Centralized and Distributed Implementations of Correct-by-construction Component-based Systems by using Source-to-source Transformations in BIP

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Implémtations Centralisée et Répartie de Systèmes Corrects par construction à base des Composants par Transformations Source-à-source dans BIP

Centralized and Distributed Implementations of Correct-by-construction Component-based Systems by using Source-to-source Transformations in BIP

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To my parents & Roukaya ...
Abstract

The thesis studies theory and methods for generating automatically centralized and distributed implementations from a high-level model of an application software in BIP. BIP (Behavior, Interaction, Priority) is a component framework with formal operational semantics. Coordination between components is achieved by using multiparty interactions and dynamic priorities for scheduling interactions. A key idea is to use a set of correct source-to-source transformations preserving the functional properties of a given application software. By application of these transformations we can generate a full range of implementations from centralized to fully distributed.

**Centralized Implementation:** the implementation method transforms the interactions of an application software described in BIP and generates a functionally equivalent program. The method is based on the successive application of three types of source-to-source transformations: flattening of components, flattening of connectors and composition of atomic components. We shown that the system of the transformations is confluent and terminates. By exhaustive application of the transformations, any BIP component can be transformed into an equivalent monolithic component. From this component, efficient standalone C++ code can be generated.

**Distributed Implementation:** the implementation method transforms an application software described in BIP for a given partition of its interactions, into a Send/Receive BIP model. Send/Receive BIP models consist of components coordinated by using asynchronous message passing (Send/Receive primitives). The method leads to 3-layer architectures. The bottom layer includes the components of the application software where atomic strong synchronization is implemented by sequences of Send/Receive primitives. The second layer includes a set of interaction protocols. Each protocol handles the interactions of a class of the given partition. The third layer implements a conflict resolution protocol used to resolve conflicts between conflicting interactions of the second layer. Depending on the given partition, the execution of obtained Send/Receive BIP model range from centralized (all interactions in the same class) to fully distributed (each class has a single interaction). From Send/Receive BIP models and a given mapping of their components on a platform providing Send/Receive primitives, an implementation is automatically generated. For each class of the partition we generate C++ code implementing the global behavior of its components.

The transformations have been fully implemented and integrated into BIP tool-set. The experimental results on non trivial examples and case studies show the novelty and the efficiency of our approach.

Key words: Component-based modeling, source-to-source transformation, correct-by-construction, distributed systems, optimization for performance.
Implémentations Centralisées et Répartie de Systèmes Corrects par construction à base des Composants par Transformations Source-à-source dans BIP

Résumé

La thèse étudie la théorie et les méthodes qui permettent de générer automatiquement des implémentations efficaces tant centralisées que distribuées à partir d’une langage de description de haut niveau pour les applications logicielles embarquées. Ce langage (modèle) appelé BIP (un acronyme pour Behavior, Interaction, Priority) est un outil de développement logiciel fondé sur une théorie saine qui permet la composition incrémentale de composants hétérogènes, ainsi que la génération de code. Une coordination entre les composantes est réalisée en utilisant des interactions multiparties et des priorités dynamiques pour planifier les interactions. L’idée de base consiste à utiliser un ensemble de transformations source-à-source correctes en préservant les propriétés fonctionnelles de l’application logiciel. En appliquant ces transformations, nous pouvons générer une gamme des implémentations centralisées, partiellement distribuées et entièrement distribuées.

Implémentations Centralisées: la méthode transforme les interactions d’un logiciel décrit dans BIP et génère un programme équivalent au niveau fonctionnel. La méthode est basée sur l’application successive de trois types de transformations source-à-source: aplatissement des composants, aplatissement des connecteurs et la composition des composants. On a montré que le système des transformations est confluent et se termine. Par une application exhaustive de ces transformations, un composant BIP peut être transformé en un composant atomique. A partir de ce composant, un code C++ efficace peut être généré.

Implémentations Distribuées: Pour une partition donnée des interactions la méthode transforme un application logiciel décrit en BIP, en un modèle Send/Receive BIP. Ce modèle BIP est constitué des composants coordonnés à l’aide des messages asynchrones (Send/Receive). La méthode conduit à une architecture de 3-couches. La couche inférieure comprend les composants du logiciel où les fortes synchronisations atomiques sont implémentées par des séquences des primitives Send/Receive. La deuxième couche comprend un ensemble de protocoles d’interaction. Chaque protocole gère un ensemble des interactions. La troisième couche implémente un protocole de résolution des conflits utilisées pour résoudre les conflits entre les interactions conflictuelles de la deuxième couche. A partir des modèles Send/Receive BIP, une implémentation C++ est générée automatiquement.

Les transformations ont été implémentées et intégrées dans la chaîne d’outil BIP. Les résultats expérimentaux sur des exemples non triviaux et des études de cas montrent la nouveauté et l’efficacité de notre approche.

Mots clé: Modélisation à base de composants, source-à-source transformation, correcte-par-construction, systèmes distribuées, optimisation pour la performance.
Several Chapters in this thesis appeared in several papers in form of articles or of Verimag technical reports.

The symbolic implementation of BIP engine in Chapter 3 appeared in the paper [JBB09] in ICE 2009 (Structured Interactions CONCUR 2009 affiliated workshop), 31st August - Bologna, Italy. A journal version is under review in Mathematical Structures in Computer Science.


The BIP into Distributed BIP methods together with the results in Chapter 5 appeared in two papers: paper [BBJ+10a] has been accepted (best paper award) for SIES 2010 (IEEE Symposium on Industrial Embedded Systems, University of Trento, Italy, July 7 - 9, 2010), and paper [BBJ+10b] has been accepted for EMSOFT 2010 (International Conference on Embedded Software, Scottsdale, Arizona, USA, October 24 - 29, 2010).
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1.1 Les Défis de la Construction Correct de Systèmes à base de Composants

Les systèmes informatiques sont omniprésents aujourd’hui, ils se touchent tous les aspects de la vie humaine. Nous pouvons les trouver dans différents types des applications de l’automobile, aux appareils ménagers banals comme les machines à laver et des fours à micro-ondes, à l’aéronautique, militaire, télécommunications, médicaux, ...
Les systèmes deviennent de plus en plus complexes et leur adoption est en augmentation exponentielle. En outre, la nécessité de prouver et de garantir leur validité, l’efficacité et la fiabilité est une tâche essentielle qui est aussi complexes. Bien que différentes techniques de génie logiciel existent pour assurer l’exactitude telles que la vérification formelle, la simulation et les tests, le renforcement des systèmes justes et fiables est toujours une tâche difficile. Les erreurs existent encore dans plus de 90% des cahiers de charge de production. En plus, le temps passé dans les débogage est très élevé et parfois sans résultats satisfaisants. A titre d’exemple, les États-Unis a tenté de remplacer son système de contrôle du trafic aérien trois fois au cours des vingt dernières années, mais malgré les milliards de dollars US dépensés, tel remplacement ne s’est pas passé.

Approche Fondée sur les Composants La technique la plus fondamentale pour résoudre des problèmes complexes est de les décomposer en plus petites. Comme les systèmes complexes peuvent être obtenus par assemblage de composants (construction des blocs), le processus de conception de ces systèmes complexes peut être réduit à l’étude des plus petits et plus simples composants. Ainsi, en utilisant un tels cadres (appelé à base de composants ) qui permettent la construction des systèmes à partir de composants prédéfinis.

Le développement de systèmes à base de composants nécessite des méthodes et des outils de différents concepts de l’architecture qui fournissent aussi une caractérisation de coordination entre les composantes. L’architecture est la structure d’un système. Il comprend des éléments et leur relation entre les propriétés visibles de l’extérieur. Cela signifie que l’architecture décrit comment les composants sont connectés et comment ils peuvent interagir. Par conséquent, le comportement global d’un système peut en principe être déduit du comportement de ses composants et son architecture. Les systèmes à base de composants fournissent descriptions claires et logiques qui en font un bon candidat pour un processus basés sur la construction. En outre, ils permettent la réutilisations des sous-systèmes, ainsi que leur modification progressive sans nécessiter de changements mondiaux, ce qui peut simplifier considérablement le processus de vérification. Néanmoins, la clarté et l’expressivité des systèmes à base de composants peut être au détriment de l’efficacité. En effet, la compilation naïve de systèmes à base de composants se traduit généralement par l’inefficacité comme une conséquence de l’interconnexion des composants.

Conception de Haut Niveau Lors de la conception des systèmes complexes, le traitement de la complexité à un haut niveau est d’une grande aide pour les concepteurs. En fait, il serait plus facile pour les concepteurs de commencer à travailler sur le système à un haut niveau, où ils n’ont pas à tenir compte de la complexité du système. Cet phase de modélisation phase de modélisation est bénéfique, car les concepteurs peuvent abstraire des détails d’implémentation et de valider le modèle abstrait par rapport à un ensemble de prescriptions visant à travers différentes techniques telles que la vérification formelle, la
Malheureusement, une fois le modèle abstrait est validé, la génération des implémentations correcte et efficace est toujours difficile, car ajoutant des détails de mise d’implémentation implique de nombreuses subtilités qui peuvent potentiellement introduire des erreurs dans le système résultant.

**Implémentation** La conception haut niveau est utilisé pour prototyper et valider de modèles complexes. Après cette phase, la conception est généralement coder par la main pour générer une implémentation du système. Toutefois, ce processus de traduction est complexe et il peut réduire la contribution de la vérification et de tests effectués sur le modèle de haut niveau. Par conséquent, la génération automatique de code est nécessaire et peut fournir un gain de temps important. Néanmoins, le principal inconvénient de cette approche automatique est son efficacité. En effet, l’efficacité d’un code généré dépend en général de l’architecture cible.

Étant donné un modèle de haut niveau, l’implémentation généré peuvent être, en général, centralisée ou distribuée. Cela dépend à la fois de la topologie des systèmes et l’architecture sur laquelle le système sera déployé. Par exemple, si l’on considère un système centralisé avec une petite quantité de calcul et beaucoup de communication, dans ce cas une implémentation centralisée se comporte mieux qu’un distribuée.

Dans d’autres cas, nous pourrions avoir à choisir une implémentation distribuée parce que le système lui-même est répartie géographiquement, ou parce que l’utilisation des plusieurs processeurs est nécessaire pour effectuer un grand puissance de calcul. Une implémentation distribuée est nécessaire pour ce genre de systèmes. Ces systèmes sont principalement utilisés pour le monde entier-web, serveur de transfert des fichiers, réseau bancaire, les réseaux pair-à-pair, les systèmes de contrôle de processus, réseaux de capteurs, grille de calcul, etc

Il est clair que la complexité sont amplifiés de manière significative dans le cas des implémentations distribuées à cause de la structure concurrente, non déterministe, et non-atomique de systèmes distribués, ainsi que la survenue d’événements imprévus physiques et informatiques telles que les failles. En outre, il est difficile de savoir comment transformer un modèle abstrait (où atomicité est supposé par la sémantique global et les détails de distribution ont été omis tout en employant primitives de synchronisation de haut niveau) en une implémentation distribués.

Dans cette thèse nous proposons une théorie et des outils pour dériver automatiquement des implémentations centralisée et distribuée correcte et efficace d’une manière systématique et automatisé à partir d’un cadre de haut niveau à base de composants. Ce résultat est obtenu, en utilisant des techniques de transformations source-à-source, correcte techniques de construction.
1.2 État de l’art de Conception et Implémentation de Systèmes à base des Composants

1.2.1 Les cadres à base de composants

Les techniques de conception à base de composants sont utilisés pour réduire la complexité des systèmes. L’idée est que les systèmes complexes peuvent être obtenus par assemblage de composants. Cela est essentiel pour le développement à grande échelle, les systèmes évolutifs d’une manière rapide et abordable. Il offre une flexibilité dans la phase de construction des systèmes en soutenant l’ajout, la suppression ou la modification des composants sans aucune ou très peu d’impact sur les autres composantes. Les composants sont généralement caractérisés par des abstractions qui ignorent les détails d’implémentation et de décrire les propriétés pertinentes de leur composition, par exemple fonctions de transfert, interfaces. La principale caractéristique de cadres de conception à base de composants est la composition. La composition est utilisée pour construire des composants complexes à partir de plus simples. Il peut être formalisé par une opération qui prend en entrée un ensemble de composants et de leurs contraintes d’intégration et fournit, en sortie, la description d’un nouveau composant plus complexe. Cette approche permet de réduire la complexité des systèmes en offrant l’effet d’accroissement dans la phase de construction. Il existe une grande littérature traitant la conception à base de composants. Les travaux suivants sont liés à cette approche:

– PtolemyII est un cadre logiciel, développé au génie électrique et informatique (RECS) UC Université de Berkeley. PtolemyII porte sur la modélisation hétérogènes à base de composants. Il préconise une vision axée sur les acteurs d’un système, où le bloc de construction de base d’un système est un acteur. Un modèle est une interconnexion hiérarchique des acteurs. Les acteurs sont des composants logiciels qui s’exécutent simultanément et se communiquent à travers des interfaces appelées ports. Un acteur peut être atomique, dans ce cas elle doit être au bas de la hiérarchie. Un acteur peut être composite, dans ce cas, il contient d’autres acteurs. La sémantique d’un modèle n’est pas déterminée par le cadre, mais plutôt par un composant logiciel dans le modèle appelé le directeur, qui met en œuvre un modèle de calcul. PtolemyII permet la simulation de modèles. Toutefois, la vérification des modèles est, actuellement, pas possible. Des travaux sont en cours pour ajouter cette possibilité. Il permet la génération de code, pour ce faire, chaque composant doit être accompagné d’un modèle («template») de code C qui sera complété par le générateur de code. En outre, PtolemyII n’a pas de notion intrinsèque de mappage entre les acteurs ou de l’utilisation de spécification déclarative dans la conception.

– IF (Intermediate Format) est un ensemble d’outils, développé à Verimag. IF est un ensemble d’outils de plateforme de modélisation et validation des composants. Il se compose d’ensemble d’automates temporisé communicants de manière asynchrone en
envoyant des signaux à travers FIFO, ou de manière synchrone avec rendez sur les ports synchronisé. IF utilise des techniques telles que la réduction d’ordre partiel et model-checking à la volée pour l’exploration d’espace d’état.

- Fractal est un modèle de composant, développé à France Télécom R & D et l’INRIA en France. Fractal est un modèle de composants générales, pour le déploiement et la gestion de systèmes logiciels complexes. Il peut être compris comme étant composé d’un membrane qui se compose d’un ensemble de composants (appelés sous-composants) et une ou plusieurs interfaces (semblable au port de modèles de composants d’autres). Les interfaces peuvent être de deux sortes: les interfaces serveur pour les appels des opérations entrants, et les interfaces clients pour les appels des opération sortants. THINK est l’un des la mise en œuvre de Fractal. Cependant Fractal et THINK ne fournissent pas d’outils ou de techniques d’analyse, que ce soit pour la simulation ou de vérification.

- Metropolis est un cadre à base de composants: composants atomiques contenant comportement (code), pièce en matériau composite contenant d’autres sous-composantes. Interfaces des composants sont constitués d’un ensemble de ports utilisés soit pour les communications asynchrones (envoyer des événements), ou pour rendez-vous (synchronisation entre les composants). Metro II fournit une interface qui produit une représentation interne de la méta-modèle. Cette représentation peut être utilisée pour, génération de code C pour la simulation, ou modèle de génération utilisé avec le modèle de spin-checker, etc.

D’autres développements traitent, d’une façon ou d’une autre, des questions liées à la modélisation basée sur les composants:

- langages de modélisation de système tel que UML et les outils associés.
- langages pour la conception de systèmes tels que SystemC, GME, Simulink/Stateflow, Autofocus.
- normes Middleware tels que Corba, JavaBeans,.NET
- les environnements de développement logiciels tels que PCTE, SWbus, Softbench, Eclipse.
- langage de coordination extension du langage de programmation tels que Linda, JavaSpaces, TSplaces, Concurrent Fortran, le CNES et polyphoniques C♯.
- cadres théoriques basés sur les algèbres de processus par exemple, le Pi-calcul ou basé sur des automates.

1.3 Résumé Chapitres

1.3.1 Chapitre 2 - BIP Cadre à Base des Composants

Ce chapitre présente plusieurs notions de base fondamentales pour l’étude des systèmes complexes, à savoir la méthodologie à base de composants, leur composition fondée des
opérateurs adéquats et bien définis, ainsi que les propriétés nécessaires à la construction de systèmes construits à partir de composants. Ce chapitre introduit ensuite le langage BIP (son architecture, sa sémantique, et ses propriétés).

1.3.2 Chapitre 3 - Transformation pour la Génération des Implémentation Centralisées

Ce chapitre présente une méthode pour générer des implémentations centralisées correctes et efficaces à partir de programmes BIP. La méthode proposée est fondée sur l’application successive de trois types de transformations source-à-source: aplatissement des composants, aplatissement des connecteurs, puis composition des composants. L’application successive des ces transformations permet de synthétiser vers un seul composant (qui est un réseau de Pétri). A partir de ce composant un code C efficace est généré. Ce chapitre décrit des applications de cette méthode sur des exemples non triviaux (encodage Vidéo MPEG4, Network Sorting Algorithm).

1.3.3 Chapitre 4 - Transformation pour la Génération des Implémentation Distribuées

Ce chapitre développe une méthode pour générer des implémentations distribuées correctes et efficaces à partir de programmes BIP. Pour cela il présente d’abord les principales difficultés rencontrées dans la génération d’implémentations distribuées. Il définit ensuite un ensemble de transformations source-à-source qui permettent:

– la rupture des atomicités des actions dans les composants atomiques en remplaçant les synchronisations fortes (rendez-vous) par des interactions asynchrones de type Send/Receive;
– la génération des contrôleurs qui coordonnent l’exécution des interactions selon une partition définie par l’utilisateur;
– l’ajout d’un algorithme distribué pour le traitement des conflits entre contrôleurs.

Le modèle Send/Receive est prouvés observationnellenement équivalents au modèle initial. En outre, des implémentations distribuées peuvent être générées à partir de modèles Send/Receive qui permettent un parallélisme réel entre les composants ainsi entre les interactions. En particulier, il est possible de générer des modules autonomes en code C qui utilisent des communications du type MPI ou socket.

1.3.4 Chapitre 5 - Chaîne d’outils

Ce chapitre présente les outils qui mettent en œuvre la théorie et les transformations définies dans ce travail. Il présente aussi une intégration de ces outils dans la phase de conception des applications en BIP, dont le but est de générer des implémentations (tant centralisées que distribuées).
1.3.5 Chapitre 6 - Conclusions et Perspectives

Ce chapitre fournit une conclusion générale et des directions de recherche futures. L’une d’entre elles réside dans la prise en compte de priorités dans les implémentations distribuées.
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Introduction

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2.1 Challenges in Building Correct Component-based Systems

Computer systems are ubiquitous today, they are touching all aspects of human life. We can find them in different types of applications from automobile, to mundane home appliances like washing machines and microwave ovens, to aeronautic, military, telecommunication, medical etc.

Systems become more and more complex and their adoption is increasing exponentially. Moreover, the need to prove and to ensure their correctness, efficiency and reliability is an essential task that is also becoming complex. Although different techniques in software engineering exist for ensuring correctness such as formal verification, simulation, and testing, building correct and reliable systems is still a time-consuming and hardly predictive task.
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Errors still exist in up to 90% of production spreadsheets. Moreover, the time spent in debugging them is considerably high and sometimes with no satisfying results. As an example, the United States has attempted to replace its air traffic control system three times in the past twenty years, but despite the billions of US dollars spent, no such replacement has happened.

**Component-based Approach**  The most basic technique to tackle complex and large problems is to decompose them into smaller ones. As complex systems can be obtained by assembling components (building blocks), the process design of these complex systems may be reduced to the study of smaller and simpler ones. Thus, using such frameworks (called *component-based*) that allow building systems from predefined given components would be a great interest.

Component-based system development requires methods and tools supporting different concepts of architecture which provide a characterization coordination between components. An architecture is the structure of a system. It involves components and their relationship between the externally visible properties. This means that the architecture describes how components are connected and how they can interact. Consequently, the global behavior of a system can in principle be inferred from the behavior of its components and its architecture. Component-based systems provide logical clear descriptions which make them a good candidate for a correct-by-construction process. In addition, they allow sub-systems to be reused as well as their incremental modification without requiring global changes, which may significantly simplify the verification process. Nonetheless, clarity and expressiveness of component-based systems may be at the detriment of efficiency. Indeed, naive compilation of component-based systems generally results in great inefficiency as a consequence of the interconnection of components [Lov77].

**High-Level Design**  When designing systems, dealing with their complexity at a high-level is a great help for the designers. In fact, it would be easier for designers to start working on the system at a high-level, where they do not have to take into account the complexity of the system. This *modeling* phase is beneficial, as designers can abstract away implementation details and validate the model with respect to a set of intended requirements through different techniques such as formal verification, simulation, and testing.

Unfortunately, once the abstract model is validated, deriving *correct* and *efficient* implementation from it is always challenging, since adding implementation details involves many subtleties that can potentially introduce errors into the resulting system.

**Implementation Level**  High-level design is used to prototype and validate new designs. After this phase, the design is typically translated by hand into code for a system implementation. However, this translation process is complex and it may reduce the contribution of the verification and testing done at the high-level model. Therefore, automatic code
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generation is necessary and may provide an important time gain. Nevertheless, the main drawback of this automatic approach is its efficiency. Indeed, the efficiency of a generated code depends in general on the target architecture.

Given a high-level model, the implementation generated could be, in general, centralized or distributed. This depends both on the topology of the systems and the architecture on which the system will be deployed. For instance, if we consider a centralized system with a small amount of computation and a high-level of communications overhead, then a centralized implementation behaves better than a distributed one. In other cases, we may have to choose a distributed implementation because the system itself is geographically distributed, or because deriving more computational power by using multiple processors is necessary. These kind of systems where distributed implementation is needed, are mainly used for world-wide-web, network-file server, banking network, peer-to-peer networks, process control systems, sensor networks, grid computing, etc.

It is clear the complexity are amplified significantly in the case of deriving distributed implementation because of inherently concurrent, non-deterministic, and non-atomic structure of distributed systems, as well as the occurrence of unanticipated physical and computational events such as faults. Moreover, it is unclear how to transform an abstract model (where atomicity is assumed through global state semantics and distribution details are omitted via employing high-level synchronization primitives) into a real distributed implementation.

Figure 2.1: High-level component-based design into implementation.
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In this thesis we propose a theory and tools for automatically derives correct and efficient centralized and distributed implementation in a systematic and ideally automated correct fashion from a high-level component-based framework. This is achieved, by using correct-by-construction source-to-source transformations techniques (see Figure 2.1).

The rest of the chapter is organised as follows. First in Section 2.2 we give a brief description of the current state of the art of component-based systems design and implementation. Then, in Section 2.3 we present the contribution of the thesis. Finally, we describe the outline of the thesis in Section 2.4.

2.2 State-of-the-Art of Component-based Systems Design and Implementation

In this section we summarize a brief description of the current state-of-the-art in component-based systems.

2.2.1 Existing component-based frameworks

Component-based design techniques are used to cope with the complexity of the systems. The idea is that complex systems can be obtained by assembling components. This is essential for the development of large-scale, evolvable systems in a timely and affordable manner. It offers flexibility in the construction phase of systems by supporting the addition, removal or modification of components without any or very little impact on other components. Components are usually characterized by abstractions that ignore implementation details and describe relevant properties to their composition, e.g. transfer functions, interfaces. The main feature of component-based design frameworks is allowing composition. This composition is used to build complex components from simpler ones. It can be formalized as an operation that takes, as input, a set of components and their integration constraints and provides, as output, the description of a new more complex component. This approach allows to cope with the complexity of systems by offering incrementality in the construction phase. There exists a large body of literature dealing with component-based design. The following works are related to this approach:

- PtolemyII [DII*99, EJL*03] is a software framework, developed at Electrical Engineering and Computer Sciences (EECS) UC Berkeley University. PtolemyII focuses on component-based heterogeneous modeling. It advocates an actor-oriented view of a system, where the basic building block of a system is an actor. A model is a hierarchical interconnection of actors. Actors are software components that run concurrently and communicate through interfaces called ports. An actor can be atomic, in which case it must be at the bottom of the hierarchy. An actor can be composite, in which case it contains other actors. The semantics of a model is not determined by the framework, but rather than by a software component in the model called di-
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rector, which implements a model of computation. PtolemyII allows the simulation of models. However, the verification of models is, currently, not possible. Work is underway to add this possibility. It allows the generation of code, to do this, each component must be accompanied by a model ("template") of C code which will be completed by the code generator. Moreover, PtolemyII has no intrinsic notion of mapping between actors or of using declarative specification in the design.

– IF (Intermediate Format) [BGO+04] is a toolset, developed at Verimag. The IF toolset is a platform for component modeling and validation. It consists of set of timed automata communicating, either asynchronously by sending signals through FIFO, or synchronously with rendez on synchronized ports. The IF toolset uses techniques such as partial order reduction and on-the-fly model checking to explore the state space of the IF specification, giving access at the semantic level, to the corresponding labeled transition system (LTS). The latter can be analysed using the tool suite CADP [FGK+96], including the minimization and comparison tool Aldebaran based on bisimulation, and the alternating-free -calculus model-checker Evaluator.

– Fractal [BCS02, BCL+06] is a component model, developed at France Telecom R&D and INRIA France. Fractal is a general component model for implementing, deploying and managing complex software systems. It can be understood generally as being composed of a membrane which consists of a set of components (called subcomponents) and one or more interfaces (similar to port in other component models). Interfaces can be of two kinds: server interfaces for incoming operation invocations, and client interfaces for outgoing operations invocations. Think [FSLM02, AHJ+09] is one of the Fractal implementation. However Fractal and THINK do not provide tools or analysis techniques, whether for simulation or verification.

– Metropolis [BWH+03] and its successor [DDM+07]. Metropolis is a component-based framework : atomic component containing behavior (code), composite component containing other sub-components. Interfaces of components consist of a set of ports used either for asynchronous communications (send events), or for rendez-vous (synchronization between components). Metro II provides a frontend which produces an internal representation from the meta-model. This representation can be used for, generation of C++ code for simulation, or generation model used with the SPIN model-checker, etc.

Other developments deal, one way or another, with issues related to component-based modeling:

– Software Design Description Languages [GS04, BFL+04], and Architecture Description Languages focusing on non-functional aspects [VPL99, AVCL02].

– Standardized system modeling languages such as UML [OMG] and associated tools.

– Languages and notations specific to system design tools such as SystemC [Pan01,
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- RHG\textsuperscript{+}01, GME [BGK\textsuperscript{+}06], Simulink/Stateow [Mat], Autofocus [HS02].
- Middleware standards such as Corba, Javabeans, .NET
- Software development environments such as PCTE, SWbus, Softbench, Eclipse.
- Coordination language extension of programming languages such as Linda, Javaspaces [FAH99], TSpaces [FLN\textsuperscript{+}03], Concurrent Fortran, nesC [GLvB\textsuperscript{+}03] and Polyphonic C\textsuperscript{♯} [BCF02].
- Theoretical frameworks based on process algebras e.g., the Pi-Calculus [Mil98] or based on automata e.g., [RC03].

2.2.2 Discussion

There are different requirements for building efficient and correct implementation for complex systems.
- Firstly, we need a component framework with the concept of component and associated composition operators for incremental description and correctness by construction.
- Secondly, this framework should be expressive enough to directly encompass all types of coordination, and hence, help designers to formulate their solutions in terms of tangible, well-founded and organized concepts instead of using dispersed coordination mechanisms such as semaphores, monitors, message passing, remote call, protocols etc.
- Thirdly, it also needs to be abstract enough by providing high-level primitives for modeling behaviors and communications. However, abstraction reduces expressiveness. Thus, the first challenge is to find the best compromise between a high level of abstraction and high expressiveness. All of these requirements lead to design complex systems in an easy and correct manner. Nonetheless, on top of abstraction and expressiveness, etc., other challenges appear, mainly how to automatically derive a correct and efficient implementation.
- For this reason, the fourth requirement, for such framework, is to provide a rigorous but not complex semantics, because complexity limitates abstraction. When having a rigorous semantics, we can define correct and automatic source-to-source transformation for deriving efficient low-level implementation from the high-level models in a correct manner. Moreover, a strong theoretical backing can be defined at the high-level models that allows formal verification of design properties.

Indeed, source-to-source transformations have been considered as a powerful means for optimizing programs [Lov77, HG06, BMFT07]. In contrast to conventional optimization techniques, they can be applied for deeper semantics-preserving transformations which are visible to programmers and subject to their direction and guidance.

In the context of component-based frameworks, we have not seen major work on source-to-source transformations, since component frameworks such as [BWH\textsuperscript{+}03, DII\textsuperscript{+}99] have well-defined denotational semantics. Nonetheless, it can be made only at the execution
level and not at source level. For example, it is not clear how to define component composition at source level from these semantics.

There also exist many component frameworks without rigorous semantics. This is particularly absent in the case of modeling, as well as for middleware and software development standards, like CORBA. They use ad-hoc mechanisms for building systems from components and offer syntax level concepts only. In this case, using ad-hoc transformations, may easily lead to inconsistencies e.g. transformations may not be confluent.

One the other hand, there are other techniques based on source-to-source transformations applied by language compilers. For example, compilation of synchronous languages [JHRC08], or optimizing communications in periodic reactive systems [CKL+05, CKL+02] use transformation techniques for optimization by flattening structure and composing Petri net behavior. Nonetheless, their underlying models are completely simple with respect to the communication primitives offered by these languages. For instance, the model considered in [CKL+05, CKL+02] is an extension of Kahn process networks allowing non-deterministic waiting on multiple input channels. The communication is binary, through point-to-point message passing on FIFO channels.

To this end, on exploring the current state of the art we have not seen a component-framework that meets the requirements above. Generally speaking, we can divide them into two categories. The first category provides high-level design and modeling, however it is still unclear how to derive correct and efficient implementation from the high-level models. In contrast, the second category provides efficient implementation, however the design process is either based on low-level primitives, or not expressive enough.

2.3 Our Contribution

We present, in this thesis, a methodology to provide automatically efficient and correct-by-construction centralized and distributed implementations starting from a high-level model of the software application in BIP. BIP (Behavior, Interaction, Priority) is a component-based framework with formal and rigorous semantics that rely on multiparty interactions for synchronizing components and dynamic priorities for scheduling between interactions.

A key idea of our methodology is to use a set of correct source-to-source transformations which preserve functional properties. Furthermore, they take into account extra-functional constraints. We propose several types of source-to-source transformations:

1. Transformation for generating centralized implementations: We define a set of transformations taking BIP models as input and transform them into functionally equivalent BIP models with different architectures. Such transformations allow in particular to generate from a hierarchical model an equivalent flat model or a single component by composing the behavior of the constituent components. From flat models monolithic C++ code can be generated. This code has been proven optimal
2. Transformation for generating distributed implementations: Coordination in BIP is achieved through multiparty interactions and scheduling by using dynamic priorities. The associated semantics is defined on a global state model. This makes reasoning about systems easy. However, it is hard to obtain distributed implementations where the primitives available for communication and coordination are less powerful. For this reason, we propose automated transformation of high-level BIP models (where high atomicity is assumed and distributed coordination is sought by multi-party synchronization primitives) into distributed implementations in a systematic and correct fashion. In general, our methodology transforms arbitrary BIP models into Send/Receive BIP models, directly implementable on distributed execution platforms. The transformation consists of:

- breaking atomicity of actions in atomic components by replacing strong synchronizations with asynchronous Send/Receive interactions;
- inserting several distributed Engines that coordinate execution of interactions according to a user-defined partition;
- augmenting the model with a distributed algorithm for handling conflicts between Engines.

The obtained Send/Receive BIP models are proven observationally equivalent to the initial models. Hence, all the functional properties are preserved by construction in the implementation. Moreover, Send/Receive BIP models can be used to automatically derive distributed implementations. Currently, it is possible to generate stand-alone C++ implementations using either TCP sockets for conventional communication, or MPI implementation, for the deployment on multi-core platforms.

This approach has been fully implemented and integrated in the BIP framework.

2.4 Organization of the Thesis

The rest of the thesis consists of five chapters. In Chapter 3 we present an overview of the BIP framework. Then, in Chapter 4 we describe a set of source-to-source transformations for generating efficient centralized implementation for deployment on single-processor platforms. In Chapter 5 we present a method using source-to-source transformation for generating efficient distributed implementation for deployment on multi-core platforms. In Chapter 6 we present the tool implementing the techniques proposed in this thesis. Finally, Chapter 7 draws conclusion and future work. The details of all chapters are as follows:

- Chapter 3 presents the basic ideas about component-based methodology, the basic notions about components, their composition using glue operators, and the necessary properties for component-based construction of systems. It introduces the BIP component framework, describing its architecture, its semantics as well as its properties.
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- Chapter 4 presents a method for generating efficient centralized implementation from high-level BIP model. The method is based on the successive application of three types of source-to-source transformations: flattening of components, flattening of connectors and composition of atomic components. We show that the system of the transformations is confluent and terminates. By exhaustive application of the transformations, any BIP component can be transformed into an equivalent monolithic component. From this component, efficient standalone C++ code can be generated. Applications of the method on two non trivial examples are also described in the chapter.

- Chapter 5 presents a method for generating efficient distributed implementations from high-level BIP model. First, in this chapter, we present the main subtitles of generation distributed implementations. Second, we define set of source-to-source transformations which transform a high-level BIP model (where atomicity is assumed through global state semantics and distribution details are omitted via employing high-level synchronization primitives) into a real distributed implementation that allows parallelism between components as well as parallel execution of interactions. Moreover, we prove that the defined transformations preserve observational equivalence. Applications of the method on three non trivial examples are also described in the chapter.

- In Chapter 6 we present a tool which implements the transformations defined in Chapter 4 and 5. Moreover, we give an overview of the integration of our tool in the design methodology of BIP for automatically deriving efficient centralized and distributed implementations from high-level BIP models.

- We conclude the thesis in Chapter 7, with an overview of the work and its future perspectives.
The BIP Component-based Framework

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BIP (Behavior, Interaction, Priority) is a component framework for modeling heterogeneous real-time systems. In the first section, we give notions about component, their composition and the necessary properties for component-based construction of systems. Then, in Section 3.2 we present the BIP language. In Section 3.3 we present the execution platforms implementing the operational semantics of BIP. Then, Section 3.4 we give an overview of the BIP tool-chain. And we finish this chapter by giving conclusions in summary.
3.1 General Overview

BIP is a component framework with a formal operational semantics given in terms of Labeled Transition Systems and Structural Operational Semantics derivation rules. Indeed, a component is a behavioral entity, having a well defined interface. It denotes an executable specification whose runs can be modeled as sequences of discrete actions.

We distinguish two kinds of components: atomic and composite. Atomic components are the basic elements in the components hierarchy. Their behavior represented as labeled transition systems.

**Definition 3.1.1 (Labeled transition system.)** A labeled transition system is a triple \( B = (Q, \Sigma, \rightarrow) \), where \( Q \) is a set of states, \( \Sigma \) is a set of labels, and \( \rightarrow \subseteq Q \times \Sigma \times Q \) is a set of labeled transitions.

For any pair of states \( q, q' \in Q \) and label \( a \in \Sigma \), we write \( q \xrightarrow{a} q' \), iff \( (q, a, q') \in T \). If such \( q' \) does not exist, we write \( q \xrightarrow{a} \).

Composite components are obtained by composing together other components (atomic or composite) using a glue operator, then, a new component can be derived (see Figure 3.1). Their behavior is the product of behaviors of the inner components, with restriction implied by the glue.

**Figure 3.1:** Components composition.

Our ultimate goal is to provide a methodology for component description and integration in a meaningful manner. The methodology must be incremental, i.e., components can be composed through a meaningful hierarchy of glues. Moreover, in order to ensure the correctness of composite components, it must provide support for compositinality and composability. In the following we give a description of these requirements:

- **Incrementally** a system can be considered as the composition of smaller components with the ability to the combination of decomposition and flattening (see Figure 3.2).

**Figure 3.2:** Incrementality of composition.
Chapter 3. The BIP Component-based Framework

- **Compositionality** the possibility of inferring a global system properties from the local properties of sub-systems (e.g inferring global deadlock-freedom from the deadlock-freedom of the individual components) (see Figure 3.3).

- **Composability** the preservation of the main properties of components during the construction of the system (see Figure 3.4).

A detailed and fully formalized of these properties are presented in [Bas08]. The BIP component framework presents the composition of behaviors using two kinds of glue, interactions and priorities. It is shown in [BS08b] that these encompass the universal glue.

### 3.2 The BIP Language

BIP [Sif05, BBS06] is a component framework for constructing systems by superposing three layers of modeling (see Figure 3.5): Behavior, Interaction, and Priority. The lowest layer consists of a set of atomic components represented by transition systems. The second layer models Interaction between components. Interactions are sets of ports specified by connectors [BS08a]. Priority, given by a strict partial order on interactions, is used to enforce scheduling policies applied to interactions of the second layer. The BIP component framework has a formal operational semantics given in terms of Labeled Transition Systems and Structural Operational Semantics derivation rules. The BIP language offers primitives and constructs for modeling and composing complex behavior from atomic components. Atomic components are communicating Petri net extended with C functions and data. Transitions are labeled with sets of communication ports. Composite components are obtained from subcomponents by specifying connectors and priorities.

A component in BIP can also be viewed as a point in a three-dimensional space represented in Figure 3.6. The dimension Behavior characterizes component behavior and the
Chapter 3. The BIP Component-based Framework

Figure 3.5: Layered component model.

Figure 3.6: Three-dimensional space construction.

space \textit{Interactions} \times \textit{Priorities} characterizes the overall structure of the system. In the following sections, we give a formal description of each of the layers, introduced here.

### 3.2.1 Ports and Interfaces

Ports are particular names defining communication points for components. As we shall see later, they are used to establish interactions between components by using connectors.

In BIP, we assume that every port has an associated distinct data variable \( x \). This variable is used to exchange data with other components, when interactions take place.

A set of ports is called an interface.

**Definition 3.2.1 (Port.)** A port \( p[x] \) is defined by

- \( p \) – the port identifier,
- \( x \) – the data variable associated with the port.
3.2.2 Atomic Component

Atomic component is a unit of behavior with an interface consisting of ports, and behavior encapsulated as a set of transitions. It consists of:

- A set of control states \( L \), denoting locations at which the components await for synchronization.
- A set of ports \( P \) used for synchronization with other components. Ports form the interface of atomic components. They are instances of predefined port-types, and may be associated with atom variables.
- A set of variables \( X \) used to store (local) data. Basic C types can be used for variables. Variables may be associated to one or more ports. A variable associated to a port can be modified as a result of an interaction involving that port.
- A behavior given by the set of transitions modeling atomic computation steps. A transition represents a step from a set of control states \( L_1 \) to \( L_2 \) labeled by a port \( p \), guard \( g \), function \( f \). A transition is denoted as \( L_1 \xrightarrow{p,g,f} L_2 \).

Here \( p \) is a port through which an interaction is sought, \( g \) a pre-condition for interaction through \( p \), and \( f \) is a computation step consisting of local state transformations. \( g \), also know as the guard of the transition, is a boolean condition on \( X \). The transition can be executed if the guard is true.

Example 1 Figure 3.7(a) shows an example of an atomic component with two ports \( p_1 \), \( p_2 \), a variable \( x \), and two control states \( l_1 \), \( l_2 \). At control state \( l_1 \), the transition labeled \( p_1 \) is enabled. When an interaction through \( p_1 \) takes place, a random value is assigned for the variable \( x \). This value is exported through the port \( p_2 \). From the control state \( l_2 \), the transition labeled \( p_2 \) can occur (the guard is true by default), the variable \( x \) is eventually modified and the value of \( x \) is printed.

Definition 3.2.2 (Atomic component.) An atomic component \( B \) is defined by

\[
B = (L, P, T, X, \{g_\tau\}_\tau \in T, \{f_\tau\}_\tau \in T),
\]

where,

- \( (L, P, T) \) is a 1-safe Petri net, that is
  - \( L = \{l_1, l_2, \ldots, l_k\} \) is a set of control states,
  - \( P \) is a set of ports,
  - \( T \subset 2^L \times P \times 2^L \) is a set of transitions,
- \( X = \{x_1, \ldots, x_n\} \) is a set of variables and for each transition \( \tau \in T \), \( g_\tau \) is a guard and \( f_\tau \) is an update function that is state transformer defined on \( X \), \( (f_\tau(X)) \).

Hereafter, we use the dot notation to denote the parameters of atomic components. For example, \( B.P \) means the set of ports of the atomic component \( B \).

In order to define the operational semantics for atomic component, let us first introduce some notations. Given a Petri net \( N = (L, P, T) \) we define the set of 1-safe markings \( \mathcal{M} \).
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as the set of functions \( m : L \rightarrow \{0, 1\} \). Given two markings \( m_1, m_2 \), we define inclusion \( m_1 \leq m_2 \) iff for all \( l \in L \), \( m_1(l) \leq m_2(l) \). Also, we define addition \( m_1 + m_2 \) as the marking \( m_{12} \) such that, for all \( l \in L \), \( m_{12}(l) = m_1(l) + m_2(l) \). Given a set of places \( K \subseteq L \), we define its characteristic marking \( m_K \) by \( m_K(l) = 1 \) for all \( l \in K \) and \( m_K(l) = 0 \) for all \( l \in L \setminus K \). Moreover, when no confusion is possible from the context, we will simply use \( K \) to denote its characteristic marking \( m_K \). Finally, for a given transition \( \tau \in T \), we define its pre-set \( \tau^- \) (resp. post-set \( \tau^+ \)) as the set of the control states which are direct predecessors (resp. successors) of this transition.

**Definition 3.2.3 (Atomic component semantics.)** The semantics of an atomic component \( B = (L,P,T,X,\{g_\tau\}_{\tau \in T}, \{f_\tau\}_{\tau \in T}) \) is defined as the labeled transition system \( S_B = (Q_B, \Sigma_B, \rightarrow_B) \) where

- \( Q_B = M \times V \) is the set of states defined by:
  - \( M = \{ m : L \rightarrow \{0, 1\}\} \) the set of 1-safe markings,
  - \( V = \{ v : X \rightarrow \mathcal{D} \} \) the set of valuations of variables \( X \),
  - \( \Sigma_B = P \times \mathcal{D}^2 \) is the set of labels,
  - \( \rightarrow_B = Q_B \times \Sigma_B \times Q_B \) is the set of transitions defined by the following rule:
This rule corresponds to the firing of behavior transition. Indeed, a transition can be taken as soon as they are enabled by the marking and the guard, and update the data valuation and the marking, according to the net flow and annotations of the transition. Moreover, it performs an instantaneous data exchange through the port $p$: the current value $v^{up}$ is sent and a new value $v^{dn}$ is received for $x_p$, before the update.

### 3.2.3 Connectors and Interactions

Composition of components allows to build a system as a set of components that interact by respecting constraints of an interaction model. Connectors are used to specify possible interaction patterns between the ports of components.

A connector is a set of ports of components which can be involved in an interaction. The number of interactions of a connector can grow exponentially to the number of ports. A connector is a macro notation for representing sets of related interactions in a compact manner.

Two types of port (synchron, trigger) are defined, in order to specify the feasible interactions of a connector. A trigger is an active port, and can initiate an interaction without synchronizing with other ports. It is represented graphically by a triangle. A synchron port is passive, hence needs synchronization with other ports, and is denoted by a circle.

A feasible interaction of a connector is a set of its ports such that either it contains some trigger, or it is maximal, i.e., consisting of all the synchron ports. Example of sets of connectors and their feasible interactions are shown in Figure 3.8. By convention, triangles represent trigger and circles represent synchron ports. In the partially ordered set of interactions, the shaded nodes denote feasible interactions. In (a), the connector consists of the ports $p_1$ and $p_2$, both are of type synchron. In this connector, the only feasible interaction is $p_1p_2$. It represents a rendezvous, meaning that both actions are necessary for the synchronization. In (b), the interaction between $p_1$ and $p_2$ is asymmetric as $p_1$ is a trigger and can occur alone, even if $p_2$ is not possible. Nevertheless, the occurrence of
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\[\gamma = \{sr_1r_2r_3\}\]

Rendezvous

\[\gamma = \{s, sr_1, sr_2, sr_3, sr_1r_2, sr_2r_3, sr_1r_2r_3\}\]

Broadcast

\[\gamma = \{s, sr_1r_2r_3\}\]

Atomic broadcast

\[\gamma = \{s, sr_1r_2, sr_1r_2r_3\}\]

Causal chain

Figure 3.9: Graphic representation of connectors

\[p_2\] requires the occurrence of \[p_1\]. The feasible interactions are \[p_1\] and \[p_1p_2\]. In (c), the interactions between \[p_1\], \[p_2\] and \[p_3\] are also asymmetric. The interactions \[p_1\] can occur alone or synchronize with either or both \[p_2\] and \[p_3\].

On the other hand, connectors sometimes need to be structured, i.e., having types associated to groups of ports. This is necessary to represent some interactions. The representation of structured connectors require connectors to be treated as expressions with typing and other operations on groups of connectors. This led to the formalization of the algebra of connectors defined in [BS07]. The Algebra of Connectors is defined to provide a compact notation for algebraic representation and manipulation of connectors. The Algebra of Connectors \(AC(P)\), introduced in [BS07], formalizes the concept of connectors supported by the BIP component model. It extends the notion of connectors to terms built from a set of ports by using a \(n\)-ary fusion operator and a unary typing operator for triggers and synchrons.

Figure 3.9 shows four different coordination schemes:

- Rendezvous means strong synchronization between port \(s\) and all \(r_i\). This is specified by a single interaction involving all the ports. This interaction can occur only if all the components are in states enabling transitions labeled respectively by \(s\), \(r_1\), \(r_2\), \(r_3\).
- Broadcast means weak synchronization, that is a synchronization involving \(s\) and any (possibly empty) subset of \(r_i\). This is specified by the set of all interactions containing \(s\).
- Atomic broadcast means that either a message is received by all \(r_i\), or by none. Two interactions are possible: \(s\), when at least one of the receiving ports is not active, and the interaction \(sr_1r_2r_3\), corresponding to strong synchronization.
- Causal chain means that for a message to be received by \(r_i\) it has to be received at the same time by all \(r_j\), for \(j < i\).

For rendezvous, the priority model is empty. For all other coordination schemes, the
**maximal progress** priority model ensures that, whenever several interactions are possible, the interaction involving a maximal number of ports has higher priority.

An interaction consists of one or more ports of the connector, a guard on the variables of the ports of the interaction, two methods $\text{up}\{\}$ and $\text{down}\{\}$ realize data transfer between the ports of the interaction. The method $\text{up}\{\}$ updates the local variables of the connector based on the values of variables associated with the ports. The method $\text{down}\{\}$ updates the variables associated with the ports based on the values of the interaction variables. This structure also allows data transfer in hierarchical connectors.

**Example 2** Figure 3.11 gives an example of hierarchical connectors allowing data transfer in hierarchical manner. Let consider an initial valuation $p_1.x \mapsto 3$, $p_2.x \mapsto 5$, $p_3.x \mapsto 8$, $p_5.x \mapsto 9$ for the bottom ports. During the upward transfer, the value 9 is propagated to $v_1$ and $v_2$, following the upward predicates $p_1 p_2$ and then $p_3 p_4 p_5$. Then, during the downward transfer the value 9 gets propagated downwards to $p_1.x$, $p_2.x$, $p_3.x$ and $p_5.x$ following $p_3 p_4 p_5$ and then $p_1 p_2$.

**Definition 3.2.4 (Connector.)** A connector $\gamma = (p[x], P, A)$ is defined as follows

- $p$ is the exported port of the connector $\gamma$,
- $P = \{p_i[x_i]\}_{i \in I}$ is the support set of $\gamma$, that is, the set of ports that $\gamma$ synchronizes,
- $A \subseteq 2^P$ is a set of interactions $a = \{p_i\}_{i \in I}$ labeled by $G, U, D$ where,
  - $G$ is the guard of $\gamma$, an arbitrary predicate $G(\{x_i\}_{i \in I})$,
  - $U$ is the upward update function of $\gamma$ of the form, $x := F_u(\{x_i\}_{i \in I})$,
  - $D$ is the downward update function of $\gamma$ of the form, $\cup p_i \{x_i := F_d(x_i)\}$.

Hereafter, we use the dot notation to denote the parameters of connectors. For example, $\gamma.A$ means the set of interactions of the connector $\gamma$.

**Figure 3.10:** An example of a connector containing one interaction in BIP.
Example 3 Figure 3.10(a) shows a connector with two ports $p_1$, $p_2$, and exported port $p_3$ (allows to define hierarchical connectors). There is one and only one interaction $p_1p_2$ feasible by this connector. Synchronization through this interaction involves two steps providing its guard $G : (p_1.x > 0) ∧ (p_2.x > 0)$ is true: 1) The computation of the upward update function $U$ by assigning to $v_1$ the maximum of the values of $p_1.x$ and $p_2.x$; 2) The computation of the downward update function $D$ by assigning the value of $v$ to $p_1.x$ and $p_2.x$. Figure 3.10(b) presents the corresponding BIP code.

Definition 3.2.5 For a set of connectors $\Gamma = \{\gamma_j\}_{j \in J}$, we define the dominance relation $\rightarrow$ on $\Gamma$ as follows:

$$\gamma_i \rightarrow \gamma_j \equiv \gamma_j.p \in \gamma_i.P$$

That is, $\gamma_i$ dominates $\gamma_j$ means that the exported port of $\gamma_j$ belongs to the support set of $\gamma_i$ (see Figure 3.11). By definition, we assume that the dominance relation has no cycle.

Let $P(\Gamma) = \{p_0 \mid \gamma = (p_0, P, A) \in \Gamma\}$ be the set of their (distinct) exported ports.

Definition 3.2.6 (Interaction tree.) Let $\Gamma = \{\gamma_j\}_{j \in J}$ a set of connectors, an interaction tree $a^i = (A_\Gamma, \rightarrow_\Gamma)$ of $\Gamma$ is defined as follows:

- $A_\Gamma \subseteq \cup\{\gamma_j.A\}_{j \in J}$ is a set of interactions,
- Let $a_1, a_2 \in A_\Gamma$, we have $a_1 \rightarrow_\Gamma a_2$ iff \( \exists \gamma_1, \gamma_2 \in \Gamma \) such that $a_1 \in \gamma_1$, $a_2 \in \gamma_2$ and $\gamma_2.p \in a_1$. Moreover, $\rightarrow_\Gamma$ should satisfy the following conditions:
  - $a^i$ must contains a uniquely defined interaction $a_0 \in A_\Gamma$, called top($a^i$) from which all other interactions are recursively dependent, that is, $\exists a_0. \forall a_i \in A_\Gamma, a_0 \rightarrow^* a_i$, and $a_0$ is unique.
  - $a^i$ must contains at most one interaction per connector. That is, if $\exists a_i \in A_\Gamma \cap \gamma.A$, then, $\forall a_k \in \gamma.A \setminus a_i, a_k \notin A_\Gamma$.

Let $\Gamma$ be a set of connectors such that $(\Gamma, \rightarrow)$ has no cycle and let $a^i = (A_\Gamma, \rightarrow_\Gamma)$ be an interaction tree of $\Gamma$. We denote by:

- $A^i(\Gamma) = \{a^i_1 \mid a^i_1$ is an interaction tree of $\Gamma\}$, the set of all interaction trees of $\Gamma$;
- $\text{bottom}(a^i) = \{a_i \mid a_i \in A_\Gamma, a_i \cap P(\Gamma) = \emptyset\}$, the leaf interactions of the tree $a^i$;
- $\text{support}(a^i) = \{p \mid p \in a_i, a_i \in \text{bottom}(a^i)\}$, the support set of ports for the leaf interactions of the tree $a^i$.

Definition 3.2.7 (Hierarchical connectors semantics.) Let $\Gamma = \{\gamma_j\}_{j \in J}$ a set of connectors. Executing of interactions in $\Gamma$ implies an execution of an arbitrary interaction tree $a^i \in A^i(\Gamma)$. Moreover, it involves transfer of data between synchronizing ports. In particular, let $\sigma_0$ an initial valuation of $\text{bottom}(a^i)$, $\sigma_0 = \{p.x \rightarrow v_p \mid p \in \text{support}(a^i)\}$. The upward valuation $U_{\gamma_i}(\sigma_0)$ is obtained by propagating values from ports in the bottom interactions into the tree $a^i$ according to upward update functions of the interactions of the tree, as long as the guard conditions allow them $G_{a^i}$. In a dual manner, we define the downward
valuation $D_{a^t}(\sigma)$ obtained by transforming a given valuation $\sigma$ on ports of interactions of $a^t$ according to their downward update functions. More precisely, guards and update functions, $G_{a^t}, U_{a^t}$ and $D_{a^t}$ are defined as follows:

$$U_{a^t} = \begin{cases} U_a & a^t = \{a\}, \\ U_{a^t} \cdot U_a & a^t = a^t \setminus \{a\}, a \in \text{bottom}(a^t). \end{cases}$$

$$G_{a^t} = \begin{cases} G_a & a^t = \{a\}, \\ G_{a^t} \land G_a & a^t = a^t \setminus \{a\}, a \in \text{bottom}(a^t). \end{cases}$$

$$D_{a^t} = \begin{cases} D_a & a^t = \{a\}, \\ D_a \cdot D_{a^t} & a^t = a^t \setminus \{a\}, a \in \text{bottom}(a^t). \end{cases}$$

Definition 3.2.8 (Flat connectors.) $\Gamma$ is a set of flat connectors, iff no connector dominates another, that is, $\forall \gamma_i, \gamma_j \in \Gamma$ we have $\gamma_i \not\rightarrow \gamma_j$.

3.2.4 Priorities

Given a system of interacting components, priorities are used to filter the enabled interactions. They are given by a set of rules, each consisting of an ordered pair of interactions or connectors. When connectors are specified in a priority, the rules apply between all the respective interactions of the connectors. Dynamic priorities can be specified by providing guard condition, which are boolean expression in C on the variables of the components involved in the interactions. The maximal progress priority is enforced implicitly by the BIP Engine: if one interaction is contained in another one, the latter has higher priority. Below is an example of priority expressed in the BIP language.
3.2.5 Composite Components

Composite components are defined recursively by composition from atomic components or other composite components using glue consisting of interaction and priority models. The interface of a composite component is defined by exporting ports of subcomponents and connectors.

Definition 3.2.9 (Component.) A composite component (or simply component) \( C \) is defined by the following grammar:

\[
C ::= B|\{C_i\}_{i \in I}, \Gamma, P
\]

where,
- \( B \) is an atomic component,
- \( \{C_i\}_{i \in I} \) is a set of constituent components,
- \( P = (\cup_{i \in I} C_i.P) \cup (\cup_{j \in J}\{\gamma_j.p\}) \), is the set of ports of the component, that is \( P \) contains the ports of the constituent components and the exported ports of the connectors,
- \( \Gamma = \{\gamma_j\}_{j \in J} \) is a set of connectors, such that,
  1. \( (\Gamma, \rightarrow) \) has no cycle,
  2. \( \cup_{j \in J}\gamma_j.P \subseteq P \) (\( P \) is defined above),
  3. Each \( \gamma \in \Gamma \) uses at most one port of every constituent component, that is, \( \forall \gamma \in \Gamma, \forall i \in I, |C_i.P \cap \gamma.P| \leq 1 \).

Notice that a component is either an atomic component \( B \) or a composite component obtained as the composition of a set of constituent components \( \{C_i\}_{i \in I} \) by using a set of connectors \( \Gamma = \{\gamma_j\}_{j \in J} \). The restriction 3) is needed to prevent simultaneous firing of two or more transitions in the same atomic component, because they may affect the same variables.

Example 4 Figure 3.12(a) shows a compound component consisting of three identical atomic components described in Figure 3.7, connected by using the connector described in Figure 3.10. Each atomic component generates an integer. Then it synchronizes with all the other atomic components. During synchronization the global maximal value is computed and each atomic component receives the maximum of the values generated. Figure 3.12(b) presents the corresponding BIP code.

This specifies the priority \( p_1 \) that, when the boolean condition \( G \) is true, interactions of connector \( c_2 \) would be preferred to those of \( c_1 \).
compound type CompoundExample

component Generator comp1

component Generator comp2

component Generator comp3

connector Singelton conn1(comp1.p1)

connector Singelton conn2(comp2.p1)

connector Singelton conn3(comp3.p1)

connector Sync2Max conn4(comp1.p2,comp2.p2)

connector Sync2Max conn5(conn4.p,comp3.p2)

export port IntPort p is conn5.p end

(b) BIP code of the compound in Figure 3.12(a)

Figure 3.12: An example of compound component in BIP.

Definition 3.2.10 (Flat component.) Composite component $C$ is flat, iff the set of constituent component $\{C_i\}_{i \in I}$ are atomic components.

The operational semantics of composite components is recursively defined on the component structure. For atomic components, their semantics coincides with the semantics of the underlying behavior. For composition, the semantics is obtained by restricting the parallel behavior according to the interaction and priority models applied.

Definition 3.2.11 (Component Semantics.) The semantics of component $C$ is a labeled transition system $S_C = (Q_C, \Sigma_C, \rightarrow_C)$ defined inductively on the structure of $C$ as follows:

1. $C$ is an atomic component, defined by an atomic behavior $B = (L, P, T, X, \{g_\tau\}_{\tau \in T}, \{f_\tau\}_{\tau \in T})$. Then, $S_C = S_B$ (see Definition 3.2.3).

2. $C$ is a composite component defined as the $\langle \{C_i\}_{i \in I}, \Gamma, P \rangle$, where $C$ is flat. Let $S_{C_i} = (Q_{C_i}, \Sigma_{C_i}, \rightarrow_{C_i})$ be the semantics of its atomic components. The labeled transition system $S_C = (Q_C, \Sigma_C, \rightarrow_C)$ is defined as:
   - $Q_C = \bigotimes_{i \in I} Q_{C_i}$ is the set of states, the Cartesian product of set of states of sub-components,
   - $\Sigma_C = A^f(\Gamma)$ is the set of labels,
   - $\rightarrow_C \subseteq Q_C \times \Sigma_C \times Q_C$ is the transition relation, defined by the following rule:
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3. if $C$ is a composite component defined as the $\{C_i\}_{i \in I}, \Gamma, P$, and $C$ is not flat. Then, the same principle as above can be defined.

Example 5 Figure 3.13 illustrates an abstract overview of the semantics of composite component. The composed behavior is shown in the right. It shows the product of the two behaviors, where the only allowed transitions are the ones with a solid arrow. The dotted transitions are not legal and shows the maximal behavior allowed by the interactions glue.

3.3 Execution Platform

The operational semantics is implemented by an Engine. In the basic implementation, the Engine computes the enabled interactions by enumerating over the complete list of interactions in the model (enumerative Engine). Another implementation is based on computing boolean representation for components and connectors by using an existing BDD package (symbolic Engine)\(^1\).

\(^1\) The implementation of the boolean functions is made using the BDD package CUDD.
3.3.1 Enumerative Engine

During the execution, on each iteration of the Engine, the enabled interactions are selected from the complete list of interactions, based on the current state of the atomic components. Then, between the enabled interactions, priority rules are applied to eliminate the ones with low priority. The main loop of the Engine consists of the following steps:

1. Each atomic component sends to the Engine its current state.
2. The Engine enumerates on the list of interactions in the model, selects the enabled ones based on the current states of the atomic components and eliminates the ones with low priority.
3. Amongst the enabled interactions, the Engine selects any one and notifies the involved atoms the transition to take.

The time to compute the enabled interactions by Engine is proportional to the number of interactions in the model.

3.3.2 Symbolic Engine

In the enumerative BIP Engine [JBB09], for each connector, the Engine needs to compute all the possible interactions, check which ones are enabled in the current global state of the system, and select a maximal enabled one to be executed. As interactions are sets of ports, their number is potentially exponential in the number of ports in the connector. Hence, in the worst case, the performance of this Engine can be extremely poor.

The boolean BIP Engine leverages on representing component behavior, connector interactions, and priorities as boolean functions. For an atomic component, all ports and control states are represented by boolean variables. This allows to encode behavior as a boolean expression of these variables. Similarly, each connector is represented by the boolean expression on its ports. The global behavior is obtained as a boolean operation on the expressions representing atomic components, connectors, and priorities.

The choice of an interaction to be executed boils down to evaluating the control states, substituting their respective boolean variables, and picking a valuation of the port variables satisfying the boolean expression that represents the global behavior.

The boolean representation of connectors replaces the costly enumeration step by efficient BDD manipulations. In comparison to the exponential cost of the enumerative Engine, this renders a more efficient Engine with evaluation that, in practice, remains linear.

3.4 The BIP Tool-Chain

The BIP tool-chain provides a set of tools for the modeling, the execution, the verification and the static transformation of BIP models. The overview of the BIP tool-chain is
shown in Figure 3.14. It includes the following tools:

- An editor, for describing textually a system in BIP language.
- A compiler, for generating a BIP model from BIP description source.
- A code generator, for generating, from a model, C++ code executable on the BIP Engine. The code-generator can also produce THINK specification [Pou10, PPRS06], from which the Think tool-chain can generate code to be executed over a choice of target platforms.
- D-Finder, is a compositional verification tool for component-based systems described in BIP language [BBNS09, BBSN08].
- Source-to-Source transformations, of which the method and the implementation are presented in this thesis, allow useful transformations which generate efficient central-
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ized [BJS09] and distributed implementation from a composite component [BBJ+10b, BBJ+10a].

- An exporter to connect with external tools such as IF toolbox or analysis tools.
- A set of translators from other languages (Lustre, MATLAB/Simulink, AADL, etc.) to BIP. For example, a Simulink-to-BIP translation [STS+10] from Simulink models into BIP which allows the validation and implementation of Simulink models. In particular, compositional and incremental generation of invariants can be applied for complex Simulink models. These compilation paths are also becoming available for Simulink models. Moreover, Simulink models can be explored using the compilation paths of BIP. Other example of translations is AADL-to-BIP translation from Architecture Analysis & Design Language (AADL) into BIP [CRBS08], allows simulation of systems specified in AADL and application to these systems of formal verification techniques developed for BIP, e.g. deadlock detection.

3.5 Summary

Component-based approach is aimed to deal with the complexity of systems. It is based on the idea of building a complex system by assembling basic components (blocks). It provides important characteristics for system construction such as reuse, incrementality, compositionality, etc. It allows not only the reuse of components but also the reuse of known properties of constituent components.

We have presented BIP, a component-based framework for modeling heterogeneous systems. The BIP component model is the superposition of three layers: the lower layer describes the behavior of a component as a transition system; the intermediate layer consists of the interactions between transitions of the layer underneath; the upper layer describes the priorities characterizing a set of scheduling policies for interactions. Such a layering offers a clear separation between components’ behaviors and the structure of the system (interactions and priorities).

BIP modeling framework allows dealing with complexity of systems by providing incremental composition of heterogeneous components. It also considers correctness-by-construction for a class of essential properties such as deadlock-freedom [GS05].

The BIP tool-chain has been developed providing automated support for component integration and generation of glue code meeting given requirements. Efficient model transformations, verification methods have also been studied and implemented in the BIP tool-chain.

We are now going to present a method for generating efficient centralized implementations from BIP models. We will also show applications of our method for MPEG4 encoder and network sorting algorithm described in the BIP language.
CHAPTER 4

Transformation for Generating Centralized Implementations

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4.1 Problem Statement

Efficient implementation of component-based systems is a non-trivial task. Moreover, clarity of models may be at the detriment of efficiency. Indeed, naive compilation of component-based systems results in great inefficiency as a consequence of the interconnection of components [Lov77]. Nowadays, it is widely admitted that modularity in component-based development incurs an additional non-negligible overhead for implementation because of extensive use of interfaces, wrappers and other implementation artifacts. For instance, the generated BIP code is modular and can be executed on a dedicated platform consisting of an Engine which orchestrates the computation of atomic components.
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by executing their interactions (described in Section 3.3). Hierarchical description allows incremental reasoning and progressive design of complex systems. Nonetheless, it may lead to inefficient programs if structure is preserved at run time. Compared to functionally equivalent monolithic C programs, BIP programs may be more than two times slower. This overhead is due to the computation of interactions between components by the Engine.

The aim of this chapter is to show that it is possible to synthesize efficient monolithic code from component-based software described incrementally. We study source-to-source transformations for BIP allowing the composition of components and thus leading to more efficient code. These are based on the operational semantics of BIP which allows to compute the meaning of a composite component as a behaviorally equivalent atomic component. Thus, we show how by incremental composition of the components contained in a composite component, a behaviorally equivalent component can be computed. This composition operation has been implemented in the BIP2BIP tool, by using three types of source-to-source transformations. A set of interacting components is replaced by a functionally equivalent component. By successive application of transformations, an atomic component can be obtained, that is a component with no interactions.

The transformation from a composite component to an atomic one is fully automated and implemented through three steps:

1. **Component flattening** which replaces the hierarchy on components by a set of hierarchically structured connectors applied on atomic components;
2. **Connector flattening** which computes for each hierarchically structured connector an equivalent flat connector;
3. **Component composition** which composes atomic components to get an atomic component.

Using such a transformation allows to combine advantages of component-based descriptions such as clarity and reuse with efficient implementation. The generated code is readable and by-construction functionally equivalent to the component-based model. We show through non trivial examples the benefits of this approach.

The rest of the chapter is organised as follows. First in Section 4.2 we define the three source-to-source transformations. In section 4.4, we provide benchmarks for two examples: a MPEG encoder and a concurrent sorting program. In Section 4.5, we finish this chapter by giving conclusions in summary.

### 4.2 Transformations

In this section, we will define the transformations which successively transform a composite component into atomic components. That is, they eliminate component hierarchy and the hierarchical connectors by computing the product behavior. The transformation
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from a composite component to an atomic one involves three steps: Component flattening, Connector flattening, Component composition. In this section, we describe the three transformations, and we illustrate them on the example shown in Figure 4.1. This example consists of two composite components ($C_1$ and $C_2$). Each one of these composite components consists of three identical atomic components described in Figure 4.1-c, connected by using the connector described in Figure 4.1-b. Each atomic component generates an integer. Then it synchronizes with all the other atomic components. During synchronization the global maximal value is computed and each atomic component receives the maximum of the values generated.

![Figure 4.1: Example.](image)

4.2.1 Components Flattening

This transformation replaces the hierarchy on components by a set of hierarchically structured connectors applied on atomic components. Consider a composite component $C$, obtained as the composition of a set of components $\{C_i\}_{i \in I}$. The purpose of this transformation is to replace each non atomic component $C_j$ of $C$ by its description. By successive applications of this transformation, the component $C$ can be modelled as the set of its atomic components and their hierarchically structured connectors (see Figure 4.2).

**Definition 4.2.1 (Component flattening.)** Consider a non atomic component $C = (\{C_i\}_{i \in I}, \Gamma, P)$ such that there exists a non atomic component $C_j \in \{C_i\}_{i \in I}$ with $C_j = (\{C_{jk}\}_{k \in K}, \Gamma_j, P_j)$. We define $C[C_j \mapsto \Gamma_j]$ as the component $C = (\{C_i\}_{i \in I} \cup \{C_{jk}\}_{k \in K} \setminus \{C_j\}, \Gamma \cup \Gamma_j, P)$. Component flattening is defined by the following function:
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Figure 4.2: Component flattening.

\[
F_c(C) = \begin{cases} 
C & \text{if } C \text{ is flat} \\
F_c(C[C_j \mapsto \Gamma_j]) & \text{if } C \text{ is not flat}
\end{cases}
\]

**Proposition 1** Component flattening is well-defined i.e., \( F_c \) is a function which produces a unique result on every input component, and terminates in a finite number of steps.

**Proof** Regarding unicity of the result, we can show that, if two constituent components respectively \( C_j \) and \( C_k \) can be replaced inside the composite component \( C \), then the replacement can be done in any order and the final result is the same. That is, formally we have \( C[C_j \mapsto \Gamma_j][C_k \mapsto \Gamma_k] = C[C_k \mapsto \Gamma_k][C_j \mapsto \Gamma_j] \). The result follows immediately from the definition and elementary properties of union on sets.

Regarding termination, every transformation step decreases the overall number of composite components by one, so component flattening eventually terminates when all the components are atomic.

By applying to Example in Figure 4.1 the transformation \( C[C_1 \mapsto \{\gamma_2, \gamma_3\}] \) then \( C[C_2 \mapsto \{\gamma_4, \gamma_5\}] \), we obtain the new component in Figure 4.3.

Finally, notice that this transformation never increases the structural complexity of the transformed component. The transformation does not change the set of atomic components as well as the set of the hierarchical connectors. Hence, it preserves the operational semantics of the original model.

### 4.2.2 Connectors Flattening

This transformation flattens hierarchical connectors. It takes two connectors \( \gamma_i \) and \( \gamma_j \) with \( \gamma_i \rightarrow \gamma_j \) (recall that the dominance relation \( \rightarrow \) is given in Definition 3.2.5 in Chapter 3) and produces an equivalent connector.

We show in Figure 4.4 the composition of two connectors \( \gamma_i \) and \( \gamma_j \). It consists in “glueing” them together on the exported port \( p_j \). For the composite connector, the update functions
are respectively, the bottom-up composition of the upward update functions, and the top-down composition of the downward update functions. This implements a general two-phase protocol for executing hierarchical connectors. First, data is synthesized in a bottom up fashion by executing upward update functions, as long as guards are true. Second, data is propagated downwards through downward update functions, from the top to the support set of the connector.

Figure 4.4: Connector glueing.

**Definition 4.2.2 (Connector glueing.)** Given connectors $\gamma_i = (p_i[x_i], P_i, A_i)$ and $\gamma_j = (p_j[x_j], P_j, A_j)$ such that $\gamma_i \rightarrow \gamma_j (p_j \in P_i)$ we define the composition $\gamma_i[p_j \rightarrow \gamma_j]$ as a connector $\gamma = (p, P, A)$ where
- $p = p_i$,
- $P = P_j \cup P_i \setminus \{p_j\}$,
- $A = A_1 \cup A_2$, where,
  - $A_1 = \{a_i \mid a_i \in A_i, p_j \notin a_i\}$,
  - $A_2 = \{a_i \setminus \{p_j\} \cup a_j \mid a_i \in A_i, p_j \in a_i, a_j \in A_j\}$

If $a \in A_1$ the guards and data transfer are inherited as such from $\gamma_i$. In the second case, if $a \in A_2$ the guard and the transfer are defined as follows:
- $G_a = G_{a_j} \land G_{a_j}[U_{a_j}/x_j]$,
- $U_a = x_i := U_{a_j}[U_{a_j}/x_j]$,
\[- D_a = (\bigcup_{p_k \in a_j} x_k := D_{a_j,x_i/x_i}) \cup (\bigcup_{p_k \in a_i \setminus \{p_j\}} x_k := D_{a_i,x_i/x_i}) \]

Intuitively, by composition, two linked connectors are glued together into a single connector. Their guards, respectively the upward and downward transfer functions are composed. Consequently, any port valuation obtained by the successive application of the upward (resp. downward) transfer predicates of the two connectors is equally obtained by the application of the upward (resp. downward) transfer predicate of the composed connector.

Let us introduce some notations. Let \( \Gamma = \{ \gamma_i = (p_i[x], P_i, A_i) \mid i \in I \} \) a set of connectors, and let \( P = \{ \{p_i\} \cup P_i \mid i \in I \} \) the set of all used ports. We call a port \( p_j \) transient in \( \Gamma \) if it is both exported by some connector \( \gamma_j \) from \( \Gamma \) and used by another connector \( \gamma_i \) from \( \Gamma \). Obviously, transient ports can be eliminated through connector glueing.

For a transient port \( p_j \) exported by a connector \( \gamma_j \), we will use the notation \( \Gamma[p_j \mapsto \gamma_j] \) to denote the new set of connectors obtained by replacing thoroughly \( p_j \) by its exporting connector \( \gamma_j \), formally: \( \Gamma[p_j \mapsto \gamma_j] = \{ \gamma \mid \gamma \in \Gamma, p_j \notin \gamma.ports, \gamma \neq \gamma_j \} \cup \{ \gamma[p_j \mapsto \gamma_j] \mid \gamma \in \Gamma, p_j \in \gamma.ports \} \). That is, all connectors (except \( \gamma_j \)) without \( p_j \) in their support set are kept unchanged, while the others are transformed according to definition 4.2.2.

**Definition 4.2.3 (Connector flattening.)** Connector flattening is defined by the following function:

\[
\mathcal{F}_\gamma(\Gamma) = \begin{cases} 
\Gamma & \text{if } \Gamma \text{ is a set of flat connectors} \\
\mathcal{F}_\gamma(\Gamma[p_j \mapsto \gamma_j]) & \text{if } \Gamma \text{ is not a set of flat connectors, } p_j \text{ is a transient port of } \Gamma 
\end{cases}
\]

**Proposition 2** Connector flattening is well-defined i.e., \( \mathcal{F}_\gamma \) produces a unique result for any set of connectors, and terminates in a finite number of steps.

**Proof** Regarding unicity of the result, if \( p_j \) and \( p_k \) are two transient ports of \( \Gamma \) defined respectively by connectors \( \gamma_j \) and \( \gamma_k \), then flattening in any order gives the same result, formally \( \Gamma[p_j \mapsto \gamma_j][p_k \mapsto \gamma_k] = \Gamma[p_k \mapsto \gamma_k][p_j \mapsto \gamma_j] \).

To show this result it is sufficient to show that any connector \( \gamma \) of \( \Gamma \), different from \( \gamma_j \) and \( \gamma_k \) gets transformed in the same way, independently of the order of application of the two transformations. This can be shown, case by case, depending on the occurrence of ports \( p_j \) and \( p_k \) in the supports of \( \gamma \), \( \gamma_j \) and \( \gamma_k \) following definition 4.2.2.

Regarding termination, flattening of connectors is applicable as long as there are transient ports. Moreover, it can be shown that, every flattening step reduces the number of transient ports by one - the one that is replaced by its definition. Hence, flattening eventually terminates when no more transient ports exist, that is, \( \Gamma \) is a set of flat connectors.
By application of the transformation $\gamma_1[p_2 \mapsto \gamma_2]$ to Example 2 in Figure 4.3, we obtain the new composite component given in Figure 4.5. If we apply successively, $\gamma_1[p_3 \mapsto \gamma_3], \gamma_1[p_4 \mapsto \gamma_4], \gamma_1[p_5 \mapsto \gamma_5]$ we obtain the new composite component given in Figure 4.6.

In a similar way to component flattening, this second transformation does not increase the structural complexity of the transformed components. The set of atomic components is preserved as such, whereas, the overall set of connectors is decreasing. However, the remaining connectors have an increased computational complexity, because they integrate the guards and the data transfer of the eliminated ones. The operational semantics is also
preserved. The effect of the eliminated connectors is "in-lined" in the remaining according to definition 3.2.7.

### 4.2.3 Components Composition

We present the third transformation which allows to obtain a single atomic component from a set of atomic components and a set of flat connectors. This transformation defines the composition of behaviors.

Intuitively, as shown in Figure 4.7, the composition operation consists in "glueing" together transitions from atomic components that are synchronized through the interaction of some connector (interaction $p_1p_2$ for this example). Guards of synchronized transitions are obtained by conjuncting individual guards and the guard of the connector. Similarly, actions of synchronized transitions are obtained as the sequential composition of the upward update function followed by the downward update function of the connector, followed by the actions of the components in an arbitrary order.

**Definition 4.2.4 (Component composition.)** Consider a component $C = (\{B_i\}_{i \in I}, \Gamma, P)$ such that $\forall i \in I B_i$ is an atomic component and $\Gamma$ is a set of flat connectors. We define the composition $\Gamma(\{B_i\}_{i \in I})$ as component $B = (L, P, T, X, \{g_\tau\}_{\tau \in T}, \{f_\tau\}_{\tau \in T})$ defined as follows:

- the set of control states $L = \bigcup_{i \in I} B_i.L$,
- the set of ports $P = \bigcup_{\gamma \in \Gamma} \{\gamma.p\}$,
- the set of variables $X = (\bigcup_{i \in I} B_i.X) \cup (\bigcup_{\gamma \in \Gamma} \gamma.p.x)$,
- each transition in $T$ corresponds to a set of interacting transitions $\{\tau_1, \ldots, \tau_k\} \subseteq \bigcup_{i \in I} T_i$ such that $\bigcup_{i=1}^k \tau_i.p = a$ ($\gamma \in \Gamma, a \in \gamma.A$). We define the transition $\tau = (l, \gamma.p, l')$ where,
  - $l = \ast \tau_1 \cup \ldots \cup \ast \tau_k$,
  - $l' = \tau_1^\ast \cup \ldots \cup \tau_k^\ast$,
  - the guard $g_\tau = \bigwedge_{i=1}^k g_{\tau_i} \land a.G$,
  - the action $X := f_\tau(X)$ with $f_\tau = a.U; a.D; (\bigcup_{i=1}^k f_{\tau_i})$.

Figure 4.8 shows the Petri net obtained by composition of the atomic components of Figure 4.6 through the interaction $r_1r_2r_3r_4r_5r_6$.

In contrast to previous transformations, component composition may lead to an exponential blowup of the number of transitions in the resulting Petri net. This situation may happen if the same interaction can be realized by combining different transitions from each one of the involved components. For instance, the interaction $p_1p_2$ can give rise to four transitions in the resulting Petri net if there are two transitions labeled by $p_1$ and $p_2$ in the synchronizing components. Nevertheless, in practice exponential explosion seldom occurs, as in atomic components each port labels at most one transition (as in the examples shown hereafter). In this case, the resulting Petri net has as many transitions as connectors in $\Gamma$. 

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\[ p_2 \] \quad [p_1, p_2] = [g_1, f_1] \quad G_{12} \quad [U_{12}, D_{12}] \quad p_1 \]

\[ g_{12} = g_{12} \land g_1 \land g_2 \quad f_{12} = U_{12}, D_{12} : (f_1 \cup f_2) \]

Figure 4.7: Component composition.

\[ t_1 := r_{12} \quad t_2 := r_{12} \quad t_3 := r_{12} \quad t_4 := r_{12} \quad t_5 := r_{12} \quad t_6 := r_{12} \]

\[ a_1 := \text{rand()} \quad a_2 := \text{rand()} \quad a_3 := \text{rand()} \quad a_4 := \text{rand()} \quad a_5 := \text{rand()} \quad a_6 := \text{rand()} \]

\[ U : x_1 := \max(\max(a_1, a_2), a_3) \quad \max(\max(a_4, a_5), a_6)) \]

\[ D : a_1 := x_1; a_2 := x_1; a_3 := x_1; a_4 := x_1; a_5 := x_1; a_6 := x_1 \]

\[ f : \text{print}(a_1) \cup \text{print}(a_2) \cup \text{print}(a_3) \cup \text{print}(a_4) \cup \text{print}(a_5) \cup \text{print}(a_6) \]

Figure 4.8: Component composition for example in Figure 4.6.
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4.3 Efficient Sequential Implementation

By exhaustive application of these transformations, an atomic component can be obtained. From the latter, the code-generator can generate efficient standalone C++ code, which can be run directly without the Engine. In particular, all the remaining non-determinism in the final atomic component is eliminated at code generation by applying an implicit priority between transitions. It should be noted that the transformations also can be applied independently, to obtain models that respond to a particular user needs. For example, one may decide to eliminate only partially the hierarchy of components, or to compose only some components.

4.4 Experimental Validation

These transformations have been implemented in the BIP2BIP tool which is currently integrated in the BIP toolset [BIP]. A detailed description of the BIP2BIP tool is given in Chapter 6.

For two examples, we compare the execution times of BIP programs before and after flattening. These examples show that it is possible to generate efficient standalone C++ code from component-based descriptions in BIP.

4.4.1 MPEG Video Encoder

In the framework of an industrial project, we have componentized in BIP an MPEG4 encoder written in C by an industrial partner. The aim of this work was to evaluate gains in scheduling and quality control of the componentized program. The results were quite positive regarding quality control [CFLS05a, CFLS05b, CFSS07] but the componentized program was almost two times slower than the handwritten C program. We have used BIP2BIP to generate automatically standalone C++ code from the BIP program as explained below (see Figure 4.10).

The BIP program consists of 11 atomic components, and 14 connectors. It uses the data and the functions of the initial handwritten C program. It is composed of two atomic
Figure 4.10: Encode component structure.
components and one composite component. The atomic component **GrabFrame** gets a frame and produces macroblocks (each frame is split into N macroblocks of 256 pixels). The atomic component **OutputFrame** produces an encoded frame. The composite component **Encode** consists of 9 atomic components and the corresponding connectors. It encodes macroblocks produced by the component **GrabFrame**.

Figure 4.11 shows execution times for the initial handwritten C code, for the BIP program and the corresponding standalone C++ code generated automatically by using the presented technique. Notice that the automatically generated C++ code and the handwritten C code have almost the same execution times. The advantages from the componentization of the handwritten code are multiple. The BIP program has been rescheduled as shown in [CFLS05a] so as to meet given timing requirements. Table 4.1 gives the size of the handwritten C code, the BIP model, as well of the generated C++ code from the BIP model \(C^{(1)}\) and the generated C++ code from the BIP model after flattening \(C^{(2)}\). The time taken by the BIP2BIP tool to generate automatically \(C^{(2)}\) is less than 1sec.

<table>
<thead>
<tr>
<th></th>
<th>Handwritten</th>
<th>BIP</th>
<th>(C^{(1)})</th>
<th>(C^{(2)})</th>
</tr>
</thead>
<tbody>
<tr>
<td>loc</td>
<td>600</td>
<td>350</td>
<td>1800</td>
<td>800</td>
</tr>
</tbody>
</table>

Table 4.1: Code size in lines-of-code (loc) for MPEG4 encoder.

### 4.4.2 Concurrent Sorting

This example is inspired from a network sorting algorithm [AKS83]. We consider \(2^n\) atomic components, each of them containing an array of \(N\) values. We want to sort all the values, so that the elements of the first component are smaller than those of the second component and so on. We solve the problem by using incremental hierarchical composition of components with particular connectors.

In Figure 4.12, we give a model for sorting the elements of 4 atomic components. The components \(C_1\) and \(C_2\) are identical. The pair \((B_1, B_2)\) is composed by using two connectors \(\gamma_1\) and \(\gamma_2\) to form the composite component \(C_1\). Each atomic component computes the minimum and the maximum of the values in its array. These values are then exported on port \(p\). The connector \(\gamma_1\) is used to compare the maximum value of \(B_1\) with the minimum value of \(B_2\), and to permute them if the maximum is bigger than the minimum value.

When the maximum value of \(B_1\) is smaller than the minimum value of \(B_2\), that is the components are correctly sorted, then the second connector \(\gamma_2\) is triggered. It is used to export the minimum value of \(B_1\) and the maximum value of \(B_2\) to the upper level. At this level the same principle is applied to sort the values of the composite components \(C_1\) and \(C_2\). This pattern can be repeated to obtain arbitrary higher hierarchies (see Figure 4.13).
Figure 4.11: Execution time for the MPEG4 encoder.
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Figure 4.12: Concurrent sorting $n = 2$.

Figure 4.13: Concurrent sorting $n = 4$. 

$G : p.\text{Max} > q.\text{Min}$
$U :$
$D : x := p.\text{Max}; p.\text{Max} := q.\text{Min}; q.\text{Min} := x;$

$G : p.\text{Max} \leq q.\text{Min}$
$U :$
$D : p.\text{Min} := Min; q.\text{Max} := Max.$
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\begin{table}
\centering
\begin{tabular}{|c|c|c|c|}
\hline
$n$ & BIP$^{(1)}$ & C$^{(1)}$ & C$^{(2)}$ \\
\hline
2 & loc 112 & 360 & 400 \\
3 & loc 120 & 400 & 620 \\
4 & loc 128 & 440 & 1100 \\
5 & loc 136 & 480 & 1850 \\
6 & loc 144 & 520 & 2850 \\
\hline
\end{tabular}
\caption{Code size in loc for concurrent sorting.}
\end{table}

Figure 4.14 shows the execution times for the hierarchically structured BIP program and for the corresponding standalone C++ code generated automatically by using the presented technique. Notice the exponentially increasing difference between the execution time of the component-based BIP program and the corresponding C++ code. In particular, component flattening and connector flattening do not provide much better performance, because the hierarchical structure is actually exploited by the BIP Engine to compute enabled interactions in an efficient manner. However, these transformations are mandatory for applying the static composition.

Notice that the overhead is due to many reasons when using the BIP Engine. First, each atomic components sends to the Engine its current state and the list of enabled ports. Second, the Engine enumerates on the list of interactions in the model, identifies all enabled ones based on the current state of the atomic components, then among them it selects one for execution and, finally, notifies atoms to take the corresponding transition. This overhead is partially eliminated in the standalone C++ code generated automatically. Indeed, the call function between components and the Engine is omitted. The time needed to select an enabled interaction is drastically reduced. Moreover, control and code optimization such as guard combination, removal of unnecessary assignments, etc., are applied.

Table 4.2 shows the size in lines of code of the BIP model, as well of the generated C++ generated from the BIP model C$^{(1)}$ and the generated C++ code from the BIP model after flattening C$^{(2)}$, for 4, 8, 16, 32 and 64 atomic components. The size of the BIP model changes only linearly with $n$. However, we notice that for this example, the size of the generated C++ code from the BIP model is much smaller than the generated C++ code from the BIP model after flattening. This is due to the use of component types and component types instantiation. In particular, for this example, the initial BIP model contains just one component type instantiated, respectively 4, 8, 16, 32, 64 times for $n = 2, 3, 4, 5, 6$. However, the BIP model after flattening, contains one component types with one instance each. The size of the generated code is directly dependent on the number of component types and not on the number of component types instance.
Figure 4.14: Execution time for concurrent sorting.
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4.5 Summary

In this chapter we presented a technique for generating efficient centralized implementation running on a single-core platforms. The technique is based on source-to-source transformations. We defined three transformations: Component flattening, Connector flattening, Component composition. The aim of these transformation is to transform a composite component into a single atomic component. From the latter an efficient code can be generated.

Moreover, we shown that it is possible to reconcile component-based incremental design and efficient code generation by applying a paradigm based on the combined use of:

1. a high level modelling notation based on well-defined operational semantics and supporting powerful mechanisms for expressing structured coordination between components;

2. semantics-preserving source-to-source transformations that progressively transform architectural constraints between components into internal computation of product components.

This paradigm opens the way to the synthesis of efficient monolithic software which is correct-by-construction by using the design methodology supported by BIP.

We are now going to present a method for generating efficient distributed implementations from BIP models. We will also show applications of our method on non trivial example described in the BIP language.
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CHAPTER 5

Transformation for Generating Distributed Implementations

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In the previous chapter, we present a method to generate an efficient centralized implementation from a high-level BIP model. Centralized implementations can be useful when the target platform consists of a single-processor. However, nowadays, most applications themselves can be geographically distributed, on the other hand, most of target platforms are multi-processor boards. Thus, a distributed and parallel implementation should be derived from the high-level BIP model. In this chapter, we propose a methodology for producing automatically efficient and correct-by-construction distributed implementations from a high-level BIP models.

5.1 Problem Statement

Deriving from a high-level model a correct and efficient distributed implementation, that allows parallelism between components as well as parallel execution between interactions, is a challenging problem. As adding implementation details involves many subtleties (e.g., inherently concurrent, non-deterministic, and non-atomic structure of distributed systems) that can potentially introduce errors to the resulting system.

In order to understand the subtleties of transformation from high-level BIP model to distributed implementations, consider the BIP model in Figure 5.1. In this model, atomic components $B_1 \cdots B_5$ synchronize through four rendezvous interactions $a_1 \cdots a_4$. In sequential implementation, interactions are executed atomically by a centralized Engine. On the contrary, introducing concurrency and distribution (and possibly multiple Engines) to this model requires dealing with more complex issues:

- (Partial observability) Suppose interaction $a_1$ (and, hence, components $B_1 \cdots B_3$) is being executed. If component $B_3$ completes its computation before $B_1$ and $B_2$, and, ports $p_4, p_5$ are enabled, then interaction $a_2$ is enabled. In such a case, distributed Engines must be designed so that concurrent execution of interactions does not intro-
duce behaviors that were not allowed by the high-level model. We address the issue of partial observability by breaking the atomicity of execution of interactions, so that a component can execute unobservable actions once a corresponding interaction is being executed [BBBS08].

- (Resolving conflicts) Suppose interactions $a_1$ and $a_2$ are enabled simultaneously. Since these interactions share component $B_3$, they cannot be executed concurrently. We call such interactions conflicting. Obviously, distributed Engines must ensure that conflicting interactions are mutually exclusive.

- (Performance) On top of correctness issues, a real challenge is to ensure that a transformation does not add considerable overhead to the implementation. After all, one crucial goal of developing distributed and parallel systems is to exploit their computing power.

### 5.2 Original BIP Model

In this chapter, we consider that the original BIP component consists only of atomic component and flat connectors. Moreover, each connector defines one and only one interaction (see Figure 5.1). Indeed, these assumptions does not impose any restrictions on the original model, since we can apply the first two transformation Component Flattening and Connector Flattening described in Section 4.2. Hence, we obtain a model which meets these assumptions. Under these assumptions, a composite component $C$ can be denoted as $C = \gamma(B_1, B_2, \ldots, B_n)$ where $B_1, B_2, \ldots, B_n$ is a set of atomic components and $\gamma$ is a set of interactions.

Recall that an interaction is defined as follows:

**Definition 5.2.1 (Interaction.)** An interaction $a$ is given by a tuple $(P, G, F)$ such that:

- $P$ is a set of ports $P = \{p_i[x_i]\}_{i \in I}$,
- $G$ is a guard (a predicate on the variables $x_i$),
- $F$ is an update function defined on the variables $x_i$.

Furthermore, let $B = (L, P, T, \{g_\tau\}_{\tau \in T}, \{f_\tau\}_{\tau \in T})$ be an atomic component. Hereafter, we assume that $T \subseteq L \times P \times L$. That is, the behavior of an atomic component is described as labeled transition system. For any pair of control states $l, l' \in L$ and a port $p \in P$, we write $l \xrightarrow{p} l'$ iff $(l, p, l') \in T$. When the communication port is irrelevant, we simply write $l \rightarrow l'$. Similarly, $l \xrightarrow{p}$ means that there exists $l' \in L$ such that $l \xrightarrow{p} l'$.

### 5.3 Conflicting Interactions

In order to understand what is a conflicting interaction, let us first recall how the execution of interactions and local code of components in a model are orchestrated by the
centralized Engine (see Figure 5.2):

1. Each atomic component sends to the Engine the set of enabled ports based on the current state of the atomic components.
2. The Engine enumerates the list of interactions in the model, selects the enabled ones.
3. The Engine selects any one from the enabled ones after eliminating the ones with low priority.
4. Finally, the Engine notifies the involved atomic components the transition to take and wait for completion.

The operational semantics of the BIP framework is handled by the centralized Engine. Nonetheless, introducing concurrency and distribution between interactions by adding multiple Engines requires dealing with more complex issues. For instance, the system must respect the global state semantics although it works in a distributed setting where components do not have a global view of the system. Moreover, suppose that interaction $a_1$ and $a_2$, in the example in Figure 5.1 are enabled simultaneously. Since these interactions share the same component $B_3$, they cannot be executed concurrently such as the example in Figure 5.3. We call such interactions conflicting. Obviously, distributed Engines must ensure that conflicting interactions are mutually exclusive.

**Definition 5.3.1 (Conflict interaction.)** Let $\gamma(B_1, \ldots, B_n)$ be a BIP component. We say that two interactions $a_1, a_2 \in \gamma$ are conflicting iff:
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Figure 5.3: Violating of BIP semantics when badly distributed interactions.

Figure 5.4: Conflict interactions.

- either, they share a common port $p$; i.e., $p \in a_1 \cap a_2$ (see Figure 5.4-a),
- or, there exist an atomic component $B_i = (L_i, P_i, T_i, \{g_{\tau_i}\}_{\tau_i \in T_i}$, $\{f_{\tau_i}\}_{\tau_i \in T_i}$), a control state $l \in L_i$, and two ports $p_1, p_2 \in P_i$ such that (1) $p_1 \in a_1$, (2) $p_2 \in a_2$, and (3) $l \xrightarrow{p_1} \land l \xrightarrow{p_2}$ (see Figure 5.4-b).

Therefore, introducing concurrency by adding multiple Engines requires to resolve conflict between conflicting interactions.

The first straightforward solution consists of generation distributed Engines which is conflict-free by construction. This is done by grouping interactions according to the transitive closure of the conflict relation in the same Engine. In this case, we do not need communications in order to safely execute interactions of the high-level model.

However, this solution has a drawback, because grouping conflicting interactions according to the transitive closure reduces drastically parallelism between interactions. In
other words, two interactions which are not in direct conflict cannot run in parallel. For instance, let us consider the example in the Figure 5.1. Assuming that \( a_1 \) is conflict only with \( a_2 \), and, interaction \( a_2, a_3, a_4 \) are pairwise conflict. This situation leads to create only one centralized Engine handling all the interactions (see Figure 5.5), and then, it is not possible to run the interaction \( a_1 \) and \( a_3 \) in parallel, despite that it is possible without violating the operational semantics of BIP.

For this reason, we need a solution which solve conflict dynamically by using some protocols. Indeed, resolving conflicts leads us to solving the committee coordination problem [CM88], where a set of professors organize themselves in different committees and two committees that have a professor in common cannot meet simultaneously. The original distributed solution to the committee coordination problem assigns one manager to each interaction [CM88]. Conflicts between interactions are resolved by reducing the problem to the dining or drinking philosophers problems [CM84], where each manager is mapped onto a philosopher. Bagrodia [Bag87] proposes an algorithm where message counts are used to solve synchronization and exclusion is ensured by using a circulating token. In a follow-up paper [Bag89], Bagrodia modifies the solution in [Bag87] by combining the use of message counts to ensure synchronization and reducing the conflict resolution problem to dining or drinking philosophers problems. Also, Perez et al [PCT04] propose an approach, Alpha-Core Protocol, that essentially implements the same idea using a lock-based synchronization mechanism. For instance, if we consider the algorithm proposed by Perez et al, this leads to embedding the protocol into the high-level BIP model as it is presented in the Figure 5.6. Notice that, using such methodology the modification of protocol requires to do a modification in the different layers.

Thus, several distributed algorithms exist in the literature for conflict resolution, moreover the performance of each of them depends on the type of the application. For this reason, we need to design our framework, so that it provides appropriate interfaces with minimal restrictions.
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5.4 Proposed Solution

With this motivation we propose a generic framework for transforming high-level BIP models into a distributed implementation that allow parallelism between components as well as parallel execution of non-conflicting interactions by embedding a solution to the committee coordination problem. To the best of our knowledge, this is the first instance of such a transformation (the related work mentioned above only focus on impossibility results, abstract algorithms, and in one instance [Bag89] simulation of an algorithm). Our method utilizes the following sequence of transformations preserving observational equivalence:

1. First, we transform the given BIP model into another BIP model that (1) operates in partial-state semantics, and (2) expresses multi-party interactions in terms of asynchronous message passing (Send/Receive primitives). Moreover, the target BIP model is structured in three layers:
   (a) The Atomic Components Layer consists of a transformation of behavioral components in the original model.
   (b) The Interaction Protocol Layer detects enabledness of interactions of the original model and executes them after resolving conflicts either locally or by the help of the third layer. This layer consists of a set of components, each hosting a user-defined subset of interactions from the original BIP model.
   (c) The Conflict Resolution Protocol Layer resolves conflicts requested by the Interaction Protocol. The Conflict Resolution Protocol implements a committee coordination algorithm and our design allows employing any such algorithm.

Figure 5.6: Resolving conflict interactions using alpha-core protocol.
We, in particular, consider three committee coordination algorithms:

i. a fully centralized algorithm,

ii. a token-based distributed algorithm,

iii. an algorithm based on reduction to distributed dining philosophers.

2. Then, we transform the 3-layer BIP model into C++ code that employs TCP sockets or MPI for communications.

The BIP composite component generated from the first phase is called Send/Receive BIP, and it is defined as the following:

**Definition 5.4.1 (Send/Receive BIP.)** We say that \( B^{SR} = \gamma^{SR}(B_{1}^{SR}, \ldots, B_{n}^{SR}) \) is a Send/Receive BIP composite component iff we can partition the set of ports in \( B^{SR} \) into three sets \( P_s, P_r, P_u \) that are respectively the set of send-ports, receive-ports, and unary interaction ports, such that:

- Each interaction \( a \in \gamma^{SR} \), is either a Send/Receive interaction \( a = (s,r_1,r_2,\ldots,r_k) \) with \( s \in P_s \) and \( r_i \in P_r \), or, a unary interaction \( a = \{p\} \) with \( p \in P_u \).
- If \( s \) is a port in \( P_s \), then there exists one and only one Send/Receive interaction \( (s,r_1,r_2,\ldots,r_k) \in \gamma^{SR} \) where all ports \( r_1,\ldots,r_k \) are receive-ports. We say that \( r_1, r_2, \ldots, r_k \) are the receive-ports associated to \( s \).
- If \( (s,r_1,\ldots,r_k) \) is a Send/Receive interaction in \( \gamma^{SR} \) and \( s \) is enabled at some global state of \( B^{SR} \), then all its associated receive-ports \( r_1,\ldots,r_k \) are also enabled at that state.
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Notice that the second condition requires that only one component can receive a "message" sent by another component. The last condition ensures that every Send/Receive interaction can take place as soon as the sender is enabled, i.e., the sender can send the message immediately.

5.5 The 3-Layer Architecture

We design our target BIP model based on the three tasks identified above, where we incorporate one layer for each task. We use the high-level BIP model in Figure 5.1 as a running example throughout this section to describe the concepts of our transformation. We assume that interaction $a_1$ is in conflict with only interaction $a_2$, and, interactions $a_2$, $a_3$, and $a_4$ are in pairwise conflict. Our 3-layer architecture consists of the following layers.

Atomic Components Layer. Atomic components in the high-level model are placed in this layer with the following additional ports per component. The send-port $o$ that shares the list of enabled ports in the component with the upper layer. Also, for each port $p$ in the original component, we include a receive-port $p$ through which the component is notified to execute the transition labeled by $p$ once the upper layers resolve conflicts and decide on which components can execute on what port. The bottom layer in Figure 5.8 includes components illustrated in Figure 5.1.
Interaction Protocol Layer. This layer consists of a set of components each hosting a set of interactions in the high-level model. Conflicts between interactions included in the same component are resolved by that component locally. For instance, interactions $a_1$ and $a_2$ (resp. $a_3$ and $a_4$) of Figure 5.1 are grouped into component $IP_1$ (resp. component $IP_2$) in Figure 5.8. Thus, the conflict between $a_1$ and $a_2$ (resp. $a_3$ and $a_4$) is handled locally in $IP_1$ (resp. $IP_2$). On the contrary, the conflicts between $a_2$ and either $a_3$ or $a_4$ must be resolved using an external algorithm that solves the committee coordination problem. Such an algorithm forms the top layer of our model. The Interaction Protocol also evaluates the guard of each interaction and executes the code associated with an interaction that is selected locally or by the upper layer. The interface between this layer and the component layer provides ports for receiving enabled ports from each component (i.e., port $o$) and notifying the components on permitted port for execution.

Conflict Resolution Protocol Layer. This layer accommodates an algorithm that solves the committee coordination problem. For instance, the external conflicts between interactions $a_2$ and $a_3$, and, interactions $a_2$ and $a_4$ are resolved by the central component $RP_1$ in Figure 5.8. We emphasize that the structure of components in this layer solely depends upon the augmented conflict resolution algorithm. Incorporating a centralized algorithm results in one component $CENT_1$ as illustrated in Figure 5.8. Other algorithms (as will be discussed in 5.6.3), such as ones that use a circulating token [Bag87] or dining philosophers [CM84, Bag89] result in different structures. The interface between this layer and the Interaction Protocol involves ports for receiving request to reserve an interaction (labeled $r$) and responding by either success (labeled $ok$) or failure (labeled $f$).

5.6 Transformations

In this section, we describe our technique for transforming a BIP model into a 3-layer distributed BIP model in detail. Construction of the three layers are described in Subsections 5.6.1, 5.6.2, and 5.6.3 respectively. Finally, we describe cross-layer interactions in Subsection 5.6.4.

5.6.1 Transformation of Atomic Components

We now present how we transform an atomic component $B$ from a given BIP model into a Send/Receive atomic component $B^{SR}$ that is capable of communicating with the Interaction Protocol in the 3-layer model. As mentioned in Section 5.5, $B^{SR}$ sends offers to the Interaction Protocol that are acknowledged by a response. An offer includes the set of enabled ports of $B^{SR}$ at the current state through which the component is ready to interact. Enabled ports are specified by a set of Boolean variables. These variables are
modified by a port update function. The function evaluates each variable when reaching a new state. When the upper layers select an interaction involving $B^{SR}$ for execution, $B^{SR}$ is notified by a response sent on the port chosen. We also include a participation number variable $n$ in $B^{SR}$, which counts the number of interactions $B^{SR}$ has participated in.

Since each response triggers an internal computation, following [BBBS08], we split each state $s$ into two states, namely, $s$ itself and a busy state $\perp_s$. Intuitively, reaching $\perp_s$ marks the beginning of an unobservable internal computation. We are now ready to define the transformation from $B$ into $B^{SR}$.

**Definition 5.6.1 (Transformation of atomic component.)** Let $B = (L, P, T, X, \{g_\tau\}_{\tau \in T}, \{f_\tau\}_{\tau \in T})$ be an atomic component. The corresponding Send/Receive atomic component is $B^{SR} = (L^{SR}, P^{SR}, T^{SR}, X^{SR}, \{g_\tau^{SR}\}_{\tau \in T^{SR}}, \{f_\tau^{SR}\}_{\tau \in T^{SR}})$ such that:

- $L^{SR} = L \cup L^{\perp}$, where $L^{\perp} = \{s \mid s \in L\}$.
- $X^{SR} = X \cup \{n\} \cup \{x_p\}_{p \in P}$, where $x_p$ is a new boolean variable associated to the port $p$, and $n$ is the number of interactions $B^{SR}$ has participated in.
- $P^{SR} = P \cup \{o\}$, where the set of variables $X^{SR}$ are associated to offer port $o$.
- For each transition $\tau = (s, p, t) \in T$, we include the following two transitions in $T^{SR}$: $\tau_1 = (\perp_s, o, s)$ and $\tau_2 = (s, p, \perp_t)$. The guards of the transition $\tau_1$ and $\tau_2$ are true. Moreover, the transition $\tau_2$ triggers the functions $f_\tau$ followed by the $f_t$. Where the function $f_t$ modifies $X^{SR}$ as follows: it sets $x_p$ to true if $(t \xrightarrow{p} ) \land (g_\tau = \text{true})$, it sets it to false otherwise, and increments $n$.

Figure 5.9 illustrates transformation of the component into its corresponding Send/Receive component.

### 5.6.2 The Interaction Protocol

Given a high-level BIP model $B = \gamma(B_1 \cdots B_n)$, one parameter to our transformation is a partition of interactions $\gamma_1, \ldots, \gamma_m$. Partitioning of interactions is a means for the designer...
to enforce load-balancing and improving the performance of the given model when running in a distributed fashion. It also determines whether or not a conflict between interactions can be resolved locally. We associate each class $\gamma_j$ of interactions to an Interaction Protocol component $IP_j$ that is responsible for (1) detecting enabledness by collecting offers from the Components Layer, (2) selecting a set of non-conflicting interactions (either locally or by the help of the Conflict Resolution Protocol), and (3) executing the selected interactions in $\gamma_j$ by notifying the corresponding atomic components. For instance, in Figure 5.8, we have two classes: $\gamma_1 = \{a_1, a_2\}$ (hosted by component $IP_1$) and $\gamma_2 = \{a_3, a_4\}$ (hosted by component $IP_2$).

Since components of the Interaction Protocol deal with interactions of the original model, they need to be aware of conflicts in the original model as defined in Definition 5.3.1. We distinguish two types of conflicting interactions according to a given partition:

- **External**: two interactions are externally conflicting if they conflict and they belong to different classes of the partition. External conflicts are resolved by the Conflict Resolution Protocol. For instance, in Figure 5.8, interaction $a_2$ is in external conflict with interactions $a_3$ and $a_4$.

- **Internal**: two interactions are internally conflicting if they conflict, but they belong to the same class of the partition. Internal conflicts are resolved by the Interaction Protocol within the component that hosts them. For instance, in Figure 5.8, interaction $a_1$ is in internal conflict with interaction $a_2$. If component $IP_1$ chooses interaction $a_1$ over $a_2$, no further action is required. Note, however, that if $IP_1$ chooses $a_2$, then it has to request its reservation from $CENT_1$, as it is in conflict with $a_3$ and $a_4$ externally.

For each Interaction Protocol we create the corresponding atomic component. The behavior of an Interaction Protocol component $IP_j$ handling a class $\gamma_j$ of interactions is constructed as follows. We refer to Figure 5.10 as a concrete example for construction of the atomic component of $IP_1$ in Figure 5.8.

**Definition 5.6.2 (Interaction protocol.)** Let $B = \gamma(B_1 \ldots B_n)$ be a composite component, and $\gamma_j = \{a_1 = (P_1, F_1, G_1), \ldots, a_k = (P_k, F_k, G_k)\}$ a set of interactions. The corresponding atomic component $IP_j = (L^{IP_j}, P^{IP_j}, T^{IP_j}, X^{IP_j}, \{g^{IP_j}_\tau\}_{\tau \in T^{IP_j}}, \{f^{IP_j}_\tau\}_{\tau \in T^{IP_j}})$ handling $\gamma_j$ is defined as follows:

- **Control states $L^{IP_j}$**: We include three types of places:
  - For each component $B_i$ involved in interactions of $\gamma_j$, we include waiting and received places $w_i$ and $rcv_i$, respectively. $IP_j$ waits in a waiting place until it receives an offer from the corresponding component. When an offer from component $B_i$ is received (along with the fresh values of the Boolean variables associated to the ports of the sender), $IP_j$ moves from $w_i$ to $rcv_i$. In Figure 5.10, since components $B_1 \cdots B_4$ are involved in interactions hosted by $IP_1$ (i.e., $a_1$ and $a_2$), we include waiting places $w_1 \cdots w_4$ and received places $rcv_1 \cdots rcv_4$. 

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Figure 5.10: Component $IP_1$ in Figure 5.8.
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- For each port \( p \) involved in interactions of \( \gamma_j \), we include a sending place \( \text{snd}_p \). The response to an offer with \( x_p = \text{true} \) is sent from this place to port \( p \) of the component that has made the offer. In Figure 5.10, places \( \text{snd}_{p_1} \cdots \text{snd}_{p_5} \) correspond to ports \( p_1 \cdots p_5 \) respectively, as they form interactions hosted by IP\(_1\) (i.e., \( a_1 \) and \( a_2 \)).
- For each interaction \( a \in \gamma_j \) that is in external conflict with another interaction, we include an engaged place \( e_a \) and a free place \( \text{fr}_a \). In Figure 5.10, only interaction \( a_2 \) is in external conflict, for which we add places \( e_{a_2} \) and \( \text{fr}_{a_2} \).
- **Variables** \( X_{IP_j} \). For each port \( p \) involved in interactions of \( \gamma_j \), we include a Boolean variable \( x_p \). The value of this variable is equal to the value of the same variable in the most recent offer received from the corresponding component. Also, for each component \( B_i \) involved in interactions of \( \gamma_j \), we include an integer \( n_i \) that stores participation number of \( B_i \), and its own variables.
- **Ports** \( P_{IP_j} \). The set of ports of IP\(_j\) is the following:
  - For each component \( B_i \) involved in interactions of \( \gamma_j \), we include an offer port \( o_i \). Each port \( o_i \) updates the values of variables \( n_i, x_p \) and the variable associated for each port \( p \) exported by \( B_i \). In Figure 5.10, ports \( o_1 \cdots o_4 \) represent offer ports for components \( B_1 \cdots B_4 \).
  - For each port \( p \) involved in interactions of \( \gamma_j \), we include a response port \( p \). In Figure 5.10, ports \( p_1 \cdots p_5 \) correspond to the ports that form interactions \( a_1 \) and \( a_2 \).
  - For each interaction \( a = (P,G,F) \in \gamma_j \) that is in external conflict, we include reservation ports \( r_a \), \( \text{ok}_a \), and \( f_a \). If \( P = \{p_i\}_{i \in I} \), the port \( r_a \) is associated to the variables \( \{n_i\}_{i \in I} \), where \( I \) is the set of components involved in interaction \( a \). In Figure 5.10, ports \( r_{a_2}, \text{ok}_{a_2} \), and \( f_{a_2} \) represent the external conflict of \( a_2 \) with interactions \( a_3 \) and \( a_4 \).
  - For each interaction \( a \in \gamma_j \) that is not in external conflict, we include a unary port \( a \). In Figure 5.10, we include unary port \( a_1 \), as \( a_1 \) is only in internal conflict with \( a_2 \).
- **Transitions** \( T_{IP_j} \). IP\(_j\) performs two tasks: (1) receiving offers from components in the lower layer and responding to them, and (2) requesting reservation of an interaction from the Conflict Resolution Protocol in case of an external conflict. The following set of transitions of IP\(_j\) performs these two tasks:
  - In order to receive offers from a component \( B_i \), we include transition \((w_i, o_i, \text{rcv}_i)\). If \( B_i \) participates in an interaction not handled by IP\(_j\), we also include transition \((\text{rcv}_i, o_i, \text{rcv}_i)\) to receive new offers when \( B_i \) takes part in such an interaction. Transitions labeled by \( o_1 \cdots o_4 \) in Figure 5.10 are of this type.
  - Requesting reservation of an interaction \( a = (P,G,F) \in \gamma_j \) that is in external conflict is accomplished by transition \((\{\text{rcv}_i\}_{i \in I} \cup \{\text{fr}_a\}, r_a, \{\text{rcv}_i\}_{i \in I} \cup \{e_a\})\), where \( I \) is the set of components involved in interaction \( a \). This transition is guarded by the predicate \( \bigwedge_{i \in I} x_{p_i} \land G \) which ensures enabledness of \( a \). Notice that this transition...
is enabled when the token for each participating component is in its corresponding receive place $rcv_i$ and the guard $G$ of the interaction is true. Execution of this transition results in moving the token from a free place to an engaged place. In Figure 5.10, transition $r_{a_2}$ is of this type, and is guarded by $x_{p_4} \land x_{p_5}$.

- For the case where the Conflict Resolution Protocol responds positively, we include the transition $(\{rcv_i\}_{i \in I} \cup \{e_a\}, ok_a, \{snd_{p_i}\}_{i \in I} \cup \{fr_a\})$. The execution of this transition triggers the function $F$ of the interaction $a$, and then, the token from the engaged place moves to the free place and the tokens from received move to sending places for informing the corresponding components. Transition $ok_{a_2}$ in Figure 5.10 occurs when interaction $a_2$ is successfully reserved by the Conflict Resolution Protocol.

- For the case where the Conflict Resolution Protocol responds negatively, we include the transition $(e_a, f_a, fr_a)$. Upon execution of this transition, the token moves from the engaged place to the free place. Transition $f_{a_2}$ in Figure 5.10 occurs when the Conflict Resolution Protocol fails to reserve interaction $a_2$ for component $IP_1$.

- For each interaction $a = \{p_i\}_{i \in I}$ in $\gamma_j$ that has only internal conflicts, let $A$ be the set of interactions that are in internal conflict with $a$, but are externally conflicting with other interactions. We include the transition $(\{rcv_i\}_{i \in I} \cup \{fr_a'\}_{a' \in A}, a, \{snd_{p_i}\}_{i \in I} \cup \{fr_a'\}_{a' \in A})$. This transition is guarded by the predicate $\wedge_{i \in I} x_{p_i}$ and moves the tokens from receiving to sending places. Tokens from $fr_a'$ places ensure that no internally conflicting interaction requested a reservation. The transition labeled by $a_1$ in Figure 5.10 falls in this category.

- Finally, for each component $B_i$ exporting $p$, we include the transitions $(snd_{p_i}, p, w_i)$. This transition notifies component $B_i$ to execute the transition labeled by port $p$. These are transitions labeled by $p_1 \cdots p_5$ in Figure 5.10.

### 5.6.3 The Conflict Resolution Protocol

As discussed earlier, the main task of the Conflict Resolution Protocol is to ensure that externally conflicting interactions are executed mutually exclusive. The Conflict Resolution Protocol can be implemented using any algorithm that solves the committee coordination problem. Our design of Conflict Resolution Protocol allows employing any such algorithm with minimal restrictions.

We adapt a variation of the idea of the message-count technique from [Bag89] as a minimal restriction to ensure that our design makes progress (see Lemma 2) and it does not interfere with exclusion algorithms. This technique is based on counting the number of times that a component interacts. Each component keeps a counter $n$ which indicates the current number of participations of the component in interactions. The Conflict Resolution Protocol ensures that each participation number is used only once. That is, each component takes part in only one interaction per transition. To this end, in the Conflict Resolution Protocol, for each component $B_i$, we keep a variable $N_i$ which stores the latest
number of participations of \(B_i\). Whenever a reserve message \(r_a\) for interaction \(a = \{p_i\}_{i \in I}\) is received by the Conflict Resolution Protocol, the message provides a set of participation numbers \(\{n_a^i\}_{i \in I}\) for all components involved in \(a\). If for each component \(B_i\), the participation number \(n_a^i\) is greater than \(N_i\), then the Conflict Resolution Protocol acknowledges successful reservation through port \(ok_a\) and the participation numbers in the Conflict Resolution Protocol are set to values sent by the Interaction Protocol. On the contrary, if there exists a component whose participation number is less than or equal to what Conflict Resolution Protocol has recorded, then the corresponding component has already participated for this number and the Conflict Resolution Protocol replies failure via port \(f_a\).

Now, since the structure and behavior of the Conflict Resolution Protocol components depend on the employed algorithm, we only specify an abstract set of minimal restrictions of this layer as follows:

- For each component \(B_i\), the Conflict Resolution Protocol maintains a variable \(N_i\) indicating the last participation number reserved for \(B_i\).
- For each interaction \(a = \{p_i\}_{i \in I}\) handled by the Conflict Resolution Protocol, we include three ports: \(r_a\), \(ok_a\) and \(f_a\). The receive-port \(r_a\) accepts reservation requests containing fresh values of variables \(n_a^i\). The send-ports \(ok_a\) and \(f_a\) accept or reject the latest reservation request, and \(N_i\) variables are updated in case of positive response.
- Each \(r_a\) message should be acknowledged by exactly one \(ok_a\) or \(f_a\) message.
- Each component of the Conflict Resolution Protocol should respect the message-count properties described above.

### 5.6.3.1 Centralized Implementation

Figure 5.11 shows a centralized Conflict Resolution Protocol for the model in Figure 5.8. In fact, the component in Figure 5.11 is the component \(CENT_1\) in Figure 5.8. A reservation request, for instance, \(r_{a_2}\), contains fresh variables \(n_{3}^{a_2}\) and \(n_{4}^{a_2}\) (corresponding to components \(B_3\) and \(B_4\)). The token representing interaction \(a_2\) is then moved from place
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\( \text{wait}_{a_2} \), to place \( \text{treat}_{a_2} \). From this state, the Conflict Resolution Protocol can still receive a request for reserving \( a_3 \) and \( a_4 \) since \( \text{wait}_{a_3} \) and \( \text{wait}_{a_4} \) still contain a token. This is where message-counts play their role. The guard of transition \( \text{ok}_{a_2} \) is \( (n_{a_2}^3 > N_3) \land (n_{a_2}^4 > N_4) \) where \( N_i \) is the last known used participation number for \( B_i \). Note that since execution of transitions are atomic in BIP, if transition \( \text{ok}_{a_2} \) is fired, it modifies variables \( N_i \) atomically (i.e., before any other transition can take place). We denote this implementation by \( \text{CENT} \).

### 5.6.3.2 Token Ring Implementation

Another example of a Conflict Resolution Protocol is inspired by the token-based algorithm due to Bagrodia [Bag87], where we add one reservation component per externally conflicting interaction. Figure 5.12 shows the respective components for the model presented in Figure 5.8. Exclusion is ensured using a circulating token carrying \( N_i \) values; i.e., the component that owns the token compares the value of the received \( n_i \) variables with the \( N_i \) variables from the token. If they are greater, an \( \text{ok} \) message is sent to the component that handles that interaction and the \( N_i \) values on the token are updated. Otherwise, a fail message is sent. Subsequently, the reservation component releases the token via port \( ST \), which is received by the next component via port \( RT \). Obviously, this algorithm allows a better level of distribution at the Conflict Resolution Protocol layer. We denote this implementation by \( \text{TR} \).

### 5.6.3.3 Implementation Based on Dining Philosophers

A third choice of Conflict Resolution Protocol algorithm is an adaption of the hygienic solution to the dining philosophers problem presented in [CM88, Bag89]. Its Send/Receive BIP implementation is presented in Figure 5.13. Similar to token ring, each externally conflicting interaction is handled by a separate component. If two interactions are conflicting, the two corresponding components share a fork carrying \( N_i \) variables corresponding to the atomic components causing the conflict. In order to positively respond to a reserve, a component has to fetch all forks shared with its neighbors. Then, it compares participation numbers received from the reservation request and from the forks and responds accordingly. After such a response, the forks become dirty. Finally, the component sends the forks if it is asked to do so. We denote this implementation by \( \text{DP} \).

### 5.6.4 Cross-Layer Interactions

In this subsection, we define the interactions of our 3-layer model. Following Definition 5.4.1, we construct Send/Receive interactions by specifying which one is the sender. Given a BIP model \( \gamma(B_1 \cdots B_n) \), a partition \( \gamma_1 \cdots \gamma_m \), and the obtained Send/Receive components \( B_1^{SR} \cdots B_n^{SR} \), Interaction Protocol components \( IP_1 \cdots IP_m \), and Conflict Resolution
Figure 5.12: Token-based Conflict Resolution Protocol for the BIP models in Figures 5.1 and 5.8.
Figure 5.13: Dining philosophers-based Conflict Resolution Protocol for the BIP models in Figures 5.1 and 5.8.
Protocol components $CENT_1 \cdots CENT_k$, we construct the Send/Receive interactions $\gamma^{SR}$ according to Definition 5.4.1 as follows:

- For each component $B_i$, $\gamma^{SR}$ contains a multicast connector formed by all ports $o_i$, where $B_i$ is the sender.
- For each Interaction Protocol component $IP_j$ and port $p$ in $IP_j$, we include a binary interaction, such that port $p$ of $IP_j$ is the sender, and, port $p$ of the corresponding component in the components layer is the receiver.
- For each interaction $a$ that is in external conflict, $\gamma^{SR}$ contains an interaction between $r_a$ ports, such that the Interaction Protocol is the sender and Conflict Resolution Protocol is the receiver. Likewise, $\gamma^{SR}$ contains interactions between $ok_a$ and $f_a$ ports.

Note that the interaction do not depend on the Conflict Resolution Protocol. The entire model obtained is denoted $B_{SR CENT}^{SR}$, $B_{TR}^{SR}$ or $B_{DP}^{SR}$ following the embedded Conflict Resolution Protocol. The interactions between the three layers of our running example are presented in Figure 5.8. The send-ports are graphically denoted by triangles and receive-ports by bullets.

5.7 Correctness

In Subsection 5.7.1, we show that our 3-layer model meets the constraints of the Send/Receive model specified in Section 5.5. In Subsection 5.7.2, we prove that a BIP model is observationally equivalent with the BIP model obtained by the transformation of Section 5.6. Finally, we prove the correctness of models embedding different implementations of Conflict Resolution Protocol in Subsection 5.7.3.

5.7.1 Compliance with Send/Receive Models

**Proposition 3** Given a BIP model $B$, the model $B^{SR}$ obtained by transformation of Section 5.6 meets the constraints of Definition 5.4.1.

**Proof** The send-ports and receive-ports are clearly determined in subsection 5.6.4 and respect the syntax presented in the two first points of definition 5.4.1. We now prove the third point, that is whenever a send-port is enabled, all its associated receive-ports are enabled.

Between the Interaction Protocol and Conflict Resolution Protocol layers, for reserve, ok and fail interactions related to $a \in \gamma$ it is sufficient to consider places $fr_a$ and $e_a$ in the Interaction Protocol layer, $wait_a$ and $treat_a$ in the Conflict Resolution Protocol layer. Initially the configuration is $(fr_a, wait_a)$ from which only the send-port $r_a$ in Interaction Protocol might be enabled, and the receive-port $r_a$ is enabled. If the $r_a$ interaction takes place, we reach the configuration $(e_a, treat_a)$, in which only send-ports $ok_a$ and $f_a$ in Conflict Resolution Protocol might be enabled, and the associated receive-ports in Interaction Protocol...
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Protocol are enabled. Then if either ok or fail interaction takes place we switch back to the initial configuration.

Between components and Interaction Protocol layers, for all interactions involving component \( B_i \), it is sufficient to consider only the places \( w_i, r_i \) and \( s_p \) for each port \( p \) exported by \( B_i \) in the Interaction Protocol. Whenever one of the places \( w_i \) or \( r_i \) is enabled in each Interaction Protocol component, the property holds for the \( o_i \) interaction. In this configuration, no place \( s_p \) might be active since it would require one of the tokens from a \( w_i \) or a \( r_i \), thus no send port \( p \) is enabled.

If there is an Interaction Protocol component such that the token associated to \( B_i \) is an a place \( s_p \), it comes either from an a or an ok labeled-transition. In the first case, no other interaction involving \( B_i \) can take place, otherwise it would be externally conflicting with a. In the second case according to the Conflict Resolution Protocol, the ok was given for the current participation number in the component \( B_i \) and no other interaction using this number will be granted. Thus in all cases, there is only one active place \( s_p \) with \( p \) exported by \( B_i \). The response can then take place and let the components continue their execution.

\[ \blacksquare \]

5.7.2 Observational Equivalence between Original and Transformed BIP Models

In this subsection, our goal is to show that \( B \) and \( B^{SR} \) are observationally equivalent. Let us first recall the definition of observational equivalence of two transition systems \( A = (Q_A, P \cup \{\beta\}, \rightarrow_A) \) and \( B = (Q_B, P \cup \{\beta\}, \rightarrow_B) \). It is based on the usual definition of weak bisimilarity [Mil95], where \( \beta \)-transitions are considered unobservable.

Notice that, a state of an atomic component is defined as a pair \( q = (s, v) \) where \( s \in L \) is the control state, \( v : X \mapsto Data \) is a valuation of the variables \( X \) of the atomic component. For simplicity of reasoning and clarity about correctness we omit the variables defined in the original atomic components, hence, functions and guards defined in the transitions and interactions is not necessary. Moreover, a state of an atomic component \( q \) become the actual control state \( l \). In this case, an interaction can be seen as a set of ports, and a composite component \( B = \gamma(B_1, \ldots, B_n) \) is a transition system \((Q,\gamma,\rightarrow)\), where \( Q = \bigotimes_{i=1}^{n} Q_i \) (\( Q_i = B_i.L \)) and \( \rightarrow \) is the least set of transitions satisfying the rule:

\[
\frac{a = \{p_i\}_{i \in I} \in \gamma \quad \forall i \in I. \quad q_i \xrightarrow{p_i} q_i'}{\forall i \notin I. \quad q_i = q_i'}
\]

\[
(q_1, \ldots, q_n) \xrightarrow{a} (q_1', \ldots, q_n')
\]

Definition 5.7.1 (Weak simulation.) A weak simulation over \( A \) and \( B \), denoted \( A \subset B \), is a relation \( R \subseteq Q_A \times Q_B \), such that we have \( \forall (q,r) \in R \) : \( a \in P : \quad q \xrightarrow{a} A q' \implies \exists r' : (q', r') \in R \wedge r \xrightarrow{\beta} B r' \) and \( \forall (q,r) \in R : \quad q \xrightarrow{\beta} A q' \implies \exists r' : (q', r') \in R \wedge r \xrightarrow{\beta} B r' \)
A weak bisimulation over $A$ and $B$ is a relation $R$ such that $R$ and $R^{-1}$ are both weak simulations. We say that $A$ and $B$ are observationally equivalent and we write $A \sim B$ if for each state of $A$ there is a weakly bisimilar state of $B$ and conversely.

We consider the correspondence between actions of $B$ and $B^{SR}$ as follows. For each interaction $a \in \gamma$, where $\gamma$ is the set of interactions of $B$, we associate either the binary interaction $ok_a$ or the unary interaction $a$, depending upon existence of an external conflict. All other interactions (offer, response, reserve, fail) are unobservable and denoted $\beta$.

We proceed as follows to complete the proof of observational equivalence. Amongst unobservable actions $\beta$, we distinguish between $\beta_1$ actions, that are communication interactions between the components layer and the Interaction Protocol (namely offer and response), and $\beta_2$ actions that are communications between the Interaction Protocol and Conflict Resolution Protocol (namely reserve and fail). We denote $q^{SR}$ a state of $B^{SR}$ and $q$ a state of $B$. A state of $B^{SR}$ from where no $\beta_1$ action is possible is called a stable state, in the sense that any $\beta$ action from this state does not change the state of the component layer.

**Lemma 1** From any state $q^{SR}$, there exists a unique stable state $[q]^{SR}$ such that $q^{SR} \xrightarrow{\beta_1} [q]^{SR}$.

**Proof** The state $[q]^{SR}$ exists since each Send/Receive component $B^{SR}_i$ can do at most two $\beta_1$ transitions: receive a response and send an offer. Since two $\beta_1$ transitions involving two different components are independent (i.e. do not change the same variable or the same place), the ordering of $\beta_1$ action does not change the final state. Thus $[q]^{SR}$ is unique. ■

We now show a property of the participation numbers. Let $B.n$ mean ‘the variable $n$ that belongs to component $B$’.

**Lemma 2** When $B^{SR}$ is in a stable state, for each couple $(i, j)$, such that $B_i$ is involved in interactions handled by $IP_j$, we have $B_i.n_i = IP_j.n_i > CENT.N_i$.

**Proof** When in stable state, all offers have been sent, thus the participation numbers in Interaction Protocol correspond to those in components $B_i.n_i = IP_j.n_i$.

Initially, for each component $B_i$, $CENT.N_i = 0$ and $B^{SR}_i.n_i = 1$ thus the property holds. The $N_i$ variables in Conflict Resolution Protocol are updated on a $ok$ transition, using values provided by the Interaction Protocol, that is by the components. We show that after each $ok_a$ transition, the property still holds. For each component $B^{SR}_i$ participant in $a$, it holds that $B^{SR}_i.n_i = CENT.N_i$ after the offer. Then, the response transitions increments participation numbers in components such that in the next stable state $B^{SR}_i.n_i > CENT.N_i$. For components $B_{i'}$ not participating in $a$, by induction we have $B^{SR}_{i'}.n_{i'} > CENT.N_{i'}$ and only participation numbers in components can be incremented. ■
Since we need to take into account participation numbers \( n_i \), we introduce an intermediate centralized model \( B^n \). This new model is a copy of \( B \) that includes in each atomic component an additional variable \( n_i \) which is incremented whenever a transition is executed. As \( B \) and \( B^n \) have identical set of states and transitions labeled by the same ports, they are observationally equivalent. (They are even strongly bisimilar.)

**Lemma 3** \( B \sim B^n \).

**Proof** We say that two states \((q, q^n)\) of \( B \) and \( B^n \) are equivalent if they have the same control states. This defines a bisimulation.

We are now ready to state and prove our central result.

**Proposition 4** \( B^{SR} \sim B^n \).

**Proof** We define a relation \( R \) between the states \( Q^{SR} \) of \( B^{SR} \) and the states \( Q \) of \( B^n \) as follows: \( R = \{(q^{SR}, q) \mid \forall i \in I : [q_i^{SR}] = q_i \} \) where \( q_i \) denotes the state of \( B^n_i \) at state \( q \) and \( [q_i^{SR}] \) denotes the state of \( B^{SR}_i \) at state \([q]^{SR}\). The three next assertions prove that \( R \) is a weak bisimulation:

(i) If \((q^{SR}, q) \in R \) and \( q^{SR} \xrightarrow{\beta} r^{SR} \) then \( (r^{SR}, q) \in R \).

(ii) If \((q^{SR}, q) \in R \) and \( q^{SR} \xrightarrow{a} r^{SR} \) then \( \exists r \in Q : q \xrightarrow{a} r \) and \( (r^{SR}, r) \in R \).

(iii) If \((q^{SR}, q) \in R \) and \( q \xrightarrow{a} r \) then \( \exists r^{SR} \in Q^{SR} : q^{SR} \xrightarrow{\beta a} r^{SR} \) and \( (r^{SR}, r) \in R \).

(i) If \( q^{SR} \xrightarrow{\beta} r^{SR} \), either \( \beta \) is a \( \beta_1 \) action and \( [q]^{SR} = [r]^{SR} \), either \( \beta \) is a \( \beta_2 \) action which does not change the state of component layer and does not enable any send-port.

(ii) The action \( a \) in \( B^{SR} \) is either a unary interaction \( a \) or a binary interaction \( ok_a \). In both cases, \( a = \{p_i\}_{i \in I} \) has been detected to be enabled in \( IP_j \) by the tokens in received places and the guard of the \( a \) or \( r_a \) transition in Interaction Protocol, with the participation numbers \( n_i \). We show that \( a \) is also enabled at state \([q]^{SR}\):
– If $a$ has only local conflicts, no move involving $B_i$ can take place in another Interaction Protocol, and no $\beta_1$ move involving $B_i$ can take place in $IP_j$ since $a$ is enabled.
– If $a$ is externally conflicting, no move involving $B_i$ has taken place in another Interaction Protocol (otherwise $ok_a$ would not have been enabled), nor in $IP_j$ since the $fr_a$ place is empty.

At stable state $[q]^{SR}$, the lemma 2 ensures that $IP_j.n_i = B_i^{SR}.n_i$. Following the definition of $R$, we have $B_i.n_i = B_i^{SR}.n_i$ when $B^n$ is at state $q$. Thus $a$ is enabled with the same participation numbers at state $q$ and in $IP_j$ at state $q^{SR}$ and $[q]^{SR}$, which implies $q \xrightarrow{a} r$.

Since the $\beta_1$ actions needed to reach the state $[q]^{SR}$ did not interfere with action $a$, we can replay them from $r^{SR}$ to reach a state $\rho^{SR}$, as shown on figure 5.14. The state $\rho^{SR}$ is not stable because of response and offers that can take place in each component participant in $B_i$. Executing these actions brings the system in state $[r]^SR$ which is clearly equivalent to $r$, and by point (i) we have $(r^{SR}, r) \in R$.

(iii) In figure 5.14, we show the different actions and states involved in this part. From $q^{SR}$, we reach $[q]^{SR}$ by doing $\beta_1$ actions. Then we execute all possible fail interactions (that are $\beta_2$ actions), so that all $fr_a$ places are empty, to reach a state $[q]^SR$. At this state, if $a$ has only local conflicts, the interaction $a$ is enabled, else the sequence $r_a ok_a$ can be executed since lemma 2 ensures that guard of $ok_a$ is true. In both cases, the interaction corresponding to $a$ brings the system in state $r^{SR}$. From this state, the responses corresponding to each port of $a$ are enabled, and the next stable state $[r]^SR$ is equivalent to $r$, thus $(r^{SR}, r) \in R$.

\[\blacksquare\]

5.7.3 Interoperability of Conflict Resolution Protocol

As mentioned in Subsection 5.6.3, the centralized implementation $CENT$ of the Conflict Resolution Protocol can be seen as a specification. We also proposed two other implementations, respectively, token-ring $TR$ and dining philosophers $DP$. However, these implementations are not observationally equivalent to the centralized implementation. More precisely, the centralized version defines the most liberal implementation: if two reservation requests $a_1$ and $a_2$ are received, the protocol may or may not acknowledge them, in a specific order. This general behavior is not implemented neither by the token ring nor by the dining philosophers implementations. In the case of token ring, the response may depend on the order the token travels through the components. In the case of dining philosophers, the order may depend on the order the token travels through the components. In the case of dining philosophers, the order may depend on places and the current status of forks.

Nevertheless, we can prove an observational equivalence if we consider weaker versions of the above implementations. More precisely, for the token ring protocol, consider the weaker version $TR^{(w)}$ which allows to release the token or provide a fail answer regardless of the values of counters. Likewise, for the dining philosophers protocol, consider the weaker version $DP^{(w)}$, where forks can always be sent to neighbors, regardless of their status and the values of counters. Clearly, a weakened Conflict Resolution Protocol is not desirable for
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a concrete implementation since they do not enforce progress. But, they play a technical role in proving the correctness of our approach. The following proposition establishes the relation between the different implementations of the Conflict Resolution Protocol.

**Proposition 5**

(i) \(\text{CENT} \sim \text{TR}^{(w)} \sim \text{DP}^{(w)}\)

(ii) \(\text{TR} \subset \text{TR}^{(w)}\), \(\text{DP} \subset \text{DP}^{(w)}\).

Let us denote by \(B_{X}^{SR}\) the 3-layer model obtained from the initial system \(B\) and embedding algorithm \(X\) in the Conflict Resolution Protocol. Also, let us denote \(Tr(B)\) the set of all possible traces of observable actions allowed by an execution of \(B\). The following proposition states the correctness of our implementation.

**Proposition 6**

(i) \(B \sim B_{X}^{SR} \sim B_{TR}^{SR} \sim B_{DP}^{SR}\)

(ii) \(Tr(B) \supseteq Tr(B_{TR}^{SR})\) and \(Tr(B) \supseteq Tr(B_{DP}^{SR})\).

**Proof** (i) The leftmost equivalence is a consequence of lemma 3 and proposition 4. The other equivalences come from proposition 5 and the fact that observational equivalence is a congruence with respect to parallel composition. (ii) The trace inclusions follows from the simulations \(\text{TR} \subset \text{TR}^{(w)}\) respectively \(\text{DP} \subset \text{DP}^{(w)}\).

5.8 Transformation from Send/Receive BIP into C++

In this section, we describe how we generate for a Send/Receive BIP component pseudo C++ code. Notice that since the behavior of these components are formalized as Petri nets, we only present generation of C++ code for a Petri net whose transitions are labeled by send-ports, receive-ports, or unary ports (see C++ Pseudo Code 1).

Initially, each component creates a TCP socket and establishes reliable connections with all components that it needs to interact (Lines 1-2). These interactions and their corresponding physical connections are determined according to the complete Send/Receive BIP model and a configuration file. This file specifies the IP address and port number of all components for final deployment. We assign one Boolean variable to each place of the given Petri net, which shows whether or not the place contains the token. Thus, the initial state of the Petri net is determined by an initial assignment of these variables (Line 3).

After initializations, the code enters an infinite loop that executes the transitions of the Petri net as follows. For each step, the code scans the list of all possible transitions and gives priority to transitions that are labeled by a send-port (Lines 6-10) or unary ports of the given Petri net (Lines 11-15). Actual emission of data is performed by an invocation of the TCP sockets system call `send()` in Line 7. Once data transmission or an internal computation is completed, tokens are removed from input places and put to output places of the corresponding transitions (Lines 8 and 13).
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C++ Pseudo Code 1 Petri net

Input: A Petri net of a Send/Receive BIP component and a configuration file.
Output: C++ code that implements the given Send/Receive Petri net

// Initializations
1: CreateTCP$ocket();
2: EstablishConnections();
3: PrepareInitialState();
4: while true do
5:   // Handling send-ports and internal computations
6:   if there exists an enabled transition labeled by a send-port then
7:     send(...);
8:     PrepareNextState();
9:     continue;
10:   end if
11:   if there exists an enabled transition labeled by a unary port then
12:     DoInternalComputation();
13:     PrepareNextState();
14:     continue;
15:   end if
16:   // Handling receiving messages
17:   select(...);
18:   recv(...);
19:   PrepareNextState();
20: end while

Finally, if no send-port is enabled and all internal computations are completed, execution stops and waits for messages from other components (Line 17). Once one of the sockets contains a new message, the component resumes its execution and receives the message (Line 18).

It is straightforward to observe that our code avoids creating deadlocks by giving priority to send-ports and unary-ports. Moreover, sending messages before doing internal computation triggers receivers components waiting for a response and increases parallelism.

Note that we also provide the generation of C++ code by using MPI for communications. Generally speaking, the same principle is applied as above, however, we use the communication primitives offered by MPI (e.g., \texttt{MPI\_Send()}, \texttt{MPI\_ISend()}, \texttt{MPI\_Recv()}, etc.) instead of TCP sockets.
5.9 Component Composition

This technique is applied to the intermediate 3-layer Send/Receive model developed in Section 5.6. It consists in composing some components from the 3-layer Send/Receive model. For instance, if we consider an Interaction Protocol handling one interaction, in this case, it is possible to merge it into one of its corresponding components without losing any parallelism. This method will be also useful in the case when generating MPI code, since an important overhead will appear due to context switching between processes. For example, if we consider a platform consisting of two cores, it is preferable to generate exactly two processes. Thus, we need to merge components from the 3-layer Send/Receive model to obtain the less number of components and without killing parallelism. Figure 5.15 illustrates an example of merging components. As input we take a set of partitioning of components and we obtain as output a new equivalent BIP model by merging with respect to the partition given as input. This can be done by applying the third transformation (Component Composition) presented in Chapter 4.

5.10 Experimental Validation

In this section, we present the results of our experiments. Recall that, our implementation automatically generates C++ code from the 3-layer BIP model developed in Sections 5.5 and 5.6, where Send/Receive interactions are implemented by TCP sockets or MPI primitives. We have implemented and integrated the transformations in the BIP toolset. The tool takes a composite BIP model in the global state semantics and a network configuration file as input and generates the corresponding C++ executable for each Send/Receive component for all layers of the intermediate BIP model (e.g, Atomic Component, Interaction Protocol, Conflict Resolution Protocol). Each executable can be run independently on a different machine or a processor core.

We denote each experiment scenario by \((i, X)\), where \(i\) is the number of interaction
partitions and $X$ is the choice among the three Conflict Resolution Protocols described in Subsection 5.6.3 (i.e., CENT, TR, or DP). For the case where partitioning of interactions results in having no external conflicts, hence, requiring no reservation component, we use the symbol ‘$-$’ to denote an empty Conflict Resolution Protocol. All experiments in this section are conducted on quad-Xeon 2.6 GHz machines with 6GB RAM running under Debian Linux. The machines are connected via a 100Mbps Ethernet network.

For three non trivial examples, Diffusing Computation, Utopar Transportation System and Bitonic Sorting, described bellow we show that different conflict resolution algorithms and partitioning may result in significantly different performance depending on the initial BIP model and the deploying of the distributed implementation over target platforms.

5.10.1 Diffusing Computation

We model a simplified version of Dijkstra-Scholten termination detection algorithm for diffusing computations [DS80] in BIP. Diffusing computation is the task of propagating a message across a distributed system; i.e., a wave that starts from an initial node and diffuses to all processes in a distributed system. Diffusing computation has numerous applications such as traditional distributed deadlock detection and reprogramming of modern sensor networks. One challenge in diffusing computation is to detect its termination. In our version, we consider a torus (wrapped around grid) topology for a set of distributed processes, where a spanning tree throughout the distributed system already exists; each process has a unique parent and the root process is its own parent. Termination detection is achieved in two phases: (1) the root of the spanning tree possesses a message and initiates a propagation wave, so that each process sends the message to its children, and (2) once the first wave of messages reaches the leaves of the tree, a completion wave starts, where a parent is complete once all its children are complete. In this setting, when the root is complete, termination is detected.

Our BIP model has $n \times m$ atomic components (see Figure 5.16 for a partial model). Each component participates in two types of interactions: (1) four binary rendezvous interactions (e.g., $a_0 \cdots a_3$) to propagate the message to its children (as in a torus topology, each node has four neighbors, hence, potentially four children), and (2) one 5-ary rendezvous interaction (e.g., $a$) for the completion wave, as each parent has to wait for all its children to complete.

Our first set of experiments is on a $4 \times 6$ torus. We apply different partitioning scenarios as illustrated in Figure 5.17. Figure 5.18 shows the time needed for 100 rounds of detecting termination of diffusing communication for each scenario. In the first two scenarios, the interactions are partitioned, so that all conflicts are internal and, hence, resolved locally by the Interaction Protocol. In case of (2, $-$), all interactions of the propagation wave are grouped into one component of the Interaction Protocol and all interactions related to the completion wave are grouped into the second component. Such grouping does not allow parallel execution of interactions. This is the main reason that the performance of (1, $-$)
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Figure 5.16: Partial BIP model for diffusing computations.

Figure 5.17: Different scenarios for diffusing computations.
and \((2, -)\) are the worst in Figure 5.18.

Next, we group all interactions involved in components \(1 \cdots 12\) into one component and the rest in a second component of the Interaction Protocol. This constitutes experiments \((2, CENT), (2, TR),\) and \((2, DP)\). Such partitioning allows more parallelism during propagation and completion waves, as an interaction in the first partition can be executed in parallel with an interaction in the second partition. This is why the performance of \((2, CENT/TR/DP)\) is better than \((1, -)\) and \((2, -)\). Now, since almost all propagation interactions conflict with each other and so do all completion interactions, in case of the dining philosophers algorithm, the conflict graph is not dense. Hence, a small number of decisions can be made in a local neighborhood of philosophers. It follows that the performance of \((2, TR)\) is quite competitive with \((2, DP)\). It can also be seen that \((2, CENT)\) performs as good as \((2, TR)\) and \((2, DP)\). This is due to the fact that there exist only two partitions, which results in a low number of reservation requests.

Figure 5.18 also shows the same type of experiments with 4 and 24 partitions. Similar to the case of two partitions, the performance of \(TR\) and \(CENT\) for 4 and 24 partitions are almost the same. However, \(CENT\) and \(TR\) outperform \(DP\). This is due to the fact that in case of \(DP\), each philosopher needs to acquire 4 forks, which requires considerable communication. On the other hand, \(TR\) does not require as much communication, as the only task it has to do is releasing and acquiring the token. Moreover, the level of parallelism in \(DP\) in case of a \(6 \times 4\) torus is not high enough to overcome the communication volume.

In the next experiment, following the lesson learned from the tradeoff between communication volume and parallelism, we design a scenario where we exploit the fact that each reservation component in \(DP\) resolves conflicts through communicating with its neighboring components. This is not the case in \(TR\). Thus, we consider a \(20 \times 20\) torus. As can be seen in Figure 5.19, the performance of \(DP\) is significantly better than \(TR\). This is solely because when we have a large number of components, in \(TR\). The token has to travel a long way in order to allow parallel execution of interactions. To the contrary, in \(DP\), the Conflict Resolution Protocol components act in their local neighborhood and although more communication is needed, it allows better concurrency, hence, higher simultaneous execution of interactions. We expect that by increasing the size of the torus, \(DP\) outperforms \(CENT\) as well.

We conclude from this example by stating the main lesson learned from our experiments:

Different partitioning schemes and choice of committee coordination algorithm for distributed conflict resolution suit different topologies and settings although they serve a common purpose. Designers of distributed applications should have access to a library of algorithms and choose the best according to parameters of the application.

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1. Execution of each interaction involves 10ms suspension of the corresponding component in the Interaction Protocol to perform and I/O command.
5.10.2 Utopar Transportation System

The second example is Utopar, an industrial case study of the European Integrated project SPEEDS\(^2\). Utopar is an automated transportation system managing various requests for transportation. The system consists of a set of autonomous vehicles, called U-cars, a centralized automatic control (Central-Station) and calling units (see Figure 5.20).

We modeled a simplified version of the Utopar Transportation System in BIP. The overall system architecture is depicted in Figure 5.21. It is a composition of an arbitrary (but fixed) number of components of three different types: U-Cars, Calling-Units and Central-Station. The Utopar system interacts with external users, i.e., the passengers. For sake of completeness, users are also represented in the Figure 1 as components, however, their behavior is not explicitly modeled.

The overall behavior of the system is obtained by composing the behavior of the inner components according to the following set of interactions:

- request: handling car requests (made by passengers) at Calling-Units;
- destination: handling destination requests (made by passengers) seating within U-Cars;

\(^2\) http://www.speeds.eu.com/
Figure 5.19: Performance of termination detection in diffusing computation in different scenarios (Torus 20×20).
Figure 5.20: Utopar transportation system.
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Figure 5.21: High-level BIP model for Utopar system.

- enter: handling the step on (resp. off) for passengers into U-Cars;
- departure: handling departure commands issued by Central-Station towards the U-Cars;
- posChanged, arrival: information provided by moving U-Cars towards the Central-Station;
- open, close: handling the opening/closing of the U-Cars doors, while parked at Calling-Units.

Our first set of experiments consists of $25 = 5 \times 5$ calling units and 4 cars. For each calling unit we group all the interactions connected on it in the same Interaction Protocol. Moreover, for each car we group all the interactions connecting the car with the central station in the same Interaction Protocol. Thus, we obtain 29 Interaction Protocol components. Using this partitioning we generate the corresponding 3-layer Send/Receive model for the three Conflict Resolution Protocols.

We simulate the target platform as follows. We consider that there exists a machine on each calling unit. Moreover, each machine is connected to their four neighbours and the communication between two neighbours machines takes $1ms$.

We generate the corresponding C++ executable for each Send/Receive component for all layers, and we embedded each C++ executable as follows. We embedded the code corresponding to the calling unit with its Interaction Protocol into the machine located on the
calling unit. Furthermore, we embedded the code corresponding to the central station into the central machine that is located in the center of the calling units. Regarding the Conflict Resolution Protocol, in the case of CENT the best choice is to embed its corresponding code into the central machine. Concerning TR and DP algorithm we embedded the code of each component of this layer in its corresponding Interaction Protocol.

Figure 5.22 shows the time needed for responding 10 requests by each calling unit. It is clear that the performance of (29, DP) is better than (29, TR) and (29, CENT). This is due to the overhead of communications for the case of TR and CENT. More precisely, regarding CENT the overhead is due to the communication between the components of Interaction Protocol layer and Conflict Resolution Protocol layer, since the centralized Conflict Resolution Protocol is placed in the central machine. Regarding TR the overhead is due to the communications between Conflict Resolution Protocol which depends on the number of components in this layer. To the contrary, in DP, the Conflict Resolution Protocol components act in their local neighborhood although more communication is needed.

Figure 5.23 also shows the same type of experiments by taking 4 cars and 49 = 7 × 7 calling units. The performance becomes worse for TR since the token has to travel a long way through the components of the Conflict Resolution Protocol layer.
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5.10.3 Bitonic Sorting

In the two previous examples, the system itself is geographically distributed, hence the generation of a distributed implementation is necessary. However, in other case the generation of a distributed implementation is necessary for deriving more computational power by using multiple processors (e.g., sorting algorithm, genetic algorithm).

The aim of this example is to show that our methodology assists developers of parallel and multi-core applications to start developing from high-level BIP models and not get involved in low-level synchronization details.

Bitonic sorting [Bat68] is one of the fastest sorting algorithms suitable for distributed implementation in hardware or in parallel processor arrays. A sequence is called bitonic if it is initially nondecreasing then it is nonincreasing. The first step of the algorithm consists in constructing a bitonic sequence. Then, by applying a logarithmic number of bitonic merges, the bitonic sequence is transformed into totally ordered sequence. We provide an implementation of the bitonic sorting algorithm in BIP using four atomic components, each one handling one part of the array. These components are connected as shown in Figure 5.24. The six interactions are non conflicting. Moreover, interactions \( I_1 \), \( I_2 \) and \( I_3 \) cannot run in parallel. The same holds for interactions \( I_4 \), \( I_5 \), \( I_6 \). Thus, to obtain maximal parallelism between interactions it sufficient to create only two components for the Interaction Protocol layer. Where, the first one handles the interactions \( I_1 \), \( I_2 \) and \( I_3 \) and

![Figure 5.23: Performance of responding 10 request per calling unit (49 = 7 × 7 calling units, and 4 cars).](image-url)
the second one handles the interactions $I_4$, $I_5$ and $I_6$. Furthermore, since all interactions are non conflicting, there is no need for the Conflict Resolution Protocol layer (detected automatically by the tool). In this example each component sends only three messages, each one containing its own array.

We run experiments for three configurations: $1c$, $4c$, $4c'$. For $1c$, we use one single-core machine, where the four atomic components along with the two IP components. For $4c$, we use two dual-core machines and place each atomic component on a different core. We also distribute the IP components over two cores, such as to reduce the network communication overhead. For $4c'$, we use the same distribution for components and IP. The results are
Table 5.1: Performance of Bitonic Sorting Algorithm.

<table>
<thead>
<tr>
<th>$k$</th>
<th>C++/Socket (generated)</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$1c$</td>
<td>$4c$</td>
<td>$4c'$</td>
</tr>
<tr>
<td>20</td>
<td>96</td>
<td>23</td>
<td>24</td>
</tr>
<tr>
<td>40</td>
<td>375</td>
<td>96</td>
<td>100</td>
</tr>
<tr>
<td>80</td>
<td>1504</td>
<td>390</td>
<td>397</td>
</tr>
<tr>
<td>160</td>
<td>6024</td>
<td>1539</td>
<td>1583</td>
</tr>
</tbody>
</table>

Table 5.2: The impact of component composition on Send/Receive models.

<table>
<thead>
<tr>
<th>$k$</th>
<th>S/R BIP</th>
<th>Merged S/R BIP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Socket</td>
<td>MPI</td>
</tr>
<tr>
<td>20</td>
<td>23</td>
<td>63</td>
</tr>
<tr>
<td>40</td>
<td>96</td>
<td>271</td>
</tr>
<tr>
<td>80</td>
<td>390</td>
<td>964</td>
</tr>
<tr>
<td>160</td>
<td>1539</td>
<td>4158</td>
</tr>
</tbody>
</table>

reported in Table 5.1 for arrays of size $k \times 10^4$ elements, and $k = 20, 40, 80, 160$. As can be seen in Table 5.1

Table 5.1 shows the performance of the automatically C++ code generated using TCP sockets. It is clear that if we consider larger arrays, then increasing the number of cores leads to a proportional performance gain. For example, the execution time for sorting an array of size $160 \times 10^4$, for the configuration $4c$ is 1539 seconds, and for the configuration $1c$ is $6024 \approx 4 \times 1539$ seconds.

The performance of case $4c$ (2 computers with two cores each) configuration is shown in Table 5.2. Observe that the performance of the C++/Socket code is approximately identical in both cases. This is because socket operations are interrupt-driven. Thus, if a component is waiting for a message, it does not consume CPU time. On the other hand, MPI uses active waiting, which results in CPU time consumption when the IP is waiting. Since we have four cores for six processes, the MPI code generated from the original Send/Receive model is much slower than the socket code. Nevertheless, as it appears in the table, reducing the number of components to one per core by merging (see Figure 5.26) allows the MPI code to reach the same speed as in the C++/socket implementation.
5.11 Summary

In this chapter we proposed a methodology for producing automatically efficient and correct-by-construction distributed implementations by starting from a high-level model of the application software in BIP. The methodology transforms arbitrary BIP models into Send/Receive BIP models, directly implementable on distributed execution platforms. The transformation consists of:

1. breaking atomicity of actions in atomic components by replacing strong synchronizations with asynchronous Send/Receive interactions;
2. inserting several distributed controllers that coordinate execution of interactions according to a user-defined partition;
3. augmenting the model with a distributed algorithm for handling conflicts between controllers.

We shown that the obtained Send/Receive BIP models are observationally equivalent to the initial models. Hence, all the functional properties are preserved by construction in the implementation. Moreover, Send/Receive BIP models can be used to automatically derive distributed implementations. Currently, it is possible to generate stand-alone C++ implementations using either TCP sockets for conventional communication, or MPI implementation, for deployment on multi-core platforms. This method is fully implemented. We report concrete results obtained under different scenarios (i.e., partitioning of the interactions and choice of algorithm for distributed conflict resolution).

In the next chapter, we present the tool which implements the transformations presented in this thesis. Moreover, we give an overview of the integration of our tool in the design methodology for BIP for automatically deriving efficient centralized and distributed from high-level BIP models.
Chapter 5. Transformation for Generating Distributed Implementations
CHAPTER 6

Tool-Chain

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6.1 BIP into Centralized Implementations Tool-Chain - BIP2BIP

The transformations from BIP into centralized implementations in Chapter 4 have been implemented in the BIP2BIP tool, which is currently integrated in the BIP toolset [BIP] as shown in Figure 6.1.

The BIP2BIP tool is written in Java. It allows transformation of parsed models. It contains the following modules implementing the presented transformations.

- Component flattening: this module transforms a composite component into an equivalent one consisting only of atomic components of the initial model and a set of connectors.
- Connector flattening: this module transforms an hierarchically structured connector into an equivalent flat one. By successive applications of this module, we obtain a new model with flat connectors.
- Component composition: this module transforms a set of atomic components and a set of flat connectors into an equivalent atomic component. By successive applications of this module, we obtain a new model consisting of a single atomic component.
Chapter 6. Tool-Chain

Figure 6.1: BIP2BIP toolset: General architecture
Chapter 6. Tool-Chain

By exhaustive application of these transformations, an atomic component can be obtained. From the latter, the code-generator can generate standalone C++ code, which can be run directly without the Engine. In particular, all the remaining non-determinism in the final atomic component is eliminated at code generation by applying an implicit priority between transitions.

It should be noted that the transformations also can be applied independently, to obtain models that respond to a particular user needs. For example, one may decide to eliminate only partially the hierarchy of components, or to compose only some components.

The performance of BIP2BIP is quite satisfactory. For example, when applied to an artificially complex BIP model, consisting of 256 atomic components, composed by using 509 connectors with 7 levels of hierarchy, it takes less than 15 seconds to generate the corresponding C++ program.

6.2 BIP into Distributed Implementations Tool-Chain - BIP2Dist

The transformations from BIP into distributed implementations in Chapter 5 have been implemented in the BIP2Dist tool (see Figure 6.2), which is also integrated in the BIP toolset.

The BIP2Dist tool is written in Java. It allows the generation of distributed implementation starting from a high-level BIP model. In the follows, we will describe the design process for that generation using BIP2Dist tool. We illustrate the process on the example shown in Figure 6.3.

1. Starting from a high-level BIP model, we flatten the hierarchy of connectors and components;
2. From the flatten model that only consists of atomic components and flat connectors, we generate 3-layer Send/Receive BIP model by choosing a partition of interactions and a Conflict Resolution Protocol;
3. From the 3-layer Send/Receive BIP model, a designer may merge some components by choosing a partition of component to merge;
4. Finally, from the obtained model we generate C++ code using TCP socket or MPI for communication according to designer demand. Moreover, this step takes as input a mapping of the components over a distributed target platform.

6.3 Summary

We have provided an overview on the implementation of the BIP2BIP and BIP2Dist tool for generating centralized and distributed implementations from BIP models. In the next chapter, we conclude the thesis with an overview of the work and its future perspectives.
Chapter 6. Tool-Chain

Figure 6.2: BIP2Dist toolset: General architecture
Figure 6.3: Design process to automatically generate distributed implementations
Chapter 6. Tool-Chain
Conclusions and Perspectives

In this chapter, we conclude the thesis describing the main objectives of the work, the goals we have achieved, the future work directions and its perspectives.

7.1 Conclusions

The thesis shows that it is possible to reconcile component-based incremental design and efficient code generation by applying a paradigm based on the combined use of:

1. A high-level modelling language, BIP, based on well-defined operational semantics and supporting powerful mechanisms for expressing structured coordination between components. The design methodology using BIP language involves the following steps:
   (a) The system (software) to be designed is decomposed into components. The decomposition can be represented as a tree which shows how the system can be obtained as the incremental composition of components. Its root is the system and its leaves correspond to atomic components;
   (b) Description of the behavior of the atomic components;
   (c) Description of composite components as the composition of atomic components by using only connectors and priorities.
Chapter 7. Conclusions and Perspectives

This is possible because BIP is expressive enough for describing any kind of coordination by using only architectural constraints [BS08b]. Along steps b) and c) it is possible by using the D-Finder tool, to generate and/or check invariants of the components and validate their properties. The methodology provides sufficient conditions for preserving the already established properties of the sub-systems along the construction.

BIP has already successfully been used for the componentization of non trivial systems such as the controller of the DALA robot [BGL+08]. This allowed building component-based models for which enhanced analysis and verification is possible by using tools such as D-Finder [BBNS09, BBSN08] for compositional verification.

2. Semantics-preserving source-to-source transformations that allows to generate automatically efficient centralized or distributed implementations. We have developed two implementation methods for BIP, sequential and distributed, which target respectively single-processor or multi-processor execution platforms.

– centralized implementations: we defined a set of source-to-source transformations that progressively transform architectural constraints between components into internal computation of product components. These transformations include flattening of hierarchical compositions and hierarchical connectors and also static composition of atomic behavior. The aim of these transformation is to transform a composite component into a single atomic component. From the latter an efficient C++ code can be generated. We show that these transformations are semantic preserving and moreover, when used in the implementation flow, they reduce overheads in execution time by reducing modularity introduced by the designer when it is not necessary at implementation level.

– distributed implementations: we defined a set of source-to-source transformations that generate automatically distributed implementation from the high-level models specified in BIP. Although BIP provides a rich set of interactions, we only considered rendezvous interactions, as they play an important role in systems whose constituents need to synchronize on some event in order to start some computation. In a distributed setting, implementation of a multi-party rendezvous results in solving the committee coordination problem [CM88], where a set of professors are organized in a set of committees and two committees can meet concurrently only if they have no professor in common; i.e., they are not conflicting. Conflict resolution is the main obstacle in distributed implementation of multi-party rendezvous interactions.

Our transformation consists of two steps. First, it takes as input a BIP model in terms of a set of components glued by rendezvous interactions and generates another BIP model which contains component glued by Send/Receive interactions in the following three layers: (1) the Atomic Components layer consists of a transformation of behavioral components in the original model, (2) the Interaction Protocol
layer detects enabledness of interactions of the original model and executes them after resolving conflicts either locally or by the help of the third layer, and (3) the Conflict Resolution Protocol layer resolves conflicts unresolved by the interaction protocol. The Conflict Resolution Protocol implements a committee coordination algorithm and our design allows employing any such algorithm. The second step of our transformation takes the intermediate three-layer BIP model as input and generates C++ executables using either TCP sockets or MPI for communications. We conducted several experiments using different algorithms in the Conflict Resolution Protocol. As predicated, our experiments show that each algorithm is suitable for a different topology, size of the distributed system, communication load, and of course the structure of the initial high-level model. Thus, the important lesson learned from our experiments is that there is no silver bullet to automate code generation for distributed applications and designers must have access to a formal framework and a library of algorithms to be able to develop correct and yet efficient distributed applications.

7.2 Perspectives

For future work, we are considering several research directions.
- According centralized and distributed implementations we plan to take into account priorities. Concerning centralized implementations, priority rules can be compiled in the form of restrictions of the guards of components. On the other hand, according to distributed implementations, we agree that priorities complicate the problem, as unlike conflict-resolution, priorities must be applied globally which requires approaches such computing a global snapshot. The rest of the following future works are according to distributed implementations;
- Another direction is introducing the notion of time in distributed semantics of BIP. Providing timing guarantees in a distributed setting has always been a challenge and BIP is not an exception;
- An important extension to explore is to allow the Conflict Resolution Protocol to incorporate different algorithms for conflict resolution simultaneously. This is because each set of conflicting interactions within the same system may react differently to different algorithms and, hence, it is desirable to handle each set with the algorithm that performs the best. In this context, we are also planning to develop and implement other algorithms, such as solutions to distributed graph matching and distributed independent set. Moreover, we need to have a better understanding of tradoffs between parallelism within the components layer and the Interaction Protocol, load balancing, and network traffic;
- Another important line of research is to measure the overhead of our transformation technique as compared to hand-written code. To this end, we plan to design
customized techniques and conduct experiments in large sensor networks where communication and computation tradeoffs play an important role in efficiency and energy consumption and the network cannot afford incorporating solutions that add significant overhead. Another potential avenue for our work is large peer-to-peer networks; – Finally, given the recent advances in the multi-core technology, we plan to customize our transformation for multi-core platforms as well. In these platforms, network communication can be replaced by simple inter-process communication and one can investigate whether it is possible to devise more effective techniques to achieve correct-by-construction concurrency and process synchronization.


[CKL+F02] Jordi Cortadella, Alex Kondratyev, Luciano Lavagno, Claudio Passerone, and Yosinori Watanabe, Quasi-static scheduling of independent tasks for reactive
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Thesis 130 Mohamad Jaber
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