3L$, muskel and JJPF to describe the “concept behind” SPPE models.

We developed this last one, whereas all the other skeleton environments presented in this section have been developed by the Parallel and Distributed Architecture Group, part of the Department of Computer Science at University of Pisa. This group has a deep background on skeleton environment, indeed the group began to work in this field from the very beginning the skeleton model were conceived. We collaborated with several researchers belonging to this group, also for the conception and the design of the results presented in this thesis.

$P^3L$ is a high-level structured explicitly parallel language developed in the nineties (31). Using $P^3L$ parallelism can be expressed only by means of a restricted set of parallel constructs each corresponding to a specific parallel form. Sequential parts are expressed by using an existing language also called the host sequential language of $P^3L$. Being a SPPE its constructs can be hierarchically composed to express more complex
parallel forms. This compositional property relies on the semantics associated with the various $P^3L$ constructs and their compositions. In fact, each of them can be thought of as a data-flow module. In $P^3L$ each module computes in parallel or sequentially a function on a given stream of input data and produces an output stream of results. The lengths of both the streams are identical and the ordering is preserved, i.e.

$$[in_1, ..., in_n] \rightarrow M \rightarrow [out_1, ..., out_n]$$

where $M$ is the data-flow module corresponding to a generic $P^3L$ construct $[in_1, ..., in_n]$ is the input stream, $[out_1, ..., out_n]$ is the output stream, $n$ is the length of both the streams and every output data item $out_i$ is obtained by applying the function computed by $M$ on the input data item $in_i$. The types of the input and the output interface of each $P^3L$ construct i.e. the types of every $in_i$ and every $out_i$ have to be declared statically. Actually the compiler performs type checking on these interfaces when the $P^3L$ constructs are to be composed. Another feature of $P^3L$ is its interface with the host sequential language. The interface has been designed to make easier portability between different host languages. In fact, sequential parts are completely encapsulated into the constructs of $P^3L$. Parameter passing between $P^3L$ constructs are handled by linguistic constructs that are external to the specific host sequential language while the data types that can be used to define the interface of the $P^3L$ constructs are a fixed subset of those usually available in the most common languages. The first $P^3L$ compiler adopted as host sequential language C and C++. The constructs included since the first $P^3L$ compiler were

- The **farm** construct which models processor farm parallelism. In this form of parallelism a set of identical workers execute in parallel the independent tasks that come from an input stream and produce an output stream of results.

- The **map** construct which models data parallel computations. In this form of parallelism each input data item from an input stream is decomposed into a set of partitions and assigned to identical and parallel workers. The workers do not need to exchange data to perform their data parallel computations. The results produced by the
workers are recomposed to make up a new data item of an output stream of results.

- The **pipe** construct which models pipeline parallelism. In this form of parallelism a set of stages execute serially over a stream of input data producing an output stream of results.

- The **loop** construct which models computations where for each input data item a loop body has to be iteratively executed until a given condition is reached and an output data item is produced.

- The **sequential** construct which corresponds to a sequential process that for each data item coming from an input stream produces a new data item of an output stream.

The sequential constructs constitute the leaves of the hierarchical composition because the computations performed by them have to be expressed in terms of the host sequential language.

**muskel** (58) is a full Java framework, providing programmers with structured ways of expressing parallel programs. The muskel environment represents a sensible evolution of the Lithium one (12). It inherits from Lithium the **normal form** (63) and macro data-flow (57 101) implementation techniques as well as the general structure of the run-time support.

**Normalization** consists in transforming the original skeleton tree (or composition) into a program that is basically a task farm with sequential workers (18). Such optimization basically substitute skeleton subtrees by skeleton subtrees providing a better performance and efficiency in the target machine resource usage than the original skeleton tree. Previous results demonstrated that full stream parallel skeleton subtrees can be collapsed to a single farm skeleton with a (possibly huge) sequential worker leading to a service time which is equal or even better that the service time of the uncollapsed skeleton tree (50).

The **muskel** macro data-flow run-time support consists in deriving a graph of macro data-flow blocks from skeleton trees and dispatching
them to computational resources running macro-actors.

**muskel** adds to Lithium a limited form of resource discovery and fault tolerance features as well as the whole Application Manager concept.

The Application Manager (AM) is an entity that takes care of assuring that the application non-functional requirement were satisfied. The requirements are specified by programmers in a performance contract. The AM actively observes the application behavior and in case of faults or performance contract violations it reacts aiming to fix the problem, as an example, in case of a computational resource fault it recruits a new resource in the computation.

Using **muskel** a programmer can implement parallel programs that match the task farm or the pipeline parallelism exploitation patterns as well as arbitrary composition of the two. Despite the limited amount of patterns supported, however, a large range of applications can be programmed, for instance all embarrassingly parallel applications, parameter sweeping applications and multistage applications.

A task farm computation can be defined just using a Farm object. The Farm constructor takes a parameter representing the computation performed by the farm workers. This computation can be either a sequential computation or another parallelism exploitation pattern (another Farm or a Pipeline one). A pipeline computation can be defined using a Pipeline object. The Pipeline constructor takes two parameters that can either be sequential computation objects or in turn parallel exploitation patterns. Pipelines with more stages can be obtained composing several Pipeline objects. Then the programmer has to add an Application Manager to the application code, and he must also specify the performance contract he pretends to be respected on the target architecture. This is done instantiating an application manager and specifying a performance contract. **muskel** supports two different kinds of contracts. The first one requires a constant parallelism degree, that is, it requires that a constant number of processing elements are dedicated to the parallel execution of our parallel program. The second one requires that a given throughput is maintained in terms of task processed per unit time. Both of these kinds of contracts can be specified before the computation of the parallel **muskel**
program actually starts and can be changed during the program execution. The management of the parallel computation in such a way that the contracts are satisfied is completely handled by an independent execution flow. Therefore, the submission of a new performance contract to the application manager immediately triggers all those (possibly additional) activities needed to satisfy the contract. The possibility to change the performance contracts during the execution of the parallel applications allows the programmer to implement some kind of application dependent dynamic execution strategy. Once the program has been specified along with its performance contract the programmer must supply the list/stream of tasks to be computed. When all the elements belonging to the list/stream have been processed, the parallel execution of the program is terminated and the relative results can be fetched.

During the computation of the parallel program the muskel run-time automatically discovers available processing elements. In case there are no enough resources to satisfy the contract, an error is signaled to the programmer.

As we stated before, in case of faults the Application Manager recruits new resources among the available ones to substitute the faulty one. In case the application manager recognizes that the performance contract specified by the programmer cannot be satisfied, it raises an Exception. Being any task to be computed a fireable macro data flow instruction, it is completely independent of any other task needed to compute the parallel application. Therefore, it can be scheduled on any one of the available resources. However, the normal form concept implemented in muskel, only generates fully independent macro data flow instructions. That is, no result of an instruction is needed to compute another instruction. In this case, most of the scheduling problems we just mentioned disappear.

**JJPF** is a parallel programming framework built on top of plain Java that can run stream parallel applications on several parallel/distributed architectures ranging from tightly coupled workstation clusters to generic workstation networks and grids. In a sense, JJPF represents our approach to old-fashioned structured parallel programming environments. It di-
directly inherits from the early versions of Lithium and muskel \cite{12}. Both Lithium and muskel exploit plain RMI Java technology to distribute computations across nodes, and rely on NFS (the network file system) to distribute the application code to the remote processing elements. JJPF, instead, is fully implemented on top of JINI and Java and relies on the Jini Extensible Remote Invocation (JERI) mechanism to distribute code across the remote processing nodes involved in stream parallel application computation. JJPF exploits the stream parallel structure of the application in such a way that several distinct goals can be achieved:

- *load balancing is achieved* across the computing elements participating in the computation

- processing elements available to participate to the computation of stream parallel application are *automatically discovered and recruited* exploiting standard Jini mechanisms

- *faulty processing elements are automatically substituted* by fresh ones (if any) in a seamless and automatic way. Therefore, the stream parallel applications computations resist to both node and network faults. Programmers do not need to add a single line of code in his application to deal with faulty nodes/network, nor it has to take any other kind of action to get advantage of this feature.

JJPF has been tested using both synthetic and real applications, on both production workstation networks and on clusters, with very nice and encouraging results. JJPF has been designed to provide programmers with an environment supporting the execution of stream parallel applications on a network of workstations, exploiting plain Java technology. Overall JJPF provides a distributed server providing a stream parallel application computation service. Programmers must write their applications in such a way they just exploit an arbitrary composition of task farm and pipeline patterns. Task farm only applications are directly executed by the distributed server, while applications exploiting composition of task farm and pipeline patterns are first processed to get their *normal form*. A distributed environment that exploits task parallel computations, permits to
Figure 8: Simplified state diagram for the generic JJPF client (left) and service (right)

implement different applications in really different applicative and hardware contexts. JJPF is based on a master-worker architecture. JJPF defines two entities: “client”, that is the application code (the master), and “service”, that consists in distributed server instances (the workers) that actually compute results out of input task data to execute client program. Figure 8 sketches the structure of the two components. The client component basically recruits available services and forks a control thread for each one of them. The control thread, in turn, fetches uncomputed task items from the task vector, delivers them to the remote service and retrieves the computed results, storing them to the result repository. Low-level activities, like resource recruiting, program deployment and data transfer are performed directly by the framework exploiting the JINI technology (4). The key concept in JJPF is that service discovery is automatically performed in the client run time support. Not a single line of code dealing with service discovery or recruiting is to be provided by application programmers. JJPF achieves automatic load balancing among the recruited services, due to the scheduling adopted in the control threads managing the remote services. Furthermore, it handles faults in service nodes automatically taking care of the tasks assigned to a service node in such a way that in case the node does not respond any more they can be rescheduled to other service nodes. This is only possible because of the
kind of parallel applications that are supported in JJPF, that is stream parallel computations. In this case, there are natural descheduling points that can be chosen to restart the computation of one of the input tasks, in case of failure of a service node. JJPF has demonstrated good scalability both in embarrassingly parallel application and in more “problematic” applications.

2.3 Open issues in structured approaches

Despite being around since long time and despite the progress made in skeletal system design and implementation, the skeleton systems did not take off as expected. Nowadays, the skeleton system usage is actually restricted to small communities grown around the teams that develop the skeleton systems. Cole focused very well the problem in his manifesto (51). Here he stated four principles that have to be tackled in skeletal systems to make them effective and successful:

I) Propagate the concept with minimal conceptual disruption  It means that skeletons must be provided within existing programming environments without actually requiring the programmers to learn entirely new programming languages. In order to make them widely used by practitioners they should not require further conceptual baggage.

II) Integrate ad-hoc parallelism  Many parallel applications are not obviously expressible as instances of skeletons. Some have phases that require the use of less structured interaction primitives. For example, Cannon’s well-known matrix multiplication algorithm (90) invokes an initial step in which matrices are skewed across processes in a manner which is not efficiently expressible in many skeletal systems. It is unrealistic to assume that skeletons can provide all the parallelism we need. We must construct our systems to allow the integration of skeletal and ad-hoc parallelism in a well-defined way.
III) Accommodate diversity

All the existing skeleton systems have a common core of simple skeletons and a variety of more exotic forms. When described informally, the core operations are straightforward. Instead, precise specification reveals variations in semantics that reflect the ways skeletons are applied in real algorithms. The result is that some algorithms, which intuitively seem to represent an instance of a skeleton, cannot be expressed in certain systems because of constraints imposed by the specification. Hence, skeletal systems should provide mechanisms to specialize skeletons, in all those cases where specialization does not radically change the nature of the skeleton, and consequently the nature of the implementation.

IV) Show the pay-back

A new technology will only gain acceptance if it can be demonstrated that adoption offers some improvement over the status quo. The structural knowledge embedded in skeletons should allow optimization within and across uses that would not be realistically achievable by hand, i.e. demonstrate that the effort required to adopt a skeletal system is immediately rewarded by some kind of concrete results: shorter design and implementation time of applications, increased efficiency, increased machine independence of the application code, etc.

The second and the third points are specifically technical whereas the first and the last one are actually a kind of “advertising” ones, in a sense. All these points, however, have impacts on both the way the skeleton systems are designed and on the way they are implemented. The Cole’s analysis is not the only one, [19] extends it adding some other features a skeleton environment have to address to be suitable for the computational grids. In particular, the authors present three more requirements for Skeletal systems:

V) Support code reuse

that is allow programmers to reuse with minimal effort existing sequential code;
VI) Handle heterogeneity  i.e. implement skeletons in such a way skeleton programs can be run on clusters/networks/grids hosting heterogeneous computing resources (different processors, different operating systems, different memory/disk configurations, etc.);

VII) Handle dynamicity  i.e. implement in the skeleton support mechanisms and policies suitable to handle typical dynamic situations, such as those arising when non-dedicated processing elements are used (e.g. peaks of load that impair load balancing strategies) or from sudden unavailability of processing elements (e.g. network faults, node reboot).

Summarizing, the next generation of Skeletal Systems, that drawing a parallel with web programming model we can refer as “Skeletons 2.0”, have to integrate ad-hoc parallelism and provide mechanisms to specialize skeletons in order to express customized form of parallel exploitation. They have to support code reuse, handle heterogeneity and dynamicity in order to be exploited in grid environments. Moreover, such features must be provided with minimal conceptual disruption, hence without requiring the programmers to learn entirely new programming languages or environments but integrating “Skeletons 2.0” principles inside the existing programming tools, possibly without changing their programming abstraction.

Some Skeletal systems have addressed the “Skeletons 2.0” principles to different degrees in different combinations. Next section reports some of the most notable among these systems.

2.3.1  Attempts to address issues

In its “manifesto” paper Murray Cole, together with the check-list of issues that next generation of skeleton system should address, sketches the eSkel library [51]. eSkel consists in Cole’s attempt to address the issues he present in his “manifesto” paper. More in detail, eSkel is a library of C functions and type definitions that extends the standard C binding to MPI with skeletal operations. Its underlying conceptual model is the SPMD distributed memory model, inherited from MPI, and its operations must
be invoked from within a program that has already initialized an MPI environment. eSkel provides programmers with some language primitives performing complex operations that can be integrated with the traditional MPI functions. eSkel implements skeletons as collective MPI operations. In (35; 51) authors describe how the manifesto issues are addressed in eSkel. eSkel also provides some code reuse facilities (check-list point V) as most C and C++ code can simply be adapted in eSkel programs. In eSkel heterogeneous architectures are supported (VI) through the usage of MPI, much in the sense heterogeneous architectures are supported through the usage of Java in muskel. However, current implementation of eSkel does not support custom, programmer defined, MPI data types in the communication primitives, that actually use MPI_INT data buffers, and therefore heterogeneous architectures can be targeted using proper MPI implementations just when all the nodes have the same type of processors. No support for dynamicity handling (VII) is provided in eSkel, however.

Some other groups involved in structured parallel programming research, developed programming systems that partially address the issues above presented. Schaeffer and his group at the University of Alberta that implemented a system were programmers can insert new parallelism exploitation patterns in the system (38). Kuchen Muesli (89) is basically a C++ library built on top of MPI providing stream parallel skeletons, data parallel objects and data parallel operations as C++ template classes. The programming interface is definitely very good, as the full power of object oriented paradigm along with templates is exploited to provide Muesli programmers with user-friendly skeletons, and consequently C++ programmers can develop parallel applications very rapidly. In particular, Muesli does not require any MPI specific knowledge/action to write a skeleton program. Therefore, point (I) is very well addressed here. Points (II) and (III) are addressed providing the programmer with a full set of (data parallel) operations that can be freely combined. The payback (IV) is mainly related to the OO techniques exploited to provide skeletons. Code reuse (V) is supported as it is supported in eSkel, as programmers can use C++/C code to build their own skeletons as well as sequential code to be used in the skeletons. Even in this case there is limited support
to heterogeneity (VI): the MPI code in the Skeleton library directly uses MPI\_BYTE buffers to implement Muesli communications, and therefore MPI libraries supporting heterogeneous architectures may be used just in case the nodes sport the same kind of processor and the same C/C++ compiler tool-set. Dynamicity handling (VII) is not supported at all in Muesli.

Gorlatch’s and its research group presented a grid programming environment HOC (73), which provides suitable ways of developing component based grid applications exploiting classical skeleton components. The implementation exploits Web Services technology. Overall, the HOC programming environment addressed principles (I) and (IV). Points (II) and (III) rely on the possibility given to programmers to insert/create new HOCs in the repository. Point (VI) is handled via Web Services. This technology is inherently multiplatform, and therefore heterogeneous target architectures can be easily used to run HOC programs. Point (V) is guaranteed as sequential code can easily (modulus the fact some XML code is needed, actually) be wrapped in Web Services. However, no support to (VII) is included in the current HOC version.

2.4 Our efforts in designing “Skeletons 2.0” systems

Even though Cole and other research groups, focused on skeleton system, designed and developed skeleton systems that own some of the features required to be a next generation skeleton system, the research for addressing the presented issues is just started. In fact, up to now tools and model that are generally recognized as the best solutions for addressing the issues presented in (51) and in (19) simply do not exist. In the Chapters 3, 4 and 5 we present some models and the concerning tools that we designed and developed in order to contribute to research for next generation skeleton systems.

More in detail, in Chapter 3 we propose a macro data-flow based approach designed supporting the integration of unstructured form of parallelization in skeleton systems, hence addressing the issue number II. To
validate the approach we modified a skeleton system that in its original form does not deal with unstructured parallelism: muskel. We extended muskel, in collaboration with the research staff that develop it, to integrate it with a methodology that can be used to implement mixed parallel programming environments providing the programmer with both structured and unstructured ways of expressing parallelism. The methodology is based on data-flow. Structured parallel exploitation patterns are implemented translating them into data-flow graphs executed by a distributed macro data-flow interpreter. Unstructured parallelism exploitation can be achieved by explicitly programming data-flow (sub)graphs. The modified muskel provides suitable ways to interact with the data-flow graphs derived from structured pattern compilation in such a way that mixed structured and unstructured parallelism exploitation patterns can be used within the same application. Two mechanisms provided to the muskel programmers for unstructured parallelism exploitation. First, we provide primitives that allow accessing the fundamental features of the data-flow graph generated out of the compilation of a skeleton program. Namely, methods to deliver data to and retrieve data from data-flow graph. We provide to programmers the ability to instantiate a new graph in the task pool by providing the input task token and to redirect the output token of the graph to an arbitrary data-flow instruction in the pool. Second, we provide the programmer with direct access to the definition of data-flow graphs, in such a way he can describe his particular parallelism exploitation patterns that cannot be efficiently implemented with the available skeletons. The two mechanisms can be jointly used to program all those parts of the application that cannot be easily and efficiently implementing using the skeletons subsystem. Unfortunately, this approach is not free from shortcomings In fact exploiting unstructured parallelism interacting directly with data-flow graph requires to programmers to reason in terms of program-blocks instead of a monolithic program. Hence, at a first sight this approach may look like the ones present in the other early macro data-flow models. Nevertheless, we want to point out that the effort required to customize an application made by a composition of existing skeleton is not comparable with the complexity
of developing it from scratch as a set of macro data-flow blocks.

In order to ease the generation of macro data-flow blocks, and therefore provide programmers with a easier way to express program-blocks, we exploited some metaprogramming techniques that are successfully used for code transformation in fields like web development and component based programming \cite{41, 79, 97}. Exploiting these techniques the programmers are no longer requested to deal with complex application structuring but simply give hints to the metaprogramming support using high-level directives. The directives are used by the support to drive the application transformation. Chapter \ref{chapter:4} presents our efforts aimed at providing metaprogramming tools and models for ease the generation of macro data-flow blocks and their run-time optimization. In particular, two results are presented. The first is “Parallel Abstraction Layer” (PAL). A java annotation \cite{8} based metaprogramming framework that restructures applications at bytecode-level at run-time in order to make them parallel. The parallelization is obtained asynchronously executing the annotated methods. Each method call is transformed in a macro data-flow block that can be dispatched and executed on the available computing resources. PAL transformations depend on the resources available at run-time, the programmers hints and the available adapters. An adapter is a specialized entity that instructs the PAL transformation engine to drive the code transformation depending on the available parallel tools and frameworks. The other result presented in the chapter concerns the integration of the Aspect Oriented Programming \cite{68, 85} mechanisms (more in detail the AspectJ framework \cite{6}) with our modified muskel skeleton framework. The first step in this direction was exploiting AspectJ to implement aspect driven program normalization \cite{18} in muskel. The second step consisted in testing the integration of muskel with AspectJ to in a more complex scenario. Hence, we exploited the aspect oriented programming support integrated in muskel in order to develop workflows which structure and processing are optimized at run-time depending on the available computational resources. Let us point out that we introduced metaprogramming techniques for easing the generation of macro data-flow blocks (in particular to address the issue number I) but
as a corollary we obtained the possibility to optimize the application and adapt it at run-time with respect to the executing environment (addressing the issues number III and VI).

The other two main issues to address are the support for code reuse (V) and the handling of dynamicity (VII). As we already discussed when we introduced muskel, it addresses this last point through the definition of the Application Manager. The dynamicity handling is a very important feature for next generation parallel programming systems, especially for the ones designed for computational Grids. Actually, muskel framework, at least in its original form, is designed to be exploited in cluster and network of workstations rather than in Grids. Indeed, some of its features limit its exploitation on Grids, in particular:

- **muskel** communicates with the resources it recruits exploiting the RMI protocol, that (at least in its original version) uses TCP ports that are typically blocked by firewall;

- the computational resources are found by **muskel** exploiting multicast communications that are often blocked by firewall;

- the recruitment of a computational resource requires to **muskel** programmers to run a proper application on the resource, hence to have an account on it;

- the Application Manager is a centralized entity. This represents a twofold limitation in Grid environment: it is a single point of failure and a bottle-neck that curb the scalability of the approach.

We addressed most of these limitations exploiting ProActive Parallel Suite \(^{108}\) to implement the macro data-flow distributed interpreters (see the experimental results presented in Chapter \(^3\)). ProActive provides mechanisms to tunnel RMI communications and ease the deployment of Grid applications. Indeed, it has been successfully used for developing applications in the Grid5000 \(^2\) platform. ProActive support for Grids has became more complete since it began to support the component based development, in particular the support for the CoreGrid Grid Component Model \(^{52}\). Indeed, several studies recognized that component tech-
nology could be leveraged to ease the development of Grid Application and a few component based model have been proposed by parallel computing scientific community for programming Grids. Component-based software development can be considered an evolutionary step beyond object-oriented design. Object-oriented techniques have been very successful in managing the complexity of modern software, but they have not resulted in significant amounts of cross-project code reuse. Furthermore, sharing object-oriented code is difficult because of language incompatibilities, the lack of standardization for inter-object communication, and the need for compile-time coupling of interfaces. Component-based software development addresses issues of language independence (seamlessly combining components written in different programming languages) and component frameworks define standards for communication among components. Finally, the composition compatibility is evaluated providing a meta-language specification for their interfaces. The GCM represents one of the main European scientific community efforts for designing and developing a grid component model. We contributed to the design of GCM and its reference implementation together with the research group that developed muskel and with several European research groups. In particular, we focused our contribution, in the context of the CoreGrid Programming model virtual institute, on GCM autonomic features. Therefore, by designing the autonomic features of GCM components, each component is able to react dynamically to changes in the executing environment. We referred to the muskel application manager approach, generalizing and extending the approach to make it suitable for components based models. Indeed, each GCM component with a complete support of autonomic features has an Autonomic Manager that observes the component behavior. In case the behavior turns out to be different from the one expected the manager trigger a component reconfiguration. In other words, GCM autonomic features provide programmers with a configurable and straightforward way to implement autonomic grid applications. Hence, they ease the development of application for the Grids. Nevertheless, they rely fully on the application programmer’s expertise for the set-up of the manage-
ment code, which can be quite difficult to write since it may involve the management of black-box components, and, notably, is tailored for the particular component or assembly of them. As a result, the introduction of dynamic adaptivity and self-management might enable the management of grid dynamism, and uncertainty aspects but, at the same time, decreases the component reuse potential since it further specializes components with application specific management code. In Chapter 5, we propose Behavioural Skeletons as a novel way to describe autonomic components in the GCM framework. Behavioural Skeletons aim to describe recurring patterns of component assemblies that can be (either statically or dynamically) equipped with correct and effective management strategies with respect to a given management goal. Behavioural Skeletons help the application designer to i) design component assemblies that can be effectively reused, and ii) cope with management complexity by providing a component with an explicit context with respect to top-down design (i.e. component nesting). We consider the Behavioural Skeletons, coupled with the CoreGRID Grid Component, a good structured parallel programming model for handling dynamicity (VII), supporting reuse both of functional and non-functional code (V). The model defines characters as the Skeleton designers and the Expert users that can design new skeletons and customize the existing ones (II and III), whereas, standard users can easily (I) exploit the existing ones.
Chapter 3

Mixing Structured and Macro-Dataflow approaches

Chapter road-map  In this chapter we describe our contribution to skeleton customization. We start with an introduction on structured programming model outlining its main advantages and recalling its main limitations. In particular, we focus on the skeleton customization issue. Namely the lack of flexibility of skeletal systems in expressing parallel form different from the ones “bundled” with the skeleton framework. Then we briefly introduce the data-flow approach we conceived to address of this limitation and we report related work: alternative approaches addressing the structured parallel programming limitations (Section 3.1). Besides, we introduce classical implementation template and more recent data-flow technologies as used to design and implement skeleton systems (Section 3.2). Then, we describe the details of our contribution, i.e. our extended version of muskel framework, discussing how skeletons customization is supported exploiting data-flow implementation (Section 3.3.1). Finally, we report the experimental results we obtained exploiting our customized muskel (Section 3.4).
3.1 Data-flow enables skeleton customization

We already introduced structured parallel programming models in the previous chapter, where we described their Pros and Cons. Let us briefly recall here their main features and limitations.

Structured parallel programming models provide the programmers with native high-level parallelism exploitation patterns that can be instantiated, possibly in a nested way, to implement a wide range of applications [12; 32; 51; 87; 88]. In particular, those programming models hide to programmers “assembly level” of parallel programming, i.e. by avoiding a direct interaction with the distributed execution environment via communication or shared memory access primitives and/or via explicit scheduling and code mapping. Rather, the high-level native, parametric parallelism exploitation patterns provided encapsulate and abstract from all these parallelism exploitation related details. In contrast, when using a traditional parallel programming system, the programmers have usually to explicitly program code for distributing and scheduling the processes on the available resources and for moving input and output data among the involved processing elements. The cost of this appealing high-level way of dealing with parallel programs is paid in terms of programming freedom. The programmer (or skeleton system user) is normally not allowed to use arbitrary parallelism exploitation patterns, but he must only use the ones provided by the system. They usually include all those reusable patterns that have efficient distributed implementations available. This is mainly aimed at avoiding the possibly for the programmers to write code that can potentially impairs the efficiency of the implementation provided for the available, native parallel patterns. This is a well-known problem (See chapter 2).

In this Chapter we discuss the methodology we conceived, designed and used to modify the muskel parallel programming environment in order to provide to programmers the possibility to mix structured and unstructured ways of expressing parallelism while preserving most of the benefits typical of structured parallel programming models. The methodology is based on the macro data-flow model. Structured parallel ex-
ploitation patterns are implemented translating them into macro data-flow graphs executed by the distributed macro data-flow interpreters. Unstructured, user-defined parallelism exploitation patterns are achieved by explicitly programming data-flow graphs. These (macro) data-flow graphs can be used in the skeleton systems in any place where predefined skeletons can be used, thus providing the possibility to seamlessly integrate both kind of parallelism exploitation within the same program. The mechanisms enabling data-flow graphs customization provide programmers the possibility to program new parallelism exploitation patterns.

The methodology has been developed together with the other authors of [21], we all contributed in a substantially equal way to the conception, design and implementation of the approach.

Macro data-flow implementation for algorithmical skeleton programming environment was introduced in late ’90 [56] and then has been used in other contexts related to skeleton programming environments [101].

Cole eSkel, we already presented in the previous chapter, addresses these problems by allowing programmers to program their own peculiar MPI code within each process in the skeleton tree. Programmers can ask to have a stage of a pipeline or a worker in a farm running on $k$ processors. Then, the programmer may use the $k$ processes communicator returned by the library for the stage/worker to implement its own parallel pipeline stage/worker process. As far as we know, this is the only attempt to integrate ad hoc, unstructured parallelism exploitation in a structured parallel programming environment. The implementation of eSkel, however, is based on process templates, rather than on data flow.

Other skeleton libraries, such as Muesli [87, 88, 89], provide programmers with a quite large flexibility in skeleton programming following a different approach. They provide a number of data parallel data structures along with elementary, collective data parallel operations that can be arbitrary nested to get more and more complex data parallel skeletons. However, this flexibility is restricted to the data parallel part, and it is anyway limited by the available collective operations.

CO2P3S [92] is a design pattern based parallel programming envi-
environment written in Java and targeting symmetric multiprocessors. In CO2P3S, programmers are allowed to program their own parallel design patterns (skeletons) by interacting with the intermediate implementation level (38). Again, this environment does not use data flow technology but implements design patterns using proper process network templates.

JaSkel (69) provides a skeleton library implementing the same skeleton set than muskel. In JaSkel, however, skeletons look much more implementation templates, according to the terminology used in Section 3.2. However, it looks like the programmer can exploit the full OO programming methodology to specialize the skeletons to his own needs. As the programmer is involved in the management of support code too (e.g. he has to specify the master process/thread of a task farm skeletons) JaSkel can be classified as a kind of “low-level, extensible” skeleton system, although it is not clear from the paper whether entirely new skeletons can be easily added to the system (actually, it looks like it is not possible at all).

3.2 Template based vs. data-flow based skeleton systems

A skeleton based parallel programming environment provides programmers with a set of predefined and parametric parallelism exploitation patterns. The patterns are parametric in the kind of basic computation executed in parallel and, possibly, in the execution parallelism degree or in some other execution related parameters. As an example, a pipeline skeleton takes as parameters the computations to be computed at the pipeline stages. In some skeleton systems these computations can be either sequential computations or parallel ones (i.e. other skeletons) while in other systems (mainly the ones developed at the very beginning of the skeleton related research activity) these computations may only be sequential ones.

The first attempts to implement skeleton programming environments all relied on the implementation template technology. As discussed in (94), in a implementation template based skeleton system each skeletons
Pipeline main (...)  
  stage1(...)  
  stage2(...)  
  stage3(...)  
  end pipeline  

  seq stage1(...)  
  {...}  

  farm stage2(...)  
  seq2(...)  
  end farm  

  farm stage3(...)  
  seq3(...)  
  end farm

Source code

Skeleton tree

Process network

Optimized process network

Figure 9: Skeleton program execution according to the implementation template approach.
is implemented using a parametric process network picked up among the ones available for that particular skeleton and for the kind of target architecture at hand in a template library (see (96), discussing several implementation templates, already appeared in bibliography, all suitable to implement task farms, that is embarrassingly parallel computations implemented according to a master-worker paradigm). The template library is designed once and for all by the skeleton system designer and summarizes his knowledge concerning implementation of the parallelism exploitation patterns modeled by skeletons. Therefore, the compilation process of a skeleton program, according to the implementation template model, can be summarized as follows:

1. the skeleton program is parsed, a skeleton tree is derived, hosting the precise skeleton structure of the application. The skeleton tree has nodes marked with one of the available skeleton, and leaves marked with sequential code (sequential skeletons).

2. the skeleton tree is traversed, in some order, and templates from the library are assigned to each one of the skeleton nodes, but the sequential ones, that always correspond to the execution of a sequential process on the target machine. During this phase, parameters of the templates (e.g. the parallelism degree or the kind of communication mechanisms used) are fixed, possibly exploiting proper heuristics associated to the library entries

3. the enriched skeleton tree is used to generate the actual parallel code. Depending on the system that may involve a traditional compilation step (e.g. in P3L when using the Anacleto compiler (47) or in ASSIST when using the astcc compiler tools (14,15)) or exploiting proper parallel libraries (e.g. in Muesli (89) and eSkel (49) exploiting MPI within a proper skeleton library hosting templates

4. the parallel code is eventually run on the target architecture, possibly exploiting some kind of loader/deploy tool.

Figure 9 summarizes the process leading from a skeleton source code to the running code exploiting template technology.
Figure 10: Skeleton program execution according to the data-flow approach.
More recently, an implementation methodology based on data-flow has been proposed (56). In this case the skeleton source code is used to compile a data-flow graph and the data-flow graph is then executed on the target architecture exploiting a suitable distributed data-flow interpreter engine. The approach has been used both in the implementation of Lithium (12; 109) and in Serot’s SKIPPER skeleton environment (100). In both cases, the data-flow approach was used to support fixed skeleton set programming environments. We adopted the very same implementation approach to develop our version of the muskel framework, modifying it in collaboration with the original developers, enriching it with a data-flow implementation to support extensible skeleton sets.

When data-flow technology is exploited to implement skeletons, the compilation process of a skeleton program can be summarized as follows:

1. The skeleton program is parsed, a data-flow graph is derived. The data-flow graph represents the pure data-flow behavior of the skeleton tree in the program.

2. For each one of the input tasks, a copy of the data-flow graph is instantiated, with the task appearing as an input token to the graph. The new graph is delivered to the distributed data-flow interpreter “instruction pool”.

3. The distributed macro data-flow interpreter fetches fireable instructions from the instruction pool and the instructions are executed exploiting the nodes in the target architecture. Possibly, optimizations are taken into account (based on proper heuristics) that try to avoid unnecessary communications (e.g. caching tokens that will eventually be reused) or to adapt the computation grain of the program to the target architecture features (e.g. delivering more than a single fireable instruction to remote nodes to decrease the impact of communication set up latency, or multiprocessing the remote nodes to achieve communication and computation overlap).

Figure [10] summarizes the process leading from skeleton source code to the running code exploiting this data-flow approach.
The two approaches just outlined appear very different, but they have been successfully used to implement different skeleton systems. Let us to point out a quite subtle difference in the two approaches.

On the one side, when using implementation templates, the process network eventually run on the target architecture is very close to the one the programmer has in mind when instantiating skeletons in the source code. In some systems the “optimization” phase of Figure 9 is actually empty and the program eventually run on the target architecture is build out of plain juxtaposition of the process networks making up the templates of the skeletons using in the program. Even in case the optimization phase do actually modify the process network structure (in Figure 9 the master/slave service process of the two consecutive farms are optimized/collapsed, for instance), the overall structure of the process network does not change too much.

On the other side, when a data-flow approach is used the process network run on the target architecture is completely different from the skeleton tree exposed by programmer in the source code. Rather, the skeleton tree is used to implement the parallel computation in a correct and efficient way, exploiting a set of techniques and mechanisms that are much more close to the techniques and mechanisms used in operating systems rather than to those used in the execution of parallel programs, both structured and unstructured. Under a slightly different perspective, this can be interpreted as follows:

- skeletons in the program “annotate” sequential code by providing the meta information required to efficiently implement the program in parallel;

- the support tools of the skeleton programming environment (the macro data-flow graph compiler and the distributed macro data-flow interpreter, in this case) “interpret” the meta information to accurately and efficiently implement the skeleton program, exploiting (possibly at run-time, when the target architecture features are known) the whole set of known mechanisms supporting implementation optimization (e.g. caches, pre-fetching, node multiprocess-
ing, etc.).

Under this perspective, the macro data-flow implementation for parallel skeleton programs opens new perspectives in the design of parallel programming systems where parallelism is dealt with as a “non-functional” feature, specified by programmers and handled by the compiling/runtime support tools in the more convenient and efficient way w.r.t. to the target architecture at hand. In the following Chapters of this thesis will be presented some techniques we exploited to provide programmers methodologies aiming the expression of non-functional requirements and their run-time enforcement.

### 3.3 muskel

We already introduced muskel and its programming model in the Chapter. There we also outlined how we modified muskel, collaborating with its original developers, in order to provide programmers with mechanisms enabling skeleton customizations. In this section we give a more detailed explanation both of the original muskel and of the enhanced version we proposed.

muskel is skeleton programming environment derived from Lithium, it provides the stream parallel skeletons of Lithium, namely stateless task farm and pipeline. These skeletons can be arbitrary nested, to program pipelines with farm stages, as an example, and they process a single stream of input tasks to produce a single stream of output tasks. muskel implements skeletons exploiting data-flow technology and Java RMI facilities. muskel programmers can express parallel computations simply using the provided Pipeline and Farm classes. For instance, to express a parallel computation structured as a two-stage pipeline where each stage is a farm, muskel programmers should write a code such as the one of Figure. The two classes f and g implement the Skeleton interface, i.e. supplying a compute method with the signature

```
Object compute(Object t)
```
computing $f$ and $g$ respectively. The Skeleton interface represents the “sequential” skeleton, that is the skeleton always executed sequentially and only aimed at wrapping sequential code in such a way such code can be used in other, non-sequential skeletons.

In order to execute the program, a muskel programmer first sets up a Manager object. Then, using proper methods, he specifies the program to execute, the performance contract required (in this case, the parallelism degree required for the execution), the input data source (the input stream manager, which is basically an iterator providing the classical boolean hasNext() and Object next() methods) and who is in charge of processing the output data (the output stream manager, just providing a void deliver(Object) method processing a single result of the program). Eventually he can ask parallel program execution simply issuing an eval call to the manager. When the call terminates, an output file is produced.

Actually, the eval method execution happens in steps. First, the manager looks for available processing elements using a simplified, multicast based peer-to-peer discovery protocol, and recruits the required remote
processing elements. Each remote processing element runs a data-flow interpreter. Then the skeleton program (the main of the example depicted in Figure 11) is compiled into a macro data-flow graph (actually capitalizing on normal form results shown in (12; 18)) and a thread is forked for each one of the remote processing elements recruited. Then the input stream is read. For each task item, an instance of the macro data-flow graph is created and the task item token is stored in the proper place (initial data-flow instruction(s)). The graph is placed in the task pool, the repository for data-flow instructions to be executed. Each thread looks for a fireable instruction in the task pool and delivers it for execution to the associated remote data-flow interpreter. The remote interpreter instance associated to the thread is initialized by being sent the serialized code of the data-flow instructions, once and for all before the computation actually starts. Once the remote interpreter terminates the execution of the data-flow instruction, the thread either stores the result token in the proper “next” data-flow instruction(s) in the task pool, or it directly writes the result to the output stream, invoking the deliver method of the output stream manager. If a remote node “fails” (e.g. due to a network failure, or to the node failure/shutdown), the manager looks for another node and starts dispatching data flow instructions to the new node instead (58). As the manager is a centralized entity, if it fails, the whole computation fails. However, the manager is usually run on the machine of the muskel user, which is assumed to be safer than the remote nodes recruited as remote interpreter instances.

The policies implemented by the muskel managers are best effort. The muskel framework tries to do its best to accomplish user requests. In case it is not possible to completely satisfy the user requests, the framework accomplishes to establish the closest configuration to the one implicitly specified by the user with the performance contract. In the example above, the framework tries to recruit 10 remote interpreters. In case only \( n < 10 \) remote interpreters are found, the parallelism degree is set exactly to \( n \). In the worst case, that is if no remote interpreter is found, the computation is performed sequentially, on the local processing element.

In the current version of muskel, the only performance contract actu
ally implemented is the **ParDegree** one, asking for the usage of a constant number of remote interpreters in the execution of the program. We do not enter in more detail in the implementation of the distributed data-flow interpreter here. The interested reader can refer to [56, 58]. Instead, we will try to give a better insight into the compilation of skeleton code into data-flow graphs.

A **muskel** parallel skeleton code is described by the grammar:

\[
P ::= \text{seq}(className) \mid \text{pipe}(P, P) \mid \text{farm}(P)
\]

where the **classNames** refer to classes implementing the **Skeleton** interface, and a macro data-flow instruction is a tuple: \( \langle id, gid, opcode, \mathcal{I}^n, \mathcal{O}^k \rangle \) where \( id \) is the instruction identifier, \( gid \) is the graph identifier (both are either integers or the special \( \text{NoId} \) identifier), \( opcode \) is the name of the **Skeleton** class providing the code to compute the instruction (i.e. computing the output tokens out of the input ones) and \( \mathcal{I} \) and \( \mathcal{O} \) are the input tokens and the output token destinations, respectively. An input token is a pair \( \langle \text{value}, \text{presenceBit} \rangle \) and an output token destination is a pair \( \langle \text{destInstructionId}, \text{destTokenNumber} \rangle \). With these assumptions, a data-flow instruction such as:

\[
\langle a, b, f, \langle \langle 123, \text{true} \rangle, \langle \text{null, false} \rangle \rangle, \langle \langle i, j \rangle \rangle \rangle
\]

is the instruction with identifier \( a \) belonging to the graph with identifier \( b \). It has two input tokens, one present (the integer 123) and one not present yet. It is not fireable, as one token is missing. When the missing token will be delivered to this instruction, coming either from the input stream or from another instruction, the instruction becomes fireable. To be computed, the two tokens must be given to the **compute** method of the \( f \) class. The method computes a single result that will be delivered to the instruction with identifier \( i \) in the same graph, in the position corresponding to input token number \( j \). The process compiling the skeleton program into the data-flow graph can therefore be more formally described as follows. We define a pre-compile function \( PC[\ ] \) as:

\[
PC[\text{seq}(f)]_{gid} = \lambda i.\{(\text{newId}()), gid, f, \langle \text{null, false} \rangle, \langle i, \text{NoId} \rangle \}
\]
\[ PC[farm(P)]_{gid} = C[P]_{gid} \]
\[ PC[\text{pipe}(P_1, P_2)]_{gid} = \lambda i.\{C[P_1]_{gid}(get\text{Id}(C[P_2]_{gid})), C[P_2]_{gid}(i)\} \]

where \( \lambda x.T \) is the usual function representation \( ((\lambda x.T)(y) = T|_{x=y}) \) and \( \text{get\text{Id}}() \) is the function returning the \( id \) of the first instruction in its argument graph, that is, the one assuming to receive the input token from outside the graph, and a compile function \( C[] \) such as:

\[ C[P] = PC[P]_{\text{newGid}()}(\text{NoId}) \]

where \( \text{newId}() \) and \( \text{newGid}() \) are stateful functions returning a fresh (i.e. unused) instruction and graph identifier, respectively. The compile function returns therefore a graph, with a fresh graph identifier, hosting all the data-flow instructions relative to the skeleton program. The result tokens are identified as those whose destination is \( \text{NoId} \). As an example, the compilation of the \textbf{main} program \( \text{pipe}(\text{farm}(\text{seq}(f)), \text{farm}(\text{seq}(g))) \) produces the data flow graph:

\[
\{(1,1,f,\langle\text{null, false}\rangle,\langle(2,1)\rangle), (2,1,g,\langle\text{null, false}\rangle,\langle\text{NoId, NoId}\rangle)\}
\]

(assuming that identifiers and token positions start from 1).

When the application manager is told to actually compute the program, via an \textbf{eval}() method call, the input file stream is read looking for tasks to be computed. Each task found is used to replace the data field of the lower \( id \) data-flow instruction in a new \( C[P] \) graph. In the example above, this results in the generation of a set of independent graphs such as:

\[
\{(1,i,f,\langle\text{task}_i, \text{true}\rangle,\langle(2,1)\rangle), (2,i,g,\langle\text{null, false}\rangle,\langle\text{NoId, NoId}\rangle)\}
\]

for all the tasks ranging from \text{task}_1 to \text{task}_n.

All the resulting instructions are put in the task pool of the distributed interpreter in such a way that the control threads taking care of “feeding” the remote data-flow interpreter instances can start fetching the fireable instructions. The output tokens generated by instructions with destination tag equal to \text{NoId} are directly delivered to the output file stream by the threads receiving them from the remote interpreter instances. Those
with a non-NoId flag are delivered to the proper instructions in the task pool that will eventually become fireable.

3.3.1 Programmer-defined skeletons

In order to introduce completely new parallelism exploitation patterns, our version of the muskel framework provides programmers with mechanisms that can be used to design plain, arbitrary macro data-flow graphs. A macro data-flow graph can be defined creating some Mdfi (macro data-flow instruction) objects and connecting them in a MdfGraph object. As an example, the code in Figure 12 is the code needed to program a data-flow graph with two instructions. The first one computes the compute method inc1 on its input token and delivers the result to the second instruction. The second one, computes the sq1 compute method on its input token and delivers the result to a generic “next” instruction (this is modeled giving the destination token tag a Mdfi.NoInstrId tag). The Dest stuff in the code is meant to represent destination of output tokens as triples hosting the graph identifier, the instruction identifier and the
destination input token targeted in this instruction. Macro data-flow instructions are build stating the manager they refer to, their identifier, the code executed (must be a Skeleton object) the number of input and output tokens and a vector with a destination for each one of the output tokens. Take into account that the simple macro data-flow graph of Figure 12 is actually the very same macro data-flow graph derived compiling a primitive muskel skeleton code such as:

```java
Skeleton main = new Pipeline(new Inc(), new Sq());
```

More complex, programmer-defined macro data-flow graph may comprehend instructions delivering tokens to an arbitrary number of other instructions, as well as instructions gathering input tokens from several distinct other instructions.

MdfGraph objects are used to create new ParCompute objects. The ParCompute objects can be used in any place were a Skeleton object is used. Therefore programmer-defined parallelism exploitation patterns can be used as pipeline stages or as farm workers, for instance. The only limitation on the graphs that can be used in a ParCompute object consists in requiring that the graph has a unique input token and a unique output token.

When executing programs with programmer-defined parallelism exploitation patterns the process of compiling skeleton code to macro data-flow graphs is slightly modified. When an original muskel skeleton is compiled, the process described above is applied. When a programmer-defined skeleton is compiled, the associated macro data-flow graph is directly taken from the ParCompute instance variables where the graph supplied by the programmer is maintained. Such graph is linked to the rest of the graph according to the rules relative to the skeleton where the programmer-defined skeleton appears. To show how the whole process works, let us suppose we want to pre-process each input tasks in such a way that for each task $t_i$ a new task

$$t'_i = h_1(f_1(t_i), g_2(g_1(f_1(t_i))))$$

is produced. This computation cannot be programmed using the stream parallel skeletons currently provided by the original muskel. Then we
Figure 13: Mixed sample MDF graph: the upper part comes from a programmer-defined MDF graph (it cannot be derived using primitive muskel skeletons) and the lower part is actually coming from a three stage pipeline with two sequential stages (the second and the third one) and a parallel first stage (the programmer-defined one).

want to process the preprocessed tasks through a two-stage pipeline, in order to produce the final result. In this case the programmer can set up a new graph using a code similar to the one shown in Figure 11 and then used that new ParCompute object as the first stage of a two-stage pipeline whose second stage happens to be the postprocessing two-stage pipeline. When compiling the whole program, the outer pipeline is compiled first. As the first stage is a programmer-defined skeleton, its macro data-flow graph is directly taken from the programmer-supplied one. The second stage is compiled according to the (recursive) procedure previously described and eventually the (unique) last instruction of the first graph is modified in such a way it sends its only output token to the very first instruction in the second stage graph. The resulting graph is outlined in Figure 13.
Making good usage of the mechanisms that allow to define new data-flow graphs, the programmer can arrange to express computations with arbitrary mixes of arbitrary data-flow graphs and graphs coming from the compilation of structured, stream parallel skeleton computations. The execution of the resulting data-flow graph is supported by the muskel distributed data-flow interpreter as the execution of any other data-flow graph derived from the compilation of a skeleton program. Therefore, the customized skeletons are efficiently executed as the skeletons “bundled” with muskel. Indeed, in data-flow based skeleton systems, as we already stated when we presented them, the optimizations do not directly depend on the skeleton structure but on the data-flow engine capability of executing the macro data-flow instruction in an efficient way.

In order to allow primitive muskel skeleton usage as code to be executed in an instruction of a programmer-defined macro data-flow graph it is sufficient to compile “on the fly” the primitive skeleton and include the result (i.e. the macro data-flow graph) of this compilation in the programmer-defined macro data-flow graph.

As a final example, consider the code of Figure 14. This code actually shows how a new Map2 skeleton, performing in parallel the same computation on all the portions of an input vector, can be defined and used. It’s worth pointing out how programmer-defined skeletons, once properly debugged and fine-tuned, can simply be incorporated in the muskel skeleton framework and used seamlessly, as the primitive muskel ones, but for the fact (as show in the code) the constructor needs the manager as a parameter. This is needed just to be able to link together the macro data-flow graphs generated by the compiler and those supplied by the programmer. This feature has been released by postponing the data-flow graph creation to the moment the graph needs to be instantiated after the arrival of a new task to compute, as at that time all the information necessary to perform graph “conjunction” is available.
public class Map2 extends ParCompute {

    public Map2(Skeleton f, Manager manager) {
        super(null);
        program = new MdfGraph(); // first build the empty graph
        Dest[] dds1 = new Dest[2]; // build the emitter instruction
        dds1[0] = new Dest(0,2);
        dds1[1] = new Dest(0,3);
        Mdfi emitter = new Mdfi(manager, 1, new MapEmitter(2), 1, 2, dds1);
        program.addInstruction(emitter); // add it to the graph
        Dest[] dds2 = new Dest[1]; // build first half map Skeleton node
        dds2[0] = new Dest(0,4);
        Mdfi if1 = new Mdfi(manager,2, f, 1, 1, dds2);
        program.addInstruction(if1); // add it to the graph
        Dest[] dds3 = new Dest[1]; // build second half map Skeleton node
        dds3[0] = new Dest(1,4);
        Mdfi if2 = new Mdfi(manager,3, f, 1, 1, dds3);
        program.addInstruction(if2); // add it to the graph
        Dest[] ddslast = new Dest[1];
        ddslast[0] = new Dest(0,Mdfi.NoInstrId);
        Mdfi collector = new Mdfi(manager,4,new MapCollector(), 2, 1, ddslast);
        program.addInstruction(collector);
        return;
    }
}

public class SampleMap {

    public static void main(String[] args) {
        Manager manager = new Manager();
        Skeleton worker = new Fdouble();
        Skeleton main = new Map2(worker,manager);

        InputManager inManager = new DoubleVectorIM(10,4);
        OutputManager outManager = new DoubleVectorOM();

        ParDegree contract = new ParDegree(10);
        manager.setInputManager(inManager);
        manager.setOutputManager(outManager);
        manager.setContract(contract);
        manager.setProgram(main);

        manager.compute();
    }
}

Figure 14: Introducing a new, programmer-defined skeleton: a map working on vectors and with a fixed, programmer-defined parallelism degree.
3.4 Experimental results

To validate our approach we conducted some test with our modified version of the *muskel* framework. The original *muskel* interpreter engine has been left basically unchanged, whereas the part supporting parallelism exploitation pattern programming has been changed to support linking of custom MDF graphs to the code produced by the compiler out of plain *muskel* skeleton trees. We used our customized version for implementing an application that can not be (at least not easily) implemented using standard (i.e. without our proposed customization support) skeleton environments.

Figure 16 summarizes the typical performance results of our enhanced interpreter. We ran several synthetic programs using the custom macro data-flow graph features introduced in *muskel*. We designed the programs in such a way the macro data-flow instructions appearing in the graph had a precise “average grain” (i.e. average ration between the time spent by the remote interpreter to compute the macro data flow instruction sent to it, and the time spent in communicating data to the remote interpreter plus the time to retrieve the computation results). For each test-bed we passed as input parameters to the developed programs 1K input tasks.

The results show that when the computational grain is small, *muskel* does not scale well, even using a very small number of remote interpreter instances. Indeed, Figure 16 clearly shows that when the computational grain is 3 the efficiency rapidly decreases, going under 0.7 even when only four computational resources are used. When the grain is 70 the efficiency goes under 0.8 only when the number of recruited computational resources is higher than 14. Finally, when the grain is high enough (about 200 times the time spent in communications actually spent in computation of MDF instructions) the efficiency is definitely close to the ideal one even using 16 or more machines.

Despite the data shown refers to some synthetic computations, actual computations (e.g. image processing ones) achieved very similar results. This because the automatic load balancing mechanism implemented in
Figure 15: Effect of middleware: scalability of the muskel prototype using plain RMI vs. the one using ProActive active objects

the muskel distributed interpreter, obtained by mean of auto scheduling techniques, perfectly optimized the execution of variable grain MDF instructions. All the experiments have been performed on a Linux (kernel 2.4.22) RLX Pentium III blade architecture, with Fast Ethernet connection among the blades, equipped with Java 1.4.1_01 run-time.

Despite measuring scalability of our modified muskel framework, we also have taken into account the possibility to use different mechanisms to support distributed data-flow interpreter execution. In particular, we investigated the possibility of implementing the muskel approach for skeleton customization on top of the ProActive framework (108) both to be able to target a different set of architectures and to demonstrate the “portability” of our approach, i.e. that it is a feasible and efficient solution not only when it exploits the muskel data-flow interpreter.

For this purpose, we conducted some experiments aimed at verifying the overhead introduced by ProActive with respect to the plain Java RMI muskel prototype, when using the secure shell (ssh) tunneling of the
RMI protocol (feature natively provided by the ProActive framework). In particular, we modified the “kernel” of the data-flow interpreter of muskel in order to make it able to exploit the ProActive active objects in place of plain RMI objects as remote data-flow interpreter instances. The results we achieved are summarized in Figure 15. The figure plots the completion times for the very same program run on a Linux workstation cluster when using plain Java RMI and when using ProActive active objects to implement the remote data-flow interpreter instances. The macro data-flow instructions, in this case, have a grain comparable to the “high grain” of instructions of Figure 16. Experiments showed that ProActive active objects are slightly less efficient but the difference is negligible. In this case, the setup time of the remote data-flow interpreter instances was not considered in the overall completion time, being paid once and forall when the system is started up.
Figure 16: Scalability of the muskel prototype and effect of computation grain.
Summarizing the Chapter

In this Chapter we discussed a methodology for extending algorithmic skeletons based parallel programming frameworks aimed at providing programmers with the possibility to freely customize the structure of their parallel applications. It is based on mechanisms allowing programmers to modify the data-flow graph derived from the compilation of skeleton based application. In particular, we discussed how we modified the muskel framework for parallel programming. The version we developed (collaborating with the team that developed the original muskel) supports extendability of the skeleton set, as advocated by Cole in his “manifesto” paper [51]. In particular, we discussed how our modified muskel supports the introduction of new skeletons, modeling parallelism exploitation patterns not originally covered by the primitive muskel skeletons. This possibility is supported by allowing programmers to define new skeletons providing the arbitrary data-flow graph executed in the skeleton and by letting muskel to seamlessly integrate such new skeletons in the primitive ones. We also presented experimental results validating our muskel approach to extend and customize its skeleton set. As far as we know, this is the most significant effort in the skeleton community to tackle problems deriving from a fixed skeleton set. Only Schaeffer and his group at the University of Alberta implemented a system were programmers can, in controlled ways, insert new parallelism exploitation patterns in the system [38], although the approach followed here is a bit different, in that programmers are encouraged to intervene directly in the run-time support implementation, to introduce new skeletons, while in our muskel new skeletons may be introduced using the intermediate macro data-flow language as the skeleton “assembly” language.
Chapter 4

Metaprogramming
Run-time Optimizations

Chapter road-map  This Chapter presents our efforts aimed at exploiting metaprogramming techniques for optimizing at run-time the execution of structured parallel applications. The approaches are based on the run-time generation of macro data-flow blocks from the application code. We start presenting the motivations (Section 4.1) of our contributions. Then we present PAL (Section 4.2), our first result in the field. PAL is a metaprogramming engine that transforms at run-time an annotated sequential java code in a parallel program, exploiting both programmer hints and executing platform information. We describe our PAL prototype implementation (Section 4.2.1) and the results of the tests we made with it (Section 4.2.2). After we discuss the motivations that convinced us to integrate the PAL approach with our version of the muskel framework (Section 4.2.3). In the following section (4.3) we describe the preliminary attempts we made integrating metaprogramming techniques in muskel. In Section 4.4 we present how we further enhanced muskel making it able to exploit metaprogramming for run-time code optimizations. In particular, how it can be exploited to optimize the parallel execution of computations expressed as workflows. In Section 4.4.2 we describe the implementation of workflows transformations and in Section 4.4.3 we present the performance results obtained. Finally, we compare the two approaches (Section 4.5) and we summarize the Chapter contributions.
4.1 Our efforts in run-time optimization

In the previous chapter we described how the macro data-flow model can be exploited in order to allow the customization of algorithmic skeletons. We showed how we modified the muskel parallel framework in order to provide programmers with mechanisms able to change skeletons structure. In this chapter we present the metaprogramming techniques we exploited both to ease the generation of the macro data-flow graph and to optimize at run-time the parallel execution of the macro data-flow blocks.

4.1.1 Metaprogramming

Code-generating programs are sometimes called metaprograms; writing such programs is called metaprogramming. Metaprograms do part of the work during compile-time that is otherwise done at run-time. Compile-time metaprogramming exploits information available at compile-time to generate temporary source code, which is merged by the compiler with the rest of the source code and then compiled. The goal of run-time metaprogramming, instead, is to achieve real-time code optimizations transforming or adapting the code whenever some information becomes available.

Compile-time metaprogramming

The most common metaprogramming tool is a compiler, which allows a programmer to write a relatively short program in a high-level language and uses it to write an equivalent assembly language or machine language program. Another still fairly common example of metaprogramming might be found in the use of Template Metaprogramming. Template metaprogramming is a metaprogramming technique in which templates are used by a compiler to generate temporary source code, which is merged by the compiler with the rest of the source code and then compiled. The output of these templates includes compile-time constants, data structures, and complete functions. The use of templates can be thought of as compile-time execution. The technique is used by a number
of languages, the most well-known being C++, but also D, Eiffel, Haskell, ML and XL. The use of templates as a metaprogramming technique requires two distinct operations: a template must be defined, and a defined template must be instantiated. The template definition describes the generic form of the generated source code, and the instantiation causes a specific set of source code to be generated from the generic form in the template. Template metaprogramming is generally Turing-complete, meaning that any computation expressible by a computer program can be computed, in some form, by a template metaprogram. Templates are different from macros. A macro, which is also a compile-time language feature, generates code in-line using text manipulation and substitution. Macro systems often have limited compile-time process flow abilities and usually lack awareness of the semantics and type system of their companion language (an exception should be made with Lisp’s macros, which are written in Lisp itself, and is not a simple text manipulation and substitution). Template metaprograms have no mutable variables that is, no variable can change value once it has been initialized, therefore template metaprogramming can be seen as a form of functional programming. In fact, many template implementations only implement flow control through recursion. Some common reasons to use templates is to implement generic programming (avoiding sections of code which are similar except for some minor variations) and especially to perform automatic compile-time optimization such as doing something once at compile-time rather than every time the program is run, for instance having the compiler unroll loops to eliminate jumps and loop count decrements whenever the program is executed. The main problem of this approach is the inefficient exploitation of the executing environment. Indeed to guarantee the code portability such optimizations are done in a generic way, for instance without exploiting specific CPU extension like SSE or 3DNow. To overwork it the application should be re-compiled once all the running architecture details are known.
Run-time metaprogramming

Run-time metaprogramming points at either the generation of programs specialized with respect to the running architecture or the adaptation of programs with respect to additional information provided by programmers, e.g. non-functional requirements. The metaprogramming related information (metadata) is processed by the metaprogramming run-time support. It exploits both such metadata and the environmental information to transforms the original code into an optimized one. Nevertheless, this solution presents a major problem: the re-compilation overhead. Indeed, re-compile the whole application from scratch on each machine it is moved for execution is computationally expansive. A viable solution consists in writing the applications using bytecode based languages, like Java and .NET. Indeed, their compilers do not translate the program into target machine language but translate it into an intermediate language (IL). The IL has greater expressiveness than the machine and the assembly languages and can be transformed in a machine-level program paying a small overhead. Furthermore, there are other advantages in implementing application, especially the distributed ones, exploiting a virtual machine based language: e.g. the possibility to run programs across different platforms at the only cost of porting the execution environment and to achieve better security (the execution engine mediates all accesses to resources made by programs verifying that the system can not be compromised by the running application).

In the past, other programming languages with the same architecture, essentially p-code, have been proposed (see for instance the introduction of [86]) but Java has been the first to have a huge impact on programming mainstream. Java approach has been recognized as successful, indeed, since the 2002 also Microsoft introduced their virtual-machine based programming languages. They are based on the Common Language Infrastructure (CLI). The core of CLI is the virtual execution system also known as Common Language Runtime (CLR). Both JVM [91] and CLR [7] implement a multi-threaded stack-based virtual machine, that offers many services such as dynamic loading, garbage collection, clearly the Just In
Time (JIT) compilation and above all a noteworthy reflection support. Features like garbage collection raise the programming abstraction level whereas dynamic loading, JIT compilation and a native multi-thread support simplify the task of programming distributed and concurrent applications. Reflection support enables programs to read its own metadata. A program reflecting on itself extract metadata (from its representation expressed in terms of intermediate language) and using that metadata can modify its own behavior. Reflection support is useful to inspect the structure of types, to access fields and even to choose dynamically the methods to invoke. Exploiting reflection support programs can change their structure and their (byte) code. The reflection support can be provided by the run-time system at different levels of complexity (36):

- **Introspection**: the program can access to a representation of its own internal state. This support may range from knowing the type of values at run-time to having access to a representation of the whole source program.

- **Intercession**: the representation of the state of the program can be changed at run-time. This may include the set of types used, values and the source code.

Both introspection and intercession require a mechanism, called reification, to expose the execution state of a program as data. The reification mechanism exposes an abstraction of some elements of the execution environment. These elements may include programming abstractions such as types or source code; they may also include other elements, like the evaluation stack (as in 3-LISP (105)), that are not modeled by the language. For compiled languages it could be harder to reflect elements of the source language: the object program runs on a machine that usually is far from the abstract machine of the source language. Enabling RTTI (Runtime Type Identification, a support that allows a program to have exact information about type of objects at run-time) in C++, for instance, requires that the run-time support contain additional code to keep track of types at run-time. Besides, the programmer would expect abstractions compatible with the structure of the programming language abstract ma-
chine (unless he is interested in manipulating the state of the machine that is target of the compilation).

Custom metadata management

The metadata readable through the advanced reflection supports are both the information about types (class, method, field names an hierarchies) and about additional, non-functional attributes. A straightforward example is the Java serialization architecture: the programmer can declare the instances of a serializable class simply by implementing the `Serializable` interface, which in fact is an empty interface. Thus, two types that differ only for the implementation of the `Serializable` interface are indistinguishable from the execution (functional) standpoint. Besides, the serialization of the instances of non-serializable types will not be allowed by the serialization support. Clearly, this “interface-based” mechanism for the metadata specification is not flexible and can not be expressed at more fine level, for instance at method-level. This limitation leads to the development of Java annotations (8). A Java annotation is a special syntax that adds metadata to Java source code. Annotations can be added to program elements such as classes, methods, fields, parameters, local variables, and packages. Unlike Javadoc tags, Java annotations are reflective in that they may be retained by the Java VM and made retrievable at run-time. The possibility to retain and retrieve this information at run-time makes the “real” difference between the Java annotations and the earlier annotation based approach. For instance, the OpenMP pragma based approach or the HPF annotation or consisting in simple directives to compiler driving the data decomposition optimization, approaches that are not designed to work with non-shared memory architectures.

The exploitation of Java annotations as a way to embed non-functional information is at the base of Attribute Oriented Programming (98; 114). Attribute Oriented Programmers use Java annotations to mark program elements (e.g. classes and methods) to indicate that they maintain the application-specific or domain-specific semantics. As an example, some programmers may define a “logging” attribute and associate it with a method to indicate the method should implement a logging function,
while other programmers may define a “web service” attribute and associate it with a class to indicate the class should be implemented as a web service. Attributes aim the separation of concerns: application’s core logic (or business logic) are clearly distinguished from application-specific or domain-specific semantics (e.g. logging and web service functions). By hiding the implementation details of those semantics from program code, attributes increase the level of programming abstraction and reduce programming complexity. The program elements associated with attributes are transformed in order to fit the programmers’ requirements.

The effectiveness of the approach is demonstrated by its rapidly diffusion, indeed some very popular and widely used programming frameworks \((70; 79)\) adopted the Attribute Oriented Programming approach as a way to embed programmers’ hints and requirements. There are also some scientific works exploiting annotations information to drive the application run-time transformation, for instance in \((45)\) authors propose a way to transform an annotated application in a multithreaded one and \((97)\) describes a way to transform a POJO in a Fractal component simply transforming the code according to the programmer annotations.

In Section 4.2 we describe how we exploited the Attribute Oriented Programming approach in our Parallel Abstraction Layer (PAL). PAL is a metaprogramming engine able to dynamically restructure parallel applications depending both on the information gathered at run-time about the running platform and on the hints specified inside the source code by programmers.

A slightly different approach that aims to a clear separation between the application business code and application management information is the Aspect Oriented Programming (AOP) model. Whereas the Attribute Oriented Programming model separates the management code from the business one exploiting a language support, Aspect Oriented Programming model requires programmers provide additional files containing a set of rules which describe the actions to perform when the application execution flow reach certain points. The main actions performed consist in code injection and code substitution. Some scientific works exploit AOP for code transformations. Sobral et al. discussed the
usage of AOP to support modular computing (53) [106] [107]. They use AOP techniques to separately solve partition, concurrency and distribution problems and eventually show how the related aspects can be used to provide a (kernel for a) general purpose, modular parallel computing framework. Other authors (33) demonstrated that AOP can be efficiently exploited in conjunction with components and patterns to derive parallel applications for distributed memory systems. It highly relies on the ability of the programmer to find out the right places to exploit aspects. In (78) another approach exploiting aspects to parallelize Java applications from the Java Grande forum using AspectJ is presented. Good results are shown in the paper, but the procedure used to exploit aspects requires entering the program details to find out possibilities for parallelization.

In the Sections 4.3 and 4.4 we describe how we integrated the AOP approach in our next generation muskel. In particular, how we exploited the AspectJ (6) tool to manage the generation of macro data-flow blocks, aimed at the parallelization of workflow computations.

Both the PAL and the AspectJ integration with muskel approaches have been published, respectively in (61) and (60). In both the cases the authors collectively contributed to the paper.

4.2 The PAL experience

The Parallel Abstraction Layer is a general-purpose approach for implementing simple parallel applications that does not require complex application structuring by programmers. Programmers are only required to insert, in the source code, some hints, eventually exploited by the PAL run-time support to transform the application code. The transformation is aimed at in enforcing an efficient parallel (even distributed) execution of the application. The general idea is outlined in Figure 17. Programmers’ hints consist in non-functional requirements, namely, requirements which specify criteria that can be used to judge the operation of a system, rather than specific behaviors. Examples of non-functional requirements includes: Efficiency, Price, Hardware Reliability, Software and tools availability and Parallelism degree. In PAL implementation they are speci-
fied through the annotation mechanisms provided by Java (8). The PAL run-time support exploits the information conveyed in the annotations to transform the original program in a parallel one. The transformed program is optimized with respect to the target parallel/distributed architecture.

Programmers are required to give some kind of “parallel structure” to the code directly at the source code level, as it happens in the algorithmic skeleton case. In our PAL implementation it can be done exploiting the java annotation mechanism. For instance, the farm semantics is obtained indicating which “parts” of code should be replicated and executed in parallel. A “part” is intended to be a piece of side-effect free code which input and output data are well-defined. Programmers are in charge of ensuring the “parts” satisfy these requirements. Each java code “part” is transformed by the PAL in a macro data-flow block that can be dispatched for execution.

PAL has a multi-level software architecture. It is depicted in Figure 19. On top, there is PAL frontend, namely the annotations provided by PAL and the host language, Java in our PAL implementation. In the bottom layer, there are the adapters and the information system: the formers foster PAL during code transformation instructing it about how to structure the application code to make it parallel and compliant with a specific
public class Mandelbrot{
    public void paint(GraphicsContext gcont) {
        // computing image size
        ... 
        Vector<PFFuture<Vector<Vector<Integer>>>> man = 
            new Vector<PFFuture<Vector<Vector<Integer>>>>>(numOfLines);
        for(int i=0;i<numOfLines;i++)
            man.add(createLines(...);
        ...
    }
}

@Parallel(parDegree=16)
public PFFuture<Vector<Vector<Integer>>> createLines (params ...){
    Vector<Vector<Integer>> v = new Vector<Vector<Integer>>();
    // compute points ...
    for (int i = 0; i<cls; i++) {
        ...
        v.add(point);
    }
    return new PFFuture<Vector<Vector<Integer>>>(v);
}

public class Main {
    ...
    public static void main(String[] args) {
        Class[] toBeTransformed = new Class[2];
        toBeTransformed[0] = Main.class;
        toBeTransformed[1] = Mandelbrot.class;
        PAL.transform(toBeTransformed,args);
        Mandelbrot mBrot = new Mandelbrot();
        BufferedImage bi = new BufferedImage(2400,1600,TYPE_INT_BGR);
        mBrot.paint(GraphicsEnvironment.getLocalGraphicsEnvironment().createGraphics(bi));
    }
}

Figure 18: Sample code using PAL

parallel framework. The latter is a set of tools aimed at run-time information gathering. Finally, the middle layer is the real metaprogramming engine that uses the information gathered in order to decide which adapter exploit among the available to enforce the non-functional requirements expressed by the programmers through annotations.

Compared with traditional skeletal environments, PAL presents three additional advantages.
• First, annotations can be ignored and the semantics of the original sequential code is preserved. This means that the programmers’ application code can be run through a classical sequential compiler (or interpreter) suite and debugged using normal debugging tools.

• Second, annotations are processed at run-time, typically exploiting reflection properties of the hosting language. As a consequence, while handling annotations, a bunch of knowledge can be exploited which is not available at compile-time (kind of machines at hand, kind of interconnection network, etc.) and this can lead to more efficient parallel implementations of the user application.

• Third, the knowledge concerning the kind of target architecture can be exploited leading to radically diverse implementation of the very same user code. As an example, if the run-time can figure out that the target architecture where the program is running happens to be a grid, it can transform the code in such a way possibly coarser grain parallelism is exploited. On the other hand, in case the run-time figures out that user asked to execute the code on a SMP target, a more efficient, possibly finer grain, multithreaded version of the code can be produced as the result of the annotation handling.
PAL enforces code optimizations via automatic application restructuring in order to exploit all the available application parallelism with respect to programmer’s annotations (non-functional application requirements). The transformation process is done at run-time, which is at the time we have the information we need to optimize the restructuring process with respect to the available parallel tools and underlying resources. The code is transformed at bytecode level thus, it does not need to recompile the application source code on the target architecture. Hence, the transformation introduces only a small overhead for the code transformations.

The generative (54) metaprogramming engine of PAL gathers at run-time information on available parallel tools and computational resources. Then, it analyzes the bytecode looking for programmer annotations (non-functional requirements) and transforms the annotated original code to an optimized, parallel one. The structure of the transformed bytecode depends on the selected parallel framework (clearly subjected to adapters availability) and on the presence and/or value of some non-functional requirements.

PAL exploits the available parallelism by asynchronously executing parts of the original code. The parts to be executed asynchronously are individuated by the annotations specified by programmers. In particular, in Java the most natural choice consists in individuating methods calls as the parts to be asynchronously executed. Asynchronous execution of method code is based on the concept of future (43, 44). When a method is called asynchronously it immediately returns a future, that is a stub “empty” object. The caller can then continue its own computations and access to the future object content (e.g. calling its methods) just when needed. If in the meanwhile the return value has already been computed, the call to reify the future succeeds immediately, otherwise it blocks until the actual return value is computed and then returns it.

In our PAL implementation, to indicate a method as “parallelizable” PAL programmers have simply to put a proper @Parallel annotation enriched with non-functional requirements, such as the required parallelism degree, on the line right before method declaration. Exploiting the an-
notation mechanism allows to keep the PAL applications very similar to normal sequential applications, actually. Hence, Programmers may simply run the application through standard Java tools to verify it is functionally correct. PAL autonomically performs at run-time activities aimed at achieving the asynchronous and parallel execution of the PAL-annotated methods and at managing any consistency related problems, without any further programmer intervention. The PAL approach also avoids the proliferation of source files and classes, that is a quite common situation in framework based programming, as it works transforming bytecode. Unfortunately, it raises several problems related to data sharing management. As an example, methods annotated with a \texttt{@Parallel} should not access class fields: they may only access their own parameters and the local method variables. This is due to the impossibility to intercept all the accesses to non-private class fields. This limitation prevent the usage of static class fields as a way for sharing data among different instances of annotated method calls, making more complex the development of application in which the computational resources running the different annotated method calls need to exchange data during the method computation. It is worth to note that this is not a limitation of the approach but depends by the Java language. Indeed having a proper language support for detecting public field changes it would not be difficult to provide a proper annotation for managing the remote accesses to fields.

4.2.1 PAL: implementation details

We implemented a PAL prototype in Java 1.5, as Java provides a manageable intermediate language (Java bytecode\textsuperscript{(110)}) and natively supports code annotations, since version 1.5. Furthermore, it owns all the properties needed by our approach (e.g. type safety and security). For this implementation we developed two distinct adapters. One for transforming the bytecode in a multithreaded one and another to transform the bytecode making it compliant with JJPF. In order to do this our PAL implementation makes better usage of ASM\textsuperscript{(40)}: a Java bytecode manipulation framework.
The current PAL prototype accepts only one kind of non-functional attribute that can be specified with the \texttt{@Parallel} annotation: \texttt{parDegree}. It denotes the number of processing elements to be used for the method execution. PAL uses such information to make a choice between the multithreaded and JJPF adapter. This choice is driven by the number of processors/cores available on the host machine: if the machine owns a sufficient number of processors the annotated bytecode directly compiled from user code is transformed in a semantically equivalent multithreaded version. Otherwise, PAL chooses to transform the compiled bytecode in a semantically equivalent JJPF version that uses several networked machines to execute the program. PAL basically transforms code in such a way the annotated methods can be computed asynchronously. The original code is “adapted” using an adapter in order to be compliant with the parallel framework associated with the adapter. In our implementation, where the only available adapter for distributed computations is the JJPF one, the methods are adapted to be run on the remote JJPF servers displaced onto the processing elements. Conversely, the \texttt{main} code invoking the \texttt{@Parallel} methods is used to implement the “client” code, i.e. the application the user runs on its own local machine. This application eventually will interact with the remote JJPF servers according to proper JJPF mechanisms and protocols. Method call parameters, the input data for the code to be executed asynchronously, are packaged in a “task”. When a server receives a task to be computed, it removes its server-descriptor from the processing elements available for JJPF. When the task computation is completed the server re-inserts its descriptor from the available ones. In other words, when a annotated method is called an empty future is immediately returned, a “task” is generated and it is inserted into the JJPF queue; eventually it is sent to one among the available processing element, which remove itself from the available resources, computes the task and returns the result that JJPF finally put inside the proper future. This implementation schema looks like very close to a classical master/slave implementation.

We could have developed an adapter for other parallel programming frameworks as targets. As an example, we could have used the Globus
toolkit. However, JJPF is very compact and required a slightly more compact amount of code to be targeted, with respect to the Globus or other grid middleware frameworks. As the principles driving the generation of the parallel code are the same both using JJPF and other grid middleware frameworks, we preferred JJPF to be able to implement a proof-of-concept adapter prototype in a very short time.

As we already stated before, our current PAL prototype has some limitations, in particular, the only parameter passing semantics available for annotated methods is the *deep-copy* one, and the program sequential semantics is not guaranteed if the class fields are accessed from inside the PAL-annotated methods.

Figure 18 shows an example of PAL prototype usage, namely a program computing the Mandelbrot set. The *Mandelbrot* class uses a `@Parallel` annotation to state that all the `createLines` calls should be computed in parallel, with a parallelism degree equal to 16. Observe that, due to some Java limitations (see below), the programmer must specify `PFFuture` as return type, and consequently return an object of this type. `PFFuture` is a template defined by the PAL framework. It represents a container needed to enable the future mechanism. The type specified as argument is the original method return type. Initially, we tried to have a more transparent mechanism for the future implementation, without any explicit Future declaration. It consisted in the run-time substitution of the return type with a PAL-type inheriting from the original one. In our idea, the PAL-type would have filtered any original type dereferentiation following the *wait-by-necessity* \(^{(42)}\) semantics. Unfortunately, we had to face two Java limitations that limit the current prototype to the current solution. These limitations regard the impossibility to extend some widely used Java BCL classes (String, Integer,...) because they are declared `final`, and the impossibility to intercept all non-private class field accesses.

In the *Main* class, the programmer just asks to transform the *Main* class and the *Mandelbrot* ones with PAL, that is, to process the relevant PAL annotations and to produce an executable IL which exploits parallelism according to the features (hardware and software) of the target architecture where the *Main* itself is being run.
4.2.2 Experimental results

To validate the PAL approach we ran some experiments with the current prototype we developed. In particular, the conducted experiments were aimed at evaluating the effectiveness of PAL approach. It has been evaluated measuring the overhead caused by raising the programming abstraction by means of PAL.

We ran tests for each adapter developed, i.e. both for the multithread adapter and for the JJPF one. In other words, the tests were covering parallel transformations suiting both multiprocessor and cluster architectures. In the former case, we used, as computing resource for the testbed, a hyper-threading bi-processors workstation (Dual Intel Xeon 2GHz, Linux kernel 2.6). In the latter case, instead, we ran the transformed application on a blade cluster (24 machines single PentiumIII-800Mhz processor with multiple Fast Ethernet network, Linux kernel 2.4). In both cases, our test application was a fractal image generator, which computes

![PAL-JJPF Efficiency Comparison](image.png)

**Figure 20:** Mandelbrot computation: efficiency comparison with different image resolution, processing element number and task computational weight.
sections of the Mandelbrot set. The Mandelbrot set is a set of points in
the complex plane, the boundary of which forms a fractal. Mathemati-
cally, the Mandelbrot set can be defined as the set of complex \( c \)-values for
which the orbit of 0 under iteration of the complex quadratic polynomial
\[ x_{n+1} = x_n^2 + c \] remains bounded. A complex number, \( c \), is in the Mandel-
brot set if, when starting with \( x_0 = 0 \) and applying the iteration repeat-
edly, the absolute value of \( x_n \) never exceeds a certain number (that num-
ber depends on \( c \) ) however large \( n \) gets. When computed and graphed on
the complex plane, the Mandelbrot Set has an elaborate boundary, which
does not simplify at any given magnification. This qualifies the boundary
as a fractal. We picked up Mandelbrot because it is a very popular bench-
mark for embarrassingly parallel computation. PAL addresses exactly
these kinds of computations, as it only allows executing remotely meth-
ods not accessing shared (static) variables nor having any kind of side
effects. On the one hand, this obviously represents a limitation, as PAL
cannot compete, as an example, with other approaches supporting plain
loop parallelization. On the other hand, huge amounts of embarrassingly
parallel applications are executed on clusters, workstation networks and
grids. Most of times, the implementation of these applications requires a
significant programming effort, despite being “easy” embarrassingly par-
allel, far more consistent than the effort required to execute the same kind
of application exploiting PAL.

To study in more detail the behavior of the transformed, parallel, ver-
sion of the Mandelbrot application in several contexts, we ran the fractal
generator setting different resolutions (600x400, 1200x800 and 2400x1600)
and task computational weights, starting from 1 up to 40 lines at time.
For each test-bed the total number of lines were fixed, hence when the
task size (number of lines to compute) increases, the total number of tasks
decreases.

The Mandelbrot application, when transformed exploiting the multi-
thread adapter, has been executed only with \texttt{parDegree} parameter set to
1 or 2 (we used a bi-processor machine for the test-bed). Nevertheless, the
multithreaded experiments achieved promising results, as the registered
efficiency with parallel degree 2 is very close to the ideal one, for all the
setting combinations (resolution and compute lines). Since in a multicore solution we have a lower communication impact than in a COW or grid solution, we can point out that this performance should be easily maintained with symmetric multiprocessors even with larger (with four, eight or more cores) processing elements.

After the test with the multithread adapter, we tested also the JJPF one for distributed architectures. We used the very same Mandelbrot source code. PAL transformed it exploiting the JJPF adapter in order to make it able to be executed on distributed workstation network. In this case, we achieved performances definitely close to the ones we achieved with hand written JJPF code (see Figure 20). The Figure shows the result of the experiments with an image resolution of 2400x1600 (other results obtained using different image resolutions gave comparable results) when a different number of processing elements are used (i.e. different values specified to the \texttt{@Parallel(parDegree=...)} annotation).

These results demonstrate that PAL performance strictly depends on the parallel tool targeted by the PAL IL transformation techniques. Actually, the overhead introduced by PAL is negligible.

4.2.3 Learning from PAL experience

Designing, developing and then testing PAL we are taught a lesson by exploiting generative metaprogramming techniques coupled with programmers high-level hints specified at source code level, it is possible to transform a java program that own some properties, enriched with some proper annotations, in a parallel program. The parallelization is obtained through the asynchronous and parallel execution of annotated methods. Annotated method code is transformed in a macro data-flow block that can be dispatched to be executed on the available computational resources. This process executed at run-time directly at intermediate language level, allows to exploit the information available to parallelize the applications with respect both to the parallel tools available on the target execution environment and to the programmer supplied non-functional requirements. A run-time transformation allows to hide most
of parallelization issues. The results we obtained are very encouraging and show that the overhead introduced by PAL is negligible. Nevertheless, the PAL prototype we developed has some limitations. The non-functional requirements are limited to the possibility to indicate the parallelism degree, the parameter passing semantic to PAL-annotated method is limited to deep-copy and the class fields are not accessible from PAL-annotated methods. Furthermore, the programmer has to include an explicit dereferentiation of objects returned by PAL-annotated methods. Finally, current PAL prototype allows only very simple forms of parallelization.

In a sense, PAL has been a proof of concept demonstrating the effectiveness of the approach. With this awareness in mind, we decided to exploit the gained experience to integrate some elements of the PAL approach in our modified muskel framework. The goal is to obtain a framework allowing programmers to develop customizable parallel structured applications which “parts” can be transformed in macro data-flow blocks optimized at run-time according to programmers directives and available hardware and software resources.

4.3 Metaprogramming muskel

PAL proved that, given the existence of a proper metaprogramming runtime support, annotations are a handy way both to indicate which parts of a program must run in parallel and to express non-functional requirements directly in the source code. Such information given as input to PAL metaprogramming engine can be actually exploited to optimize the original annotated code with respect to the running platform and the programmers’ non-functional specifications. Therefore, we decided to apply the main features of PAL approach to our modified muskel implementation. Actually, adapting them to muskel we changed a little bit the approach. Such a change is due to a few motivations. First of all because muskel provides per se a distributed macro data-flow executor whereas PAL exploits external tools for distributed program execution. Moreover, we would like to have a more flexible mechanism for macro data-flow
block generation and management. Finally, we would like to exploit a standard tool for run-time code transformation instead of using ad-hoc tools. As a consequence we decided to use integrate in muskel the AOP model and in particular the AspectJ framework.

The first step in this direction was exploiting AspectJ to implement aspect driven program normalization in muskel. We already introduced normal form and code normalization in Section 2.2.6. Let us to recall it briefly. Normalization consists in transforming an arbitrary muskel program, whose structure is a generic skeleton tree, into a new, equivalent one, whose parallel structure is a farm with a worker made up of the sequential composition of the sequential skeletons appearing in the original skeleton tree taken left to right. This second program is the skeleton program normal form and happens to perform better (with respect to the service time) than the original one in the general case and in the same way in the worst case.

As an example, the code reported in the previous chapter in Figure 11 can be transformed into the equivalent normal form code:

```java
Skeleton main = new Farm(new Seq(f,g));
```

where `Seq` is basically a pipeline whose stages are executed sequentially on a single processor.

Code normalization can be obtained explicitly inserting statements in the source code. This means that programmers must change the source code to use the normal form in place of the non-normal form version of the same program. Exploiting AspectJ we defined a proper aspect dealing with normal form transformation by defining a pointcut on the execution of the `setProgram Manager` method and associating to the pointcut the action performing normal form transformation on the source code in the aspect, such as the one of Figure 22. As a consequence, the programmers can decide whether to use the original or the normal form version of the program just picking up the standard Java compiler or the AspectJ one. The fact the program is left unchanged means the programmer may debug the original bug and have the normal form one debugged too as a consequence, provided the AOP code in the normal form aspect is cor-
public aspect Normalize {
    public boolean ContractSpecified = false;
    public boolean normalized = false;
    // contract is an integer, to simplify ...
    public int contract = 0;

    pointcut calledContract(int i): call(public void Manager.setContract(int)) && args(i);

    void around(int i): calledContract(i){
        ContractSpecified = true;
        contract = i;
        proceed(i);
    }

    pointcut callSetProgram(Skeleton c): call(public void Manager.setProgram(Skeleton)) && args(c);

    void around(Skeleton c):
        callSetProgram(c) {
            normalized = true;
            proceed(new NormalForm(c));
        }

    pointcut callEval(Manager m) : call(public void Manager.eval()) && target(m);

    before(Manager m):callEval(m){
        if(ContractSpecified)
            if(normalized)
                m.setContract(Manager.NormalizeContract(contract));
            else
                m.setContract(Manager.DefaultNormalizedContract);
        }
    }
}

Figure 21: AspectJ code handling performance contracts in muskel.

rect. Moreover, exploiting aspects as discussed above, we handled also related features by means of proper aspects. In fact, in case the programmer provided a performance contract (a parallelism degree, in the simpler case) and then used the AspectJ compiler to ask normal form execution of the program, it turns out to be quite natural imagine a further aspect handling the performance contract consequently. Figure 21 shows the AspectJ aspect handling this aspect. In this case, contracts are stored as soon as they have been issued by the programmer, with the first pointcut, then, in when normalization has been required (second pointcut) and program parallel evaluation is required, the contract is handled consequently
public aspect Normalize {
    pointcut callSetProgram(Skeleton c):
call(public void Manager.setProgram(Skeleton)) && args(c);

    void around(Skeleton c)
        : callSetProgram(c) {
    proceed(new NormalForm(c));
    }
}

Figure 22: AspectJ code modeling normal form in muskel.

(third pointcut), that is, it is either left unchanged or a new contract is derived from the original one according to some normal form related procedure.

The second step consisted in testing the integration of muskel with AspectJ to in a more complex scenario. Hence, we exploited the aspect oriented programming support integrated in muskel in order to develop workflows which structure and processing are optimized at run-time.

4.4 Workflows with muskel

Workflows represents a popular programming model for grid applications (74). In a workflow, programmers express the data dependencies that incurs among a set of blocks, possibly using a DAG. Each block processes input data to produce output data. Workflow schedulers arrange the computations for grid execution in such a way

- all the parallelism implicitly defined through the (absence of) dependencies in the DAG is exploited, and
- available grid resources (processing elements) are efficiently used.

In a sense, a programming model that eases the development of efficient workflow applications can be successfully exploited for the development
of many grid applications. For this reason, we conceived an approach aimed at the implementation of workflows on top of the muskel distributed macro data-flow interpreter. We took into account the execution of workflows on a set of input data items. The set of input data items represents the program input stream. Each item on that stream will be submitted to a full workflow computation. The results of that computation will appear as a data items onto the program output stream. Usually the workflows considered in grids are made of nodes that are computationally complex. Possibly parallel applications processing data contained in one or more input files to produce data in one or more output files (74). We considered a very simple class of workflows: those whose DAG nodes are Java “functions” processing a generic Object input parameters to produce an Object output results.

4.4.1 Aspects to implement workflows

As already stated, we considered workflows processing stream of input data to produce stream of output data. Actually, these are not classical workflows. As discussed in the following, however, classical workflows can be efficiently addressed as well as a side effect of the efficient implementation of stream parallel workflows. This allows to express both parallelism implicit in the workflow definition (and therefore exploited within the computation of a single instance of the workflow) and stream parallelism (parallelism among distinct instances of workflow computation, relative to independent input data items). In order to obtain a macro data-flow graph from the workflow abstract code, we exploited the AspectJ AOP framework (84):

- Programmers express workflows as plain Java code, with the constraint the nodes of the workflow must be expressed using Compute object calls.

- Programmers declare a Manager object passing it an Iterator providing the input tasks. The Manager object completely and transparently takes care of implementing stream parallelism using the muskel distributed macro data-flow interpreter.

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AOP pointcuts and advices are used to intercept the calls to the `compute` methods and to transform such calls into proper fireable macro data-flow instructions submitted to the `muskel` distributed data-flow interpreter.

Sample code used to model workflows is shown in Figure 23. The right part of the Figure lists the Java code modeling the workflow graphically depicted in the left part of the Figure. Multiple results are modeled returning `Vector` objects and multiple input parameters are modeled with a “vararg” `compute` method.\(^1\)

```
Vector resF = (Vector) F.compute(in.elementAt(0));
Object resG1 = G1.compute(resF.elementAt(0));
Object resG2 = G2.compute(resF.elementAt(1),
                        in.elementsAt(1));
Object resH = H.compute(resG1, resG2);
```

**Figure 23:** Sample workflow (left) and relative Java code (right)

More in detail, the calls to `compute` methods are transformed into the submission of a proper (already fireable) macro data-flow instruction to the muskel distributed macro data-flow interpreter modified in such a way a `Future` for the result is immediately returned. If one of the input arguments of the `compute` call is a `Future`, the advice intercepting the `compute` method call takes care of waiting for its actual value to be computed before submitting the macro data-flow instruction to the interpreter.

As input `Future` actual values are only required by the advice right before the workflow node is started, parallelism implicit in the workflow

---

\(^1\) **varargs** have been introduced in Java 1.5 and allow to pass a variable number of arguments (of the same type) to a method; the arguments are referred to in the method body as array elements.
is correctly delegated to the underlying muskel interpreter. As an example, consider the workflow of Figure 23. The functions G1 and G2 are evaluated (their evaluation is requested by the advice to muskel interpreter) sequentially. However, as the first one immediately returns a Future, the second one (also returning a Future) will eventually run in parallel on a distinct remote processing element as outlined in Figure 24. When the evaluation of the H node is requested, the advice intercepting the request will realize two futures are passed as input parameters and therefore it will wait before submitting the node evaluation request to the muskel interpreter up to the moment the two actual values of the “input” Futures are available. Overall, advices transforming calls to compute methods into fireable macro data-flow instructions act as the data-flow matching unit, according to classical data-flow jargon.

The approach suggested here to implement workflows on top of the muskel macro data-flow interpreter presents at least two significant advantages:

- the whole, already existing, efficient and assessed muskel macro data-flow interpreter structure is fully exploited. The muskel interpreter takes completely care of ensuring load balancing, fault tolerance (w.r.t. remote resource faults) and security;

- programmers are only asked to express workflows with elementary Java code, possibly spending some time wrapping workflow node code in Compute objects and declaring a Manager object which is used to supply input data, retrieve output data, control non functional features (e.g. parallelism degree in the execution of the workflow) and to ask the evaluation of the workflow code.

- As in PAL, transformation can be easily disabled. This means that the programmers’ application code can be run through a classical sequential compiler/interpreter suite and debugged using normal debugging tools.
4.4.2 Aspects with muskel: implementation details

In order to be able to express workflows, the programmer must write one class per workflow node. The class has to implement the Compute interface, which is a very simple interface such as:

```java
public interface Compute extends Serializable{
    public Object compute(Object... params);
}
```

The `compute` method is assumed to compute the workflow node re-
results (the returned Object) out of the input parameters params. Then the workflow can be described in a class implementing the Workflow interface, which is defined as follows:

```java
public interface Workflow {
    public Object doWorkflow(Object param);
}
```

As an example, a workflow can be described by the class:

```java
public class WorkFlow1 implements Workflow {
    public Object doWorkflow(Object task) {
        Vector resF = (Vector) F.compute(((Vector)task).elementAt(0));
        Object resG1 = G1.compute(resF.elementAt(0));
        Object resG2 = G2.compute(resF.elementAt(1),
                                 ((Vector)task).elementAt(1));
        Object resH = H.compute(resG1, resG2);
        return resH;
    }
}
```

The code style here is quite close to the style used when programming plain Java applications.

We capture the execution of the Compute calls in the workflow exploiting aspects. The pointcut is defined on the calls of the compute method of any object implementing Compute:

```java
pointcut computeRemotely(Object param[], itfs.Compute code) :
    call(Object itfs.Compute.compute(Object ... )) &&
    !within(execEngine.Engine) &&
    args(param) && target(code);
```

The advice invoked on the pointcut is an around advice such as:

```java
execEngine.Engine eng = new execEngine.Engine();
Future around(Object param[], itfs.Compute code)
    :computeRemotely(param, code) {
    for(int i=0; i<param.length; i++) {
        // reifying each parameter right before call
        if(param[i] instanceof Future) {
            param[i] = ((Future) param[i]).getValue();
        }
    }
    // deliver fireable instruction
    Object future = eng.exec(codice, param);
    // and return the corresponding Future object
    return future;
}
```
It arranges to collect the Compute class name and the input parameters and creates a macro data-flow instruction, which is submitted to the distributed muskel macro data-flow interpreter via the predefined execEngine object instance declared in the aspect class. Input tokens to the macro data-flow instruction that are Future instances rather than plain reified objects, are eventually reified on the fly within the advice. Eventually, a Future object is returned. It can be eventually used to retrieve the actual data computed by the distributed interpreter during the compute call. In particular, Future interface provides two methods: a getValue() method to get the actual value of the Future, possibly waiting for the completion of the corresponding computation, and a boolean isReady() method to test whether the computation producing the actual value of the Future is already terminated.

As a whole, the procedure just described models an asynchronous execution of the macro data-flow instructions implementing the workflow nodes. It allows to fully exploit the parallelism intrinsic to the workflow, by properly using Futures.

As already stated, we are interested not only in the exploitation of parallelism within the evaluation of a single workflow instance, but also in exploiting the parallelism among different instances of workflows run on distinct input data sets. In order to support stream parallelism, we provide the programmer with a StreamIterator manager. This manager takes as parameters an Iterator (providing the input data sets to be processed by the Workflow) and a Workflow. It provides a method to compute the whole bunch of inputs, as well as a method to get an Iterator that can be used to retrieve workflow results. Using the StreamIterator manager, the main code relative to our example can therefore be expressed as follows:

```
public static void main(String[] args) {
   // workflow to be used (userdef)
   Workflow wf = new Workflow1();

   // provide the input tasks via an iterator (userdef)
   InTaskIterator intIt =
      new InTaskIterator();

   // declare the manager
   Manager mgr = new StreamIterator(wf,intIt);

   // start parallel computation
```
The main task of the StreamIterator manager is to invoke execution of the parameter Workflow instances on all the input data sets provided by the Iterator. This is achieved exploiting a proper Thread pool and activating one thread in the pool for each independent workflow computation. Then, the AOP procedure illustrated above intercepts the calls to compute methods and arrange to run them in parallel through the muskel distributed macro data-flow interpreter.

4.4.3 Experiments

In order to prove the effectiveness of the approach, we tested it making some experiments on a distributed computing architecture (a network of workstations, actually). We directly used Java (version 1.5) accessible via plain secure shell (ssh/scp) rather than with other more sophisticated grid middleware. It is worth to point out that the tests have not been conducted to evaluate the scalability of plain muskel, that has actually already been demonstrated, as discussed in (58). Rather, the tests have been performed in order to give an estimation of the overhead introduced by aspectj transformations.

In fact, the only difference between plain muskel and the system proposed here, able to execute workflows on top of muskel, lies in the way the fireable instructions are provided to the distributed data-flow interpreter of muskel. Actually, in plain muskel, fireable instructions are retrieved from a compiled representation of a data-flow graph. In particular, each time a new token arrives to a macro data-flow instruction in
the graph (either from the input stream or as the result of the distributed computation of another macro data-flow instruction) the target data-flow instruction is checked for “fireability” and, possibly, delivered to the distributed macro data-flow interpreter. The time spent is in the sub-micro second range (only considering net time, not taking into account time spent to copy parameters in memory during the interpreter call). When executing workflows according to the approach discussed here, instead, fireable instructions are generated by means of the aspectj tool. In particular, they come from the “advice” invoked on the “pointcut” intercepting the **compute** calls. In order to estimate the overhead introduced by using these Aspect Oriented Techniques we measured the time spent to intercept the **compute** calls and to transform them in macro data-flow blocks. The measurement results are shown in the following table (times are in milliseconds):

![Graph showing efficiency of the muskel/aspect workflow prototype](image)

**Figure 25**: Efficiency of the muskel/aspect workflow prototype
<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>Average</td>
<td>23.09</td>
<td>Minimum</td>
<td>19</td>
</tr>
<tr>
<td>Standard deviation</td>
<td>3.01</td>
<td>Maximum</td>
<td>27</td>
</tr>
</tbody>
</table>

These values are relative to an Intel dual-core machine (2 GHz Core 2 Duo machine), running Mac OS/X 10.4, Java 1.5.0_07, AspectJ 1.5.4 with AspectJ tools 1.4.2 and Eclipse 3.2.2. On the same machine, delivering a fireable instruction to the macro data-flow interpreter with plain `muskel` requires a time average of 0.004 milliseconds. The difference in the times is not surprising: in the former case, we go through pure metaprogramming tools and we “interpret” each call, while in the latter we use plain (compiled) Java to handle each one of the calls.

Therefore, we can conclude the average 23 milliseconds represent the pure overhead spent each time a new fireable instruction has to be computed (i.e. each time one of the workflow `Compute` nodes is computed). The time spent in `Future` reification (i.e. filling the object placeholder with the computed value, once available), instead, is negligible (this not taking into account the time spent to wait for actual production of `Future` values, of course). This allows us to conclude that the parallel execution of workflows on top of `muskel` slightly increases the grain required to achieve almost ideal scalability.

In fact, Figure 25 shows how with suitable grain of the workflow nodes (i.e. of the `Compute` functions) efficiency close to the ideal one is achieved.

### 4.5 Differences between the two approaches

As we already stated before, both PAL and AspectJ enriched `muskel` (AEM) were conceived, designed and implemented to provide a proof-of-concept of our metaprogramming approach to structured parallel programming. Actually, they enforce code parallelization via a hints-driven code transformation. Hints are provided by programmers in the form of java annotations (PAL) and AspectJ rules AEM. Even if the two frameworks attain the same idea, they are slightly different. The main differences between the two frameworks are:
• In AEM there is a sharp-cut distinction between the “control” and “business” code, actually contained in separate files, whereas with PAL programmers write business code and annotations (that behaves as control code) inside the same file.

• PAL was conceived to exploit method-level parallelism: through a simple program enrichment process, programmers choose which Java methods-call should be transformed in asynchronous ones, i.e. PAL allows to add parallelism to legacy java code with a minimal intervention. Instead, in AEM programmers have to implement their application as a workflow.

• PAL provides a fixed number of annotations (hence a very limited number of action can be performed) that an adapter-based architecture exploits to transform bytecode at run-time. The transformation process depends, in a way, on the adapter used. In AEM the code transformation policies implementation is based on AspectJ, the most widely diffused tool for aspect oriented programming, which offers a rich set of mechanisms for customizing the “aspectization” process. As a consequence, the programmers can customize/optimize/change the transformation process by simply modifying the aspects (without a direct code update).
In this Chapter we presented two results, about the exploitation of metaprogramming techniques in structured parallel programming environment. We exploited those techniques in order to generate and optimize at run-time macro data-flow blocks without directly dealing with their low-level management. First we presented a new technique for high-level parallel programming based on the introduction of a Parallel Abstraction Layer (PAL). PAL does not introduce a new parallel programming model, but actually exploits the programmer knowledge provided through annotations to restructure at run-time the application, hiding most of parallelization issues, once it notice the information about the running platform. This process is executed directly at intermediate language level. This allows to have a portable code transformation mechanism without paying a complete code recompilation for each change in the code. In order to have a proof-of-concept of the approach we developed a PAL Java prototype and we used it to perform some experiments. The results are very encouraging and show that the overhead introduced by PAL is negligible, while keeping the programmer effort to parallelize the code negligible. Then we presented the other result we obtained integrating the AspectJ framework with our modified muskel. We described how AOP techniques can be seamlessly used to transform a very basic kind of workflows in such a way they can be executed on distributed target architectures through the muskel macro data-flow interpreter. How AOP techniques allow to completely separate the concerns relative to parallelism exploitation and application functional core. In particular, the same application code used to perform functional debugging on a single, sequential machine may be turned into parallel code by adding aspects, compiling it through AspectJ and then running it on the muskel run-time support. The way used to write workflow code is quite basic Java programming. Workflow components must implement a simple interface, and programmers are explicitly required to provide them as side effect free sequential components. The experiments conducted show that the approach is perfectly feasible and that actual speedups can be achieved provided that the workflow nodes are medium to coarse grain.
Chapter 5

Behavioural Skeletons

Chapter road-map  In this chapter we present Behavioural Skeletons, an approach, we contribute to conceive and validate, aimed at providing programmers with the ability to implement autonomic grid component-based applications that completely take care of the parallelism exploitation details by simply instantiating existing skeletons and by providing suitable, functional parameters. The model has been specifically conceived to enable code reuse and dynamicity handling. We start describing (Section 5.1) how component-based application can ease the task of developing grid applications. Then we outline the Grid Component Model (Section 5.2) with respect to its autonomic features. After we present the Behavioural Skeletons model (Section 5.4), a set of noteworthy Behavioural Skeletons (Section 5.5) and their implementation (Section 5.6). At the end of chapter we describe a set of experiment we conducted to validate the Behavioural Skeletons model (Section 5.7).

5.1 Components to simplify Grid programming

Developing grid applications is even more difficult than programming traditional parallel applications. This is due to several factors as, the heterogeneity of resources, their worldwide distribution, their dynamic recruiting and releasing. Indeed, when programming Grid applications neither the target platforms nor their status are fixed (82).
As a consequence, grid applications need to dynamically adapt to the features of the underlying architecture in order to be efficient and/or high performance (19). In recent years, several research initiatives exploiting component technology (52) have investigated the area of component adaptation, i.e. the process of changing the component for use in different contexts. This process can be either static or dynamic.

The basic use of static adaptation covers straightforward but popular methodologies, such as copy-paste, and OO inheritance. A more advanced usage covers the case in which adaptation happens at run-time. These systems enable dynamically defined adaptation by allowing adaptations, in the form of code, scripts or rules, to be added, removed or modified at run-time (37). Among them is worth to distinguish the systems where all possible adaptation cases have been specified at compile-time, but the conditions determining the actual adaptation at any point in time can be dynamically changed (23). Dynamically adaptable systems rely on a clear separation of concerns between adaptation and application logic. This approach has recently gained increased impetus in the grid community, especially via its formalization in terms of the Autonomic Computing (AC) paradigm (22, 24, 77). The AC term is emblematic of a vast hierarchy of self-governing systems, many of which consist of many interacting, self-governing components that in turn comprise a number of interacting, self-governing components at the next level down (83). An autonomic component will typically consist of one or more managed components coupled with a single autonomic manager that controls them. To pursue its goal, the manager may trigger an adaptation of the managed components to react to a run-time change of application QoS requirements or to the platform status.

In this regard, an assembly of self-managed components implements, via their managers, a distributed algorithm that manages the entire application. Several existing programming frameworks aim to ease this task by providing a set of mechanisms to dynamically install reactive rules within autonomic managers. These rules are typically specified as a collection of when-event-if-cond-then-act clauses, where event is raised by the monitoring of component internal or external activity (e.g. the
component server interface received a request, and the platform running a component exceeded a threshold load, respectively); \textit{cond} is an expression over component internal attributes (e.g. component life-cycle status); \textit{act} represents an adaptation action (e.g. create, destroy a component, wire, unwire components, notify events to another component’s manager). Several programming frameworks implement variants of this general idea, including ASSIST \cite{19,113}, AutoMate \cite{93}, SAFRAN \cite{66}, and finally the forthcoming CoreGrid Component Model (GCM) \cite{52}. The latter two are derived from a common ancestor, i.e. the Fractal hierarchical component model \cite{39}. All the named frameworks, except SAFRAN, are targeted to distributed applications on grids.

Though such programming frameworks considerably ease the development of an autonomic application for the grid (to various degrees), they rely fully on the application programmer’s expertise for the set-up of the management code, which can be quite difficult to write since it may involve the management of black-box components, and, notably, is tailored for the particular component or assembly of them. As a result, the introduction of dynamic adaptivity and self-management might enable the management of grid dynamism, and uncertainty aspects but, at the same time, decreases the component reuse potential since it further specializes components with application specific management code.

From the point of view of issues to address for designing and developing next generation structured parallel programming systems, this is a big problem. Indeed, if on the one hand making components adaptive addresses the issue of handling dynamicity (issue number VII), on the other hand it impairs the code reuse (issue number V). In this chapter we cope with this problem proposing \textit{Behavioural Skeletons} as a novel way to describe autonomic components in the GCM framework. We contributed significantly to their conception, design and implementation together with other researchers, co-authored of the papers \cite{16,17} in which we presented this model. My personal contribution has mainly concerned the definition of the task farm Behavioural Skeleton as well as the implementation of that skeleton within GridCOMP.

\textit{Behavioural Skeletons} aim to describe recurring patterns of compo-
nt assemblies that can be (either statically or dynamically) equipped with correct and effective management strategies with respect to a given management goal. Behavioural Skeletons help the application designer to i) design component assemblies that can be effectively reused, and ii) cope with management complexity by providing a component with an explicit context with respect to top-down design (i.e. component nesting).

5.2 GCM: the Grid Component Model

GCM is a hierarchical component model explicitly designed to support component-based autonomic applications in highly dynamic and heterogeneous distributed platforms, such as grids. It is currently under development by the partners of the EU CoreGRID Network of Excellence\(^1\). A companion EU STREP project, GridCOMP \(^2\) is going to complete the development of an open source implementation of GCM (preliminary versions are already available for download as embedded modules in the ProActive middleware suite)\(^3\). GCM builds on the Fractal component model \(^{39}\) and exhibits three prominent features: hierarchical composition, collective interactions and autonomic management. We participate to both the projects (CoreGrid & GridComp) and collaborate for the design and development of GCM, in particular in the context of autonomic management. The full specification of GCM can be found in \(^{52}\).

Hierarchical composition \quad \text{As in fractal, a GCM component is composed of two main parts: the membrane and the content. The membrane is an abstract entity that embodies the control behavior associated with a component, including the mediation of incoming and outgoing invocations of content entities. The content may include either the code directly implementing functional component behavior (primitive) or other components (composite). In the latter case, the included components are referred}

\(^1\)http://www.coregrid.net
\(^2\)http://gridcomp.ercim.org
\(^3\)http://www-sop.inria.fr/oasis/ProActive
as the inner components. GCM components, as Fractal ones, can be hier-
archically nested to any level. Component nesting represents the implemented by relationship. Composite components are first class citizens in GCM and, once designed and implemented, they cannot be distinguished from primitive, non-composite ones.

**Collective interactions** The Grid Component Model allows component interactions to take place with several distinct mechanisms. In addition to classical “RPC-like” use/provide ports (or client/server interfaces), GCM allows data, stream and event ports to be used in component interaction. Both static and dynamic wiring between dual interfaces is supported. Each interface may expose several operations of different types. Furthermore, collective interaction patterns (communication mechanisms) are also supported. In particular, composite components may benefit from customizable one-to-many and many-to-one functional interfaces to distribute requests arriving to one component’s port to many inner components and gather requests from many inner components to a single outgoing port.

**Autonomic management** Autonomic management aims to attack the complexity which entangles the management of complex systems (as applications for Grids are) by equipping their parts with self-management facilities. GCM is therefore assumed to provide several levels of autonomic managers in components, that take care of the non-functional features of the component programs. GCM components thus have two kinds of interfaces: functional and non-functional ones. The functional interfaces host all those ports concerned with implementation of the functional features of the component. The non-functional interfaces host all those ports needed to support the component management activity in the implementation of the non-functional features, i.e. all those features contributing to the efficiency of the component in obtaining the expected (functional) results but not directly involved in result computation. Each GCM component therefore contains an Autonomic Manager (AM), interacting with other managers in other components via the component non-
functional interfaces. The AM implements the autonomic cycle via a simple program based on the reactive rules described above. In this, the AM leverages on component controllers for the event monitoring and the execution of reconfiguration actions. In GCM, the latter controller is called the Autonomic Behaviour Controller (ABC). This controller exposes server-only non-functional interfaces, which can be accessed either from the AM or an external component that logically surrogates the AM strategy. From the point of view of autonomic features, the GCM components exhibiting just the ABC are called passive, whereas the GCM components exhibiting both the ABC and the AM are called active.

5.3 Describing Adaptive Applications

The architecture of a component-based application is usually described via an ADL (Architecture Description Language) text, which enumerates the components and describes their relationships via the used-by relationship. In a hierarchical component model, such as the GCM, the ADL describes also the implemented-by relationship, which represents the component nesting.

However, the ADL supplies a static vision of an application, which is not fully satisfactory for an application exhibiting autonomic behavior since it may autonomously change behavior during its execution. Such change may be of several types:

- **Component lifecycle.** Components can be started or stopped.

- **Component relationships.** The used-by and/or implemented-by relationships among components are changed. This may involve component creation/destruction, and component wiring alteration.

- **Component attributes.** A refinement of the behavior of some components (which does not involve structural changes) is required, usually over a pre-determined parametric functionality.

In the most general case, an autonomic application may evolve along adaption steps that involve one or more changes belonging to these three
classes. In this regard, the ADL just represents a snapshot of the launch
time configuration.

The evolution of a component is driven by its AM, which may request
management action with the AM at the next level up in order to deal
with management issues it cannot solve locally. Overall, it is a part of a
distributed system that cooperatively manages the entire application.

In the general case, the management code executing in the AM of a
component depends both on the component’s functional behavior and
on the goal of the management. The AM should also be able to cooper-
ate with other AMs, which are unknown at design time due to the nature
of component-based design. Currently, programming frameworks sup-
porting the AC paradigm (such as the ones mentioned in Section 5.1) just
provide mechanisms to implement management code. This approach has
several disadvantages, especially when applied to a hierarchical compo-
nent model:

- The management code is difficult to develop and to test since the
  context in which it should work may be unknown.
- The management code is tailored to the particular instance of the
  management elements (inner components), further restricting the
  component reusability possible.

5.4 Behavioural Skeletons

Behavioural Skeletons aim to abstract parametric paradigms of the GCM
components assembly, each of them specialized to solve one or more man-
agement goals belonging to the classical AC classes, i.e. configuration,
optimization, healing and protection.

They represent a specialization of the algorithmic skeleton concept for
component management. Behavioural Skeletons, as algorithmic skele-
tons, represent patterns of parallel computations (which are expressed in
GCM as graphs of components), but in addition they exploit skeletons’
inherent semantics to design sound self-management schemes of parallel
components.
As a byproduct, Behavioural Skeletons allow categorization of GCM designers and programmers into three classes. They are, in increasing degree of expertise and decreasing cardinality:

1. **GCM users**: they use Behavioural Skeletons together with their pre-defined AM strategy. In many cases they should just instantiate a skeleton with inner components, and get as result a composite component exhibiting one or more self-management behaviors.

2. **GCM expert users**: they use Behavioural Skeletons overriding the AM management strategy. However, the specialization does not involve the ABC and thus does not require specific knowledge about the GCM membrane implementation.

3. **GCM skeleton designers**: they introduce new Behavioural Skeletons or classes of them. To this end, the design and development of a brand new ABC might be required. This may involve the definition of new interfaces for the ABC, the implementation of the ABC itself, together with its wiring with other controllers, and the design and wiring of new interceptors. Obviously, this requires quite a deep knowledge of the particular GCM implementation.

Due to the hierarchical nature of GCM, Behavioural Skeletons can be identified with a composite component with no loss of generality (identifying skeletons as particular higher-order components (73)).

Since skeletons are fully-fledged GCM components, they can be wired and nested via standard GCM mechanisms. From the implementation viewpoint, a Behavioural Skeleton is a partially defined composite component, i.e. a component with placeholders, which may be used to instantiate the skeleton. As sketched in Figure 26, there are three classes of placeholders:

1. The functional interfaces $S$ and $C$ that are GCM membrane controllers (thus objects).

2. The AM that is a particular inner component. It includes the management plan, its goal, and exported non-functional interfaces.
3. Inner component $W$, implementing the functional behavior.

The orchestration of the inner components is implicitly defined by the skeleton type. In order to instantiate the skeleton, placeholders should be filled with suitable entities. Observe that just entities in the former two classes are skeleton specific. Indeed, the placeholders of the third class, representing the inner components implementing the functional behavior, are filled with user-defined components. The entities part of the first two classes characterize the composite component as a higher order one orchestrating the entities of the third class; like traditional skeletons are higher order functions taking as parameter user specified functions.

Behavioural Skeletons usage helps designers in two main ways. First, the application designer benefits from a library of skeletons, each of them carrying several pre-defined, efficient self-management strategies. Then, the component/application designer is provided with a framework that helps both the design of new skeletons and their implementation.

In both cases two features of Behavioural Skeletons are exploited: on the one hand, the skeletons exhibit an explicit higher-order functional semantics that delimits the skeleton usage and definition domain. On the other hand, the skeletons describe parametric interaction patterns and can be designed in such a way that parameters affect non-functional behavior but are invariant for functional behavior.

5.5 A Basic Set of Behavioural Skeletons

Here we present a basic set of Behavioural Skeletons. Despite their simplicity, they cover a significant set of parallel computations of common usage.

The presented Behavioural Skeletons springs from the idea of functional replication. Let us assume these skeletons have two functional interfaces: a one-to-many stream server $S$, and a many-to-one client stream interface $C$ (see Figure 26). The skeleton accepts requests on the server interface; and dispatches them to a number of instances of an inner component $W$, which may propagate results outside the skeleton via $C$ inter-
face. Assume that replicas of $W$ can safely lose the internal state between different calls. For example, the component has just a transient internal state and/or stores persistent data via an external database component.

**Farm**  A task farm processes a stream of tasks $\{x_0, \ldots, x_m\}$ producing a stream of results $\{f(x_0), \ldots, f(x_m)\}$. The computation of $f(x_i)$ is independent of the computation of $f(x_j)$ for any $i \neq j$ (the task farm parallel pattern is often referred to as the “embarrassingly parallel” pattern). The items of the input stream are available at different times, in general: item $x_i$ is available $t \geq 0$ time units after item $x_{i-1}$ was available. Also, in the general case, it is not required that the output stream keeps the same ordering as the input stream, i.e. item $f(x_i)$ may be placed in the output stream in position $j \neq i$. In this case, in our farm Behavioural Skeleton, a stream of tasks is absorbed by a *unicast* $S$. Then each task is computed by one instance of $W$ and the result is sent to $C$, which collects results according to a *from-any* policy. This skeleton can be equipped with a self-optimizing policy as the number of $W$ can be dynamically changed in a sound way since they are stateless. The typical QoS goal is to keep a given limit (possibly dynamically changing) of served requests in a time frame. Therefore, the AM just checks the average time tasks need to traverse the skeleton, and possibly reacts by creating/destroying instances of $W$, and wiring/unwiring them to/from the interfaces.

**Data-Parallel** the task farm Behavioural Skeleton can be conveniently and easily adapted to cover other common patterns of parallel computation. For example, data parallel computations can be captured by simply modifying the behavior associated with the $S$ and $C$ interfaces. In a data parallel computation a stream of tasks is absorbed by a *scatter* $S$. Each of the tasks appearing is split into (possibly overlapping) partitions, which are distributed to replicas of $W$ to be computed. The results computed by the $W$ are *gathered* and assembled by $C$ in a single item, which is eventually delivered onto the output stream. As in the previous case, the number of $W$ can be dynamically changed (between different requests) in a sound way since they are stateless. In addition to the previous case,
The task farm (and data parallel) Behavioural Skeleton just outlined can be easily modified to the case in which the S is an RPC interface. In this case, the C interface can be either an RPC interface or missing. Also, the stateless functional replication idea can be extended to the stateful case by requiring inner components W to expose suitable methods to serialize, read and write the internal state. A suitable manipulation of the serialized state enables the reconfiguration of workers (also in the data-parallel scenario (19)).

Anyway, in order to achieve self-healing goals some additional requirements on the GCM implementation level should be enforced. They are related to the implementation of GCM mechanisms, such as component membranes and their parts (e.g. interfaces) and messaging system. At the level of interest, they are primitive mechanisms, in which correctness and robustness should be enforced ex-ante, at least to achieve some of the described management policies.

The process of identification of other skeletons may benefit from the work done within the software engineering community, which identified some common adaptation paradigms, such as proxies (99), which may be interposed between interacting components to change their interaction relationships; and dynamic wrappers (111). Both of these can be used for self-protection purposes. As an example, a couple of encrypting proxies can be used to secure a communication between components. Wrapping can be used to hide one or more interfaces whether a component is deployed into an untrusted platform.
5.6 Autonomic Components: design and implementation

The two main characteristics of autonomic components are the ability to self-manage and to cooperate with other autonomic components to achieve a common goal, such as guaranteeing a given behavior of an entire component-based application. In the light of this, viewing the management of a single component as an atomic feature enables design of its management (to a certain extent) in isolation. The management of a single component is therefore considered a logically centralized activity. Components will be able to interact with other components according to well-defined protocols described by management interaction patterns, which are established by the component model.

5.6.1 The management of a GCM component

The management of a single component is characterized by its ability to make non-trivial decisions. Thus GCM components are differentiated as being passive or active, with the following meanings:

**Passive** A component exposes non-functional operations enabling introspection (state and sensors) and dynamic reconfiguration. These operations exhibit a parametric but deterministic behavior. The operation semantics is not underpinned by a decision making process (i.e. does not implement any optimization strategy), but can only be constrained by specific pre-conditions that, when not satisfied, may nullify an operation request. All components should implement at least a reflection mechanism that may be queried about the list and the type of exposed operations.

**Active** A component exhibits self-managing behavior, that is a further set of autonomic capabilities built on top of passive level functionality. The process incarnates the autonomic management process: monitor, analyze, plan, execute. The monitoring phase is supported by introspective operations, while the executing phase is supported by re-configuring operations described above.
In the architecture of GCM components, these two features are implemented within the Autonomic Behaviour Controller (ABC) and Autonomic Manager (AM), respectively. Since the management is a logically centralized activity, a single copy of each of them can appear in a component. Notice that, this does not prevent a parallel implementation of them for different reasons, such as fault-tolerance or performance. A passive component implements just the ABC, whereas an active component implements both the ABC and the AM. The following relationship holds

\[
\text{Comp} <: \text{PassiveComp} <: \text{ActiveComp}
\]

where \(<:\) is a subtyping relation. This is described in the GCM specification by increasing values of conformance levels [52].

**GCM Passive Autonomic Components** The ABC and the AM represent two successive levels of abstraction of component management. As mentioned above, the ABC implements operations for component reconfiguration and monitoring. The design of these operations is strictly related to membrane structure and implementation, and therefore the choice of implementing the ABC as a controller in the membrane was the more obvious and natural. Within the membrane, the ABC can access all the services exposed by sub-component controllers, such as that related to life cycle and binding, in order to implement correct reconfiguration protocols. In general, these protocols depend on component structure and behavior. However, in the case of Behavioural Skeletons they depend almost solely on the skeleton family and not on the particular skeleton. In this regard, the ABC effectively abstracts out management operations for Behavioural Skeletons.

As we presented Behavioural Skeletons based on the idea of functional replication, we show the details of these skeletons. In this case, the reconfiguration operations require the addition/removal of workers as well as the tuning of distribution/collection strategies used to distribute and collect tasks and results to and from the workers. The worker addition and/or removal operations can be used to change the parallelism
degree of the component as well to remap workers on different processing elements and/or platforms. The distribution/collection tuning operations can be used to throttle and balance the resource usage of workers, such as CPU, memory and IO. The introspection operations involve querying component status with respect to one or more pre-defined QoS metrics. The component status is generally obtained as a harmonized measure involving component status and inner component status.

In the following we describe in some detail the implementation of a reconfiguration and an introspection operation.

**add_worker(k)**  
*Semantics:* Add $k$ workers to a skeleton based on the functional replication.

1. **Stop.** The ABC requires the *Lifecycle Controller* (LC) to stop all the components. To this end, the LC retrieves from the *Content Controller* (CC) the list of inner components $W_1 \cdots W_n$, and then issues a *stop* on them.

2. **Type Inspection.** All the $W_1 \cdots W_n$ have the same type. The ABC retrieves from the CC the list of inner components $W_1 \cdots W_n$, then retrieves $\text{TypeOf}(W_1)$.

---

**Figure 26:** GCM: membrane and content (CC is the content controller, LC the lifecycle controller and BC is the binding controller).
3. **New.** One or more new inner components of type `TypeOf(W_1)` are created.

4. **Bind.** The component server interface `S` is wired to newly created `W_{n+1} \cdots W_{n+k}` inner components via the *Binding Controller* (BC). `W_{n+1} \cdots W_{n+k}`, in turn, wire their client interfaces to the component collective client interface `C`. The process requires the inspection of the types of the interfaces of `W_1` that is used again as a template for all `W_i`.

5. **Restart.** The ABC requires the LC to re-start all the components.

6. **Return.** Return a failure code if some of the previous operations failed (e.g. inner components do not implement stop/start operations); return success otherwise.

**get\_measure(m)**  
*Semantics:* Query the component about the current status of the measure `m`, which may depend on the status of the inner components (possibly involving other measures) and the membrane status.  
*Examples:* Transactions per unit time, load balancing, number of up-and-running workers, etc.

1. **Collect Workers’ Measures.** The ABC retrieves from the CC the list of inner components `W_1 \cdots W_n`, then issues a `get\_measure(m)` on each.

2. **Collect Membrane Measures.** The ABC queries membrane sensors relating to the particular metric `m`.

3. **Harmonize Measures.** Measures acquired from workers and from the membrane are harmonized by using a `m`-dependent function (e.g. average, maximum, etc.).

4. **Return.** Return a failure code if some of the previous operations failed (e.g. sensor not implemented in inner components); return monitor information otherwise.
GCM Active Autonomic components  The operations implemented in the ABC can be arbitrarily complex; however, they do not involve any decision making process. In general, each of them implements a protocol that is a simple list of actions. On the contrary, the AM is expected to enforce a contractually specified QoS. To this end the AM should decide if a reconfiguration is needed, and if so, which reconfiguration plan can re-establish contract validity (13). Furthermore, as we shall see in Section 5.6.2 the AM should also determine if the contract violation is due to the managed component or is the byproduct of other components’ malfunction. The architecture of an active GCM component is shown in Figure 27.

Figure 27: Left) GCM active component architecture. Right) ABC and AM interaction.

The AM accepts a QoS contract\(^4\), which is currently defined as pair \(\langle V, E \rangle\), where \(V\) is a set of variables representing the measures the AM can evaluate (via the ABC), and \(E\) is a mathematical expression over these variables that might include the \(\min\) and \(\max\) operator over a finite domain. The set of \(V\) determines the minimum set of measures the AM should be able to monitor to accept the contract. The \(E\) encodes the con-

\(^{4}\)the notion of QoS contract is still the subject of further investigations and possible refinements. The one discussed here is the bare minimum necessary to discuss AM behavior and implementation.
straints and goal the AM is required to pursue. This encoding can be realized in many different ways provided $E$ can be evaluated in finite time and possibly quite efficiently.

Having accepted a QoS contract, the AM iteratively checks its validity, and in the case that it appears broken, evaluates a number of pre-defined reconfiguration plans. Each reconfiguration plan consists of a sequence of actions (to be executed via the ABC), and a QoS forecast formula. This formula allows the value of a subset of $V$ after the reconfiguration to be forecast. The AM instantiates in turn all reconfiguration plans obtaining, for each plan, a set of forecast values. A plan is marked as valid if the set of $V$ updated with forecast values satisfies the QoS contract. Among the valid plans, the AM heuristically chooses the reconfiguration plan to be executed. If no reconfiguration plan is valid, an exception is raised.

As is clear, the main difficulty in the AM definition is the specification of a reconfiguration plan. In the general case, the reconfiguration plans, and especially their forecast formula, are strictly related to the behavior of a particular component. As discussed in Section 5.3, Behavioural Skeletons enable the definition of reusable reconfiguration plans by categorizing and restricting component behavior in families and skeletons.

### 5.6.2 Cooperative management

The ultimate goal of QoS management is to guarantee programmer intentions despite software and environmental instabilities and malfunctions. To this end, the management of a whole system should be coordinated to achieve a common goal. In general, we envisage a component-based system as a graph, whose nodes are components, and edges are relations among them, such as data dependency, management, geographic locality, etc. Different relations can be kept distinct by a proper labeling of edges. Here we restrict the focus to two relations which are of particular interest for GCM: used by and the implemented by (see Section 5.3). Since the GCM is a hierarchical model, the nesting relation naturally defines the implemented by relationship. In particular, the application structure along the nesting relation describes a tree whose nodes represent components.
(leaves are primitive components) and edges represent their nesting. In this case, the management of a composite component \( C \) is cooperatively performed by the \( \text{AM}_C \) of the component itself and the \( \text{AM}_{C_i} \) of the child components \( C_i, i = 1..n \). In the case where inner components are passive, the cooperation is really one of control by the outer component: services exposed by the \( \text{ABC}_{C_i} \) are called by the \( \text{ABC}_C \).

Conceptually, non-functional properties modeling run-time behavior of the whole hierarchy can be synthesized in a bottom-up fashion: the behavior of a composite component depends on the behavior of its nested components. Management actions and QoS contracts should be projected along the tree in a top-down fashion: the users usually would like to declare a global goal they expect from an application. This matches the idea of submitting a contract at the root of tree. A fully autonomic system should automatically split the global goal into sub-goals that should then be forced on inner components.

On the whole, each GCM component enforces local decisions. When a contract violation is detected, its AM tries autonomously to re-establish the contract to a valid status by re-configuring its membrane or inner components. In the event that it cannot (no valid plan), it raises an event to its father component, thus increasing the extent of the reconfiguration. The overall behavior enforces the maximum locality of reconfigurations, which is a highly desirable property in a distributed system, since it eases the mapping of components onto the network of platforms that usually exhibit a hierarchical nature in terms of uniformity of resources and latency/bandwidth of networks (cluster of clusters).

Observe that cooperation between components is unavoidable even in very simplistic applications. Let us consider an example:

**Producer-filter-consumer** Let us assume that the application sketched in Figure 28 has the final goal to generate, render, and display a video with a given minimum number of frames/sec \( (FPS > k) \). The contract is split into three identical contracts since the property should be enforced on all stages in order to hold globally. The rendering (filter) has been parallelized since it is the most CPU-demanding stage. Two common prob-
lems of such applications are a transient overload of platform where $W_1 \cdots W_n$ are running, or an increased complexity of scene to be rendered. These events may lead to a violation of QoS contract at the AM$_F$. In this case, it may increase the number of workers (mapped on fresh machines) to deal with the insufficient aggregate power of already running resources. In many cases this will locally solve the problem. However, a slightly more sophisticated contract should consider also the input and output channels. In particular the filter stage might be not rendering enough frames because it does not receive enough scenes to render. In this case the AM$_F$ can detect the local violation, but cannot locally solve the problem. As a matter of fact, no plan involving a change of parallelism degree can solve this problem. AM$_F$ can just signal the problem to a higher level AM$_A$, which can try to remap the input channel to a faster link, or simply signal to the end user that the contract is not satisfied.

5.7 Experiments

In order to validate the Behavioural Skeletons approach, we conducted some experiments with the current prototype of the GCM. It is under development in the GridCOMP STREP project (3). The prototype, which is being developed on top of ProActive middleware (108), includes almost all of the features described in this chapter. All the experimental data
Figure 29: Reconfiguration overhead: Stop.

Figure 30: Reconfiguration overhead: New.
are measured on the application shown in Figure 28 that we already presented in the previous section. It basically is a three-stages pipeline in which the second stage consists in a farm of workers processing the images coming from the first stage, and delivering them to the third stage. The experiments mainly aim to assess the overhead due to management and reconfiguration of GCM components. For the sake of reproducibility, the experiments have been run on a cluster instead of a more heterogeneous grid. The cluster includes 31 nodes (1 Intel P3@800MHz core per node) wired with a fast Ethernet. Workers are allocated in the cluster in a round robin fashion with up to 3 workers per node (for a total of 93 workers). Note however, the very same experimental code can run on any distributed platform supported by the ProActive middleware.

Figures 29, 30, and 31 respectively show the time spent on the farm Behavioural Skeleton (filter) for the stop, new and restart Autonomic Behavioural Controller (ABC) services described in Section 5.6.1. This time consists in application overhead, since in current implementation none of the workers can accept new tasks during the process. In the figures, a point $k$ in the X-axis describes the overhead due to stop/new/restart in the
adaptation of the running program from a \( k \) to \( k + 1 \) worker configuration. As highlighted by the curves in Figure 29 and 31, the overhead of stop and restart is linear with respect to the number of workers involved in the operations. This is mainly due to a linear time barrier within the Life cycle Controller (LCC), which is an inherent part of the underlying ProActive middleware. Indeed, in the current implementation the LCC sequentially stops all the workers. Note that adaptation process does not strictly require such a barrier. Both stopping all the workers and linear time synchronization are peculiarities of the current GCM implementation on top of the ProActive middleware, and not of the farm Behavioural Skeleton, which can be implemented avoiding both problems. In addition, the creation of a new worker can be executed, at least in principle, outside the critical path by using a speculative creation.

Figure 30 shows the time spent for the new Autonomic Behavioural Controller (ABC) operation (see Section 5.6.1). Again, in this case, the time is overhead. The experiment measures the time required for the creation of a single worker, and thus the times measured are almost independent of the number of workers pre-existing the new one.

As highlighted by the Figure 30 and 31, the overhead of the new and restart operations is much higher in the case where a fresh platform is involved (number of workers less than 32). The difference is mainly due to the additional time for Java remote class loading. In fact, when a worker is created, if the classes it needs are not present (in the machine that is running it), they are copied locally then loaded in the cluster node main memory and compiled. Clearly, performing such operations require time, hundreds of milliseconds. Rather, if the classes are already present, already loaded in main memory or even already compiled in machine target code by the Java JIT, performing these reconfiguration operations is noticeably cheaper.

The results of the last experiment are presented in Figure 32. It describes the behavior of the application over quite a long run (two hours, approximately) that includes several self-triggered reconfigurations. In this case the application is provided with a Quality of Service (QoS) contract that enforces the production of a minimum of 1.5 results per second
During the run, an increasing number of platforms are externally overloaded with an artificial load (we started the compilation of some complex software written in C++). The top half of the figure reports the measured average throughput of the filter stage (the second, actually), and the QoS contract. The bottom half of the figure reports the number of overloaded machines along the run, and the corresponding increase of workers of the filter stage. Initially the throughput of the filter stage is abundantly higher than requested ($\sim 3.5$ tasks/s); but it decreases when more machines are overloaded. As soon as the contract is violated, the Autonomic Manager reacts by adding more workers.
Summarizing the Chapter

The challenge of autonomicity in the context of component-based development of grid software is substantial. Building into components autonomic capability typically impairs their reusability. In this Chapter we proposed Behavioural Skeletons as a compromise: being skeletons they support reuse, while their parameterization allows the controlled adaptivity needed to achieve dynamic adjustment of QoS while preserving functionality. We also presented a significant set of skeletons and we discussed how Behavioural Skeletons can be implemented in the framework of the GCM component model. Behavioural Skeletons provide the programmer with the ability to implement autonomic managers completely taking care of the parallelism exploitation details by simply instantiating existing skeletons and by providing suitable, functional parameters. Finally, we discussed the experimental results achieved when running an application exploiting instances of our Behavioural Skeletons and we showed how the skeletons used may take decisions at the appropriate time to maintain the application behavior within the limits stated by the user with a specific performance contract. The whole experiments have been performed using GCM components and Behavioural Skeletons, as being designed and implemented in the framework of the CoreGRID and GridCOMP projects. To our knowledge, no other similar results are available yet.
Chapter 6

Conclusions

Over the years, a lot of models and tools for parallel programming have been proposed. This great deal of efforts is mainly due to the difficulties in coordinating several, possibly hundreds or thousands, activities in an easy way but allowing an efficient exploitation of computational resources. In fact, to date does not exist a universal approach working better than others in every situation. Actually, there are several good approaches based on different perspectives and abstraction levels. Nevertheless, starting from the second half of nineties, with the advent of computational Grids, parallel programming difficulties became greater and greater and also the most promising approaches trail along. Indeed, programming the Grids is even more difficult than traditional parallel programming. This is because the computers belonging to a Grid can be heterogeneous, separated by firewalls, unsafe and managed by different administration policies. To address these additional difficulties most of the models and tools conceived and developed for parallel programming have to be re-thought and adapted. In particular, Structured Parallel Programming models, and the derived environment have been proved to be very effective approach for programming parallel applications, but some well-known issues prevent them from achieving significant popularity in the wider parallel and grid programming community.

In this thesis we presented an organic set of tools and models con-
ceived, designed and developed or properly modified to address most of these issues.

We started discussing how we modified the muskel framework for supporting the issue related to the lack of extendability of the skeleton systems. We discussed how our customized muskel supports the introduction of new skeletons, modeling parallelism exploitation patterns not originally covered by the primitive muskel skeletons. This possibility is supported by allowing muskel users (the programmers) to define new skeletons providing the arbitrary data flow graph executed in the skeleton and by letting our muskel version to seamlessly integrate such new skeletons with the primitive ones. We also presented experimental results validating our muskel approach to extend and customize its skeleton set. We ran several test programs using the custom features introduced in muskel. When grain is small, muskel does not scale well, even using a very small number of remote interpreter instances. When the computational grain is high enough the efficiency is definitely close to the ideal one. Despite the data shown in this thesis refer to synthetic computations, the tests we conducted using actual computations achieved very similar results. This because the automatic load balancing mechanism implemented in the muskel distributed interpreter through auto scheduling perfectly optimized the execution of variable grain macro data-flow instructions. As far as we know, this is the most significant effort in the skeleton community to tackle problems deriving from a fixed skeleton set. Only Schaeffer and his group at the University of Alberta implemented a system were programmers can, in controlled ways, insert new parallelism exploitation patterns in the system (38), although the approach followed here is a bit different, in that programmers are encouraged to intervene directly in the run-time support implementation, to introduce new skeletons, while in muskel new skeletons may be introduced using the intermediate macro data flow language as the skeleton “assembly” language. Unfortunately, programmers using this approach, in order to program unstructured parallel application, have to interact directly with data-flow graph. It requires to programmers to reason in terms of program-blocks instead of a monolithic program. In order to ease the
generation of macro data-flow blocks and in general to provide mecha-
nism easing the use of structured parallel programming environment, we
exploited some metaprogramming techniques.

We exploited some metaprogramming techniques based both on As-
pect Oriented Programming (AOP) and on Attribute Oriented Program-
ming (@OP). We showed how these techniques can be seamlessly ex-
ploited to transform sequential applications into parallel ones. In par-
ticular, we showed how annotations and aspect can be exploited to drive
the sequential application transformation into a macro data-flow graph
that can be executed on distributed architectures. The exploitation of
@OP and AOP techniques allows to completely separate the concerns
relative to parallelism exploitation and application functional code. In
particular, the same application code used to perform functional debug-
ging on a single, sequential machine may be easily turned into parallel
code. To validate the @OP approach we implemented PAL, a java anno-
tation based metaprogramming framework that restructures applications
at bytecode-level at run-time in order to make them parallel. PAL trans-
formations depend on: i) the resources available at run-time, ii) the hints
provided by programmers and iii) the available adapters. An adapter is a
specialized entity that instructs the PAL transformation engine to drive
the code transformation depending on the available parallel tools and
frameworks. Experimental results show that the PAL is an effective and
efficient approach for handling resource heterogeneity and dynamicity.
Actually, run-time code transformation brings to a very good exploita-
tion of computational resources. For this implementation we developed
two distinct adapters. The first adapter we developed foster the byte-
code transformation making the original code a multithreaded one. The
other adapter supports the bytecode transformation that makes the orig-
inal code compliant with JJPF, a structured parallel programming frame-
work we developed some years ago. PAL demonstrated that, given the
existence of a proper metaprogramming run-time support, annotations
are a handy way both to indicate which parts of a program must run in
parallel and to express non-functional requirements directly in the source
code. Therefore, we decided to apply the main features of PAL approach
to our modified muskel implementation. Actually, adapting them to muskel we changed a little bit the approach. Such a change is due to a few motivations. First of all because muskel provides per se a distributed macro data-flow executor whereas PAL exploits external tools for distributed program execution. Moreover, we would like to have a more flexible mechanism for macro data-flow block generation and management. Finally, we would like to exploit a standard tool for run-time code transformation instead of using ad-hoc tools, like the one we developed for PAL. As a consequence we decided to use integrate in muskel the AOP model and in particular the AspectJ framework. The integration has been performed in two steps, in the first step we integrated the AOP mechanisms in order to achieve very simple code transformation. The second step consisted in testing the integration of muskel with AspectJ to in a more complex scenario. Hence, we exploited the aspect oriented programming support we integrated in muskel in order to develop workflows which structure and processing are optimized at runtime. In order to prove the effectiveness of the approach in muskel, we conducted some experiments on a network of workstations. The only difference between plain muskel and the system proposed here to execute workflows lies in the way fireable instructions are provided to the distributed data-flow interpreter of muskel. Indeed, in plain muskel, fireable instructions are taken from a compiled representation of a data-flow graph. Each time a new token arrives to a macro data-flow instruction in the graph the target data-flow instruction is checked for “fireability” and, possibly, delivered to the distributed macro data-flow interpreter. The time spent is in the sub-micro second range (net time, not taking into account time spent to copy parameters in memory during the interpreter call). When executing workflows according to the approach discussed here, instead, fireable instructions is generated at run-time by the AOP engine. We measured the overhead when exploiting the AOP approach, it is approximately 23 milliseconds per workflow node.

These two results presented are feasible approaches for programming cluster or networks of workstation but are not suitable for computational Grids, where component models are preferable. This is due to several
motivations we described in deep in this thesis. Provide parallel programming models for Grids are important because they are becoming the dominant type of parallel architectures. Moreover, due to their heterogeneous and distributed nature, they represent a very good test-bed for testing parallel programming models dealing with dynamicity handling. The \textit{muskel} framework, handle dynamicity exploiting the \textit{Application Manager}: an entity that observes the behavior of the parallel application and in case of problems reacts aiming to fix them. This approach has proved to be effective. Nevertheless, some of the implementation choices done when \textit{muskel} was developed limit its exploitation on Grids. Therefore, we decided to generalize and extend the \textit{muskel Application Manager} approach to make it suitable for components models, in order to be able to port the approach in existing component models. We ported the \textit{muskel} approach in the Grid Component Model. Actually, the \textit{Application Manager} approach form the base of the autonomic features of GCM: each self-optimizing GCM component contains an \textit{Application Manager} that in GCM is called \textit{Autonomic Manager}. Nevertheless, \textit{Autonomic Manager} rely fully on the application programmer’s expertise for the setup of the management code, which can be quite difficult to write since it is tailored for the particular component or assembly of them. As a result, the introduction of dynamic adaptivity might enable the management of grid dynamism but, at the same time, decreases the component reuse potential since it further specializes components with application specific management code. In order to address this problem, we proposed the \textit{Behavioural Skeletons} as a novel way to describe autonomic components in the GCM framework. Behavioural Skeletons aim to describe recurring patterns of component assemblies that can be equipped with correct and effective management strategies with respect to a given management goal. The Behavioural Skeletons model provides a way for handling dynamicity, supporting reuse both of functional and non-functional code. We presented a significant set of skeletons and we discussed how behavioural skeletons can be implemented in the framework of the GCM component model. Behavioural skeletons provide the programmer with the ability to implement autonomic managers completely taking care of the paral-
lelism exploitation details by simply instantiating existing skeletons and by providing suitable, functional parameters. To validate our Behavioural Skeletons we conducted some experiments with the current prototype of the GCM that is currently under development in the GridCOMP STREP project [3]. We discussed the experimental results achieved when running an application exploiting instances of our Behavioural Skeletons and we showed how the skeletons used may take decisions at the appropriate time to maintain the application behaviour within the limits stated by the user with a specific performance contract.

**Future Works**

New efforts for future work can be invested in different directions, as suggested by the results offered by this thesis.

Concerning the macro data-flow based skeleton customizations, new mechanisms for modifying the macro data-flow graph can be conceived, possibly simpler than the existing one. Just as a note, currently we are developing a graphic tool that allows programmers (muskel users) to design their macro data-flow graphs and then compile them directly to Java code as required by muskel.

Several other annotations and aspects can be designed and implemented for easing the run-time generation of macro data-flow blocks. Possibly supporting several types of non-functional requirements. Regarding PAL, many adapters, even more complex than existing one can be developed. In particular, adapters for widely-used frameworks for Grid programming, like Globus or ProActive. Another interesting possibility can be the porting of the adapters model in our customized muskel, perhaps making possible the transformation, at run-time, of the macro data-flow blocks generated by muskel in GCM components.

In this thesis we presented a reduced set of Behavioural Skeletons, other skeletons can be conceived, designed and implemented. As an example, a Behavioural Skeleton supporting the non-functional replication management for easing the development of fault-tolerant component applications. Furthermore, a lot of research can be conducted on the
distributed (cooperative) self-management of component applications, in particular regarding to the methodologies for splitting the user specified QoS contracts.
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