Linearization in parallel pCRL

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Abstract

We describe a linearization algorithm for parallel pCRL processes similar to the one implemented in the linearizer of the $\mu$CRL Toolset. This algorithm finds its roots in formal language theory: the 'grammar' defining a process is transformed into a variant of Greibach Normal Form. Next, any such form is further reduced to linear form, i.e., to an equation that resembles a right-linear, data-parametric grammar. We aim at proving the correctness of this linearization algorithm. To this end we define an equivalence relation on recursive specifications in $\mu$CRL that is model independent and does not involve an explicit notion of solution. © 2001 Elsevier Science Inc. All rights reserved.

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1. Introduction

In this paper we address the issue of linearization of recursive specifications in the specification language micro Common Representation Language ($\mu$CRL, [16,20]). The language $\mu$CRL has been developed under the assumption that an extensive and mathematically precise study of the basic constructs of specification languages is fundamental to an analytical approach of much richer (and more complicated) specification languages such as SDL [33], LOTOS [24], PSF [26,27] and CRL [32]. Moreover, it is assumed that $\mu$CRL and its proof theory provide a solid basis for the design and construction of tools for analysis and manipulation of distributed systems.

The language $\mu$CRL offers a uniform framework for the specification of data and processes. Data are specified by equational specifications: one can declare sorts and functions working upon these sorts, and describe the meaning of these functions by equational axioms. Processes are described in process algebraic style, where the particular process syntax stems from ACP [2,4,14], extended with data-parametric ingredients: there are con-
structs for conditional composition, and for data-parametric choice and communication. As is common in process algebra, infinite processes are specified by means of (finite systems of) recursive equations. In µCRL such equations can also be data-parametric. As an example, for action a and adopting standard semantics for µCRL, each solution for the equation $X = a \cdot X$ specifies (or “identifies”) the process that can only repeatedly execute a, and so does each solution for $Y(n)$ where $Y(n)$ is defined by the data-parametric equation $Y(n) = a \cdot Y(n + 1)$ with $n \in \text{Nat}$. An interesting subclass of systems of recursive equations consists of those that contain only one linear equation. Such a system is called an Linear Process Equation (LPE). Here, linearity refers both to the form of recursion allowed, and to a restriction on the process operations allowed. The above examples $X = a \cdot X$ and $Y(n) = a \cdot Y(n + 1)$ are both LPEs. The restriction to LPE format still yields an expressive setting (for example, it is not hard to show that each computable process over a finite set of actions can be simply defined using an LPE containing only computable functions over the natural numbers, cf. [30]). Moreover, in the design and construction of tools for µCRL, LPEs establish a basic and convenient representation format. This applies, for example, to tools for generation of labeled transition systems, or tools for optimization, deadlock checking, or simulation. The LPE format stems from [7], in which the notion of a process operator is distinguished, and a proof technique for dealing with convergent LPEs is defined. Furthermore, there is a strong resemblance between LPEs and specifications in UNITY [9,11]. The restriction to linear systems has a long tradition in process algebra. For instance, restricting to so-called linear specifications, i.e., linear systems that in some distinguished model have a unique solution per variable, various completeness results were proved in a simple fashion (cf. [5,28]). However, without data-parametric constructs for process specification, the expressiveness is limited: only regular processes can be defined.

The language µCRL is considered to be a specification language because it contains ingredients that facilitate in a straightforward, natural way the modeling of distributed, communicating processes. In particular, it contains constructs for parallelism, encapsulation and abstraction. On the other hand, as mentioned above, LPEs constitute a basic fragment of µCRL in terms of expressiveness and tool support. This explains our interest in transforming any system of µCRL equations into an equivalent LPE, i.e., our interest to linearize µCRL process definitions. In this paper we do not consider full µCRL as the source language for linearization, and allow only a restricted use of the above-mentioned constructs. In [7], pico CRL (pCRL) was defined as a fragment of µCRL. Essentially, pCRL restricts µCRL to the basic operations of process algebra, with data parametric choice, sequential composition and conditionals. Typically, in an LPE only pCRL syntax occurs. Now, as the source language for linearization we take parallel pCRL, an extension of pCRL in which a restricted use of more involved operations, such as $\parallel$ (parallel composition), is allowed. For example, in parallel pCRL the $\parallel$ may not occur in the scope of a recursion. Very often distributed processes have a straightforward definition in parallel pCRL. In [8], a linearization procedure was sketched for a fragment of µCRL, which is similar to parallel pCRL, by means of an informal explanation and examples.

We define the linearization algorithm on an abstract level, but in a very detailed manner. We do not concern ourselves with the question if and in what way systems of recursive equations over parallel pCRL define processes as their unique solutions (per variable). Instead, we argue that the transformation is correct in a more general sense: we show that linearization “preserves all solutions”. This means that if a particular parallel pCRL system of recursive equations defines a series of solutions for its variables in some model, then the LPE resulting from linearization has (at least) the same solutions for the associated process
terms. Consequently, if the resulting LPE is such that one can infer that these solutions are *unique* in some particular (process) model, then both systems define the same processes in that model. In our algorithm, most transformation steps satisfy a stronger property: the set of solutions is the same before and after the transformation. Both the detailed description of the linearization algorithm itself, and the preservation of solutions, which technically speaking is a notion of *implication* between process terms over different μCRL systems, can be considered the contribution of this paper.

To the best of our knowledge, a first description of a transformation of (non-parallel) pCRL into an LPE like format was given in [6]. Transformation procedures from BPA to Greibach Normal Forms were outlined in [1] and presented in [23]. The implications and equivalences of regular systems of recursive equations and recursive program schemes w.r.t. their full sets of solutions were extensively studied by Courcelle in [12,13] and Benson and Guessarian in [3]. The definitions in these papers have a lot in common with our approach, but they could not be directly applied to the μCRL setting.

**Structure of the paper.** In Section 2 we discuss parallel pCRL. Furthermore, we define implication and equivalence between pCRL process terms defined over different pCRL specifications. Sections 3–5 fully describe the linearization procedure. In Section 3 we describe in detail the first part of this transformation, which yields process definitions in so-called extended Greibach normal form. In Section 4 we define the LPE format, and describe the transformation from extended Greibach normal form into this format. Then, in Section 5 we consider the effect of the typical parallel pCRL operations on LPEs. The paper is ended with some conclusions in Section 6. In particular, we provide some comments on our transformation, and relate our approach to other work.

### 2. Description of μCRL and parallel pCRL

In this section we first recall some general information about μCRL. Then we consider (recursive) process definitions in detail, and define various notions of equivalence, among which equivalence between process terms defined over different μCRL specifications. Next, we shortly discuss guardedness and dependency in process definitions. Finally, we introduce pCRL and parallel pCRL as fragments of μCRL.

#### 2.1. Theory of μCRL

First we define the signature and axioms for booleans which are quite standard and can be found for instance in [10, p. 116]. We use equational logic to prove boolean identities. Booleans are obligatory in any μCRL specification.

**Definition 1.** The signature of $\text{Bool}$ consists of constants $t, f$, unary operation $\neg$ and binary operations $\land, \lor, \leftrightarrow$.

**Note 1 (Booleans).** We use infix notation $\neg, \land, \lor, \leftrightarrow$ for $\neg$, $\land$, $\lor$, $\leftrightarrow$ respectively.

**Definition 2.** The axioms of $\text{Bool}$ are the ones presented in Table 1.
Table 1
Axioms of $\text{Bool}$

<table>
<thead>
<tr>
<th>Axiom</th>
<th>Axiom</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x \land y = y \land x$</td>
<td>$x \lor y = y \lor x$</td>
</tr>
<tr>
<td>$(x \land y) \land z = x \land (y \land z)$</td>
<td>$(x \lor y) \lor z = x \lor (y \lor z)$</td>
</tr>
<tr>
<td>$x \land x = x$</td>
<td>$x \lor x = x$</td>
</tr>
<tr>
<td>$x \land (x \lor y) = x$</td>
<td>$x \lor (x \land y) = x$</td>
</tr>
<tr>
<td>$(x \land y) \lor (x \lor z) = x \land (y \lor z)$</td>
<td>$(x \lor y) \land (x \lor z) = x \lor (y \land z)$</td>
</tr>
<tr>
<td>$x \land \neg x = \mathbf{f}$</td>
<td>$x \lor \mathbf{t} = \mathbf{t}$</td>
</tr>
<tr>
<td>$x \leftrightarrow y = (x \land y) \lor (\neg x \land \neg y)$</td>
<td>$x \land \mathbf{t} = \mathbf{t}$</td>
</tr>
</tbody>
</table>

Next we define the generalized equational theory of $\mu$CRL by defining its signature and the axioms. The axioms are taken from, or inspired by Refs. [18,19].

**Note 2 (Vector notation).** Tuples occur a lot in the language, so we use a vector notation for them. Expression $\bar{d}$ is an abbreviation for $d^1, \ldots, d^n$, where $d^k$ are data variables. Similarly, if type information is given, $d: \bar{D}$ is an abbreviation for $d^1:D^1, \ldots, d^n:D^n$ for some natural number $n$. In case $n=0$ the whole vector vanishes as well as brackets (if any) surrounding it. For instance $a(\bar{d})$ is an abbreviation for $a$ in this case (here $a$ is an action, this notion is introduced below). For all vectors $d$ and $\bar{e}$ we have $\bar{d}, \bar{e} = \bar{d}, \bar{e}$. Thus $\bar{d}, \bar{e}$ is an abbreviation for $d^1, \ldots, d^n, e_1, \ldots, e^n$. We also write $\bar{d}, \bar{D}$ & $e:E$ for $d^1:D^1, \ldots, d^n:D^n, e:E$.

For any vector of variables $\bar{d}$, $f(\bar{d})$ is an abbreviation for $f^1(\bar{d}), \ldots, f^m(\bar{d})$ for some $m \in \text{Nat}$ and $\bar{f} = f_1, \ldots, f^n$, where each $f^k(\bar{d})$ is a data term that may contain elements of $\bar{d}$ as free variables. As with vectors of variables, in case $m=0$ the vector of data terms vanishes. We often use $\bar{t}$ to express a data term vector without explicitly denoting its variables.

**Definition 3.** The signature of $\mu$CRL consists of data sorts (or ‘data types’) including $\text{Bool}$ as defined above, and a distinct sort $\text{Proc}$ of processes. Each data sort $D$ is assumed to be equipped with a binary function $eq : D \times D \rightarrow \text{Bool}$. (This requirement can be weakened by demanding such functions only for data sorts that are parameters of communicating actions.) The operational signature of $\mu$CRL is parameterized by the set of action labels $\text{ActLab}$ and a partial commutative and associative function $\gamma : \text{ActLab} \times \text{ActLab} \rightarrow \text{ActLab}$ such that $\gamma(a_1, a_2) \in \text{ActLab}$ implies that $a_1$, $a_2$ and $\gamma(a_1, a_2)$ have parameters of the same sorts. The process operations are the ones listed below:

- Actions $a(\bar{t})$ parameterized by data terms $\bar{t}$, where $a \in \text{ActLab}$ is an action label. More precisely, $a$ is an operation $a : \bar{D} \rightarrow \text{Proc}$.
- Constants $\delta$ and $\tau$ of sort $\text{Proc}$.
- Binary operations $+, \cdot, ||, \Box, |$ defined on $\text{Proc}$, where $|$ is defined using $\gamma$.
- Unary $\text{Proc}$ operations $\delta H, \tau I, \rho R$ for each set of action labels $H$, $I \subseteq \text{ActLab}$ and action label renaming function $R : \text{ActLab} \rightarrow \text{ActLab}$ such that $a$ and $R(a)$ have parameters of the same sorts.
- A ternary operation $\preceq : \text{Proc} \times \text{Bool} \times \text{Proc} \rightarrow \text{Proc}$.
- Binders $\sum_{d:D}$ defined on $\text{Proc}$, for each data variable $d$ of sort $D$.

The partial function $\gamma$ is called a communication function. If $\gamma(a, b) = c$, this indicates that actions with labels $a$ and $b$ can synchronize, becoming action $c$, provided that the data parameters of these actions are equal. The constant $\delta$ represents a deadlocked process and
the constant \( \tau \) represents some internal or hidden activity. The \textit{choice} operator + and the \textit{sequential composition} operator \( \cdot \) are well known. The \textit{merge} operator \( \parallel \) represents \textit{parallel composition}. The \( \| \) (left merge) and \( | \) (communication merge) are auxiliary operations used to equationally define \( \| \). The \textit{encapsulation} operator \( \partial_H(q) \) blocks actions in \( q \) with action labels in the set \( H \), which is especially used to enforce actions to communicate. The \textit{hiding} operator \( \tau_1(q) \) with a set of action labels \( I = \{ a, b, \ldots \} \) hides actions with these labels in \( q \) by renaming them to \( \tau \). The \textit{renaming} operator \( \rho_R(q) \) where \( R \) is a function from action labels to action labels renames each action with label \( a \) in \( q \) an action with label \( R(a) \).

The operator \( p_1 < c > p_2 \) is the \textit{if} \( c \) then \( p_1 \) else \( p_2 \) operator, where \( c \) is an expression of type \( \text{Bool} \). The \textit{sum} operator \( \sum_{d:D} p \) expresses a (potentially infinite) summation \( p[d := d_0] + p[d := d_1] + \cdots \) if data domain \( D = \{ d_0, d_1, \ldots \} \), and \( p[d := d_i] \) is the term \( p \) with all free occurrences of \( d \) replaced by \( d_i \).

**Definition 4.** Axioms of \( \mu \text{CRL} \) are the ones presented in Tables 2–7. We assume that

- \( + \) binds weaker, and \( \cdot \) binds stronger than other operations.
- \( x, y, z \) are variables of sort \( \text{Proc} \).
- \( c, c_1, c_2 \) are variables of sort \( \text{Bool} \).
- \( d, d^1, d^n, d', \ldots \) are data variables (but \( d \) in \( \sum_{d:D} p \) is not a variable).
- \( b \) stands for either \( \text{a}(\overrightarrow{d}) \), or \( \tau \), or \( \delta \).
- \( \overrightarrow{d} = \overrightarrow{d'} \) is an abbreviation for \( \text{eq}(d^1, d'^1) \land \cdots \land \text{eq}(d^n, d'^n) \), where \( \overrightarrow{d} = d^1, \ldots, d^n \) and \( \overrightarrow{d'} = d'^1, \ldots, d'^n \).
- The axioms where \( p \) and \( q \) occur are schemas ranging over all terms \( p \) and \( q \) of sort \( \text{Proc} \), including those in which \( d \) occurs freely.
- The axiom (SUM2) is a scheme ranging over all terms \( r \) of sort \( \text{Proc} \) in which \( d \) does not occur freely.

The axioms in Table 7 (actually only (SC3)) are used only for the parallel composition elimination (Section 5). Note that due to (SC3), the axioms (CM6), (CM9), (CT2), (CD2), (Cond9') and (SUM7') become derivable. The axioms (B1) and (B2) are not used in the transformations described in this paper, so these transformations are also sound in models where these two axioms do not hold.

We use many sorted equational logic for processes and booleans, while other data types can have slightly different proof rules, which may include induction principles, quantifier introduction principles, etc. The proof theory of \( \mu \text{CRL} \) consists of proof rules for the data sorts, the rules of equational logic for the booleans, and the rules of generalized equational logic [18] for the processes. Note that the rules of generalized equational logic do not allow us to substitute terms containing free variables if they become bound. For example, in axiom (SUM1) we cannot substitute \( \text{a}(d) \) for \( x \).

**Definition 5.** Two process terms \( p_1 \) and \( p_2 \) are (unconditionally) \textit{equivalent} (notation \( p_1 = p_2 \)) if \( p_1 = p_2 \) is derivable from the axioms of \( \mu \text{CRL} \) and boolean identities by using many sorted generalized equational logic \( \{ \mu \text{CRL}, \text{BOOL} \} \vdash p_1 = p_2 \). Here \( \text{BOOL} \) is used to refer to the specification of the booleans, and the use of equational logic for deriving boolean identities.

Two process terms \( p_1 \) and \( p_2 \) are \textit{conditionally equivalent} if \( \{ \mu \text{CRL}, \text{BOOL}, \text{DATA} \} \vdash p_1 = p_2 \). Here \( \text{DATA} \) is used to refer to the specification of all data sorts involved, and all proof rules that may be applied.
2.2. Systems of recursion equations

We assume a fixed and infinite set \( \text{Procnames} = \{X, Y, Z, \ldots \} \) of process names with type information associated to them. We extend the sort \( \text{Proc} \) of processes by allowing the process names in \( P \subseteq \text{Procnames} \) as variables of type \( D \rightarrow \text{Proc} \). The terms in the
Table 5
Axioms for sums in μCRL

<table>
<thead>
<tr>
<th>Axiom</th>
<th>Notation</th>
</tr>
</thead>
<tbody>
<tr>
<td>[ \sum_{d:D} x = x ]</td>
<td>(SUM1)</td>
</tr>
<tr>
<td>[ \sum_{e:D} r = \sum_{d:D} (r[e := d]) ]</td>
<td>(SUM2)</td>
</tr>
<tr>
<td>[ \sum_{d:D} p = \sum_{d:D} p + p ]</td>
<td>(SUM3)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (p + q) = \sum_{d:D} p + \sum_{d:D} q ]</td>
<td>(SUM4)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (p \cdot x) = \left( \sum_{d:D} p \right) \cdot x ]</td>
<td>(SUM5)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (p \downarrow x) = \left( \sum_{d:D} p \right) \downarrow x ]</td>
<td>(SUM6)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (p</td>
<td>x) = \left( \sum_{d:D} p \right)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (x</td>
<td>p) = x</td>
</tr>
<tr>
<td>[ \sum_{d:D} (\partial_H(p)) = \partial_H \left( \sum_{d:D} p \right) ]</td>
<td>(SUM8)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (\tau_I(p)) = \tau_I \left( \sum_{d:D} p \right) ]</td>
<td>(SUM9)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (\rho_R(p)) = \rho_R \left( \sum_{d:D} p \right) ]</td>
<td>(SUM10)</td>
</tr>
<tr>
<td>[ \sum_{d:D} (p \triangleleft c \triangledown \delta) = \left( \sum_{d:D} p \right) \triangleleft c \triangledown \delta ]</td>
<td>(SUM12)</td>
</tr>
</tbody>
</table>

signature of μCRL extended with P are further called (μCRL) process terms and the set of all of them is denoted by Terms(P). The free data variables in a process term are those not bound by \( \sum_{d:D} \) occurrences. We write DVar for the set of all free and bound data variables that can occur in a term.

**Definition 6.** A process equation is an equation of the form \( \mathsf{X}(\overrightarrow{d_X}:\overrightarrow{D_X}) = q_\mathsf{X} \), where \( \mathsf{X} \) is a process name with a list of data parameters \( \overrightarrow{d_X}:\overrightarrow{D_X} \), and \( q_\mathsf{X} \) is a process term, in which only the data variables from \( \overrightarrow{d_X} \) may occur freely. We write \( \mathsf{rhs}(\mathsf{X}) \) for \( q_\mathsf{X} \), \( \mathsf{pars}(\mathsf{X}) \) for \( \overrightarrow{d_X} \), and \( \mathsf{type}(\mathsf{X}) \) for \( \overrightarrow{D_X} \).

**Definition 7.** Let \( P \subseteq \text{Procnames} \) be a finite set of process names such that each process name is uniquely typed. A (finite) non-empty set \( G \) of process equations over Terms(P) is called a (finite) system of process equations if each process name in \( P \) occurs exactly once at the left. The set of process names (with types) that appear within \( G \) is denoted as \( |G| \) (so, \( |G| = P \)). We use \( \mathsf{rhs}(\mathsf{X}, G) \), \( \mathsf{pars}(\mathsf{X}, G) \) and \( \mathsf{type}(\mathsf{X}, G) \) to refer to the corresponding parts of the equation for \( \mathsf{X} \) in \( G \).
Table 6
Axioms for renaming operators in CRL

\[ \partial_H(b) = b \text{ if } b = \tau \text{ or } (b = a(d) \text{ and } a \not\in H) \]  
(D1)
\[ \partial_H(b) = \delta \text{ otherwise} \]  
(D2)
\[ \partial_H(x + y) = \partial_H(x) + \partial_H(y) \]  
(D3)
\[ \partial_H(x \cdot y) = \partial_H(x) \cdot \partial_H(y) \]  
(D4)
\[ \partial_H(x \triangleleft c \triangleright \delta) = \partial_H(x) \triangleleft c \triangleright \delta \]  
(D5)
\[ \tau_I(b) = b \text{ if } b = \delta \text{ or } (b = a(d) \text{ and } a \not\in I) \]  
(T1)
\[ \tau_I(b) = \tau \text{ otherwise} \]  
(T2)
\[ \tau_I(x + y) = \tau_I(x) + \tau_I(y) \]  
(T3)
\[ \tau_I(x \cdot y) = \tau_I(x) \cdot \tau_I(y) \]  
(T4)
\[ \tau_I(x \triangleleft c \triangleright \delta) = \tau_I(x) \triangleleft c \triangleright \delta \]  
(T5)
\[ \rho_R(\delta) = \delta \]  
(RD)
\[ \rho_R(\tau) = \tau \]  
(RT)
\[ \rho_R(a(d)) = R(a)(d) \]  
(R1)
\[ \rho_R(x + y) = \rho_R(x) + \rho_R(y) \]  
(R2)
\[ \rho_R(x \cdot y) = \rho_R(x) \cdot \rho_R(y) \]  
(R3)
\[ \rho_R(x \triangleleft c \triangleright \delta) = \rho_R(x) \triangleleft c \triangleright \delta \]  
(R4)

Table 7
Axioms for standard concurrency in \( \mu \)CRL

\[ x|y|z = x|y|z \]  
(SC1)
\[ x|y = y|x \]  
(SC2)
\[ (x|y)|z = x|(y|z) \]  
(SC3)
\[ x|(y|z) = (x|y)|z \]  
(SC4)

Although the original definition of a \( \mu \)CRL specification allows us to have the same process names with different types, we do not treat this possibility here as it would make the explanation only more long-winded.

**Definition 8.** Let \( G \) be a finite system of process equations, \( X \) be a process name in it, and \( \vec{t} \) be a data term vector of type \( type(X, G) \). Then the pair \((X(\vec{t}), G)\) is called a process definition. We use the abbreviation \((X, G)\) for \((X(pars(X, G)), G)\).

**Example 9.** Both \( G_1 = \{X = a \cdot Y, Y = b \cdot X, Z = X||Y\} \) and \( G_2 = \{T(n: Nat) = a(even(n)) \cdot T(S(n))\} \) with \( even : Nat \rightarrow Bool \) as expected and \( S : Nat \rightarrow Nat \) the successor function, are examples of systems of process equations. All of \((X, G_1), (T, G_2), (T(m), G_2)\) are process definitions.

**Definition 10.** Process term \( q \) directly depends on process name \( X \) if this name occurs in \( q \). Process name \( X \) directly depends on process name \( Y \) in a system of process equations \( G \) if \( rhs(X, G) \) directly depends on \( Y \). Process term \( q \) depends on \( X \) in \( G \) if it either directly depends on it, or there is a sequence of process names \( Y_1, \ldots, Y_n = X \) such that \( q \) directly depends on \( Y_1 \) and for each \( i < n \), \( Y_i \) directly depends on \( Y_{i+1} \). Process name \( X \) depends on \( Y \) in \( G \) if \( rhs(X, G) \) depends on it.

We note that the combination of the given data specification with a system \( G \) of process equations determines a \( \mu \)CRL specification in the sense as defined in [20]. Such a specifica-
tion depends on a finite subset \textbf{act} of ActLab and on \textbf{comm}, an enumeration of \( \gamma \) restricted to the labels in \textbf{act}. So a finite system \( G \) implicitly describes a finitary based language.

For a consistent (meaningful) specification, i.e., a \textit{Statically Semantically Correct} specification, it is necessary that all objects are specified only once, that all typing is respected and that the communications in \textbf{comm} are specified in a functional way. Furthermore, the \textit{eq} functions for the data sorts should have the following properties:

\[
\{ \text{DATA, eq}(d, e) = t \} \vdash d = e \quad \text{and} \quad \{ \text{DATA, } d = e \} \vdash \text{eq}(d, e) = t
\]

All data sorts that are introduced during the linearization must have \textit{eq} functions satisfying these properties.

### 2.3. Equivalence of process definitions

We introduce \textit{equivalence} over systems of process equations in a stepwise manner. Let \( G_1 \) and \( G_2 \) be systems of process equations, and assume that the common data sorts of \( G_1 \) and \( G_2 \) are equally defined. Then \( \text{DATA}(G_1, G_2) \) represents all data specifications occurring in \( G_1 \) and \( G_2 \) and all proof rules adopted for these data. We first define (conditional) implication between process terms, and then the equivalence.

In the following definition, derivabilities of the form \( \{ \mu \text{CRL}, \text{BOOL, DATA} \} \cup G_1 \vdash \phi \) are required. In this case, the axioms from \( \mu \text{CRL}, \text{BOOL and DATA} \) may be used to derive \( \phi \), as well as the process equations in \( G_1 \). However, we restrict derivability by requiring that the (data-parametric) process names from \( G_1 \) are considered as (data-parametric) \textit{constants}. For example, if \( G_1 = \{ X = a \cdot X \} \), we may use \( X = a \cdot X \) as an axiom in \( \{ \mu \text{CRL, BOOL, DATA} \} \cup \{ X = a \cdot X \} \vdash \phi \), but \( X \) may not be used as a variable that can be instantiated (e.g., \( \{ \mu \text{CRL, BOOL, DATA} \} \cup \{ X = a \cdot X \} \not\vdash a = a \cdot a \)).

**Definition 11.** Let \( G_1, G_2 \) be systems of process equations with \( |G_1| = \{ X_1, \ldots, X_n \} \) and \( |G_2| = \{ Y_1, \ldots, Y_m \} \). Let furthermore \textit{DATA} be such that it contains \( \text{DATA}(G_1, G_2) \), i.e., \textit{DATA} contains all data sorts and associated proof rules of \( \text{DATA}(G_1, G_2) \).

We say that \((X_1(\vec{t}_1), G_1)\) \textit{conditionally implies} \((Y_1(\vec{t}_2), G_2)\) (notation \((X_1(\vec{t}_1), G_1) \Rightarrow_{c} (Y_1(\vec{t}_2), G_2)\)) for some (possibly open) data term vectors \( \vec{t}_1, \vec{t}_2 \) over \textit{DATA} if for \( j = 1, \ldots, m \) there is a set of mappings \( g_{Y_j} : \text{type}(Y_j) \rightarrow \text{Terms}(\{X_1, \ldots, X_n\}) \) such that

\[
\{ \mu \text{CRL, BOOL, DATA} \} \cup G_1 \vdash X_1(\vec{t}_1) = g_{Y_1}(\vec{t}_2)
\]

and

\[
\forall j \in 1 \ldots m \left( \{ \mu \text{CRL, BOOL, DATA} \} \cup G_1 \vdash g_{Y_j}(d'_j) = \text{rhs}(Y_j) \left[ \forall k \in 1 \ldots m \ Y_k(t') := g_{Y_k}(t') \right] \right)
\]

If \textit{DATA} identities are not used in these derivations we say that \((X_1(\vec{t}_1), G_1)\) \textit{unconditionally implies} \((Y_1(\vec{t}_2), G_2)\) (notation \((X_1(\vec{t}_1), G_1) \Rightarrow (Y_1(\vec{t}_2), G_2)\)). In case \((X(\text{pars}(X, G_1)), G_1)\) \textit{conditionally implies} \((Y(\text{pars}(Y, G_2)), G_2)\) we say that \((X, G_1)\) \textit{conditionally implies} \((Y, G_2)\) (notation \((X, G_1) \Rightarrow (Y, G_2)\)).

The adjective “conditional” could be replaced by “data-dependent”, but we did not do this because it is used similarly in the guardedness definition (See Section 2.4).

We state without proof:
Lemma 12. Let $G_1$ and $G_2$ be systems of process equations, and let the set $H$ of process equations be such that $G_i \cup H$ is a system of process equations ($i = 1, 2$). If $G_1 \Rightarrow G_2$, then $G_1 \cup H \Rightarrow G_2 \cup H$, and if $G_1 \Rightarrow_c G_2$, then $G_1 \cup H \Rightarrow_c G_2 \cup H$.

Definition 13. Process definition $(X(t_1), G_1)$ is equivalent to process definition $(Y(t_2), G_2)$ (notation $(X(t_1), G_1) = (Y(t_2), G_2)$) if both $(X(t_1), G_1) \Rightarrow (Y(t_2), G_2)$ and $(Y(t_2), G_2) \Rightarrow (X(t_1), G_1)$. Similarly, if $(X(pars(X, G_1)), G_1) = (Y(pars(Y, G_2)), G_2)$ we say that $(X, G_1)$ is equivalent to $(Y, G_2)$. The conditional equivalence (notation $=c$) is defined in the same way.

Finally, $G_1 = G_2$ if $|G_1| = |G_2|$ and for all $X \in |G_1|$, $(X, G_1) = (X, G_2)$.

Example 14. Let $G_1 = \{X = a \cdot Y, Y = b \cdot X\}$ and $G_2 = \{X = a \cdot b \cdot X\}$. We can show that $(X, G_1) = (X, G_2)$. The implication from left to the right can be shown by choosing $g_X = X$. The reverse direction can be shown by choosing $g_X = X$ and $g_Y = b \cdot X$.

Example 15. Let $G_1 = \{X(b) = a(b) \cdot X(\neg b)\}$ and $G_2 = \{Y(n:Nat) = a(even(n)) \cdot Y(S(n))\}$. We can show that $(X(t), G_1) \Rightarrow_c (Y(0), G_2)$ by choosing $g_Y(n) = X(even(n))$. In this case we need to show that $X(t) = g_Y(0)$ (which follows from even(0) = t) and that $X(even(n)) = a(even(n)) \cdot X(even(S(n)))$. This latter identity follows from $X(b) = a(b) \cdot X(\neg b)$ and the data identity even(S(n)) = \neg even(n). If we assume the existence of a function $n : \text{Bool} \rightarrow \text{Nat}$, defined by $n(t) = 0$ and $n(f) = 1$, we can prove that $(X(b), G_1) \Rightarrow_c (Y(n(b)), G_2)$ using the same function $g_Y(n)$ and the data identities even(n(b)) = b and even(S(n(b))) = \neg b, both of which seem reasonable.

We do not have any of the reverse implications: consider the model with carrier set Nat, in which $a(b)$ is interpreted as 1, and sequential composition as +. Then $Y(0)$ has many solutions, whereas $X(t)$ has none.

Below we argue that the basic Definition 11 characterizes preservation of solutions. For more details on this subject we refer to [31].

Proposition 16. Let $G_1, G_2$ be systems of process equations with $|G_1| = \{X_1, \ldots, X_n\}$ and $|G_2| = \{Y_1, \ldots, Y_m\}$. Let $(X_1(t_1), G_1) \Rightarrow_c (Y_1(u_1), G_2)$ and let $\mathcal{M}$ be a model of $\mu\text{CRL}$, $\text{Bool}$, $\text{DATA}$ and $G_1$. If $P \in \mathcal{M}$ is a solution for $X_1(t_1)$, then $P$ is also a solution for $Y_1(u_1)$.

Proof. Let $P_1 \in \mathcal{M}$ be a solution for $X_1(t_1)$. So, there are processes $P_i$ ($i = 1, \ldots, n$) that solve the equations of $G_1$ for $X_i(t_i)$. By $(X_1(t_1), G_1) \Rightarrow_c (Y_1(u_1), G_2)$ there are functions $g_{Y_i}$ ($i = 1, \ldots, m$) such that $\mathcal{M} \models X_1(t_1) = g_{Y_i}(u_i)$. Furthermore, the derivability of $g_{Y_{j'}}(d_{j'}) = \text{rhs}(Y_j) \left[vk Y_k(t') = g_{Y_k}(t')\right] (j = 1, \ldots, m)$ yields that $P_1$ is also a solution for $Y_1(u_1)$ in $G_2$. □

The following lemma shows that by applying a $\mu\text{CRL}$ axiom to the right-hand side of an equation we get an equivalent system.
Lemma 17. Let $p_1$, $p_2$ be process terms such that $p_1 = p_2$. Let $G$ be a system of process equations, and $X$ be a process name in it such that $p_1$ is a subterm of $\text{rhs}(X, G)$. Let $G'$ consist of equations in $G$, but in the equation defining $X$ an occurrence of $p_1$ is replaced by $p_2$. Then $G = G'$.

The following lemma shows that by replacing a subterm of the right-hand side of an equation by a fresh process name, and adding the equation for it, we get an equivalent process definition for each process name in the original system.

Lemma 18. Let $G$ be a system of process equations, and $X$ be a process name in it. Let $p$ be a subterm of $\text{rhs}(X, G)$ with free data variables $d^1:D^1, \ldots, d^n:D^n = \vec{d}:\vec{D}$ in it. Let $Y$ be a process name, $Y \notin |G|$. Let $G'$ consist of equations in $G$, but in the equation defining $X$ an occurrence of $p$ is replaced by $Y(\vec{d})$, and the equation $Y(\vec{d}:\vec{D}) = p$ is added to $G$. Then for any $Z \in |G|$ we have $(Z, G) = (Z, G')$.

Proof. To prove that $(Z, G) \Rightarrow (Z, G')$ we take $g_Z(\text{pars}(Z)) = Z(\text{pars}(Z))$ for all $Z \in |G|$, and $g_Y = p$. To prove the other direction we just take $g_Z(\text{pars}(Z)) = Z(\text{pars}(Z))$ for all $Z \in |G|$. □

The following lemma shows that under certain conditions we can substitute a process name by its right-hand side in a right-hand side of an equation.

Lemma 19. Let $G$ be a system of process equations, and $X$ be a process name in it. Let $Y(\vec{t})$ be a subterm of $\text{rhs}(X, G)$ for some $Y \neq X$. Let $G'$ consist of equations in $G$, but in the equation defining $X$ an occurrence of $Y(\vec{t})$ is replaced by $\text{rhs}(Y, G)[\text{pars}(Y, G) := \vec{t}]$. Then we have that $G = G'$.

Proof. In both directions we take the mappings $g_X$ to be the identity mappings. □

The following lemma says that we can add dummy data parameters to a process equation, or remove such parameters.

Lemma 20. Let $G$ be a system of process equations, and $X$ be a process name in it with parameters $d^1, \ldots, d^n$. Suppose that $d^i$ does not occur freely in $\text{rhs}(X, G)$. Let $G'$ be as $G$, but the process name $X$ is replaced by $X'$ and $\text{pars}(X', G') = d^1, \ldots, d^{i-1}, d^{i+1}, \ldots, d^n$. Then for all $Y \in |G| \land Y \neq X$ we have $(Y, G) = (Y, G')$, and $(X(d^1, \ldots, d^n), G) = (X'(d^1, \ldots, d^{i-1}, d^{i+1}, \ldots, d^n), G')$.

Proof. In both directions we take the mappings $g_Y$ (for $Y \neq X$) to be the identity mappings. In one direction $g_{X'}(d^1, \ldots, d^{i-1}, d^{i+1}, \ldots, d^n) = X(d^1, \ldots, d^n)$ and $g_X(d^1, \ldots, d^n) = X'(d^1, \ldots, d^{i-1}, d^{i+1}, \ldots, d^n)$. □

In many cases we are interested in a process definition $(X, G)$ for a fixed process name $X$. The following lemma states that we can drop a defining equation for a process name $Y \neq X$, in cases when the $X$ does not depend on $Y$, and $Y$ does not depend on itself, under the condition that the resulting set of equations will form a system of process equations (Definition 7).
Lemma 21. Let $G$ be a system of process equations, and $X$, $Y$ be process names in it such that $X$ does not depend on $Y$, and $Y$ does not depend on itself. Let $G'$ contain all equations in $G$ except the defining equation for $Y$. If $G'$ is a system of process equations, then we have $(X, G) = (X, G')$.

Proof. In the direction from left to the right we use the identity mapping for $g_Z$. In the reverse direction we use the same mapping, but $g_Y = rhs(Y, G)$. □

2.4. Guardedness

In this paper we use a slightly different notion of guardedness as the one used in [19].

Definition 22. An occurrence of a process name $X$ in a process term $p$ is completely guarded if there is a subterm $p'$ of $p$ of the form $q \cdot p''$ containing this occurrence of $X$, where $q$ is a process term containing no process names.

A process term is called completely guarded if every occurrence of a process name in it is completely guarded. Note that a term that contains no process names is completely guarded.

A system of process equations $G$ is completely guarded if for any $X \in |G|$, $rhs(X, G)$ is a completely guarded term.

Definition 23. A process definition $(X, G)$ is (unconditionally) guarded if there is a process definition $(X', G')$ such that $G'$ is a completely guarded system of process equations, and $(X, G) = (X', G')$.

Definition 24. Let $G$ be a system of process equations. A Process Name Unguarded-Dependency Graph (PNUDG) is an oriented graph with the set of nodes $|G|$, and edges defined as follows: $X \rightarrow Y$ belongs to the graph if $Y$ is not completely guarded in $rhs(X, G)$.

Lemma 25. If the PNUDG of a finite system of process equations $G$ is acyclic, then $G$ is guarded.

Proof. Given a system $G$ we replace each unguarded occurrence of a process name by its right-hand side. By Lemma 19 we get an equivalent system. Due to the fact that PNUDG is acyclic, we need to perform the replacement only finitely many times, and after that we get a completely guarded system. □

The following example shows that the converse of Lemma 25 does not hold.

Example 26. System $G$ consisting of one equation $X = X \triangleleft f \triangleright \delta$ is guarded, but its PNUDG contains the cycle $X \rightarrow X$.

2.5. Parallel pCRL

We define (parallel) pCRL processes as a subset of $\mu$CRL processes.
Definition 27. Let $G$ be a system of process equations. A process term in $\text{Terms}(G)$ is called a pCRL process term in $G$ if it has the syntax

$$
p ::= a(\vec{t}) \mid \delta \mid Y(\vec{t}) \mid p + p \mid p \cdot p \mid \sum_{d:D} p \mid p \triangleleft c \triangleright p \tag{1}
$$

and can directly depend only on process names whose right-hand sides are also pCRL process terms. A process name is called a pCRL process name if its right-hand side is a pCRL process term.

Definition 28. Let $G$ be a system of process equations. A process term in $\text{Terms}(G)$ is called a parallel pCRL process term in $G$ if it has the syntax

$$
q ::= Y(\vec{t}) \mid q \parallel q \mid \partial_H(q) \mid \tau_I(q) \mid \rho_R(q), \tag{2}
$$

and directly depends only on process names whose right-hand sides are pCRL or parallel pCRL process terms. It is called a parallel pCRL process name if its right-hand side is a parallel pCRL process term.

Example 29. Referring to $G_1$ and $G_2$ as defined in the previous Example 9, $X + a$ is a pCRL process term in $G_1$, and $X \parallel X$ and $X \parallel Y$ are parallel pCRL process terms in $G_1$. Furthermore, $P(S(n))$ with $n$ a variable of sort Nat and $a(even(0)) \cdot P(0)$ are pCRL process terms in $G_2$. Finally, $X \parallel a$ is not a (parallel) pCRL process term in $G_1$.

In the following definition we define what a parallel pCRL process definition is. For this definition we assume that we have a $\mu$CRL specification that is Statically Semantically Correct (cf. [20]), that is, in which the data types, actions, communication functions and processes are all well-defined. The first two restrictions posed in the definition below distinguish parallel pCRL as a subset of $\mu$CRL. The third one is present to disallow parallel process names on which the head process name does not depend, and to exclude the presence of certain independent equations in a system. This is not a severe restriction and it simplifies the algorithm presented in Section 5.

Definition 30. Let $G$ be a finite system of process equations, and $(X, G)$ be a process definition. $(X, G)$ is called a parallel pCRL process definition if

- all of the process names in $G$ are either pCRL or parallel pCRL process names;
- no parallel pCRL process name depends on itself;
- process name $X$ depends on all parallel pCRL process names in $G$, but not on itself.

It is called a pCRL system of process equations if all process names in it are pCRL process names.

It follows from Definitions 30 and 28 that for every (parallel) pCRL process definition $(X, G)$, either $X$ is a pCRL process name, or it depends on a pCRL process name in $G$.

Example 31. Referring to $G_1$ as defined in Example 9, $(Z, G_1)$ is a parallel pCRL process definition, but $(X, G_1)$ is not.
3. Transformation to Extended Greibach Normal Form

As the input for the linearization procedure we take a (parallel) pCRL process definition \((X, G)\) such that PNUDG of \(G\) is acyclic. The system of process equations \(G\) can be partitioned in two parts: \(G_1\) and \(G_2\), where \(G_1\) has pCRL equations, and \(G_2\) parallel pCRL equations. \(G_2\) can be empty, in which case \(X\) is a pCRL process name. Otherwise \(X\) is a parallel pCRL process name.

In this section we transform \(G_1\) into a system of process equations \(G'_1\) in Extended Greibach Normal Form (EGNF). The resulting system will contain process equations for all process names in \(|G_1|\) with the same names and types of data parameters involved, as well as, possibly, other process equations. After that we need to linearize the process definition \((X, G')\), where \(G' = G'_1 \cup G_2\). The transformation to EGNF is done via an intermediate form called pre-Extended Greibach Normal Form (pre-EGNF). Adopting the algorithm from Proposition 7.12 in [12] to the setting of pCRL yields a transformation procedure to pre-EGNF. However, for practical purposes such an approach introduces far too many equations, and therefore we define a more refined procedure below.

From this point on we assume that \(\vec{a}(\vec{t})\) with possible indices can also be an abbreviation for \(\tau\). This is done to make the normal form representations more concise.

**Definition 32.** A pCRL process equation is in pre-EGNF iff it is of the form:

\[
X(\overrightarrow{d} : D) = \sum_{i \in I} \sum_{e_i : E_i} p_i(\overrightarrow{d}, \overrightarrow{e_i}) \prec c_i(\overrightarrow{d}, \overrightarrow{e_i}) \triangleright \delta
\]

where \(p_i(\overrightarrow{d}, \overrightarrow{e_i})\) are terms of the following syntax:

\[
p ::= \vec{a}(\vec{t}) \mid \vec{Y}(\vec{t}) \mid \vec{a}(\vec{t}) \cdot p \mid \vec{Y}(\vec{t}) \cdot p
\]

A pCRL process equation is in EGNF iff it is of the form:

\[
X(\overrightarrow{d} : D) = \sum_{i \in I} \sum_{e_i : E_i} a_i(\overrightarrow{f_i}(\overrightarrow{d}, \overrightarrow{e_i})) \cdot p_i(\overrightarrow{d}, \overrightarrow{e_i}) \prec c_i(\overrightarrow{d}, \overrightarrow{e_i}) \triangleright \delta
\]

\[
+ \sum_{j \in J} \sum_{e_j : E_j} a_j(\overrightarrow{f_j}(\overrightarrow{d}, \overrightarrow{e_j})) \prec c_j(\overrightarrow{d}, \overrightarrow{e_j}) \triangleright \delta
\]

where \(I\) and \(J\) are disjoint, and all \(p_i(\overrightarrow{d}, \overrightarrow{e_i})\) are terms of the following syntax:

\[
p ::= \vec{Y}(\vec{t}) \mid \vec{Y}(\vec{t}) \cdot p.
\]

Finally, a finite system of process equations is in (pre-)EGNF iff all its equations are.

**Note 3 (Sum notation).** Apart from functions \(\sum_{d : D} p\) that are included in the syntax of process terms, we use the following abbreviations. Expression \(\sum_{d : D} p\) is an abbreviation for \(\sum_{d_1 : D_1} \cdots \sum_{d_n : D_n} p\). In case \(n = 0\), \(\sum_{d : D} p\) is an abbreviation for \(p\). Expression \(\sum_{i \in I} p_i\), where \(I\) is a finite set, is an abbreviation for \(p_{i_1} + \cdots + p_{i_n}\) such that \(\{i_1, \ldots, i_n\} = I\). In case \(I = \emptyset\), \(\sum_{i \in I} p_i\) is an abbreviation for \(\delta\).

**Note 4 (Conditions).** As follows from the above definition, any process equation in pre-EGNF or EGNF must have a condition in each summand. However, this is not a necessary restriction. In case a summand \(q\) does not have a condition, it is an abbreviation for \(q \triangleleft t \triangleright \delta\).
3.1. Preprocessing

We first transform $G_1$ into $G_1'$. This can be seen as a preprocessing step that possibly renames bound data variables. For instance $\sum_d:D((\sum_e:E a(e)) \cdot b(d))$ is replaced by $\sum_d:D((\sum_e:E a(e)) \cdot b(d))$, where $e$ is a fresh variable. We replace each equation $X(\overrightarrow{d}X) = p_X$ in $G_1$ with the equation $X(\overrightarrow{d}X) = S_0((\overrightarrow{d}X), p_X)$, where $S_0 : DVar \times Terms(|G_1|) \rightarrow Terms(|G_1|)$ is defined in the following way:

$$S_0(S, (\overrightarrow{p}1, \ldots, \overrightarrow{p}n)) \rightarrow f(S_0(S, \overrightarrow{p}1), \ldots, S_0(S, \overrightarrow{p}n)) \text{ if } f \text{ is not } \sum_d:D$$

$$S_0(S, \sum_d:D p) \rightarrow \begin{cases} 
\sum_d:D S_0(S \cup \{d\}, p) & \text{if } d \notin S \\
\sum_e:D S_0(S \cup \{e\}, p[d := e]) & \text{if } d \in S
\end{cases}$$

where $e$ is a fresh variable.

**Proposition 33.** Let $G_1'$ be the result of applying the preprocessing to $G_1$. Then $G_1' = G_1$.

**Proof.** The statement follows from Lemma 17 if we apply axiom (SUM2). \square

As can easily be seen, the preprocessing step does not increase the size or the number of equations in the system.

3.2. Reduction by simple rewriting

By applying term rewriting we get an equivalent set of process equations to the given one, but with terms in right-hand sides having the more restricted form as presented in Table 8.

The rewrite rules that we apply to the right-hand sides of the equations are listed in Table 9. The symbols $\sum_d:D$ are treated in this rewrite system as function symbols, not as binders. This is justified by the fact that we have renamed all nested bound variables, which allows the use of first order term rewriting. We call the function induced by the rewrite rules $rewr : Terms(|G|) \rightarrow Terms(|G|)$ for a given system of process equations $G$.

Before applying the rewriting we eliminate all terms of the form $\overrightarrow{c} \triangleright \overrightarrow{d}$ with the third argument being different from $\delta$ with the following rule:

$$y \neq \delta \Rightarrow x \triangleright \overrightarrow{c} y \rightarrow x \triangleright \overrightarrow{c} \triangleright \delta + y \triangleright \neg \overrightarrow{c} \triangleright \delta$$  \hspace{1cm} (RCOND3)
Table 9
Rewrite rules defining \( \text{rewr} \)

<table>
<thead>
<tr>
<th>Rule</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( x + \delta \rightarrow x )</td>
<td>(RA6)</td>
</tr>
<tr>
<td>( \delta \cdot x \rightarrow \delta )</td>
<td>(RA7)</td>
</tr>
<tr>
<td>( \sum_{d:D} \delta \rightarrow \delta )</td>
<td>(RSUM1)</td>
</tr>
<tr>
<td>( \sum_{d:D} (x + y) \rightarrow \sum_{d:D} x + \sum_{d:D} y )</td>
<td>(RSUM4)</td>
</tr>
<tr>
<td>( \left( \sum_{d:D} x \right) \cdot y \rightarrow \sum_{d:D} (x \cdot y) )</td>
<td>(RSUM5)</td>
</tr>
<tr>
<td>( \left( \sum_{d:D} x \right) \langle c \rangle \delta \rightarrow \sum_{d:D} x \langle c \rangle \delta )</td>
<td>(RSUM12)</td>
</tr>
<tr>
<td>( \delta \langle c \rangle \delta \rightarrow \delta )</td>
<td>(RCOND0)</td>
</tr>
<tr>
<td>( (x \langle e_1 \rangle \delta) \langle e_2 \rangle \delta \rightarrow x \langle e_1 \land e_2 \rangle \delta )</td>
<td>(RCOND4)</td>
</tr>
<tr>
<td>( (x + y) \langle c \rangle \delta \rightarrow x \langle c \rangle \delta + y \langle c \rangle \delta )</td>
<td>(RCOND7)</td>
</tr>
<tr>
<td>( (x \langle c \rangle \delta) \cdot y \rightarrow (x \cdot y) \langle c \rangle \delta )</td>
<td>(RCOND6)</td>
</tr>
</tbody>
</table>

The rewriting is performed modulo the following rules:

\[
\begin{align*}
    x + y &= y + x \\
    x + (y + z) &= (x + y) + z \\
    (x \cdot y) \cdot z &= x \cdot (y \cdot z)
\end{align*}
\]

The optimization rules presented in Table 10 are not needed to get the desired restricted syntactic form, but can be used to simplify the terms. They could be applied with higher priority than the rules in Table 9 to achieve possible reductions. Note that the rule (RSCA') could lead to optimizations only in cases when \( x \) is completely guarded, and \( y \) or \( z \) are not.

**Proposition 34.** The commutative/associative term rewriting system of Table 9 is strongly terminating.

**Proof.** Termination can be proved by using the following order on the operations: \_ &lt; \_ &gt; \cdot &gt; _ &lt; c &gt; \delta > \sum > +. □

**Lemma 35.** For any process term \( p \) not containing \( p_1 \langle c \rangle p_2 \), where \( p_2 \not\equiv \delta \), we have that \( \text{rewr}(p) \) has the syntax defined in Table 8.

Table 10
Optimization rules

<table>
<thead>
<tr>
<th>Rule</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( x + x \rightarrow x )</td>
<td>(RA3)</td>
</tr>
<tr>
<td>( x \langle c \rangle x \rightarrow x )</td>
<td>(RCOND0)</td>
</tr>
<tr>
<td>( x \langle f \rangle y \rightarrow x )</td>
<td>(RCOND1)</td>
</tr>
<tr>
<td>( x \langle e_1 \rangle y \rightarrow y )</td>
<td>(RCOND2)</td>
</tr>
<tr>
<td>( x \langle e_1 \rangle \delta + x \langle e_2 \rangle \delta \rightarrow x \langle e_1 \lor e_2 \rangle \delta )</td>
<td>(RCOND5)</td>
</tr>
<tr>
<td>( (x_1 \langle c \rangle x_2) \cdot (y_1 \langle c \rangle y_2) \rightarrow x_1 \cdot y_1 \langle c \rangle x_2 \cdot y_2 )</td>
<td>(RSCA)</td>
</tr>
<tr>
<td>( x \cdot (y \langle c \rangle z) \rightarrow x \cdot y \langle c \rangle x \cdot z )</td>
<td>(RSCA')</td>
</tr>
</tbody>
</table>
Proof. Let $q = \text{rewr}(p)$. It can be seen from the rewrite rules that they preserve the syntax in Definition 27. Suppose $q$ does not satisfy the syntax defined in Table 8. The following possibilities exist, and all of them imply that $q$ is reducible:

- $q = \delta \cdot p_1$. Can be reduced by (RA7).
- $q = (p_1 \谟 c \谟 \delta) \cdot p_2$. Can be reduced by (RCOND6).
- $q = (\sum_{d:D} p_1) \cdot p_2$. Can be reduced by (RSUM5).
- $q = \delta + p_1$. Can be reduced by (RA6).
- $q = \delta \谟 c \谟 \delta$. Can be reduced by (RCOND0').
- $q = (p_1 + p_2) \谟 c \谟 \delta$. Can be reduced by (RCOND7).
- $q = (p_1 \谟 c_1 \谟 \delta) \谟 c_2 \谟 \delta$. Can be reduced by (RCOND4).
- $q = (\sum_{d:D} p_1) \谟 c \谟 \delta$. Can be reduced by (RSUM12).
- $q = \delta \cdot p_1$. Can be reduced by (RA7).
- $q = (p_1 \谟 c \谟 \delta) \cdot p_2$. Can be reduced by (RCOND6).
- $q = (\sum_{d:D} p_1) \cdot p_2$. Can be reduced by (RSUM5).
- $q = \delta + p_1$. Can be reduced by (RA6).
- $q = \delta \谟 c \谟 \delta$. Can be reduced by (RCOND0').
- $q = (p_1 + p_2) \谟 c \谟 \delta$. Can be reduced by (RCOND7).
- $q = (p_1 \谟 c_1 \谟 \delta) \谟 c_2 \谟 \delta$. Can be reduced by (RCOND4).
- $q = (\sum_{d:D} p_1) \谟 c \谟 \delta$. Can be reduced by (RSUM12).
- $q = \delta \cdot p_1$. Can be reduced by (RA7).
- $q = \delta \谟 c \谟 \delta$. Can be reduced by (RCOND0').
- $q = (p_1 + p_2) \谟 c \谟 \delta$. Can be reduced by (RCOND7).
- $q = (p_1 \谟 c_1 \谟 \delta) \谟 c_2 \谟 \delta$. Can be reduced by (RCOND4).
- $q = (\sum_{d:D} p_1) \谟 c \谟 \delta$. Can be reduced by (RSUM12).

□

Proposition 36. Let $G^2_1$ be the result of applying the rewriting to $G^1_1$. Then $G^2_1 = G^1_1$.

Proof. Taking into account that $G^1_1$ does not contain nested occurrences of bound variables, each rewrite rule is a consequence of the axioms of μCRL. By Lemma 17 we get $G^2_1 = G^1_1$. □

As the result of applying simple rewriting the number of equations obviously remains the same. The process terms may grow with a constant factor, but the number of occurrences of action labels and process names does not increase. The data terms and the number of their occurrences may grow with a constant factor, too.

3.3. Adding new process equations

In this step we reduce the complexity of terms in the right-hand sides of the $G^2_1$ equations even further by the introduction of new process equations. In some cases we take a subterm of a right-hand side and substitute it by a fresh process name parameterized by (at least) all free variables that appear in that subterm. As the result we get a system of process equations $G^3_1$ with equations in pre-EGNF. Such a transformation can be done for all equations $X(d_X:D_X) = p_X$ by replacing them with $X(d_X:D_X) = S_1(d_X:D_X, p_X)$.

$$S_1(S, a(t)) \rightarrow a(t)$$

$$S_1(S, \delta) \rightarrow \delta$$

$$S_1(S, X(t)) \rightarrow X(t)$$

$$S_1(S, p_1 \cdot p_2) \rightarrow S_2(S, p_1 \cdot p_2)$$

$$S_1(S, p_1 + p_2) \rightarrow S_1(S, p_1) + S_1(S, p_2)$$

$$S_1(S, p \谟 c \谟 \delta) \rightarrow S_2(S, p) \谟 c \谟 \delta$$

$$S_1\left(S, \sum_{d:D} p\right) \rightarrow \sum_{d:D} S_1(S & d:D, p)$$

$$S_2(S, a(t)) \rightarrow a(t)$$

$$S_2(S, \delta) \rightarrow (Y := \text{fresh\_var}); \text{add}(Y = \delta)$$
Here \( \text{fresh\_var} \) represents a fresh process name, and \( \text{add} \) represents addition of the equation to the resulting system. Thus formally, \( S_1 \) and \( S_2 \) induce operations \( \hat{S}_1 \) and \( \hat{S}_2 \) that operate on sets of equations and are defined in the expected way (those operations actually transform the system of recursive equations). In the following we provide a simple example of the transformation.

**Example 37.** Let \( G = \{ X(d:D) = a(d) \cdot (b(d) + X(f(d))) \} \) be a given system of process equations. After applying the transformation we get the system \( G' = \{ X(d:D) = a(d) \cdot Y(d), Y(d:D) = b(d) + X(f(d)) \} \) which is in pre-EGNF.

**Proposition 38.** The functions \( S_1 \) and \( S_2 \) are well defined.

**Proof.** Using the order on the operations \( S_1 > +, S_1 > \sum, S_2 > \cdot \) it can be shown that the infinite recursion is not possible for any admissible arguments given. \( \square \)

**Lemma 39.** All process equations in \( G_3^1 \) are in pre-EGNF.

**Proof.** It is easy to see that \( S_2 \) produces terms that satisfy the syntax (3) from Definition 32. The transformation \( S_1 \) can add only \( +, \sum \) or \( \prec \succ \) operations to them at the correct places. The only interesting transformation to consider is \( S_1(S, \sum_{d:D} p) \rightarrow \sum_{d:D} S_1(S & d:D, p) \), as we need to show that \( p \) is not of the form \( p_1 + p_2 \). This follows from the fact that \( p \) satisfies the syntax defined in Table 8. \( \square \)

**Proposition 40.** For any process name \( X \) in \( G_1^3 \) we have \( (X, G_1^3) = (X, G_1^2) \).

**Proof.** The statement follows from Lemma 18. \( \square \)

The transformation described in this section does not increase the size of terms. The number of processes may increase linearly in the size of terms in the original system.

### 3.4. Guarding

Next we transform the equations of \( G_1^3 \) in such a way that each sequential term starts with an action (or \( \tau \)). To this end, we define the function \( \text{guard} : \text{DVar} \times \text{Terms}(|G|) \rightarrow \text{Terms}(|G|) \) in the following way:

\[
\text{guard} \left( S, \sum_{i \in I} \sum_{e_i : E_i} p_i < c_i \succ \delta \right) = \text{rewr} \left( \sum_{i \in I} \sum_{e_i : E_i} \text{guard}(S \cup \{ e_i \}, p_i < c_i \succ \delta) \right)
\]
\( \text{guard}(S, a(\vec{t})) = a(\vec{t}) \)

\( \text{guard}(S, Y(\vec{t})) = \text{guard}(S, S_0 (S \setminus \{\text{pars}(Y)\}, \text{rhs}(Y)) [\text{pars}(Y) := \vec{t}]) \)

\( \text{guard}(S, p_1 \cdot p_2) = \text{rewr}'(\text{guard}(S, p_1) \cdot p_2) \)

Here we use functions \( \text{rewr} \) and \( S_0 \) from the previous subsections. The function \( \text{rewr}' \) represents the rewrite system of \( \text{rewr} \) extended with the following rule.

\[(x + y) \cdot z \rightarrow x \cdot z + y \cdot z \quad \text{(RA4)}\]

**Proposition 41.** For any finite system \( G_1^3 \) with acyclic PNUDG, and any process name \( X \) in it, the function \( \text{guard} \) is well-defined on \( \text{rhs}(X, G_1^3) \).

**Proof.** Let \( n \) be the number of equations in \( G_1^3 \), and \( m \) be the maximal number of process names in sequences \( p_i \) for all \( i \in I \). Suppose that \( \text{guard} \) is applied more than \( n \cdot m \) times on a term. This means that a process name \( Y \) is substituted more than once, which contradicts to the fact that PNUDG is acyclic. \( \square \)

We define the system \( G_1^4 \) in the following way. For each equation

\[ X(d; D) = \sum_{i \in I} \sum_{e_i : E_i} p_i(d, \vec{e}_i) \Leftarrow c_i(d, \vec{e}_i) \triangleright \delta \]

in \( G_1^3 \) we put

\[ X(d; D) = \text{guard} \left( \{d; D\}, \sum_{i \in I} \sum_{e_i : E_i} p_i(d, \vec{e}_i) \Leftarrow c_i(d, \vec{e}_i) \triangleright \delta \right) \]

into \( G_1^4 \).

**Lemma 42.** The equations in \( G_1^4 \) are in pre-EGNF and all sequential process terms in the right-hand sides of its equations start with an action.

**Proof.** Due to Proposition 41 we can apply induction on the definition of \( \text{guard} \). The second and third clauses of the definition are trivial. The first one is brought to the desired form by applying (RCOND4) and (RSUM4) from Table 9. The fourth clause is brought to the desired form by applying (RA4), and then (RSUM5) and (RCOND6) from Table 9. \( \square \)

**Proposition 43.** Let \( G_1^3 \) and \( G_1^4 \) be defined as above. Then \( G_1^3 = G_1^4 \).

**Proof.** According to Lemmas 19 and 17 all transformations performed by \( \text{guard} \) lead to equivalent systems. We note that care has been taken to rename some data variables during the substitution (in the third clause of \( \text{guard} \) definition) in order to make the substitution and the following applications of the axioms sound. \( \square \)

The transformation performed in this step does not increase the number of equations, but their sizes may grow exponentially, due to application of (RA4). An example of such an exponential growth is given below.
Example 44. Let \( n \) be a natural number and let the system of process equations \( G \) contain the following \( n \) equations.

\[
\begin{align*}
X_0 &= a + b \\
& \vdots \\
X_n &= X_{n-1} \cdot a + X_{n-1} \cdot b
\end{align*}
\]

By induction on \( n \) it is easy to show that after applying guarding we get \( X_n = \sum_{p \in \{a, b\}^{n+1}} p \) where \( \{a, b\}^n \) is the set of all strings of length \( n \) consisting of \( a \) and \( b \) occurrences. Indeed, for \( n = 0 \) this is trivial. For \( n > 0 \) we get

\[
\begin{align*}
X_n &= \left( \sum_{p \in \{a, b\}^n} p \right) \cdot a + \left( \sum_{p \in \{a, b\}^n} p \right) \cdot b \\
&= \sum_{p \in \{a, b\}^n} (p \cdot a) + \sum_{p \in \{a, b\}^n} (p \cdot b) \\
&= \sum_{p \in \{a, b\}^{n+1}} p
\end{align*}
\]

This example shows that the term in the right-hand side of the equation for \( X_n \) contains \( 2^n \) summands after the transformation.

3.5. Postprocessing

Finally, we transform all equations of \( G_1^4 \) into EGNF. This transformation can be seen as a simple postprocessing step in which we eliminate all actions that appear not leftmost in the right-hand sides in the equations. This elimination is obtained by introducing a new process name \( X_a \) for each action \( a \) that occurs inside the process terms \( p_i \), with parameters corresponding to those of the action. Thus we add equations \( X_a(d_a; \overrightarrow{D_a}) = a(d_a) \) to the system, and replace the occurrences of the action \( a(\vec{t}) \) by \( X_a(\vec{t}) \).

**Proposition 45.** Let the system \( G'_1 \) of process equations be obtained after the postprocessing of the system \( G_1^4 \) as described above. Then for all \( X \in |G_1^4| \) we have \( (X, G'_1) = (X, G_1^4) \) and \( G'_1 \) is in EGNF.

**Proof.** According to Lemma 18 this transformation is correct and leads to a system that obviously is in EGNF. \( \square \)

As a possible optimization during the postprocessing step, the following slightly different strategy can be applied. If we encounter a subterm \( a \cdot Y \) in \( p_i \), we replace it by a new process name (with the parameters for both \( a \) and \( Y \)), and add the equation for it to the system. This optimization goes along the lines of a so-called regular linearization procedure (see Conclusion), which is a more general case of such an optimization.

**Summary.** In this section we described the transformation of a finite system \( G = G_1 \cup G_2 \) with acyclic PNUDG and \( G_2 \) containing all parallel pCRL process equations into a system \( G' = G'_1 \cup G_2 \) with \( G'_1 \) in EGNF. For each \( X \in |G_1| \),
(X, G_1) = (X, G_1^1) ("Preprocessing", by Proposition 33)
= (X, G_1^2) ("Rewriting", by Proposition 36)
= (X, G_1^3) ("Adding new equations", by Proposition 40)
= (X, G_1^4) ("Guarding", by Proposition 43)
= (X, G_1') ("Postprocessing", by Proposition 45)

By Lemma 12 it follows that (X, G) = (X, G') for each X ∈ |G|.

4. From EGNF to LPE

In this section we transform the system of process equations \( G' = G_1' \cup G_2 \) where \( G_1' \) is in EGNF (cf. Definition 32) into \( G'' = G_1'' \cup G_2' \), where

- \( G_1'' \) consists of a single linear process equation with a specially constructed parameter list;
- if \( G_2 \) is not empty, it is transformed into \( G_2' \) with the same set \(|G_2|\) of process names, but taking the effect of the transformation from \( G_1' \) into \( G_1'' \) into account (references to \( G_1' \) process identifiers may have to be adapted).

Definition 46. A process equation is called a linear process equation (LPE) if it is of the form

\[
X(d; D) = \sum_{i \in I} \sum_{e_i; E_i} a_i(\vec{f}_i(d; e_i)) \cdot X(\vec{g}_i(d, e_i)) \prec c_i(d, e_i) \triangleright \delta \\
+ \sum_{j \in J} \sum_{e_j; E_j} a_j(\vec{f}_j(d, e_j)) \prec c_j(d, e_j) \triangleright \delta
\]

where \( I \) and \( J \) are disjoint sets of indices.

We note that the transformation described in this section is uni-directional, i.e., is formulated in terms of \( \Rightarrow_c \). We again give counterexamples for the associated reverse implications.

4.1. Formal parameters harmonization

In this section we make the formal parameters of all (non-parallel) pCRL process names in \( G_1' \) to be the same, and adapt the parallel pCRL equations in \( G_2 \) in an appropriate way. This is done to be able to compress all (non-parallel) pCRL equations in one process equation. The harmonization is defined by the following steps.

(1) We rename the data variables with the same names, but different types in different processes. This can be easily done (see Section 3.1).
(2) We create the common list of data parameters \( d; D \) by taking the set of all data parameters in the pCRL equations, and giving some order to it.
(3) For each pCRL process name \( X \) in \( G_1' \) we define a mapping \( M_X \) from its parameter list \( D_X \) to the common parameter list \( D \). This mapping is such that each newly created
parameter is a constant. (Recall that a correct μCRL specification contains constants for each declared data sort.)

(4) Then we replace all left-hand sides of the pCRL process equations $X(dx; D_X)$ by $X(d; D)$, and all pCRL process name occurrences $Y(\vec{r})$ in the right-hand sides of all the equations in $G'$ by $Y(M_Y(\vec{r}))$.

**Proposition 47.** Let the system $G^5_1 \cup G^1_2$ of process equations be obtained after harmonization of the system $G'_1 \cup G_2$ as described above. Then for all $X \in |G'_1|$, $(X(d; D), G^5_1) = (X(dx; D_X), G'_1)$, and for all $X \in |G_2|$, $(X, G^5_1 \cup G^1_2) = (X, G'_1 \cup G_2)$.

**Proof.** By Lemma 20 it follows that this transformation yields an equivalent system of equations. □

We remark that a more optimal strategy than ‘global harmonization’ is to merge as many data parameters as possible. This can be achieved by renaming parameters of some processes so that they match the parameters of other processes, and therefore are not introduced in the general parameter list. In this case the number of parameters of some type $s$ in the general list will be the maximal number of parameters of this type in an equation. A drawback of this optimization is the fact that we may lose parameter name information for some process names.

4.2. Making one process equation

Let $G^5_1$ be a system of $n$ pCRL process equations in EGNF with the same formal parameters.

$$X^1(d; D) = \sum_{i \in I^1} \sum_{e_i : E^1_i} a^1_i(\vec{f}^1_i(d, e_i)) \cdot p^1_i(d, e_i) \times c^1_i(d, e_i) \triangleright \delta$$

$$+ \sum_{j \in J^1} \sum_{e_j : E_j} a^1_j(\vec{f}^1_j(d, e_j)) \times c^1_j(d, e_j) \triangleright \delta$$

$$\vdots$$

$$X^n(d; D) = \sum_{i \in I^n} \sum_{e_i : E^n_i} a^n_i(\vec{f}^n_i(d, e_i)) \cdot p^n_i(d, e_i) \times c^n_i(d, e_i) \triangleright \delta$$

$$+ \sum_{j \in J^n} \sum_{e_j : E^n_j} a^n_j(\vec{f}^n_j(d, e_j)) \times c^n_j(d, e_j) \triangleright \delta$$

We define the system $G^6_1$ as a single EGNF process equation in the following way:

$$X(s::State, d; D) = \sum_{i \in I^1} \sum_{e_i : E^1_i} a^1_i(\vec{f}^1_i(d, e_i)) \cdot S(p^1_i(d, e_i)) \times c^1_i(d, e_i) \land s = 1 \triangleright \delta$$

$$+ \sum_{j \in J^1} \sum_{e_j : E^1_j} a^1_j(\vec{f}^1_j(d, e_j)) \times c^1_j(d, e_j) \land s = 1 \triangleright \delta$$
+ \cdots 
+ \sum_{i \in I} \sum_{e_i : E_i} a_i^n(f^n_i(d_i, e_i)) \cdot S(p^n_i(d_i, e_i)) \triangleleft c_i^n(d_i, e_i) \wedge s = n \triangleright \delta 
+ \sum_{j \in J} \sum_{e_j : E_j} a_j^n(f^n_j(d_j, e_j)) \triangleleft c_j^n(d_j, e_j) \wedge s = n \triangleright \delta 

where \( S(X^s(\vec{a})) = X(s, \vec{a}) \), and \( S(X^s(\vec{a}) \cdot p) = X(s, \vec{a}) \cdot S(p) \).

The data type \textit{State} is an enumerated data type with equality predicate. Natural numbers are normally used for \textit{State}, though a finite data type is, of course, sufficient.

Let the system \( G_1^5 \cup G_1^6 \) of process equations be obtained after harmonization of the system \( G_1' \cup G_2 \) as described above. Then for all \( X \in |G_1'| \), \((X(d: \vec{D}), G_1^5) = (X(d: \vec{D}), G_1')\), and for all \( X \in |G_2| \), \((X, G_1^5 \cup G_1^6) = (X, G_1' \cup G_2)\). During the current step we construct the system \( G_1^6 \) consisting of the single equation for \( X \) and the set \( G_2^1 \) being \( G_1^6 \) with all pCRL process terms \( X' \) replaced by \( X(i, \vec{a}) \) for each \( 1 \leq i \leq n \).

\textbf{Proposition 48.} \textit{Let} \( G_1^5 \) be a system of \( n \) process equations in EGNF, each with formal parameters \( d: \vec{D} \), and let State enumerate \( 1, \ldots, n \). Let furthermore \( G_1^5 \cup G_1^2 \) be a system of parallel pCRL process equations and \( G_1^6 \cup G_2^2 \) be the result of the transformation described above. Then for any \( s: \text{State} \), data term vector \( \vec{a} \), and any \( X \in |G_1^5| \), \((X(s, \vec{a}), G_1^6) =_c (X(s, \vec{a}), G_1^5) \).

\textbf{Proof.} The equivalence is easy to derive with the following functions: \( g_X^i(\vec{a}) = X(i, \vec{a}) \) for each \( i: \text{State} \), and \( g_X(s, \vec{a}) = X(s, \vec{a}) \). Note that identities of sort \textit{State} are used in the derivations. \( \square \)

4.3. Introduction of a stack

The final step in the linearization of pCRL processes consists of the introduction of a stack parameter which allows us to model a sequential composition of process names with parameters as a single process term. In the case that such sequential compositions do not occur in the equation, we do not apply this step. For the particular transformation described here, it is necessary that the process equation to be transformed is data-parametric. This need not be the case after application of all preceding transformation steps. For instance the equation \( X = a \cdot X \cdots X + b \) does not have a data parameter. In this case we need to add a dummy data parameter (over a singleton data type, cf. Lemma 20) to apply the following transformation.

Let \( G_1^6 \) be a single pCRL process equation in EGNF:

\[ X(d: \vec{D}) = \sum_{i \in I} \sum_{e_i : E_i} a_i(f_i(d, e_i)) \cdot X(t_i^1) \cdots X(t_i^m) \triangleleft c_i(d, e_i) \triangleright \delta \]

\[ + \sum_{j \in J} \sum_{e_j : E_j} a_j(f_j(d, e_j)) \triangleleft c_j(d, e_j) \triangleright \delta \]

We define \( G_1'' \) by the single process equation for \( Z \) in the following way:
The result of an insertion to another value of this type.

Implementing data types in a standard stack data type with constructors \( \{ \} \) representing the empty stack, and \( \text{push}(\vec{t}, \vec{e}) \) inserting the new element \( \vec{e} \) to the top of the stack \( \vec{t} \). We use the equality predicate on stacks, but a predicate that checks if a stack is empty can be used instead. The function \( \text{get}_i(\vec{t}) \) returns the \( i \)th element of the top of \( \vec{t} \), and the function \( \text{pop}(\vec{t}) \) returns the stack value \( \vec{t} \) without its top element. See [22] for details on implementing data types in \( \mu \text{CRL} \). To prove the following proposition we use an induction principle on the data type \( \text{Stack} \), namely that every value of type stack is either empty or the result of an insertion to another value of this type.

During the current step we construct the system \( G''_1 \) consisting of the single equation for \( X \) and the set \( G'_2 \) with all \( p \text{CRL} \) process terms \( X(\vec{t}) \) replaced by \( Z(\vec{t}) \).

**Proposition 49.** Let systems \( G^6_1 \) and \( G''_1 \) as described above be given. Then for any data term vector \( \vec{t} \) we have \( (X(\vec{t}), G^6_1) \Rightarrow_c (Z(\vec{t}), \vec{t}), G''_1) \). Let furthermore \( G^6_1 \cup G'_2 \) be a system of parallel \( p \text{CRL} \) process equations and \( G''_1 \cup G'_2 \) be the result of the transformation described above. Then for any \( X \in |G'_2| \), \( (X, G^6_1 \cup G'_2) \Rightarrow_c (X, G''_1 \cup G'_2) \).

**Proof.** We define \( g_2(\vec{t}, \vec{e}) = X(\vec{t}) \cdot \text{get}(\vec{t}) \). To prove the implication we consider two cases. First, if the stack \( \vec{t} \) is empty we have \( g_2(\vec{t}, \vec{e}) = X(\vec{t}) \).

It can be shown by induction on \( n \) that

\[
g_2(\text{push}(\vec{t}^1, \ldots, \text{push}(\vec{t}^n, \vec{t}^0)), \vec{e}) = X(\vec{t}^1) \cdot \cdots \cdot X(\vec{t}^n)
\]

When we apply this \( g_2 \) to the equation for \( Z \) and use the identities of the sort \( \text{Stack} \), we get an identity which is the same for the equation for \( X \).

In the second case, if the stack \( \vec{t} = \text{push}(\vec{t'}, \vec{t}^0) \) for some stack value \( \vec{t'} \) and data term vector \( \vec{t}^0 \), we have \( g_2(\vec{t}, \vec{e}) = X(\vec{t'}) \cdot g_2(\vec{t'}, \vec{e}) \). By induction on \( n \) it can be shown that

\[
g_2(\text{push}(\vec{t}^1, \ldots, \text{push}(\vec{t}^n, \vec{t}^0)), \vec{t}^0) = X(\vec{t}^1) \cdot \cdots \cdot X(\vec{t}^n) \cdot g_2(\vec{t'}, \vec{t}^0)
\]

When we apply this \( g_2 \) to the equation for \( Z \) and use the identities of the sort \( \text{Stack} \), we get the following identity:

\[
X(\vec{t}) \cdot g_2(\vec{t'}, \vec{t}^0)
\]

\[
= \sum_{i \in I} \sum_{e_i : E_i} a_i(\vec{t}_i, \vec{e}_i) \cdot X(\vec{t}_i) \cdot \cdots \cdot X(\vec{t}^n) \cdot g_2(\vec{t'}, \vec{t}^0) < c_i(\vec{d}, \vec{e}_i) \triangleright \delta
\]

\[
+ \sum_{j \in J} \sum_{e_j : E_j} a_j(\vec{f}_j, \vec{e}_j) \cdot g_2(\vec{t'}, \vec{t}^0) < c_j(\vec{d}, \vec{e}_j) \triangleright \delta
\]

where \( \text{get}(\vec{t}) = \text{get}_1(\vec{t}), \ldots, \text{get}_n(\vec{t}) \).

The data type \( \text{Stack} \) is a standard stack data type with constructors \( \{ \} \) representing the empty stack, and \( \text{push}(\vec{t}, \vec{e}) \) inserting the new element \( \vec{e} \) to the top of the stack \( \vec{t} \). We use the equality predicate on stacks, but a predicate that checks if a stack is empty can be used instead. The function \( \text{get}_i(\vec{t}) \) returns the \( i \)th element of the top of \( \vec{t} \), and the function \( \text{pop}(\vec{t}) \) returns the stack value \( \vec{t} \) without its top element. See [22] for details on implementing data types in \( \mu \text{CRL} \). To prove the following proposition we use an induction principle on the data type \( \text{Stack} \), namely that every value of type stack is either empty or the result of an insertion to another value of this type.
This identity is derivable from the equation for $X$ by applying axioms (A4), (SUM5) and (Cond6).

The following example [29] shows that the reverse implication does not hold in every model. It is easy to see that if data parameters do not matter, the stack is isomorphic to a counter which can be implemented by means of natural numbers.

**Example 50.** Let $G_1 = \{X = a \cdot X \cdot X\}$ and $G_2 = \{Z(n: \text{Nat}) = a \cdot Z(\text{succ}(n))\}$. Consider the model with integers $\mathbb{Z}$ as the carrier set, and the operations $\cdot \to +, a \to -1$. The equation in $G_1$ has the unique solution $X = 1$, while the equation in $G_2$ has infinitely many solutions $Z(n) = n + c$, where $c \in \mathbb{Z}$. For a more elaborated model that includes interpretations of other $\mu$CRL operations see Example 52.

**Summary.** This section is about the transformation of a finite system $G' = G'_1 \cup G''$ with acyclic PNUDG and $G'_1$ in EGNF into a system $G'' = G''_1 \cup G''_2$ with $G''_1$ an LPE and $G''_2$ appropriately updated. For each $X \in |G'|$,

\[
(X, G') = (X', G''_1 \cup G''_2) \quad (\text{"Harmonization", by Proposition 47})
\]

\[
=_{c} (X'', G''_1 \cup G''_2) \quad (\text{"One equation", by Proposition 48})
\]

\[
\Rightarrow_{c} (X''', G'') \quad (\text{"One LPE", by Proposition 49})
\]

Here the primed versions of $X$ represent the possible updates of parameters, as prescribed by the propositions mentioned.

5. **From parallel pCRL to LPE**

As the result of the previous section we have obtained $G'' = G''_1 \cup G''_2$, where $G''_1$ is an LPE and $G''_2$ a (possibly empty) set of parallel pCRL process equations. In this section we show that the parallel part of $G''$ can be eliminated. First we take a general point of view, and show that LPEs are closed under the parallel pCRL process operations, viz. parallel composition, encapsulation, hiding, and renaming (see Definition 28). Then we show that with these results and those from Sections 3 and 4, the transformation of $G''$ into a single LPE can be carried out. We note that the transformation described in this section is uni-directional, and we give counterexamples for the associated reverse implications.

5.1. **Parallel composition of LPEs**

Let $G$ be a system of process equations in which each of $(X(dX), G)$ and $(Y(dY), G)$ is defined by an LPE, and that contains an equation $Z(dX, dY) = X(dX) \parallel Y(dY)$. Assume that the LPEs for $X$ and $Y$ have no common data variables, and are defined in the following way:

\[
X(dX; D_X) = \sum_{i \in I} \sum_{e_i: E_i} a_i(f_i(dX, e_i)) \cdot X(g_i(dX, e_i)) \leq c_i(dX, e_i) \triangleright \delta
\]

\[
+ \sum_{j \in J} \sum_{e_j: E_j} a_j(f_j(dX, e_j)) \leq c_j(dX, e_j) \triangleright \delta
\]
\[
\gamma(\overrightarrow{dy}:\overrightarrow{dY}) = \sum_{i \in I'} \sum_{e_i \in E_i} a_i \left( \overrightarrow{f_i} (\overrightarrow{dx}, \overrightarrow{e_i}) \right) \cdot \gamma(\overrightarrow{g_i} (\overrightarrow{dx}, \overrightarrow{e_i})) < c_i (\overrightarrow{dx}, \overrightarrow{e_i}) \triangleright \delta \\
+ \sum_{j \in J'} \sum_{e_j \in E_j} a_j \left( \overrightarrow{f_j} (\overrightarrow{dy}, \overrightarrow{e_j}) \right) \cdot \gamma(\overrightarrow{dY}) < c_j (\overrightarrow{dy}, \overrightarrow{e_j}) \triangleright \delta
\]

where \( I \cap J = I' \cap J' = \emptyset \). We construct the equation for \( Z(\overrightarrow{dx}:\overrightarrow{dX}, \overrightarrow{dy}:\overrightarrow{dY}) \), being equal to \( X(\overrightarrow{dx})||Y(\overrightarrow{dY}) \), as follows.

\[
Z(\overrightarrow{dx}:\overrightarrow{dX}, \overrightarrow{dy}:\overrightarrow{dY}) = \sum_{i \in I} \sum_{e_i \in E_i} a_i \left( \overrightarrow{f_i} (\overrightarrow{dx}, \overrightarrow{e_i}) \right) \cdot Z(\overrightarrow{g_i} (\overrightarrow{dx}, \overrightarrow{e_i}), \overrightarrow{dY}) < c_i (\overrightarrow{dx}, \overrightarrow{e_i}) \triangleright \delta \\
+ \sum_{j \in J} \sum_{e_j \in E_j} a_j \left( \overrightarrow{f_j} (\overrightarrow{dy}, \overrightarrow{e_j}) \right) \cdot \gamma(\overrightarrow{dY}) < c_j (\overrightarrow{dy}, \overrightarrow{e_j}) \triangleright \delta
\]

\[
\gamma(a_k, a'_k)(\overrightarrow{f_k} (\overrightarrow{dx}, \overrightarrow{e_k})) \cdot Z(\overrightarrow{g_k} (\overrightarrow{dx}, \overrightarrow{e_k}), \overrightarrow{dY}) < c_k (\overrightarrow{dx}, \overrightarrow{e_k}) \triangleright \delta
\]

where \( P \gamma Q = \{ (p, q) \in P \times Q \mid \gamma(a_p, a'_q) \) is defined\).

**Proposition 51.** Let \( G' \) contain the equations for \( X, Y \) and \( Z \) defined above. Let \( G \) contain the equations for \( X \) and \( Y \), and the equation \( Z(\overrightarrow{dx}:\overrightarrow{dX}, \overrightarrow{dy}:\overrightarrow{dY}) = X(\overrightarrow{dx})||Y(\overrightarrow{dY}) \). Then \( (Z, G) \Rightarrow (Z, G') \).

**Proof.** We use the identity mapping for \( g_X, g_Y, g_Z \). Then the equations for \( X \) and \( Y \) are proven trivially because they are the same in \( G \) and \( G' \). To prove the equation for \( Z \) first
apply the axiom (CM1) to get \( Z = (X(\overrightarrow{d_x})||Y(\overrightarrow{d_y}) + Y(\overrightarrow{d_y})||X(\overrightarrow{d_x})) + X(\overrightarrow{d_x})|Y(\overrightarrow{d_y}) \). Then we replace \( X \) and \( Y \) in the left hand sides of \( \parallel \) and in both sides of \( | \) by their right-hand sides. After that we apply the axioms (CM4), (SUM6), (Cond8), (CM2) and (CM3) to eliminate \( \parallel \), and the axioms (CM8), (CM9), (SUM7), (SUM7'), (Cond9), (Cond9'), (CM5), (CM6), (CM7), (CF1), (CF2), (CT1), (CT2), (CD1), and (CD2) to eliminate \( | \). Note that before applying the axioms for sums we might need to apply (SUM2), and after elimination \( \parallel \) and \( | \) we might need to apply (A7) and (A6). After that we apply the identity \( x||y = y||x \), which is derivable from axioms (CM1), (A1) and (SC3), to replace all occurrences of \( Y(\overrightarrow{t_x})||X(\overrightarrow{t_y}) \) by \( X(\overrightarrow{t_x})||Y(\overrightarrow{t_y}) \), and finally we replace all \( X(\overrightarrow{t_x})||Y(\overrightarrow{t_y}) \) by \( Z(\overrightarrow{t_x}) \) using the equation for \( Z \) in \( G \). As the result we get the equation for \( Z \) in \( G' \). □

In the following example we present a model of \( \mu \)CRL based on the trace model [15], but in which the sequential composition operation is commutative and idempotent. This model is used in Example 53 to show that the reverse implication of Proposition 51 does not hold in every model.

**Example 52.** Let \( \text{ActLab} \) be a finite set of action labels and \( \gamma \) be the totally undefined function. Consider the model with carrier set \( \left( 2^{(\text{ActLab} \setminus \{ \emptyset \})} \cup \{ T, \bot \} \right) \), and the operations defined as follows:

- For each \( a \in \text{ActLab} \) \( a(\overrightarrow{t}) \rightarrow \{ \{a\} \} \).
- \( \delta \rightarrow T \) and \( \tau \rightarrow \bot \).
- \( + \rightarrow \cup \), where \( S \cup T = T \cup S = S \) and \( S \cup \bot = \bot \cup S = \bot \).
- \( \cdot, ||, \mid \rightarrow \ast \), where \( S \ast S' = \{ s \cup s' \mid s \in S \land s' \in S' \} \), \( S \ast T = T \ast S = T \) and \( S \ast \bot = \bot \ast S = S \).
- \( \partial_H \rightarrow e_H \), where \( e_H([[\{a\}]]) = [[\{a\}]] \) if \( a \notin H \), \( e_H([[\{a\}]]) = T \) if \( a \in H \), \( e_H(S \cup S') = e_H(S) \cup e_H(S') \), \( e_H(S \ast S') = e_H(S) \ast e_H(S') \), \( e_H(T) = T \), \( e_H(\bot) = \bot \).
- \( \tau_I \rightarrow h_I \), where \( h_I \) is defined in a similar way as \( e_H \).
- \( \sum_{d:D} \rightarrow id \), where \( id \) is the identity mapping.
- \( x \ast c \ast y \rightarrow if(c, x, y) \), where \( if(c, x, y) \) is the if-then-else mapping.

**Example 53.** Let \( G = (X = a \cdot X, Y = b \cdot Y, Z = X \parallel Y) \) and \( G' = (X = a \cdot X, Y = b \cdot Y, Z = a \cdot Z + b \cdot Z) \). In the model defined in Example 52 the equations for \( X \) in both \( G \) and \( G' \) have the following solutions:

\[
\{ \{a\} \}, \quad \{ \{a, b\} \}, \quad \{ \{a\}, \{a, b\} \}, \quad T
\]

while the equations for \( Y \) have the following solutions:

\[
\{ \{b\} \}, \quad \{ \{a, b\} \}, \quad \{ \{b\}, \{a, b\} \}, \quad T
\]

The equation for \( Z \) in \( G \) has two solutions \( \{ \{a, b\} \} \) and \( T \), while the equation for \( Z \) in \( G' \) has five solutions \( \{ \{a, b\} \}, \{ \{a\}, \{a, b\} \}, \{ \{b\}, \{a, b\} \}, \{ \{a\}, \{b\}, \{a, b\} \} \) and \( T \).

5.2. **Encapsulation, hiding and renaming of LPEs**

Let \( G \) be an LPE defining \( X \) as in the previous section, \( A \) be a set of action labels, and \( R \) be a renaming function. We construct LPEs for \( Z_1 \) being equal to \( \partial_A(X) \), \( Z_2 \) being equal to \( \tau_A(X) \), and \( Z_3 \) being equal to \( \rho_R(X) \), in the following way:
\[ Z_1(d_X;D_X) = \sum_{i \in I} \sum_{e_i;E_i} a_i(f_i(d_X, e_i)) \cdot Z_1(g_i(d_X, e_i)) < c_i(d_X, e_i) \triangleright \delta \]
\[ + \sum_{j \in J} \sum_{e_j;E_j} a_j(f_j(d_X, e_j)) < c_j(d_X, e_j) \triangleright \delta \]

Here and in the equations below we assume that \( I_1 = \{ i \in I \mid a_i \notin A \} \) and \( J_1 = \{ j \in J \mid a_j \notin A \} \).

\[ Z_2(d_X;D_X) = \sum_{i \in I} \sum_{e_i;E_i} a_i(f_i(d_X, e_i)) \cdot Z_2(g_i(d_X, e_i)) < c_i(d_X, e_i) \triangleright \delta \]
\[ + \sum_{j \in J} \sum_{e_j;E_j} a_j(f_j(d_X, e_j)) < c_j(d_X, e_j) \triangleright \delta \]
\[ + \sum_{i \in I} \sum_{e_i;E_i} \tau \cdot Z_2(g_i(d_X, e_i)) < c_i(d_X, e_i) \triangleright \delta \]
\[ + \sum_{j \in J} \sum_{e_j;E_j} \tau < c_j(d_X, e_j) \triangleright \delta \]

\[ Z_3(d_X;D_X) = \sum_{i \in I} \sum_{e_i;E_i} R(a_i)(f_i(d_X, e_i)) \cdot Z_3(g_i(d_X, e_i)) < c_i(d_X, e_i) \triangleright \delta \]
\[ + \sum_{j \in J} \sum_{e_j;E_j} R(a_j)(f_j(d_X, e_j)) < c_j(d_X, e_j) \triangleright \delta \]

**Proposition 54.** Let \( G'_1 \) contain the equations for \( X \) and \( Z_1 \) defined above, \( G'_2 \) contain the equations for \( X \) and \( Z_2 \) defined above, and \( G'_3 \) contain the equations for \( X \) and \( Z_3 \) defined above. Let \( G_1 \) contain the equations for \( X \) and \( Z_1(d_X;D_X) = \partial_A(X(d_X)) \), \( G_2 \) contain the equations for \( X \) and \( Z_2(d_X;D_X) = \tau_A(X(d_X)) \), and \( G_3 \) contain the equations for \( X \) and \( Z_3(d_X;D_X) = \rho_R(X(d_X)) \). Then we have \( G_1 \Rightarrow G'_1, G_2 \Rightarrow G'_2 \) and \( G_3 \Rightarrow G'_3 \).

**Proof.** To prove the implications we use the identity mappings for \( g_X, g_{Z_1}, g_{Z_2} \) and \( g_{Z_3} \). The equations for \( X \) are proven trivially. For the other equations we substitute \( X \) by its right-hand side and apply the axioms (D3), (SUM8), (D5), (D4), (D1), (D2), (A7), and (A6) to push \( \partial_A \) inside; the axioms (T3), (SUM9), (T5), (T4), (T1), and (T2) to push \( \tau_A \) inside; the axioms (R3), (SUM10), (R5), (R4), (R1), (RT), and (RD) to push \( \rho_R \) inside. After that we use the equations for \( Z_1, Z_2, Z_3 \) in \( G_1, G_2, G_3 \) respectively to eliminate the operators \( \partial_A, \tau_A \) and \( \rho_R \) completely and arrive at equations for \( Z_1, Z_2, Z_3 \) in \( G'_1, G'_2, G'_3 \) respectively. \( \square \)

The following examples show that the reverse implications of the latter proposition do not hold in every model.

**Example 55.** Let \( G_1 = \{ X = a \cdot X + b \cdot X, Z_1 = \partial_{[b]}(X) \} \) and \( G'_1 = \{ X = a \cdot X + b \cdot X, Z_1 = a \cdot Z_1 \} \). Consider the model from Example 52. The equations for \( X \) in both \( G_1 \) and \( G'_1 \) have the following solutions:
\[ \{\{a, b\}, \{a, a, b\}, \{b, a, b\}, \{\{a\}, \{a, b\}\}, \{a\}, \{a, b\}\} \]

The equation for \( Z_1 \) in \( G_1 \) has two solutions \( \{\{a\}\} \) and \( \top \), while the equation for \( Z_1 \) in \( G_1' \) has four solutions \( \{\{a\}\}, \{\{a, b\}\}, \{\{a\}, \{a, b\}\} \) and \( \top \).

**Example 56.** Let \( G_2 = \{ X = a \cdot X, Z_2 = \tau_{|a|}(X) \} \) and \( G_2' = \{ X = a \cdot X, Z_2 = \tau \cdot Z_2 \} \). Consider the branching bisimulation model [15]. The equation for \( Z_2 \) in \( G_2 \) has the unique solution \( Z_2 = \tau \), while the equation for \( Z_2 \) in \( G_2' \) has infinitely many solutions \( Z_2 = \tau \cdot p \), where \( p \) is any element of the model.

**Example 57.** Let \( G_3 = \{ X = a \cdot X + b \cdot X, Z_3 = \rho_R(X) \} \) and \( G_3' = \{ X = a \cdot X + b \cdot X, Z_3 = a \cdot Z_3 \} \), where \( R(a) = R(b) = a \). Consider the model from Example 52. The equation for \( Z_3 \) in \( G_3 \) has two solutions \( \{\{a\}\} \) and \( \top \), while the equation for \( Z_3 \) in \( G_3' \) has four solutions \( \{\{a\}, \{a, b\}\}, \{\{a\}, \{a, b\}\} \) and \( \top \).

### 5.3. Towards an LPE

Let \( G'' = G''_1 \cup G''_2 \) be a system of process equations with \( G''_1 \) an LPE and \( G''_2 \) containing parallel pCRL process equations. If \( G''_2 \) is empty, we are done. Otherwise, let \( (X, G'') \) be the process definition to be transformed. We substitute the right-hand sides for all parallel pCRL process names (other than \( X \)) in \( G''_2 \) and obtain the set \( G''_2 \) with a single process equation for \( X \), such that \( (X, G'') = (X, G''_1 \cup G''_2) \). We finish the description of our transformation of \( G'' \) into a single LPE by describing how \( G''_2 \) can be integrated with \( G''_1 \).

A general strategy is to apply an innermost/outermost reduction along the lines of Propositions 51 and 54, occasionally adding or replacing process equations.

We consider a typical case (but note that many variants are conceivable):

\[
\begin{align*}
G''_1 &= \{ Y(d\vec{Y}; D\vec{Y}) = p_Y \} \\
G''_2 &= \{ X(d\vec{X}; D\vec{X}) = \tau_I(\partial_H(Y(\vec{t})) Y(\vec{u})) \}
\end{align*}
\]

and proceed in a stepwise manner. First we reduce the \( ||\) -occurrence, so transform \( G''_2 \) into

\[
\begin{align*}
G''_3 &= \{ X(d\vec{X}; D\vec{X}) = \tau_I(\partial_H(Z(\vec{t}), \vec{u})) , Z(d\vec{Y}; D\vec{Y}, e\vec{Y}; D\vec{Y}) = Y(d\vec{Y}) Y(e\vec{Y}) \}
\end{align*}
\]

where \( e\vec{Y} \) is a fresh copy of \( d\vec{Y} \). With Lemma 18 it follows that for all \( Y \in |G''| \), \( (Y, G'') = (Y, G''_1 \cup G''_2) \). According to Proposition 51, there exists a system \( H \) with \( Z \) defined by a number of linear equations in the process names \( Z \) and \( Y \) such that \( (Z, G''_1 \cup G''_2) \Rightarrow_c (Z, H) \), and for the remaining process names \( Y \in |G''| \), \( (Y, G'') = (Y, G''_1 \cup G''_2) \Rightarrow_c (Y, H) \). Comparing the newly created system \( H \) of process equations with \( G'' \), we see that it contains one parallel pCRL operation less, and one more pCRL process equation consisting of the linear equation for \( Z \). Next, with Propositions 47 and 48 this system can be transformed into a system \( H' \) that contains a single LPE, say over process name \( U \), and the equation \( X(d\vec{X}; D\vec{X}) = \tau_I(\partial_H(U(\vec{u}))) \) where application of these propositions prescribes the value vector \( \vec{u} \). With Proposition 54 we can resolve the encapsulation and hiding operation in a similar fashion. This yields a system of process equations \( H'' \) that consists of an LPE over process name \( V \) and the equation \( X(d\vec{X}; D\vec{X}) = V(\vec{v}) \), and \( (X, H') \Rightarrow (X, H'') \). Now the last step of this final transformation is the conclusion \( (X, H'') = (V(\vec{v}), G_{\text{lin}}) \), where \( G_{\text{lin}} \) contains only the LPE for \( V \).
The description above illustrates the last part of our transformation. Without further proof we state the following result.

**Proposition 58.** Let \( G'' = G_1'' \cup G_2' \) be a system of process equations as described above (\( G_1'' \) an LPE, and \( G_2' \) containing parallel pCRL process equations). Then \( G'' \) can be transformed via innermost/outermost reduction into a system \( G_{\text{lin}} \) that contains one single LPE, and that satisfies \((X, G'') \Rightarrow_c (X'(\bar{t}X'), G_{\text{lin}})\) for a certain value vector \( \bar{t}X' \).

6. Conclusions

We described a transformation of parallel pCRL process definitions into a linear format, and argued that this transformation is correct. Our correctness argument is not tied to some particular model, and also applies to process definitions that do not necessarily imply that the models have unique solutions. Furthermore, this transformation is idempotent in the following sense: applying the transformation to an LPE yields the same LPE.

The algorithm underlying the transformation into LPE format basically matches the one that is currently implemented in the \( \mu \)CRL toolset [17]. Of course, during the process of linearization many optimizations are conceivable, some of which can only be applied in a certain context. We have already mentioned some optimization rewrite rules (Table 10) that can be applied during one of the linearization steps. Another optimization can be performed in the cases where a new process name is introduced. There can be a choice of what parameters to use for the new process name in order to fetch the complicated structure of data terms involved. Furthermore, there are many (minor) optimizations, such as the rewriting of conditions or the elimination of constant parameters. Due to the fact that the LPE format provides such a simple process structure, we feel that this type of optimizations can be best performed after the transformation into the LPE format. Such optimizations include rewriting of data terms, eliminations of redundant variables and constants, abstract interpretation, and so on.

There are two particular optimizations that we want to mention here in more detail: *regular linearization* and *clustering of actions*. The first of these is based on [25], and applies to the situation where regularity follows from the absence of termination in a recursion, like in \( X = a \cdot X \cdot X \). Restricting to standard process semantics for \( \mu \)CRL, an LPE that specifies the same behavior is \( X = a \cdot X \). However, this optimization is model dependent, as there can be models in which the two equations have different sets of solutions. For some other cases, also dealt with in [25] and used in the \( \mu \)CRL toolset, these optimizations can be justified on a general level using the equivalence of systems of process equations. For example, the system \( G_1 = \{X = a \cdot Y \cdot X, Y = b\} \) can be transformed into \( G_2 = \{X = a \cdot Z, Z = b \cdot X\} \), and we can prove that \((X, G_1) = (X, G_2)\), thus showing that this transformation is sound in every model. As for ‘clustering of actions’, we refer to Definition 2.7, Theorem 2.8 and Theorem A.4 in [21]. The transformation allows us to optimize an LPE to a form in which every action label occurs at most twice (either as a termination action or not). The constructed LPE is equivalent to the original one. During the transformation the sums \( \sum_{i \in I} \) and \( \sum_{j \in J} \) which in Definition 46 represent the abbreviations for alternative compositions, are changed to the ‘real’ sums over enumerated data types. We note that both these latter optimizations are implemented in the current version of the \( \mu \)CRL toolset.

In the future we plan to work on extending the linearization procedure to cover the full syntax of \( \mu \)CRL. Furthermore, the procedure can be extended to handle the timed version of
the language. Finally, additional extensions to the language like interrupts, process creation and priorities could be investigated, as these seem to be useful for applications.

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References


