Abstract Graph Transformation

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Abstract

Graphs may be used as representations of system states in operational semantics and model checking; in the latter context, they are being investigated as an alternative to bit vectors. The corresponding transitions are obtained as derivations from graph production rules.

In this paper we propose an abstraction technique in this framework: the state graphs are contracted by collecting nodes that are sufficiently similar (resulting in smaller states and a finite state space) and the application of the graph production rules is lifted to this abstract level. Since graph abstractions and rule applications can all be computed completely automatically, we believe that this can be the core of a practically feasible technique for software model checking.

Key words: Graph Transformation, Abstract Interpretation, Model Checking, Verification

1 Introduction

We study state-based models of system behaviour, in particular for software systems. Our eventual aim is to develop tools to support the verification of software through such models. For the basic modelling formalism we rely on *graph transformation*, which is a long-standing field of research with a rich underlying theory (see, e.g., [23] for an overview). Systems in many domains and on different levels of design detail lend themselves naturally to a behavioural specification in which states are modelled by graphs (with directed labelled edges over a finite alphabet) and transitions correspond to applications of graph transformation rules. We have presented in [20] some first results on applying model checking techniques in this

setting. The use of graph transformation pays off here, since the notions of independence and composition implied by the theory can be brought to bear immediately to the problems of state space explosion.

A powerful notion in model checking is *state abstraction*. It is generally recognized that some kind of abstraction, be it through slicing, bounding, predicate abstraction, symmetry recognition or a combination of these, is indispensible in software model checking in order to keep state space size under control. In [8,7] we have studied model checking in a setting where the states are abstractions of restricted classes of graphs (multisets and single-linked lists, respectively), in which nodes with sufficiently similar structure are combined. In the current paper we combine these two strands of research, by extending the abstraction to general graphs and lifting the theory of graph transformation to the resulting abstracted graphs. Technically, we base the abstraction of the graphs on the prior work in [16]; the contribution of the current paper is to lift the graph transformations to this level.

It should be noted that this kind of abstraction has already been studied, to great effect, in the context of abstract interpretation, viz. in the theory of *shape graphs*; e.g., by Sagiv, Reps, Wilhelm and others in [24,25,12,22]. There is therefore every reason to believe that this will give rise to an equally effective verification method in our setting of graph transformation-based model checking.

Approach. Any set of graph transformation rules, applied to a concrete start graph, gives rise to a (possibly infinite) concrete transition system. In this paper we define the application of the same rules, but now to graph abstractions — which, following the work cited above, we call shapes — in such a way that all transitions between the concrete states (transformations of concrete graphs) give rise to transitions between the abstract states (transformations of shapes). Thus we have an over-approximation of the concrete transition system, on the basis of which we can make certain predictions about the actual system behaviour. Moreover, for every abstract transition there is at least one underlying concrete transition, meaning that we do not have spurious abstract transitions.

We will use a running example of a circular buffer used to store data values. The buffer consists of an n-linked circular structure of C-nodes and a central B-node pointing to the (current) first and last cell through f- and l-edges. A cell can contain an object, modelled by a v-edge to an O-node, or be empty, modelled by a e-edge back to the B-node. Fig. 1 shows an example buffer of four cells, two of which are empty. The shape of this buffer combines the (structurally similar) empty C-nodes and the O-nodes, and additionally specifies *multiplicities* on the nodes and incoming edges. ¹ The ⁼1 on the incoming edge of the O-node, e.g., indicates that each concrete O-instance has exactly one incoming v-edge, which can come from

¹ In this paper we assume that graphs are deterministic — defined below — which means that outgoing multiplicities are not needed. We write the edge multiplicities on the *opposite* end of the arrows than is usual in class diagrams.

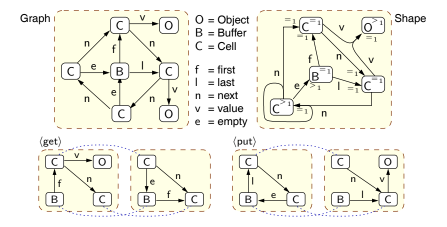


Fig. 1. Example circular buffer with four cells, its shape, and two production rules

either of the C-nodes.

To transform this example graph, Fig. 1 also shows two rules $\langle \text{put} \rangle$ and $\langle \text{get} \rangle$, each consisting of two graphs: a left hand side (LHS) and a right hand side (RHS). The rules describe the insertion and removal of objects, where for simplicity the nodes modelling the objects are actually created at insertion and deleted at removal. The effect of a rule is defined relative to a *matching* of the LHS, which is an injective graph morphism into the host graph. The images of those elements not in the RHS are subsequently removed from the host graph, whereas elements that are *fresh* in the RHS are added.

Given an initial graph and a set of production rules, we obtain a transition system by recursively applying all rules to all graphs. For instance, Fig. 2 shows the transition system for the graph and rules in Fig. 1. We propose to use such transition systems as the basis for model checking; first results are reported in [20]. However, for this technique to become practically feasible we need to address the following issues (among others):

- The models should be generic in the size of the data structures. As it is, for our example we get a different model if we start with a 5-cell buffer, etc.
- The models should be finite. As it is, if we add a rule to our example that may add a cell to a circular buffer when it is completely full, then the size of the graphs becomes unbounded and the state space becomes infinite.

By lifting graph transformations to the level of shapes we achieve both these goals. In fact, what we achieve is a completely automatic technique for state abstraction, in a setting where the models are inherently dynamic — that is, nodes and edges can be created and deleted at run-time. We believe that this is a promising basis for software verification, complementary to existing model checking techniques.

The abstract states will be *canonical* shapes, which is a sub-class satisfying some normalisation constraints. Their transformation is a three-step process.

Materialisation. This involves identifying the sub-shape where the rule applies

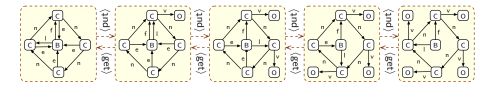


Fig. 2. Concrete transition system of the circular buffer

(using the matching) and extracting an explicit, concrete copy of it. This is necessary to accurately mimic the effect of the transformation. The same principle can be found in [24], from where we took the term "materialisation", but also in our own work [8,7], where it is called "extraction."

Transformation. The transformation of a materialised shape is much like an ordinary graph transformation. We will show that this type of transformation both preserves and reflects transformations of the corresponding instance graphs.

Normalisation. The result of the transformation, though still an abstract graph, is typically outside the sub-class of *canonical* shapes. Therefore, we have to massage it to fit it back into that class. This may introduce additional non-determinism: an arbitrary shape typically gives rise to multiple canonical shapes.

Structure of the paper. In Sect. 2 we define the basic notions of graphs and graph transformations, and we recall the abstraction as defined in [16]. The materialisation, transformation and normalisation steps are described in Sections 3–5; in Sect. 6 we combine them and complete the framework. Finally, Sect. 7 summaries the paper and discusses related work. Proofs of the theorems are given in the full report version, [19].

2 Definitions

Graphs and their transformations. In this section we define the basic graph formalism that we will use. In the following, \mathcal{L} denotes a fixed, finite set of labels.

Definition 2.1 (graph and morphism) A graph over \mathcal{L} is a tuple $G = \langle N, E \rangle$ where N is a set of nodes and $E \subseteq N \times \mathcal{L} \times N$ a set of labelled edges. G is called deterministic if $(v, a, w), (v, a, w') \in E$ implies w = w'.

If $G = \langle N_G, E_G \rangle$ and $H = \langle N_H, E_H \rangle$ are graphs over \mathcal{L} , a morphism $\phi: G \to H$ is a function $\phi: N_G \to N_H$, extended to E_G by $\phi((v, a, w)) = (\phi(v), a, \phi(w))$, such that $\phi(E_G) \subseteq E_H$.

An example deterministic graph was given in Fig. 1. Note that the node labels (B, C etc.) are actually not part of the formal definition; in fact they are superfluous (they can be derived from the edge labels), we have just included them for the sake of readability. In the following, $\mathbf{Gra}_{\mathcal{L}}$ denotes the class of graphs and $\mathbf{DGra}_{\mathcal{L}}$ the class of deterministic graphs. Given a edge $e = (v, a, w) \in E$ we call v the source, a the label and w the target of e. They denoted src(e), lab(e), and tgt(e)

respectively.

A bijective morphism $\phi: G \to H$ is called an *isomorphism*, and G and H are called *isomorphic* (denoted $G \cong H$) if there exists an isomorphism between them.

In the following definitions, we present production rules and their applications constructively, instead of the standard algebraic characterisation [3].

Definition 2.2 (production rule) A graph production rule is a pair of graphs P = (L, R) with $L, R \in \mathbf{DGra}_{\mathcal{L}}$, called the left hand side (LHS) and right hand side (RHS), respectively. We also sometimes regard P itself as a single graph given by the union $L \cup R$, and we distinguish the following sets:

- $N^{\text{del}} = N_L \setminus N_R$ and $E^{\text{del}} = E_L \setminus E_R$, the elements to be deleted;
- $N^{\text{use}} = N_L \cap N_R$ and $E^{\text{use}} = E_L \cap E_R$, the elements used (but not changed);
- $N^{\mathsf{new}} = N_R \setminus N_L$ and $E^{\mathsf{new}} = E_R \setminus E_L$, the elements to be created.

Two example production rules were given in Fig. 1. The set of production rules over \mathcal{L} is denoted $\mathbf{Prod}_{\mathcal{L}}$. The *application* of a production rule P=(L,R) to a graph G entails finding a *matching* $m:L\to G$, which is an injective morphism from the LHS to the graph (also satisfying some other conditions, introduced below), and then removing from G the images of N^{del} and E^{del} and adding to the resulting graph the elements in N^{new} and E^{new} . Care must be taken, however, to ensure that the new elements do not coincide with elements already in G. For this purpose, when discussing the application of a rule P to a graph G we will always assume P and G to be disjoint, i.e., $N_P \cap N_G = \emptyset$. This assumption can be satisfied without loss of generality by taking an isomorphic copy of P (and the result of the transformation does not depend on which isomorphic copy we take, modulo isomorphism).

Definition 2.3 (graph transformation) Let $P = (L, R) \in \mathbf{Prod}_{\mathcal{L}}$ and $G \in \mathbf{Gra}_{\mathcal{L}}$ be disjoint. A matching for P in G is an injective morphism $m: L \to G$ such that the following conditions hold for all $e \in E_G$:

- (i) If $src(e) \in m(N^{\mathsf{del}})$ or $tgt(e) \in m(N^{\mathsf{del}})$, then $e \in m(E^{\mathsf{del}})$;
- (ii) If $src(e) \in m(N^{\text{use}})$ and $\exists (m^{-1}(src(e)), lab(e), w) \in E^{\text{new}}$, then $e \in m(E^{\text{del}})$.

If m is a matching for P in G, the transformation of G according to P and m is defined by $((N_G \setminus m(N^{\text{del}})) \cup N^{\text{new}}, (E_G \setminus m(E^{\text{del}})) \cup E^{\text{new}})$. We write $G \xrightarrow{P,m} H$ to denote that m is a matching for P in G and H is the resulting transformed graph.

Application condition 1 is called the *dangling edge condition*; it is standard in the so-called *double pushout approach* to graph transformation (cf. [3]). Condition 2 could be called *preservation of determinism*; it is the most straightforward way to ensure that transformations remain in **DGra** (see Sect. 7 for a brief discussion). Example transformations (without the matchings) were shown in Fig. 2.

Proposition 2.4 Let $P \in \mathbf{Prod}_{\mathcal{L}}$, $G \in \mathbf{DGra}_{\mathcal{L}}$. If $G \xrightarrow{P,m} H$ then $H \in \mathbf{DGra}_{\mathcal{L}}$.

Multiplicities and shapes. A multiplicity is an interval of natural numbers. Formally, we define the set of multiplicities as $\mathbf{M} = \{(i,j) \in \mathbb{N} \times (\mathbb{N} \cup \{\star\}) \mid i \leq j\}$, where \star is used to denote infinity (i.e., $i < \star$ for all $i \in \mathbb{N}$). We use μ to range over multiplicities. We write =i for (i,i), >i for $(i+1,\star)$ and $\geq i$ for (i,\star) . The lower bound of a multiplicity $\mu \in \mathbf{M}$ is denoted by $\lfloor \mu \rfloor$ and the upper bound $\lceil \mu \rceil$; thus $\lfloor (i,j) \rfloor = i$ and $\lceil (i,j) \rceil = j$. Multiplicity μ is called positive if $\lfloor \mu \rfloor > 0$. We write $i \in \mu$ if $\lfloor \mu \rfloor \leq i \leq \lceil \mu \rceil$; based on this we define inclusion, $\mu_1 \subseteq \mu_2$, as $\forall i: i \in \mu_1 \Rightarrow i \in \mu_2$. A given set X has multiplicity μ , denoted $X:\mu$, if $|X| \in \mu$. The following defines two operations over multiplicities, where $\mu, \mu_1, \mu_2 \in \mathbf{M}$ and $i \in \mathbb{N}$ (note that $\star - i = \star + i = \star$ for all $i \in \mathbb{N}$):

$$\mu_1 + \mu_2 = (\lfloor \mu_1 \rfloor + \lfloor \mu_2 \rfloor, \lceil \mu_1 \rceil + \lceil \mu_2 \rceil)$$

$$\mu - i = (\max(0, |\mu| - i), \lceil \mu \rceil - i) \quad \text{if } \lceil \mu \rceil \ge i.$$

The following expresses some algebraic properties of these various concepts.

Proposition 2.5 Let $\mu \in \mathbf{M}$, and let A, B be arbitrary finite sets.

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(i) If A : \mu then (A \setminus B) : \mu - |A \cap B|.

(ii) If i < \lceil \mu \rceil then (\mu - i) + \exists i \subseteq \mu.
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Multiplicities are used as basic ingredients for the definition of *shapes*. These are graphs where a multiplicity is associated with each node, stating how many concrete nodes it represents, and with each pair of node v and label a, stating how many incoming a-edges each instance of v has. Formally:

Definition 2.6 (shape) A shape is a tuple $S = \langle N, E, nd, in \rangle$ with $\langle N, E \rangle \in \mathbf{Gra}_{\mathcal{L}}$ (sometimes denoted by G_S), and

- $nd: N \to \mathbf{M}$ a node multiplicity function;
- $in: N \to \mathcal{L} \to \mathbf{M}$ an incoming edge multiplicity function.

S is called deterministic if the following property holds:

• for all $v \in N$ such that $nd(v) = {}^{=}1$ and all $a \in \mathcal{L}$, $|\{w \mid (v, a, w) \in E\}| \le 1$ and $|\{w \mid (w, a, v) \in E\}| \le \lceil in(v)(a) \rceil$.

An example deterministic shape was shown in Fig. 1. We use $\mathbf{Sha}_{\mathcal{L}}$ to denote the class of shapes over \mathcal{L} , and $\mathbf{DSha}_{\mathcal{L}}$ for the deterministic shapes. Each shape stands for a number of *instances*, which are concrete (deterministic) graphs. In this sense, a shape is comparable to a *type graph*; however, the multiplicities provide far more control over the structure of the instances. The relation between a shape and its instances is defined by the following notion of *shaping*.

Definition 2.7 (shaping) Given a graph $G \in \mathbf{DGra}_{\mathcal{L}}$ and a shape $S \in \mathbf{Sha}_{\mathcal{L}}$, a shaping of G into S is a morphism $s : G \to G_S$ such that:

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(i) for all v \in N_S, s^{-1}(v) : nd(v);
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- (ii) for all $v \in N_G$ and $a \in \mathcal{L}$, $\{w \in N_G \mid (w, a, v) \in E_G\} : in(s(v))(a)$;
- (iii) for all $v \in N_G$ and $a \in \mathcal{L}$, if $\exists (s(v), a, w) \in E_S$ then $\exists (v, a, w') \in E_G$.

We write $s: G \to S$ to denote that s is a shaping of G into S. It is important to note that, due to possible inconsistencies between multiplicity constraints, not all shapes have instances. If a shape admits instances we call it *consistent*. In [16] we have shown that the notion of consistency is decidable for arbitrary (finite) $S \in \mathbf{Sha}_{\mathcal{L}}$.

A graph typically has (shapings into) many shapes; for instance, by changing the multiplicities of a shape into more permissive ones (i.e., that extend the old ones), all shapings remain preserved. In fact, shapes are interrelated by so-called *abstraction morphisms*.

Definition 2.8 (abstraction morphism) Let $S, T \in \mathbf{Sha}_{\mathcal{L}}$. An abstraction morphism α from S to T (written $\alpha : S \to T$) is a morphism $\alpha : G_S \to G_T$ with:

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(i) for all v \in N_T, nd_T(v) \supseteq \sum nd_S(\alpha^{-1}(v));
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- (ii) for all $v \in N_S$ and $a \in \mathcal{L}$, $in_T(\alpha(v))(a) \supseteq in_S(v)(a)$.;
- (iii) for all $v \in N_S$ and $a \in \mathcal{L}$, $\exists (\alpha(v), a, w) \in E_T$ implies $\exists (v, a, w') \in E_S$.

The following proposition states that (as expected) any instance of a shape is also an instance of a more abstract shape.

Proposition 2.9 Let $G \in \mathbf{DGra}_{\mathcal{L}}$ and $S, T \in \mathbf{Sha}_{\mathcal{L}}$. If $s: G \to S$ is a shaping and $\alpha: S \to T$ an abstraction, then $\alpha \circ s$ is a shaping of G into T.

3 Materialisation

As discussed in the introduction, we will lift the application of graph production rules to shapes. We do this in two steps: first we *materialise* the shape, then we transform the materialised graph as if it were a concrete graph. Materialisation is done relative to a prospective matching of the rule's LHS. Since such a matching is not a shaping (the LHS is only a *fragment* of a graph and so the cardinality constraints in the shape are not necessarily met) we have to define first what kind of objects they are.

Definition 3.1 Let $L \in \mathbf{DGra}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$. A pre-shaping p of L in S is a graph morphism $p: L \to G_S$ with the additional property that the upper bounds of the node and edge cardinalities are satisfied; i.e.,

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• for all v \in N_S, |p^{-1}(v)| \leq \lceil nd_S(v) \rceil;
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• for all $v \in N_G$ and $a \in \mathcal{L}$, $|\{w \in N_G \mid (w, a, v) \in E_G\}| \leq \lceil in_S(p(v))(a) \rceil$.

A pre-shaping p is called concrete if the following additional properties hold:

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• for all v \in N_L, nd_S(p(v)) = {}^{=}1;
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• for all
$$(v, a, w) \in E_L$$
, $(p(v), a, w') \in E_S$ implies $w' = p(w)$.

Pre-shapings extend injective morphisms from a graphs-to-graphs notion to a graphs-to-shapes notion. Concreteness means that the morphism maps only to nodes and edges that are uniquely identifiable in any concrete instance.

Proposition 3.2 Let $L, G \in \mathbf{DGra}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$. If $f: L \to G$ is an injective

morphism and $s: G \to S$ a shaping, then $s \circ f$ is a pre-shaping of L into S.

The intuition is that the existence of a pre-shaping $p: L \to S$ indicates that L may be a *fragment* of an instance of S. We do not currently have a result that supports that intuition; that is, we do not know if or when the existence of p implies that there is an instance G with a (proper) shaping $s: G \to S$ and an embedding $m: L \to G$ such that $p = s \circ m$. We conjecture, however, that the results of [16] can easily be extended so as to reduce this property (for a given L and S) to an integer program, thus giving a decision procedure. For concrete pre-shapings, on the other hand, we have the following further property, depicted graphically in Fig. 3:

Proposition 3.3 Let $L \in \mathbf{DGra}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$ and let $c: L \to S$ be a concrete pre-shaping. For any $G \in \mathbf{DGra}_{\mathcal{L}}$ with a shaping $s: G \to S$, there is an injective morphism $m: L \to G$ such that $c = s \circ m$.



Fig. 3: Visualisation of Prop. 3.3

Given a LHS L, a shape S and a preshaping $p: L \rightarrow S$, the *materialisation* of S relative to p is defined by disjointly adding a copy of L to S, connecting it to S where necessary, and adapting the node multiplicities of S to account for

the extraction of one or more instances from them. W.l.o.g. we assume $N_L \cap N_S = \emptyset$; we define a function $\alpha_p: (N_L \cup N_S) \to N_S$ by

$$\alpha_p = p \cup id_S .$$

 α_p is extended to edges as usual. The materialisation of S relative to p is defined by $S^{+p}=\langle N^{+p},E^{+p},nd^{+p},in^{+p}\rangle$ with

$$N^{+p} = N_L \cup N_S$$

$$E^{+p} = \alpha_p^{-1}(E_S) \setminus \{(v, a, w) \mid v \in N_L, \exists (v, a, w') \in E_L : w' \neq w\}$$

$$nd^{+p} : v \mapsto \begin{cases} nd_S(v) - |p^{-1}(v)| & \text{if } v \in N_S \\ = 1 & \text{otherwise} \end{cases}$$

$$in^{+p} : v \mapsto in_S(\alpha_p(v)) .$$

An example materialisation is shown in Fig. 4. The first thing to show is the relation between L, S and S^{+p} . (See also Fig. 5.)

Proposition 3.4 Let $L \in \mathbf{DGra}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$, and let $p: L \to S$ be a preshaping. α_p gives rise to an abstraction morphism from S^{+p} to S, and id_L gives rise to a concrete pre-shaping of L into S^{+p} , such that $p = \alpha_p \circ id_L$.

The materialisation satisfies the following characteristic property (see Fig. 5):

Proposition 3.5 Let $L, G \in \mathbf{DGra}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$. For an arbitrary injective morphism $m: L \to G$ and a shaping $s: G \to S$, let $p = s \circ m$; then there is a shaping $t: G \to S^{+p}$ with $s = \alpha_p \circ t$ and $t \circ m = id_L$.

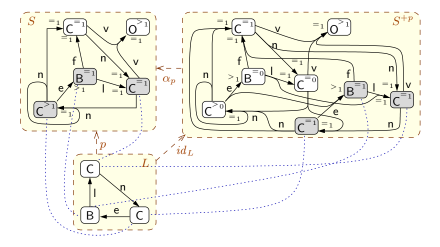


Fig. 4. Materialisation of the shape in Fig. 1 w.r.t. the LHS of \(\rangle \text{put} \)

4 Transformation

In this section we prove the correctness of the abstraction we have defined, in the sense that a transformation of a shape with respect to a singular pre-shaping simulates a transformation of the underlying instance graphs and vice-versa.

First we extend the transformation definition (see Def. 2.3) to shapes.

Definition 4.1 (shape transformation) Let $P = (L, R) \in \mathbf{Prod}_{\mathcal{L}}$ and $S \in \mathbf{Sha}_{\mathcal{L}}$ be disjoint. An abstract matching for P in S is a concrete pre-shaping $c: L \to S$ such that $c: L \to G_S$ is a (concrete) matching for P in the graph part of S. If c is an abstract matching for P in S, then the transformation of S according to P and S is defined by S is S and S such that

$$\begin{split} N_T &= (N_S \backslash c(N^{\mathsf{del}})) \cup N^{\mathsf{new}} \\ E_T &= (E_S \backslash c(E^{\mathsf{del}})) \cup E^{\mathsf{new}} \\ nd_T(v) &= \left\{ \begin{matrix} nd_S(v) & \text{if } v \in N_S \\ = 1 & \text{otherwise} \end{matrix} \right. \\ in_T(v)(a) &= \left\{ \begin{matrix} in_S(v)(a) - |\{w \mid (w, a, v) \in c(E^{\mathsf{del}})\}| \\ + = |\{w \mid (w, a, v) \in E^{\mathsf{new}}\}| & \text{if } v \in N_S \end{matrix} \right. \\ &= |\{w \mid (w, a, v) \in E^{\mathsf{new}}\}| & \text{otherwise} \end{split}$$

We write $S \xrightarrow{P,c} T$ to denote that c is an abstract matching for P in S and T is the resulting transformed shape.

The following are two of the crucial theorems of this paper, providing the connection between abstract and concrete transitions.

Theorem 4.2 Let $P = (L, R) \in \mathbf{Prod}_{\mathcal{L}}$, $S, T \in \mathbf{Sha}_{\mathcal{L}}$ and assume $S \xrightarrow{P,c} T$. For any shaping $s: G \to S$, there exists a matching m for P in G such that $c = s \circ m$, and for $G \xrightarrow{P,m} H$ there is a shaping $t: H \to T$.

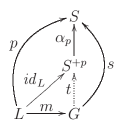


Fig. 5. Visualisation of Propositions 3.4 and 3.5

Theorem 4.3 Let $P = (L, R) \in \mathbf{Prod}_{\mathcal{L}}$, $G \in \mathbf{DGra}_{\mathcal{L}}$ and assume $G \xrightarrow{P,m} H$. For any shaping $s: G \to S$ such that $c = s \circ m$ is concrete, $S \xrightarrow{P,c} T$ with a shaping $t: H \to T$.

5 Normalisation

Materialisation and transformation are two essential ingredients of abstract graph transformations. However, there is a third ingredient still missing for an effective technique: namely, we need to identify a *canonical abstraction level*, on which there exist only a finite number of shapes and to which the target graph of each transformation will be re-normalised. Failing this, due to materialisation the graphs under transformation will become ever larger and more concrete, so that the state space is still infinite and the advantages of abstraction are lost.

For this canonical abstraction level, we will rely on the ideas developed in [16]. First of all, we select a collection of *base multiplicities* $\underline{\mathbf{M}} = \{=0, =1, >1\}$ (chosen in such a way that every finite set has exactly one base multiplicity). $\underline{\mathbf{M}}^{>0} = \underline{\mathbf{M}} \setminus \{=0\}$ denotes the set of *positive* base multiplicities. Next, we define the following notion of similarity $\sim_S \subseteq N_S \times N_S$ over nodes of a shape S:

(1)
$$v_1 \sim_S v_2 \Leftrightarrow in_S(v_1) = in_S(v_2) \wedge lab(src_S^{-1}(v_1)) = lab(src_S^{-1}(v_2))$$
.

Hence, two nodes are similar if they have the same incoming edge multiplicities and outgoing edge labels.

Definition 5.1 (canonical shape) A shape $S \in \mathbf{Sha}_{\mathcal{L}}$ is called canonical if

- (i) S is deterministic;
- (ii) for all $v \in N$, $nd(v) \in \mathbf{M}^{>0}$;
- (iii) for all $(v, a, w) \in E$, $in(v)(a) \in \underline{\mathbf{M}}^{>0}$;
- (iv) for all $v, w \in N$, $v \sim_S w$ implies v = w.

In words, a shape is canonical if it is deterministic, specifies positive base multiplicities for all nodes and edges (Clauses 2 and 3) and contains no non-trivially similar nodes (Clause 4). The class of canonical shapes is denoted $\mathbf{CSha}_{\mathcal{L}}$. An important fact from [16] is that $\mathbf{CSha}_{\mathcal{L}}$ is finite for every finite set \mathcal{L} .

We use the term *canonical* because, as we have shown in [18], there is an automatic way to obtain the *most concrete* canonical shape can(G) of a given deter-

ministic graph G. For an arbitrary shape S, on the other hand, there is typically not a *single* canonical shape that "covers" S in the sense of being more abstract (see Def. 2.8). Instead, we define a function norm such that norm(S) is a *set* of canonical shapes, which is optimal in a sense (shown below).

To normalise multiplicities, we take all (non-empty) intersections of the multiplicities occurring in S with $\underline{\mathbf{M}}$. This is defined as follows (where $\mu \in \mathbf{M}$ and $f: X \to \mathbf{M}$):

$$\mu/\underline{\mathbf{M}} = \{ \mu' \in \underline{\mathbf{M}} \mid \exists i : i \in \mu \land i \in \mu' \}$$
$$f/\underline{\mathbf{M}} = \{ g : X \to \underline{\mathbf{M}} \mid \forall x \in X : g(x) \in f(x)/\underline{\mathbf{M}} \} .$$

The function $norm: \mathbf{Sha}_{\mathcal{L}} \to \mathbf{2}^{\mathbf{CSha}_{\mathcal{L}}}$ is then defined as follows:

$$norm: S \mapsto \{part(T) \mid T \in \mathbf{DSha}_{\mathcal{L}}, T \triangleleft S, T \text{ consistent}\}$$
.

where the property $T \triangleleft S$ is defined as the conjunction of the following conditions:

$$N_{T} \subseteq \{(v,f) \mid v \in N_{S}, \ f \in in_{S}(v)/\underline{\mathbf{M}}\}$$

$$E_{T} \subseteq \{((v,f),a,(w,g)) \mid (v,a,w) \in E_{S}, \ g(a) \neq {}^{=}0\}$$

$$nd_{T} \in \{h: N_{T} \to \underline{\mathbf{M}}^{>0} \mid \forall v \in N_{S}: nd_{S}(v) \subseteq \sum_{(v,f) \in N_{T}} h((v,f))\}$$

$$in_{T} = \{((v,f),f) \mid (v,f) \in N_{T}\}$$
and $part(S) = T$ is defined by:
$$N_{T} = N_{S}/\sim_{S}$$

$$E_{T} = \{([v]_{\sim_{S}}, a, [w]_{\sim_{S}}) \mid (v,a,w) \in E_{S}\}$$

$$nd_{T} = \{([v]_{\sim_{S}}, (\sum_{v \sim_{S}w} nd_{S}(w))/\underline{\mathbf{M}}) \mid v \in N_{S}\}$$

$$in_{T} = \{([v]_{\sim_{S}}, in_{S}(v)) \mid v \in N_{S}\}$$

 $T \triangleleft S$ means that T is essentially obtained from S by assigning normalised incoming edge multiplicities and positive normalised node multiplicities to the nodes of S. This may result in S-nodes disappearing (if they otherwise would have multiplicity =0) or being split (if there is a choice of incoming edge multiplicities). The conditions on T ensure that it satisfies Clauses 2 and 3 of Def. 5.1. part(S), on the other hand, combines \sim_S -similar nodes, and so ensures Clause 4 of the definition provided that S already satisfies Clauses 1–3.

An example can be found in Fig. 6, which shows the normalisation of the shape obtained by transforming S using the materialisation in Fig. 4. This normalisation contains four shapes, two of which (on the right hand side) contain a sub-structure consisting of one or more n-linked C-nodes disconnected from the rest of the buffer. Such a structure does not model any graph occurring on the concrete level; it is an example of the ambiguity introduced by abstraction.

The canonical shape of an arbitrary deterministic graph is defined through a mapping $can: \mathbf{DGra}_{\mathcal{L}} \to \mathbf{CSha}_{\mathcal{L}}$, defined by

$$(2) can: G \mapsto part(S_G^{inst})$$

where $S_G^{inst} = (N_G, E_G, nd, in)$ is the "instance shape" of G, defined such that nd assigns =1 to all nodes $v \in N$ and $in(v)(a) = \mu$ is the unique multiplicity in $\underline{\mathbf{M}}$

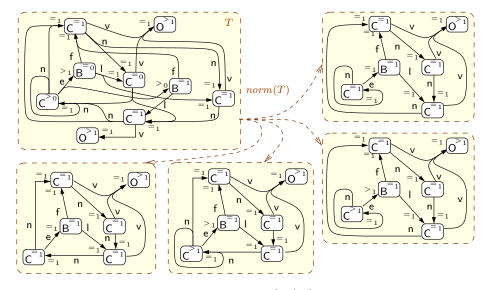


Fig. 6. Normalisation of the shape T with $S^{+p} \xrightarrow{\langle \mathsf{put} \rangle, id_L} T$ (with S and p as in Fig. 4)

such that $(tgt_G^{-1}(v) \cap lab_G^{-1}(a)) : \mu$. For instance, the shape in Fig. 1 is the image under can of the graph in that figure. The following results are recalled from [18].

Theorem 5.2 For arbitrary $G \in \mathbf{DGra}_{\mathcal{L}}$, $can(G) \in \mathbf{CSha}_{\mathcal{L}}$ and $\exists s : G \rightarrow can(G)$.

Theorem 5.3 For arbitrary $S \in \mathbf{Sha}_{\mathcal{L}}$, $norm(S) = \{can(G) \mid \exists s : G \to S\}$.

6 Transitions

In this section we pull together the results above, by defining concrete (graph) transition systems and abstract (shape) transition systems and stating their relation.

Definition 6.1 (transition system) *Let* Π *be a set of production rules.*

- A graph transition is a triple $G \xrightarrow{P} H$ with $G, H \in \mathbf{DGra}_{\mathcal{L}}$ and $P \in \Pi$ such that $G \xrightarrow{P,m} H$ for some m. A graph transition system is a tuple (\mathbf{G}, \to) where \to is the graph transition relation and $\mathbf{G} \subseteq \mathbf{DGra}_{\mathcal{L}}$ is closed under \to (i.e., $G \in \mathbf{G}$ and $G \xrightarrow{P} H$ implies $H \in \mathbf{G}$).
- A shape transition is a triple $S \xrightarrow{P} T$ with $S, T \in \mathbf{CSha}_{\mathcal{L}}$ and $P = (L, R) \in \Pi$ such that S^{+p} is consistent, $S^{+p} \xrightarrow{P, id_L} S'$ and $T \in can(S')$ for some pre-shaping $p: L \to S$. A shape transition system is a tuple (\mathbf{S}, \to) where \to is the shape transition relation and $\mathbf{S} \subseteq \mathbf{CSha}_{\mathcal{L}}$ is closed under \to .

Given a set of production rules Π and a graph $G \in \mathbf{DGra}$, we write $GTS(\Pi, G)$ for the smallest graph transition system including G; likewise, given $S \in \mathbf{CSha}_{\mathcal{L}}$ we write $STS(\Pi, S)$ for the smallest shape transition system including S. For instance, Fig. 2 shows the graph transition system $GTS(\Pi, G)$ where $\Pi = \{\langle \mathsf{put} \rangle, \langle \mathsf{get} \rangle\}$ and G is the graph of Fig. 1. Fig. 7 shows $STS(\Pi, can(G))$, where we have used some notational conventions to represent multiplicities: thin arrows and nodes are

singular (node/incoming edge multiplicity $^=1$) whereas fat ones are multiple ($^>1$). The arrows in Fig. 7 indicate $\langle put \rangle$ -applications; for each arrow there is an implicit $\langle get \rangle$ -application in the reverse direction. The darker (shaded) area is the fragment of the state space that actually is the image of the concrete transition system.

We now come to the main result, which states that the abstraction defined by can is finite and conservative, and in a weak sense does not over-approximate.

Theorem 6.2 Let Π be a set of production rules and $I \in \mathbf{DGra}$; let $GTS(\Pi, I) = (\mathbf{G}, \rightarrow)$ and $STS(\Pi, can(I)) = (\mathbf{S}, \rightarrow)$.

- (i) $can(\mathbf{G}) \subseteq \mathbf{S}$ and \mathbf{S} is finite;
- (ii) For all $G, H \in \mathbf{G}$, $G \xrightarrow{P} H$ implies $can(G) \xrightarrow{P} can(H)$.
- (iii) For all $S, T \in \mathbf{S}$ such that $S \xrightarrow{P} T$, there are $G', H' \in \mathbf{DGra}$ such that S = can(G') and $G' \xrightarrow{P} H'$.

This theorem implies that we can verify safety properties, where the propositions are graph predicates in the fragment of first-order logic that is reflected by our abstraction — characterised in [16] as a fragment of 2-variable logic. Typical examples of such properties are *state invariants*, such as:

- The buffer is either empty (i.e., no cell reachable from the first contains an object), or the first cell contains an object;
- If the buffer is empty, then the last cell is the predecessor of the first;
- If a cell contains an object, either it is the last or the next also contains an object.

Examples of valid properties that can *not* be verified, i.e., that appear to be violated on the abstract level but are in fact true in the concrete system (so-called "false negatives") are:

- All cells of the circular buffer are connected;
- \(\rho\text{put}\rangle\) can only be executed infinitely often if \(\rho\text{get}\rangle\) is also done infinitely often;
- Objects are removed in the order they were inserted.

List reversal. To enable a better comparison with existing approaches, the remainder of this section is devoted to an example that has been used several times before in heap structure analysis; see, e.g., [24,21]. ²

```
while (a.head != nil) do
Cell tmp := b.head;
b.head := a.head;
a.head := a.head.next;
b.head.next := tmp;
dod
```

The program uses a data structure consisting of List-nodes pointing via a head-edge to a list of Cell-nodes linked by next-edges; there is a unique nil-node modelling the end of the list. Fig. 8 shows a straightforward, line-by-line translation of this program into graph transformation rules. The vari-

ables and fields are represented by edges and their values by nodes. There is a

² Note, however, that we present this only for the sake of comparison and *not* because we consider this kind of sequential program analysis to be the strength of our method; the approaches in *op. cit.* are superior here. See also Sect. 7.

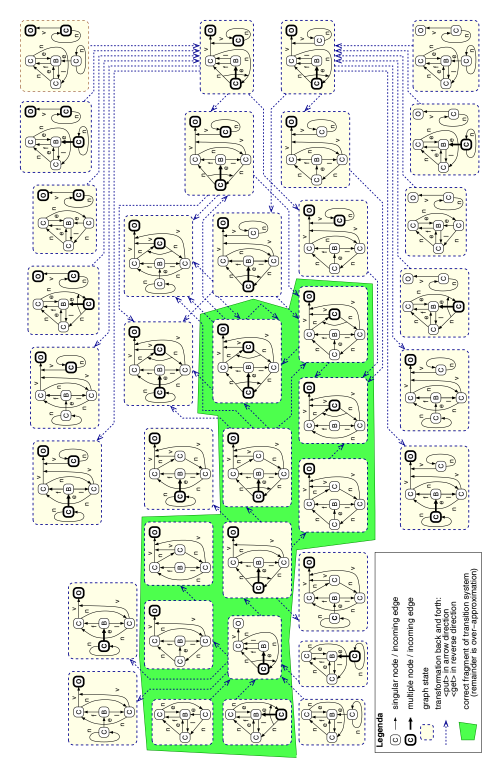


Fig. 7. Abstract buffer transition system

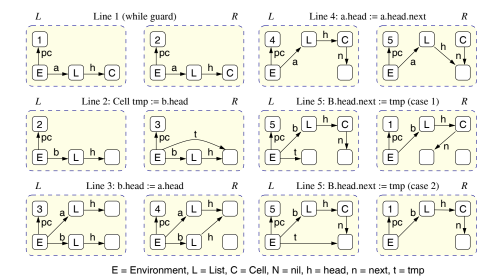


Fig. 8. Small-step transformation rules for the list reversal program.

central, E-labelled node that stands for the run-time environment, to which the local variable edges are attached and which maintains a pc-labelled "program counter" edge. Line 5 needs two rules, to distinguish the case where b.head.next already equals tmp (which may occur if the list a originally has only a single element); this is because our matchings are required to be injective (see Def. 2.2). We can now use standard graph transformation results to combine these rules into "large-step" ones that describe the combined effect of the loop body. The two resulting rules are shown in Fig. 9 (omitting the program counter, which now always equals 1).

We show in Fig. 10 the complete transition system generated by the large-step rule $\langle \text{swap} \rangle$ ($\langle \text{single} \rangle$ is never enabled from the chosen start state). The transition system is smaller than the one we would get from the small-step rules (see, e.g., [24]): the graph transformation theory has paid off here. The possible runs of the transition system all terminate in S_7 , which represents the reversed list, now pointed to by b, whereas a is empty. An example property verifiable in the transition system that the two lists are always kept separate: no Cell node is ever shared.

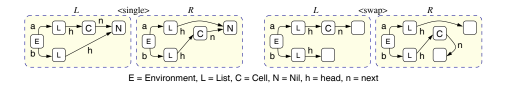


Fig. 9. Large-step rules for the list reversal program.

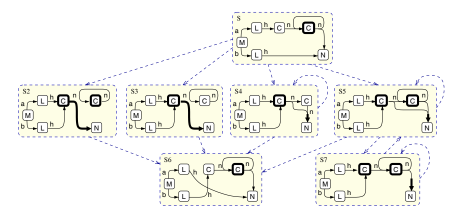


Fig. 10. Abstract transition system of the list reversal program.

7 Conclusions

We have presented a technique for the push-button construction of a finite abstract model of operational semantics, on the basis of a graph production system consisting of a set of graph transformation rules. As pointed out in the introduction, the contribution with respect to previous work is that this paper works out the transformation itself and the ensuing abstract transition system (Sections 4 and 6): the shape model was presented before. Given the fact that, as argued elsewhere (see, e.g., [2,6,9,13]), graph transformations are a very suitable formalism to model the behaviour of software systems, especially in the face of dynamic evolution, the results of this paper form an important step in creating a practically feasible method for the verification of such systems.

Related work. In addition to the more or less related work mentioned above, there are some lines of research that should be described in some more detail.

First and foremost among these is the work on shape graphs in [24,25], already discussed in the introduction. As remarked in Footnote 2, the analysis methods presented there are superior in the sense that, for the purpose of sequential program analysis, they yield more accurate predictions. For instance, through instrumentation it is possible to tune the notion of "shape" to particular properties to be checked, and the notion of 3-valued logic on the level of system assertions makes it possible to state some "negatives" with certainty. The basic contribution of this paper is that we develop essentially the same ideas (though originating from another direction altogether, namely model checking) in the context of the general, domain-independent formalism of graph transformation. This means that, on the one hand, the results can be applied in all those settings where graph production systems are being used for specifying or modelling system behaviour (which include parallel and distributed systems, and design-level semantic models, see above); and on the other hand, long-standing results from graph transformation, for instance regarding rule composition and independence, can be re-used in the context of abstraction.

More broadly speaking, our approach can be seen as an instance of abstract interpretation, pioneered by Cousot and Cousot [4]; see also [5] for a discussion of the use of abstract interpretation in model checking. In terms of [15], our shapes form a distinctness domain; however, in that terminology our abstract domain consists not of individual shapes but of *sets* of shapes (modulo isomorphism), and the abstract ordering is set inclusion. We therefore do have a Galois connection; but then, since we are not interested in computing fixpoints of computations but rather in expressing temporal properties of behaviour, we do not currently derive much benefit from this fact.

Another related area is the assertional approach for local reasoning on memory structures developed in, e.g., separation logic [14,21]. Here, too, an abstraction of a graph-based memory representation is taken as the basic model upon which verification is carried out. Although the core formalism is quite different in this case, one possible way to combine strengths is to investigate assertional semantics for graph transformation rules.

In the context of graph transformation, the closest related work is [1] on approximation of graph transition systems using *unfolding*, a technique that is generalised from Petri nets. Instead of constructing individual states, an unfolding combines all states into a single structure, in which transitions are modelled as purely local changes. Since eventually such local changes tend to propagate to a global level, the unfolding is *cut off* after a certain number of steps, at which point an over-approximation of the remaining behaviour is taken. Essentially, this approach promises the same capabilities for generic and infinite-state system verification as ours; once tool support for both is in place, a more detailed comparison should prove very interesting.

Future work. There is a host of smaller and larger improvements to be made.

- The current framework has a number of limitations in the graphs and transformation rules that are supported: graphs are deterministic, matchings have a dangling edge condition and have to be injective, and negative application conditions (cf. [11]) are not allowed. We conjecture that all of these restrictions can be lifted to some degree, at the price of some complications in the theory. For instance, rather than forbidding transformations that would violate the determinism, as we currently do in the definition of concrete matchings, one could take the pushout in the category of deterministic graphs, which essentially means determinising the graph after transformation, i.e., recursively merging outgoing edges with the same label.
- Graph transformations enjoy a very elegant algebraic characterisation (see, e.g., [3]), which we have ignored in the current paper. In particular, our abstract shape transformation have no underlying notion of a morphism or span of morphisms; instead they are based on *ad hoc* constructions. Consequently, there is no way to lift the results of this paper to other graph formalisms (for instance typed, at-

- tributed, or hypergraphs) or other types of abstraction without redoing the proofs.
- The connection with the work on shape graphs, discussed above, can be exploited further by transferring more of the results achieved there to the setting of graph transformation. Prime candidate among these is the notion of abstraction refinement through instrumentation.

Notwithstanding the fact that there is ample room for improvement, the constructions worked out in this paper are mature enough for implementation. We plan to extend the tool GROOVE (see [17]), which has the capability of generating concrete state spaces from graph production systems for the purpose of model checking (see [20]), with the necessary functionality to deal with shapes. As a proof-of-concept, we have "hand-crafted" the examples presented in this paper into GROOVE production rules mimicking the abstract behaviour.

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