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INSTITUT NATIONAL DE RECHERCHE EN INFORMATIQUE ET EN AUTOMATIQUE

*Compositional Verification of Asynchronous  
Concurrent Systems using CADP (extended version)*

Hubert Garavel — Frédéric Lang — Radu Mateescu

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# Compositional Verification of Asynchronous Concurrent Systems using CADP (extended version)

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**Abstract:** During the last decades, concurrency theory successfully developed salient concepts to formally model and soundly reason about distributed and parallel systems. In practice, however, most attempts at analyzing large systems face severe complexity issues, especially state explosion, which prevents to exhaustively enumerate reachable state spaces. Compositionality is the most promising approach to fight state explosion. In this report, we focus on finite-state verification techniques for asynchronous message-passing systems, highlighting the existence of multiple, diverse compositional techniques such as: compositional model generation, semi-composition and projection, automatic generation of projection interfaces, formula-dependent model generation, and partial model checking. These approaches have been implemented in the framework of the CADP (*Construction and Analysis of Distributed Processes*) software toolbox and applied to large-scale, industrial systems. A key point is the ability to combine several compositional techniques, as no single technique is sufficient to address all kinds of systems.

**Key-words:** Bisimulation, Concurrency theory, Formal method, Labeled Transition System, Model checking, Model generation, Model minimization, Mu-calculus, Network of automata, Partial model checking, Process algebra, Projection interface, Semi-composition, Temporal logic, Verification

# Vérification compositionnelle de systèmes concurrents asynchrones avec CADP

**Résumé :** Au cours des dernières décennies, la théorie de la concurrence a donné naissance à des concepts importants pour modéliser et raisonner formellement sur les systèmes parallèles et distribués. En pratique cependant, la plupart des tentatives d'analyser de gros systèmes se confronte à de sérieux problèmes de complexité, en particulier l'explosion d'états, qui empêche d'énumérer de manière exhaustive les espaces d'états atteignables. La compositionnalité est l'approche la plus prometteuse pour lutter contre l'explosion d'états. Dans ce rapport, nous nous concentrons sur les techniques de vérification d'états finis pour les systèmes asynchrones avec passage de messages, mettant en exergue l'existence de multiples et diverses techniques compositionnelles telles que : génération compositionnelle de modèle, semi-composition et projection, génération automatique d'interfaces de projection, génération de modèle dépendant de la formule, et model checking partiel. Ces approches ont été implémentées dans le contexte de la boîte à outils logiciels CADP (*Construction and Analysis of Distributed Processes*) et appliquées à des systèmes industriels de grande échelle. Un élément clé est la possibilité de combiner plusieurs techniques compositionnelles, une technique unique étant insuffisante pour traiter tous les types de systèmes.

**Mots-clés :** Algèbre de processus, Bisimulation, Génération de modèle, Interface de projection, Logique temporelle, Méthode formelle, Minimisation de modèle, Model checking, Model checking partiel, Mu-calcul, Réseau d'automates, Semi-composition, Système de transitions étiquetées, Théorie de la concurrence, Vérification

# 1 Introduction

Concurrency theory, which can be traced back to the early seventies, has produced over the years a large corpus of fundamental results for the modeling and analysis of distributed and parallel systems. Among the major theoretical successes, one can mention: *process calculi* and *process algebras* [77, 1, 107, 26, 18, 108, 109, 19], which explore the consequences of introducing parallel composition operators as first-class citizens in specification languages; *structural operational semantics* [118, 76, 1, 120, 119], which cleanly formalize the semantics of process calculi in terms of lower-level models, such as *labelled transition systems* [116]; *bisimulations* and related preorder/equivalence relations [107, 116, 108, 136, 135, 137], which enable a precise and meaningful comparison of concurrent systems; and *temporal logics* and  $\mu$ -*calculus* [121, 82, 97], which provide declarative means to specify system properties.

Beyond these high-level concepts and formalisms, concurrency theory also led to concrete algorithmic advances, such as finite- and infinite-state techniques for analyzing state spaces, efficient *equivalence checking* algorithms for computing bisimulations [115, 73, 20], and powerful *model checking* algorithms for temporal logics and  $\mu$ -calculus [34, 12].

Such advances have been implemented in many academic and industrial tools, which have been applied to a full range of problems, often with success. However, in most cases, the automated analysis of realistic, large-scale systems faces difficult complexity issues, especially the *state-explosion* problem that occurs when the set of reachable states of a system is too large to be analyzed, either in extension or in comprehension. Although careful specification decisions may help contain this complexity to a certain extent [113, Chapter 6][71], in general algorithmic approaches are needed to fight against such complexity while maintaining the goal of exhaustive verification; examples of such approaches are based on partial order [114, 81, 66, 133, 67, 68, 117, 54] or symmetry [78, 36, 35, 2] reductions. Probably, the most general and most promising approaches are *compositional verification* techniques, which rely on the “divide-and-conquer” paradigm to breakdown the complexity of the systems under study.

The present report focuses on compositionality in the setting of finite-state verification techniques for *asynchronous*, *action-based* systems. Concretely, this encompasses systems described using process calculi or networks of communicating finite-state automata, with underlying semantic models such as LTSs (labelled transition systems) as regards functional verification, as well as (extensions of) Markov chains as regards quantitative performance evaluation. In this context, the term *asynchronous* refers to classes of systems, the concurrent components of which may either synchronize or evolve independently, contrary to the so-called *synchronous* systems [74], in which concurrent components evolve together at periodic instants, e.g., cadenced by some clock. The term *action-based* refers to system models in which the only observable events are attached to transitions; such events are called *actions* (or *transition labels*) and may correspond, in practice, to inputs received and outputs sent by the system, as well as internal transitions performed by the system. Action-based models differ from so-called *state-based* models, in which only the internal contents of system states (namely, variables and other information stored in memory) can be observed. Although action-based and state-based can be seen as dual concepts from a mathematical point of view, they are practically different for at least two reasons: (1) action-based approaches correspond to “black box” observation while state-based approaches correspond to “white box” observation, and (2) compositional verification techniques for both kinds of approaches are quite different algorithmically.

As a matter of fact, state-based approaches are currently predominant in the scientific publications and university textbooks on model checking, whereas action-based approaches are less widespread and mostly appear in academic software tools dedicated to concurrency theory and process calculi. The present report follows this latter line, and addresses both equivalence checking and model checking, whereas state-based approaches often deal exclusively with model checking issues. It gathers numerous scientific results produced in the framework of the CADP verification toolbox [58] during the last 15–20 years, and presents a unifying overview of these results by situating them in a global, coherent landscape.

This paper is organized as follows. Section 2 gives definitions that will be used throughout the paper. Section 3 introduces *property-independent* compositional approaches; such approaches (among which *compositional model generation*, *semi-composition*, and *behavioral interfaces*) are closely related to equivalence checking and preserve a given equivalence relation, as well as all temporal logic properties adequate with respect to this equivalence relation. Section 4 presents *property-dependent* compositional approaches (among

which *property-dependent reductions* and *partial model checking*), which are specifically related to model checking and preserve the truth value of a particular set of properties (temporal logic or  $\mu$ -calculus formulas) of interest. Section 5 discusses high-level strategies for effective compositional verification, among which *smart reduction* techniques. Section 6 details how the aforementioned theoretical principles are implemented within the CADP toolbox, with a particular emphasis on the SVL language that provides a powerful, user-friendly means to specify compositional verification scenarios. Section 7 presents a concrete application in which (most of) the compositional capabilities of CADP can be illustrated. Finally, Section 8 concludes the paper and lists some open issues and directions for future work. The comparison to related work is not gathered into one unique section, but done at appropriate places throughout the paper.

## 2 Definitions

### 2.1 Vector notations

**Definition 1 (Vector)** A vector  $\vec{v}$  of size  $n$  is a total function on  $[1, n]$ . We use the following notations:

- We write  $|\vec{v}|$  for the size of the vector  $\vec{v}$ .
- For  $i \in [1, |\vec{v}|]$ , we write  $\vec{v}[i]$  for  $\vec{v}$  applied to  $i$ , denoting the element of  $\vec{v}$  stored at index  $i$ .
- We write  $(e_1, \dots, e_n)$  for the vector  $\vec{v}$  of size  $n$  such that  $(\forall i \in [1, n]) \vec{v}[i] = e_i$ . In particular,  $()$  denotes a vector of size 0.
- We write  $e^n$  for the vector  $\vec{v}$  of size  $n$  such that  $(\forall i \in [1, n]) \vec{v}[i] = e$ .
- Given two vectors  $\vec{v}_1$  and  $\vec{v}_2$ , we write  $\vec{v}_1 \# \vec{v}_2$  for the concatenation of  $\vec{v}_1$  and  $\vec{v}_2$ , i.e., the vector  $\vec{v}$  of size  $|\vec{v}_1| + |\vec{v}_2|$  such that for all  $i \in [1, |\vec{v}_1|]$ ,  $\vec{v}[i] = \vec{v}_1[i]$  and for all  $i \in [|\vec{v}_1| + 1, |\vec{v}_1| + |\vec{v}_2|]$ ,  $\vec{v}[i] = \vec{v}_2[i - |\vec{v}_1|]$ .
- Given an element  $e$  and a vector  $\vec{v}$ , we also write  $e :: \vec{v}$  for  $(e) \# \vec{v}$ .

**Definition 2 (Vector projection)** Given  $I = \{i_1, \dots, i_m\} \subseteq [1, n]$  with  $i_1 < \dots < i_m$  ( $0 \leq m \leq n$ ), we write  $\vec{v}|I$  for the projection of  $\vec{v}$  on to  $I$ , defined as the vector of size  $m$  such that  $(\forall j \in [1, m]) (\vec{v}|I)[j] = \vec{v}[i_j]$ . We write  $\vec{v}|\bar{I}$  for  $\vec{v}|([1, n] \setminus I)$ .

### 2.2 Labelled Transition Systems

We consider systems whose behavioural semantics can be represented using an LTS (*Labelled Transition System*).

**Definition 3 (LTS)** Let  $\mathcal{A}$  denote a finite set of elements called labels. An LTS is a tuple  $(\Sigma, A, \longrightarrow, s_0)$ , where:

- $\Sigma$  is a set of states,
- $A \subseteq \mathcal{A}$  is a set of labels (or actions),
- $\longrightarrow \subseteq \Sigma \times A \times \Sigma$  is the (labelled) transition relation,
- and  $s_0 \in \Sigma$  is the initial state.

We consider a particular label written  $\tau$  and called the invisible label, which denotes internal actions. All labels different from  $\tau$  are called the visible labels.

For an LTS  $S = (\Sigma, A, \longrightarrow, s_0)$ , we may write  $s \xrightarrow{a} s'$  instead of  $(s, a, s') \in \longrightarrow$ . We write  $\longrightarrow^*$  for the reflexive and transitive closure of  $\longrightarrow$ . Given a regular expression  $r$  built upon labels, we write  $s \xrightarrow{r} s'$  if there exist  $a_1, \dots, a_m$  and  $s_1, \dots, s_m$  ( $m \geq 0$ ) such that  $s \xrightarrow{a_1} s_1 \xrightarrow{a_2} \dots \xrightarrow{a_m} s_m$ , the sequence  $a_1 \dots a_m$  belongs to the regular language denoted by  $r$ , and  $s_m = s'$ .

**Definition 4 (LTS inclusion)** Let  $S_i = (\Sigma_i, A_i, \longrightarrow_i, s_i^0)$  for  $i \in \{1, 2\}$ . We say that  $S_1$  is included in  $S_2$ , written  $S_1 \subseteq S_2$ , if  $s_1^0 = s_2^0$ ,  $\Sigma_1 \subseteq \Sigma_2$ ,  $A_1 \subseteq A_2$ , and  $\longrightarrow_1 \subseteq \longrightarrow_2$ .

**Definition 5 (Equality modulo reachability)** Let  $S_i = (\Sigma_i, A_i, \longrightarrow_i, s_i^0)$  for  $i \in \{1, 2\}$ . The reachable part of  $S_i$  is the LTS  $(\Sigma'_i, A'_i, \longrightarrow'_i, s_i^0)$ , where:

- $\longrightarrow'_i = \{(s, a, s') \in \longrightarrow_i \mid s_i^0 \longrightarrow_i^* s\}$
- $A'_i = \{a \in A_i \mid \exists (s, a, s') \in \longrightarrow'_i\}$
- $\Sigma'_i = \{s_i^0\} \cup \{s' \in \Sigma_i \mid \exists (s, a, s') \in \longrightarrow'_i\}$

$S_1$  is equal to  $S_2$  modulo reachability, written  $S_1 \stackrel{r}{=} S_2$ , if the reachable parts of  $S_1$  and  $S_2$  are equal.

### 2.3 Networks of LTSs

To deal with compositions of LTSs, we use a general model called *network of LTSs* (or *network* for short) [85], which is inspired from the MEC [11] and FC2 [24] synchronization vectors.

**Definition 6 (Network of LTSs)** A network of LTSs, written  $N$ , is a pair  $(\vec{S}, V)$ , where  $\vec{S}$  is a vector of LTSs (called component LTSs), and  $V$  is a set of synchronization rules. Each synchronization rule has the form  $(\vec{t}, a)$ , where  $a \in \mathcal{A}$  is a label and  $\vec{t}$  is a vector comprising labels and occurrences of a special symbol  $\bullet \notin \mathcal{A}$ , such that  $|\vec{t}| = |\vec{S}|$ . Each  $\vec{t}$  is called a synchronization vector. The size of the network  $N$  is defined as  $|\vec{S}|$ . Let  $\vec{S}[i] = (\Sigma_i, A_i, \longrightarrow_i, s_i^0)$  ( $i \in [1, n]$ ). To  $N$  one can associate a “compound” LTS  $\text{lts}(N)$  that is the parallel composition of component LTSs. Each  $(\vec{t}, a) \in V$  defines transitions labelled by  $a$ , obtained either by synchronization (if more than one index  $i$  is such that  $\vec{t}[i] \neq \bullet$ ) or by interleaving (otherwise) of component LTS transitions. Formally,  $\text{lts}(N) = (\Sigma, A, \longrightarrow, \vec{s}_0)$ , where:

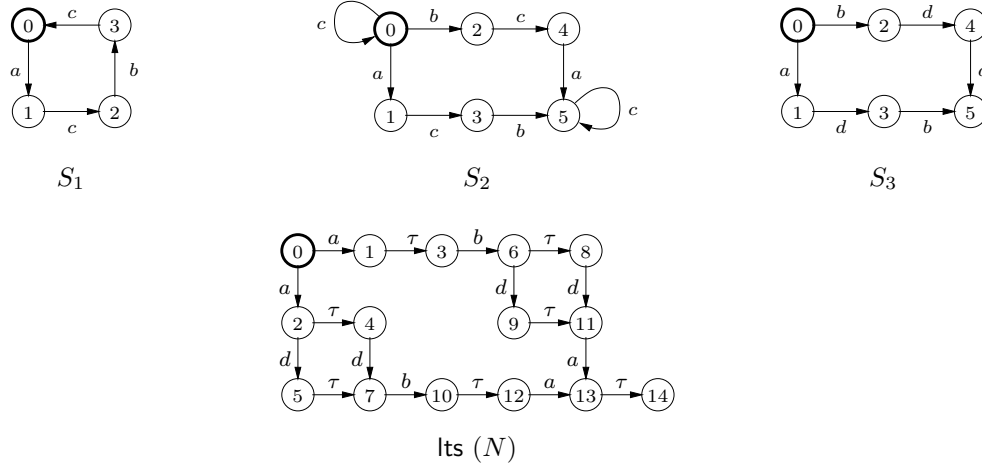
- $\Sigma = \Sigma_1 \times \dots \times \Sigma_n$ ,
- $A = \{a \mid \exists (\vec{t}, a) \in V\}$ ,
- $\vec{s}_0 = (s_1^0, \dots, s_n^0)$ , and
- $\longrightarrow$  is the relation satisfying  $\vec{s} \xrightarrow{a} \vec{s}'$  if and only if there exists  $(\vec{t}, a) \in V$  such that for all  $i \in [1, n]$ :

$$\begin{cases} \vec{s}'[i] = \vec{s}[i] & \text{if } \vec{t}[i] = \bullet \\ \vec{s}[i] \xrightarrow{\vec{t}[i]}_i \vec{s}'[i] & \text{otherwise} \end{cases}$$

We define  $\text{acv}(\vec{t}) = \{i \mid i \in [1, n] \wedge \vec{t}[i] \neq \bullet\}$ , the set of active LTS indices of the synchronization vector  $\vec{t}$ .

**Example 1** Let  $a, b, c$ , and  $d$  be labels, and  $S_1, S_2$ , and  $S_3$  be the LTSs defined in Figure 1 (top), where the initial states are denoted by bold circles. Let  $N = ((S_1, S_2, S_3), V)$  with  $V = \{((a, a, \bullet), a), ((a, \bullet, a), a), ((b, b, b), b), ((c, c, \bullet), \tau), ((\bullet, \bullet, d), d)\}$ . The first two synchronization rules of  $N$  express a nondeterministic synchronization on  $a$  between either  $S_1$  and  $S_2$ , or  $S_1$  and  $S_3$ . The third rule expresses a multiway synchronization on  $b$ . The fourth rule yields an internal ( $\tau$ ) transition. The fifth rule expresses full interleaving of transitions labelled by  $d$ . The compound LTS corresponding to  $N$  is depicted in Figure 1 (bottom).



Figure 1: Compound LTS corresponding to  $N$  defined in Example 1

$B ::= S$	LTS
$B_1 \parallel [A] B_2$	
$B_1 \parallel\parallel B_2$	
$B_1 \parallel B_2$	parallel compositions
<b>rename</b> $a_1 \rightarrow a'_1, \dots, a_n \rightarrow a'_n$ <b>in</b> $B_0$	rename
<b>hide</b> $A$ <b>in</b> $B_0$	hide
<b>cut</b> $A$ <b>in</b> $B_0$	cut

where  $A \subseteq \mathcal{A} \setminus \{\tau\}, \{a_1, \dots, a_n\} \subseteq \mathcal{A} \setminus \{\tau\}, (\forall i \neq j) a_i \neq a_j, \{a'_1, \dots, a'_n\} \subseteq \mathcal{A}$

Figure 2: Composition expressions

## 2.4 Composition expressions

Alternatively to networks, which compose LTSs in a flat manner, LTSs can be composed together using higher-level algebraic operators, such as parallel composition, label hiding, label renaming, and transition cutting, which we define below. Many operators available in the literature can be considered, e.g., from  $\mu\text{CRL}$  [72], CCS [108], CSP [125], LOTOS [79], E-LOTOS [80], or LNT [28] without theoretical difficulty, but for the sake of brevity, this paper considers only expressions comprising LTSs composed with LOTOS-like parallel composition, hiding, renaming, and cutting operators.

**Definition 7 (Composition expression)** *The syntax of composition expressions, written  $B, B_0, B_1, \dots$ , is given in Figure 2. Each composition expression  $B$  has semantics in terms of an LTS written  $\text{lts}(B)$ , defined as follows. By convention, given a composition expression  $B_i$ , we write  $(\Sigma_i, A_i, \longrightarrow_i, s_i^0)$  for  $\text{lts}(B_i)$ .*

1. The composition expression “ $S$ ” denotes the LTS  $S$ , whose semantics are defined by  $\text{lts}(S) = S$ .
2. The composition expression “ $B_1 \parallel [A] B_2$ ” denotes the parallel composition of  $B_1$  and  $B_2$  with synchronization on the labels belonging to  $A$ . Its semantics are defined by  $\text{lts}(B_1 \parallel [A] B_2) = (\Sigma_1 \times \Sigma_2, A_1 \cup A_2, \longrightarrow, (s_1^0, s_2^0))$ , where  $\longrightarrow$  is the smallest relation satisfying the following SOS (Structural Operational Semantics) [1] rules:

$$\frac{s_1 \xrightarrow{a}_1 s'_1 \quad a \notin A}{(s_1, s_2) \xrightarrow{a} (s'_1, s_2)} \quad \frac{s_2 \xrightarrow{a}_2 s'_2 \quad a \notin A}{(s_1, s_2) \xrightarrow{a} (s_1, s'_2)}$$

$$\frac{s_1 \xrightarrow{a}_1 s'_1 \quad s_2 \xrightarrow{a}_2 s'_2 \quad a \in A}{(s_1, s_2) \xrightarrow{a} (s'_1, s'_2)}$$

The notations “ $B_1 \parallel B_2$ ” and “ $B_1 \parallel B_2$ ” are respective shorthands for “ $B_1 \parallel [\emptyset] B_2$ ” and “ $B_1 \parallel [A \setminus \{\tau\}] B_2$ ”.

3. The composition expression “**rename**  $a_1 \rightarrow a'_1, \dots, a_n \rightarrow a'_n$  **in**  $B_0$ ” denotes  $B_0$  in which each visible action  $a_i$  ( $i \in [1, n]$ ) is replaced by the corresponding  $a'_i$ . For convenience, we may write  $\theta$  instead of  $a_1 \rightarrow a'_1, \dots, a_n \rightarrow a'_n$ . To  $\theta = a_1 \rightarrow a'_1, \dots, a_n \rightarrow a'_n$  we associate a total function  $\bar{\theta} : A \rightarrow A$  defined as follows:

$$\bar{\theta}(a) = \begin{cases} a' & \text{if } (\exists i \in [1, n]) a = a_i \wedge a' = a'_i \\ a & \text{otherwise} \end{cases}$$

The semantics of rename are defined by  $\text{lts}(\text{rename } \theta \text{ in } B_0) = (\Sigma_0, (A_0 \setminus \{a_1, \dots, a_n\}) \cup \{a'_1, \dots, a'_n\}, \longrightarrow, s_0^0)$ , where  $\theta = a_1 \rightarrow a'_1, \dots, a_n \rightarrow a'_n$  and  $\longrightarrow$  is the smallest relation satisfying the following SOS rule:

$$\frac{s \xrightarrow{a}_0 s'}{\bar{\theta}(a) \xrightarrow{\bar{\theta}(a)} s'}$$

4. The composition expression “**hide**  $A$  **in**  $B_0$ ” denotes  $B_0$  where labels in  $A$  are renamed into the invisible label  $\tau$ , i.e., turned into an unobservable action. Its semantics are defined by  $\text{lts}(\text{hide } A \text{ in } B_0) = (\Sigma_0, (A_0 \setminus A) \cup \{\tau\}, \longrightarrow, s_0^0)$ , where  $\longrightarrow$  is the smallest relation satisfying the following SOS rules:

$$\frac{s \xrightarrow{a}_0 s' \quad a \in A}{s \xrightarrow{\tau} s'} \quad \frac{s \xrightarrow{a}_0 s' \quad a \notin A}{s \xrightarrow{a} s'}$$

If  $A = \{a_1, \dots, a_n\}$ , then **hide**  $A$  **in**  $B$  is equivalent to **rename**  $a_1 \rightarrow \tau, \dots, a_n \rightarrow \tau$  **in**  $B$ .

5. The composition expression “**cut**  $A$  **in**  $B_0$ ” denotes  $B_0$  where every transition labelled by an element of  $A$  is deleted. Its semantics are defined by  $\text{lts}(\text{cut } A \text{ in } B_0) = (\Sigma_0, A_0 \setminus A, \longrightarrow, s_0^0)$ , where  $\longrightarrow$  is the smallest relation satisfying the following SOS rule:

$$\frac{s \xrightarrow{a}_0 s' \quad a \notin A}{s \xrightarrow{a} s'}$$

It follows that  $\xrightarrow{a} \subseteq \xrightarrow{a}_0$ .

Note that a composition expression is a syntactic algebraic expression structured as a tree, which we call the *component hierarchy*. We could further enrich composition expressions with operators borrowed from other process calculi than LOTOS.

There exists a mapping from composition expressions to networks: every composition expression  $B$  can be translated into a network of LTSs, by flattening the component hierarchy. This network is written  $\text{net}(B)$  and defined as the couple  $(\text{ind}(B), \text{sync}(B))$ , where  $\text{ind}(B)$  and  $\text{sync}(B)$  denote respectively the vector of

component LTSs and the synchronization rules corresponding to  $B$ . These functions are defined as follows:

$$\begin{aligned}
\text{ind}(S) &= (S) \\
\text{ind}(B_1 \parallel [A] B_2) &= \text{ind}(B_1) \# \text{ind}(B_2) \\
\text{ind}(\mathbf{hide} A \mathbf{in} B_0) &= \text{ind}(B_0) \\
\text{ind}(\mathbf{cut} A \mathbf{in} B_0) &= \text{ind}(B_0) \\
\text{ind}(\mathbf{rename} \theta \mathbf{in} B_0) &= \text{ind}(B_0) \\
\text{sync}(S) &= \{(a, a) \mid a \in A\} \text{ where } S = (\Sigma, A, \longrightarrow, s_0) \\
\text{sync}(B_1 \parallel [A] B_2) &= \\
&\{(\vec{t}_1 \# \bullet \mid \text{ind}(B_2), a) \mid (\vec{t}_1, a) \in \text{sync}(B_1) \wedge a \notin A\} \cup \\
&\{(\bullet \mid \text{ind}(B_1), \vec{t}_2, a) \mid (\vec{t}_2, a) \in \text{sync}(B_2) \wedge a \notin A\} \cup \\
&\{(\vec{t}_1 \# \vec{t}_2, a) \mid (\vec{t}_1, a) \in \text{sync}(B_1) \wedge (\vec{t}_2, a) \in \text{sync}(B_2) \wedge a \in A\} \\
\text{sync}(\mathbf{rename} \theta \mathbf{in} B_0) &= \\
&\{(\vec{t}_0, \bar{\theta}(a)) \mid (\vec{t}_0, a) \in \text{sync}(B_0)\} \\
\text{sync}(\mathbf{hide} A \mathbf{in} B_0) &= \\
&\{(\vec{t}_0, \tau) \mid (\vec{t}_0, a) \in \text{sync}(B_0) \wedge a \in A\} \cup \\
&\{(\vec{t}_0, a) \mid (\vec{t}_0, a) \in \text{sync}(B_0) \wedge a \notin A\} \\
\text{sync}(\mathbf{cut} A \mathbf{in} B_0) &= \\
&\{(\vec{t}_0, a) \mid (\vec{t}_0, a) \in \text{sync}(B_0) \wedge a \notin A\}
\end{aligned}$$

**Example 2** *The network of Example 1 corresponds to the composition expression  $\mathbf{hide} c \mathbf{in} (S_1 \parallel [a, b, c] (S_2 \parallel [b] S_3))$ .*

Networks are more general than composition expressions, as illustrated by the following example.

**Example 3** *Networks enable  $m$ -among- $n$  synchronization [61], where any  $m$  LTSs among the  $n$  LTSs of the composition synchronize on a given label. For instance, 2-among-3 synchronization on  $a$  between LTSs  $S_1$ ,  $S_2$ , and  $S_3$  can be achieved using the network  $((S_1, S_2, S_3), \{(a, a, \bullet), (a, \bullet, a), (\bullet, a, a)\})$ . This type of synchronization cannot be described using classical process algebraic operators [61].*

## 2.5 LTS Equivalences

Equivalence relations on LTSs characterize semantic equivalences between systems. Several relations are available in the literature, differing mainly in their treatment of invisible labels. We focus here on a few of them, namely strong bisimulation, branching bisimulation and its divergence-sensitive variant, and  $\tau^*.a$  equivalence.

**Definition 8 (Strong bisimulation [116])** *A strong bisimulation is a symmetric relation  $R \subseteq \Sigma \times \Sigma$  such that if  $(s_1, s_2) \in R$  then: for all  $s_1 \xrightarrow{a} s'_1$ , there exists  $s'_2$  such that  $s_2 \xrightarrow{a} s'_2$  and  $(s'_1, s'_2) \in R$ .*

*Two states  $s_1$  and  $s_2$  are strongly bisimilar if there exists a strong bisimulation  $R$  such that  $(s_1, s_2) \in R$ . Two LTSs are strongly bisimilar if their initial states are strongly bisimilar.*

**Definition 9 (Branching bisimulation [136, 137])** *A branching bisimulation is a symmetric relation  $R \subseteq \Sigma \times \Sigma$  such that if  $(s_1, s_2) \in R$  then: for all  $s_1 \xrightarrow{a} s'_1$ , either  $a = \tau$  and  $(s'_1, s_2) \in R$ , or there exists a sequence  $s_2 \xrightarrow{\tau^*} s'_2 \xrightarrow{a} s''_2$  such that  $(s_1, s'_2) \in R$  and  $(s'_1, s''_2) \in R$ .*

*Two states  $s_1$  and  $s_2$  are branching bisimilar if there exists a branching bisimulation  $R$  such that  $(s_1, s_2) \in R$ . Two LTSs are branching bisimilar if their initial states are branching bisimilar.*

Branching bisimulation does not distinguish between inaction and a cycle of internal actions. Divergence-sensitive branching bisimulation is introduced to take into account cycles of internal actions.

**Definition 10 (Divergence-sensitive branching bisimulation [136, 137])** A divergence-sensitive branching bisimulation (or divbranching bisimulation for short) is a branching bisimulation  $R$  such that if  $(s_1^0, s_2^0) \in R$  and there is an infinite sequence  $s_1^0 \xrightarrow{\tau} s_1^1 \xrightarrow{\tau} s_1^2 \xrightarrow{\tau} \dots$  with  $(s_1^i, s_2^0) \in R$  for all  $i \geq 0$ , then there is an infinite sequence  $s_2^0 \xrightarrow{\tau} s_2^1 \xrightarrow{\tau} s_2^2 \xrightarrow{\tau} \dots$  such that  $(s_1^i, s_2^j) \in R$  for all  $i, j \geq 0$ .

Two states  $s_1$  and  $s_2$  are divbranching bisimilar if there exists a divbranching bisimulation  $R$  such that  $(s_1, s_2) \in R$ . Two LTSs are divbranching bisimilar if their initial states are divbranching bisimilar.

**Definition 11 ( $\tau^*.a$  equivalence [51])** A  $\tau^*.a$  equivalence is a symmetric relation  $R \subseteq \Sigma \times \Sigma$  such that if  $(s_1, s_2) \in R$  then: for each sequence  $s_1 \xrightarrow{\tau^*.a} s'_1$  where  $a$  is a visible label, there exists a sequence  $s_2 \xrightarrow{\tau^*.a} s'_2$  such that  $(s'_1, s'_2) \in R$ .

Two states  $s_1$  and  $s_2$  are  $\tau^*.a$  equivalent if there exists a  $\tau^*.a$  equivalence  $R$  such that  $(s_1, s_2) \in R$ . Two LTSs are  $\tau^*.a$  equivalent if their initial states are  $\tau^*.a$  equivalent.

Other equivalence relations are defined in the literature and possibly used in our framework of composition expressions and networks, such as observation equivalence [108], safety equivalence [23], trace equivalence (also known as *language equivalence*), and weak trace equivalence [26]. For the sake of conciseness, we do not give their definitions in this paper.

Equivalence relations and their associated preorders can be used to compare the behaviour of two systems. Equivalence relations can also be used to compute a “*canonical form*” for each LTS, by identifying and merging equivalent states and by replacing equivalent transition sequences by a single, representative one. Such an operation is often called LTS *minimization*, or *reduction* if applied partially. In the case of strong and branching bisimulations, the resulting LTS is indeed the smallest (in both the number of states and the number of transitions) of its equivalence class. In the case of  $\tau^*.a$  equivalence, the representative of every sequence of transitions matching  $\tau^*.a$  is a single transition labelled by  $a$ , i.e., the  $\tau$ -transitions are eliminated. This yields an LTS that is not necessarily minimal in the number of transitions, but always minimal in the number of states (and obviously in the number of  $\tau$ -transitions). Note that there exist other equivalence relations whose canonical forms may be dramatically larger (in the number of states) than the smallest elements of their equivalence classes. This is the case of trace and weak trace equivalences, whose canonical forms require determinization. This illustrates an advantage of bisimulations and equivalence relations preserving the branching structure of LTSs over relations dealing with traces.

**Definition 12** Given an LTS equivalence relation  $R$  and an LTS  $S$ , we write  $\text{reduce}_R(S)$  for the LTS corresponding to  $S$  minimized modulo  $R$ . Given a network  $N$  (respectively a composition expression  $B$ ) and an LTS equivalence relation  $R$ , we write  $\text{reduce}_R(N)$  (respectively  $\text{reduce}_R(B)$ ) as a shorthand for  $\text{reduce}_R(\text{lts}(N))$  (respectively  $\text{reduce}_R(\text{lts}(B))$ ).

## 2.6 Congruence results

**Definition 13 (Congruence for networks)** An LTS equivalence relation  $R$  is a congruence for networks if and only if for all networks  $N = (\vec{S}, V)$ , for all  $i \in [1, n]$  where  $n$  is the size of  $N$ , and for all  $S_i$  equivalent to  $\vec{S}[i]$  modulo  $R$ , then  $\text{lts}(\vec{S}, V)$  is equivalent modulo  $R$  to  $\text{lts}(\vec{S}', V)$  where  $\vec{S}'$  is the same as  $\vec{S}$  in which  $\vec{S}[i]$  has been replaced by  $S'_i$ .

**Proposition 1** Strong bisimulation and trace equivalence are congruences for networks [85].

**Proposition 2** Branching bisimulation, divergence-sensitive branching bisimulation, observation equivalence, safety equivalence, and weak trace equivalence [26], are congruences for networks provided that the synchronization rules satisfy the following constraints [85] regarding the internal transitions of component LTSs, for all  $i \in [1, n]$ :

- Internal transitions should not be synchronized:

$$(\vec{t}, a) \in V \wedge \vec{t}[i] = \tau \implies \text{acv}(\vec{t}) = \{i\}$$

- *Internal transitions should not be renamed:*

$$(\vec{t}, a) \in V \wedge \vec{t}[i] = \tau \implies a = \tau$$

- *Internal transitions should not be cut:*

$$\xrightarrow{\tau}_i \neq \emptyset \implies (\exists(\vec{t}, \tau) \in V) \vec{t}[i] = \tau$$

In the sequel, we consider only networks satisfying the constraints defined in Proposition 2. Note that these constraints are natural, and that they are satisfied by the networks obtained from composition expressions by translation.

Note also that  $\tau^*.a$  equivalence is neither a congruence for networks nor for composition expressions. However, it can be used compositionally as a pre-reduction for weaker congruence relations, such as safety and weak trace equivalences.

## 3 Property-independent compositional approaches

### 3.1 Basic compositional LTS generation

In its simplest forms [49, 95, 126, 140, 128, 129, 134, 125], compositional verification (also called incremental reduction [126], incremental reachability analysis [128, 129], compositional state space generation [134], or inductive compression [125]) consists in replacing each component LTS by an *abstraction*, simpler than the original component but still preserving the properties to be verified on the whole system.

The approach for compositionality presented in this section deals with the construction of a minimal (or, at least, reduced) LTS modulo a given equivalence relation such as strong, branching, or observational equivalence, starting from a composition expression or a network. The chosen abstraction is thus LTS minimization modulo an appropriate equivalence relation: instead of  $\text{lts}((S_1, \dots, S_n), V)$ , we compute  $\text{lts}((\text{reduce}_R(S_1), \dots, \text{reduce}_R(S_n)), V)$ . For weak equivalence relations, such as branching bisimulation, this may yield substantial reductions of the initial LTS, and consequently facilitate its subsequent analysis.

This abstraction is sound as long as the equivalence relation  $R$  is a congruence for the composition expression. Minimization can also be applied at any intermediate level in the composition, thus opening the way for various compositional minimization strategies. This will be detailed in Section 5.

Although basic compositional LTS generation has been applied successfully to some complex systems (e.g., [50, 29] in the case of the LOTOS language [79]), it may be counter-productive in some other cases: generating the LTS of each component separately may lead to state explosion, whereas the generation of the whole system at once might succeed if components constrain each other when composed in parallel. Indeed, there may be many states of a component that, although useful in a general environment, are useless (i.e., never explored) in a particular environment.

### 3.2 Interfaces and projections

Enhanced compositional verification approaches [69, 31, 139, 32, 33, 70, 83, 30, 63] have been proposed to generate the LTS of a component by taking into account *interface constraints* (also known as *environment constraints* or *context constraints*). These constraints express the behavioural restrictions imposed on the considered component by synchronization with its neighbour components. Taking into account the environment of a component permits local elimination of states and transitions unreachable in the LTS of the whole system.

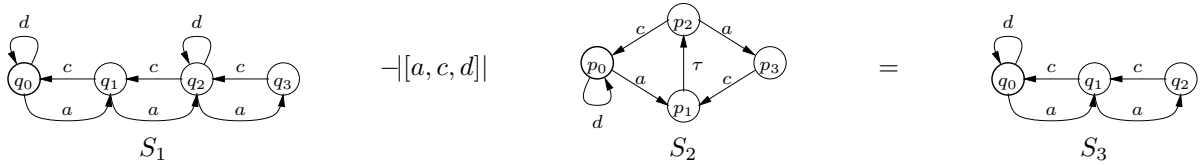
In general, interface constraints are expressed in the form of an LTS called *interface*. There are two approaches to restrict the behaviour of a component w.r.t. an interface. In the first one [30, 31, 32, 33, 63], the component is composed in parallel with the interface, which must have been transformed beforehand so that the composition does not affect the global behaviour of the system (a property known as *context transparency*).

In the second approach, the component is constrained using a specific *semi-composition* operator [69, 70, 83], which cuts the component states and transitions that cannot be reached when considering the traces of

the interface as the only possible interactions between the component and its environment. We consider the second approach. The semi-composition operator is defined as follows.

**Definition 14 (Semi-composition)** Let  $S_i = (\Sigma_i, A_i, \longrightarrow_i, s_i^0)$  ( $i = 1, 2$ ) be two LTSs,  $A$  be a set of visible labels, and  $(\Sigma_{||}, A_{||}, \longrightarrow_{||}, s_{0||}) = \text{Its}(S_1 || [A] S_2)$ . The semi-composition of  $S_1$  and  $S_2$ , written “ $S_1 \text{--[} A \text{]} S_2$ ”, is the LTS  $(\Sigma, A_1, \longrightarrow, s_1^0)$ , where  $\longrightarrow = \longrightarrow_1 \cap \{(s_1, a, s'_1) \mid (s_1^0, s_2^0) \longrightarrow_{||}^* (s_1, s_2) \xrightarrow{a}_{||} (s'_1, s'_2)\}$  and  $\Sigma = \{s \mid (\exists a, s') s \xrightarrow{a} s' \vee s' \xrightarrow{a} s\}$ .  $A$  is called the synchronization set and the pair  $(A, S_2)$  is called the interface<sup>1</sup>. We say that a label  $a \in A_1$  is controlled by the interface  $(A, S_2)$  if  $a \in A$ . We write  $S_1 \text{--[} A \setminus \{\tau\} \text{]} S_2$ .

**Example 4** The following holds:



Transitions  $q_2 \xrightarrow{d} q_2, q_2 \xrightarrow{a} q_3$ , and  $q_3 \xrightarrow{c} q_2$  do not belong to  $S_3$  because they are not reachable in  $S_1 || [a, c, d] S_2$ . Therefore, state  $q_3$  is unreachable in  $S_3$ .

The following properties ensure the applicability of semi-composition.

**Proposition 3** Semi-composition cannot increase the size of the LTS to which it is applied:  $(\forall A, S_1, S_2) \text{Its}(S_1 \text{--[} A \text{]} S_2) \subseteq S_1$ .

**Proposition 4**  $(\forall A, S_1, S_2) \text{Its}(S_1 \text{--[} A \text{]} S_2) = S_1$  if the visible traces of “hide  $A \setminus A$  in  $S_1$ ” are included in the visible traces of “hide  $A \setminus A$  in  $S_2$ ”.

**Proposition 5 (Laws of semi-composition [83])**

$$S_1 || [A] S_2 \stackrel{r}{=} (S_1 \text{--[} A \text{]} S_2) || [A] S_2 \quad (1)$$

$$(S_1 || [A] S_3) || [A'] S_2 \stackrel{r}{=} ((S_1 \text{--[} A' \cap (A \cup (A_1 \setminus A_3)) \text{]} S_2) || [A] S_3) || [A'] S_2 \quad (2)$$

where  $A_1$  is the set of labels of  $S_1$ , and  
 $A_3$  is the set of labels of  $S_3$

$$(\text{hide } A \text{ in } S_1) || [A'] S_2 \stackrel{r}{=} (\text{hide } A \text{ in } (S_1 \text{--[} A' \setminus A \text{]} S_2)) || [A'] S_2 \quad (3)$$

Proposition 5 defines how semi-composition can be used to reduce  $S_1$  given an LTS  $S_2$  in its environment, by removing the unreachable states and transitions, without losing any temporal property of the system. Note that, unlike the approach of Cheung & Kramer, which requires that the interface be context transparent — and thus be transformed into a deterministic LTS using a well-known but expensive algorithm — no restriction is made here on the shape of  $S_2$ .

**Proposition 6**  $S_1 \text{--[} A \text{]} S_2 = S_1 \text{--[} A \text{]} S'_2$  if “hide  $A \setminus A$  in  $S_2$ ” and “hide  $A \setminus A$  in  $S'_2$ ” have the same visible traces.

This proposition implies that the uncontrolled labels can be hidden in the interface and that the resulting interface can then be minimized modulo any relation preserving visible traces (e.g., *safety equivalence* [23]), which permits reduction of the number of states to explore while calculating semi-composition. Safety minimization is less expensive than determinization and, unlike determinization which can induce a dramatic growth of the LTS, yields an LTS that never contains more states than the input. Minimization of the interface is not mandatory but helps reduce the cost of semi-composition.

<sup>1</sup>This definition of semi-composition is simpler but equivalent to that given in [83].

### 3.3 Automatic generation of interfaces

Interfaces can be either written by the user (and possibly checked automatically [83]) or generated automatically. Although automated generation has the neat advantage to relieve users from the burden of calculating appropriate constraints, existing automated interface generation techniques often undergo two main limitations: first, these techniques are specific to a given parallel composition operator and thus not directly applicable in the framework of concurrent languages featuring different and/or more general parallel composition operators; second, as already observed in [31], they may fail to capture effective interface constraints due to deficiencies in their analysis of synchronizations between components.

In this section, we propose to generate interfaces automatically from networks of LTSs, following the approach proposed in [86]. The network of LTS intermediate representation permits the derivation of effective interface constraints imposed on a given component by a subset of its neighbours automatically, independently of the component hierarchy and of the nature of the parallel composition operators. This permits combination of constraints induced by distant components, and improvement of the accuracy of interfaces by exploiting more precisely the synchronizations between components. For this reason, we qualify as *refined* the interfaces generated using this technique.

A method was previously proposed in [83] to compute automatically an exact interface in the framework of composition expressions built upon parallel composition and label hiding. In this method, defined inductively based on the semi-composition laws described in Proposition 5, two component LTSs  $S_1$  and  $S_2$  are selected and a synchronization set  $A$  is computed such that  $S_1$  can be replaced by  $S_1 \dashv[A] S_2$  without changing the compound LTS. The interface  $(A, S_2)$  built using this method generally does not give the best account of environment constraints.

Here, we propose to generate automatically interfaces that give a better account of environment constraints, using *refined interface generation* [86]. This method uses the information available in the intermediate network model to compute an interface capturing the constraints imposed on a given component  $P$  in a concurrent system by one or several neighbour components. This interface can then be semi-composed with  $P$  on-the-fly, so as to restrict  $P$ 's behaviour.

**Definition 15 (Refined interface generation)** *Given a network  $N = (\vec{S}, V)$  of size  $n$ , an index  $k \in [1, n]$ , and a nonempty set of indices  $I \subseteq [1, n] \setminus \{k\}$ , the refined interface of  $\vec{S}[k]$  capturing constraints induced by  $\{\vec{S}[i] \mid i \in I\}$ , written  $\text{refint}(N, k, I)$ , is the couple  $(\mathcal{A}, S_I)$ , where:*

$$S_I = \text{lts}(\vec{S}|I, V_I)$$

$$V_I = \{(\vec{t}|I, \vec{t}[k]) \mid (\exists a) (\vec{t}, a) \in V \wedge \vec{t}[k] \neq \bullet\} \cup \{(\vec{t}|I, \tau) \mid (\exists a) (\vec{t}, a) \in V \wedge \vec{t}[k] = \bullet\}$$

The *refint* operation may create synchronization rules of the form  $(\bullet^n, a)$ , which induce a self-looping transition labelled by  $a$  in each state of the interface. Some of these synchronization rules can be eliminated as follows:

- Every synchronization rule of the form  $(\bullet^n, \tau)$  can merely be removed, as  $\tau$  loops do not affect the visible traces of the interface.
- Every synchronization rule of the form  $(\bullet^n, a)$  where  $a \neq \tau$  can be removed and then  $a$  removed from the synchronization set if the set of synchronization rules does not contain another rule with the same label  $a$  as right-hand side. Indeed, for all  $\vec{S}, S', A$ , and  $V$  in which  $a$  does not occur as a right-hand side,  $S' \dashv[A] \text{lts}(\vec{S}, V \cup (\bullet^n, a)) = S' \dashv[A \setminus \{a\}] \text{lts}(\vec{S}, V)$ .

The refined interface thus consists of a product of the LTSs  $\vec{S}[i]$  ( $i \in I$ ), synchronized by synchronization rules derived systematically from the synchronization rules of  $N$ , each rule  $(\vec{t}, a)$  being transformed into a rule  $(\vec{t}|I, \vec{t}[k])$  if  $\vec{t}[k] \neq \bullet$ , or  $(\vec{t}|I, \tau)$  otherwise. Therefore, whenever a transition  $\vec{q} \xrightarrow{a} \vec{q}'$  can be fired in  $\text{lts}(N)$  using a synchronization rule  $(\vec{t}, a)$  with  $\vec{t}[k] \neq \bullet$ , then the participating transition  $\vec{q}[k] \xrightarrow{\vec{t}[k]} \vec{q}'[k]$  of  $\vec{S}[k]$  is also a transition of  $\vec{S}[k] \dashv[A] S_I$ . Conversely, transitions of  $\vec{S}[k]$  that cannot participate in any mandatory synchronization with  $S_I$  are eliminated by the semi-composition  $\vec{S}[k] \dashv[A] S_I$ .

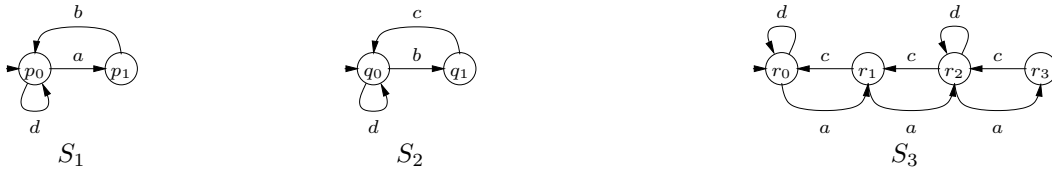
**Example 5** Consider the network  $N$  displayed at the left below, with arbitrary LTSs  $S_1, \dots, S_4$ . The refined interface of  $S_1$  capturing constraints induced by  $S_3$  and  $S_4$ , written  $\text{refint}(N, 1, \{3, 4\})$ , has the LTS corresponding to the network displayed at the right below. Note the projection on  $S_3$  and  $S_4$ , and observe that the right-hand sides of synchronization rules in the result are the elements of column  $S_1$ , where  $\bullet$  is renamed into  $\tau$ .

$$\text{refint} \left( \left( \left( \begin{array}{c} (S_1, S_2, S_3, S_4), \\ \left\{ \begin{array}{l} ((a_1, a_2, a_3, a_4), a), \\ ((\bullet, b_2, b_3, \bullet), b), \\ ((c_1, c_2, \bullet, \bullet), c) \end{array} \right\} \end{array} \right), 1, \{3, 4\} \right) = \text{Its} \left( \left( \begin{array}{c} (S_3, S_4), \\ \left\{ \begin{array}{l} ((a_3, a_4), a_1), \\ ((b_3, \bullet), \tau), \\ ((\bullet, \bullet), c_1) \end{array} \right\} \end{array} \right) \right)$$

**Proposition 7** Let  $N$  be a network of LTSs of size  $n$ ,  $k \in [1, n]$ ,  $I \subseteq [1, n] \setminus \{k\}$ , and  $(A, S_I) = \text{refint}(N, k, I)$ . If  $\vec{S}'$  is the vector of LTSs of size  $n$  defined by  $(\forall i \in [1, n] \setminus \{k\}) \vec{S}'[i] = \vec{S}[i]$  and  $\vec{S}'[k] = \vec{S}[k] \dashv [A] S_I$ , then  $\text{Its}(\vec{S}, V) \stackrel{r}{=} \text{Its}(\vec{S}', V)$ .

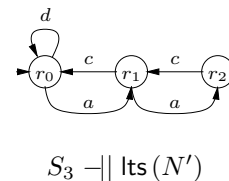
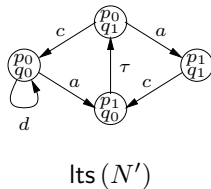
The following examples show that refined interfaces may constrain components better than the interfaces generated by the method proposed in [83].

**Example 6** Let  $B = S_1 \parallel [a, b, d] (S_2 \parallel [c, d] S_3)$  with  $S_1, S_2$ , and  $S_3$  defined as follows:



According to the semi-composition laws of Proposition 5,  $S_3$  can be replaced in  $B$  either by  $S_3 \dashv [a, d] S_1$ , or by  $S_3 \dashv [c, d] S_2$ , but both expressions result in  $S_3$  itself. Yet, one can see that actions  $a$  and  $c$  are executed with some alternation in  $B$ , due to the mandatory synchronization on  $b$  between  $S_1$  and  $S_2$ . As a consequence, state  $r_3$  is not reachable in  $B$ . To capture such a constraint, it should be possible to build an interface that takes simultaneously into account the constraints induced by both  $S_1$  and  $S_2$ , even though there is no sub-expression of  $B$  containing  $S_1$  and  $S_2$  only (parallel composition is not associative). This is not possible using the method described in [83]<sup>2</sup>, but this can be done using refined interface generation. To this aim, the expression  $B = S_1 \parallel [a, b, d] (S_2 \parallel [c, d] S_3)$  is translated into the network  $N$  displayed below and then  $S_3$  may be restricted using the refined interface  $\text{Its}(N') = \text{refint}(N, 3, \{1, 2\})$  that takes simultaneously both  $S_1$  and  $S_2$  into account, where  $N'$  and  $\text{Its}(N')$  are displayed below.  $S_3 \dashv \parallel \text{Its}(N')$ , also displayed below, is strictly included in  $S_3$  as the unreachable state  $r_3$  and transitions  $r_2 \xrightarrow{a} r_3, r_3 \xrightarrow{c} r_2$ , and  $r_2 \xrightarrow{d} r_2$  are deleted.

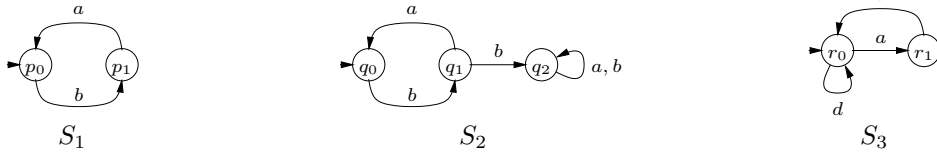
$$N = \left( \begin{array}{c} (S_1, S_2, S_3), \\ \left\{ \begin{array}{l} ((a, \bullet, a), a), \\ ((b, b, \bullet), b), \\ ((\bullet, c, c), c), \\ ((d, d, d), d) \end{array} \right\} \end{array} \right) \quad N' = \left( \begin{array}{c} (S_1, S_2), \\ \left\{ \begin{array}{l} ((a, \bullet), a), \\ ((b, b), \tau), \\ ((\bullet, c), c), \\ ((d, d), d) \end{array} \right\} \end{array} \right)$$



<sup>2</sup>This limitation holds similarly for the approach described in [31].



**Example 7** Let  $B = S_1 \parallel [a, b] (S_2 \parallel [a] S_3)$  with  $S_1, S_2$ , and  $S_3$  defined as follows:



According to the semi-composition laws of Proposition 5,  $S_2$  can be replaced by  $S_2 \dashv [a] S_1$ , but this expression yields  $S_2$  itself. Yet, it is clear from  $S_1$  and the synchronizations in  $B$  that state  $q_2$  of  $S_2$  is unreachable in  $B$ , as two successive  $b$  actions cannot be fired without an  $a$  in between. A better interface should permit to take into account the environment constraints due to synchronizations on  $b$ , even though every  $b$  of  $S_1$  does not necessarily synchronize with a  $b$  of  $S_2$ . Such an interface can be obtained using refined interface generation. To this aim, the expression  $B = S_1 \parallel [a, b] (S_2 \parallel [a] S_3)$  is translated into the network  $N$  displayed below and then  $S_2$  may be restricted using the refined interface  $\text{lts}(N') = \text{refint}(N, 2, \{1\})$  that takes  $S_1$  into account, where  $N'$  and  $\text{lts}(N')$  are displayed below. In practice,  $\text{lts}(N')$  can be minimized modulo safety equivalence, yielding an LTS with 2 states and 3 transitions.  $S_2 \dashv \text{lts}(N')$  is isomorphic to  $S_1$ .

$$N = \left( \begin{array}{c} (S_1, S_2, S_3), \\ \left\{ \begin{array}{l} ((a, a, a), a), \\ ((b, b, \bullet), b), \\ ((b, \bullet, b), b), \\ ((\bullet, \bullet, d), d) \end{array} \right\} \end{array} \right) \quad N' = \left( \begin{array}{c} (S_1), \\ \left\{ \begin{array}{l} ((a), a), \\ ((b), b), \\ ((b), \tau), \\ ((\bullet), \tau) \end{array} \right\} \end{array} \right)$$

This example shows that by taking a better account of the synchronization structure of the system, the *refint* operation permits refinement of the interface with respect to that obtained using equation (2), turning the set of visible traces of the interface from  $a^*$  with  $b$  uncontrolled using the method of [83] to  $a^* + b + (ba^+)^*$  using refined interface generation. The latter set of traces does not contain any trace with two consecutive  $b$ 's, thus disabling the transition  $q_1 \xrightarrow{b} q_2$  in  $S_2$  and making state  $q_2$  and transitions  $q_2 \xrightarrow{a} q_2$ ,  $q_2 \xrightarrow{b} q_2$  also unreachable.

Algorithmically, refined interface generation has the same complexity as the synchronization product of the LTSs taken into account in the environment. In practice, the cost of computing the interface can be reduced by minimizing the component LTSs participating in the interface modulo safety equivalence. This is correct, because safety equivalence is a congruence for networks. In addition, well-known partial order reductions preserving visible traces can be used to further reduce interfaces on-the-fly during their construction.

So far, refined interface generation required that each component of the concurrent system under verification (usually written in a high-level language) was replaced by its LTS. This apparently contradicts the claim that refined interfaces can be used to restrict processes on the fly. However, one can see in Definition 15 that the states and transitions of the component to be restricted ( $\vec{S}[k]$ ) are not needed for interface generation. In practice, only the visible labels of  $\vec{S}[k]$  are needed to compute the synchronization rules of the network from higher-level operators. To do so,  $\vec{S}[k]$  can be replaced by an abstraction consisting of an arbitrary (and much smaller) LTS containing the same set of labels. In fact, the method remains correct if the abstraction contains a superset of  $\vec{S}[k]$ 's labels, although the reduction obtained on  $\vec{S}[k]$  by semi-composition generally increases while the set of labels of the abstraction gets closer to the exact set of labels of  $\vec{S}[k]$ . In principle, the abstraction can be generated automatically by examining the gates (or channels) occurring in the high-level specification and the types of their data, and to enumerate data of this type appropriately.

## 4 Property-dependent compositional approaches

There are two ways of expressing and checking properties on a given concurrent system: one (*equivalence checking*) is based on behavioural relations and the other (*model checking*) is based on properties expressed, e.g., as formulas of some temporal logic.

The approach for compositionality presented in the previous section was based on equivalences, and independent from any particular set of properties, in the sense that it preserves all properties compatible with this equivalence. In this section, we consider alternative property-based approaches, which exploit compositionality while preserving a given set of temporal logic formulas to be verified. We develop this idea along two complementary ways:

- The first way aims at checking a set of temporal formulas on a (monolithic) LTS after reducing it as much as possible w.r.t. each formula. This is done by identifying the maximal set of actions that do not influence the truth value of a formula, hiding these actions in the LTS, and minimizing the LTS modulo an equivalence relation preserving the formula. The minimization can be carried out using the compositional equivalence-based approach of Section 3, which therefore becomes specialized for a given formula.
- The second way aims at checking one temporal formula on the parallel composition of several LTSs by taking them into account individually following the *partial model checking* method [8]. This is done by combining the formula with one of the LTSs and then checking the resulting formula recursively on the parallel composition of the remaining LTSs, until all of them have been taken into account. At intermediate stages, simplifications must be applied to the formula to keep it as small as possible.

These two ways of verification can be combined naturally by applying maximal hiding before starting partial model checking, and thus cumulating their benefits. One of our achievements is the reformulation of partial model checking in terms of graph manipulations, which paves the way towards the connection between compositional model checking and compositional state space construction.

We first revisit briefly the definitions of the temporal logics considered and the associated on-the-fly model checking methods. Then, we present the property-preserving reductions based on maximal hiding, and the partial model checking approach on networks of LTSs.

## 4.1 Temporal logics

Temporal logics [121, 122] were introduced more than three decades ago as a means for reasoning about the behaviour of concurrent systems. These formalisms consist of a small set of temporal operators expressing the logical precedence of states or events over time. Since the early proposals, a plethora of temporal logics emerged in the literature (see [48, 110, 25] for surveys), having different interpretation models, expressiveness, conciseness, and complexity of model checking. Regarding the interpretation models, two orthogonal criteria are most often used for classifying temporal logics: state-based (resp. action-based) logics specify properties of states (resp. actions, or events) of the system under study, and linear-time (resp. branching-time) logics are interpreted over execution sequences (resp. trees) of the system. Traditionally, state-based logics are interpreted on Kripke structures (KSs), in which states are labelled with atomic propositions, and action-based logics are interpreted on LTSs. The table below shows the most representative temporal logics in each of the four classes induced by the two criteria.

	Linear	Branching
State	LTL [122]	CTL [44]
Action	ALTL [64]	ACTL [111]

Besides these “pure” temporal logics, many other temporal logics combining linear-time and branching-time operators have been proposed, e.g., CTL\* [45] and ACTL\* [112] in the state- and action-based setting, respectively. Variants of these logics combining state- and action-based properties were also defined, having as interpretation models Kripke transition systems (KTSs) [110], i.e., state spaces containing relevant information on both states and transitions. A class of very expressive temporal logics are those comprising fixed point operators, such as the modal  $\mu$ -calculus ( $L\mu$ ) [82], which subsumes virtually all other temporal logic operators, and therefore can be seen as an assembly language for reasoning on LTSs.

Finally, to increase their expressiveness and facilitate their usage for value-passing systems, temporal logics were extended with various constructs: regular expressions over sequences (e.g., ELTL [138], RCTL [16], PDL- $\Delta$  [127]), automata (e.g., BRTL [75], ECTL\* [131]), or data manipulation (e.g., extended

$\mu$ -calculi [42, 124], FORSPEC [9]). Also, extensions with both regular expressions and data handling are proposed in the state-based, linear-time setting (e.g., EAGLE [14]) and the action-based, branching-time setting (e.g., MCL [105]).

Given that we work in the compositionality framework underpinned by process algebras, we focus on action-based, branching-time temporal logics, which are suitable for LTSs and adequate w.r.t. bisimulation relations. More precisely, we use the dataless fragment of MCL (*Model Checking Language*) [105], defined in Figure 3. Action formulas (denoted by  $\alpha$ ) are built over the set of actions by using Boolean connectors in a way similar to ACTL [112], which is a slight extension w.r.t. the original definition of  $L\mu$  [82]. Derived action operators can be defined as usual:  $\mathbf{true} = \neg\mathbf{false}$ ,  $\alpha_1 \wedge \alpha_2 = \neg(\neg\alpha_1 \vee \neg\alpha_2)$ , etc. Regular formulas (denoted by  $\beta$ ) are built from action formulas by using the testing ( $\langle \rangle$ ), concatenation ( $\cdot$ ), choice ( $|$ ), and transitive reflexive closure ( $*$ ) operators. Derived regular operators can be defined as usual:  $\varepsilon = \mathbf{false}^*$  is the empty sequence operator,  $\beta^+ = \beta.\beta^*$  is the transitive closure operator, etc. State formulas (denoted by  $\varphi$ ) are built from Boolean connectors, the possibility modality ( $\langle \rangle$ ) and the infinite looping operator ( $\langle \rangle @$ ) containing regular formulas as in PDL [53] and PDL- $\Delta$  [127], and the minimal fixed point operator ( $\mu$ ) defined over propositional variables  $X$  belonging to a set  $\mathcal{X}$ . Derived state operators can be defined as usual:  $\mathbf{true} = \neg\mathbf{false}$ ,  $\varphi_1 \wedge \varphi_2 = \neg(\neg\varphi_1 \vee \neg\varphi_2)$ ,  $[\beta]\varphi = \neg\langle\beta\rangle\neg\varphi$  is the necessity modality, and  $[\beta]\dashv = \neg\langle\beta\rangle @$  is the saturation operator.  $\nu X.\varphi = \neg\mu X.\neg\varphi[\neg X/X]$  is the maximal fixed point operator ( $\varphi[\neg X/X]$  stands for  $\varphi$  in which all free occurrences of  $X$ , i.e., not bound by a fixed point operator, have been negated). State formulas are assumed to be syntactically monotonic [82], i.e., each free occurrence of a propositional variable  $X$  inside a formula  $\mu X.\varphi$  or  $\nu X.\varphi$  must be preceded by an even number of negations.

<b>Action formulas:</b>	
$\alpha ::= b$	$\llbracket b \rrbracket_A = \{b\}$
$\mathbf{false}$	$\llbracket \mathbf{false} \rrbracket_A = \emptyset$
$\neg\alpha_1$	$\llbracket \neg\alpha_1 \rrbracket_A = A \setminus \llbracket \alpha_1 \rrbracket_A$
$\alpha_1 \vee \alpha_2$	$\llbracket \alpha_1 \vee \alpha_2 \rrbracket_A = \llbracket \alpha_1 \rrbracket_A \cup \llbracket \alpha_2 \rrbracket_A$
<b>Regular formulas:</b>	
$\beta ::= \alpha$	$\llbracket \alpha \rrbracket_A = \{(s, s') \in \Sigma \times \Sigma \mid (\exists b \in A) s \xrightarrow{b} s' \wedge b \in \llbracket \alpha \rrbracket_A\}$
$\varphi?$	$\llbracket \varphi? \rrbracket_A = \{(s, s) \in \Sigma \times \Sigma \mid s \in \llbracket \varphi \rrbracket_M\}$
$\beta_1.\beta_2$	$\llbracket \beta_1.\beta_2 \rrbracket_A = \llbracket \beta_1 \rrbracket_A \circ \llbracket \beta_2 \rrbracket_A$
$\beta_1 \beta_2$	$\llbracket \beta_1 \beta_2 \rrbracket_A = \llbracket \beta_1 \rrbracket_A \cup \llbracket \beta_2 \rrbracket_A$
$\beta_1^*$	$\llbracket \beta_1^* \rrbracket_A = \llbracket \beta_1 \rrbracket_A^*$
<b>State formulas:</b>	
$\varphi ::= \mathbf{false}$	$\llbracket \mathbf{false} \rrbracket_M \rho = \emptyset$
$\neg\varphi_1$	$\llbracket \neg\varphi_1 \rrbracket_M \rho = \Sigma \setminus \llbracket \varphi_1 \rrbracket_M \rho$
$\varphi_1 \vee \varphi_2$	$\llbracket \varphi_1 \vee \varphi_2 \rrbracket_M \rho = \llbracket \varphi_1 \rrbracket_M \rho \cup \llbracket \varphi_2 \rrbracket_M \rho$
$\langle\beta\rangle\varphi_1$	$\llbracket \langle\beta\rangle\varphi_1 \rrbracket_M \rho = \{s \in \Sigma \mid (\exists s' \in \Sigma) (s, s') \in \llbracket \beta \rrbracket_A \wedge s' \in \llbracket \varphi_1 \rrbracket_M \rho\}$
$\langle\beta\rangle @$	$\llbracket \langle\beta\rangle @ \rrbracket_M \rho = \{s \in \Sigma \mid (\forall k \geq 0) (\exists s' \in \Sigma) (s, s') \in \llbracket \beta \rrbracket_A^k\}$
$X$	$\llbracket X \rrbracket_M \rho = \rho(X)$
$\mu X.\varphi_1$	$\llbracket \mu X.\varphi_1 \rrbracket_M \rho = \bigcap \{U \subseteq \Sigma \mid \llbracket \varphi_1 \rrbracket_M (\rho \circ [U/X]) \subseteq U\}$

Figure 3: Syntax and semantics of dataless MCL

The interpretation  $\llbracket \alpha \rrbracket_A$  of an action formula on the set of actions of an LTS  $M = (\Sigma, A, T, s_0)$  denotes the subset of actions satisfying  $\alpha$ . An action  $b$  satisfies a formula  $\alpha$  (also denoted by  $b \models_A \alpha$ ) if and only if  $b \in \llbracket \alpha \rrbracket_A$ . A transition  $s_1 \xrightarrow{b} s_2$  such that  $b \models_A \alpha$  is called an  $\alpha$ -transition. The interpretation  $\llbracket \beta \rrbracket_A$  of a regular formula on an LTS denotes a relation between the states that are source and target of transition sequences whose concatenated actions form a word belonging to the regular language defined by  $\beta$ . The testing operator specifies state formulas that must hold in the intermediate states of a transition sequence. A propositional context  $\rho : \mathcal{X} \rightarrow 2^\Sigma$  is a partial function mapping propositional variables to subsets of states. The notation  $\rho \circ [U/X]$  stands for a propositional context identical to  $\rho$  except for variable  $X$ , which is mapped to the state subset  $U$ . The interpretation  $\llbracket \varphi \rrbracket_M \rho$  of a state formula on an LTS  $M$  and

a propositional context  $\rho$  (which assigns a set of states to each propositional variable occurring free in  $\varphi$ ) denotes the subset of states satisfying  $\varphi$  in that context. The Boolean connectors are interpreted as usual in terms of set operations. The possibility modality  $\langle\beta\rangle\varphi_1$  (resp. the necessity modality  $[\beta]\varphi_1$ ) denotes the states for which some (resp. all) of their outgoing transition sequences satisfying  $\beta$  lead to states satisfying  $\varphi_1$ . The infinite looping operator  $\langle\beta\rangle@$  (resp. the saturation operator  $[\beta]\dashv$ ) denotes the states having some (resp. no) outgoing transition sequence consisting of an infinite concatenation of sub-sequences satisfying  $\beta$ . The minimal fixed point operator  $\mu X.\varphi_1$  (resp. the maximal fixed point operator  $\nu X.\varphi_1$ ) denotes the least (resp. greatest) solution of the equation  $X = \varphi_1$  interpreted over the complete lattice  $(2^\Sigma, \emptyset, \Sigma, \cap, \cup, \subseteq)$ . A state  $s$  satisfies a closed formula  $\varphi$ , denoted by  $s \models_M \varphi$ , if and only if  $s \in \llbracket\varphi\rrbracket_M$  (the propositional context  $\rho$  can be omitted since  $\varphi$  does not contain free variables). An LTS  $M = (\Sigma, A, T, s_0)$  satisfies a closed formula  $\varphi$ , denoted by  $M \models \varphi$ , if and only if  $s_0 \models_M \varphi$ .

Regular operators can be eliminated by translating possibility modalities and infinite looping operators into plain  $L\mu$  by applying the identities below [46] (dual identities hold for necessity modalities and saturation operators):

$$\begin{aligned} \langle\varphi'?\rangle\varphi &= \varphi' \wedge \varphi & \langle\beta_1.\beta_2\rangle\varphi &= \langle\beta_1\rangle\langle\beta_2\rangle\varphi \\ \langle\beta_1|\beta_2\rangle\varphi &= \langle\beta_1\rangle\varphi \vee \langle\beta_2\rangle\varphi & \langle\beta^*\rangle\varphi &= \mu X.(\varphi \vee \langle\beta\rangle X) \\ \langle\beta\rangle@ &= \nu X.\langle\beta\rangle X \end{aligned}$$

Although regular formulas do not increase expressiveness, they make possible a much more concise and intuitive description of properties than their fixed point counterparts. For example, the *fair reachability* [123] (i.e., by skipping cycles) of a reception *rcv* after each emission *snd* possibly followed by a finite number of transmission errors *err*, can be specified in MCL as follows:

$$[\text{true}^*.\text{snd}.((\neg\text{rcv})^*.\text{err})^*]\langle\text{true}^*.\text{rcv}\rangle\text{true}$$

whereas in plain  $L\mu$  a more verbose formula is needed:

$$\nu X.([\text{snd}]\nu Y.(\mu Z.(\langle\text{rcv}\rangle\text{true} \vee \langle\text{true}\rangle Z) \wedge \nu W.([\text{err}]Y \wedge [\neg\text{rcv}]W)) \wedge [\text{true}]X)$$

To obtain efficient model checking algorithms, i.e., linear in the size of the LTS (number of states and transitions) and the size of the formula (number of operators) we consider only alternation-free [46] fixed point formulas, i.e., without mutually recursive minimal and maximal fixed point operators. Note that infinite looping operators whose regular formulas contain star operators yield after translation fixed point formulas of alternation depth 2, such as  $\langle a^*.b \rangle@ = \nu X.\mu Y.(\langle b \rangle X \vee \langle a \rangle Y)$ , which contains two mutually recursive variables  $X$  and  $Y$  of different fixed point sign. Although the  $L\mu_2$  fragment of alternation depth 2 has in general a model checking complexity quadratic in the size of the LTS [46], the particular formulas obtained from infinite looping operators can be checked with a linear complexity, as shown in the next subsection.

## 4.2 Model checking

Given an LTS  $M = (\Sigma, A, T, s_0)$  and a closed state formula  $\varphi$ , the model checking problem consists in determining if  $M \models \varphi$ , which amounts to verify if the initial state  $s_0$  satisfies  $\varphi$ , i.e.,  $s_0 \in \llbracket\varphi\rrbracket$ . In the explicit-state setting, there are basically two approaches to solve this problem: the *global* (or enumerative) approach evaluates the interpretation of  $\varphi$  on  $M$ , i.e., the subset of states satisfying  $\varphi$ , and checks if  $s_0$  belongs to it, whereas the *local* (or on-the-fly) approach evaluates the truth value of  $\varphi$  on  $s_0$  based on the truth values of its subformulas on the descendant states of  $s_0$ . The global approach requires the entire construction of the LTS before carrying out verification, whereas the local approach explores it in a demand-driven way during verification. In practice, local model checking is suitable for the early phases of the design process, when errors are likely to occur more frequently and are detected quickly by exploring relatively small parts of the LTS; at later phases, when the model becomes stable and no errors are detected anymore, global model checking is more appropriate for verifying invariant properties since the underlying algorithms are slightly more efficient.

The alternation-free  $\mu$ -calculus fragment (consisting of formulas without mutually recursive variables of different fixed point signs) is equipped with global [38] and local [3] model checking algorithms having a

linear complexity  $O(|\varphi| \cdot (|\Sigma| + |T|))$ . For checking MCL formulas, we adopt the on-the-fly approach proposed in [3], based on Boolean Equation Systems (BESs), and we generalize it to deal efficiently with formulas of particular shapes frequently encountered in practice [102]. A BES is a tuple  $B = (X, M_1, \dots, M_n)$ , where  $X \in \mathcal{X}$  is a Boolean variable and  $M_i$  are equation blocks ( $i \in [1, n]$ ). Each block  $M_i = \{X_j \stackrel{\sigma_i}{=} op_j \mathbf{X}_j\}_{j \in [1, m_i]}$  is a set of minimal (resp. maximal) fixed point equations of sign  $\sigma_i = \mu$  (resp.  $\sigma_i = \nu$ ). The right-hand side of each equation  $j$  is a pure disjunctive or conjunctive formula obtained by applying a Boolean operator  $op_j \in \{\vee, \wedge\}$  to a set of variables  $\mathbf{X}_j \subseteq \mathcal{X}$ . The Boolean constants **F** and **T** abbreviate the empty disjunction  $\vee \emptyset$  and the empty conjunction  $\wedge \emptyset$ .

The *main* variable  $X$  must be defined in block  $M_1$ . A variable  $X_j$  depends upon a variable  $X_l$  if  $X_l \in \mathbf{X}_j$ . A block  $M_i$  depends upon a block  $M_k$  if some variable of  $M_i$  depends upon a variable defined in  $M_k$ . A block is *closed* if it does not depend upon any other blocks. A BES is *alternation-free* if there are no cyclic dependencies between its blocks; in this case, the blocks are assumed to be sorted topologically such that a block  $M_i$  only depends upon blocks  $M_k$  with  $k > i$ .

The semantics of a formula  $op_i\{X_1, \dots, X_k\}$  w.r.t.  $\mathbf{Bool} = \{\mathbf{F}, \mathbf{T}\}$  and a context  $\delta : \mathcal{X} \rightarrow \mathbf{Bool}$ , which must initialize all variables  $X_1, \dots, X_k$ , is the boolean value defined as follows:

$$\llbracket op_i\{X_1, \dots, X_k\} \rrbracket \delta = \delta(X_1) op_i \dots op_i \delta(X_k)$$

The semantics of an equation block  $M_i = \{X_j \stackrel{\sigma_i}{=} op_j \mathbf{X}_j\}_{j \in [1, m_i]}$  w.r.t. a context  $\delta$  is the  $\sigma_i$ -fixed point of a vectorial functional  $\Phi_{i\delta} : \mathbf{Bool}^{m_i} \rightarrow \mathbf{Bool}^{m_i}$ :

$$\left[ \left[ \{X_j \stackrel{\sigma_i}{=} op_j \mathbf{X}_j\}_{j \in [1, m_i]} \right] \right] \delta = \sigma_i \Phi_{i\delta}$$

where

$$\Phi_{i\delta}(b_1, \dots, b_{m_i}) = (\llbracket op_j \mathbf{X}_j \rrbracket (\delta \circ [b_1/X_1, \dots, b_{m_i}/X_{m_i}]))_{j \in [1, m_i]}$$

The notation  $\delta \circ [b_1/X_1, \dots, b_n/X_n]$  stands for a context identical to  $\delta$  except for variables  $X_1, \dots, X_n$ , which are assigned values  $b_1, \dots, b_n$ , respectively.

The semantics of an alternation-free BES is the value of its main variable  $X$  given by the solution of  $M_1$ , i.e.,  $\delta_1(X)$ , where the contexts  $\delta_i$  are calculated as follows:

$$\begin{aligned} \delta_n &= \llbracket M_n \rrbracket \square \\ \delta_i &= (\llbracket M_i \rrbracket \delta_{i+1}) \circ \delta_{i+1} \text{ for } i \in [1, n-1] \end{aligned}$$

Note that the context for interpreting  $M_n$  is empty (since  $M_n$  is closed) and a block  $M_i$  is interpreted in the context of all blocks  $M_k$  with  $k > i$ .

A block  $M_i$  is *acyclic* if there are no cyclic dependencies between the variables defined in  $M_i$ . A variable  $X_j$  is called *disjunctive* (resp. *conjunctive*) if  $op_j = \vee$  (resp.  $op_j = \wedge$ ). A block  $M_i$  is disjunctive (resp. conjunctive) if each of its variables either is disjunctive (resp. conjunctive), or it depends upon at most one variable defined in  $M_i$ , all its other dependencies being constants or variables defined in other blocks.

The on-the-fly resolution of an alternation-free BES  $B = (X, M_1, \dots, M_n)$  consists in computing the value of  $X$  by exploring the right-hand sides of the equations in a demand-driven way, without explicitly constructing the blocks. To each block  $M_i$  is associated a resolution routine  $R_i$  responsible for computing the values of the variables defined in  $M_i$ . When a variable  $X_j$  defined in  $M_i$  is computed by a call  $R_i(X_j)$ , the values of other variables  $X_l$  defined in other blocks  $M_k$  may be needed; these values are computed by calls  $R_k(X_l)$  of the routine associated to  $M_k$ . This process always terminates because there are no cyclic dependencies between blocks (the call stack of resolution routines has a size bounded by the depth of the dependency graph between blocks). Since a variable  $X_j$  defined in  $M_i$  may be required several times during the resolution process, the computation results must be kept persistent between subsequent calls of  $R_i$  in order to obtain a linear-time overall resolution. Compared to other on-the-fly resolution algorithms like LMC [43], which consists of a single monolithic routine handling the whole BES, the above scheme yields simpler resolution algorithms (dedicated to blocks of equations having the same fixed point sign), and is easier to optimize (by designing more efficient algorithms for blocks with particular structure) [102].

The on-the-fly model checking procedure for MCL formulas is described in detail in [104, 105]. We briefly illustrate it here for the dataless MCL fragment using the example on Figure 4. The LTS shown on

Figure 4(a) models the behaviour of a two-place buffer made by connecting two one-place cells in sequence. States are represented by numbers, the initial state being 0. The buffer accepts as input an infinite stream containing two kinds of messages, denoted by 0 or 1 (actions  $put_0$  and  $put_1$ ) and delivers it as output (actions  $get_0$  and  $get_1$ ) in the same order. The move of a message from the first to the second cell of the buffer is modeled by a  $\tau$ -transition.

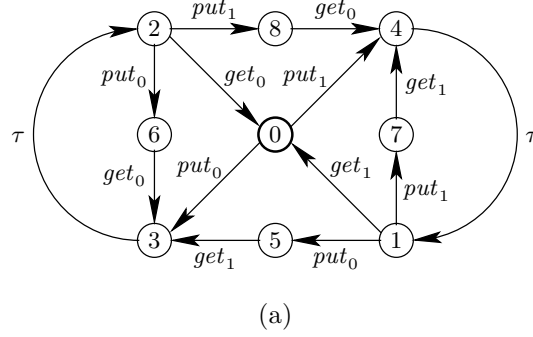
On this LTS, we want to check that after every message 0 accepted by the buffer, it is possible that the buffer cyclically accepts a message 1 and delivers it. This property is expressed by the MCL formula on top of Figure 4(b), which contains a PDL box modality followed by an infinite looping operator of alternation depth 2. The subsequent lines in Figure 4(b) illustrate the various translation steps that bring this formula into an equational form suitable for evaluation using BESs: (i) the formula is first translated into PDL with recursion [104], which consists of two blocks of fixed point equations defining propositional variables using PDL formulas; (ii) the PDL modalities in the equations are split to eliminate concatenation operators in the regular formulas, and the equation block corresponding to the infinite looping operator is transformed specifically by adding an extra equation and flipping its fixed point sign; (iii) the star operators in the regular formulas are replaced by fixed point equations, yielding a specification in HML with recursion [92], which contains only single-step modal formulas; (iv) the interpretation of this specification on the LTS is encoded as a BES made of two corresponding equation blocks  $M_1$  and  $M_2$  defining Boolean variables of the form  $X_s$  (resp.  $Y_s, Z_s, W_s$ ), which are true if and only if the state  $s$  satisfies the propositional variable  $X$  (resp.  $Y, Z, W$ ). In this way, the local model checking of the formula on the initial state of the LTS amounts to solve the variable  $X_0$  of block  $M_1$ .

This evaluation is carried out by using local resolution algorithms that explore the Boolean graph [3] associated to the BES in a forward way, as shown by the excerpt on Figure 4(c). Since the block  $M_1$  is conjunctive and  $M_2$  is disjunctive, they are solved in a memory-efficient manner by the algorithms proposed in [102] and [105], respectively. These algorithms, based on a depth-first search of the Boolean graph, store only Boolean variables and not dependencies between them, and hence, only the states and not the transitions of the LTS. When exploring the Boolean graph of  $M_2$ , each cycle containing a variable  $Y_s$  is interpreted as a constant **true**, since it corresponds to a sequence satisfying the infinite looping operator in the initial formula. This is done in linear-time by computing the strongly connected components in the Boolean graph of  $M_2$  and keeping track of those containing variables  $Y_s$ , which are similar to the marked states in Büchi automata. The thick arrows in Figure 4(c) define the diagnostic of the BES resolution, which is a minimal Boolean subgraph (w.r.t. graph inclusion) containing the variable of interest and whose variables have the same solution as those of the BES [101]. On this example, the LTS satisfies the MCL formula, and the positive diagnostic (witness), illustrated partially on Figure 4(c), contains an execution sequence leading to every  $put_0$ -transition in the LTS, followed by a cycle passing through  $put_1$ - and  $get_1$ -transitions.

### 4.3 Property-dependent reductions

A property-based specification of a system consists usually of several properties expressed as temporal logic formulas, each one describing a certain aspect of the system. This kind of specification has naturally two benefits [96]: it is *abstract*, in the sense that it may characterize a family of systems sharing the same observable behaviour (e.g., the abstract properties of mutual exclusion and inevitable progress towards the critical section must be satisfied by all mutual exclusion protocols, regardless their implementation details); and it is *modular*, meaning that one can add, eliminate, or replace a property in the specification without changing the other properties and their interpretation on the system under study.

In practice, since the model checking complexity grows with the size of the formula (number of operators), to increase efficiency it is desirable to verify formulas as small as possible. In our framework, a given dataless MCL formula  $\varphi$  should be first decomposed into a set of smaller “temporal” formulas (i.e., dominated by a modal or fixed point operator), each one being checked separately on the system. A simple way to achieve this is by transforming the formula in conjunctive normal form (CNF), by considering as literals the maximal temporal subformulas of  $\varphi$ ; then, for each conjunct, each of its disjuncts is verified in turn (by stopping the process when some disjunct is satisfied by the system), the whole process terminating when all conjuncts have been checked to hold on the system, or some conjunct not satisfied by the system is encountered. This decomposition and evaluation process can be easily automated. However, in practice a property-based



MCL formula:

$$[\text{true}^*.put_0] \langle \text{true}^*.put_1.\text{true}^*.get_1 \rangle @$$

Translation to PDL with recursion:

$$\begin{cases} X =_\nu [\text{true}^*.put_0] Y \\ Y =_\nu \langle \text{true}^*.put_1.\text{true}^*.get_1 \rangle Y \end{cases}$$

Conversion of infinite looping block:

$$\begin{cases} X =_\nu [\text{true}^*][put_0] Y \\ Y =_\mu Z \\ Z =_\mu \langle \text{true}^* \rangle \langle put_1 \rangle \langle \text{true}^* \rangle \langle get_1 \rangle Y \end{cases}$$

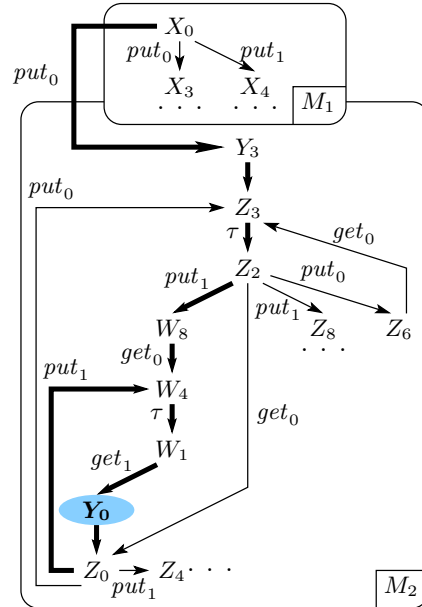
Translation to HML with recursion:

$$\begin{cases} X =_\nu [put_0] Y \wedge [\text{true}] X \\ Y =_\mu Z \\ Z =_\mu \langle put_1 \rangle W \vee \langle \text{true} \rangle Z \\ W =_\mu \langle get_1 \rangle Y \vee \langle \text{true} \rangle W \end{cases}$$

Model checking encoding as BES:

$$\begin{cases} X_s =_\nu \bigwedge_{s \xrightarrow{put_0} s'} Y_{s'} \wedge \bigwedge_{s \rightarrow s''} X_{s''} \\ Y_s =_\mu Z_s \\ Z_s =_\mu \bigvee_{s \xrightarrow{put_1} s'} W_{s'} \vee \bigvee_{s \rightarrow s''} Z_{s''} \\ W_s =_\mu \bigvee_{s \xrightarrow{get_1} s'} Y_{s'} \vee \bigvee_{s \rightarrow s''} W_{s''} \end{cases}_{s \in S}$$

(b)



(c)

Figure 4: Example of on-the-fly model checking using BESs. (a) LTS of a two-place buffer accepting and delivering messages of two kinds, denoted by 0 and 1. (b) MCL formula and the translation of its evaluation on the LTS in terms of a BES resolution. (c) Local resolution of the resulting BES by forward exploration of its Boolean graph (excerpt).

specification is often a list of temporal formulas that are supposed to hold on the system (which corresponds to a CNF formula with monome conjuncts), and this is the case we focus on here, by considering the verification of a single temporal formula.

Given an LTS and a temporal formula, the performance of verification can be improved by reducing the LTS before checking the formula, modulo a relation that must preserve the truth value of the formula. The standard approach is to reduce the LTS modulo an equivalence relation adequate w.r.t. the temporal logic to which belongs the formula. Of course, to improve the overall performance of the verification process (LTS generation, reduction, and model checking), the computational complexity of the LTS reduction should be smaller than that of checking the formula on the initial LTS; this can be achieved by using the compositional LTS construction and minimization described in Section 3. This approach relies on existing adequacy results between behavioural equivalences and temporal logics (see, e.g., [48] for a survey of adequacy in the action-based setting). In our framework, a useful result is the adequacy of ACTL-X (Action-based CTL without the next state operator) [112] w.r.t. divergence-sensitive branching bisimulation. This result also holds for  $\mu$ ACTL-X [47], which extends ACTL-X with fixed point operators.

Nevertheless, the reduction of the LTS modulo an equivalence relation preserves *all* temporal formulas adequate w.r.t. that relation, whereas in practice one is interested in preserving only the properties forming the specification of the system. A better solution would be to specialize the reduction for each temporal formula considered: given that each formula references specific aspects of the system, the other aspects (referenced by the other properties in the specification) could be abstracted away when checking that formula, thus yielding more potential for reduction. An early attempt for property-dependent reduction in the action-based setting was made with the selective  $\mu$ -calculus [13], which synthesizes an equivalence relation preserving a given formula. However, this attempt has two practical limitations: it imposes a particular formulation of properties (using special modalities indexed by action sets), and the synthesized equivalence relations are variants of the  $\tau^*.a$  equivalence (which is not a congruence w.r.t. parallel composition and hence not appropriate for compositional reasoning, and it preserves only safety and weak liveness properties).

Another, more recent, attempt for property-dependent reduction [106] takes a modal  $\mu$ -calculus formula and synthesizes the maximal set of actions which can be hidden (i.e., renamed into  $\tau$ ) in the LTS without disturbing the interpretation of the formula. This approach keeps in the LTS only the relevant information determining the truth value of the formula, therefore yielding a high potential for reduction. After applying this maximal hiding, the LTS can be reduced modulo an adequate equivalence relation before checking the formula.

**Definition 16 (Hiding set)** *Let  $\alpha$  be an action formula interpreted over a set of actions  $A$ . The hiding set of  $\alpha$  w.r.t.  $A$  is defined as follows:*

$$h_A(\alpha) = \begin{cases} \llbracket \alpha \rrbracket_A & \text{if } \tau \models_A \alpha \\ A \setminus \llbracket \alpha \rrbracket_A & \text{if } \tau \not\models_A \alpha \end{cases}$$

*The hiding set of a state formula  $\varphi$  w.r.t.  $A$ , denoted by  $h_A(\varphi)$ , is defined as the intersection of  $h_A(\alpha)$  for all action subformulas  $\alpha$  of  $\varphi$ .*

**Definition 17 (Hiding)** *Let  $A$  be a set of actions and  $H \subseteq A$ . The hiding of an action  $b \in A$  w.r.t.  $H$  is defined as follows:*

$$\text{hide}_H(b) = \begin{cases} b & \text{if } b \notin H \\ \tau & \text{if } b \in H \end{cases}$$

*The hiding of an LTS  $M$  w.r.t.  $H$  is defined as follows:*

$$\text{hide}_H(M) = \text{lts}(\mathbf{hide } H \text{ in } M)$$

The following lemma [106] states that, given an action formula  $\alpha$ , the fact of hiding an action w.r.t. the hiding set of  $\alpha$  does not disturb the interpretation of  $\alpha$  on that action.

**Lemma 1** *Let  $\alpha$  be an action formula interpreted over a set of actions  $A$ . A subset  $H \subseteq A$  does not disturb the interpretation of  $\alpha$  on the actions of  $A$  after hiding them w.r.t.  $H$  if the following property holds:*

$$\forall b \in A. (b \models_A \alpha \Leftrightarrow \text{hide}_H(b) \models_A \alpha)$$

*Then: a subset  $H \subseteq A$  satisfies this property if and only if  $H \subseteq h_A(\alpha)$ .*



Lemma 1 ensures that, for an action formula  $\alpha$ , its hiding set  $h_A(\alpha)$  is the maximal set of actions that can be hidden in the LTS without disturbing the interpretation of  $\alpha$ . To make possible LTS reductions prior to (or simultaneously with) the verification of a state formula  $\varphi$ , it is desirable to hide as many actions as possible in the LTS, i.e., all actions in  $h_A(\varphi)$ . The following proposition [106], which makes use of Lemma 1, guarantees that this hiding preserves the interpretation of  $\varphi$ .

**Proposition 8 (Maximal Hiding)** *Let  $M = (\Sigma, A, \longrightarrow, s_0)$  be an LTS,  $\varphi$  be a state formula of  $L\mu$ , and  $H \subseteq h_A(\varphi)$ . Then:*

$$\llbracket \varphi \rrbracket_M \rho = \llbracket \varphi \rrbracket_{\text{hide}_H(M)} \rho$$

for any propositional context  $\rho$ .

For instance, consider the MCL formula below, expressing the inevitable execution of a *recv* action after every *send* action:

$$\varphi = [\text{true}^*. \text{send}] \mu X. ((\text{true}) \text{true} \wedge [\neg \text{recv}] X)$$

When checking  $\varphi$  on an LTS, one can hide all actions in  $h_A(\varphi) = h_A(\text{send}) \cap h_A(\neg \text{recv}) = (A \setminus \llbracket \text{send} \rrbracket_A) \cap \llbracket \neg \text{recv} \rrbracket_A = (A \setminus \{\text{send}\}) \cap (A \setminus \{\text{recv}\}) = A \setminus \{\text{send}, \text{recv}\}$ , i.e., all actions other than *send* and *recv*, without changing the interpretation of the formula.

For a given formula  $\varphi$ , after hiding all actions in  $h_A(\varphi)$ , one can improve the performance of verification by reducing the LTS modulo an equivalence relation that preserves the interpretation of  $\varphi$ . Two such relations are considered in [106]: strong bisimulation, which preserves all  $L\mu$  formulas, and divergence-sensitive branching bisimulation (divbranching for short), which is adequate w.r.t.  $\mu\text{ACTL-X}$  [47] and also with the  $L_\mu^{\text{dsbr}}$  fragment of  $\mu$ -calculus proposed in [106]. Strong bisimulation minimization, although it has a lower potential for reduction compared to weak equivalences (since strong bisimulation does not give a special treatment to  $\tau$ -transitions), can nevertheless improve the performance of verification when the LTS obtained after hiding contains a high percentage of  $\tau$ -transitions (execution time divided by 4 and memory consumption divided by 2 [106]). Regarding divbranching bisimulation, the performances can be improved significantly either by minimization (time divided by 20 and memory divided by 5), or by on-the-fly  $\tau$ -confluence reduction (time divided by 10) [106].

More generally, maximal hiding for a given formula can be applied in conjunction with the compositional LTS construction and minimization approach described in Section 3. This specialization is likely to yield better results (especially w.r.t. the peak memory consumption of the whole verification process) than the compositional reduction modulo an equivalence relation, as shown in Section 7.

## 4.4 Partial model checking

*Partial model checking* is a dual approach to compositional model generation, where one needs to check a formula on a network of automata. Partial model checking was first proposed by Andersen [8, 4, 5] for concurrent components running asynchronously and composed using CCS parallel composition and restriction operators. For a modal  $\mu$ -calculus formula  $\varphi$  and a composition expression  $S_1 \parallel \dots \parallel S_n$ , Andersen uses an operation  $\varphi \parallel S_1$  called *quotienting* of the formula  $\varphi$  w.r.t. the component  $S_1$ , so that  $S_1 \parallel \dots \parallel S_n$  satisfies  $\varphi$  if and only if the smaller composition  $S_2 \parallel \dots \parallel S_n$  satisfies  $\varphi \parallel S_1$ . In addition, simplifications can (and must) be applied to  $\varphi \parallel S_1$  to reduce its size. Partial model checking is the incremental application of quotienting and simplifications, so that state explosion is avoided if the size of intermediate formulas can be kept sufficiently small.

Partial model checking has been adapted and used successfully in various contexts, such as state-based models [7, 6], synchronous state/event systems [22], and timed systems [17, 27, 90, 91, 93]. It has also been specialised for security properties [99]. More recently, it has been generalised to the full CCS process algebra, with an application to the verification of parameterized systems [15]. These various developments of partial model checking, although successful, were relatively scarce, which may be explained by the complexity of the method: obtaining a fully operational partial model checker requires a significant implementation effort and extensive experiments for fine-tuning and optimization.

In this section, we focus on partial model checking of the modal  $\mu$ -calculus applied to (untimed) concurrent asynchronous components [87, 88]. By considering only binary associative parallel composition operators

(such as CCS and CSP parallel compositions), previous works [4, 5, 15] are not directly applicable to more general operators, such as  $m$ -among- $n$  synchronization and parallel composition by synchronization interfaces (where all components containing a given action in their synchronization interface must synchronize on that action) [61], present in the E-LOTOS standard and variants [28, 80]. Our first contribution in this section is thus a generalisation of partial model checking to networks of LTSs, which subsumes all the above-mentioned parallel composition operators. Regarding the communication of data values, our approach is applicable to classical (i.e., with static communication) value-passing process algebras equipped with early operational semantics. This framework encompasses a significant fragment of the  $\pi$ -calculus (containing channel mobility and bounded process creation), which can be translated into classical value-passing process algebras [103].

In realistic cases, partial model checking handles huge formulas and components, thus requiring efficient implementations. Our second contribution is a reformulation of quotienting as a synchronous product (which can itself be represented in the network model) between a graph representing the formula (called a *formula graph*) and the behaviour graph of a component, thus enabling efficient implementation using existing tools dedicated to graph manipulations. Our third contribution is the reformulation of formula simplifications as a combination of graph reductions (including minimisations modulo equivalence relations and bisimulations) and partial evaluation of the formula graph using a BES (*Boolean Equation System*) [3].

In this section, we consider plain  $L\mu$  formulas in disjunctive form, i.e., expressed using negation  $\neg$ , variables, and the disjunctive operators  $\text{false}$ ,  $\langle \_ \rangle$ ,  $\vee$ , and  $\mu$ . We also suppose that the formulas are *well-formed block-labelled*, as defined below.

**Definition 18 (Block-labelled formula)** A block-labelled  $\mu$ -calculus formula  $\varphi$  is a formula in which each propositional variable  $X$  is labelled by a natural number  $k$ , called its block number.

Given a block-labelled  $\mu$ -calculus formula  $\varphi$ , we write  $\text{fv}(\varphi)$  for the set of variables free in  $\varphi$ , and  $\text{bv}(\varphi)$  for the set of variables bound in  $\varphi$ . We call a closed formula any formula  $\varphi$  such that  $\text{fv}(\varphi) = \emptyset$ . We assume that all bound variables have distinct names, and for  $X^k \in \text{bv}(\varphi)$ , we write  $\varphi[X^k]$  for the (unique) subformula of  $\varphi$  of the form  $\mu X^k.\varphi_0$ . Given  $\varphi_1$  and  $\varphi_2$ , we write  $\varphi_1[\varphi_2/X^k]$  for substituting all free occurrences of  $X^k$  in  $\varphi_1$  by  $\varphi_2$  (while implicitly applying  $\alpha$ -conversion<sup>3</sup> to maintain the unicity of bound variables).

Intuitively, a block-labelling is well-formed if the  $\mu$ -calculus formula can be converted into an equivalent set of  $\mu$ -calculus equations partitioned into blocks, so that all variables having the same block number are defined in the same block and if  $k < k'$  then the equations within block number  $k$  occur before the equations within block number  $k'$ . The well-formedness conditions are the following:

1. All occurrences of a given variable are labelled by the same block number.
2. All variables sharing the same block number have the same fixed point sign<sup>4</sup>.
3. For all  $X^k \in \text{bv}(\varphi)$  and  $Y^{k'} \in \text{fv}(\varphi[X^k])$  it holds that  $k' \leq k$ .

By convention, we assume without loss of generality that the even block numbers are associated to variables of sign  $\mu$  and odd block numbers are associated to variables of sign  $\nu$ .

Initially, every unlabelled formula  $\varphi$  can be turned into the well-formed block-labelled formula  $\text{bl}(\varphi, \text{true}, 0, [])$ , where  $\text{bl}(\psi, b, k, \gamma)$  is defined as follows,  $b$  denoting a boolean that is true only if  $\psi$  is in the context of an even number of negations,  $k$  the current block number, and  $\gamma$  a mapping from variables to block numbers:

$$\begin{aligned}
\text{bl}(\text{false}, b, k, \gamma) &= \text{false} \\
\text{bl}(X, b, k, \gamma) &= X^{\gamma(X)} \\
\text{bl}(\neg\varphi_0, b, k, \gamma) &= \neg\text{bl}(\varphi_0, \neg b, k, \gamma) \\
\text{bl}(\varphi_1 \vee \varphi_2, b, k, \gamma) &= \text{bl}(\varphi_1, b, k, \gamma) \vee \text{bl}(\varphi_2, b, k, \gamma) \\
\text{bl}(\langle a \rangle \varphi_0, b, k, \gamma) &= \langle a \rangle \text{bl}(\varphi_0, b, k, \gamma) \\
\text{bl}(\mu X.\varphi_0, b, k, \gamma) &= \begin{cases} \mu X^k.\text{bl}(\varphi_0, \text{true}, k, \gamma \odot [X \mapsto k]) & \text{if } b = \text{true} \\ \mu X^{k+1}.\text{bl}(\varphi_0, \text{true}, k+1, \gamma \odot [X \mapsto k+1]) & \text{otherwise} \end{cases}
\end{aligned}$$

<sup>3</sup> $\alpha$ -conversion is a well-known equivalence relation between terms that differ only in variable names, originally defined in the framework of the  $\lambda$ -calculus and applied implicitly during substitution to avoid variable capture.

<sup>4</sup>Considering a formula  $\varphi$  in disjunctive form, the sign of variable  $X^k$  is  $\mu$  if  $\varphi[X^k]$  occurs in  $\varphi$  in the context of an even number of negations, and  $\nu$  otherwise.

We write  $\text{blocks}(\varphi)$  for the set of block numbers occurring in  $\varphi$ . A block-labelled formula  $\varphi$  is alternation-free if  $k' = k$  for all  $X^k \in \text{bv}(\varphi)$  and  $Y^{k'} \in \text{fv}(\varphi[X^k])$ .

To check a closed formula  $\varphi$  on a network  $N = (\vec{S}, V)$ , one can choose a component LTS  $\vec{S}[i]$ , compute the quotient of the formula  $\varphi$  with respect to  $\vec{S}[i]$ , and check the resulting quotient formula on the smaller (at least in the number of component LTSs, but also hopefully in the compound LTS size) network  $N_{\overline{\{i\}}}$ .

**Definition 19 (Quotient formula)**  $N = (\vec{S}, V)$  being a network of size  $n$ , we assume a function  $\alpha(\vec{t}, a)$  that assigns a unique unused label to each  $(\vec{t}, a) \in V$ . Given  $\varphi$  a closed formula and  $i \in [1, n]$ , the quotient formula is written  $\varphi \parallel_i^0 s_i^0$  and defined as follows:

$$\begin{aligned} \text{false} \parallel_i^B s &= \text{false} \\ X^k \parallel_i^B s &= \varphi[X^k] \parallel_i^B s \\ (\neg\varphi_0) \parallel_i^B s &= \neg(\varphi_0 \parallel_i^B s) \\ (\varphi_1 \vee \varphi_2) \parallel_i^B s &= (\varphi_1 \parallel_i^B s) \vee (\varphi_2 \parallel_i^B s) \\ (\mu X^k . \varphi_0) \parallel_i^B s &= \begin{cases} X_s^k & \text{if } X_s^k \in B \\ \mu X_s^k . (\varphi_0 \parallel_i^{B \cup \{X_s^k\}} s) & \text{otherwise} \end{cases} \\ \langle a \rangle \varphi_0 \parallel_i^B s &= \bigvee_{(\vec{t}, a) \in V} \left( \begin{array}{l} (i \notin \text{acv}(\vec{t}) \wedge \langle a \rangle (\varphi_0 \parallel_i^B s)) \vee \\ (\{i\} \subsetneq \text{acv}(\vec{t}) \wedge \bigvee_{s \xrightarrow{\vec{t}[i]}_i s'} \langle \alpha(\vec{t}, a) \rangle (\varphi_0 \parallel_i^B s')) \vee \\ (\{i\} = \text{acv}(\vec{t}) \wedge \bigvee_{s \xrightarrow{\vec{t}[i]}_i s'} (\varphi_0 \parallel_i^B s')) \end{array} \right) \end{aligned}$$

This definition follows and generalises Andersen's [4] (specialised for CCS) to networks. The main difference is the definition of  $\langle a \rangle \varphi_0 \parallel_i^B s$ , CCS composition corresponding to vectors  $((a, \bullet), a)$ ,  $((\bullet, a), a)$ , or  $((a, \bar{a}), \tau)$ ,  $a$  and  $\bar{a}$  being an action and its CCS *co-action*, making the use of special labels  $\alpha(\vec{t}, a)$  not necessary. A minor difference is that we use  $\mu$ -calculus terms instead of equations<sup>5</sup>. Any sub-formula produced by quotienting has the same block number as the original sub-formula, reflecting the order of equation blocks in Andersen's definition. The set  $B$  keeps track of new variables already introduced in the quotient formula. Quotienting is well-defined, because formulas are finite, every  $\varphi[X^k]$  has the form  $\mu X^k . \varphi_0$  (the formula is in disjunctive form), and the size of the set  $B$  is bounded by  $|\text{bv}(\varphi)| \times |\Sigma_i|$ . Note that well-formedness of the block-labelling is preserved by quotienting, because for every variable  $X_s^k \in \text{bv}(\varphi \parallel_i^0 s_0)$  we have  $X^k \in \text{bv}(\varphi)$  and for every variable  $Y_{s'}^{k'} \in \text{fv}((\varphi \parallel_i^0 s_0)[X_s^k])$  we have  $Y^{k'} \in \text{fv}(\varphi[X^k])$ , and therefore  $k' \leq k$ .

**Example 8** The  $\mu$ -calculus formula  $\mu X^0 . \langle a \rangle \text{true} \vee \langle b \rangle X^0$  (existence of a path of zero or more  $b$  leading to an  $a$ ) can be rewritten to disjunctive form as  $\mu X^0 . \langle a \rangle \neg \text{false} \vee \langle b \rangle X^0$ . Quotienting of this formula with respect to  $S_3$  in the network  $N$  introduced in Example 1 yields the formula  $\mu X_0^0 . \langle a \rangle \neg \text{false} \vee \langle \alpha_a \rangle \neg \text{false} \vee \langle \alpha_b \rangle \mu X_2^0 . \langle a \rangle \neg \text{false} \vee \text{false}$ . In other words, an action  $a$  can be reached after a (possibly empty) sequence of  $b$  actions in the network  $N$  if and only if an action  $a$ , or an action  $\alpha_a$ , or an action  $\alpha_b$  followed by an action  $a$ , can be reached immediately in  $N_{\overline{\{3\}}}$ , given the behaviour of  $S_3$  depicted in Figure 1.

Interestingly, quotienting can be implemented as a network that realises a product between an LTS encoding the formula (called a *formula graph*) and a component LTS of the network under verification. In practice, the EXP.OPEN tool of CADP can be used for that purpose.

We now introduce the notion of *formula graph*, and then we define the network that implements quotienting.

**Definition 20 (Circuit)** Let  $S = (\Sigma, A, \longrightarrow, s_0)$  be an LTS and  $T \subseteq \longrightarrow$  be a subset of its transitions. The states of  $T$  are defined as the set  $\text{st}(T) = \{s, s' \in \Sigma \mid (s, a, s') \in T\}$ .  $T$  is a circuit of  $S$  if for all  $s, s' \in \text{st}(T)$  there is a sequence of transitions belonging to  $T$  from  $s$  to  $s'$ . A state  $s \in \text{st}(T)$  is a root of the circuit  $T$  if there is a sequence of transitions from  $s_0$  to  $s$  that does not traverse any transition of  $T$ .

<sup>5</sup>Note that terms will be compiled into graphs, thus enabling the sharing of sub-formulas that is also possible using equations.

**Definition 21 (Formula graph)** A formula graph is an LTS  $(\Sigma, A, \longrightarrow, s_0)$  such that:

1. Every label  $\sigma \in A$  has either form  $\vee, \neg, \langle a \rangle$  (for some  $a$  belonging to a fixed set of action names), or  $\mu^k$  (for some  $k \in \mathbb{N}$ ). To avoid confusion, in the sequel, we will use  $\sigma$  to denote the label of a formula graph and  $a$  to denote a label of the network on which the formula is to be checked.
2. If  $s_0 \xrightarrow{\delta} s \xrightarrow{\mu^k} s'$  for some  $\delta \in A^*$  and  $k \in \mathbb{N}$ , then  $k$  is even if and only if  $\delta$  contains an even number of occurrences of the label  $\neg$ .
3. If  $s \in \Sigma$  is a root of a circuit then (a) the circuit contains a  $\mu^k$ -transition and (b) if the first  $\mu^k$ -transition traversed on the circuit starting in  $s$  has block number  $k'$  then every  $\mu^k$ -transition belonging to the circuit satisfies  $k \geq k'$ .

Every formula graph can be decoded into a closed formula as follows.

**Definition 22 (Decoding a formula graph)** A formula graph  $P = (\Sigma, A, \longrightarrow, s_0)$  encodes the modal  $\mu$ -calculus formula  $\text{dec}_s(P, s_0, \emptyset)$ , where  $\text{dec}_s(P, s, E)$  is defined as follows ( $E \subseteq \Sigma$ ). In our decoding every variable is uniquely identified by the source state  $s$  and the block number  $k$  of the  $\mu$ -transition, and written  $\bar{s}^k$ .

$$\text{dec}_s(P, s, E) = \bigvee_{s \xrightarrow{\sigma} s' \in P} \text{dec}_t(P, s \xrightarrow{\sigma} s', E)$$

where

$$\begin{aligned} \text{dec}_t(P, s \xrightarrow{\vee} s', E) &= \text{dec}_s(P, s', E) \\ \text{dec}_t(P, s \xrightarrow{\neg} s', E) &= \neg \text{dec}_s(P, s', E) \\ \text{dec}_t(P, s \xrightarrow{\langle a \rangle} s', E) &= \langle a \rangle \text{dec}_s(P, s', E) \\ \text{dec}_t(P, s \xrightarrow{\mu^k} s', E) &= \begin{cases} \bar{s}^k & \text{if } s \in E \\ \mu \bar{s}^k . \text{dec}_s(P, s', E \cup \{s\}) & \text{otherwise} \end{cases} \end{aligned}$$

This definition implies that a deadlock state decodes as **false** (empty disjunction). Function  $\text{dec}_s$  is well-defined. In particular, it terminates because every cyclic path contains a label of the form  $\mu^k$ . By recording in the set  $E$  the source states of traversed  $\mu^k$ -transitions, we thus avoid infinite traversals of cycles. In practice, formula graphs need not be decoded.

**Definition 23 (Encoding a formula into a formula graph)** The formula graph corresponding to a well-formed block-labelled formula  $\varphi$  in disjunctive form is an LTS written  $\text{enc}(\varphi)$ , whose states are identified with sub-formulas of  $\varphi$ . The initial state of the formula graph is  $\varphi$ , **false** is a deadlock state, and each sub-formula has transitions as follows:

$$\begin{array}{lll} X^k \xrightarrow{\vee} \varphi[X^k] & \neg \varphi_0 \xrightarrow{\neg} \varphi_0 & \langle a \rangle \varphi_0 \xrightarrow{\langle a \rangle} \varphi_0 \\ \varphi_1 \vee \varphi_2 \xrightarrow{\vee} \varphi_1 & \varphi_1 \vee \varphi_2 \xrightarrow{\vee} \varphi_2 & \mu X^k . \varphi_0 \xrightarrow{\mu^k} \varphi_0 \end{array}$$

Although the states of a formula graph are identified by formulas, only the transition labels are required for decoding. In figures, states will be simply identified by numbers.

Note that the formula graph obtained by encoding a formula satisfies the conditions given in Definition 21. Condition (2) is a direct consequence of the block-labelling convention stated in Definition 18. Condition (3) comes from the fact that the roots of the circuits are the states associated to formulas of the form  $\mu X^k . \psi$  such that  $X^k$  occurs free in  $\psi$ . In particular, subcondition (b) is a consequence on the third well-formedness condition given in Definition 18.

**Example 9** The formula graph corresponding to the formula  $\mu X^0 . (\langle a \rangle \text{true}) \vee \langle b \rangle X^0$  introduced in Example 8 is depicted in Figure 5 (a).

Using this encoding, the quotient of a formula with respect to the  $i$ th LTS of a network can be computed as a synchronous product using a network called *quotient formula network*.

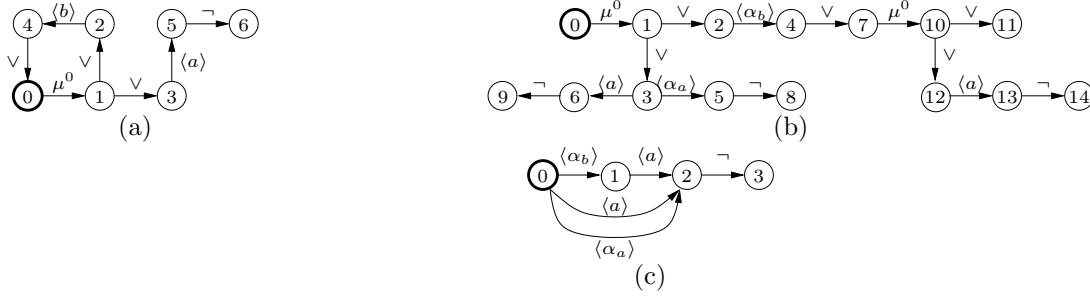


Figure 5: Examples of formula graphs

**Definition 24 (Quotient formula network)** Let  $\varphi$  be a modal  $\mu$ -calculus formula in disjunctive form,  $N = (\vec{S}, V)$  be a network of size  $n$ , and  $i \in [1, n]$ . The quotient formula network of  $\varphi$  with respect to  $\vec{S}[i]$  is defined as the network  $((\text{enc}(\varphi), \vec{S}[i]), V//_i)$ , where  $V//_i$  denotes the following set of rules:

$$\begin{array}{l}
 \{ ((\sigma, \bullet), \sigma) \mid \sigma \in \{\neg, \vee\} \cup \{\mu^k \mid k \in \text{blocks}(\varphi)\} \} \cup \\
 \{ ((\langle a \rangle, \bullet), \langle a \rangle) \mid (\vec{t}, a) \in V \wedge i \notin \text{acv}(\vec{t}) \} \cup \\
 \{ ((\langle a \rangle, \vec{t}[i]), \langle \alpha(\vec{t}, a) \rangle) \mid (\vec{t}, a) \in V \wedge \{i\} \subsetneq \text{acv}(\vec{t}) \} \cup \\
 \{ ((\langle a \rangle, \vec{t}[i]), \vee) \mid (\vec{t}, a) \in V \wedge \{i\} = \text{acv}(\vec{t}) \}
 \end{array}$$

Note that the LTS corresponding to the quotient formula network is a formula graph. We have proven [88] that our encoding of closed formulas into formula graphs is sound, so that the formula can be recovered from the formula graph into which the formula is encoded (Proposition 9 below), and that the LTS corresponding to the quotient formula network indeed encodes the quotient correctly (Proposition 10 below).

**Proposition 9 (Soundness of formula encoding)** If  $\varphi$  is a closed formula in disjunctive form, then  $\text{dec}_s(\text{enc}(\varphi), \varphi, \emptyset)$  is equivalent to  $\varphi$ .

**Proposition 10 (Correctness of the quotient formula network)** The LTS corresponding to the quotient formula network of  $\varphi$  with respect to  $\vec{S}[i]$  encodes a formula that is equivalent to  $\varphi //_i^0 s_i^0$ , where  $s_i^0$  is the initial state of  $\vec{S}[i]$ .

**Example 10** Consider the network  $N$  of Example 1 and the formula of Example 9. Quotienting of the formula with respect to  $S_3$  involves the following set of rules:

$$\{ ((\neg, \bullet), \neg), ((\vee, \bullet), \vee), ((\mu^0, \bullet), \mu^0), ((\langle a \rangle, \bullet), \langle a \rangle), ((\langle a \rangle, a), \langle \alpha_a \rangle), ((\langle b \rangle, b), \langle \alpha_b \rangle) \}$$

It yields the formula graph depicted in Figure 5 (b). This graph encodes as expected the quotient formula of Example 8, which can be evaluated on  $N_{\overline{\{3\}}}$ .

Working with formulas in disjunctive form is crucial: branches in the formula graph denote disjunctions between sub-formulas (*or-nodes*). During composition between the formula graph and a component LTS, the impossibility to synchronize on a modality  $\langle a \rangle$  (no transition labelled by  $\vec{t}[i]$  in the current state of the component LTS) denotes invalidation of the corresponding sub-formula, which merely disappears, in conformance with the equality  $\text{false} \vee \varphi_0 = \varphi_0$ .

The quotient of a formula graph with  $n$  states with respect to an LTS with  $m$  states may have up to  $n \times m$  states. Hence, as observed by Andersen [4], simplifications are needed to keep intermediate quotiented formulas at a reasonable size. We present in Figure 6 several simplifications applying to formula graphs, as conditional rules of the form “ $l \rightsquigarrow r$  (*cond*)” where  $l$  and  $r$  are transition relations and *cond* is a Boolean condition.  $l$ ,  $r$ , and *cond* are expressed using variables representing either states (written  $s, s_1, s_2, \dots$ ) or labels (written  $\sigma, \sigma_1, \sigma_2, \dots$ ), such that every variable occurring in  $r$  or in *cond* must also occur in  $l$ . It means that all transitions matching the left-hand side so that *cond* is satisfied can be replaced by the transitions of the right-hand side.

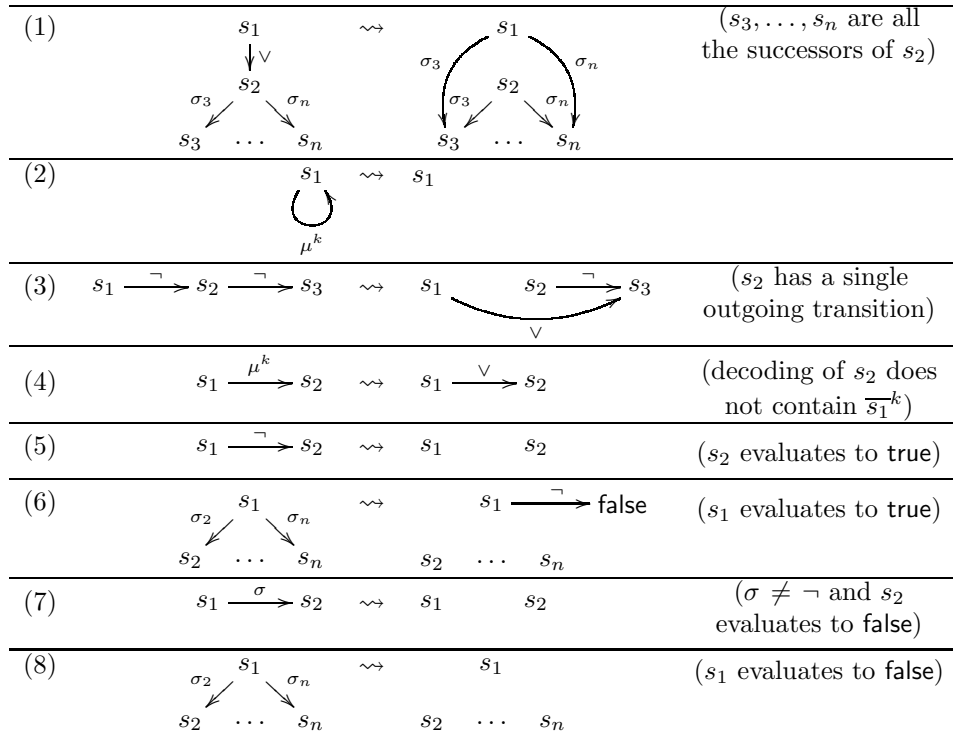


Figure 6: Simplification rules applying to formula graphs

**Elimination of  $\vee$ -transitions** (1). This rule allows transitions generated by synchronization rules of the form  $((\langle a \rangle, \vec{t}[i]), \vee)$  in the quotient formula network to be eliminated. This elimination can be achieved efficiently by applying reduction modulo  $\tau^*.a$  equivalence [51],  $\vee$ -transitions being interpreted as internal ( $\tau$ ) transitions.

**Elimination of unguarded variables** (2). When combined with the previous rule, this rule allows unguarded variable occurrences to be eliminated. Indeed, an unguarded variable is characterized by a (possibly empty) sequence of  $\vee$ -transitions connecting the target and source of a  $\mu$ -transition. The elimination of this sequence of  $\vee$  transitions then produces a self-looping transition labelled by  $\mu$ , which can be thereafter eliminated using the current rule.

**Elimination of double-negations** (3). This rule can be used to simplify formulas of the form  $\neg\neg\varphi$ , which often occur in quotient formulas. For instance, a double-negation is introduced in the quotient of the formula  $\neg\langle a \rangle\neg\varphi'$  (which is the disjunctive form of  $[a]\varphi'$ ) with respect to an LTS that offers an action synchronizing with  $a$  (thus having the modality disappear if the synchronization is binary).

**Elimination of  $\mu$ -transitions** (4). In this rule, the transition from  $s_1$  to  $s_2$  denotes the binder of a propositional variable  $\overline{s_1^k}$ . If this variable does not occur free in the sub-formula denoted by state  $s_2$ , then the  $\mu$ -transition can be replaced by an  $\vee$ -transition, which can be subsequently eliminated using rule (1). Determining whether  $\overline{s_1^k}$  occurs free would require to decode the formula graph, which should be avoided in practice. For this reason, we only consider the following sufficient conditions, which can be checked in linear-time:

- $s_1$  and  $s_2$  are not in the same strongly connected component (i.e., there is no path from  $s_2$  to  $s_1$ ), or
- $s_1$  is not the initial state and has a single predecessor  $p$ , and either (1)  $p$  has a single outgoing transition (which necessarily goes to  $s_1$ ) and this transition is labelled by  $\mu^{k'}$ , or (2)  $p$  satisfies the same condition

as  $s_1$ , recursively (this recursion is well-founded as long as it is applied to states reachable from the initial state)

**Evaluation of constant sub-formulas** (5–8). These four rules apply when some state denotes a sub-formula that evaluates to a constant in any context. This can be determined by using the following BES, which implements partial evaluation of the formula. This BES consists of blocks  $T^k$  and  $F^k$  ( $k \in [0, n]$ ) of respective signs  $\mu$  and  $\nu$ ,  $n$  being the greatest block number in the formula graph. Blocks are ordered so that  $k < k'$  implies  $T^k$  (resp.  $F^k$ ) occurs before  $T^{k'}$  (resp.  $F^{k'}$ ):

$$\begin{aligned} T^k : & \left\{ T_s^k = \mu \bigvee_{s \xrightarrow{\nu} s'} T_{s'}^k \vee \bigvee_{s \xrightarrow{\neg} s'} F_{s'}^k \vee \bigvee_{s \xrightarrow{\mu^{k'}} s'} T_{s'}^{k'} \right\}_{s \in \Sigma} \\ F^k : & \left\{ F_s^k = \nu \bigwedge_{s \xrightarrow{\nu} s'} F_{s'}^k \wedge \bigwedge_{s \xrightarrow{\langle \beta \rangle} s'} F_{s'}^k \wedge \bigwedge_{s \xrightarrow{\neg} s'} T_{s'}^k \wedge \bigwedge_{s \xrightarrow{\mu^{k'}} s'} F_{s'}^{k'} \right\}_{s \in \Sigma} \end{aligned}$$

We consider only the variables reachable from  $T_{s_0}^0$  or  $F_{s_0}^0$ ,  $s_0$  being the initial state of the formula graph. A state  $s$  denotes true (resp. false) if the Boolean variables  $T_s^k$  (resp.  $F_s^k$ ) evaluate to true in all (reachable) blocks  $k$ . Due to the presence of modalities, there may be states  $s$  and blocks  $k$  such that  $T_s^k$  and  $F_s^k$  are both false, indicating that the corresponding sub-formula is not constant. Intuitively,  $T_s^k$  expresses that  $s$  evaluates to true in block  $k$  (i.e.,  $\overline{s}^k$  evaluates to true) if one of its successors following a transition labelled by  $\vee$  or  $\mu^{k'}$  evaluates to true, or one of its successors following a transition labelled by  $\neg$  evaluates to false. Variable  $F_s^k$  expresses that state  $s$  evaluates to false in block  $k$  (i.e.,  $\overline{s}^k$  evaluates to false) if all its successors following transitions labelled by  $\vee$ ,  $\mu^{k'}$ , or modalities (by applying the identity  $\langle a \rangle \text{false} = \text{false}$ ) evaluate to false and all its successors following transitions labelled by  $\neg$  evaluate to true. Regarding fixed point signs, observe that for the formula  $\mu X^k.X^k$  (which is equivalent to the constant false),  $F_{\mu X^k.X^k}^k$  and  $T_{\mu X^k.X^k}^k$  are defined respectively by the greatest fixed point equation  $F_{\mu X^k.X^k}^k = \nu F_{\mu X^k.X^k}^k$  and the least fixed point equation  $T_{\mu X^k.X^k}^k = \mu T_{\mu X^k.X^k}^k$ . This BES has the solution  $F_{\mu X^k.X^k}^k = \text{true}$ ,  $T_{\mu X^k.X^k}^k = \text{false}$ , reflecting that  $\mu X^k.X^k$  is constantly false, as expected.

Repeated application of quotienting progressively eliminates modalities, until none of them remains in the formula graph, which then necessarily evaluates to a constant equal to the result of evaluating the formula on the whole network.

**Sharing of equivalent sub-formulas.** In addition to the above eight rules, reducing a formula graph modulo strong bisimulation does not change its semantics. Strong bisimulation reduction can thus decrease the size of intermediate formula graphs.

**Example 11** After applying the above simplifications to the formula graph of Example 10, we obtain the (smaller) formula graph depicted in Figure 5 (c), which corresponds to the formula  $\langle \langle a \rangle \text{true} \rangle \vee \langle \langle \alpha_a \rangle \text{true} \rangle \vee \langle \langle \alpha_b \rangle \langle a \rangle \text{true} \rangle$ .

**Example 12** The graph corresponding to  $\mu X^0.(\langle a \rangle \mu Y^0. \langle b \rangle X^0) \vee \langle c \rangle X^0$  reduces as expected to a deadlock state representing the constant false (left as an exercise).

Note that the simplification of a formula graph produces a formula graph. In particular, the parity of the number of occurrences of the label  $\neg$  on paths leading to a  $\mu^k$ -transition is not changed by any rule, including rule (3) which eliminates negations by pair. Also, the simplifications do not create new circuits and every  $\mu^k$ -transition eliminated by rule (4) cannot be the first  $\mu^k$ -transition occurring on any circuit.

All the simplifications that we propose in this paper correspond more or less to simplifications already proposed by Andersen [4], but we apply them directly on formula graphs instead of systems of  $\mu$ -calculus equations.

The paper [88] presents a specialization of this technique to the alternation-free  $\mu$ -calculus formulas (using alternation-free BES), and how this specialisation can be again generalised to handle also useful fairness operators of alternation 2 (in particular the infinite looping operator  $\langle \_ \rangle @$ ) in linear time without developing the complex machinery to evaluate general alternation-2  $\mu$ -calculus formulas.

## 5 High-level strategies and smart reduction

The efficiency of compositional model generation (including formula graph generation in the case of partial model checking) depends on the order in which the components are gathered and minimized. In practice, the order can be specified by the user, but since it is not always possible to predict whether an order will be more or less efficient than another without trying them and comparing the results, the user generally has to rely on intuition. This burden can be relieved using pre-defined strategies, which can be either based on the component hierarchy, or on heuristics to automatically determine efficient composition orders, based on the interactions between components.

### 5.1 Pre-defined strategies based on component hierarchy

We present three strategies for compositional model generation modulo an equivalence relation  $R$ , namely *leaf reduction*, *root leaf reduction*, and *node reduction*.

**Definition 25 (Leaf and root leaf reductions)** *Given a composition expression  $B$  and an equivalence relation  $R$  that is a congruence for composition expressions, leaf reduction computes another composition expression  $\text{leaf\_reduce}_R(B)$  which has the same hierarchy as  $B$ , such that:*

- *hiding and cutting operators are propagated as far as possible in the composition expression, and*
- *the LTSs occurring in  $\text{leaf\_reduce}_R(B)$  are minimized modulo the equivalence relation  $R$  after hiding and cutting.*

Formally,  $\text{leaf\_reduce}_R(B)$  is a shorthand for  $\text{leaf\_reduce}_R(B, \emptyset, \emptyset)$ , where  $\text{leaf\_reduce}_R(B, A_h, A_c)$  is defined as follows,  $A_h$  denoting a set of labels to be hidden in  $B$  and  $A_c$  denoting a set of labels to be cut in  $B$ . The sets  $A_h$  and  $A_c$  satisfy the invariant  $A_h \cap A_c = \emptyset$ .

$$\begin{aligned}
\text{leaf\_reduce}_R(S, A_h, A_c) &= \text{reduce}_R(\mathbf{hide } A_h \mathbf{ in cut } A_c \mathbf{ in } S) \\
\text{leaf\_reduce}_R(B_1 \parallel [A] B_2, A_h, A_c) &= \\
&\quad \mathbf{hide } A_h \cap A \mathbf{ in} \\
&\quad \text{leaf\_reduce}_R(B_1, A_h \setminus A, A_c) \\
&\quad \parallel [A] \\
&\quad \text{leaf\_reduce}_R(B_2, A_h \setminus A, A_c) \\
\text{leaf\_reduce}_R(\mathbf{rename } \theta \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \mathbf{rename } \theta \mathbf{ in leaf\_reduce}_R(B_0, \bar{\theta}^{-1}(A_h), \bar{\theta}^{-1}(A_c)) \\
&\quad \text{where } \bar{\theta}^{-1}(A) = \{a \mid \bar{\theta}(a) \in A\} \\
\text{leaf\_reduce}_R(\mathbf{hide } A \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \text{leaf\_reduce}_R(B_0, A_h \cup A, A_c \setminus A) \\
\text{leaf\_reduce}_R(\mathbf{cut } A \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \text{leaf\_reduce}_R(B_0, A_h \setminus A, A_c \cup A)
\end{aligned}$$

The root leaf reduction is the same as leaf reduction, except that the LTS corresponding to the composition expression  $B'$  is finally generated and minimized. Therefore, the final result of root leaf reduction is a (minimal) LTS instead of a composition expression:

$$\text{root\_leaf\_reduce}_R(B) = \text{reduce}_R(\text{leaf\_reduce}_R(B))$$

**Proposition 11** *If  $R$  is a congruence for composition expressions, then  $\text{leaf\_reduce}_R(B)$  and  $\text{root\_leaf\_reduce}_R(B)$  are both equivalent to  $B$  modulo  $R$ .*

Note however that  $B$ ,  $\text{leaf\_reduce}_R(B)$ , and  $\text{root\_leaf\_reduce}_R(B)$  are not the same operationally: the sizes of intermediate LTSs differ in both expressions.



**Proposition 12** *For strong, branching, and divbranching bisimulations, the largest intermediate LTS generated to compute  $\text{root\_leaf\_reduce}_R(B)$  is not larger than the largest intermediate LTS generated to compute  $B$ .*

The *node reduction* is another strategy, which makes more intermediate steps than *leaf* and *root leaf reduction*: an intermediate LTS is generated and minimized after each parallel composition.

**Definition 26 (Node reduction)** *Given a composition expression  $B$  and an equivalence relation  $R$  that is a congruence for composition expressions, node reduction computes an LTS  $\text{node\_reduce}_R(B)$ . Formally,  $\text{node\_reduce}_R(B)$  is a shorthand for  $\text{node\_reduce}_R(B, \emptyset, \emptyset)$ , where  $\text{node\_reduce}_R(B, A_h, A_c)$  is defined as follows,  $A_h$  denoting a set of labels to be hidden in  $B$  and  $A_c$  denoting a set of labels to be cut in  $B$ . The sets  $A_h$  and  $A_c$  satisfy the invariant  $A_h \cap A_c = \emptyset$ .*

$$\begin{aligned}
\text{node\_reduce}_R(S, A_h, A_c) &= \text{reduce}_R(\mathbf{hide } A_h \mathbf{ in cut } A_c \mathbf{ in } S) \\
\text{node\_reduce}_R(B_1 \parallel [A] \parallel B_2, A_h, A_c) &= \\
&\quad \text{reduce}_R(\mathbf{hide } A_h \cap A \mathbf{ in} \\
&\quad \quad \text{node\_reduce}_R(B_1, A_h \setminus A, A_c) \\
&\quad \quad [A] \\
&\quad \quad \text{node\_reduce}_R(B_2, A_h \setminus A, A_c)) \\
\text{node\_reduce}_R(\mathbf{rename } \theta \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \text{reduce}_R(\mathbf{rename } \theta \mathbf{ in node\_reduce}_R(B_0, \bar{\theta}^{-1}(A_h), \bar{\theta}^{-1}(A_c))) \\
&\quad \text{where } \bar{\theta}^{-1}(A) = \{a \mid \bar{\theta}(a) \in A\} \\
\text{node\_reduce}_R(\mathbf{hide } A \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \text{node\_reduce}_R(B_0, A_h \cup A, A_c \setminus A) \\
\text{node\_reduce}_R(\mathbf{cut } A \mathbf{ in } B_0, A_h, A_c) &= \\
&\quad \text{node\_reduce}_R(B_0, A_h \setminus A, A_c \cup A)
\end{aligned}$$

**Proposition 13** *If  $R$  is a congruence for composition expressions, then  $\text{node\_reduce}_R(B)$  is equivalent to  $B$  modulo  $R$ .*

Unlike root leaf reduction (which does not compute intermediate compositions before computing the LTS corresponding to  $B$ ), there is no guarantee that computing the node reduction of  $B$  is less expensive (in time or memory) than computing the LTS corresponding to  $B$ . Indeed, the LTSs corresponding to the intermediate compositions may be larger (even much larger) than the LTS corresponding to  $B$ . However, there exist cases where node reduction is efficient.

## 5.2 Heuristic-based strategy: Smart reduction

Heuristics to automatically determine efficient composition orders, based on the component interactions, have been proposed in [128] for concurrent finite state machines communicating via named channels. More recently, such heuristics have been refined and implemented in a prototype tool for LTSs synchronizing on their common alphabets [40]. In both works, the components to be composed are selected using two metrics: an estimate of the proportion of internal transitions in the composition (the higher, the more the composition graph being expected to be reducible), and an estimate of the proportion of transitions that interleave.

A limitation of the above works lies in the restricted composition operators, which are generally insufficient to capture the semantics of state-of-the-art software description languages: The parallel composition operators used in [128] do not support multiway synchronization (more than two components synchronizing all together), and none of the parallel composition operators used in [128, 40] enable nondeterministic synchronization (e.g., two clients competing to synchronize with a server on some label).

In this section, we present a refinement of the compositional model generation techniques of [128, 40], called *smart reduction* [41], which relies on networks of LTSs. The first advantage is that it makes compositional model generation applicable to a wider variety of operators, provided that they can be compiled into networks. The second advantage is that networks enable any composition order, which is not in general the

case in process algebraic models, where some composition orders, possibly including the optimal order, may not be represented using the available algebraic operators.

The LTS corresponding to a network can be generated incrementally, by alternating compositions of well-chosen subsets of the components and minimizations modulo an equivalence relation. We call *aggregation* a composition followed by a minimization of the result, and *compositional aggregation* this incremental technique. Contrary to composition expressions, networks enable all possible aggregations as they are not constrained by a component hierarchy.

**Definition 27 (Aggregation)** *As in Definition 19, we assume a function  $\alpha(\vec{t}, a)$  that associates to each  $(\vec{t}, a) \in V$  a unique label distinct from all others. Given a network  $N$  of size  $n$  and a set of indices  $I \subseteq [1, n]$ , we define the network  $\text{agg}(N, I)$  obtained by aggregation of components inside  $I$  as follows:*

$$\begin{aligned} \text{agg}(N, I) &= (\text{reduce}_R(N_{|I}) :: \vec{S}|I, V_{\text{agg}}) \\ \text{where } N_{|I} &= (\vec{S}|I, V_{|I}) \\ V_{\text{agg}} &= \left\{ \begin{array}{l} (a :: \vec{t}|I, a) \quad | \quad (\vec{t}, a) \in V \wedge \text{acv}(\vec{t}) \subseteq I \\ (\alpha(\vec{t}, a) :: \vec{t}|I, a) \quad | \quad (\vec{t}, a) \in V \wedge \emptyset \subsetneq (I \cap \text{acv}(\vec{t})) \subsetneq \text{acv}(\vec{t}) \\ (\bullet :: \vec{t}|I, a) \quad | \quad (\vec{t}, a) \in V \wedge (I \cap \text{acv}(\vec{t})) = \emptyset \end{array} \right\} \cup \\ \text{and } V_{|I} &= \left\{ \begin{array}{l} (\vec{t}|I, a) \quad | \quad (\vec{t}, a) \in V \wedge \text{acv}(\vec{t}) \subseteq I \\ (\vec{t}|I, \alpha(\vec{t}, a)) \quad | \quad (\vec{t}, a) \in V \wedge \emptyset \subsetneq (I \cap \text{acv}(\vec{t})) \subsetneq \text{acv}(\vec{t}) \end{array} \right\} \cup \end{aligned}$$

The components inside  $I$  ( $\vec{S}|I$ ) are aggregated into  $\text{reduce}_R(N_{|I})$  and the components outside  $I$  ( $\vec{S}|I$ ) are kept non-aggregated. The synchronization rules  $V_{|I}$  of the auxiliary network  $N_{|I}$  are obtained by projection of  $V$  on to  $I$ , whereas the synchronization rules  $V_{\text{agg}}$  are obtained by synchronization of the labels in  $V_{|I}$  with the projection of  $V$  on to  $\bar{I}$ .

For convenience, since  $I$  denotes the set of components to be aggregated, we call  $I$  an *aggregation*.

The rules of  $V_{|I}$  and  $V_{\text{agg}}$  are organized in three subsets:

- The first subset, containing those rules of the form  $(\vec{t}|I, a)$  in  $V_{|I}$  and  $(a :: \vec{t}|I, a)$  in  $V_{\text{agg}}$ , represents the synchronization rules that involve only components inside  $I$ . In this case,  $\vec{t}|I$  is a vector of  $\bullet$ , which expresses that transitions of  $\text{reduce}_R(N_{|I})$  obtained by synchronization of components inside  $I$  do not need to synchronize with components outside  $I$ .
- The second subset, containing those rules of the form  $(\vec{t}|I, \alpha(\vec{t}, a))$  in  $V_{|I}$  and  $(\alpha(\vec{t}, a) :: \vec{t}|I, a)$  in  $V_{\text{agg}}$ , represents the synchronization rules that involve both components inside  $I$  and components outside  $I$ . This expresses that transitions of  $\text{reduce}_R(N_{|I})$  obtained by synchronization of components inside  $I$  still need to synchronize with components outside  $I$ . The special label  $\alpha(\vec{t}, a)$  is an intermediate label used for this synchronization. Note that in general,  $a$  cannot be used instead of  $\alpha(\vec{t}, a)$  because (1)  $a$  can be the internal label  $\tau$  (even though  $\vec{t}$  synchronizes only visible labels), which cannot be synchronized, and (2) in general there can be several rules with the same label  $a$  (in particular when synchronization is nondeterministic involved) and using  $a$  could create unexpected synchronizations.
- The third subset, containing those rules of the form  $(\bullet :: \vec{t}|I, a)$  in  $V_{\text{agg}}$ , represents the synchronization rules that involve only components outside  $I$ . These rules do not impose synchronization constraints on components inside  $I$ , thus explaining why  $V_{|I}$  has no rule in the third subset.

**Proposition 14** *If  $R$  is a congruence for networks, then  $\text{lts}(\text{agg}(N, I))$  is equivalent modulo  $R$  to  $\text{lts}(N)$ .*

**Definition 28 (Compositional aggregation algorithm)** *A compositional aggregation algorithm to generate and minimize modulo  $R$  the LTS corresponding to a network of LTSs  $((S_1, \dots, S_n), V)$  of size  $n$  is defined iteratively as follows:*

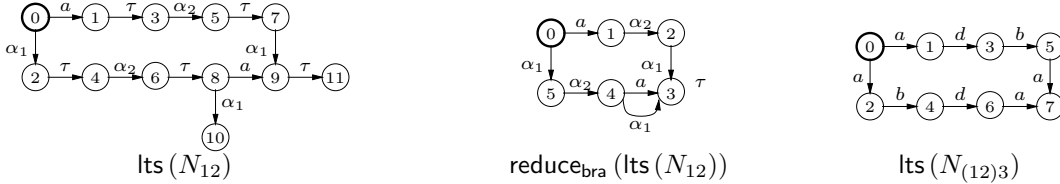
1.  $N := (\text{reduce}_R(S_1), \dots, \text{reduce}_R(S_n), V)$

2. Select a set  $I$  of indices containing at least those of two of the component LTSs of  $N$ . A strategy for selection will be addressed in the next section.
3.  $N := \text{agg}(N, I)$ .
4. If  $N$  still contains more than two LTSs, then continue in step 2. Otherwise return  $\text{reduce}_R(N)$ .

**Proposition 15** *The compositional aggregation algorithm terminates and if  $R$  is a congruence for networks, then it returns an LTS that is equivalent to its input network modulo  $R$ .*

Equivalence modulo  $R$  follows from Proposition 14, which ensures that  $\text{lts}(N)$  remains invariantly equivalent to the input modulo  $R$  until the end of the algorithm. Termination follows from the fact that the size of  $\text{agg}(N, I)$  is the size of  $N$  plus 1 minus the size of  $I$  (which is at least 2). Since  $N$  is substituted by  $\text{agg}(N, I)$  at each step, this guarantees that the size of  $N$  strictly decreases at each step. These results establish the correctness of compositional aggregation.

**Example 13** *We write  $\text{reduce}_{\text{bra}}(S)$  for minimization of  $S$  modulo branching bisimulation. The compound LTS corresponding to the network of Example 1, whose components  $S_1$ ,  $S_2$ , and  $S_3$  are already minimal, can be generated modulo branching bisimulation by first composing  $S_1$  and  $S_2$ , then  $S_3$  as follows. First, build  $N_{12} = N_{\{1,2\}} = ((S_1, S_2), V_{12})$  for the aggregation of  $S_1$  and  $S_2$ , where  $V_{12}$  is the set of rules  $\{((a, a), a), ((a, \bullet), \alpha_1), ((b, b), \alpha_2), ((c, c), \tau)\}$ ,  $\alpha_1$  is the label  $\alpha((a, \bullet), a, a)$ , and  $\alpha_2$  is the label  $\alpha((b, b), b)$ . Compute the intermediate LTS  $S_{12} = \text{reduce}_{\text{bra}}(\text{lts}(N_{12}))$  (see below). Second, build  $N_{(12)3} = \text{agg}(N, \{1, 2\}) = ((S_{12}, S_3), V_{(12)3})$ , where  $V_{(12)3}$  is the set  $\{((a, \bullet), a), ((\alpha_1, a), a), ((\alpha_2, b), b), ((\tau, \bullet), \tau), ((\bullet, d), d)\}$ . Return  $\text{reduce}_{\text{bra}}(\text{lts}(N_{(12)3}))$  (see  $\text{lts}(N_{(12)3})$  below), which is branching equivalent to  $\text{lts}(N)$  (see Figure 1). Note that both LTSs  $\text{lts}(N_{12})$  and  $\text{lts}(N_{(12)3})$  are smaller than  $\text{lts}(N)$ .*



Other aggregation orders are possible, yielding different intermediate graphs. Figure 7 gives the sizes of those intermediate graphs corresponding to the different aggregation orders. The largest intermediate graph size of each order is indicated in bold type. This table shows that the aggregation order described above (order 1) is optimal in terms of largest intermediate graph (12 states and 12 transitions).

The most difficult issue in compositional aggregation concerns step 2 of the algorithm, namely to select, if possible automatically, an aggregation  $I$  that avoids state explosion. We present *smart reduction*, which corresponds to compositional aggregation using a heuristic based on metrics evaluated against possible aggregations. The metrics use an estimate of the number of compound transitions in  $I$ , defined as follows.

**Definition 29 (Estimate number of compound transitions)** *The estimate number of compound transitions in  $I$  generated by synchronization vector  $\vec{t}$ , written  $ET(I, \vec{t})$ , is defined below:*

$$ET(I, \vec{t}) = \begin{cases} 0 & \text{if } I \cap \text{acv}(\vec{t}) = \emptyset \\ \prod_{i \in I \setminus \text{acv}(\vec{t})} |\Sigma_i| \times \prod_{i \in I \cap \text{acv}(\vec{t})} |\vec{t}[i]_i| & \text{otherwise} \end{cases}$$

Informally,  $ET(I, \vec{t})$  counts, for vector  $\vec{t}$ , the number of transitions going out of every product state of  $I$ , including unreachable states. This count equals 0 if  $\vec{t}$  does not involve components inside  $I$  (first line). Otherwise,  $N_I$  has a rule of the form  $(\vec{t}|I, b)$ . This rule generates a transition in a state of  $N_I$  without condition on the states of the components  $\vec{S}[i]$  such that  $i \in I \setminus \text{acv}(\vec{t})$  — thus justifying the first

<b>Order 1</b> : $(S_1, S_2)$ then $S_3$	states	transitions
$\text{comp}(S_1, S_2)$	<b>12</b>	<b>12</b>
$\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_2))$	6	7
$\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_2)), S_3)$	8	8
$\text{reduce}_{\text{bra}}(\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_2)), S_3))$	7	7
<b>Order 2</b> : $(S_1, S_3)$ then $S_2$	states	transitions
$\text{comp}(S_1, S_3)$	<b>19</b>	<b>23</b>
$\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_3))$	17	22
$\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_3)), S_2)$	15	17
$\text{reduce}_{\text{bra}}(\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_3)), S_2))$	7	7
<b>Order 3</b> : $(S_2, S_3)$ then $S_1$	states	transitions
$\text{comp}(S_2, S_3)$	<b>18</b>	<b>34</b>
$\text{reduce}_{\text{bra}}(\text{comp}(S_2, S_3))$	<b>18</b>	<b>34</b>
$\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_2, S_3)), S_1)$	15	17
$\text{reduce}_{\text{bra}}(\text{comp}(\text{reduce}_{\text{bra}}(\text{comp}(S_2, S_3)), S_1))$	7	7
<b>Order 4</b> : $(S_1, S_2, S_3)$	states	transitions
$\text{comp}(S_1, S_2, S_3)$	<b>15</b>	<b>17</b>
$\text{reduce}_{\text{bra}}(\text{comp}(S_1, S_2, S_3))$	7	7

Figure 7: Sizes of intermediate graphs for all aggregation orders; given  $S_1, \dots, S_m$  some components of the current network  $N$  and  $I$  the set of their indices,  $\text{comp}(S_1, \dots, S_m)$  denotes  $\text{lts}(N|_I)$

product in the definition of  $ET(I, \vec{t})$  — and provided that the states of the components  $\vec{S}[i]$  such that  $i \in I \cap \text{acv}(\vec{t})$  have a transition labelled  $\vec{t}[i]$  — thus justifying the second product. In general, the exact number of compound transitions in  $I$  generated by synchronization vector  $\vec{t}$  is below this count since some states may be unreachable.

We now define our metrics on networks, which is the sum of two terms, namely the *hiding metrics* and the *interleaving metrics*, defined below.

**Definition 30 (Hiding metrics)** *The hiding metrics is defined by  $HM(I) = HR(I)/|I|$ , where  $HR(I)$  (the hiding rate) is defined as follows:*

$$HR(I) = \frac{\sum_{(\vec{t}, \tau) \in V \wedge \text{acv}(\vec{t}) \subseteq I} ET(I, \vec{t})}{1 + \sum_{(\vec{t}, a) \in V} ET(I, \vec{t})}$$

Informally,  $HR(I)$  represents an estimate of the proportion of transitions in  $N|_I$  that are internal. Those transitions are necessarily created by rules involving only components inside  $I$ . The addition of 1 in its divisor avoids division by 0 in pathological cases. The divisor  $|I|$  of  $HM(I)$  aims at favouring smaller aggregations, which are likely to yield smaller intermediate LTSs.

Using  $HM(I)$  is justified in the context of weak equivalence relations (e.g., branching bisimulation), because internal transitions are often eliminated by the corresponding minimizations. Although to a lesser extent, it is also justified in the context of strong bisimulation, because hiding enables abstracting away from labels that otherwise would differentiate the behaviour of equivalent states.

However, in both cases,  $HM(I)$  is not sufficient to avoid intermediate explosion due to the aggregation of loosely synchronized LTSs. To palliate this, the hiding metrics will be combined with the *interleaving metrics* defined as follows.

**Definition 31 (Interleaving metrics)** *Let  $\vec{t}@i$  denote the synchronization vector of size  $n$  defined by  $(\vec{t}@i)[i] = \vec{t}[i]$  and  $(\forall j \in [1, n] \setminus \{i\}) (\vec{t}@i)[j] = \bullet$ .*

*The interleaving metrics is defined by  $IM(I) = (1 - IR(I))/|I|$ , where the interleaving rate  $IR(I)$  is defined as follows:*

$$IR(I) = \frac{\sum_{(\vec{t}, a) \in V} ET(I, \vec{t})}{1 + \sum_{(\vec{t}, a) \in V} \sum_{i \in I \cap \text{acv}(\vec{t})} ET(I, \vec{t} @ i)}$$

Intuitively, the value  $IR(I)$  is the quotient between an estimate of the number of compound transitions of  $I$  and the number of compound transitions that  $I$  would have if all components were fully interleaving. For an aggregation  $I$  of fully interleaving components, we thus have  $IR(I)$  very close to 1. It is a refinement of the *interleaving count* defined in [40], which uses the proportion of fully interleaving (i.e., non-synchronized) component LTS transitions out of the total number of component LTS transitions. We believe that  $IR(I)$  is more accurate, because it also measures the *partial* interleaving of synchronized transitions with the remainder of the aggregation.

**Definition 32 (Combined metrics)** *Taking into account both the hiding rate and the interleaving rate, we use here the combined metrics  $CM(I) = HM(I) + IM(I)$ .*

Smart reduction selects the aggregation to which the metrics gives the highest value (high proportion of internal transitions and low interleaving). To avoid the combinatorial explosion of the number of aggregations, we proceed as in [40] and only consider: (1) aggregations whose size is bounded by a constant (definable by the user), and (2) aggregations that are *connected*. An aggregation is connected if for each pair of distinct components  $S_i, S_j$  in the aggregation,  $S_i$  and  $S_j$  are *connected*, which is defined recursively as follows: either there is a synchronization rule  $(\vec{t}, a)$  such that  $\{i, j\} \subseteq \text{acv}(\vec{t})$  (i.e.,  $S_i$  and  $S_j$  are synchronized) or, recursively, the aggregation contains a third (distinct) LTS  $S_k$  connected to both  $S_i$  and  $S_j$ .

**Example 14** *The metrics evaluate as follows on the network of Examples 1 and 13:*

$I$	$\{1, 2\}$	$\{1, 3\}$	$\{2, 3\}$	$\{1, 2, 3\}$
$HM(I)$	0.211	0	0	0.129
$IM(I)$	0.357	0.227	0.124	0.249
$CM(I)$	0.568	0.227	0.124	0.378

*As expected, the combined metrics gives the highest value to  $\{1, 2\}$ , therefore designating it as the best aggregation in the first step. Note that, more generally in this example, a comparison between the graph sizes in Figure 7 and the values in the above table shows that one aggregation is more efficient than another whenever the combined metrics gives it a higher value.*

### 5.3 Combining smart reduction and partial model checking

The order in which the components of the network are composed and minimized has also an influence on the efficiency of partial model checking. The only difference is that we aim to generate compositionally the LTS corresponding to the quotient formula network obtained by composition of the formula graph with the network of LTSs under verification. Therefore, the smart reduction heuristics can be used to select the next aggregation to be computed. The reduction that has to be applied at each step depends whether the selected aggregation contains the formula graph (corresponding to the computation of a quotient) or not (corresponding to a simple compositional model generation step). If the formula graph belongs to the aggregation, then the simplifications designed for partial model checking must be applied. Otherwise, the aggregation can be minimized modulo any appropriate equivalence relation that is adequate with the formula under verification.

Currently, we use a simple strategy: the heuristics always selects an aggregation limited to two components, which always includes the formula graph, so that the formula graph simplifications are applied at each step. More elaborate strategies could be designed in the future.

## 6 Implementation in the CADP toolbox

All the above concepts and algorithms for compositional verification have been implemented in the framework of CADP [58]. CADP (*Construction and Analysis of Distributed Processes*) is a tool platform implementing salient results of concurrency theory, with a particular focus on applicability and scalability of the proposed approaches. Related tool platforms for asynchronous, action-based systems are, e.g., the *Edinburgh Concurrency Workbench* [37] and *Concurrency Workbench of the New Century*<sup>6</sup> for CCS, the *Failures-Divergences Refinement* tools [10, 65] for CSP, the *micro Common Representation Language*<sup>7</sup> toolset [72] for ACP, and LTSmin<sup>8</sup>, which supports several input languages [21].

The CADP<sup>9</sup> toolbox is one of the pioneering platforms in concurrency theory, as its development was undertaken in the mid-80s. Since then, the toolbox has grown to include nearly 50 tools; moreover, CADP tries as much as possible to interface with tools developed by other research teams, so as to provide comprehensive solutions for analyzing complex concurrent models.

Many tools of CADP address compositional verification issues, or are needed to do so. In the present section, we give an orderly presentation of these tools sorted according to their functionalities.

### 6.1 Labelled transition systems and their generation

CADP implements the theoretical concept of LTS in two different ways:

- For *explicit* LTSs that are defined *in extension* by giving the list of their states and transitions, CADP provides a file format named BCG (*Binary-Coded Graphs*). This format enables large LTSs to be stored on disk in a compact manner (typically, two bytes per transition). It is equipped with programming libraries for reading and writing BCG files, as well as a collection of tools that query and transform BCG files, e.g., to compute statistics such as the branching factor, to display the list of labels attached to transitions, to relabel transitions using string substitutions defined by regular expressions, etc. Notice that CADP supports other LTS formats (such as AUT, FC2, and SEQ), which are textual and, thus, readable by humans; however, BCG remains the format of choice for large explicit LTSs.
- For *implicit* LTSs that are defined *in comprehension* by giving their initial state and their transition function, CADP provides the OPEN/CAESAR Application Programming Interface [56]. This interface enables algorithms to be written for on-the-fly LTS exploration, and provides sufficient abstraction capabilities to make such algorithms independent from the details of the particular source language used to specify these LTSs. Above the OPEN/CAESAR interface, many tools have been developed, some of them will be described in this section.

These LTSs can be produced in different ways using the CADP tools. They can be generated from specifications written in a data-passing process calculus such as LOTOS (using the CAESAR.ADT and CAESAR compilers [55, 60]) or from the parallel composition of several LTSs (using the EXP.OPEN tool [85], which implements the parallel, hiding, and renaming operators of all mainstream process calculi, and supports the Arnold-Nivat synchronization vectors as well). These LTSs can also be generated from other concurrent languages, such as LNT [28], FSP [94, 89], CHP [98, 59] that are automatically translated to LOTOS and/or EXP.OPEN input formalism.

### 6.2 Equivalence checking: Minimization and comparison of labelled transition systems

The CADP toolbox also provides tools for minimizing (i.e., computing the quotient) of an LTS modulo various behavioral equivalence relations — especially bisimulations — and to compare two LTSs with each other modulo various behavioral equivalence or preorder relations. The development of such tools has been a

<sup>6</sup><http://www.cs.sunysb.edu/~cwb>

<sup>7</sup>[mcr12.org](http://mcr12.org)

<sup>8</sup>[fmt.cs.utwente.nl/tools/ltsmin](http://fmt.cs.utwente.nl/tools/ltsmin)

<sup>9</sup><http://cadp.inria.fr>

continuous effort since the early days of CADP. The ALDEBARAN tool [52] was developed for this purpose, and was later replaced by version 1 of another tool named BCG\_MIN.

At present, CADP provides three equivalence-checking tools: version 2 of BCG\_MIN [39], which can minimize an explicit LTS encoded in the BCG format, BCG\_CMP, which can compare two explicit LTSs encoded in the BCG format, and BISIMULATOR [102], which can compare an implicit LTS that implements the OPEN/CAESAR interface against an explicit LTS encoded in the BCG format. On powerful machines, these tools have been able to successfully handle LTS with one billion explicit states and several billion transitions. The comparison of LTSs can generate diagnostics, i.e., discriminating sequences of transitions (or more generally, graph fragments) that explain why two LTSs are not equivalent or included one in the other.

### 6.3 Semi-composition and interfaces

To implement the concept of semi-composition, CADP provides the PROJECTOR tool, which takes as input an implicit LTS (represented using the OPEN/CAESAR Application Programming Interface) and an interface (encoded in the BCG format), and produces as output a BCG file containing the semi-composition of the LTS and the interface. The PROJECTOR tool has been continuously enhanced: over the years, three successive versions of the tool have been developed. The current version is PROJECTOR 3. PROJECTOR was used successfully in the verification of an industrial protocol [132].

### 6.4 Model checking: Local and partial evaluation

There are many model checkers in the world, with many academic prototypes and a few industrial-strength tools. Most model checkers follow the *state-based approach*, in which temporal logic formulas are built upon atomic properties that can query the values of variables contained in each state of the system, e.g., computing the set of states in which a given variable  $X$  is positive. The state-based approach relies on a “white-box” assumption, i.e., the assumption that the contents of all variables of the system under verification are observable.

In concurrency theory, however, such a “white-box” assumption does not hold in general, and an *action-based approach* is used instead: the mainstream semantic models of concurrency theory (e.g., the concepts of LTS or Markov chains) follow a “black box” assumption, in which all information is attached to transitions and no information is attached to states, except the difference between the initial state(s) and other states. Practically, in order to remain compatible with equivalence checking (especially, bisimulations), the atomic properties of the temporal logic used must refer to actions (i.e., transition labels) rather than state contents. Such design constraint is materialized, e.g., by the difference that exists between the *modal  $\mu$ -calculus* [25], which is action-based, and the *propositional  $\mu$ -calculus* [82], which is state-based.

The CADP toolbox provides advanced tools for model checking, but does it in a way that remains compatible with the foundations of concurrency theory. Namely, the CADP tools follow the action-based approach rather than the state-based one. All the model checking tools of CADP work on the fly, on implicit LTSs rather than explicit LTSs, thus enabling to halt state-space generation as soon as the property under evaluation is found to be false or true. Moreover, all the model checking tools of CADP can produce diagnostics (also called witnesses or counterexamples), i.e., (explicit) fragments from the implicit LTS in order to explain why a property is true or false.

The first model checking tools added to CADP in the 90s were specialized for simple classes of properties. For instance, the EXHIBITOR tool searches for traces matching a given regular expression, and the TERMINATOR tool searches for deadlocks (i.e., states with no outgoing transition) using Holzmann’s *bitstate hashing* algorithm.

Later, more generic model checkers have been added to CADP. The EVALUATOR 3 model checker [104] is based on the alternation-free modal  $\mu$ -calculus, supplemented for convenience with regular expressions on action names and execution sequences. To produce its verdicts, EVALUATOR 3 relies on the CAESAR\_SOLVE engine [102], a library for solving Boolean equation systems on the fly and generating appropriate diagnostics. For many years, EVALUATOR 3 has been intensively used within CADP and applied to a number of large case studies, many of which in an industrial context.

Recently, EVALUATOR 4 [105] has been introduced as the new default model checker of CADP. EVALUATOR 4 generalizes EVALUATOR 3 by accepting a more expressive logic named MCL, in which one can specify properties and dependencies on the typed data values contained in action names. Using MCL, one can express meaningful and practically relevant properties (such as: “on any execution path, the sequence number attached to each message is constantly increasing”) that cannot be expressed using conventional logics, which consider actions as members of some finite alphabet, rather than complex data structures containing both communication channel names as well as typed value exchanged between concurrent processes. To evaluate the rich formula language of MCL, the concept of *Parameterized Boolean Equation Systems* [100] has been invented and implemented in the EVALUATOR 4 tool.

Finally, in order to support partial model checking, the PMC<sup>10</sup> tool has been developed and made available as a CADP companion tool.

## 6.5 The SVL language for compositional verification

Although compositional verification is firmly grounded in strong theoretical results and although it is likely to bring significant reductions in complexity, this is not sufficient to make it usable on a practical scale, especially by novice users. Indeed, applying compositional verification to any non-trivial system raises a number of difficult, low-level, yet unavoidable issues, among which: invoking all the required CADP tools in the proper order with appropriate command-line options, splitting models into components and logic formulas into subformulas, recombining system components and subformulas into larger agglomerates, and managing tens (and often hundreds) of intermediate files, which need to be archived, given unique names, and to be deleted as soon as they are no longer useful.

To address these issues, the CADP toolbox is equipped with a unique solution named SVL (*Script Verification Language*)<sup>11</sup> [57, 84]. SVL is both a high-level scripting language proposed to CADP end-users, as well as a compiler that translates SVL scripts into Bourne shell scripts, which can be directly executed on any POSIX system and perform all low-level tasks of invoking the CADP tools and managing intermediate files. SVL enables the description of systems consisting of:

- Explicit LTSs encoded in one of the formats supported by CADP, namely, AUT, BCG, FC2, or SEQ;
- Implicit LTSs represented using the OPEN/CAESAR Application Programming Interface and generated from high-level languages such as LOTOS, LNT, FSP, or any concurrent language that can be translated to these;
- Explicit LTSs obtained from implicit LTSs by exhaustively enumerating all their reachable states and transitions;
- Parallel composition of LTSs combined together using the parallel operators of mainstream process calculi (e.g., ACP/ $\mu$ CRL [72], CCS [108], CSP [125], LOTOS [79], E-LOTOS [80], LNT [28]) or Arnold-Nivat synchronization vectors;
- Semi-composition of an LTS projected over another LTS playing the role of an interface;
- Modification of an LTS by applying operations that hide or rename given labels, or that cut transitions labelled with given labels (i.e., labels that match specified regular expressions);
- Minimization (total or partial) of an LTS modulo some behavioural equivalence, e.g., bisimulation relations.

Additionally, SVL provides language constructs to perform the following verification tasks:

- Equivalence checking: comparison of two systems to decide whether they are equivalent or included one in the other modulo behaviour equivalence or preorder relations;

<sup>10</sup><http://convecs.inria.fr/software/pmc>

<sup>11</sup><http://cadp.inria.fr/man/svl.html>



Scenario	TFTP A		TFTP B	
	read	write	read	write
<i>A</i>		✓		
<i>B</i>	✓			
<i>C</i>		✓		✓
<i>D</i>	✓			✓
<i>E</i>	✓		✓	

Table 1: The five scenarios of the TFTP/UDP case study

- Model checking: search for deadlocks or livelocks in a system and, more generally, evaluation of a temporal logic formula on a system;
- Compositional verification: SVL has so-called *meta-operators* that recursively apply (total or partial) minimizations to the sub-components of a system, using various compositional reduction strategies, among which smart reduction.

Finally, SVL implements various expert strategies to make verification tractable. For instance, the LTSs representing interfaces are automatically minimized modulo safety equivalence before being used for semi-composition; also, when branching minimization fails to handle a large LTS, then SVL first tries to apply strong minimization before restarting branching bisimulation, and so on. Examples of SVL scripts are provided in Section 7.

## 7 Applications and experimental results

We now illustrate how the various compositional verification techniques presented in this paper and available in CADP can be applied to a common case study, an airplane-ground communication protocol studied by Airbus and based upon TFTP/UDP (*Trivial File Transfer Protocol/User Datagram Protocol*) [62]. The modelled system consists of two TFTP entities (A and B), connected by an UDP link that can be modelled as a lossy FIFO channel.

Since the state space of this case study is very large, five scenarios named *A* to *E* are considered [62], depending on whether each TFTP entity may write and/or read a file (see Table 1). We used the same five scenarios in our study. All of them are specified in LNT [28], in files named `SCENARIO_A.lnt` to `SCENARIO_E.lnt`, as the parallel composition of eight concurrent components named `TFTP_A`, `TFTP_B`, `MEDIUM_A`, `MEDIUM_B`, `RCV_A`, `RCV_B`, `SND_A`, and `SND_B`. All experiments were done on a 64-bit computer with 148 gigabytes of memory.

### 7.1 Generation of minimized LTS

**Direct generation of the compound LTS.** Our aim is to generate a compound LTS that is as small as possible, either for strong or for (div)branching bisimulation. A first way to obtain this LTS is to generate directly and then minimize the compound LTS. For scenario *A* and divbranching equivalence, we use the following SVL script:

```
"SCENARIO_A.bcg" = divbranching reduction of
generation of "SCENARIO_A.lnt"
```

The LTS is small enough to be generated directly, and contains 12,885,069 states and 51,305,563 transitions. Its reduction modulo divbranching bisimulation yields an LTS with 1,620,754 states and 7,060,163 transitions, i.e., between 7 and 8 times less states and transitions, but the reduction already requires more than 1 gigabyte of memory. For the other scenarios, the LTSs obtained are much larger, which justifies resorting to compositional methods.

	Non-minimized		Minimized	
	States	Trans.	States	Trans.
TFTP_A	30,865	263,986	704	4,542
TFTP_B	24,846	213,118	504	3,421
MEDIUM_{A,B}	4,642	9,296	801	5,440
SND_A, RCV_B	1	4	1	4
SND_B, RCV_A	1	4	1	4

Table 2: LTS sizes for scenario *A*, before and after minimization modulo strong, branching, or divbranching bisimulations (component LTSs do not contain internal actions)

	Scenario <i>B</i>		Scenario <i>C</i>		Scenario <i>D</i>		Scenario <i>E</i>	
	States	Trans.	States	Trans.	States	Trans.	States	Trans.
TFTP_A	719	4,610	704	4,542	719	4,610	719	4,610
TFTP_B	504	3,421	1,058	7,164	1,058	7,164	1,079	7,274
MEDIUM_{A,B}	801	5,440	801	5,440	801	5,440	801	5,440
SND_A, RCV_B	1	4	1	7	1	5	1	6
SND_B, RCV_A	1	3	1	7	1	6	1	6

Table 3: Component LTS sizes (states and transitions) once minimized for strong, branching, or divbranching bisimulation (which coincide on this example) for scenarios *B* to *E*

**Component LTS generation.** The first step consists in generating and reducing the LTSs of the components. We use the LNT.OPEN and GENERATOR tools of CADP to generate the LTSs of the components, and BCG\_MIN to minimize them. This can be done easily using SVL statements of the following form:

"TFTP\_A.bcg" = **divbranching reduction of**  
**generation of "SCENARIO\_A.lnt" : TFTP\_A**

or equivalently:

```
% DEFAULT_PROCESS_FILE="SCENARIO_A.lnt"
"TFTP_A.bcg" = divbranching reduction of generation of TFTP_A
```

For scenario *A*, we obtain the LTS sizes of Table 2. Left columns report LTS sizes before reduction and right columns report LTS sizes after reduction modulo strong bisimulation — which coincides with (div)branching bisimulation since components do not have internal actions. For the 4 other scenarios, Table 3 provides the sizes after strong reduction of the LTSs corresponding to the eight components.

**Using an automatically generated interface.** Although the component LTSs are not very large, an interface can be used to constrain the largest component, namely TFTP\_A, for the sake of illustration. In SCENARIO\_A, the refined interface that takes into account constraints imposed on TFTP\_A by SND\_A, generated using EXP.OPEN, has the same number of states and transitions as SND\_A. On-the-fly semi-composition of TFTP\_A with this interface yields an LTS containing 119 states and 320 transitions, which avoids generating the LTS containing 30,865 states and 263,986 transitions. By being two orders of magnitude smaller, this constrained LTS can be minimized more efficiently. Note, however, that this does not decrease the size of the compound LTS, which remains unchanged as expected.

**Generation and minimization of the global compound LTS using root leaf reduction.** Once component LTSs are generated and minimized, they can be composed, yielding a non-minimal (but partially reduced) LTS that can then be further minimized modulo an equivalence relation. This is done by root leaf reduction, the simplest strategy proposed in CADP. All these steps can be achieved using a single SVL statement, in which the LTS composition expression is the same as the composition of processes defined in

Scen.	non-minimized		strong bisimulation		(div)branching bisimulation	
	States	Transitions	States	Transitions	States	Transitions
<i>A</i>	1,905,716	7,950,203	1,785,841	7,695,534	1,620,754	7,060,163
<i>B</i>	851,171	3,503,318	827,396	3,548,292	754,112	3,262,499
<i>C</i>	34,003,384	151,810,170	33,430,548	144,216,908	29,278,308	128,562,334
<i>D</i>	39,908,375	177,531,725	33,205,455	152,203,388	29,140,913	135,963,503
<i>E</i>	19,016,593	77,698,491	18,322,082	78,628,709	15,895,904	69,420,304

Table 4: Size of the LTS corresponding to the whole system for each scenario (component LTSs minimized), non-minimized, once minimized modulo strong bisimulation, and once minimized modulo branching or divbranching bisimulation (which coincide on this example)

Scenario	Generation		Strong minimization		(Div)branching minimization	
	time	memory	time	memory	time	memory
<i>A</i>	19	36	33	167	34	170
<i>B</i>	8	23	12	73	15	74
<i>C</i>	670	434	1,060	3,138	1,138	3,204
<i>D</i>	805	503	1,193	3,766	1,237	3,841
<i>E</i>	326	246	501	1,716	493	1,743

Table 5: Generation and minimization: time (in seconds) and memory (in megabytes) for the LTS corresponding to each scenario

file SCENARIO\_A.lnt:

```
% DEFAULT_PROCESS_FILE="SCENARIO_A.lnt"
"SCENARIO_A.bcg" = root leaf divbranching reduction of
  par
    SEND_A, RECEIVE_A → hide GET_A, PUT_A in TFTP_A end hide
  || SEND_B, RECEIVE_B → hide GET_B, PUT_B in TFTP_B end hide
  || SEND_A, RECEIVE_B → MEDIUM_A
  || SEND_B, RECEIVE_A → MEDIUM_B
  || RECEIVE_A → RECEIVE_A
  || RECEIVE_B → RECEIVE_B
  || SEND_A → SEND_A
  || SEND_B → SEND_B
  end par
```

Table 4 provides LTS sizes after composition, non-minimized and minimized modulo strong, branching, and divbranching bisimulation. Note that branching and divbranching bisimulations coincide, because the global compound LTS does not contain cycles of internal transitions. LTS reductions are not dramatic here, essentially because only a few labels are hidden. Table 5 provides the time and memory used to generate the global compound LTS from the component LTSs, and to minimize it modulo strong bisimulation and (div)branching bisimulation.

**Compositional generation of the state space using other strategies.** More elaborate strategies (namely node reduction and smart reduction) consist in composing the minimized LTSs in a certain order, and minimizing the intermediate LTSs after hiding synchronized labels. For each of these strategies, Table 6 reports both the memory peak and the size of the largest LTS generated for scenario *A*. These strategies can be easily implemented by replacing the `root leaf` keywords in the above SVL script by either `node` or `smart`. Both node and smart reductions are good strategies for this example. The largest LTS generated by smart reduction has the same size as for node reduction, although the composition order differs slightly. In both cases, memory peak (156 megabytes instead of one gigabyte) and largest intermediate LTS size (less

	Largest LTS size		Memory peak (megabytes)
	States	Trans.	
root leaf reduction	1,905,716	7,950,203	170
node reduction	1,783,372	7,827,859	156
smart reduction	1,783,372	7,827,859	156

Table 6: Compositional strategies for Scenario *A* (branching bisimulation)

Scenario	Largest LTS	Time	Memory
<i>A</i>	1,810,925	50	158
<i>B</i>	811,893	25	70
<i>C</i>	32,574,552	1,605	4,492
<i>D</i>	38,567,594	1,907	996
<i>E</i>	17,986,105	755	1,812

Table 7: Smart divbranching reduction (without label hiding) results for the TFTP/UDP case study: largest LTS sizes (in the number of states), CPU time (in seconds), and memory peak (in megabytes)

than 2 million states instead of more than 12 millions) are about six times lower than for direct generation. Table 7 reports largest LTS size (in the number of states), CPU time (in seconds), and peak of memory (in megabytes) used to minimize the LTSs corresponding to the five scenarios modulo divbranching equivalence using smart reduction.

## 7.2 Model checking

We consider the MCL properties named *A01* to *A28*, studied in [62], as well as an additional alternation-2 fairness property *A29* not checked in [62]. These properties are reported in Appendix A (page 55).

**Standard model checking.** We first model check the properties on the BCG graphs corresponding to the five scenarios, minimized modulo divbranching equivalence using the methods presented in Section 7.1. This is done using the tool BCG\_OPEN and the model checker EVALUATOR 3 [104] of CADP. These tools are automatically invoked by SVL script fragments of the following form, where in the formula part (i.e., between symbols “ $\models$ ” and “;”), double quotes enclose character strings denoting a label, such as “REINIT\_A”, and single quotes enclose regular expressions denoting a label, such as ‘RECEIVE\_A.\*’.

```

property A06
  An internal error must cause the transfer to abort. The following formula
  ensures that there is no sequence of transitions in which a reception or a
  transmission can occur after an internal error but before a reinitialisation
  and/or the transmission of an error.
is
  "SCENARIO.bcg"  $\models$ 
  [
    true* .
    "INTERNAL_ERROR_A" .
    not ("REINIT_A" or "SEND_A !ERROR")* .
    'RECEIVE_A.*' or ('SEND_A.*' and not "SEND_A !ERROR")
  ] false;
  expected true
end property

```

Table 8 reports time and memory results for model checking. Some properties being irrelevant to some scenarios (e.g., they concern a read or write operation absent in the corresponding scenario), they are not

Prop	Scenario A		Scenario B		Scenario C		Scenario D		Scenario E	
	time	memory	time	memory	time	memory	time	memory	time	memory
A01	13	106	5	49	700	1,803	714	1,807	330	946
A02	16	110	5	50	840	1,900	867	1,908	386	4
A03	11	100	4	45	582	1,700	599	1,714	257	877
A04	13	106	5	49	700	1,803	711	1,807	330	946
A05	3	34	1	16	52	663	55	693	28	361
A06	12	101	4	47	635	1,734	661	1,755	283	898
A07	12	101	4	47	633	1,734	658	1,755	282	898
A08	11	100	4	46	587	1,704	603	1,718	260	880
A09a							662	1,777	309	927
A09b					692	1,805				
A10					1,018	1,998			410	1,015
A11					884	1,924	1,432	2,239	425	1,032
A12					69	861	57	710	29	393
A13							1,169	2,242	511	1,176
A14	30	135			1,488	2,301			636	1,218
A15			6	46	67	827	55	715	314	921
A16									39	427
A17					171	941	133	923		
A18			4	46	86	856	75	808	265	877
A19			32	96	3,789	3,441	4,925	4,020	1,262	1,687
A20	3	40			143	999			43	465
A21	86	188			2,303	2,782			1,094	1,571
A22			1	16			321	1,191	45	441
A23			22	88			3,062	3,239	1,157	1,660
A24	3	34			120	859				
A25	79	192			2,718	3,028				
A26	13	105			651	1,760			306	921
A27	20	105			1,158	2,091			503	1,112
A28			7	56	1,259	2,147	1,133	2,087	495	1,088
A29	13	106	5	49	698	1,803	714	1,807	331	946

Table 8: Standard model checking results for the TFTP/UDP case study: CPU time (in seconds) and memory (in megabytes)

checked, which explains the shaded cells. As can be seen from the results, a TRUE/FALSE verdict is returned for all verifications, because the model on which they are applied is not too large. However, some of them use much time and memory, e.g., model checking property A19 on scenario D takes 1 hour and 20 minutes and uses more than 4 gigabytes of memory.

**On-the-fly model checking.** We then model check the properties on the fly, i.e., without generating the compound LTS but directly on the composition expression, the transitions being fired on demand by the model checker. This is done still by using EVALUATOR 3, but replacing BCG\_OPEN by EXP.OPEN. These tools are automatically invoked using an SVL fragment similar to the above, but replacing "SCENARIO.bcg" by "SCENARIO.exp", which stores the corresponding composition expression. The time and memory results are reported in Table 9. Note that most often, on-the-fly model checking takes more time and memory than standard model checking, because it includes the resources needed by EXP.OPEN to compute fireable transitions, which is more expensive than enumerating the already computed transitions of a BCG graph. However, in a few cases, on-the-fly model checking is faster and uses less memory, in particular when the portion of the LTS to be explored is smaller than the entire compound LTS. For instance, the time and memory used to check property A15 on scenario D decreases from 55 seconds and 715 megabytes using standard model checking down to 15 seconds and 156 megabytes using on-the-fly model checking.

**Model checking using maximal hiding and smart reduction.** Our next experiment concerns the combination of maximal hiding with smart reduction. For formulas A08, A09a, A09b, A14, and A16, the maximal hiding set is empty, and hence we do not consider them in this experiment. For each of the relevant formulas and for each scenario, the maximal hiding set is generated (using an ad-hoc option of EVALUATOR 4) and then used to minimize the system modulo divbranching bisimulation with smart reduction, and the property is verified on the minimized LTS using EVALUATOR. Maximal hiding is not yet integrated in the SVL scripting language, but this is planned on a short term. Table 10 reports the time

Prop	Scenario A		Scenario B		Scenario C		Scenario D		Scenario E	
	time	memory	time	memory	time	memory	time	memory	time	memory
A01	28	199	10	89	1,324	2,947	1,590	3,351	772	1,530
A02	31	207	12	93	1,640	3,156	2,010	3,631	883	1,612
A03	22	182	8	80	1,210	2,737	1,365	3,162	668	1,386
A04	26	199	10	89	1,400	2,947	1,598	3,351	770	1,530
A05	1	10	1	7	1	7	1	7	1	10
A06	23	187	9	85	1,306	2,808	1,540	3,249	667	1,428
A07	23	187	9	85	1,299	2,808	1,687	3,249	674	1,428
A08	22	186	8	80	1,220	2,745	1,620	3,170	625	1,390
A09a							1,679	3,290	695	1,488
A09b					1,415	2,955				
A10					2,112	3,354			929	1,674
A11					1,722	3,206	3,583	4,444	997	1,711
A12					76	620	8	133	6	101
A13							3,297	4,499	1,446	2,094
A14	54	267			2,681	3,988			1,443	2,107
A15			11	118	55	521	15	156	705	1,524
A16									40	186
A17					315	667	217	569		
A18			9	85	86	476	35	255	599	1,391
A19			53	207	6,159	6,352	9,393	8,753	2,697	3,104
A20	1	31			224	837			39	261
A21	131	374			4,004	4,958			2,293	2,817
A22			1	35			147	427	43	191
A23			39	170			5,605	6,909	2,345	3,039
A24	1	41			148	427				
A25	133	391			4,163	5,480				
A26	25	195			1,383	2,857			687	1,477
A27	38	228			2,323	3,534			1,196	1,871
A28			15	102	2,538	3,654	2,615	4,032	1,277	1,821
A29	26	198	11	88	1,524	2,942	1,738	3,350	700	1,525

Table 9: On-the-fly model checking results for the TFTP/UDP case study: CPU time (in seconds) and memory (in megabytes)

(in seconds) and memory peak (in megabytes). Table 11 reports the largest intermediate LTS size (in the number of states) reached during each experiment. The percentages (columns labelled with %) indicate the ratio between the size of the largest LTS generated during this experiment and the size given in Table 7 of the largest LTS generated during smart divbranching reduction without label hiding. All experiments show a gain, sometimes impressive. For instance, the number of states of the largest LTS generated to check property A12 on scenario D represents only 3 % of the maximum number of states generated by smart divbranching reduction without label hiding. The gain in CPU time and memory peak is also significant.

**Partial model checking.** As the last experiment, we apply the partial model checking approach described in Section 5.2. Table 12 gives, for each scenario and each property, the time in seconds and the peak of memory in megabytes used by partial model checking. The symbol “ $\star$ ” corresponds to verifications that either take too long (exceeding a 12 hour limit) and/or use too much memory (more than 16 gigabytes). Note that most of the time and memory are used by formula simplifications, as compared to the rather low complexity of the synchronous product operation used for quotienting. Partial model checking is not yet integrated in the SVL scripting language, so that tools have to be invoked manually at the moment.

These results confirm that partial model checking may be much more efficient (up to several hundred times less memory in this example) than both standard and on-the-fly model checking. This is particularly true for some formulas having the form “[ $R$ ] false” and “ $\langle R \rangle$  true”, where  $R$  is a regular expression; Such formulas denote the absence (respectively the existence) of a sequence of transitions matching  $R$ . In these examples, the quotient evaluates to true (in the case of formulas of the form “[ $R$ ] false”) or false (in the case of formulas of the form “ $\langle R \rangle$  true”) before all component LTSs have been taken into account in the quotient, because it is possible to determine that none of the paths possible in the parts of the system already explored may yield a path satisfying  $R$  in the compound system. As an illustration, Table 13 gives details on the verification of formula A09b on Scenario C. This formula has the form “[ $R$ ] false” and evaluates to true after the partial model checking steps reported in Table 13.

Prop	Scenario A		Scenario B		Scenario C		Scenario D		Scenario E	
	time	mem.	time	mem.	time	mem.	time	mem.	time	mem.
A01	16	101	14	39	247	1,381	301	1,565	134	701
A02	16	75	10	35	245	1,036	165	765	136	570
A03	6	16	6	6	70	299	160	74	38	159
A04	16	100	10	42	247	1,231	157	996	133	724
A05	11	52	10	20	130	678	83	481	71	377
A06	14	52	12	20	183	690	116	488	100	390
A07	14	52	12	20	183	690	116	488	100	390
A10					272	857			84	588
A11					153	996	227	372	129	439
A12					71	306	169	78	39	164
A13							125	537	103	439
A15			14	45	304	1,199	325	1,334	166	575
A17					125	744	310	489		
A18			9	25	159	1,418	136	508	91	1,040
A19			10	26	303	1,388	243	451	128	1,056
A20	10	70			166	1,973			83	633
A21	13	70			237	1,973			122	633
A22			11	26			229	513	93	653
A23			12	26			267	513	117	653
A24	12	84			164	1,110				
A25	15	84			211	1,110				
A26	13	95			175	1,405			97	762
A27	17	66			1,001	4,492			626	2,262
A28			7	8	1,186	3,953	270	771	454	1,365
A29	15	100	9	42	247	1,231	157	996	133	724

Table 10: Maximal hiding results for the TFTP/UDP case study: time (in seconds) and memory peak (in megabytes)

	Scenario A		Scenario B		Scenario C		Scenario D		Scenario E	
	size	%	size	%	size	%	size	%	size	%
A01	440,679	24 %	152,392	18 %	7,446,443	22 %	9,910,844	25 %	4,207,603	23 %
A02	436,352	24 %	213,665	26 %	7,446,443	22 %	4,379,978	11 %	4,207,603	23 %
A03	293,563	16 %	29,192	3 %	4,793,248	14 %	1,260,400	3 %	2,694,634	14 %
A04	436,352	24 %	213,665	26 %	7,446,443	22 %	4,379,978	11 %	4,207,603	23 %
A05	285,256	15 %	103,739	12 %	4,845,485	14 %	2,798,880	7 %	2,740,106	15 %
A06	341,953	18 %	122,465	15 %	5,561,875	17 %	3,104,457	8 %	3,174,438	17 %
A07	341,953	18 %	122,465	15 %	5,561,875	17 %	3,104,457	8 %	3,174,438	17 %
A10					4,793,248	14 %			2,694,634	14 %
A11					4,793,248	14 %	1,260,400	3 %	2,694,634	14 %
A12					4,819,226	14 %	1,319,036	3 %	2,740,106	15 %
A13							3,201,847	8 %	3,219,910	17 %
A15			305,590	37 %	8,678,547	26 %	8,591,743	22 %	4,097,150	22 %
A17					3,728,837	11 %	2,333,189	6 %		
A18			94,094	11 %	4,793,248	14 %	1,863,667	4 %	2,694,634	14 %
A19			96,945	11 %	4,793,248	14 %	1,863,667	4 %	2,694,634	14 %
A20	293,563	16 %			4,793,248	14 %			2,694,634	14 %
A21	293,563	16 %			4,793,248	14 %			2,694,634	14 %
A22			115,924	14 %			1,474,645	3 %	3,174,438	17 %
A23			115,924	14 %			1,474,645	3 %	3,174,438	17 %
A24	351,927	19 %			5,561,875	17 %				
A25	351,927	19 %			5,561,875	17 %				
A26	351,927	19 %			5,561,875	17 %			3,174,438	17 %
A27	293,563	16 %			4,793,248	14 %			2,694,634	14 %
A28			37,012	4 %	4,793,248	14 %	1,721,736	4 %	2,694,634	14 %
A29	436,352	24 %	213,665	26 %	7,446,443	22 %	4,379,978	11 %	4,207,603	23 %

Table 11: Maximal hiding results for the TFTP/UDP case study: largest LTS size (in the number of states) and percentage w.r.t. the largest LTS size generated by smart reduction without label hiding (see Table 7)

The fairness formula A29 also evaluates more efficiently using partial model checking. This formula, specified in MCL as “ $(\text{true}^*.A_1.(\neg(A_1 \vee A_2)))^*.A_3.(\neg A_1)^*.A_2 @$ ”, denotes the existence of a cyclic sequence of transitions matching the regular expression “ $\text{true}^*.A_1.(\neg(A_1 \vee A_2))^*.A_3.(\neg A_1)^*.A_2$ ”, where  $A_1$ ,  $A_2$ , and  $A_3$  are particular actions. It evaluates to false on all scenarios. The first steps of partial model checking for this formula on Scenario E are detailed in Table 14.

Prop	Scenario A		Scenario B		Scenario C		Scenario D		Scenario E	
	time	memory	time	memory	time	memory	time	memory	time	memory
A01	2	6	3	6	3	24	2	27	3	23
A02	3	6	3	6	6	25	7	28	6	10
A03	1	6	1	6	1	6	1	6	1	6
A04	3	6	3	6	3	6	3	29	3	7
A05	5	6	5	6	5	6	5	6	5	10
A06	3	6	3	6	3	6	3	7	3	6
A07	3	6	3	6	3	6	3	6	3	6
A08	3	6	3	6	3	6	3	6	3	6
A09a							7	28	3	6
A09b					8	6				
A10					3	6			3	6
A11					3	6	1	7	3	6
A12					*	*	*	*	*	*
A13							*	*	*	*
A14	3	6			3	23			3	15
A15			5	15	*	*	*	*	7	59
A16									1	8
A17					*	*	*	*		
A18			1	6	7	11	3	6	1	6
A19			3	6	3	90	3	13	3	55
A20	3	9			6	21			6	25
A21	3	6			3	25			3	25
A22			12	7			2,712	1,271	1,007	650
A23			3	6			9	9	6	40
A24	13	9			3,189	1,786				
A25	3	6			6	40				
A26	3	6			3	15			3	10
A27	3	6			3	6			3	6
A28			3	6	3	22	3	6	3	6
A29	2	7	2	7	6	9	3	7	5	9

Table 12: Partial model checking results for the TFTP/UDP case study: time (in seconds) and memory peak (in megabytes)

Step	States	Transitions
Initial formula graph	13	62
Simplification & reduction	7	56
Quotient wrt. TFTP_A	125	1,964
Simplification & reduction	60	1,512
Quotient wrt. TFTP_B	9,166	69,490
Simplification & reduction	5,308	50,799
Quotient wrt. MEDIUM_B (encodes true)	2	1

Table 13: Successive steps for the partial model checking of property A09b

Step	States	Transitions
Initial formula graph	19	151
Simplification & reduction	7	139
Quotient wrt. TFTP_B	903	20,388
Simplification & reduction	896	20,099
Quotient wrt. TFTP_A	26,369	197,480
Simplification & reduction (encodes false)	1	0

Table 14: Successive steps for the partial model checking of property A29

In a few other cases, partial model checking leads to combinatorial explosion (properties A12, A13, A15, and A17) while other model checking techniques perform efficiently. We illustrate this with the verification



Step	Time	Memory	States	Transitions
Initial formula graph			8	56
Simplification	0	4	8	56
Reduction	0	66	4	52
Quotient wrt. TFTP_A	0	66	210	5,687
Simplification	0	4	136	3,665
Reduction	0	66	134	3,587
Quotient wrt. TFTP_B	0	66	21,172	168,172
Simplification	0	6	21,015	168,172
Reduction	1	66	14,042	119,789
Quotient wrt. MEDIUM_B	14	66	1,648,096	10,327,294
Simplification	35	267	1,648,089	10,327,294
Reduction	72	234	1,551,338	14,773,975
Quotient wrt. MEDIUM_A	686	540	40,572,824	229,050,227

...

Table 15: First successive steps for the partial model checking of property  $A12$  (time in seconds and memory in megabytes)

of formula  $A12$  on scenario  $C$ . This formula has the form “ $\langle R \rangle \text{true}$ ” and evaluates to true. The first steps of partial model checking are detailed in Table 15, which provides the time and memory used to complete each step. The reduction step includes both pre-reduction modulo  $\tau^*.a$  equivalence (i.e., elimination of  $\tau$ -transitions) and minimization modulo strong bisimulation. Note that this may produce a graph that is not minimal in the number of transitions, although always minimal in the number of states. This explosion seems inherent to the structure of the system and the formula, as intermediate quotients need to capture a large part of the behaviour before the truth value of the formula can be determined. These experiments show that partial model checking is complementary to, but does not replace, other model checking techniques.

## 8 Conclusion

This report has given a comprehensive panorama of compositional verification techniques that can be used in an asynchronous, action-based modeling setting. Although such techniques are inherently complex and require significant implementation efforts, they often achieve impressive state-space reduction, successfully tackling verification problems that could not be addressed otherwise. All the approaches presented in this report have been implemented, a task for which the CADP toolbox proved to be a suitable experimentation platform.

It is worth noticing that most of the results presented here are firmly rooted in concurrency theory. They build upon major advances of concurrency theory, namely: (1) formally-defined parallel composition operators that use transition labels as the criterion to decide when concurrent processes must synchronize and when they can interleave; (2) formally-defined abstraction operators that can hide or rename transition labels; (3) preorder and equivalence relations, such as bisimulations, that can be used to compare the behaviors of system models, that can be efficiently computed, and that satisfy the key property of congruence, meaning that such relations are “compatible” with the parallel composition operators; and (4) the existence of modal  $\mu$ -calculus and action-based temporal logics that, under certain conditions, are preserved by these equivalence relations and exhibit suitable properties with respect to parallel composition operators. Although implementing compositional approaches is intrinsically difficult, all these results give concurrency theory a unique advantage to fight state-explosion issues on the long term.

Regarding future work, there are two grand challenges to be met: (i) as the size of models under verification steadily increases, the efficiency of compositional approaches (i.e., the state-space reductions that they allow) should progress as well, and (ii) automation of compositional verification should increase, making it easily available to non-expert users. The latter point is particularly difficult, as there are multiple approaches

to compositional verification, as evidenced in the present report. Given the multiple sources of complexity, not a single compositional approach is sufficient in itself, as complexity can only be mastered by combining different approaches and strategies. In this respect, action-based approaches to compositional verification are probably easier to automate than state-based ones, because the former benefit from the existence of equivalence relations that preserve full classes of properties, whereas the latter must take into account (especially with the *assume-guarantee* paradigm) each particular property of the system under verification.

Among the open issues, one may wonder whether the principles of semi-composition and interfaces (as described in Section 3) could be transposed to formula-dependent approaches (as described in Section 4). Such a combination would be promising, but remains to be investigated. Also, it is worth noticing that all the approaches presented in the present report are based upon the decomposition of a system into concurrent components, according to parallel composition operators used to model that system; orthogonal decompositions could also be considered, in particular those taking into account data structures, especially for systems that are architected around large amounts of shared data.

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## A Temporal logic specifications for the TFTP case study

This appendix presents the temporal logic specifications used in this paper for the TFTP case study. Properties A01 to A28 were taken from [130]. Property A29 was taken from [88]. The 29 properties are given here using the SVL property notation. In the formula part (i.e., between symbols “|=” and “;”), double quotes enclose character strings denoting a label (or part of a label), such as "REINIT\_A" in property A06; single quotes enclose regular expressions denoting a label, such as 'RECEIVE\_A.\*' in property A06; and braces enclose typed expression tuples denoting a label by specifying its various components, such as {RECEIVE\_A !"ACK" ?N : Nat} in property A14.

The following constants are stored in a file named `macros.mcl`, which is automatically included in all the properties below.

```

macro FILE_SIZE_A ()      = 2 of Nat end_macro
macro FILE_SIZE_B ()      = 2 of Nat end_macro
macro MAX_RETRIES_A ()    = 2 of Nat end_macro
macro MAX_RETRIES_B ()    = 4 of Nat end_macro
macro MIN_RETRIES_AB ()   = 2 of Nat end_macro

```

### **property** A01

*The TFTP automaton has two output ports, ARM\_TIMER and STOP\_TIMER, that respectively start and stop the timer used to decide when an incoming message should be considered as lost. The following formula ensures that between two consecutive STOP\_TIMER actions, there must be an ARM\_TIMER action. Otherwise said, it states that, from every reachable state (i.e., after firing any sequence true\* of transitions from the initial state), there exists no sequence of transitions containing two STOP\_TIMER actions with no ARM\_TIMER action in between.*

**is**

```

"SCENARIO.bcg" |=
  [
    true* .
    "STOP_TIMER_A" .
    not ("ARM_TIMER_A")* .
    "STOP_TIMER_A"
  ] false;
expected true
end property

```

**property A02**  
*Between two consecutive ARM\_TIMER actions, there must be a STOP\_TIMER action, a timeout, or a reception. The following formula ensures this by guaranteeing that there exists no sequence of transitions containing two ARM\_TIMER actions without a STOP\_TIMER, a timeout, or a reception in between.*

is

```
"SCENARIO.bcg" |=
  [
    true* .
    "ARM_TIMER_A" .
    not ("STOP_TIMER_A" or "TIMEOUT_A" or 'RECEIVE_A.*')* .
    "ARM_TIMER_A"
  ] false;
  expected true
end property
```

**property A03**  
*The timer cannot be active between two transfers. The following formula ensures there is no ACTIVE\_TIMER\_BETWEEN\_TRANSFERS action reachable.*

is

```
"SCENARIO.bcg" |=
  [ true* . "ACTIVE_TIMER_BETWEEN_TRANSFERS_A" ] false;
  expected true
end property
```

**property A04**  
*A timeout cannot occur between a STOP\_TIMER action and an ARM\_TIMER action. The following formula states that there is no sequence of transitions in which a timeout can follow a STOP\_TIMER action without an ARM\_TIMER action in between.*

is

```
"SCENARIO.bcg" |=
  [
    true* .
    "STOP_TIMER_A" .
    not ("ARM_TIMER_A")* .
    "TIMEOUT_A"
  ] false;
  expected true
end property
```

**property A05**  
*A timeout cannot occur before the first message is sent. The following formula ensures that there is no sequence of transitions in which a timeout occurs before the first send action.*

is

```
"SCENARIO.bcg" |=
  [ not ('SEND_A.*')* . "TIMEOUT_A" ] false;
  expected true
end property
```

**property A06**

An internal error must cause the transfer to abort. The following formula ensures that there is no sequence of transitions in which a reception or a transmission can occur after an internal error but before a reinitialisation and/or the transmission of an error.

```
is
"SCENARIO.bcg" |=
  [
    true* .
    "INTERNAL_ERROR_A" .
    not ("REINIT_A" or "SEND_A !ERROR")* .
    'RECEIVE_A.*' or ('SEND_A.*' and not "SEND_A !ERROR")
  ] false;
  expected true
end property
```

**property A07**

An invalid packet must cause the transfer to abort. The following formula ensures that there is no sequence of transitions in which a reception or a transmission can occur after an invalid packet was received but before a reinitialisation and/or the transmission of an error.

```
is
"SCENARIO.bcg" |=
  [
    true* .
    "INVALID_PACKET_A" .
    not ("REINIT_A" or "SEND_A !ERROR")* .
    'RECEIVE_A.*' or ('SEND_A.*' and not "SEND_A !ERROR")
  ] false;
  expected true
end property
```

**property A08**

When a TFTP protocol entity receives an error, it must abort the current transfer. The original formula ensured that receiving an error cannot be followed by sending an error. It stated that there exists no sequence of transitions in which TFTP entity A can send an error after receiving one, without performing any other action in between. Actually, we used a simpler formula (given below) which, in the TFTP verification setting, is equivalent to the former.

```
is
"SCENARIO.bcg" |=
  [ true* . "RECEIVE_A !ERROR" . "SEND_A !ERROR" ] false;
  expected true
end property
```

**property A09a**

If both protocol entities initiate a transfer at the same time, they must abort upon receiving the other protocol entity's request, in particular, when a transfer request (either read or write) is received after sending a read request. The following formula ensures that there exists no sequence of transitions in which sending a read request and receiving a request can be followed by the transmission of a message until there has been a reinitialisation (transfer succeeded or aborted).

```
is
"SCENARIO.bcg" |=
  [
```

```

    true* .
    'SEND_A !RRQ.*' .
    true . (* ARM_TIMER_A *)
    'RECEIVE_A !RRQ.*' or 'RECEIVE_A !WRQ.*' .
    not ("REINIT_A")* .
    'SEND_A.*')
  ] false;
  expected true
end property

```

**property A09b**

*If both protocol entities initiate a transfer at the same time, they must abort upon receiving the other protocol entity's request, in particular, when a transfer request (either read or write) is received after sending a write request. The following formula ensures that there exists no sequence of transitions in which sending a read request and receiving a request can be followed by the transmission of a message until there has been a reinitialisation (transfer succeeded or aborted).*

```

is
"SCENARIO.bcg" |=
  [
    true* .
    'SEND_A !WRQ.*' .
    true . (* ARM_TIMER_A *)
    'RECEIVE_A !RRQ.*' or 'RECEIVE_A !WRQ.*' .
    not ("REINIT_A")* .
    'SEND_A.*')
  ] false;
  expected true
end property

```

**property A10**

*A process cannot switch from sending data fragments to sending acknowledgements without having received a write request or sent a read request. The following formula ensures that there is no sequence of transitions in which sending a data fragment can be followed by sending an acknowledgement without first either sending a read request or receiving a write request in between.*

```

is
"SCENARIO.bcg" |=
  [
    true* .
    'SEND_A !DATA.*' .
    not ('SEND_A !RRQ.*' or 'RECEIVE_A !WRQ.*')* .
    'SEND_A !ACK.*')
  ] false;
  expected true
end property

```

**property A11**

*A process cannot switch from sending acknowledgements to sending data fragments without having received a read request or sent a write request. The following formula ensures that there is no sequence of transitions in which sending an acknowledgement can be followed by sending a data fragment without first either sending a write request or receiving a read request in between.*

```

is
  "SCENARIO.bcg" |=
    [
      true* .
      'SEND_A !ACK.*' .
      not ('SEND_A !WRQ.*' or 'RECEIVE_A !RRQ.*')* .
      'SEND_A !DATA.*'
    ] false;
  expected true
end property

```

**property A12**

*In the case where TFTP protocol entity A is transferring a file, it must be possible for its transfer to finish.*

```

is
  "SCENARIO.bcg" |=
    < true* . "SUCCESS_A" > true;
  expected true
end property

```

**property A13**

*During the dallying phase, it is possible to begin a new transfer upon receiving a read or write request. The following formula states that for each of the two phases (corresponding to waiting for two timeouts to occur after sending the final acknowledgement), the reception of a request before the timeout occurs can be answered.*

```

is
  "SCENARIO.bcg" |=
    forall X : Nat among {0...1} .
      <
        true* .
        {RECEIVE_A !"DATA" ?N : Nat ?any !TRUE} .
        not ({SEND_A !"ACK" !N})* .
        {SEND_A !"ACK" !N} .
        (not (TIMEOUT_A or REINIT_A)* . TIMEOUT_A) {X} .
        not (TIMEOUT_A or REINIT_A)*
      >
      (
        <
          {RECEIVE_A !"WRQ" ?any} .
          not {RECEIVE_A ...}* .
          {SEND_A !"ACK" !0 of Nat}
        > true
        or
        <
          {RECEIVE_A !"RRQ" ?any} .
          not {RECEIVE_A ...}* .
          {SEND_A !"DATA" !1 of Nat ...}
        > true
      );
  expected true
end property

```

**property A14**

In order to avoid the Sorcerer's Apprentice bug, all resent acknowledgements must be ignored. The following formula ensures that there exists no sequence of transitions in which the same acknowledgement is answered twice without an intervening reinitialisation.

```

is
"SCENARIO.bcg" |=
  [
    true* .
    {RECEIVE_A !"ACK" ?N : Nat} .
    {SEND_A !"DATA" !N + 1 ...} .
    not (REINIT_A or {RECEIVE_A !"ACK" !N})* .
    {RECEIVE_A !"ACK" !N} .
    {SEND_A !"DATA" !N + 1 ...}
  ] false;
  expected true
end property

```

**property A15**

Re-sent data fragment can be acknowledged, to the limit set by the value of the maximum number of retries. The following formula states that after a data fragment is received, it can be received and acknowledged again (only ARM\_TIMER\_A and STOP\_TIMER\_A actions can be performed by TFTP entity A in the meantime) up to MIN\_RETRIES\_AB times (where MIN\_RETRIES\_AB is the minimum of MAX\_RETRIES\_A and MAX\_RETRIES\_B).

```

is
"SCENARIO.bcg" |=
  forall N : Nat among {1 ... FILE_SIZE_A()} .
  <
    true* .
    {RECEIVE_A !"DATA" !N ...} .
    (not ('*_A.*') or '.*TIMER_A.*')* .
    {SEND_A !"ACK" !N} .
    (
      (not ('*_A.*') or '.*TIMER_A.*')* .
      {RECEIVE_A !"DATA" !N ...} .
      (not ('*_A.*') or '.*TIMER_A.*')* .
      {SEND_A !"ACK" !N}
    ) {MIN_RETRIES_AB()}
  > true;
  expected true
end property

```

**property A16**

Every resent read request must be answered, to the limit set by the value of the maximum number of retries. The following formula states that after a read request is received for the first time, it can be received and answered again up to MIN\_RETRIES\_AB times.

```

is
"SCENARIO.bcg" |=
  [
    not {RECEIVE_A !"RRQ" ...}* .
    {RECEIVE_A !"RRQ" ?N : Nat} .
    not {RECEIVE_A ...}* .
    {SEND_A !"DATA" !1 of Nat ...}
  ]
  <

```

```

      (
        not (REINIT_A or {RECEIVE_A !"RRQ" !N}) * .
        {RECEIVE_A !"RRQ" !N} .
        {SEND_A !"DATA" !1 of Nat ...}
      ) {MIN_RETRIES_AB()}
    } true;
  expected true
end property

```

**property A17**

*Every write request must be acknowledged, to the limit set by the value of the maximum number of retries. The following formula states that after a write request is received for the first time, it can be received and answered again up to MIN\_RETRIES\_AB times.*

is

```

"SCENARIO.bcg" |=
  [
    not {RECEIVE_A !"WRQ" ...} * .
    {RECEIVE_A !"WRQ" ?N : Nat} .
    not ({RECEIVE_A ...}) * .
    {SEND_A !"ACK" !0 of Nat}
  ]
  <
    (
      not (REINIT_A or {RECEIVE_A !"WRQ" !N}) * .
      {RECEIVE_A !"WRQ" !N} .
      {SEND_A !"ACK" !0 of Nat}
    ) {MIN_RETRIES_AB()}
  > true;
  expected true
end property

```

**property A18**

*An acknowledgement can be resent as many times as allowed by the value of the maximum number of retries. The following formula states that after an acknowledgement is sent for the first time, it can be sent again (regardless of the reason) up to MAX\_RETRIES\_A times within the same transfer.*

is

```

"SCENARIO.bcg" |=
  forall N : Nat among {0 ... FILE_SIZE_A()} .
  [
    not ({SEND_A !"ACK" !N}) * .
    {SEND_A !"ACK" !N}
  ]
  <
    (
      not ('.*[WR]RQ.*' or {SEND_A !"ACK" !N}) * .
      {SEND_A !"ACK" !N}
    ) {MAX_RETRIES_A()}
  > true;
  expected true
end property

```

**property A19**



An acknowledgement cannot be resent more times than allowed by the value of the maximum number of retries. The following formula states that there is no sequence of transitions in which sending the same acknowledgement can occur more than `MAX_RETRIES_A` times within the same transfer.

```

is
"SCENARIO.bcg" |=
  forall N : Nat among {0 ... FILE_SIZE_A()} .
  [
    true* .
    {SEND_A !"ACK" !N} .
    (
      not ('.*[WR]RQ.*' or {SEND_A !"ACK" !N})* .
      {SEND_A !"ACK" !N}
    ) {MAX_RETRIES_A() + 1}
  ] false;
  expected true
end property

```

**property A20**

A data fragment can be resent as many times as allowed by the value of the maximum number of retries. The following formula states that after a data fragment is sent for the first time, it can be sent again (regardless of the reason) up to `MAX_RETRIES_A` times within the same transfer.

```

is
"SCENARIO.bcg" |=
  forall N : Nat among {1 ... FILE_SIZE_A()} .
  [
    not ({SEND_A !"DATA" !N ...})* .
    {SEND_A !"DATA" !N ...}
  ]
  <
  (
    not ('.*[WR]RQ.*' or {SEND_A !"DATA" !N ...})* .
    {SEND_A !"DATA" !N ...}
  ) {MAX_RETRIES_A()}
  > true;
  expected true
end property

```

**property A21**

A data fragment cannot be resent more times than allowed by the value of the maximum number of retries. The following formula states that there is no sequence of transitions in which sending the same data fragment can occur more than `MAX_RETRIES_A` times within the same transfer.

```

is
"SCENARIO.bcg" |=
  forall N : Nat among {1 ... FILE_SIZE_A()} .
  [
    true* .
    {SEND_A !"DATA" !N ...} .
    (
      not ('.*[WR]RQ.*' or {SEND_A !"DATA" !N ...})* .
      {SEND_A !"DATA" !N ...}
    ) {MAX_RETRIES_A() + 1}
  ] false;
  expected true
end property

```

**property A22**

*A read request can be resent as many times as allowed by the value of the maximum number of retries. The following formula states that after a read request is sent for the first time, it can be sent again (regardless of the reason) up to MAX\_RETRIES\_A times within the same transfer.*

is

```
"SCENARIO.bcg" |=
  [
    not ({SEND_A !"RRQ" ...}) * .
    {SEND_A !"RRQ" ?N : Nat}
  ]
  <
    (
      not (REINIT_A or {SEND_A !"RRQ" !N}) * .
      {SEND_A !"RRQ" !N}
    ) {MAX_RETRIES_A()}
  > true;
  expected true
end property
```

**property A23**

*A read request cannot be resent more times than allowed by the value of the maximum number of retries. The following formula states that there is no sequence of transitions in which sending the same read request can occur more than MAX\_RETRIES\_A without a reinitialisation.*

is

```
"SCENARIO.bcg" |=
  [
    true * .
    {SEND_A !"RRQ" ?N : Nat} .
    (
      not (REINIT_A or {SEND_A !"RRQ" !N}) * .
      {SEND_A !"RRQ" !N}
    ) {MAX_RETRIES_A() + 1}
  ] false;
  expected true
end property
```

**property A24**

*A write request can be resent as many times as allowed by the value of the maximum retries. The following formula states that after a write request is sent for the first time, it can be sent again (regardless of the reason) up to MAX\_RETRIES\_A times within the same transfer.*

is

```
"SCENARIO.bcg" |=
  [ not ({SEND_A !"WRQ" ...}) * . {SEND_A !"WRQ" ?N : Nat} ]
  <
    (
      not (REINIT_A or {SEND_A !"WRQ" !N}) * .
      {SEND_A !"WRQ" !N}
    ) {MAX_RETRIES_A()}
  > true;
  expected true
end property
```

**property A25**

A write request cannot be resent more times than allowed by the value of the maximum number of retries. The following formula states that there is no sequence of transitions in which sending the same write request can occur more than `MAX_RETRIES_A` without a reinitialisation.

```

is
  "SCENARIO.bcg" |=
    [
      true* .
      {SEND_A !"WRQ" ?N : Nat} .
      (
        not (REINIT_A or {SEND_A !"WRQ" !N})* .
        {SEND_A !"WRQ" !N}
      ) {MAX_RETRIES_A() + 1}
    ] false;
  expected true
end property

```

**property A26**

Data fragments must be sent in proper order. The following formula states that any data fragment numbered  $X$  cannot be followed by a data fragment numbered  $Y$ , where  $Y < X$ , unless there has been a reinitialisation in between.

```

is
  "SCENARIO.bcg" |=
    [
      true* .
      {SEND_A !"DATA" ?X : Nat ...} .
      not (REINIT_A)* .
      {SEND_A !"DATA" ?Y : Nat ... where Y < X}
    ] false;
  expected true
end property

```

**property A27**

Between the transmission of two successive data fragments, there must be the reception of the corresponding acknowledgement. The following formula states that there is no sequence of transitions in which the transmission of a data fragment numbered  $X$  can be followed by the transmission of a data fragment numbered  $X + 1$  without the reception of the acknowledgement numbered  $X$  in between.

```

is
  "SCENARIO.bcg" |=
    [
      true* .
      {SEND_A !"DATA" ?X : Nat ?any !FALSE} .
      not ({RECEIVE_A !"ACK" !X})* .
      {SEND_A !"DATA" !X + 1 ...}
    ] false;
  expected true
end property

```

**property A28**

*Between the transmission of two successive acknowledgements, there must be the reception of the corresponding data fragment. The following formula states that there is no sequence of transitions in which the transmission of an acknowledgement numbered  $X$  can be followed by the transmission of an acknowledgement numbered  $X + 1$  without the reception of the data fragment numbered  $X + 1$  in between.*

```

is
  "SCENARIO.bcg" |=
    [
      true* .
      {SEND_A !"ACK" ?X : Nat} .
      not ({RECEIVE_A !"DATA" !X + 1 ...})* .
      {SEND_A !"ACK" !X + 1}
    ] false;
  expected true
end property

```

**property A29**

*There exists a cycle of transitions satisfying the regular expression written in between angles below.*

```

is
  "SCENARIO.bcg" |=
    <
      true* .
      "STOP_TIMER_A" .
      not ("STOP_TIMER_A" or "ARM_TIMER_A")* .
      "TIMEOUT_A" .
      not ("STOP_TIMER_A")* .
      "ARM_TIMER_A"
    > @;
  expected false
end property

```



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