

The System **BiOC** for Reasoning about Biological Models in Action Language \mathcal{C}^*

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Abstract

We elaborate upon the usage of action language \mathcal{C} for representing and reasoning about biological models. First, we provide a simple extension of \mathcal{C} allowing for variables and show its usefulness in modeling biochemical reactions according to the well-known model of **BIOCHAM**. Second, we show how the biological action description language \mathcal{C}_{TAID} can be mapped onto \mathcal{C} . Finally, we describe a toolbox for using action languages, including among them, a compiler mapping \mathcal{C} and \mathcal{C}_{TAID} to logic programs under answer sets semantics along with a web-service integrating different front- and back-ends for addressing dynamical systems by means of action description languages via answer set programming. This is accompanied by an empirical evaluation with existing systems for processing action description languages.

1 Introduction

We elaborate upon *action languages* [13] for qualitative modeling of biological networks. Action languages are formal models used for reasoning about the effects of actions, while being close to natural language. Central to this approach to formalizing actions is the concept of a transition system, which constitutes its semantic underpinning. The first action language for representing and reasoning about biological networks was introduced in [19, 2, 18] and further extended in [6] leading to action language \mathcal{C}_{TAID} .

In what follows, we extend the overall approach in several ways while centering it on the classical action language \mathcal{C} [15]. To begin with, we provide a simple extension of \mathcal{C} allowing for variables and show its usefulness in modeling biochemical reactions according to the well-known model of **BIOCHAM**. Similar to the approach taken in the dlv^k system based on action language \mathcal{K} [7], we delegate the treatment of variables to Answer Set Programming (ASP; [1]) in order to be able to use ASP grounders for variable instantiation. Second, we provide a translation

mapping the biologically motivated action description language \mathcal{C}_{TAID} onto \mathcal{C} and give a result fixing the formal correspondence. This allows us to further develop \mathcal{C}_{TAID} within a broader and well-established framework, avoiding further dedicated implementations. Moreover, it provides \mathcal{C}_{TAID} with access to further implementations of \mathcal{C} , like *CCalc* [14] or *CPlan* [4] (even though they cannot harness existing ASP grounders for variable treatment). Finally, we describe a toolbox for using action languages, including among them, a compiler mapping \mathcal{C} and \mathcal{C}_{TAID} to logic programs under answer sets semantics along with a web-service integrating different front- and back-ends for addressing dynamical systems by means of action description languages via answer set programming. Our tools are designed for an easy and flexible integration with existing open source tools via pipes, in particular, ASP grounders and solvers, as well as further front- and back-ends. This is accompanied by an empirical evaluation with existing systems for processing action description languages.

2 Background

Answer Set Programming. Our language is built from a set \mathcal{F} of *function* symbols (including the natural numbers), a set \mathcal{V} of *variable* symbols, and a set \mathcal{P} of *predicate* symbols. The set \mathcal{T} of *terms* is the smallest set containing \mathcal{V} and all expressions of the form $f(t_1, \dots, t_n)$, where $f \in \mathcal{F}$ and $t_i \in \mathcal{T}$ for $1 \leq i \leq n$. The set \mathcal{A} of *atoms* contains expressions of the form $p(t_1, \dots, t_n)$, where $p \in \mathcal{P}$ and $t_i \in \mathcal{T}$ for $1 \leq i \leq n$. A *literal* is an atom a or its negation $\neg a$; both can be preceded by default negation, denoted as *not* a and *not* $\neg a$, respectively. For $a \in \mathcal{A}$, we let $\bar{a} = \neg a$ and $\overline{\bar{a}} = a$. A *logic program* over \mathcal{A} is a set of *rules* as

$$a \leftarrow b_1, \dots, b_m, \text{not } c_{m+1}, \dots, \text{not } c_n \quad (1)$$

where a, b_i, c_j are literals over \mathcal{A} for $0 < i \leq m < j \leq n$. For a rule r as in (1), let $\text{head}(r) = a$, $\text{body}(r)^+ = \{b_1, \dots, b_m\}$, and $\text{body}(r)^- = \{c_{m+1}, \dots, c_n\}$. Given an expression $e \in \mathcal{T} \cup \mathcal{A}$, let $\text{var}(e)$ denote the set of all variables occurring in e ; analogously, $\text{var}(r)$ gives all variables in rule r . The *ground instantiation* of a program P is defined as $\text{grd}(P) = \{r\theta \mid r \in P, \theta : \text{var}(r) \rightarrow \mathcal{U}\}$, where

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$\mathcal{U} = \{t \in \mathcal{T} \mid \text{var}(t) = \emptyset\}$; analogously, $\text{grd}(\mathcal{A}) = \{a \in \mathcal{A} \mid \text{var}(a) = \emptyset\}$ is the set of all ground atoms. A set $X \subseteq \text{grd}(\mathcal{A}) \cup \overline{\text{grd}(\mathcal{A})}$ is a (consistent) *answer set* of a program P over \mathcal{A} , if X is the \subseteq -smallest model of

$$\{\text{head}(r) \leftarrow \text{body}(r)^+ \mid r \in \text{grd}(P), \text{body}(r)^- \cap X = \emptyset\}.$$

Action Language \mathcal{C} . Action languages use *fluents* to describe the states of a system and *actions* influence the values of fluents. In \mathcal{C} (and \mathcal{C}_{TAID}), *static laws* describe properties between fluents that need to be satisfied in every state of the system. *Dynamic laws* describe the effects of actions, that is, how the system evolves when actions are executed.

More formally, we consider *action language \mathcal{C}* [13] over a Boolean *action signature* $\langle B, F, A \rangle$, where B is the set $\{f, t\}$ of truth values, F is a set of *fluent names*, and A is a set of *action names*. In \mathcal{C} , an *action description* D_C over a signature $\langle B, F, A \rangle$ consists of *static* and *dynamic laws*:

$$\text{(caused } \varphi \text{ if } \psi) \quad (2)$$

$$\text{(caused } \varphi \text{ if } \psi \text{ after } \omega) \quad (3)$$

where φ and ψ are propositional combinations of fluent names and ω is a propositional combination of fluent and action names. Every action description D_C induces a unique transition system $\mathcal{T}_C(D_C) = \langle S, V, R \rangle$, where S is a set of *states*, V is a function determining fluents values in state s , and R is a relation containing all possible transitions between states. A *trajectory* $s_0, A_1, s_1, \dots, s_{n-1}, A_n, s_n$ in a transition system $\langle S, V, R \rangle$ is a sequence of sets of actions $A_i \subseteq A$ and states $s_i \in S$ where $(s_{i-1}, A_i, s_i) \in R$ for $0 \leq i \leq n$. Intuitively, a trajectory represents one possible history (or simply path) within a transition system. In [13], several syntactic extensions are defined. For instance, the rule (ω **may cause** φ **if** ψ) is a shorthand for **(caused φ if φ after $\psi \wedge \omega$)**. Similarly, **(inertial φ)** is a shorthand for **(caused φ if φ after φ)**. We refer to [13] for more detailed definitions.

Besides an action description language, both \mathcal{C} and \mathcal{C}_{TAID} define a *query language*. We implemented \mathcal{R} [13] as the query language for \mathcal{C} and the query mechanisms described in [6] for \mathcal{C}_{TAID} . In this paper, we focus only on the transition systems and our toolbox realizing the different encodings, so we omit a detailed description of query languages and the different reasoning modes, like explanation, prediction, planning, etc.

3 Encoding Action Language \mathcal{C}

For implementing action language \mathcal{C} , we build upon the translation to ASP described in [17]. Let D_C be an action description over signature $\langle B, F, A \rangle$. We require D_C to be *definite*, that is, the heads φ of laws **(caused φ if ψ)** and **(caused φ if ψ after ω)** are fluent literals (or the constant

\perp). Furthermore, ψ is a conjunction of fluent literals and ω is a conjunction of fluent and/or action literals. In what follows, we denote φ by f , ψ by $g_1 \wedge \dots \wedge g_m$ and ω by $l_1 \wedge \dots \wedge l_n$.

We define a logic program $lp_n(D_C)$ whose answer sets correspond to trajectories of length n in the transition system induced by D_C . $lp_n(D_C)$ contains atoms $a(t)$ and $f(t)$ for each $a \in A$, $f \in F$ and $t = 0, \dots, n$. For each static law **(caused f if $g_1 \wedge \dots \wedge g_m$)** in D_C , $lp_n(D_C)$ contains for each $t = 0, \dots, n$ a rule $f(t) \leftarrow \text{not } g_1(t), \dots, \text{not } g_m(t)$. Analogously, each dynamic law **(caused f if $g_1 \wedge \dots \wedge g_m$ after $l_{m+1} \wedge \dots \wedge l_n$)** in D_C , adds to $lp_n(D_C)$ for each $t = 0, \dots, n-1$ a rule

$$f(t+1) \leftarrow \text{not } \overline{g_1(t+1)}, \dots, \text{not } \overline{g_m(t+1)}, l_{m+1}(t), \dots, l_n(t).$$

Furthermore, $lp_n(D_C)$ contains

$$\begin{aligned} \neg a(t) &\leftarrow \text{not } a(t), & \neg e(0) &\leftarrow \text{not } e(0), \\ a(t) &\leftarrow \text{not } \neg a(t), & e(0) &\leftarrow \text{not } \neg e(0) \end{aligned}$$

for each $a \in A$, $t = 0, \dots, n$ and each $e \in F$.

Our implementation of the encoding allows to use variables when writing rules in \mathcal{C} . This is done by delegating the grounding of variables to the grounding process of the underlying logic program. To this end, we start by extending the syntax of \mathcal{C} by a trailing keyword **where** followed by domain predicates for binding the variables occurring in the actual causal laws. To be precise, the causal laws in (2) and (3) are extended as follows:

$$\text{(caused } \varphi \text{ if } \psi \text{ where } \delta) \quad (4)$$

$$\text{(caused } \varphi \text{ if } \psi \text{ after } \omega \text{ where } \delta) \quad (5)$$

where φ, ψ , and ω are as defined in (2) and (3), except for containing variables, and δ is a combination of non-fluent and non-action atoms such that $\text{var}(\varphi) \cup \text{var}(\psi) \cup \text{var}(\omega) \subseteq \text{var}(\delta)$. Intuitively, δ captures static domain information used for binding the variables in φ, ψ, ω . The concept of a definite action description generalizes in the obvious way, restricting δ to conjunctions of non-fluent and non-action atoms. Now, given such a definite action description D_C , the variable-tolerating extension of $lp_n(D_C)$ is obtained from $lp_n(D_C)$ by extending the body of each resulting logic programming rule by d_1, \dots, d_o whenever the causal law contains the condition **where** $d_1 \wedge \dots \wedge d_o$.

Let us illustrate the practical impact of this pragmatic extension by modeling the Biochemical Abstract Machine (BIOCHAM; [9, 5]), used to build biochemical systems. The biological background is indeed very easy. A modeled scenario consists of different chemical reactions that specify relations between different compounds. *Reactants* are compounds that need to be present that a reaction can take place and *products* are compounds that will be present after a reaction took place. One can model this scenario using \mathcal{C} with the following rules. At first, our syntax requires to specify a preamble where actions and fluents are defined:

```
<action> occurs(R) <where> reaction(R).
<fluent> present(P) <where> compound(P).
```

Strings enclosed in `<>` are keywords, variables start with uppercase letters and lines end with a dot. That is, for every term t belonging to the extension of the predicate `reaction`, we introduce the actions `occurs(t)`. For every term t belonging to the extension of the predicate `compound`, we introduce the fluents `present(t)`.

We now can define the dynamics of the system:

```
<caused> present(P)
  <after> occurs(R)
<where> reaction(R), compound(P), product(P,R).

<caused> <false>
  <after> occurs(R), -present(P)
<where> reaction(R), compound(P), reactant(P,R).

occurs(R) <may cause> -present(P)
<where> reaction(R), compound(P), reactant(P,R).

<inertial> present(P) <where> compound(P).
<inertial> -present(P) <where> compound(P).
```

The first rule states that a compound P is present after a reaction R occurred producing P . The second rule is a constraint enforcing all compounds P to be present if a reaction occurs where P is a reactant of. Note that negation is denoted as `-` and `<false>` as well as `<true>` are keywords for the two Boolean constants. The third rule models a certain non-determinism: The semantics of BIOCHAM defines that after a reaction occurs, it remains unclear whether the reactants are still present or not. The reason is that the semantics abstract from concentrations of compounds. That is, we consider two cases: In one transition we assume that the compound P was fully consumed, modeled as `-present(P)`. The other transition is that P remains to be present. The last two rules state that compounds that are not affected by reactions do not change their value¹.

Let us briefly detail how variables are passed through the encoding proposed in [17]. For this, consider the first rule of the above BIOCHAM example that is translated to the following logic rule:

```
present_fluent(P,T+1)
:- occurs_action(R,T), reaction(R),
   compound(P), product(P,R), time(T).
```

Apart from the time-parameter T , we attach variable P to the fluent `present` and R to the action `occurs`. The domain information given in the `<where>` statement is then passed as grounding information to the logic program rule. The last pending issue is to specify the domains, eg.

¹These rules represents the frame axiom: Compounds that are not consumed remain present, absent compounds that where not produced remain absent.

```
compound(a). compound(b). reaction(r1).
reactant(a,r1). product(b,r1).
```

The database is represented as a logic program. It can be seen as static knowledge attached to the modeled dynamic behavior of the system. In most cases, the database only contains facts. In the example, we are now able to reason about a scenario with two compounds and one reaction. An encoding of a simple version of the biological textbook example of the *Mitogen-activated protein kinase (MAPK)*² including 23 products and 30 reactions, yields a problem instance containing 147 facts. One of the advantages of using variables is the resulting elaboration tolerance, that is, the biological system can be easily exchanged or extended by modifying the database, while leaving the specification of the dynamics untouched.

4 Mapping \mathcal{C}_{TAID} to \mathcal{C}

As with \mathcal{C} , an *action description* in \mathcal{C}_{TAID} is given relative to an action signature $\langle B, F, A \rangle$. The major conceptual difference between \mathcal{C}_{TAID} and \mathcal{C} is that the latter implicitly treats actions to be exogenous. That is, all actions might occur at every time-point as long as their effects do not lead to a contradiction. For biological purposes, this behavior is inappropriate. Unlike this, \mathcal{C}_{TAID} allows for specifying explicit conditions when actions are executed or not. For example, using \mathcal{C}_{TAID} 's *triggering rule*, we can describe properties when (re)actions must be executed immediately. Furthermore, \mathcal{C}_{TAID} offers the following constructs: *Inhibition rules* express when actions must not be executed and *allowance rules* express that actions might occur, but are not forced to. A *default* expresses that a fluent takes a certain value unless it is known otherwise. *No-concurrency constraints* allow to control the parallel execution of actions. In a more formal way, an action description in \mathcal{C}_{TAID} contains expressions of the following form:

$(a \text{ causes } \varphi \text{ if } \psi), (\varphi \text{ if } \psi), (\varphi \text{ triggers } a),$
 $(\varphi \text{ allows } a), (\varphi \text{ inhibits } a), (\text{default } f),$
 $(\text{noconcurrency } \omega),$

where a is an action and ω is either an action or a conjunction of action literals, φ and ψ are conjunctions of fluent literals and f is a fluent literal. We refer to [6] for a more detailed description of \mathcal{C}_{TAID} .

We now describe our translation of \mathcal{C}_{TAID} into \mathcal{C} . To this end, we need to extend the action signature to accommodate some control information. To be precise, we add the fluents $ih(a)$, $tr(a)$, $ex(a)$, $al(a)$ for each action name a . Intuitively, these fluents signal properties reflecting the behavior of inhibition, triggering, and allowance rules. With

²<http://en.wikipedia.org/wiki/MAPK>

them, we can define the mapping of rules in \mathcal{C}_{TAID} to rules in \mathcal{C} as follows.

Definition 1 Let $D_{\mathcal{C}_{TAID}}$ be an action description in \mathcal{C}_{TAID} over action signature $\langle B, F, A \rangle$. The corresponding action description $D_{\mathcal{C}}$ in \mathcal{C} over action signature

$$\langle B, F \cup \bigcup_{a \in A} \{ih(a), tr(a), ex(a), al(a)\}, A \rangle \quad (6)$$

is defined as follows:

1. For each action name $a \in A$, action description $D_{\mathcal{C}}$ contains the static laws (**caused** $\neg ih(a)$ **if** $\neg ih(a)$), (**caused** $\neg tr(a)$ **if** $\neg tr(a)$), (**caused** $\neg ex(a)$ **if** $\neg ex(a)$) and (**caused** $\neg al(a)$ **if** $\neg al(a)$).
2. For each dynamic law (a **causes** φ **if** ψ) in $D_{\mathcal{C}_{TAID}}$, where $\varphi = f_1 \wedge \dots \wedge f_m$, $D_{\mathcal{C}}$ contains the laws (**caused** f_i **if** \top **after** $\psi \wedge a$) for each f_i where $1 \leq i \leq m$.
3. For each static law (φ **if** ψ) in $D_{\mathcal{C}_{TAID}}$, where $\varphi = f_1 \wedge \dots \wedge f_m$, $D_{\mathcal{C}}$ contains the laws (**caused** f_i **if** ψ) for each f_i where $1 \leq i \leq m$.
4. For each allowance rule (φ **allows** a) in $D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** $al(a)$ **if** φ) and (**caused** \perp **if** \top **after** $\neg al(a) \wedge a$).
5. For each triggering rule (φ **triggers** a) in $D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** $tr(a)$ **if** φ), (**caused** $ex(a)$ **if** \top **after** a), (**caused** \perp **if** $\neg ex(a)$ **after** $tr(a) \wedge \neg ih(a)$) and (**caused** \perp **if** $ex(a)$ **after** $\neg tr(a)$).
6. For each inhibition rule (φ **inhibits** a) in $D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** $ih(a)$ **if** φ) and (**caused** \perp **if** \top **after** $ih(a) \wedge a$).
7. For each constraint (**noconcurrency** ω) in $D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** \perp **if** \top **after** ω).
8. For each default rule (**default** f) in $D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** f **if** f).
9. For each $f \in F$, such that (**default** f) $\notin D_{\mathcal{C}_{TAID}}$, $D_{\mathcal{C}}$ contains (**caused** f **if** f **after** f) and (**caused** $\neg f$ **if** $\neg f$ **after** $\neg f$).

The symbols \top and \perp denote the Boolean constants for t and f in B .

The rules in 1. state that $ih(a)$, $tr(a)$, $ex(a)$, and $al(a)$ are set to be *false* by default. As described in 4.–6., they are only set *true* when certain properties hold. There is a direct correspondence between static and dynamic rules in \mathcal{C}_{TAID} and \mathcal{C} (cf. 2. and 3.) except the fact that conjunctions in heads are split in order to get a definite action description.

An allowance rule is expressed using a static rule setting $al(a)$ and a dynamic rule that can be viewed as a constraint eliminating transitions where action a occurred while $al(a)$ was *false* (cf. 4.). Rules given in 5. express triggering rules: whenever a trigger is applicable, $tr(a)$ is set and every execution of an action a causes $ex(a)$ to be true. The second dynamic rule eliminates transitions where the conditions for a triggering rule were satisfied but a was not executed, that is, $ex(a)$ is *false*. Since \mathcal{C}_{TAID} gives inhibition rules priority over triggering rules, the constraint is only applicable if $\neg ih(a)$ is satisfied. The third dynamic rule eliminates transitions where a is executed without having an applicable trigger. Inhibition rules are mapped in the same way as allowance rules (cf. 6.). No-concurrency constraints and defaults in \mathcal{C}_{TAID} have a direct correspondence to rules in \mathcal{C} (cf. 7. and 8.). Given that fluents are implicitly inertial³ in \mathcal{C}_{TAID} but not in \mathcal{C} , for each fluent there is a dynamic rule in $D_{\mathcal{C}}$ that expresses inertial behavior (cf. 9.).

We can show the following result:

Theorem 1 Let $D_{\mathcal{C}_{TAID}}$ be an action description in \mathcal{C}_{TAID} over action signature $\langle B, F, A \rangle$ and let $D_{\mathcal{C}}$ be the corresponding action description in \mathcal{C} over the action signature in (6) generated from $D_{\mathcal{C}_{TAID}}$ using the mapping in Definition 1. Then, each trajectory in the transition system $\mathcal{T}_{\mathcal{C}}(D_{\mathcal{C}})$ (as defined in [13]), corresponds to a unique trajectory in the transition system induced by $D_{\mathcal{C}_{TAID}}$ (as defined in [6]) and vice versa.

Problem descriptions in \mathcal{C}_{TAID} can now be dealt with the general-purpose language \mathcal{C} . That is, we do not need a rather complicated (re)definition of semantics in order to describe transition systems having a biological background using \mathcal{C}_{TAID} . It can now be seen as another layer of interface on top of the action description language \mathcal{C} .

In the following sections we describe how the different encodings can be used in our toolchain and how they perform compared to other implementations.

5 The BioC System

Our approach to representing and reasoning about biological models is as follows: at first, the biological model needs to be specified in the action description language of \mathcal{C} or \mathcal{C}_{TAID} . This description is compiled into a logic program as described above and subsequently dealt with using an ASP system, usually composed of a grounder and a solver. Once the logic program is solved, the answers of the solver need to be put back in correspondence to the original problem specification. Finally, the obtained data needs to

³That is, fluents that are neither defaults nor affected directly or indirectly by dynamic rules do not change their value in a transition.

Accessible via web-interface

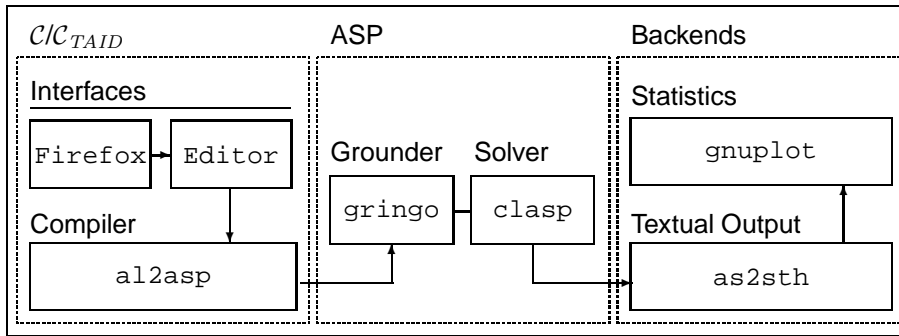


Figure 1. Overview of our system architecture

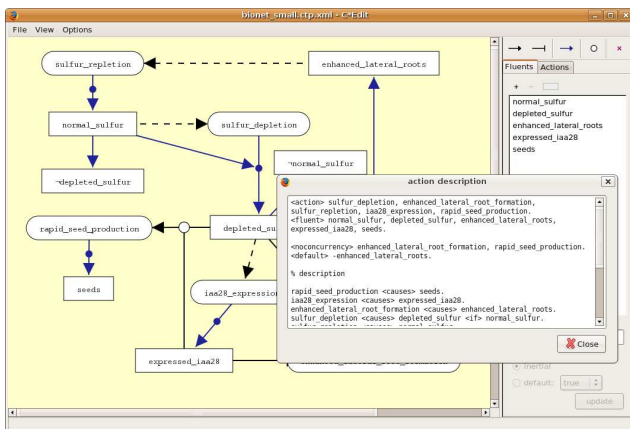


Figure 2. Graphical user interface for C_{TAID} .

be interpreted in a biologically meaningful way by a human expert. An overview of our system is given in Figure 1.

Interfaces To begin with, we have a closer look at the interfaces to our system. Our system is able to handle the discussed action descriptions in \mathcal{C} and C_{TAID} . For action descriptions in \mathcal{C} , one has to write down the rules in an editor, as shown in the BIOCHAM example. This is of course also possible using C_{TAID} . Since C_{TAID} has a much more biological orientation than \mathcal{C} , we offer another interface for C_{TAID} that is more intuitive for users having a purely biological background: A graphical interface that was built as a Firefox browser extension. It allows for building rules as a graph whose nodes (fluents and actions) and edges (causal relationships) correspond to the underlying expressions of C_{TAID} . An example is shown in Figure 2. Since this paper has more a technical orientation, we are not detailing a biological example using C_{TAID} .

Compiler Once the description is done, it is passed to our compiler `al2asp`. As mentioned before, this program is able to handle the described languages and their different encodings that need to be given via command line options:

```
al2asp -l c           direct  $\mathcal{C}$  to ASP encoding
al2asp -l c_taid     direct  $C_{TAID}$  to ASP encoding
al2asp -l c_taid2c   $C_{TAID}$  to  $\mathcal{C}$  encoding
```

While the two first commands yield a logic program⁴, the last one outputs rules in \mathcal{C} .⁵

`al2asp` is implemented in C++ and freely available at [3]. Notably, `al2asp` relies on scanner and parser generators `flex` and `bison++`, making it easily amenable to language extensions.

An `al2asp` generated logic program containing variables appears incomplete. The additional logic program providing the binding information must be concatenated to the output of `al2asp` in order to get the resulting logic program that can be grounded. This ground program expresses the transition system described by the original description in \mathcal{C} .

ASP Tools Reconsidering Figure 1, the resulting logic program is dealt with by an ASP system, consisting of a grounder and a solver component. As discussed, the logical representation of an action description may contain object variables that are passed on to the grounder. Our grounder, `gringo` [12]⁶, systematically replaces all variables by ground terms, while aiming at producing a compact propositional program. The resulting program is then passed to the ASP solver, `clasp` [11, 10]⁷, which computes the stable models (see [1] for details) of the program. Each such model represents a valid trajectory in the transition system induced by the original action description.

Backends The action description for the BIOCHAM system combined with the underlying domain induces the transition system given in Figure 3. Identifiers `a` and `b` are

⁴The direct C_{TAID} to ASP encoding implements a slightly modified encoding according to the one given in [6] that is not discussed in this paper.

⁵One can just reuse the tool to complete the encoding: `al2asp -l c_taid2c <file.desc> | al2asp -l c`.

⁶<http://www.cs.uni-potsdam.de/gringo>

⁷<http://www.cs.uni-potsdam.de/clasp>

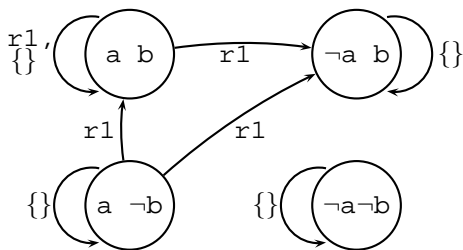


Figure 3. BIOCHAM Transition system.

shorthands for fluents `present(a)` and `present(b)`, `r1` is a shorthand for action `occurs(r1)`. `{}` denotes the empty action, that is, no action is executed in a transition labeled like this. Note that the loop at node `{a, b}` describes two transitions.

Given that fluent and action names are changed in the logical encoding (ie. an additional time parameter appears as additional argument) as well as the obtained solutions appear in an unsorted way, the output of an ASP system must be transformed in a more readable and problem-oriented format. To this end, we offer different possibilities to present the output using the program `as2sth`: One possibility is a textual representation of the trajectories that gives a detailed overview of actions and states involved in a given solution. To illustrate this, recall our BIOCHAM example and consider the answer sets representing all 8 trajectories of length 1. Our interface displays them as follows.

```
## ANSWER 1 #####
0 A + occurs(r1)
0 F + present(a)
0 F - present(b)
1 F - present(a)
1 F + present(b)
## ANSWER 2 #####
...
## SUMMARY #####
models: 8
```

The first column denotes the timestep, the second one the type of the logic literal (action or fluent), the third one the value of the literal (true or false) and the last one the original name as used in the action description.

This method becomes inapplicable when the number of solutions increases, which is the case in most of the biological applications. To this end, another possibility is to generate `csv` output that can be processed with external programs like database systems, statistical tools, etc.

A third possibility is to use our built in `gnuplot` interface: We currently provide some statistical post-processing counting fluent values and actions at each time step in all trajectories. For example, let us assume that a fluent f appears to be true at a certain timestep t in all trajectories. When presenting all occurrences of fluents (or actions) in a graphical way, one can easily see that fluent f is essential

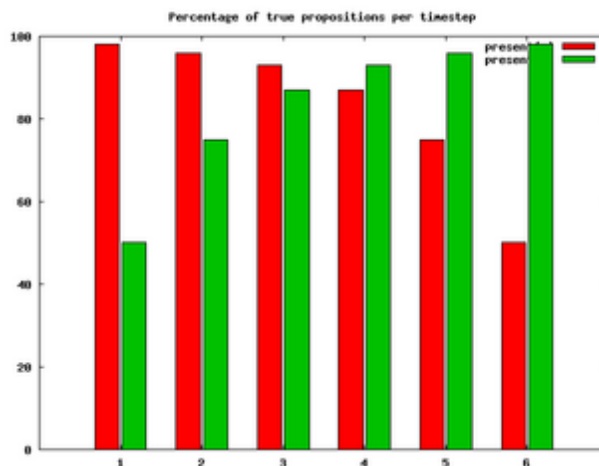


Figure 4. Solutions of the BIOCHAM example.

for having solutions.⁸ Although our simple BIOCHAM example focuses on the transition system (having no queries at all), the graphical representation can already be useful to get an idea. Reconsider the transition system given in Figure 3. Figure 4 is the graphical representation of all 128 trajectories having a length of 6. Y-Axis denotes the percentage of true propositions (resp. the presence of compounds), and X-Axis denotes the timesteps. The two different bars represent the compounds `a` and `b`. It is easy to see that there is a direct correspondence between the presence of compound `a` and `b`. While `a` tends to decrease, `b` tends to increase.⁹ It is nearly impossible to gain such information by only looking at the calculated trajectories.

Toolchain Access The whole reasoning tool is accessible in two ways. The first possibility is to download the tools described in Figure 1 from [3] and to run them on a local machine. We are building up a graphical tool wrapping the underlying command-line execution of the described tools. By now, given that the tools are available on a Linux machine, a user may start the different programs via pipelining by hand. For example, if we have a our BIOCHAM description in \mathcal{C} given in a file named `bcham.alc`, the domain specification given in a file named `bcham.stat` and want to display the chart as given in Figure 4 using `gnuplot`, you invoke on your local system the following commands:

```
> al2asp -l c bcham.alc | cat - bcham.stat \
gringo -c n=5 | clasp 0 | as2sth --csv | \
asplot present(a) present(b) && gnuplot *.plt
```

To provide a user-friendly method, we built up a web-based interface at [3], where the described tools are fully

⁸This can also be seen as a cautious reasoning mode.

⁹Indeed, this outcome seems to be trivial since this is exactly the relation between the two compounds that was modeled before. But on larger scale examples it is possible to identify relations that were not given explicitly in the action description.

encapsulated as a server application to use system without installing local applications. The C_{TAID} Firefox-Plugin is able to access the web interface directly by sending the underlying action description to the web server. We added several examples on our web interface, where one can see how descriptions and queries to the system look like and how a user is able to access the different backends.

6 Benchmarks

In this section, our core tools (`al2asp`, `gringo` and `clasp`) will be empirically compared to the systems $CCalc$ [14] and dlv^k [7] since all of them use input languages based on \mathcal{C} . Unfortunately, $CPlan$ [4] is no longer maintained and the authors provided a windows executable only which was not usable in our benchmark setting.

The benchmarks were carried out on an Intel Core2Duo 6400 with 2.13GHz and 2 GB RAM running a 32-bit version of Ubuntu GNU/Linux. For our tests, we used `al2asp` v0.4, `gringo` v1.0.0 and `clasp` v1.0.5 with default settings. $CCalc$ was used in version 2.0 and among the provided SAT solvers $grasp$ was used. Although $grasp$ does not provide the current state of the art SAT solving techniques, it was the only solver in our tests that produced all solutions. Regarding dlv^k , we used release 2007-10-11 with default settings.

Concerning planning problems, one is often interested in finding only the first solution. This issue is different in our approach, in most of the biological applications we need to consider all solutions. For example, recall Figure 4 where we need to process all answer sets in order to do statistical analysis. To this end, we consider both cases when comparing the systems, finding one, and finding all solutions.

Unfortunately, our current biological applications get solved too fast to make systems compareable. Being not generic¹⁰, a comparison of different systems using our biological problems is not yet feasible. We use crafted artificial problems instead to compare performance of systems.

The first one is the well known *blocks world* problem. We used the dlv^k encoding and problem instances from [8]. Due to advances in computer hardware, these old instances are solved too fast to get reasonable runtimes. That is why we came up with five additional instances (p6 - p10, see Table 1) which are still demanding for the systems running on today's hardware.

Our second benchmark suite *lights out*¹¹ is very similar to the *bomb in the toilet* problem: All of a variable number of light bulbs has to be switched off. The problem comes in two flavors, either with concurrent execution of actions

¹⁰Unlike most artificial problems, we do not have parameters controlling the size of problem instances.

¹¹Idea taken from General Gameplaying Competition 2008.

No.	Instance	length	BioC	$CCalc$	dlv^k
1	11nc	10	0.10	0.14	17.81
2	12nc	15	0.20	0.19	—
3	13nc	20	2.12	0.26	—
4	14nc	25	—	0.39	—
5	11c	1	8.63	—	2.43
6	12c	1	17.39	—	5.24
7	13c	1	26.41	—	8.09
8	14c	1	35.43	—	10.56
Average Time (Sum Timeouts)			12.90 (3)	0.24 (12)	8.82 (9)
Average Penalized Time			86.29	300.12	230.51

Table 2. Lights out experiments (one solution)

allowed or with concurrency disabled. The optimal¹² plan length in the latter case is equal to the number of light bulbs. It is easy to see that this problem leads to $n!$ many optimal plans regarding n bulbs that only differ in the sequence of switching off bulbs. Due to this behavior, we omit computing all solutions as in the blocks world setting.

The results of the *blocks world* benchmarks are listed in Table 1 and the *lights out* results are in Table 2. For every problem instance, we measured the time in seconds of three separate solving processes and computed the average which is shown in each systems column. A dash indicates that a system was unable to compute a solution in less than 600 seconds. The column labeled *length* denotes the length of the shortest possible plan(s) for the problem instance which is passed as a parameter to the different systems. The last row in the tables lists penalized average times. In contrast to normal average times, the penalized ones take timeouts into account. Although the system might have taken much longer to find a solution, the penalized average is computed as if the system found a solution after 600 seconds.

Results show that compared to the other systems our system performs quite well and appears to be robust. In the *blocks world* example, it was the only system that could enumerate all solutions in reasonable time. As mentioned, this issue is especially valuable because our biological applications often need all solutions to be computed. But also when only one solution has to be found, our system outperformed both $CCalc$ and dlv^k . $CCalc$'s performance was comparable to ours until the problems became too hard in benchmark number 9.

Regarding the *lights out* problem, $CCalc$ performs surprisingly well when concurrent execution of actions is not allowed. It computes a solution almost instantly, while dlv^k has difficulties even in the smallest instance. Although being quite fast with a few light bulbs, the runtime of our system rises rapidly as soon as more than twenty bulbs are involved. When allowing concurrency in this example, dlv^k is the fastest system. $CCalc$ seems to have great problems

¹²Optimal means that there is at least one solution at bound t , but no solution can at bound $t - 1$.

No.	Instance	length	BiOC - one	CCalc - one	dlv ^k - one	BiOC - all	CCalc - all	dlv ^k - all
1	p01	05	0.31	0.51	0.06	0.32	0.51	0.06
2	p02	06	0.22	0.35	0.05	0.22	0.42	0.05
3	p03	08	1.19	1.21	1.21	1.21	8.23	4.57
4	p04	09	3.89	3.88	1.17	4.05	5.74	15.19
5	p05	11	5.04	5.39	2.92	4.92	11.24	22.98
6	p06	13	4.21	3.88	21.15	4.78	408.23	—
7	p07	14	8.76	7.08	42.04	11.81	364.22	—
8	p08	16	36.88	14.28	—	133.49	—	—
9	p09	16	39.39	129.15	—	41.75	—	—
10	p10	17	66.68	—	—	85.83	—	—
Average Time (Sum Timeouts)			17.49 (0)	19.47 (3)	10.54 (9)	20.61 (0)	122.87 (9)	9.58 (15)
Average Penalized Time			17.49	77.52	187.38	20.61	266.01	304.79

Table 1. Blocks world experiments computing one and all solutions

with the huge¹³ number of light bulbs which was used in the problem instances and is unable to find a solution in any instance. Our system performs quite well in this benchmark, though not as well as dlv^k .

7 Discussion

Although we motivate (and apply) our approach in a biological setting, many features are readily applicable to representing and reasoning about dynamical systems in general. Centering our approach on \mathcal{C} has several benefits. First, \mathcal{C} is a rich and well-studied formalism. Second, it constitutes a mainstream implementation line for action languages. To this end, we provided a translation of the biologically motivated action language \mathcal{C}_{TAID} to \mathcal{C} and devise several tools for dealing with action descriptions in \mathcal{C} (and \mathcal{C}_{TAID}). Among them, we implemented the compiler `al2asp` allowing for translating action descriptions in \mathcal{C} (and \mathcal{C}_{TAID}) to logic programs under answer sets semantics. This approach is similar to the one taken by dlv^k for processing action language \mathcal{K} . Both approaches exploit the grounding and solving capacities of ASP, offering uniform (and thus instance independent) problem encodings and easy variable handling. Our approach is supported by a variety of pragmatic yet indispensable tools for addressing real world applications. Compared to other planning systems, we are able to compete with, and sometimes even outperform current systems. Finally, our tools (as well as their source code) and the benchmark problems are freely available at [3].

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¹³25.000 bulbs in instance *llc* up until 100.000 bulbs in instance *l4c*.