Forgetting in CTL Using an Resolution Approach

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Abstract

Computation Tree Logic (CTL) is one of the central formalisms in formal verification. This paper presents a resolution-based method for computing the forgetting in CTL which has been proposed in another paper for KR this year. The method is an extension of the resolution calculus used to decide the satisfiability of CTL formula. An important feature inherited from the satisfiability of resolution-based method is guided by transforming a CTL formula into the normal form, i.e. the set of $\mathrm{SNF}_{\mathrm{CTL}}^g$ clauses. The $\mathrm{SNF}_{\mathrm{CTL}}^g$ language extend CTL with index for Existential quantifier. We use binary bisimulation relation to relate CTL and SNF_{CTL}^g , this shows that how to transform the CTL formula into SNF_{CTL}^g clauses and then return to CTL formula after finishing the computing process. Besides, we extend the original resolution rules by adding EF imply rules, which connects the next state and future state. Furthermore, it is shown that our algorithm is correct and the time and space complexity of our algorithm are $O((m+1)2^{4(n+n')}$.

Introduction

As a logical notion, forgetting was first formally defined in propostional and first order-logics by Lin and Reiter (Lin and Reiter 1994). Over the last twenty years, researchers have developed forgetting notions and theories not only in classical logic but also in other non-classical logic systems (Eiter and Kern-Isberner 2019), such as forgetting in logic programs under answer set/stable model semantics (Zhang and Foo 2006; Eiter and Wang 2008; Wong 2009; Wang et al. 2012; Wang, Wang, and Zhang 2013), forgetting in description logic (Wang et al. 2010; Lutz and Wolter 2011; Zhao and Schmidt 2017a) and knowledge forgetting in modal logic (Zhang and Zhou 2009; Su et al. 2009; Liu and Wen 2011; Fang, Liu, and Van Ditmarsch 2019). In application, forgetting has been used in planning (Lin 2003), conflict solving (Lang and Marquis 2010; Zhang, Foo, and Wang 2005), createing restricted views of ontologies (Zhao and Schmidt 2017a), strongest and weakest definitions (Lang and Marquis 2008), SNC (WSC) (Lin 2001) and so on.

Computation Tree Logic (CTL) (Clarke and Emerson 1981) is one of the main logical formalisms for program specification and verification. Though forgetting has been extensively investigated from various aspects of different logical systems. The existing forgetting methods in propositional logic, answer set programming, description logic and modal logic are not directly applicable in CTL. Similar with that in (Zhang and Zhou 2009), we have studied the forgetting in CTL from the semantic forgetting point of view in "the theory paper". And it is shown that our definition of forgetting satisfies those four postulates of forgetting.

Although we have proposed an model-based approach to compute forgetting in CTL in "the theory paper", but both time and space complexity are 2-exponential. It is urgent to find an efficient algorithm.

For one thing, the existing algorithm of computing forgetting in different logics talked above are not directly applicable in CTL. For instance, in propositional forgetting theory, forgetting atom q from φ is equivalent to a formula $\varphi[q/\top] \vee \varphi[q/\bot]$, where $\varphi[q/X]$ is a formula obtained from φ by replacing each q with X ($X \in \{\top, \bot\}$). This method cannot be extended to a CTL formula. Consider a CTL formula $\psi = \mathsf{AG}p \land \neg \mathsf{AG}q \land \neg \mathsf{AG}\neg q$. If we want to forget atom q from ψ by using the above method, we would have $\psi[q/\top] \vee \psi[q/\bot] \equiv \bot$. This is obviously not correct since after forgetting q this specification should not become inconsistent.

For another, as far as I know the existing methods to compute forgetting include the classical one talked above and resolution-based approachs in propositional logic (Lin and Reiter 1994; Wang 2015) and Ackermann-based approach (second-order elimination) in description logic (Zhao and Schmidt 2017b). However, the resolution and Ackermann-based methods need a specific normal form of the formula, it is hard to obtain such normal form in CTL. Although any CTL formula can be transformed into a set of ${\rm SNF}_{\rm CTL}^g$ clauses, but it do introduce the *index* and new atoms. Both the two problems are we should solve.

In this paper we extend the Resolution Calculus in (Zhang, Hustadt, and Dixon 2014) by eliminating the

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atoms introduced in the transformation process and combining the CTL with SNF_{CTL}^g by using the *binary bisimulation relation* (one is the set of atoms and another one is the set of indexes). Such a bisimulation relation is an extension of the set-based bisimulation talked in "the theory paper" by takeing *index* into account.

The paper is structured as follows: Section 2 introduces the notation and technical preliminaries. In section 3 we give a more precise definition of the problem. As key contributions, Section 4, introduces the resolution-based approach. Conclusion closes the paper.

Preliminaries

We start with some technical and notational preliminaries. Throughout this paper, we fix a finite set \mathcal{A} of propositional variables (or atoms), and use V, V' for subsets of \mathcal{A} . In the following several parts, we will introduce the structure we will use for CTL, syntactic and semantic of CTL and the normal form SNF $_{\text{CTL}}^g$ (Separated Normal Form with Global Clauses for CTL) of CTL (Zhang, Hustadt, and Dixon 2009).

Model structure in CTL

In general, a transition system can be described by a *model* structure (or Kripke structure) (see (Baier and Katoen 2008) for details). A model structure is a triple $\mathcal{M}=(S,R,L)$, where

- S is a finite nonempty set of states ¹,
- $R \subseteq S \times S$ and, for each $s \in S$, there is $s' \in S$ such that $(s, s') \in R$,
- L is a labeling function $S \to 2^{\mathcal{A}}$.

Given a model structure $\mathcal{M}=(S,R,L)$, a path π_{s_i} starting from s_i of \mathcal{M} is an infinite sequence of states $\pi_{s_i}=(s_i,s_{i+1}s_{i+2},\dots)$, where for each j $(0\leq i\leq j)$, $(s_j,s_{j+1})\in R$. By $s'\in\pi_{s_i}$ we mean that s' is a state in the path π_{s_i} . A state $s\in S$ is initial if for any state $s'\in S$, there is a path π_s s.t $s'\in\pi_s$. If s_0 is an initial state of \mathcal{M} , then we denote this model structure \mathcal{M} as (S,R,L,s_0) .

For a given model structure $\mathcal{M}=(S,R,L,s_0)$ and $s\in S$, the *computation tree* $\mathrm{Tr}_n^{\mathcal{M}}(s)$ of \mathcal{M} (or simply $\mathrm{Tr}_n(s)$), that has depth n and is rooted at s, is recursively defined as (Browne, Clarke, and Grumberg 1988), for $n\geq 0$,

- $Tr_0(s)$ consists of a single node s with label s.
- $\operatorname{Tr}_{n+1}(s)$ has as its root a node m with label s, and if $(s,s')\in R$ then the node m has a subtree $\operatorname{Tr}_n(s')$.

A K-structure (or K-interpretation) is a model structure $\mathcal{M}=(S,R,L,s_0)$ associating with a state $s\in S$, which is written as (\mathcal{M},s) for convenience in the following. In the case $s=s_0$ is an initial state of \mathcal{M} , the K-structure is *initial*.

Syntax and semantics of CTL

In the following we briefly review the basic syntax and semantics of the CTL (Clarke, Emerson, and Sistla 1986). The *signature* of the language \mathcal{L} of CTL includes:

- a finite set of Boolean variables, called *atoms* of \mathcal{L} : \mathcal{A} ;
- constant symbols: \bot and \top ;
- the classical connectives: \vee and \neg ;
- the path quantifiers: A and E;
- the temporal operators: X, F, G U and W, that means 'neXt state', 'some Future state', 'all future states (Globally)', 'Until' and 'Unless', respectively;
- parentheses: (and).

The (existential normal form or ENF in short) formulas of \mathcal{L} are inductively defined via a Backus Naur form:

$$\phi ::= \bot \mid \top \mid p \mid \neg \phi \mid \phi \lor \phi \mid \mathsf{EX}\phi \mid \mathsf{EG}\phi \mid \mathsf{E}[\phi \cup \phi] \quad (1)$$

where $p \in \mathcal{A}$. The formulas $\phi \wedge \psi$ and $\phi \rightarrow \psi$ are defined in a standard manner of propositional logic. The other form formulas of \mathcal{L} are abbreviated using the forms of (1).

We are now in the position to define the semantics of \mathcal{L} . Let $\mathcal{M}=(S,R,L,s_0)$ be a model structure, $s\in S$ and ϕ a formula of \mathcal{L} . The *satisfiability* relationship between (\mathcal{M},s) and ϕ , written $(\mathcal{M},s)\models\phi$, is inductively defined on the structure of ϕ as follows:

- $(\mathcal{M}, s) \not\models \bot$ and $(\mathcal{M}, s) \models \top$;
- $(\mathcal{M}, s) \models p \text{ iff } p \in L(s);$
- $(\mathcal{M}, s) \models \phi_1 \lor \phi_2$ iff $(\mathcal{M}, s) \models \phi_1$ or $(\mathcal{M}, s) \models \phi_2$;
- $(\mathcal{M}, s) \models \neg \phi \text{ iff } (\mathcal{M}, s) \not\models \phi;$
- $(\mathcal{M}, s) \models \text{EX}\phi \text{ iff } (\mathcal{M}, s_1) \models \phi \text{ for some } s_1 \in S \text{ and } (s, s_1) \in R;$
- $(\mathcal{M}, s) \models EG\phi$ iff \mathcal{M} has a path $(s_1 = s, s_2, ...)$ such that $(\mathcal{M}, s_i) \models \phi$ for each $i \geq 1$;
- $(\mathcal{M}, s) \models \mathrm{E}[\phi_1 \mathrm{U} \phi_2]$ iff \mathcal{M} has a path $(s_1 = s, s_2, \ldots)$ such that, for some $i \geq 1$, $(\mathcal{M}, s_i) \models \phi_2$ and $(\mathcal{M}, s_j) \models \phi_1$ for each $1 \leq j < i$.

Similar to the work in (Browne, Clarke, and Grumberg 1988; Bolotov 1999), only initial K-structures are considered to be candidate models in the following, unless otherwise noted. Formally, an initial K-structure \mathcal{K} is a model of a formula ϕ whenever $\mathcal{K} \models \phi$. We denote $Mod(\phi)$ the set of models of ϕ . The formula ϕ is satisfiable if $Mod(\phi) \neq \emptyset$. Given two formulas ϕ_1 and ϕ_2 , $\phi_1 \models \phi_2$ we mean $Mod(\phi_1) \subseteq Mod(\phi_2)$, and by $\phi_1 \equiv \phi_2$, we mean $\phi_1 \models \phi_2$ and $\phi_2 \models \phi_1$. In this case ϕ_1 is equivalent to ϕ_2 . The set of atoms occurring in ϕ_1 , is denoted by $Var(\phi_1)$. ϕ_1 is V-irrelevant, written $IR(\phi_1,V)$, if there is a formula ψ with $Var(\psi) \cap V = \emptyset$ such that $\phi_1 \equiv \psi$.

The normal form of CTL

It has proved that any CTL formula φ can be transformed into a set T_{φ} of SNF $_{\text{CTL}}^g$ (Separated Normal Form with Global Clauses for CTL) clauses in polynomial time such that φ is satisfiable iff T_{φ} is satisfiable (Zhang, Hustadt, and Dixon 2008). An important difference between CTL formulae and SNF $_{\text{CTL}}^g$ is that SNF $_{\text{CTL}}^g$ is an extension of the syntax of CTL to use indices. These indices can be used to preserve a particular path context. The language of SNF $_{\text{CTL}}^g$ clauses is defined

¹Indeed, every state is identified by a configuration of atoms i.e., which holds in that state.

over an extension of CTL. That is the language is based on: (1) the language of CTL; (2) a propositional constant **start**; (3) a countably infinite index set Ind; and (4) temporal operators: $E_{\langle ind \rangle}X$, $E_{\langle ind \rangle}F$, $E_{\langle ind \rangle}G$, and $E_{\langle ind \rangle}U$.

Before talk about the sematic of this language, we introduce the SNF_{CTL}^g clauses at first. The SNF_{CTL}^g clauses consists of formulae of the following forms.

$$\begin{array}{lll} \operatorname{AG}(\operatorname{\mathbf{start}} \supset \bigvee_{j=1}^k m_j) & (initial\ clause) \\ & \operatorname{AG}(true \supset \bigvee_{j=1}^k m_j) & (global\ clause) \\ & \operatorname{AG}(\bigwedge_{i=1}^n l_i \supset \operatorname{AX} \bigvee_{j=1}^k m_j) & (\operatorname{A}-\operatorname{step\ clause}) \\ & \operatorname{AG}(\bigwedge_{i=1}^n l_i \supset \operatorname{E}_{\langle ind \rangle} \operatorname{X} \bigvee_{j=1}^k m_j) & (\operatorname{E}-\operatorname{step\ clause}) \\ & \operatorname{AG}(\bigwedge_{i=1}^n l_i \supset \operatorname{AF}l) & (\operatorname{A}-\operatorname{sometime\ clause}) \\ & \operatorname{AG}(\bigwedge_{i=1}^n l_i \supset \operatorname{E}_{\langle ind \rangle} \operatorname{F}l) & (\operatorname{E}-\operatorname{sometime\ clause}). \end{array}$$

where $k \geq 0$, n > 0, **start** is a propositional constant, l_i $(1 \leq i \leq n)$, m_j $(1 \leq j \leq k)$ and l are literals, that is atomic propositions or their negation and ind is an element of Ind (Ind is a countably infinite index set). By clause we mean the classical clause or the SNF $_{\text{CTL}}^g$ clause unless explicitly stated.

Formulae of SNF $_{\text{CTL}}^g$ over $\mathcal A$ are interpreted in Ind-model structure $\mathcal M=(S,R,L,[.],s_0)$, where S,R,L and s_0 is the same as our model structure talked in 2.1 and $[.]:\operatorname{Ind} \to 2^{(S*S)}$ maps every index $ind \in \operatorname{Ind}$ to a successor function [ind] which is a functional relation on S and a subset of the binary accessibility relation R, such that for every $s\in S$ there exists exactly a state $s'\in S$ such that $(s,s')\in [ind]$ and $(s,s')\in R$. An infinite path $\pi_{s_i}^{\langle ind\rangle}$ is an infinite sequence of states $s_i,s_{i+1},s_{i+2},\ldots$ such that for every $j\geq i$, $(s_j,s_{j+1})\in [ind].$

Similarly, an *Ind-structure* (or *Ind-interpretation*) is a Indmodel structure $\mathcal{M}=(S,R,L,[_],s_0)$ associating with a state $s\in S$, which is written as (\mathcal{M},s) for convenience in the following. In the case s is an initial state of \mathcal{M} , the Indstructure is *initial*.

The semantics of $\mathrm{SNF}_{\mathrm{CTL}}^g$ is then defined as shown next as an extension of the semantics of CTL defined in Section 2.2. Let φ and ψ be two $\mathrm{SNF}_{\mathrm{CTL}}^g$ formulae and $\mathcal{M}=(S,R,L,[_],s_0)$ be an Ind-model structure, the relation " \models " between $\mathrm{SNF}_{\mathrm{CTL}}^g$ formulae and \mathcal{M} is defined recursively as follows:

- $(\mathcal{M}, s_i) \models \mathbf{start} \text{ iff } s_i = s_0;$
- $(\mathcal{M}, s_i) \models \mathbf{E}_{\langle ind \rangle} \mathbf{X} \psi$ iff for the path $\pi_{s_i}^{\langle ind \rangle}$, $(\mathcal{M}, s_{i+1}) \models \psi$;

- $(\mathcal{M}, s_i) \models \mathbf{E}_{\langle ind \rangle} \mathbf{G} \psi$ iff for every $s_j \in \pi_{s_i}^{\langle ind \rangle}$, $(\mathcal{M}, s_j) \models \psi$;
- $(\mathcal{M}, s_i) \models \mathbf{E}_{\langle ind \rangle}[\varphi \mathbf{U}\psi]$ iff there exists $s_j \in \pi_{s_i}^{\langle ind \rangle}$ such that $(\mathcal{M}, s_j) \models \psi$ and for every $s_k \in \pi_{s_i}^{\langle ind \rangle}$, if $i \leq k < j$, then $(\mathcal{M}, s_k) \models \varphi$;
- $(\mathcal{M}, s_i) \models \mathbf{E}_{\langle ind \rangle} \mathbf{F} \psi \text{ iff } (\mathcal{M}, s_i) \models \mathbf{E}_{\langle ind \rangle} [\top \mathbf{U} \psi].$

The semantics of the remaining operators is analogous to that given previously but in the extended Ind-model structure $\mathcal{M}=(S,R,L,[_],s_0)$. A $\mathrm{SNF}^g_{\mathrm{CTL}}$ formula φ is satisfiable, iff for some Ind-model structure $\mathcal{M}=(S,R,L,[_],s_0), (\mathcal{M},s_0)\models\varphi$, and unsatisfiable otherwise. And if $(\mathcal{M},s_0)\models\varphi$ then (\mathcal{M},s_0) is called a Ind-model of φ , and we say that (\mathcal{M},s_0) satisfies φ . By $T\wedge\varphi$ we mean $\bigwedge_{\psi\in T}\psi\wedge\varphi$, where T is a set of formulae. Other terminologies are similar with those in CTL sub-section.

Problem Definition

In order to define our problem, *i.e.* forgetting in CTL, we review our definition of V-bisimulation.

Definition 1 Let $V \subseteq A$ and $K_i = (M_i, s_i)$ (i = 1, 2) be K-structures (Ind-structures). Then $(K_1, K_2) \in B$ if and only if

- (i) $L_1(s_1) V = L_2(s_2) V$,
- (ii) for every $(s_1, s_1') \in R_1$, there is $(s_2, s_2') \in R_2$ such that $(\mathcal{K}_1', \mathcal{K}_2') \in \mathcal{B}$, and
- (iii) for every $(s_2, s_2') \in R_2$, there is $(s_1, s_1') \in R_1$ where $\mathcal{K}'_i = (\mathcal{M}_i, s_i')$ with $i \in \{1, 2\}$.

Proposition 1 Let $i \in \{1,2\}$, $V_1, V_2 \subseteq \mathcal{A}$, $s_i's$ be two states and $\pi_i's$ be two pathes, and $\mathcal{K}_i = (\mathcal{M}_i, s_i)$ (i = 1, 2, 3) be K-structures (Ind-structures) such that $\mathcal{K}_1 \leftrightarrow_{V_1} \mathcal{K}_2$ and $\mathcal{K}_2 \leftrightarrow_{V_2} \mathcal{K}_3$. Then:

- (i) $s'_1 \leftrightarrow_{V_i} s'_2$ (i = 1, 2) implies $s'_1 \leftrightarrow_{V_1 \cup V_2} s'_2$;
- (ii) $\pi'_1 \leftrightarrow_{V_i} \pi'_2$ (i = 1, 2) implies $\pi'_1 \leftrightarrow_{V_1 \cup V_2} \pi'_2$;
- (iii) for each path π_{s_1} of \mathcal{M}_1 there is a path π_{s_2} of \mathcal{M}_2 such that $\pi_{s_1} \leftrightarrow_{V_1} \pi_{s_2}$, and vice versa;
- (iv) $\mathcal{K}_1 \leftrightarrow_{V_1 \cup V_2} \mathcal{K}_3$;
- (v) If $V_1 \subseteq V_2$ then $\mathcal{K}_1 \leftrightarrow_{V_2} \mathcal{K}_2$.

Now we give the formal definition of forgetting in CTL from the semantic forgetting point view.

Definition 2 (Forgetting) Let $V \subseteq A$ and ϕ a CTL formula. A CTL formula ψ with $Var(\psi) \cap V = \emptyset$ is a result of forgetting V from ϕ , if

$$Mod(\psi) = \{ \mathcal{K} \text{ is initial } | \exists \mathcal{K}' \in Mod(\phi) \& \mathcal{K}' \leftrightarrow_V \mathcal{K} \}.$$
 (2)

Where K and K' are K-structures.

Note that if both ψ and ψ' are results of forgetting V from ϕ then $Mod(\psi) = Mod(\psi')$, i.e. , ψ and ψ' have the same models. In the sense of equivalence the forgetting result is unique (up to equivalence).

Similar with the V-bisimulation between K-structures, we define the $\langle V,I \rangle$ -bisimulation between Ind-structures as follows:

Definition 3 (binary bisimulation relation) Let $\mathcal{M}_i = (S_i, R_i, L_i, [_]_i, s_0^i)$ with $i \in \{1, 2\}$ be two Ind-structures, V be a set of atoms and $I \subseteq Ind$. The $\langle V, I \rangle$ -bisimulation $\beta_{\langle V, I \rangle}$ between initial Ind-structures is a set that satisfy $((\mathcal{M}_1, s_0^1), (\mathcal{M}_2, s_0^2)) \in \beta_{\langle V, I \rangle}$ if and only if $(\mathcal{M}_1, s_0^1) \leftrightarrow_V (\mathcal{M}_2, s_0^2)$ and $\forall j \notin I$ there is

- (i) $\forall (s, s_1) \in [j]_1$ there is $(s', s_1') \in [j]_2$ such that $s \leftrightarrow_V s'$ and $s_1 \leftrightarrow_V s_1'$, and
- (ii) $\forall (s', s_1') \in [j]_2$ there is $(s, s_1) \in [j]_1$ such that $s \leftrightarrow_V s'$ and $s_1 \leftrightarrow_V s_1'$.

We call this relation as binary bisimulation relation, denoted as $\langle V,I \rangle$ -bisimulation. Apparently, this definition is similar with our concept V-bisimulation except that this $\langle V,I \rangle$ -bisimulation has introduced the index.

Proposition 2 Let $i \in \{1, 2\}$, $V_1, V_2 \subseteq \mathcal{A}$, $I_1, I_2 \subseteq Ind$ and $\mathcal{K}_i = (\mathcal{M}_i, s_0^i)$ (i = 1, 2, 3) be Ind-structures such that $\mathcal{K}_1 \leftrightarrow_{\langle V_1, I_1 \rangle} \mathcal{K}_2$ and $\mathcal{K}_2 \leftrightarrow_{\langle V_2, I_2 \rangle} \mathcal{K}_3$. Then:

- (i) $\mathcal{K}_1 \leftrightarrow_{\langle V_1 \cup V_2, I_1 \cup I_2 \rangle} \mathcal{K}_3$;
- (ii) If $V_1 \subseteq V_2$ and $I_1 \subseteq I_2$ then $\mathcal{K}_1 \leftrightarrow_{\langle V_2, I_2 \rangle} \mathcal{K}_2$.

Proof: (i) By Proposition 1 we have $\mathcal{K}_1 \leftrightarrow_{V_1 \cup V_2} \mathcal{K}_3$. For (i) of Definition 3 we can prove it as follows: $\forall (s,s_1) \in [j]_1$ there is a $(s',s_1') \in [j]_2$ such that $s \leftrightarrow_{V_1} s'$ and $s_1 \leftrightarrow_{V_1} s'_1$ and there is a $(s'',s_1'') \in [j]_3$ such that $s' \leftrightarrow_{V_2} s''$ and $s_1' \leftrightarrow_{V_2} s_1''$, and then we have $\forall (s,s_1) \in [j]_1$ there is a $(s'',s_1'') \in [j]_3$ such that $s \leftrightarrow_{V_1 \cup V_2} s''$ and $s_1 \leftrightarrow_{V_1 \cup V_2} s''_1$. The (ii) of Definition 3 can be proved similarly.

(ii) This can be proved from (i).

The Calculus

Resolution in CTL is a method to decide the satisfiability of a CTL formula. In this part, we will explore a resolution-based method to compute forgetting in CTL. We use the transformation rules Trans(1) to Trans(12) and resolution rules (SRES1), ..., (SRES8), RW1, RW2, (ERES1), (ERES2) in (Zhang, Hustadt, and Dixon 2009).

The key problems of this method include (1) How to fill the gap between CTL and $\mathrm{SNF}_{\mathrm{CTL}}^g$ since there is index for existential quantifier in $\mathrm{SNF}_{\mathrm{CTL}}^g$; and (2) How to eliminate the irrelevant atoms, which we want to forget and introduced by the transformation rules, in the formula. We will resolve these two problems by $\langle V,I \rangle$ -bisimulation and *eliminate* operator respectively. For convenient, we use $V \subseteq \mathcal{A}$ denote the set we want to forget, $V' \subseteq \mathcal{A}$ with $V \cap V' = \emptyset$ the set of atoms introduced in the transformation process below, φ the CTL formula, T_{φ} be the set of $\mathrm{SNF}_{\mathrm{CTL}}^g$ clause obtained from φ by using transformation rules and $\mathcal{M} = (S, R, L, [_], s_0)$ unless explicitly stated. Let T, T' be two sets of formulae, I a set of indexes and $V'' \subseteq \mathcal{A}$, by $T \equiv_{\langle V'',I \rangle} T'$ we mean that $\forall (\mathcal{M}, s_0) \in \mathit{Mod}(T)$ there is a (\mathcal{M}', s_0') such that $(\mathcal{M}, s_0) \leftrightarrow_{\langle V'',I \rangle} (\mathcal{M}', s_0')$ and $(\mathcal{M}', s_0') \models T'$ and vice versa.

The algorithm of computing the forgetting in CTL is as Algorithm 1. The main idea of this algorithm is to change the CTL formula into a set of ${\rm SNF}_{\rm CTL}^g$ clauses at first (the Transform process), and then compute all the possible resolutions

on the specified set of atoms (the Resolution process). Third, eliminating, which include *Instantiate*, *Connect* and *Removing_atoms* sub-processes, all the irrelevant atoms which dose not be eliminated by the resolution. Changing the result obtained before into a CTL formula at last, this will include three sub-processes: $Removing_index$ (removing the index in the formula), $Replacing_atoms$ (replacing the atoms in V' with an formula) and T_{CTL} (removing the **start** in T). To describe our algorithm clearly, we illustrate it with the following example.

Example 1 Let $\varphi = A((p \land q) \cup (f \lor m)) \land r$ and $V = \{p\}$.

In the following context we will show how to compute the $F_{\text{CTL}}(\varphi,V)$ step by step using our algorithm.

```
Input: A CTL formula \varphi and a set V of atoms Output: ERes(\varphi, V)

1 // T_{\varphi} = \emptyset // the initial set of SNF_{CTL}^g clauses of \varphi;

2 V' = \emptyset // the set of atoms introduced in the process of transforming \varphi into SNF_{CTL}^g clauses;

3 T_{\varphi}, V' \leftarrow Transform(\varphi);

4 Res \leftarrow Resolution(T_{\varphi}, V \cup V');

5 Inst_{V'} \leftarrow Instantiate(Res, V');

6 Com_{EF} \leftarrow Connect(Inst_{V'});

7 RemA \leftarrow Removing\_atoms(Com_{EF}, Inst_{V'});

8 NI \leftarrow Removing\_index(RemA);

9 Rp \leftarrow Replacing\_atoms(NI);

10 return \bigwedge_{\psi \in R_{PCTL}} \psi.
```

Algorithm 1: Computing forgetting - A resolution-based method

The Transform process

The *Transform* process, denoted as $Transform(\varphi)$, is to transform the CTL formula into a set of SNF_{CTL}^g clauses by using the rules Trans(1) to Trans(12) in (Zhang, Hustadt, and Dixon 2009)).

The transformation of any CTL formula φ into the set T_{φ} is a sequence $T_0, T_1, \ldots, T_n = T_{\varphi}$ of sets of formulae with $T_0 = \{ \operatorname{AG}(\operatorname{\mathbf{start}} \supset p), \operatorname{AG}(p \supset \operatorname{\mathbf{simp}}(\operatorname{\mathbf{nnf}}(\varphi))) \}$ such that for every $i \ (0 \le i < n), T_{i+1} = (T_i \setminus \{\psi\}) \cup R_i$ (Zhang, Hustadt, and Dixon 2009)), where p is a new atom not appearing in φ, ψ is a formula in T_i not in $\operatorname{SNF}_{\operatorname{CTL}}^g$ clause and R_i is the result set of applying a matching transformation rule to ψ . Note that throughout the transformation formulae are kept in negation normal form.

Proposition 3 Let φ be a CTL formula, then $\varphi \equiv_{\langle V',I \rangle} T_{\varphi}$.

Proof: (sketch) This can be proved from T_i to T_{i+1} $(0 \le i < n)$ by using one transformation rule on T_i .

This means that φ has the same models with T_{φ} excepting that the atoms in V' and the relations [i] with $i \in I$.

```
Input: A CTL formula \varphi
   Output: A set T_{\varphi} of SNF_{\text{CTL}}^g clauses and a set V' of
1 T_{\varphi} = \emptyset // the initial set of SNF<sup>g</sup><sub>CTL</sub> clauses of \varphi;
2 OldT = \{ \mathbf{start} \supset z, z \supset \mathbf{simp}(\mathbf{nnf}(\varphi)) \};
3 V' = \{z\};
4 while OldT \neq T_{\varphi} do
        OldT = T_{\varphi};
        R = \emptyset;
6
        X = \emptyset;
7
        if Chose a formula \psi \in OldT that dose not a
8
         SNF_{CTL}^g clause then
              Using a match rule Rl to transform \psi into a set
              R of SNF_{CTL}^g clauses;
              X is the set of atoms introduced by using Rl;
10
              V' = V' \cup X;
             T_{\varphi} = OldT \setminus \{\psi\} \cup R;
        end
14 end
```

Algorithm 2: $Transform(\varphi)$

Example 2 By the *Transform* process, the result T_{φ} of the Example 1 can be listed as follows:

```
 \begin{array}{lll} \textbf{1.start} \supset z & 2. \top \supset \neg z \lor r & 3. \top \supset \neg x \lor f \lor m \\ \textbf{4.} \top \supset \neg z \lor x \lor y & 5. \top \supset \neg y \lor p & 6. \top \supset \neg y \lor q \\ \textbf{7.} z \supset \mathsf{AF} x & 8. y \supset \mathsf{AX}(x \lor y). \end{array}
```

Besides, the set of new atoms introduced in this process is $V' = \{x, y, x\}.$

The Resolution process

The Resolution process is to compute all the possible resolutions of T_{φ} on $V \cup V'$, denoted as $Resolution(T_{\varphi}, V \cup V')$. A derivation on a set $V \cup V'$ of atoms and T_{φ} is a sequence $T_0, T_1, T_2, \ldots, T_n = Res$ of sets of $\mathrm{SNF}_{\mathrm{CTL}}^g$ clauses such that $T_0 = T_{\varphi}$ and $T_{i+1} = T_i \cup R_i$ where R_i is a set of clauses obtained as the conclusion of the application of a resolution rule to premises in T_i . Note that all the T_i $(0 \le i \le n)$ are set of $\mathrm{SNF}_{\mathrm{CTL}}^g$ clauses. Besides, if there is a T_i containing $\mathrm{start} \supset \bot$ or $\top \supset \bot$, then we have $\mathrm{F}_{\mathrm{CTL}}(\varphi, V) = \bot$. Given two clauses C and C', we call C and C' are resolvable, the result denoted as res(C, C'), if there is a resolution rule using C and C' as the premises on some given atom. And the pseudocode of algorithm Resolution is as Algorithm 3.

Proposition 4 Let φ be a CTL formula, then $T_{\varphi} \equiv_{\langle V \cup V', \emptyset \rangle} Res$.

Proof:(sketch) This can be proved from T_i to T_{i+1} ($0 \le i < n$) by using one resolution rule on T_i .

Proposition 3 and Proposition 4 mean that $\varphi \equiv_{\langle V \cup V', I \rangle} Res$, this resolve part of the problem (1).

Example 3 The resolutions of T_{φ} obtained from Example 2

```
Input: A set T_{\varphi} of SNF_{\text{CTL}}^g clauses and a set V' of
   Output: A set Res of SNF_{CTL}^g clauses
1 S = \{C | C \in T_{\varphi} \text{ and } Var(C) \cap (V \cup V') = \emptyset\};
\Pi = T \setminus S;
3 for (p \in V \cup V') do
        \Pi' = \{C \in \Pi | p \in Var(C)\};
        \Sigma = \Pi \setminus \Pi';
        for (C \in \Pi' \text{ s.t. } p \text{ appearing in } C \text{ positively}) do
              for (C' \in \Pi' \text{ s.t. } p \text{ appearing in } C' \text{ negatively }
              and C, C' are resolvable) do
                   \Sigma = \Sigma \cup \{res(C, C')\};
                   \Pi' = \Pi' \cup \{C'' = res(C, C') | p \in C''
                   Var(C'')};
             end
        end
        \Pi = \Sigma;
13 end
4 Res = \Pi \cup S;
```

Algorithm 3: Resolution(T, V')

on $V \cup V'$ are listed as follows:

```
(1)start \supset r
                                        (1,2,SRES5)
 (2)start \supset x \lor y
                                        (1,4,SRES5)
 (3)\top \supset \neg z \lor y \lor f \lor m
                                        (3,4,SRES8)
 (4)y \supset AX(f \lor m \lor y)
                                        (3, 8, SRES6)
 (5) \top \supset \neg z \lor x \lor p
                                        (4,5,SRES8)
 (6) \top \supset \neg z \lor x \lor q
                                     (4, 6, SRES8)
 (7)y \supset AX(x \lor p)
                                     (5,7,SRES6)
 (8)y \supset AX(x \lor q)
                                     (5, 8, SRES6)
                                     (3,(2), SRES5)
 (9)start \supset f \lor m \lor y
 (10)start \supset x \lor p
                                     (5,(2), SRES5)
 (11)start \supset x \lor q
                                     (6,(2), SRES5)
(12) \top \supset p \vee \neg z \vee f \vee m
                                      (5, (3), SRES8)
(13) \top \supset q \vee \neg z \vee f \vee m
                                      (6, (3), SRES8)
(14)y \supset AX(p \lor f \lor m)
                                      (5, (4), SRES6)
(15)y \supset AX(q \lor f \lor m)
                                      (6, (4), SRES6)
(16)start \supset f \lor m \lor p
                                      (5, (9), SRES5)
(17)start \supset f \lor m \lor q
                                      (6, (9), SRES5)
```

The Elimination process

For resolving problem (2), we should pay attention to the following properties that obtained from the transformation and resolution rules at first:

- (GNA) for all atom p in $Var(\varphi)$, p do not positively appear in the left hand of the SNF $_{CTL}^g$ clause;
- (PI) for each atom p ∈ V', if p appearing in the left hand of a SNF^g_{CTL} clause, then p appear positively.

This *Elimination* process include three sub-processes: *Instantiate*, *Connect* and *Removing_atoms*. We will describe those sub-processes carefully blow.

The Instantiation process An *instantiate formula* ψ of set V'' of atoms is a formula such that $Var(\psi) \cap V'' = \emptyset$. Given a formula of the form $p \supset \psi$ with p is an atom not in $V'' \cup Var(\psi)$, if ψ is an instantiate formula of set V'' then we call p is instantiated by ψ . A key point to compute forgetting is eliminateing those irrelevant atoms, for this purpose we define the follow instantiation process.

Definition 4 (instantiation) Let V'' = V' and $\Gamma = Res$, then the process of instantiation is as follows:

- (i) for each global clause $C = \top \supset D \lor \neg p \in \Gamma$, if there is one and on one atom $p \in V'' \cap Var(C)$ and $Var(D) \cap (V \cup V'') = \emptyset$ then let $C = p \supset D$ and $V'' := V'' \setminus \{p\}$;
- (ii) find out all the possible instantiate formulae $\varphi_1, ..., \varphi_m$ of $V \cup V''$ with $p \supset \varphi_i \in \Gamma$ $(1 \le i \le m)$;
- (iii) if there is $p \supset \varphi_i$ for some $i \in \{1, ..., m\}$, then let $V'' := V'' \setminus \{p\}$;
- (iv) for $\bigwedge_{j=1}^m p_j \supset \varphi \in \Gamma$ ($i \in \{1, ..., n\}$), if there is $\alpha \supset p_1, ..., \alpha \supset p_n \in \Gamma$ and φ is an instantiate formula of $V \cup V''$, then let $\Gamma_1 := \Gamma \cup \{\alpha \supset \varphi\}$. if $\Gamma_1 \neq \Gamma$ then let $\Gamma := \Gamma_1$ go to step (i), else return $V \cup V''$.

Where p, p_i $(1 \le i \le m)$ are atoms and α is a conjunction of literals or **start**.

Intuitively, this process iteratively removes the atoms in V' that can be represented by the formula of $Var(\varphi)\setminus (V''\cup V)$. We denote this process as $Instantiate(\Gamma,V')$, which can be described as the following Algorithm 4. After this process we obtain a set of atoms that do not has been instantiated by any instantiate formula of $V\cup V''$ in this process.

Example 4 By using the instantiation process on result of Example 3, we obtain that x is instantiated by $f \vee m$ at first since there is $\top \supset \neg x \vee f \vee m \in T_{\varphi}$ with $x \in V'$ and $Var(f \vee m) \cap (V \cup V') = \emptyset$, then $V'' = \{y, z\}$.

Similarly, due to $\top \supset \neg y \lor q \in T_{\varphi}$ and $y \supset \mathsf{AX}(q \lor f \lor m) \in T_{\varphi}$, then y can be instantiated by $q \land \mathsf{AX}(q \lor f \lor m)$. And z can be instantiated by r. Therefore $V'' = \emptyset$ That is $\mathit{Instantiate}(Res, V') = V$, which means all the introduced atoms are instantiated.

By instantiation operator, we guarantee those atoms in $V \cup V''$ are really irrelevant, *i.e.* should be forgot.

The Connect process Let P be a conjunction of literals, l, l_1 be literals, in which $Var(l_1) \in V \cup V'$, and C_i ($i \in \{2,3,4\}$) be classical clauses. Let $A = \{true \supset \neg l \lor \neg l_1 \lor C_2, l \supset C_3 \lor C_2\}$, $\alpha = P \supset ((\neg C_3 \land \neg C_2) \supset (\mathsf{E}_{\langle ind \rangle} \mathsf{X}(C_3 \land \neg (C_2 \lor C_4) \supset \mathsf{AXAF}(C_3 \lor C_2))))$, $\beta = P \supset ((\neg C_3 \land \neg C_2) \supset (\mathsf{AX}(C_3 \land \neg (C_2 \lor C_4) \supset \mathsf{AXAF}(C_3 \lor C_2))))$ and $\gamma = P \supset ((\neg C_3 \land \neg C_2) \supset (\mathsf{E}_{\langle ind \rangle} \mathsf{X}(C_3 \land \neg (C_2 \lor C_4) \supset \mathsf{E}_{\langle ind \rangle} \mathsf{XE}_{\langle ind \rangle} \mathsf{F}(C_3 \lor C_2))))$, we add following new rules, we call it \mathbf{EF} imply.

```
\begin{split} \textbf{(EF1)}\{P \supset \mathsf{AF}l, P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X}(l_1 \vee C_4)\} \cup A \to \alpha \\ \textbf{(EF2)}\{P \supset \mathsf{AF}l, P \supset \mathsf{AX}(l_1 \vee C_4)\} \cup A \to \beta \\ \textbf{(EF3)}\{P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{F}l, P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X}(l_1 \vee C_4)\} \cup A \to \gamma \\ \textbf{(EF4)}\{P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{F}l, P \supset \mathsf{AX}(l_1 \vee C_4)\} \cup A \to \gamma. \end{split}
```

```
Input: A set \Gamma of SNF_{\mathtt{CTL}}^g clauses \varphi and V,V'\subseteq \mathcal{A}
                    Output: A set of atoms
       1 Let V'' := V';
      2 Let V_1 = \emptyset;
      3 Let \Gamma_1 := \emptyset;
      4 Let \Gamma_2 := \Gamma;
                 while (\Gamma_1 \neq \Gamma_2 \text{ or } V_1 \neq V'') do
                                              \Gamma_1 := \Gamma_2;
                                              V_1 := V''';
                                              for (C \in \Gamma_2) do
                                                                       if (C is a global clause) then
                                                                                                 Let C := D \vee \neg p;
                                                                                                 if (p \in V'' \cap Var(C)) and
                                                                                                 Var(D) \cap V == \emptyset) then
                                                                                                                         C := p \supset D;
                                                                                                                         V'' := V'' \setminus \{p\};
   14
                                                                                                 end
   15
                                                                       end
                                              end
                                              for (C \in \Gamma_2) do
                                                                       if (C == p \supset \varphi \text{ and } p \in V'' \text{ and } \varphi \in V''' \text{ and } \varphi \in V'' \text{ 
                                                                        Var(\varphi) \cap V \cup V'' == \emptyset) then
19
                                                                              V'' := V'' \setminus \{p\};
 20
21
22
23
                                                                       end
                                               end
                                              for (C \in \Gamma_2) do
                                                                      if (C == \bigwedge_{j=1}^{m} p_j \supset \varphi \text{ and }
                                                                        Var(\varphi) \cap V \cup V'' == \emptyset) then
                                                                                               if (there is \alpha \supset p_1, \ldots, \alpha \supset p_m \in \Gamma_2) then
 24
25
26
27
28
                                                                                                    | \Gamma_2 := \Gamma_2 \cup \{\alpha \supset \varphi\};
                                                                                                 end
                                                                       end
                                              end
 29 end
 30 return V \cup V''.
```

Algorithm 4: Computing *Instantiate*(Γ , V')

By Connect(Instantiate(Res, V')) we mean using (EF1) to (EF4) on Res and replacing $P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X}(\neg l \lor C_2 \lor C_4)$ with $P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X}(\neg l \lor C_2 \lor C_4) \lor \alpha$ for rule (EF1), replacing $P \supset \mathsf{AX}(\neg l \lor C_2 \lor C_4)$ with $P \supset \mathsf{AX}(\neg l \lor C_2 \lor C_4) \lor \beta$ for rule (EF2) and replacing $P \supset \mathsf{AX}(\neg l \lor C_2 \lor C_4) \lor \beta$ with $P \supset \mathsf{AX}(\neg l \lor C_2 \lor C_4) \lor \gamma$ for other rules when l, C_2, C_3 and C_4 are instantiate formulae of $\mathsf{Sub}(Res, V')$ and $\mathsf{Var}(l_1) \in V \cup V'$. The reason why we specify l, C_2, C_3 and C_4 are instantiate formulae of $\mathsf{Sub}(Res, V')$ in this process will be explained later.

Proposition 5 Let $\Gamma = Res$, we have $\Gamma \equiv_{\langle V', \emptyset \rangle}$ Connect(Instantiate(Γ, V')).

Proof: It is obvious from the (EF1) to (EF4).

We prove the (EF1), other rules can be proved similarly. Let $T_{i+1} = T_i \cup \{\varphi\}$, where $\{\varphi\}$ is obtained from T_i by using rule (EF1) on T_i , i.e. $\varphi = P \supset ((\neg C_3 \land \neg C_2) \supset (\mathbb{E}_{\langle ind \rangle} \mathsf{X}(C_3 \land \neg (C_2 \lor C_4) \supset \mathsf{AXAF}(C_3 \lor C_2))))$. It is apparent that $T_{i+1} \models T_i$ and $T_i \models P \supset \mathbb{E}_{\langle ind \rangle} \mathsf{X}(\neg l \lor C_2 \lor C_4)$.

We will show that $\forall (\mathcal{M}, s_0) \in Mod(T_i)$ there is an initial Ind-structure (\mathcal{M}', s'_0) such that $(\mathcal{M}', s'_0) \models T_{i+1}$ and $(\mathcal{M}', s_0') \leftrightarrow_{\langle V', \emptyset \rangle} (\mathcal{M}, s_0)$

 $\forall (\mathcal{M}, s) \models T_i \text{ we suppose } (\mathcal{M}, s) \models P \land \neg C_3 \land \neg C_2 \text{ and }$ $(\mathcal{M}, s_1) \models C_3 \land \neg C_2 \land \neg C_4 \text{ with } (s, s_1) \in [ind] \text{ (due to oth-}$ er case can be proved easily). Then we have $(\mathcal{M}, s) \nvDash l$ (by $(\mathcal{M}, s) \models l \supset C_3 \vee C_2$) and $(\mathcal{M}, s_1) \models l_1$ (by $(\mathcal{M}, s) \models l_2$ $P \supset \mathbb{E}_{\langle ind \rangle} X(l_1 \vee C_4)$). If $(\mathcal{M}, s_1) \nvDash AXAF(C_3 \vee C_2)$ then we have $(\mathcal{M}, s_1) \models l$ due to $(\mathcal{M}, s) \models AG(l \supset C_3 \vee C_2)$ and $(\mathcal{M}, s) \models AFl$. And then $(\mathcal{M}, s_1) \models \neg l_1$ by $(\mathcal{M}, s) \models$ $AG(l \supset \neg l_1 \lor C_2)$. It is contract with our supposing. Then $(\mathcal{M}, s_1) \models AXAF(C_3 \lor C_2).$

The Removing_atoms process For eliminating those irrelevant atoms, we define the following Removing operator.

Definition 5 (Removing_atoms) Let T be a set of formulae, $C \in T$ and V a set of atoms, then the elimination operator is defined as:

$$\operatorname{Removing_atoms}(C,V) = \left\{ \begin{array}{ll} \top, & if \ \mathit{Var}(C) \cap V \neq \emptyset \\ C, & else. \end{array} \right.$$

Which means that if the formula C containing the atoms in V then let C is true, else itself. For convenience, for any set T of formula we have Removing_atoms(T, V) = $\{Removing_atoms(r, V)|r \in T\}.$

Proposition 6 Let $V'' = V \cup V'$, $\Gamma = \text{Instantiate}(Res, V')$ and Γ_1 = Removing_atoms (Connect(Γ), Γ), then $\Gamma_1 \equiv_{\langle V'', \emptyset \rangle} Res \ and \ \Gamma_1 \equiv_{\langle V'', I \rangle} \varphi.$

Proof: Note the fact that for each clause $C = T \supset H$ in $Connect(\Gamma)$, if $\Gamma \cap Var(C) \neq \emptyset$ then there must be an atom $p \in \Gamma \cap Var(H)$. It is apparent that $Connect(\Gamma) \models \Gamma_1$, we will show $\forall (\mathcal{M}, s_0) \in Mod(\Gamma_1)$ there is a (\mathcal{M}', s_0) such that $(\mathcal{M}', s_0) \models Connect(\Gamma)$ and $(\mathcal{M}, s_0) \leftrightarrow_{\langle \Gamma, \emptyset \rangle} (\mathcal{M}', s_0)$. Let $C = T \supset H$ in $Connect(\Gamma)$ with $\Gamma \cap Var(C) \neq \emptyset$, $\forall (\mathcal{M}, s_0) \in Mod(\Gamma_1)$ we construct (\mathcal{M}', s_0) as (\mathcal{M}, s_0) except for each $s \in S$, if $(\mathcal{M}, s) \nvDash T$ then L'(s) = L(s), else:

- (i) if $(\mathcal{M}, s) \models H$, then L'(s) = L(s);
- (ii) else if $(\mathcal{M}, s) \models T$ with $p \in Var(H) \cap V$, then if p appearing in H negatively, then if C is a global (or an initial) clause then let $L'(s) = L(s) \setminus \{p\}$ else let $L'(s_1) = L(s_1) \setminus \{p\}$ for (each (if C is an A-step or A-sometime clause)) $(s, s_1) \in R$, else if C is a global (or an initial) clause then let $L'(s) = L(s) \cup \{p\}$ else let $L'(s_1) = L(s_1) \cup \{p\}$ for (each (if C is a A-step or A-sometime clause)) $(s, s_1) \in R$.
- (iii) for other clause $C = Q \supset H$ with $p \in Var(H) \cap \Gamma$, we can do it as (ii).

It is apparent that $(\mathcal{M}, s_0) \leftrightarrow_{\langle \Gamma, \emptyset \rangle} (\mathcal{M}', s_0)$, we will show that $(\mathcal{M}', s_0) \models Connect(\Gamma)$ from the following two

(1) For (ii) talked-above, we show it from the form of SNF_{CTL}^g clauses. Supposing C_1 and C_2 are instantiate formula of Γ :

- (a) If C is a global clause, i.e. $C = \top \supset p \vee C_1$ with C_1 is a disjunction of literals (we suppose pappearing in C positively). If there is a $C' = \top \supset$ $\neg p \lor C_2 \in Connect(\Gamma)$, then there is $\top \supset C_1 \lor C_2 \in$ $Connect(\Gamma)$ by the resolution $((\mathcal{M}, s) \models C_2$ due to we have suppose $(\mathcal{M}, s) \nvDash C$). It is apparent that $(\mathcal{M}', s_0) \models C \wedge C'$.
- (b) If $C = T \supset \mathrm{E}_{\langle ind \rangle} \mathrm{X}(p \vee C_1)$. If there is a C' = $T' \supset E_{\langle ind \rangle} X(\neg p \lor C_2) \in Connect(\Gamma)$, then there is $T \wedge T' \supset E_{\langle ind \rangle} X(C_1 \vee C_2) \in Connect(\Gamma)$ by the resolution $((\mathcal{M}, s) \models \mathbb{E}_{\langle ind \rangle} \mathbf{X} C_2$ due to we have suppose $(\mathcal{M}, s) \nvDash C$). It is apparent that $(\mathcal{M}', s_0) \models C \wedge C'.$
- (c) Other cases can be proved similarly.
- (2) (iii) can be proved as (ii) due to the fact we point at the beginning.

Therefore, we have $\Gamma_1 \equiv_{\langle V'',\emptyset \rangle} Res$ by Proposition 2 and Proposition 5.

And then
$$\Gamma_1 \equiv_{\langle V'',I \rangle} \varphi$$
 follows.

Example 5 After removing the clauses that include atoms in $V = \{p\}$, the following clauses have been left:

In this case, if we do not specify $l,\,C_2,\,C_3$ and C_4 are instantiate formulae of Sub(Res, V'), it is easy to check that all results including $P\supset \mathrm{E}_{\langle ind \rangle}\mathrm{X}(\neg l \lor C_2 \lor C_4)$ and $P\supset$ $AX(\neg l \lor C_2 \lor C_4)$ obtained from the *Connect* process will be deleted in the Removing_atoms process.

Remove the Index and start

The Removing_index(RemA) process is to change the set RemA obtained above into a set of formulas without the index by using the equations in Proposition 7.

Proposition 7 Let P, P_i and φ_i be CTL formulas, then

- $\begin{array}{l} \mbox{\it (i)} \ \bigwedge_{i=1}^n (P \supset \mathbf{E}_{\langle ind \rangle} \mathbf{X} \varphi_i) \equiv_{\langle \emptyset, \{ind \} \rangle} P \supset \mathbf{E} \mathbf{X} \bigwedge_{i=1}^n \varphi_i, \\ \mbox{\it (ii)} \ \bigwedge_{i=1}^n (P_i \quad \supset \quad \mathbf{E}_{\langle ind \rangle} \mathbf{X} \varphi_i) \quad \equiv_{\langle \emptyset, \{ind \} \rangle} \\ \ \bigwedge_{e \in 2^{\{0, \dots, n\}} \backslash \{\emptyset\}} (\bigwedge_{i \in e} P_i \supset \mathbf{E} \mathbf{X} (\bigwedge_{i \in e} \varphi_i)), \end{array}$
- $\begin{array}{c} \textit{(iii)} \ \bigwedge_{i=1}^n (P \supset \mathsf{E}_{\langle ind \rangle} \mathsf{F} \varphi_i) \equiv_{\langle \emptyset, \{ind \} \rangle} P \supset \bigvee \mathsf{EF}(\varphi_{j_1} \land \mathsf{EF}(\varphi_{j_2} \land \mathsf{EF}(\cdots \land \mathsf{EF} \varphi_{j_n}))), \textit{ where } (j_1, \ldots, j_n) \textit{ are } \\ \end{array}$ sequences of all elements in $\{0, \ldots, n\}$,
- (iv) $P \supset (C \vee E_{\langle ind \rangle} X \varphi_1) \wedge P \supset E_{\langle ind \rangle} X \varphi_2 \equiv_{\langle \emptyset, \{ind \} \rangle}$ $P \supset ((C \land \text{EX}\varphi_2) \lor \text{EX}(\varphi_1 \land \varphi_2)),$
- (v) $P \supset (C \vee \mathbb{E}_{\langle ind \rangle} \mathbf{X} \varphi_1) \vee P \supset \mathbb{E}_{\langle ind \rangle} \mathbf{X} \varphi_2 \equiv_{\langle \emptyset, \{ind \} \rangle}$ $P \supset (C \vee \text{EX}(\varphi_1 \vee \varphi_2)).$

Proof: It is easy to check.

Lemma 1 (*NI-BRemain*) Let I be the set of indexes appearing in RemA, we have RemA $\equiv_{\langle \emptyset, I \rangle}$ Removing_index(RemA).

Proof: It is easy checking that from the definition of *Removing_index* by Proposition 7.

In our Example 5 we do not need this process since there is no index in the set of formulae. Let T be a set of SNF_{CTL}^g clauses, then we define the following operator:

$$T_{\text{CTL}} = \{C | C' \in T \text{ and } C = D \text{ if } C' \text{ is the form } AG(\text{start} \supset D), \text{ else } C = C'\}.$$

Then $T \equiv T_{\text{CTL}}$ by $\varphi \equiv \text{AG}(\text{start} \supset \varphi)$ (Bolotov 2000).

The last step of our algorithm is to eliminate all the atoms in V' which has been introduced in the Transform process. Let $\Gamma = Instantiate(Res, V')$ and $\Gamma_1 = Removing_atoms(Connect(\Gamma))$, then $Replacing_atoms(Removing_index(\Gamma_1))$ is obtained from $Removing_index(\Gamma_1)$ by doing the following three steps for each $p \in (V' \setminus \Gamma)$:

- replacing each $p \supset \varphi_1 \lor \cdots \lor p \supset \varphi_n$ with $p \supset \bigvee_{i=1}^n \varphi_i$;
- replacing $p\supset \varphi_1\wedge\cdots\wedge p\supset \varphi_m$ with φ_j are instantiate formulae of Γ $(j\in\{1,\ldots,m\})$ with $p\supset \psi$, where $\psi=\bigwedge_{j=1}^m\varphi_j$ and p do not appear in φ_j , .
- For any formula $C \in \Gamma_1$, replacing every p in C with ψ .

Recall that any atom in V' introduced in the Transform process is a name of the sub-formula of φ (Bolotov 2000). Apparently, this process is just a process of replacing each atom with an equivalent formula. Then we have:

 $\begin{array}{lll} \textbf{Proposition 8} \ \textit{Let} & \Gamma_1 &= \text{Instantiate}(\textit{Res}, V'), \\ \Gamma_2 &= \text{Removing_atoms} & (\text{Connect}(\Gamma_1), \Gamma_1) & \textit{and} \\ \Gamma_3 &= \text{Replacing_atoms}(\text{Removing_index}(\Gamma_2)), & \textit{then} \\ \Gamma_2 \equiv_{\langle V' \setminus \Gamma_1, I \rangle} & \Gamma_3 \textit{ and } \varphi \equiv_{\langle V \cup V', \emptyset \rangle} & (\Gamma_3)_{\textit{CTL}}. \end{array}$

Example 6 By using the *Replacing_atoms* process on result of Example 5 directly since there is not index in those clauses, we obtain that x is replaced by $f \lor m$ at first, then y is replaced by $q \land AX(q \lor f \lor m)$ and z is replaced by $r \land (f \lor m \lor q) \land (f \lor m \lor (q \land AX(f \lor m \lor q))) \land AF(f \lor m)$.

An example for Connect process

In order to show the necessity of the Connect process, we see the following example at first.

Example 7 Let $\psi = AF(p \land q) \land EX \neg p$ and $V = \{p\}$. By the processes Transform and Resolution, we can obtain $V' = \{f, z\}$ and the following set Res of SNF_{CTL}^g clauses.

$$\begin{array}{lll} \mathbf{start} \supset z & z \supset \mathsf{AF}f & z \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X} \neg p \\ & \top \supset \neg f \lor p & \top \supset \neg f \lor q & z \supset \mathsf{E}_{\langle ind \rangle} \mathsf{X} \neg f \end{array}$$

On the one hand, according to our Algorithm 1, we have Instantiate(Res, V') = V since f can be instantiated by q and z can be instantiated by AFf.

In the *Connect* process, by using **EF1** rule on the Res we have $\alpha = z \supset (\neg q \supset (\mathbb{E}_{\langle ind \rangle} \mathbf{X}(q \supset \mathbf{AXAF}q)))$ and replace $z \supset \mathbb{E}_{\langle ind \rangle} \mathbf{X} \neg f \in Res$ with $z \supset \mathbb{E}_{\langle ind \rangle} \mathbf{X} \neg f \lor \alpha$

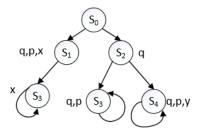


Figure 1: A model (\mathcal{M}, s_0) of φ

since l, C_2 , C_3 and C_4 are instantiate formulae. Apparently, $z \supset \operatorname{E}_{\langle ind \rangle} \operatorname{X} \neg f \lor \alpha \equiv z \supset q \lor \operatorname{E}_{\langle ind \rangle} \operatorname{X} (\neg f \lor \neg q \lor \operatorname{AXAF} q)$. After the *Removing_atoms* process, we have the following set *RemA* of formulae:

$$\begin{array}{ll} \mathbf{start} \supset z & z \supset \mathsf{AF}f \\ \top \supset \neg f \lor q & z \supset q \lor \mathsf{E}_{\langle ind \rangle} \mathsf{X}(\neg f \lor \neg q \lor \mathsf{AXAF}q) \end{array}$$

Removing the indexes appearing in the *RemA*, we obtain the following set NI:

$$\begin{array}{ll} \mathbf{start} \supset z & z \supset \mathsf{AF} f \\ \top \supset \neg f \lor q & z \supset q \lor \mathsf{EX} (\neg f \lor \neg q \lor \mathsf{AXAF} q) \end{array}$$

Replacing the atoms in V' that have been instantiated, i.e. f is replaced with q and z is replaced with $AFq \land (q \lor EX(\neg q \lor AXAFq))$, we have

$$Rp = \{ \mathbf{start} \supset \mathsf{AF}q \land (q \lor \mathsf{EX}(\neg q \lor \mathsf{AXAF}q)) \}.$$

As all the formulas ${\mathcal F}$ in the T_φ are the form ${\rm AG}{\mathcal F},$ hence we have:

$$\mathit{Rp}_{\mathsf{CTL}} = \{\mathsf{Af}q \land (q \lor \mathsf{ex}(\neg q \lor \mathsf{axaf}q))\}.$$

i.e. $ERes(\varphi, V) = AFq \land (q \lor EX(\neg q \lor AXAFq))$. In this case, we can easily check that $ERes(\varphi, V) \equiv_{\langle V, \emptyset \rangle} \varphi$.

On the other hand, if we do not using the *Connect* process, we can easily obtain the result of *ERes*, i.e. $ERes(\varphi, V) = AFq \land EX(\neg q)$. It is apparent that $ERes(\varphi, V) \not\equiv_{\langle V, \emptyset \rangle} \varphi$. This can proved by model (\mathcal{M}, s_0) as in Figure 1 since $(\mathcal{M}, s_0) \models \varphi$ and $(\mathcal{M}, s_0) \not\models ERes(\varphi, V)$.

This example shows that why we introduce the **EF** imply rules. Intuitively, the result of replacing the atoms that have been instantiated in V' with an instantiate formula is more stronger than our method, because by the Removing_atoms process, we have removing some clauses, such as $C = \top \supset$ $\neg f \lor p$, that contain f. The original one is $f \supset p \land q$, but after removing C we only obtain that $f \supset q$. In this example, there is a clause $z \supset \text{EX} \neg f \in Res$, after replacing f with q, we obtain $z \supset \text{EX} \neg q$. However, if we do not removing C (i.e. $f \supset p \land q$), then we have $z \supset EX(\neg q \lor \neg p)$, this is weaker than $z \supset \text{EX} \neg q$. In fact, for any model (\mathcal{M}, s_0) of φ there is not necessary $q \notin L(s)$ for some next state s of s_0 and if there is $q \in L(s)$ for all next states s, then there must be a next state s of s_0 with $p \notin L(s)$ s.t. for all next state s' of s there is $(\mathcal{M}, s') \models AFq$ (see Fig. 1). This is what the meanning of the *Connect* process.

The Correction and Complexity of the Algorithm

In the case that formula dose not include index, we use model structure $\mathcal{M}=(S,R,L,s_0)$ to interpret formula instead of Ind-model structure.

The correction means that the result $ERes(\varphi, V)$ obtained from our Algorithm is $F_{CTL}(\varphi, V)$, i.e. input φ and V to Algorithm 1 output the result of forgetting V from φ .

Theorem 1 (Resolution-based CTL-forgetting) *Let* $V'' = V \cup V'$ and $\Gamma_1 = \text{ERes}(\varphi, V)$, then

$$\begin{aligned} &(i) \;\; \mathbf{F}_{\mathsf{CTL}}(\varphi, V'') \equiv \Gamma_1; \\ &(ii) \;\; \mathbf{F}_{\mathsf{CTL}}(\varphi, V) \equiv \bigwedge_{\psi \in \Gamma_1} \psi. \end{aligned}$$

Proof: (i) (⇒)
$$\forall (\mathcal{M}, s_0) \in Mod(F_{\text{CTL}}(\varphi, V''))$$

⇒ $\exists (\mathcal{M}', s_0') \in Mod(\varphi) \text{ s.t. } (\mathcal{M}, s_0) \leftrightarrow_{V''} (\mathcal{M}', s_0')$
⇒ $\exists (\mathcal{M}_1, s_1) \in Mod(\Gamma_1) \text{ s.t. } (\mathcal{M}_1, s_1) \leftrightarrow_{V''} (\mathcal{M}', s_0')$
from Proposition 8
⇒ $(\mathcal{M}, s_0) \leftrightarrow_{V''} (\mathcal{M}_1, s_1)$
⇒ $(\mathcal{M}, s_0) \models \Gamma_1$ (IR(Γ₁, V''))
 $(\Leftarrow) \forall (\mathcal{M}_1, s_1) \in Mod(\Gamma_1)$
⇒ $\exists (\mathcal{M}', s_0') \in Mod(\varphi) \text{ s.t. } (\mathcal{M}_1, s_1) \leftrightarrow_{V''} (\mathcal{M}', s_0')$
⇒ $(\mathcal{M}_1, s_1) \models F_{\text{CTL}}(\varphi, V'')$ (IR(F_{CTL}(φ, V''), V'') and $\varphi \models F_{\text{CTL}}(\varphi, V'')$)
(ii) It is obtained from (i) since IR(φ, V').

Then we can obtain the result of forgetting of Example 4:

$$\begin{split} \mathbf{F}_{\text{CTL}}(\varphi,\{p\}) &\equiv r \wedge (f \vee m \vee q) \wedge \text{AF}(f \vee m) \wedge \\ (f \vee m \vee (q \wedge \text{AX}(f \vee m \vee q))) \wedge \text{AG}((q \wedge \text{AX}(f \vee m \vee q))) \wedge \\ &\supset \text{AX}(f \vee m \vee (q \wedge \text{AX}(f \vee m \vee q))))). \end{split}$$

Proposition 9 Let φ be a CTL formula and $V \subseteq A$. The time and space complexity of Algorithm 1 are $O((m+1)2^{4(n+n')})$. Where $|Var(\varphi)| = n$, |V'| = n' (V' is set of atoms introduced in transformation) and m is the number of indices introduced during transformation.

Proof: It follows from the lines 19-31 of the algorithm 1, which is to compute all the possible resolution. The possible number of SNF_{CTL}^g clauses under the give V, V' and Ind is $(m+1)2^{4(n+n')}+(m*(n+n')+n+n'+1)2^{2(n+n')+1})$.

Related work

Resolution-based satisfiability of CTL

Deciding the satisfiability with resolution calculus in Propositional Linear Temporal Logic (PLTL) was firstly introduced in (Fisher 1991) and further discussed in (Fisher 1997; Fisher, Dixon, and Peim 2001). The main idea is that transforming any PLTL formula into the normal form, called Separated Normal Form (SNF) by introducing a new connective **start** that holds only at the beginning of time.

After that the Resolution-based satisfiability in CTL was proposed by Bolotov in (Bolotov 2000) at first and then be refined by Zhang in (Zhang, Hustadt, and Dixon 2009; 2014). In those papers, the main idea is also to transform any CTL formula into the normal form SNF_{CTL}^g . But the CTL is a kind of branch time temporal logic, they introduced the "index" besides **start** for that purpose.

All in all, a complete set of transformation and resolution rules had been proposed for both PLTL and CTL. And it shows that the transformation is satisfiability preserving and also for the result obtained from using the resolution rules on the normal form.

Using Resolution Computing forgetting

Resolution, a kind of methods of Second-order quantifier elimination, has been used to compute the forgetting or uniform interpretation in different propositional logic (Wang 2015) and Modal logic (Herzig and Mengin 2008). In those case, the formula is required to be a paradigm with a particular form-"CNF" (the definition of CNF in Modal logic can be found in (Herzig and Mengin 2008)).

As have said above that the normal form used to resolution is an extension of CTL by introducing the **start** and "index". In this article, we propose the $\langle V,I \rangle$ -bisimulation to solve the "index" problem. Besides, in order to eliminate those atoms introduced in the transformation, we proposed the four EF imply rules.

Conclusion and Future Work

This article proposed a resolution-based algorithm to compute the forgetting in CTL. Our method extend the resolution calculus in (Zhang, Hustadt, and Dixon 2014) by adding processes including removing those irrelevant atoms and transforming the result into CTL formula. For this purpose, a kind of binary bisimulation relation, called $\langle V,I\rangle$ -bisimulation, has been defined and four EF imply rules have been proposed. More important, our algorithm is correct, i.e. return the result of forgetting some set of atoms, and the time and space complexity of Algorithm 1 are $O((m+1)2^{4(n+n')})$. Besides, examples show how to compute forgetting using our algorithm.

In the future we will implement this algorithm (part of it has been implement actually).

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