

# **Evolog - Actions and Modularization in Lazy-Grounding Answer Set Programming**

**DIPLOMARBEIT**

zur Erlangung des akademischen Grades

**Diplom-Ingenieur**

im Rahmen des Studiums

**Logic and Computation**

eingereicht von

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Wien, 9. Juni 2025

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DIPLOMA THESIS

submitted in partial fulfillment of the requirements for the degree of

**Diplom-Ingenieur**

in

**Logic and Computation**

by

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Michael Langowski, BSc.

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# Danksagung

Die vorliegende Abschlussarbeit ist das Ergebnis eines langen, oft zähen, und von vielen Ablenkungen - angefangen von den Anforderungen einer Vollzeit-Anstellung bis hin zu kleineren Ärgernisse wie einer Pandemie - geprägten Prozesses. Als ich die letzte Prüfung meines Studiums abschloss, war ich sicher, die Arbeit in wenigen Monaten fertigstellen zu können. Nun, mehr als ein Jahr später, ist es Zeit, allen Beteiligten für ihre Geduld und konstante Unterstützung zu danken. An erster Stelle steht hier Antonius Weinzierl, dessen Entwicklung von Alpha erst meine darauf aufbauende Arbeit ermöglichte, und der mir als Diskussionspartner und Advocatus Diavoli für die Formulierung und Abgrenzung meiner Vision stets zur Verfügung stand. Mindestens ebenso großer Dank gebührt meinem Betreuer Prof. Thomas Eiter, der in meinem langsamen Fortschritt stets den Fortschritt betrachtete und hier größere Geduld bewies als ich selbst. Seinem stets offenen Ohr und bereitwillig gegebenem Feedback ist es zu verdanken, dass aus wenig zusammenhängenden Textschnipseln ein kohärenter Text wachsen konnte.

Mein herzlicher Dank für ihren moralischen Support durch alle Entstehungsstufen dieser Arbeit und ihren Glauben an mich und den Erfolg meines Vorhabens geht an Tanja Travnicek. Darüber hinaus danke ich meinen Freunden und meiner Familie für ihre beständige Unterstützung während meines gesamten Studiums - es war eine lange Reise. *Ich habe fertig.*





# Acknowledgements

This thesis is the result of a long and oftentimes arduous process characterized by distractions ranging from small things like a global pandemic to the demands of full-time employment. When I finished the last exam from my curriculum, I was convinced I'd finish this thesis within months. Now, more than a year later, it's time to thank all those who contributed to this success. First here is Antonius Weinzierl without whose pioneering work on the Alpha system this work wouldn't have been possible. He was also readily available as a discussion partner and sounding board in the early stages of the project and helped me greatly in formulating my vision. At least equally significant is the support I got from my advisor, Prof. Thomas Eiter, who, in my slow progress always saw progress first, and demonstrated more patience than I often had myself. His readily given Feedback and support is to thank for some vaguely related text snippets finally growing into a coherent text.

My heartfelt thanks for her moral support and confidence in me and the eventual success of my project goes to Tanja Travnicek. In addition, I want to thank my friends and family for their continuing support through all stages of my studies - It has been a long road. *I'm done.*



# Kurzfassung

*Answer Set Programming (ASP)* ist ein seit Jahrzehnten etablierter Formalismus in der Logikprogrammierung, der insbesondere bei der Modellierung und Lösung NP-schwerer Planungsprobleme Anwendung findet. Effiziente Solver für ASP sind seit über 20 Jahren verfügbar. Aktuelle Trends in ASP befassen sich zunehmend mit der Interaktion mit externen Daten, sowohl zum Lesen und Verarbeiten (wie im Stream Reasoning bis hin zu Interaktion mit Machine-Learning-Software), als auch zum Ausführen von Aktionen, d.h. der Entwicklung von ASP-Programmen mit Seiteneffekten (side-effects). Das primäre Ziel der vorliegenden Arbeit ist es, eine neue Methode zur Modellierung von Regeln mit Seiteneffekten in ASP Programmen zu entwickeln, die sich leicht in einem ASP-Solver wie dem von Weinzierl et al. entwickelten *Lazy-Grounding-Solver Alpha* implementieren lässt. Ein weiteres Ziel ist die Entwicklung eines leichtgewichtigen Konzeptes zur Programmmodularisierung, das die zuvor erwähnte Unterstützung von Aktionen ergänzt. Die Notwendigkeit hierfür begründet sich aus der Tatsache, das Modellieren von Seiteneffekten in einer vollständig deklarativen Sprache zwangsläufig einige Einschränkungen struktureller Natur für Programme mit Aktionen mit sich bringt. Die mit dem Modularisierungskonzept gegebene Möglichkeit, seiteneffekt-freie Unterprogramme als eigenständige Einheiten auszuführen, sollte diese Einschränkungen im Praxiseinsatz ausgleichen. Die resultierende erweiterte Answer-Set-Programming-Sprache wird *Evolog* genannt. Die formale Semantik von Aktionen wird bezüglich eines *Frame*, d.h. eines Modells der Seiteneffekte einer Aktion in der Außenwelt in Form einer Interpretationsfunktion für *Action-Atome*, modelliert. Basierend auf diesem Modell der Außenwelt werden Kriterien definiert, nach denen ein Evolog-Programm (semantisch) gültig ist. Schließlich wird das Konzept eines *Evolog-Modells* definiert, d.h. eines Stable-Models eines Evolog-Programms, das konsistent bezüglich einem gegebenen Frame ist. Darüber hinaus wird gezeigt, dass Evolog eine *conservative* Erweiterung der üblichen Stable-Model-Semantik ist, insofern als jedes reguläre ASP-Programm auch ein gültiges Evolog-Programm in allen Frames ist. Die Modularisierung wird basierend auf Alpha's bestehendem Konzept eines *externen Atoms* modelliert, d.h. eines Atoms, das eine seiteneffektfreie Funktion abbildet, die in einer beliebigen Programmiersprache (in diesem Fall Java) implementiert ist. Modul-Atome sind definiert als externe Atome, deren Interpretationsfunktion darin besteht, einen ASP-Solver mit einem fixen Programm (der Modul-Implementierung) und den Eingabetermen des externen Atoms als zusätzlichem Input aufzurufen. Um das Schreiben von Code, der Module benutzt, effizienter zu gestalten, wird das (rein syntaktische) Konzept eines *Listen-Terms*

eingeführt. Um Evolog die Eignung von Evolog als tatsächliche Programmiersprache zu demonstrieren, wird eine einfache Kommandozeilenapplikation vorgestellt, die Graphen aus XML-Dateien einliest, 3-Colorings dieser Graphen berechnet und die berechneten Colorings in XML-Dateien serialisiert. Die Applikation wird sowohl in Bezug auf die Entwicklungsfreundlichkeit als auch auf die Ausführungsleistung diskutiert. Basierend auf der Demonstration anhand der XML-3-Coloring-App werden mögliche Erweiterungen der Sprache, sowie zukünftige Wege zur Steigerung der Solver-Performance angesprochen.

# Abstract

*Answer Set Programming (ASP)* has been an established formalism in logic programming, especially used for NP-hard planning problems, for decades, with efficient solvers available for more than 20 years. Current trends in ASP increasingly deal with interaction with external data, both for reading and processing (such as in stream reasoning or even interacting with machine learning software) as well as performing actions, i.e. influencing the state of the outside world. This thesis' primary concern is exploring a new way of representing actions in a manner suitable for easy implementation in a *lazy-grounding* ASP solver such as Alpha by Weinzierl et al. Secondary to that, the other goal of this work is to specify and implement a light-weight program modularization concept to go along with the aforementioned action support. The inspiration here is that, since inevitably modelling side-effects in a fully declarative language forces us to impose some restrictions on programs with actions, being able to run action-free subprograms as encapsulated units should alleviate any such strictures in software development. The resulting extended answer set programming language is called *Evolog*. We define the formal semantics of actions with regards to a *frame*, i.e. a formalization of the effects of an action in the outside world, that is modelled as an interpretation function for *action atoms*. Based on this model of the outside world, the notion of what constitutes a (semantically) valid Evolog program is defined. Finally, we define an *Evolog Model*, i.e. a stable model of an Evolog program that is consistent with a given frame. We further show that Evolog is a *conservative* extension to traditional ASP in the sense that every regular ASP program is also a valid Evolog program in all frames. Modularization is modelled around Alpha's existing concept of an *external atom*, i.e. an atom that models a side-effect free function which is implemented in an arbitrary programming language (Java in this case). We define module atoms as external atoms whose interpretation function is given as calling an ASP solver with a fixed program (the module body) and the input terms of the external atom as additional input. To aid in efficiently writing code around modules, the purely syntactical concept of a *list term* is introduced. In order to demonstrate Evolog's suitability as an actual application programming language, a simple command-line application for reading graphs from XML files, calculating 3-colorings of said graphs and serializing calculated colorings to XML files, is showcased and discussed in terms of both ease of development, as well as evaluation performance. Based on our demonstration of the XML-3-coloring application, we formulate possible refinements to the language as well as future paths for improvement in solver performance.

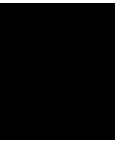


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# Introduction

This thesis deals with extending an existing solver for Answer Set Programming (ASP) programs, namely *Alpha* [WBB<sup>+</sup>19], with facilities to both execute *actions* from within ASP programs, as well as to re-use program parts by means of a modularization system. Answer Set Programming is a logic programming formalism that goes back to the seminal work of Gelfond and Lifschitz in 1988 [GL88]. Through its declarative semantics and concise syntax, it enables short and expressive programs to describe even NP-hard problems. Since its inception, ASP has been used with great success in diverse domains [EGL16] such as logistics [RGA<sup>+</sup>12] [AJO<sup>+</sup>21], automated music composition [EGPASB20] and spaceflight [NBG<sup>+</sup>01] as well as in various industry settings [FFS<sup>+</sup>18]. While ASP has enjoyed great success as a domain-specific language in the sense that it can express and efficiently solve complex reasoning problems, most applications require some kind of translation code written in a general-purpose programming language such as Python or Java to transform real-world input data into facts of an ASP program. Similarly, the answer sets an ASP solver outputs need to be translated back into the data exchange format of the problem domain, e.g. HTTP responses, XML files, etc. The aim of this work is to provide an extension to ASP that enables the development of complete applications entirely within the extended ASP, eliminating the need to use additional programming languages or write translation code. The vision is to, for example, write an application that receives XML encoded input, solves an NP-hard reasoning problem based on that input, and writes its results in an XML-based encoding, all in ASP. This chapter gives an intuitive introduction to ASP through examples, examines the state of the art on ASP extensions dealing with actions and side-effects, as well as program modularization, and finally formulates the specific goals for this thesis.

## 1.1 Answer Set Programming

At its core, an ASP program is a collection of conditional rules along the lines of "*if A holds true, then B must also hold*" as well as negative rules, so-called *constraints*, which prohibit certain conditions, e.g. "*If A has property p, then it cannot have property q*". Example 1.1.1 demonstrates the basic idea of ASP based on a program for course planning in a (very simplified) school setting.

**Example 1.1.1.** Listing 1.1 shows a knowledge base written in ASP, which encodes facts - things that are always true, such as "*There is a subject called maths*" - and rules, e.g. "*A teacher T that is qualified to teach subject S, can be assigned to teach S*", about a simplified school domain.

Listing 1.1: School planning in ASP

```
1      % The curriculum consists of different courses.
2      subject(german).
3      subject(english).
4      subject(maths).
5      subject(biology).
6      subject(history).
7
8      % Each teacher can teach one or more subjects
9      teacher(bob).
10     can_teach(bob, english).
11     can_teach(bob, maths).
12     teacher(alice).
13     can_teach(alice, maths).
14     can_teach(alice, history).
15     teacher(claire).
16     can_teach(claire, german).
17     can_teach(claire, history).
18     teacher(joe).
19     can_teach(joe, biology).
20     can_teach(joe, history).
21
22     % Assign subjects to teachers
23     { teaches(T, S) } :- teacher(T), can_teach(T, S).
24     % such that..
25     % Every teacher teaches at least one subject
26     :- teacher(T), not teaches(T, _).
27     % Every subject is taught by exactly one teacher
28     :- subject(S), not teaches(_, S).
29     :- teaches(T1, S), teaches(T2, S), T1 != T2.
```

Evaluating the above program using an *answer set solver*, i.e. an interpreter for the ASP language, yields the following collection of so-called *answer sets*:

- $A_1 = \{teaches(bob, english), teaches(bob, maths), teaches(alice, history), teaches(claire, german), teaches(joe, biology)\}$
- $A_2 = \{teaches(bob, english), teaches(alice, maths), teaches(claire, german), teaches(claire, history), teaches(joe, biology)\}$
- $A_3 = \{teaches(bob, english), teaches(alice, maths), teaches(claire, german), teaches(joe, biology), teaches(joe, history)\}$
- $A_4 = \{teaches(bob, english), teaches(alice, maths), teaches(alice, history), teaches(claire, german), teaches(joe, biology)\}$

Intuitively, each answer set constitutes a valid solution to the problem specified in the program in the sense that, if one adds the propositions from the answer set to the original program, all rules are fulfilled, while no constraint is violated. What sets ASP apart from many other logic programming formalisms such as Prolog [WDE<sup>+</sup>23] is its *multi-model semantics*, i.e. that it can express more than one solution to a problem, even when there are different solutions that are mutually exclusive.

Apart from the established use of ASP as a formalization language for decision or optimization problems, more recent developments in the field are increasingly targeted towards working with external data from different sources in ASP programs. A prominent example from this direction of research is the stream reasoning framework LARS [BDTE18]. Another recent development, NeurASP [YIL23] even generates facts out of perception results from a neural network, thereby enabling advanced artificial intelligence applications like reasoning about content in an image. Closely related to the concept of processing external data is the notion of actually influencing the outside world, for instance by writing data to a network buffer, from an ASP program, while still preserving declarative semantics. The DLV-extension Acthex [BEFI10] is an example of a system with basic action support. The goal of this thesis is to implement action support in the lazy-grounding solver Alpha [WBB<sup>+</sup>19], along with a basic modularization concept, thus enabling the development of arbitrary programs fully within ASP.

## 1.2 Actions and Modularization in ASP - Motivation

To get a clearer idea of the value of being able to write a complete application exclusively in ASP, consider the program from Example 1.2.1. It demonstrates a (very simple) parser component that may be easier to implement in a declarative language than some (typically imperative) general purpose language such as Java or Python. With most current ASP solvers, if one wanted to write such a declarative parsing component, reading of parser inputs (typically from some file or stream) and writing of parser output would have to

be done in some other language. Especially in applications where input and output are relatively simple but parsing and data transformation logic are more complex, being able to implement the entire application in a single language would greatly streamline development. Furthermore, to re-use code parts – for example some generic parser as exhibited in Example 1.2.1 – one would currently have to keep that code in a separate ASP file and manually avoid conflicts in naming of predicates without any language-level support for code encapsulation and modularization.

**Example 1.2.1.** Listing 1.2 shows a simple ASP program that parses binary strings and calculates the corresponding decimal numbers. It uses external atoms implemented in Java for basic string operations: The predicate *str\_x\_xs* takes inspiration from list syntax in Haskell and splits off the first character of a given string, e.g. *str\_x\_xs*["abc"]("a", "bc"), while *stdlib\_string\_length* and *stdlib\_string\_matches\_regex* test the length of a string and whether it matches some regular expression, respectively.

Listing 1.2: Parsing binary strings in ASP

```
1 encoding_scheme("1", "0", "[01]+").
2
3 % Helper - binstring_intm_decoded is the "internal"
4 % predicate we're using to recursively add up the
5 % bit values
6 binstring_intm_decoded(START_STR, START_STR, 0) :-
7     binstr(START_STR).
8
9 % Handle the case where the current bit is set
10 binstring_intm_decoded(START_STR, CURR_STR, CURR_VALUE) :-
11     binstring_intm_decoded(START_STR, LAST_STR,
12         LAST_VALUE),
13     &str_x_xs[LAST_STR](HIGH_CODE, CURR_STR),
14     &stdlib_string_length[LAST_STR](LAST_LEN),
15     CURR_VALUE = LAST_VALUE + 2 ** (LAST_LEN - 1),
16     encoding_scheme(HIGH_CODE, _, REGEX),
17     &stdlib_string_matches_regex[START_STR, REGEX].
18
19 % Handle the case where the current bit is not set
20 binstring_intm_decoded(START_STR, CURR_STR, CURR_VALUE) :-
21     binstring_intm_decoded(START_STR, LAST_STR,
22         LAST_VALUE),
23     &str_x_xs[LAST_STR](LOW_CODE, CURR_STR),
24     &stdlib_string_length[LAST_STR](LAST_LEN),
25     CURR_VALUE = LAST_VALUE,
26     encoding_scheme(_, LOW_CODE, REGEX),
27     &stdlib_string_matches_regex[START_STR, REGEX].
```

```

27      % These are the final values
28      bin_number(BIN, DEC) :-
29          binstring_intm_decoded(BINSTR, "", DEC).

```

In order to overcome the aforementioned hurdles and be able to write applications with input from and output to external data sources as one piece in a single ASP-based language, we propose an extension of "classical" ASP in Section 1.3. Section 1.4 gives an overview of the state of the art on action support and approaches to program modularizations in current ASP solving systems. Concluding this introduction, Section 1.5 gives an outline of the rest of this thesis.

## 1.3 Problem Statement

The main goal of this thesis is for ASP programs to be able to express interactions with external data sources which potentially have an effect on the state of the outside world. Closely related to that goal is the demand for mechanisms enabling code reuse and structuring of applications - the larger an application gets, the more the necessity for some form of modularization grows. To that end, this thesis aims to deliver the contributions listed in the following paragraphs.

**Triggering actions from programs** Most program flows follow a chain of events, each a consequence of its predecessor, e.g. "If there exists a file A, read it. If reading was successful, do something with the content. If the operation succeeds, write the result to file B". It is highly desirable to be able to write this kind of program in a declarative, logic-based language that can leverage the strengths of ASP for the "business logic" part. Specifically, the proposed action semantics should deliver

- declarative programs, i.e. order in which actions occur in code does not affect semantics,
- actions behaving in a functional fashion, i.e. an action always gives the same result for the same input. Especially, actions have to be idempotent in the sense that, for an ASP rule that is associated with some action, the result of the action never changes, no matter how often the rule fires.
- transparent action execution, i.e. every action that is executed during evaluation of a program must be reflected in an answer set.

**Program Modularization** While not formally connected, triggering actions from programs and modularization (i.e. plugable and re-usable sub-programs), intuitively complement each other in the sense that modules enable us to define clear boundaries between "data acquisition" (i.e. I/O-centered) and "data processing" parts of a program. Introducing a simple, easy-to-use module system is therefore the second goal of this work.

**Incremental Evaluation and Lazy Grounding** Experiences from existing systems for ASP application development such as ASAP [SW15] or ACTHEX [BEFI10] show that, in order to achieve the evaluation performance necessary for use in real-world applications, ASP application code needs to be evaluated in an incremental fashion (rather than iteratively re-evaluating the whole program) whenever possible. The lazy-grounding architecture employed by ASP systems such as Alpha [WBB<sup>+</sup>19] offers an intuitive solution.

**Application-programming Proof of Concept** Based on the extended version of the Alpha solver incorporating Action and Modularization support, the final goal for this thesis is to test the extended solver by developing an application making use of the newly introduced concepts.

## 1.4 State of the Art

This work aims to blend action support with modularization in the context of lazy-grounding ASP solving - all three of these areas have seen a substantial amount of research in the past.

The question of how to formally represent actions and consequences of actions in logic programs has long been investigated, giving rise to so-called *action languages*, which relate outcomes of actions to their causes and preconditions. Most interesting from an Answer Set Programming point of view is the work of Gelfond and Lifschitz in [GL93], where a translation from an action language into stable model programs is introduced. In terms of actually performing (rather than just modelling) actions from ASP programs, both Clingo [GKKS14] and DLVHEX, through the ACTHEX [BEFI10] extension, offer their own flavours of support for triggering actions from programs. While Clingo does not directly support actions as a dedicated feature - and therefore offers no strictly enforced semantics for this - similar behavior can be achieved using external functions and the reactive solving features first introduced in oClingo [GGKS11]. ACTHEX has thoroughly defined semantics for actions. In the ACTHEX model, answer set search and action execution are separate steps, where executability of actions is only determined after answer sets are calculated. While this gives users a high degree of flexibility in working with actions, it does not directly lend itself to the idea of a general purpose language where program behavior may be influenced by continuous two-way communication between a program and its environment.

With regards to Modularity, i.e. the process of "assembling" an ASP program from smaller building blocks (i.e. modules), a comprehensive semantics for so-called *nonmonotonic modular logic programs* has been introduced in [Kre18] and [DTEFK09]. While it does not impose any restrictions on language constructs used in modules and recursion within and between modules, it also comes with rather high computational complexity and no easily available implementations so far. A more "lightweight" approach to modularization are *Templates* [IIP<sup>+</sup>04]. As the name implies, this purely syntactic approach aims to

define isolated sub-programs that can be generically used to avoid code duplication throughout an application and is conceptually similar to the well-known generics in object-oriented languages such as C++ and Java. Templates are rewritten into regular ASP rules using an "explosion" algorithm which basically "instantiates" the template by generating the needed body atoms (and rules deriving them) wherever templates are used. While easy to implement and flexible, a potential disadvantage of this concept is that - due to its purely syntactical nature - programmers need to be on the watch for potential bugs arising from unintended cyclic dependencies or recursive use of templates (leading to potential non-termination of the explosion algorithm) themselves. Yet another powerful toolset for modular application development is provided by Clingo's *multi-shot-solving* [GKKS19] features. Clingo allows for parameterized sub-programs which are then repeatedly grounded in a process that is conceptually similar to the notion of module instantiation in [OJ08] and solved during solving of the overall program. However, as this "contextual grounding" needs to be programmatically controlled by an external application through Clingo's API, the inherent flexibility and usefulness for incremental solving of this approach is counterweighed by a high level of proficiency with and knowledge of the Clingo system necessary to leverage these capabilities. The concept of *lazy grounding*, i.e. interleaving of the - traditionally sequential - grounding and solving steps present in most prevalent ASP solvers, is relatively new. It has been spearheaded by the GASP [DPDPR09] and ASPERIX [LN09] solvers which avoid calculating the full grounding of an input program by performing semi-naive bottom-up evaluation along the input's topologically sorted (non-ground) dependency graph. While efficient in terms of memory use, this approach cannot stand up to the solving performance of systems like DLV or Clingo which employ their knowledge of all possible ground rules to perform conflict-driven nogood learning (CDNL) as part of their solving algorithm to great effect. Alpha [WBB<sup>+</sup>19], a more recent lazy-grounding solver, aims to bridge this gap in performance by employing CDNL-style solving techniques [Wei17] incrementally on partially ground program parts as part of its central ground-and-solve loop.

Conceptually, the common ingredient linking the - on first glance not directly connected - areas of actions in ASP, program modularization, and lazy grounding is a need for detailed static program analysis prior to solving, be it to detect potentially invalid action sequences, calculate module instantiation orders, or for up-front evaluation of stratified program parts in a lazy-grounding context. In addition, both actions and modularization can greatly benefit from - or even depend on - incremental evaluation facilities of a solver for efficient operation. Since lazy-grounding by its very definition embodies an incremental evaluation approach, it seems only natural to incorporate actions and modularization into a lazy-grounding solver's input language in order to provide ASP programmers with a powerful tool for application development. Alpha, with its good solving performance compared to other lazy-grounding systems, support for a large part of the current ASP-Core2 language standard, and active development status, appears the natural choice as the technical backbone of this work.

### 1.5 Thesis Roadmap

The core part of this work is the formal specification of the Evolog language in Chapter 3, where we formally define Evolog’s action semantics and modularization concept and make some observations on the relationship between Evolog programs versus regular ASP programs. Chapter 4 gives an overview of how the formal specifications from Chapter 3 were implemented in the Alpha ASP solver, along with some examples of actual programs written in Evolog. In Chapter 5, we showcase an example application that was developed using Alpha with its Evolog extension. The application reads graphs encoded as XML data from user-supplied files, calculates three-colorings of given graphs, and writes the obtained colorings to XML files. Finally, Chapter 6 reflects on the experiences gained for hands-on software development from developing the aforementioned showcase application. We try to gauge the practical applicability of the implementation and highlight challenges yet to be addressed. Chapter 7 wraps up the thesis, reflecting on the goals achieved and potential paths for future development.



# Preliminaries

## 2.1 Answer Set Programming

When speaking of ASP, we nowadays mostly refer to the language specified by the ASP-Core2 standard [CFG<sup>+</sup>20]. It uses the *stable model semantics* by Gelfond and Lifschitz [GL88] as a formal basis and enhances it with support for advanced concepts such as disjunctive programs, aggregate literals and weak constraints. This chapter describes the input language supported by the Alpha solver, which will serve as the basis on which we will define the Evolog language.

### 2.1.1 Syntax

**Definition 2.1.1** (Integer numeral). An *integer numeral* in the context of an ASP program is a string matching the regular expression:

$$(-) ? [0-9]^+$$

The set of all valid integer numerals is denoted as *INT*.

**Definition 2.1.2** (Identifier). An *identifier* in the context of an ASP program is a string matching the regular expression:

$$[a-z][a-zA-Z0-9\_]*$$

The set of all valid identifiers is denoted as *ID*.

**Definition 2.1.3** (Variable Name). A *variable name* in the context of an ASP program is a string matching the regular expression:

$$[A-Z][a-zA-Z0-9\_]*$$

The set of all valid variable names is denoted as *VAR*.

**Definition 2.1.4** (Term). A *term* is inductively defined as follows:

- Any *constant*  $c \in (INT \cup ID)$  is a term.
- Any *variable*  $v \in VAR$  is a term.
- Given terms  $t_1, t_2$ , any *arithmetical expression*  $t_1 \oplus t_2$  with  $\oplus \in \{+, -, *, /, **\}$  is a term.
- Given terms  $t_1, t_2$ , any *interval expression*  $t_1 \dots t_2$  is a term.
- For function symbol  $f \in ID$  and argument terms  $t_1, \dots, t_n$ , the *functional expression*  $f(t_1, \dots, t_n)$  is a term.

**Definition 2.1.5** (Subterms). Given a term  $t$ , the set of *subterms* of  $t$ ,  $st(t)$ , is defined as follows:

- If  $t$  is a *constant* or *variable*,  $st(t) = \{t\}$ .
- If  $t$  is an *arithmetical expression*  $t_1 \oplus t_2$ ,  $st(t) = st(t_1) \cup st(t_2)$ .
- If  $t$  is an *interval expression*  $t_1 \dots t_2$ ,  $st(t) = st(t_1) \cup st(t_2)$ .
- If  $t$  is a *functional expression* with argument terms  $t_1, \dots, t_n$ ,  $st(t) = st(t_1) \cup \dots \cup st(t_n)$ .

A term is called *ground* if it is variable-free, i.e. none of its subterms is a variable.

**Definition 2.1.6** (Basic Atom). Given a predicate symbol  $p \in ID$  and argument terms  $t_1, \dots, t_n$ , the expression

$$p(t_1, \dots, t_n)$$

is called a *atom*. An atom is ground if all of its argument terms are ground. A ground atom with predicate  $p$  is called an *instance* of  $p$ .

**Definition 2.1.7** (Comparison Atom). Given terms  $t_1$  and  $t_2$  and comparison operator  $\odot$  where  $\odot \in \{<, \leq, =, \geq, >, \neq\}$ , the expression

$$t_1 \odot t_2$$

is called a *comparison atom*. Syntactically, a comparison atom is a regular atom where the predicate symbol (i.e. comparison operator) is written in infix- rather than prefix-notation.

**Definition 2.1.8** (External Atom). Given an *external predicate name*  $ext$ , *input terms*  $t_1, \dots, t_n$  and *output terms*  $t_{n+1}, \dots, t_m$ , the expression

$$\&ext[t_1, \dots, t_n](t_{n+1}, \dots, t_m)$$

is called an *external atom*. Syntactically, external atoms are regular atoms where  $\&ext$  is the predicate symbol and  $t_1, \dots, t_m$  are argument terms.

**Definition 2.1.9** (Literal). A literal in ASP is an atom  $a$  or ("default"-)negated atom  $\text{not } a$ . Literals wrapping comparison- or external atoms are called *fixed interpretation literals*. Given a literal  $l$ , the expression  $\text{pred}(l)$  refers to the predicate of  $l$  (e.g.  $\text{pred}(p(a)) = p/1$ ).

**Definition 2.1.10** (Rule, Program). A *rule* is an expression of form

$$a_H \leftarrow b_1, \dots, b_n.$$

for  $n \geq 0$ , where the *rule head*  $a_H$  is an atom and the *rule body*  $b_1, \dots, b_n$  is a set of literals. An ASP *program* is a set of rules. A rule with an empty body is called a *fact*. A rule is *ground* if both its head atom and all of its body literals are ground. By the same reasoning, a program is ground if all of its rules are ground.

Given a rule  $r$ , we refer to the head of  $r$  as  $h(r)$  and the body of  $r$  as  $b(r)$ . Furthermore,  $b^+(r)$  is used to reference the set of *positive body literals* of  $r$ , while  $b^-(r)$  references the *negative body literals*.

**Definition 2.1.11** (Constraint). A *constraint* is a special form of rule, written as a rule with an empty head, i.e.

$$\leftarrow b_1, \dots, b_n.$$

It is syntactic sugar for

$$q \leftarrow b_1, \dots, b_n, \text{not } q.$$

where  $q$  is a propositional constant not occurring in any other rule in the program.

### 2.1.2 Semantics

**Definition 2.1.12** (Herbrand Universe). The Herbrand Universe  $HU_P$  of a Program  $P$  is the set of all ground terms that can be constructed with respect to Definitions 2.1.1, 2.1.2 and 2.1.4. Note that most papers use stricter definitions of the Herbrand Universe where  $HU_P$  consists only of terms constructible from constants occurring in  $P$ . The broader definition used here is chosen for ease of definition with respect to some of the extensions introduced in Section 3.1.

**Definition 2.1.13** (Herbrand Base). The Herbrand Base  $HB_P$  of a Program  $P$  is the set of all ground atoms that can be constructed from the Herbrand Universe  $HU_P$  according to definition 2.1.6.

**Definition 2.1.14** (Herbrand Interpretation). A Herbrand Interpretation is a special form of first order interpretation where the domain of the interpretation is a Herbrand Universe and the interpretation of a term is the term itself, i.e. the corresponding element of  $HU_P$ . Intuitively, Herbrand Interpretations constitute listings of atoms that are true in a given program. Since the domain of a Herbrand Interpretation is always the Herbrand Universe  $HU_P$ , we only need to give a predicate interpretation for the predicates occurring in a program  $P$  in order to fully specify a Herbrand Interpretation. We can therefore denote Herbrand Interpretations as sets of atoms  $I \subseteq HB_P$ .

### Grounding

Given a program  $P$  containing variables, *grounding* refers to the process of converting  $P$  into a semantically equivalent propositional, i.e. variable-free, program.

**Definition 2.1.15** (Substitution, adapted from [Wei17]). A substitution  $\sigma : VAR \mapsto (ID \cup INT)$  is a mapping from variables to constants. For a atom  $a$ , applying a substitution results in a substituted atom  $a\sigma$  in which variables are replaced according to  $\sigma$ . Substitutions are applied to rules by applying them to every individual atom or literal within the rule. By the same mechanism, we can apply substitutions to programs by applying the to all rules.

**Definition 2.1.16** (Grounding). Given a rule  $r$ , the *grounding* of  $r$ ,  $grnd(r)$ , is a set of substitutions  $S$ , such that the set of ground rules resulting from applying the substitutions in  $S$  is semantically equivalent to  $r$ . In a slight abuse of terminology, *grounding* in this work also refers to the set of ground rules resulting from applying  $S$  as well as the process of finding said set.

### Stable Model Semantics

**Definition 2.1.17** (Fixed interpretation literals). Fixed interpretation literals, i.e. comparison- and external literals, respectively, are interpreted by means of a program-independent oracle function  $f_O : H_U(P)^* \mapsto \{\top, \perp\}$ , i.e. a fixed interpretation literal with argument terms  $t_1, \dots, t_n$  has the same truth value under all interpretations.

**Definition 2.1.18** (Truth of Atoms and Literals). A positive ground literal  $l$  with atom  $a$  is true w.r.t. a Herbrand Interpretation  $I$ , i.e.  $I \models l$  if

- $a$  is a basic atom contained in  $I$ , i.e.  $a \in I$ ,
- $a$  is a fixed interpretation literal with terms  $t_1, \dots, t_n$  and  $f_O(t_1, \dots, t_n) = \top$ .

For a negative ground literal  $not\ a$ , the reverse holds, i.e.  $I \models not\ a$  if

- $a$  is a basic atom not contained in  $I$ , i.e.  $a \notin I$ ,
- $a$  is a fixed interpretation literal with terms  $t_1, \dots, t_n$  and  $f_O(t_1, \dots, t_n) = \perp$ .

A set of literals  $L$  is true w.r.t. an interpretation  $I$  if  $I \models l$  holds for every literal  $l \in L$ .

**Definition 2.1.19** (Positive Logic Program). A *positive* logic program is a program according to Definition 2.1.10, where all rule bodies are positive, i.e. no rule body contains a negated atom.

**Definition 2.1.20** (Immediate Consequence Operator, adapted from [EIK09]). Given a Herbrand Interpretation  $I$  and a ground positive logic program  $P$ , the immediate consequence operator  $T_P(I)$  defines a monotonic function  $T_P : 2^{HB_P} \mapsto 2^{HB_P}$  such that

$$T_P(I) = \{h(r) \mid r \in P \wedge I \models b(r)\}$$

i.e. the result set of applying  $T_P$  with a Herbrand Interpretation  $I$  contains the heads of all rules whose body is true under  $I$ .

**Definition 2.1.21** (Least Model of positive logic programs). The least model  $LM(P)$  of a (ground) positive logic program  $P$  is the least fixpoint of the  $T_P$  operator of  $P$ , i.e. the set toward which the sequence  $\langle T_P^i \rangle$ , with  $i \geq 0$ ,  $T_P^0 = \emptyset$  and  $T_P^i = T_P(T_P^{i-1})$  for  $i \geq 1$ , converges. The existence of said fixpoint and its characterisation as limit of  $\langle T_P^i \rangle$  follow from the fixpoint theorems of Knaster, Tarski and Kleene, respectively.

**Definition 2.1.22** (Gelfond-Lifschitz Reduct, adapted from [GL88] and [EIK09]). Given a ground ASP program  $P$  and a Herbrand Interpretation  $I$ , the *Gelfond-Lifschitz-Reduct* ("GL-reduct")  $P^I$  of  $P$  with respect to  $I$  is the program obtained by:

- removing from  $P$  all rules  $r$  that are "blocked", i.e.  $I \not\models l$  for some literal  $l \in b^-(r)$
- and removing the negative body of all other rules.

Note that  $P^I$  is a positive logic program.

**Definition 2.1.23** (Answer Set [GL88] [EIK09]). A Herbrand Interpretation  $I$  of an ASP program  $P$  is an *answer set* or *stable model* of  $P$  iff it is the least model  $LM(P^I)$  of the GL-reduct  $P^I$  of  $P$ . We denote the set of Answer Sets of a program  $P$  as  $AS(P)$ .

## 2.2 Lazy-grounding Answer-Set Solving

The theoretical notion underpinning lazy-grounding answer-set solving is that of a *Computation Sequence*, which is formalized in Definition 2.2.1. Intuitively, a computation sequence is a sequence of (ground) rules firing, such that, at the end, the atoms derived by the rules from the computation sequence (together with all facts of the program) constitute an answer set.

**Definition 2.2.1** (Computation Sequence, adapted from [Wei17] and [LN09]). Let  $P$  be an ASP program and  $S = (A_0, \dots, A_\infty)$  a sequence of assignments, i.e. herbrand interpretations denoted by a set of atoms assumed to be true, then  $S$  is called a *computation sequence* iff

- $A_0 = \emptyset$
- $\forall i \geq 1 : A_i \subseteq T_P(A_{i-1})$ , i.e. every  $A_i$  is a consequence of its predecessor in the sequence,

- $\forall i \geq 1 : A_{i-1} \subseteq A_i$ , i.e.  $S$  is monotonic,
- $A_\infty = \cup_{i=0}^\infty A_i = T_P(A_\infty)$ , i.e.  $S$  converges toward a fixpoint and
- $\forall i \geq 1 : \forall a \in A_i \setminus A_{i-1}, \exists r \in P : h(r) = a \wedge \forall j \geq i - 1 : A_j \models a$ , i.e. applicability of rules is persistent.

$A_\infty$  is an answer set of  $P$  iff  $S$  is a computation sequence. Note that there may exist an arbitrary number of computation sequences leading to the same answer set.

It is easily observed that, in order to calculate computation sequences, one does not need a full grounding of the input program. Since facts are ground by definition, rules that only depend on facts can be grounded (and evaluated) based on facts alone. The same holds for rules depending on facts and rules from the previous step, etc. Example 2.2.1 demonstrates how a computation sequence for a very simple positive program can be calculated using a naive algorithm based on repeated application of the  $T_P$ -operator.

**Example 2.2.1** (Lazy-grounding a positive logic program). In order to illustrate the use of computation sequences for lazy-grounding, we consider the program  $P$  in Listing 2.1. We denote the rule on line 2 as  $r_1$  and the one on line 3 as  $r_2$ .

Listing 2.1: A positive, non-ground program

1	$p(a) . p(b) . p(c) . q(b) . q(c) . q(d) .$
2	$r(X) :- p(X), q(X) .$
3	$t(X, Y) :- r(X), r(Y), X \neq Y .$

Starting from the set of facts on line 1, which we denote as  $F$ , we now look for rules that only depend on predicates in  $F$  (i.e. body literals only have predicates for which we already know ground instances). The only applicable rule is  $r_1$  for which we can construct ground instances  $r(b) :- p(b), q(b)$  and  $r(c) :- p(c), q(c)$ . Continuing this process, now based on the set  $F \cup \{r(b), r(c)\}$ , we find ground instances  $t(b, c) :- r(b), r(c), b \neq c$  and  $t(c, b) :- r(c), r(b), c \neq b$  of  $r_2$ . We thus arrive at the following computation sequence:

$$\begin{aligned}
A_0 &= \{p(a), p(b), p(c), q(b), q(c), q(d)\}, \\
A_1 &= \{p(a), p(b), p(c), q(b), q(c), q(d), r(b), r(c)\}, \\
A_2 &= \{p(a), p(b), p(c), q(b), q(c), q(d), r(b), r(c), t(b, c), t(c, b)\}
\end{aligned}$$

The last element of the sequence,  $A_2$  is the sole answer set of  $P$ .

While the intuitive approach from Example 2.2.1 clearly works for positive programs, things get more complicated when negation is involved. Consider the following program:

1	$p(a) . p(b) . p(c) . q(c) .$
2	$s(X) \text{ :- } p(X) , q(X) .$
3	$t(X) \text{ :- } p(X) , \text{ not } s(X) .$

In order to arrive at the correct answer set  $A = \{p(a), p(b), p(c), q(c), s(c), t(a), t(b)\}$ , one has to first evaluate all rules that could potentially derive instances of  $s/1$ , before starting to evaluate the last rule, in order to end up with a valid computation sequence. Evaluation orders for this kind of programs can be calculated using the notion of *stratification* [ABW88] which is described in detail in Section 2.2.1.

### 2.2.1 Structural Dependency Analysis and Stratified Evaluation

In a nutshell, a *stratifiable* logic program is a program, for which we can calculate a partition into sub-programs, such that, when evaluating the sub-programs sequentially, one always ends up with a correct computation sequence. Definition 2.2.2 formally characterizes stratifiable programs.

**Definition 2.2.2** (Stratifiable answer set program, as stated in [Lan19], adapted from [ABW88], [EIK09]). Given a program  $P = \{r_0, \dots, r_n\}$ ,  $P$  is called *stratifiable* iff there is a partition  $P = P_{S_0} \cup \dots \cup P_{S_n}$  so that for each  $0 \leq i \leq n$  the following holds:

- For every positive body literal  $l_b$  of every rule in  $P_{S_i}$ , every rule that derives instances of the predicate  $pred(l_b)$ , i.e. where the predicate of the head atom  $pred(a_H)$  is  $pred(l_b)$ , is contained in some  $P_{S_j}$  with  $j \leq i$ .
- For every negative body literal  $not\ l_b$  of every rule in  $P_{S_i}$ , every rule that derives instances of the predicate  $pred(l_b)$ , i.e. where the predicate of the head atom  $pred(a_H)$  is  $pred(l_b)$ , is contained in some  $P_j$  with  $j < i$ .

The individual subprograms  $P_{S_i}$  are called strata. A *stratification*  $S = \{P_{S_0}, \dots, P_{S_n}\}$  refers to the set of all strata making up a partition which satisfies the criteria above.

The least model, i.e. the sole answer set, of a stratifiable program can be computed by sequentially applying the immediate consequence operator to every stratum, see Definition 2.2.3.

**Definition 2.2.3** (Least model of stratifiable programs, adapted from [EIK09]). Let  $P_{strat}$  be a stratifiable program for which a stratification  $S = \{P_{S_0}, \dots, P_{S_n}\}$  exists and let  $T_{P_{S_i}}$  be the immediate consequence operator for the sub-program defined by stratum  $P_{S_i}$ . Furthermore, for a model  $M$ , let  $pr(M)$  be the program consisting of the facts representing the atoms in  $M$ . Then the least model  $LM(P_{strat})$  of  $P_{strat}$  is defined as follows. The sequence  $\langle M_{S_i} \rangle$ ,  $0 \leq i \leq n$  with  $M_{S_0} = lfp(T_{P_{S_0}})$  and  $M_{S_i} = lfp(T_{(P_{S_i} \cup pr(M_{S_{i-1}}))})$  for all  $1 \leq i \leq n$  defines the (iterative) least model for each stratum. The least model

$LM(P_{strat})$  of program  $P_{strat}$  is then the iterative least model of the highest stratum  $M_{S_n}$ , i.e. the end of the sequence  $\langle M_{S_i} \rangle$ .

Since the evaluation mode described in Definition 2.2.3 does not require any backtracking, i.e. the result of each step is part of the final result, we observe that the intermediate results of the immediate consequence operator yield a computation sequence for  $P$ . Example 2.2.2 illustrates evaluation of a short stratifiable program.

**Example 2.2.2** (Evaluating a stratified program). Consider the program  $P$  from Listing 2.2.

Listing 2.2: A stratifiable program.

```

1 p(a) . p(b) . p(c) . q(c) . q(d) .
2
3 s(X) :- p(X), q(X) .
4 t(X) :- p(X), not s(X) .
5 u(X) :- q(X), not p(X) .
6
7 v(X, Y) :- t(X), u(Y), X != Y.
```

A possible stratification  $S$  of  $P$  could look as follows:

$$\begin{aligned}
S_0 &= \{p(a). p(b). p(c). q(c). q(d). \\
&\quad s(X) \leftarrow p(X), q(X).\} \\
S_1 &= \{t(X) \leftarrow p(X), \text{not } s(X). \\
&\quad u(X) \leftarrow q(X), \text{not } p(X).\} \\
S_2 &= \{v(X, Y) \leftarrow t(X), u(Y), X \neq Y.\}
\end{aligned}$$

In this case, the least model of  $P$  can be calculated using the method from Definition 2.2.3.



The individual results of the  $T_P$  operator constitute a computation sequence.

$$\begin{aligned}
T_{P_{S_0}}^1 &= T_{P_{S_0}}(\emptyset) = \{p(a), p(b), p(c), q(c), q(d)\} \\
T_{P_{S_0}}^2 &= T_{P_{S_0}}(T_{P_{S_0}}^1) = \{p(a), p(b), p(c), q(c), q(d), s(c)\} \\
T_{P_{S_0}}^3 &= T_{P_{S_0}}(T_{P_{S_0}}^2) = \{p(a), p(b), p(c), q(c), q(d), s(c)\} = \text{lpf}(T_{P_{S_0}}) \\
\\
T_{P_{E_1}} &= T_{P_{S_1 \cup \text{pr}(\text{lpf}(T_{P_{S_0}}))}} \\
T_{P_{E_1}}^1 &= T_{P_{E_1}}(\emptyset) = \{p(a), p(b), p(c), q(c), q(d), s(c)\} \\
T_{P_{E_1}}^2 &= T_{P_{E_1}}(T_{P_{E_1}}^1) = \{p(a), p(b), p(c), q(c), q(d), s(c), t(a), t(b), u(d)\} \\
T_{P_{E_1}}^3 &= T_{P_{E_1}}(T_{P_{E_1}}^2) = \{p(a), p(b), p(c), q(c), q(d), s(c), t(a), t(b), u(d)\} = \text{lpf}(T_{P_{E_1}}) \\
\\
T_{P_{E_2}} &= T_{P_{S_2 \cup \text{pr}(\text{lpf}(T_{P_{E_1}}))}} \\
T_{P_{E_2}}^1 &= T_{P_{E_2}}(\emptyset) = \{p(a), p(b), p(c), q(c), q(d), s(c), t(a), t(b), u(d)\} \\
T_{P_{E_2}}^2 &= T_{P_{E_2}}(T_{P_{E_2}}^1) = \{p(a), p(b), p(c), q(c), q(d), s(c), t(a), t(b), u(d), v(a, d), v(b, d)\} \\
T_{P_{E_2}}^3 &= T_{P_{E_2}}(T_{P_{E_2}}^2) = \{p(a), p(b), p(c), q(c), q(d), s(c), t(a), t(b), u(d), v(a, d), v(b, d)\} = \text{lpf}(T_{P_{E_2}}) \\
\\
LM(P) &= \text{lpf}(T_{P_{E_2}})
\end{aligned}$$

However, in general, ASP programs are not stratifiable. Example 2.2.3 shows a typical non-stratifiable program.

**Example 2.2.3** (A non-stratifiable program). Consider a variant of the well-known 3-coloring problem for undirected graphs, where some vertices "stay blank", i.e. they are excluded from coloring. Essentially, prior to calculating actual colorings, we remove all ignored vertices and their corresponding edges from the graph and color only the resulting subgraph. Program  $P_{col}$  in Listing 2.3 shows a possible encoding of this problem.

Listing 2.3: Graph 3-coloring with excluded vertices

```

1 vertex(a) . vertex(b) . vertex(c) . vertex(d) .
2 edge(a, b) . edge(a, c) . edge(a, d) .
3 edge(b, c) . edge(b, d) . edge(c, d) .
4 edge(X, Y) :- edge(Y, X) .
5
6 exclude_vertex(d) .
7 exclude_edge(V1, V2) :- edge(V1, V2), exclude_vertex(V1) .
8 exclude_edge(V1, V2) :- exclude_edge(V2, V1) .
9
10 coloring_vertex(V) :- vertex(V), not exclude_vertex(V) .
11 coloring_edge(V1, V2) :-

```

```

12         edge(V1, V2), not exclude_edge(V1, V2).
13
14 % Guess colors
15 red(V) :- coloring_vertex(V), not green(V), not blue(V).
16 green(V) :- coloring_vertex(V), not red(V), not blue(V).
17 blue(V) :- coloring_vertex(V), not red(V), not green(V).
18
19 % Filter invalid guesses
20 :- coloring_vertex(V1), coloring_vertex(V2),
21     coloring_edge(V1, V2), red(V1), red(V2).
22 :- coloring_vertex(V1), coloring_vertex(V2),
23     coloring_edge(V1, V2), green(V1), green(V2).
24 :- coloring_vertex(V1), coloring_vertex(V2),
25     coloring_edge(V1, V2), blue(V1), blue(V2).

```

Clearly,  $P_{col}$  is not stratifiable - the color assignment rules cyclically depend on each other through negated body literals. In other words, the rules on lines 15 to 17 are *mutually exclusive*, i.e. a solver has to *choose* which of the three should fire for a given ground instance of *coloring\_vertex*/1.

The program from Example 2.2.3 can not be evaluated by iterative fixpoint-calculation as in Example 2.2.2 because it is not stratifiable. However, it is still possible to evaluate at least a part of it using this method, rather than a computationally more complex "trial-and-error"-approach. This partial evaluation approach uses *splitting sets*, which are characterized in Definition 2.2.4. Intuitively, a splitting set  $S$  is a set of atoms, such that if an atom  $a$  is in  $S$ , also all atoms it depends on (i.e. body atoms of rules deriving  $a$ ) are in  $S$ . By the splitting set theorem, a splitting sets lets us partition a program  $P$  such that it can be evaluated incrementally, by first evaluating the bottom  $B_U(P)$  and then the top  $T_U(P)$ . Since this section deals with non-ground programs specifically, we also give an adapted definition for splitting sets of nonground programs, see Definition 2.2.5. Example 2.2.4 applies the splitting set theorem to the program from Example 2.2.3.

**Definition 2.2.4** (Splitting Set, adapted from [LT94]). Given a program  $P$ , a set of atoms  $U$  is a *splitting set* of  $P$  if for every rule  $r$ , where  $h(r) \in U$ , also the atoms corresponding to all body literals of  $r$  are in  $U$ . The set of rules corresponding to  $U$ , i.e. the rules defining the atoms in  $U$ , is called *bottom* of  $P$  with respect to  $U$ , denoted as  $B_U(P)$ . Consequently,  $P \setminus B_U(P)$  is called *top* of  $P$ , which is denoted as  $T_U(P)$ .

**Theorem 2.2.1** (Splitting Set Theorem, adapted from [LT94]). Given a program  $P$  and splitting set  $U$  which splits  $P$  into bottom  $B_U(P)$  and top  $T_U(P) = P \setminus B_U(P)$ . Then a set of atoms  $A$  is an answer set if and only if  $A = X \cup Y$  for sets  $X$  and  $Y$  where

- $X \in AS(B_U(P))$ , i.e.  $X$  is an answer set of bottom  $B_U(P)$  and

- $Y \in AS(T_U(P) \cup X)$ , i.e.  $Y$  is an answer set of the program resulting from adding  $X$  as facts to  $T_U(P)$

**Definition 2.2.5** (Nonground splitting set). Given a non-ground program  $P$ , we denote splitting sets as sets of predicates rather than atoms. For a given predicate  $p$ , we denote as  $dep(p)$  the *dependencies* of  $p$ :

$$dep(p) = \{q \mid \exists r \in P : pred(h(r)) = p \wedge \exists l \in b(r) : pred(l) = q\}$$

Intuitively, the dependencies of a predicate  $p$  are all predicates of which all ground instances must be known in order to compute all ground instances of  $p$ . A set  $S$  of predicates  $p_1, \dots, p_n$  is a splitting set of  $P$  if the following holds.

$$\forall p : p \in S \Rightarrow dep(p) \in S$$

A non-ground splitting set  $S_{ng}$  is semantically equivalent to the ground splitting set  $S_{grnd}$  one gets by taking all ground instances of all predicates in  $S_{ng}$  from the ground program  $grnd(P)$ .

**Example 2.2.4** (Partial evaluation using splitting sets). Given program  $P_{col}$  from Listing 2.3 in Example 2.2.3, let  $S$  be the following (non-ground) splitting set:

$$S = \{coloring\_vertex/1, coloring\_edge/2, vertex/1, exclude\_vertex/1, edge/2, exclude\_edge/2\}$$

Using  $S$ , we can split  $P_{col}$  into bottom  $B_S(P_{col})$ , which is shown in Listing 2.4, and top  $T_S(P_{col})$  in Listing 2.5.

Listing 2.4: The bottom part of  $P_{col}$

```

1 vertex(a). vertex(b). vertex(c). vertex(d).
2 edge(a, b). edge(a, c). edge(a, d).
3 edge(b, c). edge(b, d). edge(c, d).
4 edge(X, Y) :- edge(Y, X).
5
6 exclude_vertex(d).
7 exclude_edge(V1, V2) :- edge(V1, V2), exclude_vertex(V1).
8 exclude_edge(V1, V2) :- exclude_edge(V2, V1).
9
10 coloring_vertex(V) :- vertex(V), not exclude_vertex(V).
11 coloring_edge(V1, V2) :-
12     edge(V1, V2), not exclude_edge(V1, V2).
```

Looking at  $B_S(P_{col})$ , we can observe that this sub-program of  $P_{col}$  is now stratified, and can therefore be evaluated using the straight-forward computation employed in

Example 2.2.2. We arrive at the following model for  $B_S(P_{col})$ :

$$LM(B_S(P_{col})) = \{coloring\_vertex(b), exclude\_vertex(d), edge(b, d), \\ edge(a, b), edge(c, d), edge(c, b), coloring\_edge(b, c), \\ coloring\_edge(a, c), coloring\_edge(b, a), edge(a, d), \\ coloring\_edge(c, a), vertex(c), vertex(a), edge(d, b), \\ exclude\_edge(d, b), exclude\_edge(c, d), exclude\_edge(a, d), \\ coloring\_vertex(c), exclude\_edge(b, d), coloring\_vertex(a), \\ edge(a, c), edge(b, c), edge(b, a), edge(d, c), coloring\_edge(a, b) \\ coloring\_edge(c, b), vertex(d), exclude\_edge(d, a), vertex(b) \\ edge(c, a), exclude\_edge(d, c), edge(d, a)\}$$

Listing 2.5: The top part of  $P_{col}$

```

1 % Guess colors
2 red(V) :- coloring_vertex(V), not green(V), not blue(V).
3 green(V) :- coloring_vertex(V), not red(V), not blue(V).
4 blue(V) :- coloring_vertex(V), not red(V), not green(V).
5
6 % Filter invalid guesses
7 :- coloring_vertex(V1), coloring_vertex(V2),
8     coloring_edge(V1, V2), red(V1), red(V2).
9 :- coloring_vertex(V1), coloring_vertex(V2),
10    coloring_edge(V1, V2), green(V1), green(V2).
11 :- coloring_vertex(V1), coloring_vertex(V2),
12    coloring_edge(V1, V2), blue(V1), blue(V2).

```

In order to compute the answer sets of the complete program, by the splitting set theorem, we only have to compute the answer sets of  $T_S(P_{col}) \cup pr(LM(B_S(P_{col})))$ . A simple (but computationally expensive) way to do this would be to perform an exhaustive search over all combinations of ground instances of the rules from lines 2 to 4 and drop all model candidates where any of the constraints fire. We finally get the following answer sets (filtered for predicates  $red/1$ ,  $green/1$  and  $blue/1$ ):

$$\begin{aligned}
A_1 &= \{blue(c), green(b), red(a)\} \\
A_2 &= \{blue(b), green(c), red(a)\} \\
A_3 &= \{blue(b), green(a), red(c)\} \\
A_4 &= \{blue(c), green(a), red(b)\} \\
A_5 &= \{blue(a), green(b), red(c)\} \\
A_6 &= \{blue(a), green(c), red(b)\}
\end{aligned}$$

Expanding on Example 2.2.4, it is clear that a stratifiable bottom with respect to some splitting set always exists - every program has at least the empty set as a trivial splitting set. We call the maximal stratifiable sub-program that only depends on facts (i.e. is a bottom w.r.t some splitting set) Common Base Program (CBP), see Definition 2.2.6.

**Definition 2.2.6** (Common Base Program, adapted from [Lan19]). Given a splitting set  $S$  of program  $P$ , the bottom  $B_S(P)$  is called *common base program*, i.e.  $CBP(P)$  if it is stratified and maximal in the sense that adding any further rule to  $B_S(P)$  would destroy the property of  $B_S(P)$  of being stratified.

What remains is to solve the remaining, non-stratifiable part of a program. State-of-the-art two-phased solvers such as Clingo employ solving techniques based on Conflict-driven Nogood Learning (CDNL) for this [GKS12]. The basic idea of CDNL is to encode a ground program as a set of constraints called *nogoods*, i.e. sets of literals that must not be contained in any answer set. Based on these nogoods, truth assignments over  $H_B(P)$  are "guessed" and conflicting guesses are used to potentially learn new nogoods using techniques originating in SAT-solving. However, these techniques normally require an input program to be fully ground, and therefore do not lend themselves to lazy-grounding approaches. Alpha uses a solving approach that is inspired by CDNL, but makes a number of modifications to apply the same principles in a lazy-grounding scenario. Section 2.2.2 gives a highlevel overview of a CDNL-based answer set search procedure that has first been implemented in Alpha [Wei17].

### 2.2.2 Employing CDNL-based answer set search for lazy-grounding solvers

Just like in traditional two-phased solvers, the lazy-grounding CDNL-approach uses *nogoods* to encode rules and constraints. However, since a full grounding of the program to solve is not available, some modifications are necessary. Without a full grounding, a solver must be able to distinguish between atoms in an assignment (i.e. answer set candidate) that are *true because some rule fired*, and atoms that *must be true because some constraint does not permit otherwise* (but haven't been derived by a rule). Lazy-grounding solvers introduce a third truth value, *must-be-true* (MBT), to reflect this. For an assignment to be an answer set, it must not contain any MBT values. Definitions 2.2.7 and 2.2.8 formally describe nogoods and their use to represent rules in this solving strategy.

**Definition 2.2.7** (Alpha-Assignment, Nogood [Wei17]). An *Alpha-Assignment*, in the following just *assignment* is a sequence  $A = \{sl_1, \dots, sl_n\}$  of literals  $l_1 \dots, l_n$  preceded by a *sign*  $s \in \{T, M, F\}$ . Intuitively, a signed literal  $sl_i$  assigns the truth value  $s$  to literal  $l$ . Possible truth values are *true* ( $T$ ), *false* ( $F$ ) and *must-be-true* ( $M$ ). Must-be-true is used as truth value for literals that are deemed by the solver to necessarily be true (e.g. because of a constraint precluding any other truth value), but there is no known firing rule (yet) which would allow assigning  $T$ . The *boolean projection*  $A^B$  of an assignment  $A$

is defined as:

$$A^B = \{Ta \mid Ta \in A \vee Ma \in A\} \cup \{Fa \mid Fa \in A\}$$

A *nogood*, in the variant used here, is a set of signed literals intuitively characterizing a set of literals that may not occur in an assignment with the given truth values:

$$ng = \{sl_1, \dots, sl_n\}, s \in \{T, F\}$$

Different from how nogoods are defined in classic two-phased solvers, a nogood may optionally have a specifically designated "head" literal. Given a nogood where  $sl_i$  is the head literal, we denote this as:  $ng = \{sl_1, \dots, sl_n\}_i$ . For a signed literal  $sl$ , we denote the literal with inverse sign as  $\overline{sl}$ . An assignment  $A$  *satisfies* a nogood  $ng$  if  $ng \not\subseteq A^B$ . Furthermore, an assignment  $A$  is a *solution* to a set  $\Delta$  of nogoods if  $\{a \mid Ta \in A^B\} \cap \{a \mid Fa \in A^B\} = \emptyset$ ,  $\{a \mid Ta \in A^B\} \cup \{a \mid Fa \in A^B\} = A^B$  and  $\forall ng \in \Delta : ng \not\subseteq A^B$ .

**Definition 2.2.8** (Nogood representation of rules [Wei17]). In order to represent a ground body of a rule, new atoms are introduced, denoted  $\beta(r, \sigma)$  for a rule  $r$  and variable substitution  $\sigma$ .

Given a rule  $r$  with head  $h(r) = a_0$ , positive body  $b^+(r) = a_1, \dots, a_k$  and negative body  $b^-(r) = a_{k+1}, \dots, a_n$ , the *nogood representation* of  $r$ ,  $ng(r)$  is defined as follows:

$$\begin{aligned} ng(r) = & \{ \{F\beta(r, \sigma), Ta_1, \dots, Ta_k, Fa_{k+1}, \dots, Fa_n\}_1, \{Fa_0, T\beta(r, \sigma)\}, \\ & \{T\beta(r, \sigma), Fa_1\}, \dots, \{T\beta(r, \sigma), Fa_k\}, \\ & \{T\beta(r, \sigma), Ta_{k+1}\}, \dots, \{T\beta(r, \sigma), Ta_n\} \} \end{aligned}$$

**Lazy Grounding and Solving Procedure [LW17]** In order to find answer sets in the lazy-grounding search procedure used by Alpha, a grounding component (grounder) always calculates ground substitutions for all rules that are *applicable* with respect to a (partial) assignment  $A$ . A rule  $r$  is said to be applicable w.r.t  $A$ , if, given a ground substitution  $\sigma$   $b^+(r\sigma) \subseteq A^B$  and  $b^-(r\sigma) \cap \{a \mid Ta \in A^B\} = \emptyset$ , i.e. a ground rule is applicable, if its positive body is satisfied and its negative body is not contradicted by an assignment. When solving a program, the grounder constructs an initial partial assignment from all facts and the model of the previously solved CBP. Based on this partial assignment, ground instances for applicable rules are computed and translated into nogoods. The partial assignment and generated nogoods are then passed to a solving component (solver). The solver then first applies propagation, i.e. extends assignment  $A$  based on unit nogoods as described in Definition 2.2.9. Then, if a conflict between the (possibly extended) assignment and the current set of nogoods exists, the conflict is analyzed, a new nogood describing the conflict cause is added, and the solver *backtracks* (i.e. retracts enough of the assignment to get to a conflict-free state). If there are no conflicts, and applicable rules exist, the solver guesses (based on a heuristic), which rule to apply. If there are no applicable rules or unassigned atoms, and no atom is assigned *must-be-true*, the solver has found an answer set. In this case, the answer set is stored, a

new nogood which prevents finding the same answer set again is added, and the solver backtracks. Algorithm 2.1 gives a highlevel summary of the answer set search procedure.

**Definition 2.2.9** (Weak and strongly unit nogoods [Wei17]). Let  $ng$  be a nogood  $ng = \{sl_1, \dots, s_n\}$  and  $A$  an assignment. Then

- $ng$  is *weakly unit* under  $A$  for  $sl$  if  $ng \setminus A^B = \{sl\}$  and  $\overline{sl} \notin A^B$
- $ng$  is *strongly unit* under  $A$  for  $sl$  if  $sl$  is designated the "head" of  $ng$ ,  $ng \setminus A = \{sl\}$  and  $\overline{sl} \notin A$

Intuitively, if a nogood is unit under an assignment  $A$ , it forces a truth value for the single literal  $sl$  which is not yet assigned (as otherwise, the nogood would be violated). Here, the notions of weak and strong unitness are used to distinguish between nogoods forcing a literal to be set to *must-be-true* (if implied by a weakly unit nogood) and *true* (if implied by a strongly unit nogood)

---

**Algorithm 2.1:** An answer set search procedure for lazy-grounding solvers [LW17].

---

**Input** : A non-ground program  $P$   
**Output** : The answer sets  $AS(P)$  of  $P$

- 1 Initialize  $AS = \emptyset$ , assignment  $A$ , nogood storage  $\Delta$ .
- 2 Run lazy grounder, obtain initial nogoods  $\Delta$  from facts.
- 3 **while** *search space not exhausted* **do**
- 4     Propagate on  $\Delta$ , extending  $A$ .
- 5     **if** *conflicting nogood exists* **then**
- 6         Analyze conflict, learn new nogood, backtrack.
- 7     **else if** *propagation extended A* **then**
- 8         Run lazy grounder w.r.t  $A$  to obtain new nogoods to extend  $\Delta$ .
- 9     **else if** *applicable rule exists* **then**
- 10         Guess rule to fire according to heuristic.
- 11     **else if** *unassigned atom exists* **then**
- 12         Assign all unassigned atoms to false.
- 13     **else if** *no atom assigned must-be-true in A* **then**
- 14          $AS \leftarrow AS \cup \{A\}$  Answer set found, add enumeration nogood and backtrack.
- 15     **else**
- 16         Backtrack.
- 17     **end**
- 18 **end**
- 19 **return**  $AS$

---





# The Evolog Language

The Evolog language extends (non-disjunctive) ASP as defined in the ASP-Core2 standard [CFG<sup>+</sup>20] with facilities to communicate with and influence the "outside world" (e.g. read and write files, capture user input, etc.) as well as program modularization and reusability features, namely *actions* and *modules*.

## 3.1 Actions in Evolog

Actions allow for an ASP program to encode operations with *side-effects* while maintaining fully declarative semantics. Actions are modelled in a functional style loosely based on the concept of monads as it is used in Haskell [Wad95]. Intuitively, to maintain declarative semantics, actions need to behave as pure functions, meaning the result of executing an action (i.e. evaluating the respective function) must be reproducible for each input value across all executions. On first glance, this seems to contradict the nature of IO operations, which inherently depend on some state, e.g. the result of evaluating a function *getFileHandle(f)* for a file *f* will be different depending on whether *f* exists, is readable, etc. However, at any given point in time - in other words, in a given state of the world - the operation will have exactly one result (i.e. a file handle or an error will be returned). A possible solution to making state-dependent operations behave as functions is therefore to make the state of the world at the time of evaluation part of the function's input. A function  $f(x)$  is then turned into  $f'(s, x)$  where  $s$  represents a specific world state. The rest of this section deals with formalizing this notion of actions.

### 3.1.1 Syntax

**Definition 3.1.1** (Action Rule, Action Program). An *action rule*  $R$  is of form

$$a_H : @t_{act} = act_{res} \leftarrow l_1, \dots, l_n.$$

where

- $a_H$  is an atom called *head atom*,
- $t_{act}$  is a functional term called *action term*,
- $act_{res}$  is a term called *(action-)result term*
- and  $l_1, \dots, l_n$  are literals constituting the *body* of  $R$ .

An *action program*  $P$  is a set of (classic ASP-)rules and action rules.

**Example 3.1.1.** Listing 3.1 shows a typical "Hello World"-program written using the action rule syntax outlined in Definition 3.1.1. It consists of only one rule, which has a head atom  $hello\_result(R)$ , the action term  $stream\_write[STDOUT, TEXT]$  and action result term  $R$ . The body consists of a regular atom  $hello\_text(TEXT)$  and an external atom  $\&stdout(STDOUT)$ .

Listing 3.1: A "Hello World"-program in Evolog.

```

1 hello_text("Hello World!") .
2 hello_result(R) : @streamWrite[STDOUT, TEXT] = R :-
    hello_text(TEXT), &stdout(STDOUT) .

```

#### 3.1.2 Semantics

To properly define the semantics of an action program according to the intuition outlined at the start of this section, we first need to formalize our view of the "outside world" which action rules interact with. We call the world in which we execute a program a *frame* - formally, action programs are always evaluated *with respect to a given frame*. The behavior of actions is specified in terms of *action functions*. The semantics (i.e. interpretations) of action functions in a program are defined by the respective frame.

#### Action Rule Expansion

To get from the practical-minded action syntax from Definition 3.1.1 to the formal representation of an action as a function of some state and an input, we use the helper construct of an action rule's *expansion* to bridge the gap. Intuitively, the expansion of an action rule is a syntactic transformation that results in a more verbose version of the original rule called *application rule* and a second rule only dependent on the application rule called *projection rule*. A (ground) application rule's head atom uniquely identifies the ground instance of the rule that derived it. As one such atom corresponds to one action executed, we call a ground instance of an application rule head in an answer set an *action witness*.

**Definition 3.1.2** (Action Rule Expansion). Given a non-ground action rule  $R$  with head atom  $a_H$ , action term  $f_{act}(i_1, \dots, i_n)$  and body  $B$  consisting of literals  $l_1, \dots, l_m$ ,

the expansion of  $R$  is a pair of rules consisting of an *application rule*  $R_{app}$  and *projection rule*  $R_{proj}$ .  $R_{app}$  is defined as

$$a_{res}(f_{act}, S, I, f_{act}(S, I)) \leftarrow l_1, \dots, l_n.$$

where  $S$  and  $I$  and function terms called *state*- and *input*-terms, respectively. An action rule's state term has the function symbol *state* and terms  $fn(l_1), \dots, fn(l_m)$ , with the expression  $fn(l)$  for a literal  $l$  denoting a function term representing  $l$ . The (function-)term representation of a literal  $p(t_1, \dots, t_n)$  with predicate symbol  $p$  and terms  $t_1, \dots, t_n$  uses  $p$  as function symbol. For a negated literal *not*  $p(t_1, \dots, t_n)$ , the representing function term is *not*( $p(t_1, \dots, p_n)$ ). The action input term is a "wrapped" version of all arguments of the action term, i.e. for action term  $f_{act}(t_1, \dots, t_n)$ , the corresponding input term is *input*( $t_1, \dots, t_n$ ). The term  $f_{act}(S, I)$  is called *action application term*.

The projection rule  $R_{proj}$  is defined as

$$a_H \leftarrow a_{res}(f_{act}, S, I, v_{res}).$$

where  $a_H$  is the head atom of the initial action rule  $R$  and the (sole) body atom is the action witness derived by  $R_{app}$ , with the application term  $f_{act}(S, I)$  replaced by a variable  $v_{res}$  called *action result variable*.

Looking at the head of an action application rule of format  $a_{res}(f_{act}, S, I, t_{app})$  with action  $f_{act}$ , state term  $S$ , input term  $I$  and application term  $t_{app}$ , the intuitive reading of this atom is "The result of action function  $f_{act}$  applied to state  $S$  and input  $I$  is  $t_{app}$ ", i.e. the action application term  $t_{app}$  is not a regular (uninterpreted) function term as in regular ASP, but an actual function call which is resolved using an interpretation function provided by a *frame* during grounding.

### Grounding of Action Rules

Grounding, in the context of answer set programming, generally refers to the conversion of a program with variables into a semantically equivalent, variable-free, version. Action application terms as introduced in Definition 3.1.2 can be intuitively read as variables, in the sense that they represent the result of applying the respective action function. Consequently, all action application terms are replaced with the respective (ground) result terms defined in the *frame* with respect to which the program is grounded.

**Definition 3.1.3** (Frame). Given an action program  $P$  containing action application terms  $A = \{a_1, \dots, a_n\}$ , a frame  $F$  is an interpretation function such that, for each application term  $f_{act}(S, I) \in A$  where  $S \in H_U(P)^*$  and  $I \in H_U(P)^*$ ,  $F(f_{act}) : H_U(P)^* \times H_U(P)^* \mapsto H_U(P)$ .

**Definition 3.1.4** (Grounding of action rules). Grounding of Evolog rules (and programs) always happens *with respect to a frame*. Given a frame  $F$ , an expanded action rule  $r_a$  and a (grounding) substitution  $\sigma$  over all body variables of application rule  $r_{a_{app}}$ , during grounding, every ground action application term  $t_{app}\sigma$  resulting from applying substitution  $\sigma$  is replaced with its interpretation according to  $F$ .

Example 3.1.2 demonstrates the expansion of an action rule as well as a compatible example frame for the respective action.

**Example 3.1.2** (Expansion and Frame). Consider following Evolog Program  $P$  which contains an action rule with action  $a$ :

$$\begin{aligned} & p(a). q(b). r(c). \\ & h(X, R) : @a(X, Z) = R \leftarrow p(X), q(Y), r(Z). \end{aligned}$$

The expansion of  $R$  is:

$$\begin{aligned} & a_{res}(a, state(p(X), q(Y), r(Z)), input(X, Z), a(state(p(X), q(Y), r(Z)), input(X, Z))) \leftarrow \\ & \quad p(X), q(Y), r(Z). \\ & h(X, R) \leftarrow a_{res}(a, state(p(X), q(Y), r(Z)), input(X, Z), R). \end{aligned}$$

Furthermore, consider following frame  $F$ :

$$F(a) = \{a(state(p(a), q(b), r(c)), input(a, c)) \mapsto success(a, c)\}$$

which assigns the result  $success(a, c)$  to the action application term (i.e. function call  $a(state(p(a), q(b), r(c)), input(a, c))$ ).

Then, the ground program  $P_{grnd}$  after action rule expansion is

$$\begin{aligned} & p(a). q(b). r(c). \\ & a_{res}(a, state(p(a), q(b), r(c)), input(a, c), success(a, c)) \leftarrow p(a). q(b). r(c). \\ & h(a, success(a, c)) \leftarrow a_{res}(a, state(p(a), q(b), r(c)), input(a, c), success(a, c)). \end{aligned}$$

The sole model of  $P$  with respect to frame  $F$  is

$$\begin{aligned} M = \{ & p(a), q(b), r(c), \\ & a_{res}(a, state(p(a), q(b), r(c)), input(a, c), success(a, c)) \\ & h(a, success(a, c)) \} \end{aligned}$$

### Evolog Models

Having introduced action rule expansions as well as frames, we now use these to extend the stable model semantics to Evolog programs.

**Definition 3.1.5** (Supportedness of Actions). Let  $r_{app}$  be a non-ground action application rule with head  $H = a_{res}(f_{act}, S, I, f_{act}(S, I))$ ,  $F$  a frame, and  $H_{grnd} = a_{res}(f_{act}, S_{grnd}, I_{grnd}, r)$  a ground instance of  $H$  with  $r$  being an arbitrary ground term. Then,  $H_{grnd}$  is *supported by  $F$* , if and only if  $F$  contains a mapping of form  $f_{act}(S_{grnd}, I_{grnd}) \mapsto r$ , i.e.  $r$  is a valid result of action function  $f_{act}$  with arguments  $S_{grnd}$  and  $I_{grnd}$  according to frame  $F$ . We call a ground instance of an action rule *supported by a frame* if the head of the corresponding application rule in the rule's expansion is supported by that frame.

**Definition 3.1.6** (Evolog-Reduct). Given a ground Evolog program  $P$ , a frame  $F$  and a set of ground atoms  $A$ , the *Evolog Reduct* of  $P$  with respect to  $F$  and  $A$   $P_F^A$  is obtained from  $P$  as follows:

1. Remove all rules  $r$  from  $P$  that are "blocked", i.e.  $A \not\models l$  for some negative body literal  $l \in b^-(r)$ .
2. Remove all action application rules from  $P$  which are not supported by  $F$ .
3. Remove the negative body from all other rules.

Note that the reduct outlined in Definition 3.1.6 extends the classic GL-reduct (see Definition 2.1.22) just by adding a check on action supportedness.

**Definition 3.1.7** (Evolog Model). A herbrand interpretation  $I$  of an Evolog Program is an *Evolog Model* ("answer set") of an Evolog program  $P$  with respect to a frame  $F$  if and only if it is a minimal classical model of its Evolog-Reduct  $P_F^A$ . We denote the set of Evolog Models of a program  $P$  as  $EM(P)$ .

### 3.1.3 Restrictions on Program Structure

While the action semantics outlined so far addresses the requirements for both declarativity and functionality of actions outlined in the introduction (see 1.3), we haven't yet addressed the demand for transparency, i.e. that every action that is executed must be reflected in an answer set of the respective program. With just the semantics outlined in Section 3.1.2, it would be possible to write programs such as the one shown in Example 3.1.3 where an action rule can fire, but the program is unsatisfiable.

**Example 3.1.3** (Unsatisfiable Program with Side-effects). The program below contains an action rule that can fire (because  $p(a)$  is true), but is also unsatisfiable due to the constraint in the last line.

$$\begin{aligned} & p(a). \\ & q(X) \leftarrow p(X). \\ & act\_done(X, R) : @act(X) = R \leftarrow p(X). \\ & \leftarrow q(X), act\_done(X, \_). \end{aligned}$$

This kind of programs raises some hard problems for implementations - given the contract that every side-effect of (i.e. action executed by) a program must be reported in an answer set, a solver evaluating the program from Example 3.1.3 would have to "retract" action  $act/1$  after finding that the program is unsatisfiable. Since it is generally not possible to "take back" side-effects (e.g. when some message is sent over a network broadcast to an unknown set of recipients), the only practical way to deal with this is to impose some conditions on programs with actions. Definition 3.1.8 details this notion and introduces *transparency* as a necessary condition for an Evolog program to be considered valid.

**Definition 3.1.8** (Action Transparency). An action rule  $r_a$  of an Evolog program  $P$  is *transparent* if, for every expanded ground instance  $gr_a$  it holds that, if  $gr_a$  fires, then the head  $h(gr_{a_{app}})$  of the respective application rule  $gr_{a_{app}}$  is contained in an answer set of  $P$ .

For an Evolog program to be valid, all its action rules must be transparent.

It follows from Definition 3.1.8 that only satisfiable programs can be transparent. Furthermore, note that Definition 3.1.8 aims to be as permissive as possible in terms of program structure. In general, it can not be assumed that transparency of all rules in a program can be guaranteed up-front. Implementations may therefore impose further restrictions on what is considered a valid Evolog program.

## 3.2 Program Modularization in Evolog

Modules aim to introduce new ways to re-use and unit-test to ASP code by enabling programmers to put frequently used sub-programs into *modules* which can be used in multiple programs.

Conceptually, an Evolog Module is a special kind of external atom (see 2.1.8) that refers to an ASP solver being called with some program and input.

### 3.2.1 Syntax

Definition 3.2.1 outlines the syntax for an Evolog module, i.e. a reusable sub-program the can be used in other programs.

**Definition 3.2.1** (Module Definition). The following EBNF describes a *module definition* in Evolog.

```

module_definition :
    "#module" ID "(" input_predicate "=>"
                    output_predicates ")"
    "{" statements "}";

input_predicate : predicate;
output_predicates :
    ("*" | ("{" predicate ("," predicate)* "}"));
predicate: ID "/" INTEGER;
statements : (fact | classic-rule | constraint)*;

```

In the context of the above grammar,

- ID are identifiers according to Definition 2.1.2,
- INTEGER are integers according to Definition 2.1.1,

- and `classical-rule` refers to a regular, i.e. non-evolog, ASP rule according to Definition 2.1.10

Intuitively, a module definition establishes a name for the sub-program in question, and specifies how input and output are translated to and from the module program. Definition 3.2.2 introduces *module atoms*, i.e. atoms in ASP rules that refer to modules.

**Definition 3.2.2** (Module Atom). Given a *module name*  $mod$ , a positive integer  $k$ , *input terms*  $t_1, \dots, t_n$  and *output terms*  $t_{n+1}, \dots, t_m$ , the expression

$$\#mod\{k\}[t_1, \dots, t_n](t_{n+1}, \dots, t_m)$$

is called a *module atom*. Syntactically, module atoms are regular atoms where  $\#mod$  is the predicate symbol and  $t_1, \dots, t_m$  are argument terms. The integer  $k$  is called the *answer set limit*. It denotes the number of answer sets of the module program to return. If  $k$  is omitted, all answer sets are returned.

Example 3.2.1 shows an encoding of the graph 3-coloring problem as an Evolog module.

**Example 3.2.1** (3-Coloring Module). The following module definition encodes the 3-coloring problem for a graph  $G = (V, E)$  as an Evolog module. Note that the module program assumes its input to be a single fact of form `graph(V, E)`, where  $V$  and  $E$  hold the lists of vertices and edges of  $G$ , respectively.

```

1 #module 3col(graph/2 => {col/2}) {
2     % Unwrap input
3     vertex_element(V, TAIL) :- graph(list(V, TAIL), _).
4     vertex_element(V, TAIL) :-
5         vertex_element(_, list(V, TAIL)).
6     vertex(V) :- vertex_element(V, _).
7     edge_element(E, TAIL) :- graph(_, list(E, TAIL)).
8     edge_element(E, TAIL) :-
9         edge_element(_, list(E, TAIL)).
10    edge(V1, V2) :- edge_element(edge(V1, V2), _).
11
12    % Make sure edges are symmetric
13    edge(V2, V1) :- edge(V1, V2).
14
15    % Guess colors
16    red(V) :- vertex(V), not green(V), not blue(V).
17    green(V) :- vertex(V), not red(V), not blue(V).
18    blue(V) :- vertex(V), not red(V), not green(V).
19
20    % Filter invalid guesses
21    :- vertex(V1), vertex(V2),

```

```
22         edge(V1, V2), red(V1), red(V2) .
23     :- vertex(V1), vertex(V2),
24         edge(V1, V2), green(V1), green(V2) .
25     :- vertex(V1), vertex(V2),
26         edge(V1, V2), blue(V1), blue(V2) .
27
28     col(V, red) :- red(V) .
29     col(V, blue) :- blue(V) .
30     col(V, green) :- green(V) .
31 }
```

### 3.2.2 Semantics

Intuitively, a module atom is a specialized external atom, and thus follows the general semantics of *fixed interpretation literals* (see 2.1.17). Rather than mapping to some annotated Java Method as is the case with regular external atoms, a module atom is interpreted by calling an ASP solver.

In order to make input terms of a module atom available to the module implementation program as facts, a translation step is necessary. Definition 3.2.3 describes this process in more detail.

**Definition 3.2.3** (Module Input Translation). Given a module definition  $M$  with input predicate  $p$  of arity  $k$  and implementation program  $P_M$  and a ground module atom  $a_m$  referencing  $M$  with input terms  $t_1, \dots, t_k$ , then the ASP program that is solved to interpret  $a_m$  is obtained by adding a fact  $p(t_1, \dots, t_k)$  to  $P_M$ . We denote this conversion of input terms to an atom as the "*factification*" function  $fct(t_1, \dots, t_k)$ .

Similar to module input translation, it is also necessary, to translate answer sets of an instantiated module program into output terms of the corresponding module atom. Every answer set gets converted into one set of terms, each - together with the input terms - constituting one ground instance of the module atom. Predicates get converted into function symbols, and atoms become list items. Definition 3.2.4 introduces the notion of a list term, which is used to represent lists of terms in ASP.

**Definition 3.2.4** (List Term). A list term is defined inductively as follows:

- The constant denoting the empty list, *empty\_list* is a list term.
- If  $t$  is a term and  $l$  is a list term, then  $s_{lst}(t, l)$  is a list term.

In the above,  $s_{lst}$  is a reserved function symbol denoting list terms.



**Definition 3.2.5** (Module Output Translation). Given a module definition with output predicates  $p_1, \dots, p_n$ , implementation program  $P_M$  and an answer sets  $A$  of  $P_M$ , the output terms of a corresponding module atom are obtained by converting the answer set into a set of terms. For every output predicate specified in the module definition, a function term is created with the predicate symbol as function symbol and a single list term as argument. The list term is constructed by collecting all atoms of the respective predicate in  $A$  into a list. We denote this conversion of an answer set as the "*termification*" function  $trm(A)$ .

Using the conversions from Definitions 3.2.3 and 3.2.5, we can now define the interpretation function for module atoms.

**Definition 3.2.6** (Module Atom Interpretation). Given (non-ground) module atom  $a_M$  with input terms  $t_1, \dots, t_k$  and output terms  $t_1, \dots, t_m$ , a module definition  $M$  with input predicate  $p$  and output predicates  $p_1, \dots, p_m$ , implementation program  $P_M$ , and ground substitution  $\sigma$ , the ground instance  $grnd(a_M)$  obtained by applying  $\sigma$  to  $a_M$  is true iff  $\{\exists A \in AS(P_M \cup \text{fact}(\sigma(t_1, \dots, t_k))) : trm(A) = \sigma(t_1, \dots, t_m)\}$ , i.e. the module atom is true if and only if there exists an answer set of the module program together with the factified input terms that, when termified, equals the output terms of the module atom.

**Corollary 3.2.1** (Fixed Module Interpretation). It follows from Definition 3.2.6 that module literals are *fixed interpretation literals* according to Definition 2.1.17, where the interpretation function is the calculation described in Definition 3.2.6. Since the implementation of a module does not change at runtime, the answer sets of the module program, i.e. the output terms of a corresponding module atom, only depend on the input terms supplied by a specific ground instance. The truth value of a given module literal  $l$  is therefore independent of the rest of the program  $l$  occurs in.

Example 3.2.2 demonstrates the use of modules to find solutions to an application of the bin-packing problem.

**Example 3.2.2** (Bin-packing Module). Suppose a group of computer scientists  $G = \{\text{bob}, \text{alice}, \text{dilbert}, \text{claire}, \text{cate}, \text{bill}, \text{carl}, \text{mary}\}$  plan to attend a conference on LLM-based AI agents in the neighboring town of Buzzwordville, which is 150 km distant. They have a number of cars at their disposal - Claire's camper van which seats 7 people, Dilbert's sedan seating 5, and Bob's roadster with 2 seats. Each car has different fuel efficiency, and the group wants to minimize the amount of fuel used.

To complicate matters even more, there are a number of additional constraints:

- For insurance reasons, each car must be driven by its owner.
- Alice and Carl are a couple and must travel together.
- Dilbert and Mary are not on speaking terms and must not travel together.

It is intuitively clear, that the problem at hand is a specialization of the bin-packing problem (with differently sized bins). We therefore introduce the Evolog module `bin_packing(instance2) ~ {item_packed2}`, which calculates solutions to the bin-packing decision problem, i.e. assignments of items to bins, such that all items are packed, and no bin capacity is exceeded. Listing 3.2 shows the module in question.

Listing 3.2: Bin-packing module

```
1 #module bin_packing(instance/2 => {item_packed/2}) {
2   % Unpack input lists.
3   bin_element(E, TAIL) :-
4       instance(lst(E, TAIL), _).
5   bin_element(E, TAIL) :-
6       bin_element(_, lst(E, TAIL)).
7   bin(B, S) :-
8       bin_element(bin(B, S), _).
9
10  item_element(E, TAIL) :-
11      instance(_, lst(E, TAIL)).
12  item_element(E, TAIL) :-
13      item_element(_, lst(E, TAIL)).
14  item(I, S) :-
15      item_element(item(I, S), _).
16  % For every item, guess an assignment to each bin.
17  { item_packed(I, B) : bin(B, _) } :- item(I, _).
18  % An item may only be assigned to one bin at a time
19  :- item_packed(I, B1),
20      item_packed(I, B2),
21      B1 != B2.
22  % We must not exceed the capacity of any bin.
23  capacity_used(B, C) :-
24      C = #sum{S : item(I, S),
25              item_packed(I, B)}, bin(B, _).
26  :- capacity_used(B, C),
27      C > S,
28      bin(B, S).
29  % Every item must be packed.
30  item_packed_somewhere(I) :-
31      item_packed(I, _).
32  :- item(I, _),
33      not item_packed_somewhere(I).
34 }
```

Since the implementation from 3.2 only solves the basic decision problem, we need additional code to represent the constraints in this specific variant of the problem and

derive a cost value for each valid assignment. Listing 3.3 shows a validation module that accepts a bin assignment and verifies it against the additional constraints.

Listing 3.3: Custom Constraints

```

1 #module assignment_valid(assignment/1 => {assignment/1}) {
2     assignment_element(1st(E, TAIL), E, TAIL) :-
3         assignment(1st(E, TAIL)).
4     assignment_element(ASSGN, E, TAIL) :-
5         assignment_element(ASSGN, _, 1st(E, TAIL)).
6     assigned(I, B, A) :-
7         assignment_element(A, item_packed(I, B), _).
8
9     owner_of(claire, van).
10    owner_of(bob, roadster).
11    owner_of(dilbert, sedan).
12
13    % For insurance reasons, each car must be
14    % driven by its owner, i.e. for each car,
15    % if it is used, the owner must be assigned to it.
16    car_in_use(C) :- assigned(_, C, A).
17    :- car_in_use(C), owner_of(P, C),
18        assigned(P, C2, A), C2 != C.
19
20    % Alice and Carl are a couple and
21    % must travel together.
22    :- assigned(alice, C1, A),
23        assigned(carl, C2, A), C1 != C2.
24
25    % Dilbert and Mary are not on speaking
26    % terms and must not travel together.
27    :- assigned(dilbert, C, A), assigned(mary, C, A).
28 }
```

Finally, we calculate the cost of driving one kilometer for each assignment, and collect all assignments with the lowest possible cost value using predicate `optimal_cost_assignment/2`. Listing 3.4 shows the full program, excluding the module definitions given in Listings 3.2 and 3.3.

Listing 3.4: Main Program

```

1 person(bob).
2 person(alice).
3 person(dilbert).
4 person(claire).
5 person(cate).
```

```

6 | person(bill).
7 | person(carl).
8 | person(mary).
9 |
10 | car(van, 7).
11 | car(roadster, 2).
12 | car(sedan, 5).
13 |
14 | distance(150).
15 | cost_per_km(van, 5).
16 | cost_per_km(roadster, 8).
17 | cost_per_km(sedan, 3).
18 |
19 | owner_of(claire, van).
20 | owner_of(bob, roadster).
21 | owner_of(dilbert, sedan).
22 |
23 | % Create input instance for bin-packing module
24 | instance(BINS, ITEMS) :-
25 |     BINS = #list{bin(C, S) : car(C, S)},
26 |     ITEMS = #list{item(P, 1) : person(P)}.
27 |
28 | % Calculate assignments and unfold list terms
29 | assignment_validation_output(ASSGN_VALID) :-
30 |     instance(BINS, ITEMS),
31 |     #bin_packing[BINS, ITEMS](ASSGN),
32 |     #assignment_vali[ASSGN](ASSGN_VALID).
33 | assignment(A) :-
34 |     assignment_validation_output(
35 |         lst(assignment(A), lst_empty)).
36 | assignment_element(lst(E, TAIL), E, TAIL) :-
37 |     assignment(lst(E, TAIL)).
38 | assignment_element(ASSGN, E, TAIL) :-
39 |     assignment_element(ASSGN, _, lst(E, TAIL)).
40 | assigned(I, B, A) :-
41 |     assignment_element(A, item_packed(I, B), _).
42 |
43 | % Calculate the total travelling cost for each assignment
44 | assignment_cost_per_km(A, C) :-
45 |     C = #sum{ CKM : cost_per_km(CAR, CKM),
46 |         assigned(_, CAR, A) },
47 |     assignment(A).
48 |

```

```

49 % Find the minimum cost per km over all eligible
50 % assignments and assignments with that cost
51 optimal_cost_per_km(C) :-
52     C = #min{CKM : assignment_cost_per_km(_, CKM)}.
53 optimal_cost_assignment(C, A) :-
54     optimal_cost_per_km(C), assignment_cost_per_km(A,
55         C).

```

The program from Listing 3.4 has a single answer set, with 16 instances of `optimal_cost_assignment/2`, i.e. 16 possible assignments of people to cars which minimize the cost of a driven kilometer (Note that we use the notation  $[a, b, c]$  for a list term holding terms  $a, b, c$ , and write terms of form *item\_packed*(*PERSON*, *CAR*) as tuples (*PERSON*, *CAR*), for brevity):

- `optimal_cost_assignment(8, [(alice, sedan), (bill, sedan), (bob, sedan), (carl, sedan), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, sedan), (bob, sedan), (carl, sedan), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, sedan), (bob, van), (carl, sedan), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, sedan), (bob, van), (carl, sedan), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, van), (bob, sedan), (carl, sedan), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, van), (bob, sedan), (carl, sedan), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, van), (bob, van), (carl, sedan), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, sedan), (bill, van), (bob, van), (carl, sedan), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, van), (bill, sedan), (bob, sedan), (carl, van), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, van), (bill, sedan), (bob, sedan), (carl, van), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])`
- `optimal_cost_assignment(8, [(alice, van), (bill, sedan), (bob, van), (carl, van), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])`

- *optimal\_cost\_assignment*(8, [(alice, van), (bill, sedan), (bob, van), (carl, van), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])
- *optimal\_cost\_assignment*(8, [(alice, van), (bill, van), (bob, sedan), (carl, van), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])
- *optimal\_cost\_assignment*(8, [(alice, van), (bill, van), (bob, sedan), (carl, van), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])
- *optimal\_cost\_assignment*(8, [(alice, van), (bill, van), (bob, van), (carl, van), (cate, sedan), (claire, van), (dilbert, sedan), (mary, van)])
- *optimal\_cost\_assignment*(8, [(alice, van), (bill, van), (bob, van), (carl, van), (cate, van), (claire, van), (dilbert, sedan), (mary, van)])

### 3.3 Relationship between Evolog- and Stable Model Semantics

The extensions to the usual ASP programming language described in the previous sections extend the original formalism with a notion of an outside world (through frames in the context of which, actions can be expressed) as well as a grouping mechanism for sub-programs into modules.

**Theorem 3.3.1** (Extension). Every ASP program  $P$  is a valid and semantically equivalent Evolog program in the sense that - for any given frame  $F$ , the Evolog Models of  $P$  are the same as its Answer Sets according to Stable Model Semantics 2.1.23.

*Proof.* First, we assert that every "regular" ASP program  $P$  is also a syntactically valid Evolog program. This follows directly from the definitions - since Evolog only adds syntactic support for actions (see Definition 3.1.1), but does not restrict regular ASP syntax, it follows that a syntactically correct ASP program is also a syntactically correct Evolog program.

Next, we show that for every regular ASP program  $P$  and any frame  $F$ , the Evolog Models of  $P$  are the same as its Stable Models, i.e.  $EM(P) = AS(P)$ :

Given any set of ground Atoms  $A$  from the Herbrand Base  $HB_P$  of  $P$ ,  $P$  is an Answer Set according to Stable Model Semantics if it is a minimal model of the GL-reduct 2.1.22  $P^A$  of  $P$  w.r.t.  $A$ . We recall that  $P^A$  is constructed as follows (see also see Definition 2.1.22):

- remove from  $P$  all rules  $r$  that are "blocked", i.e.  $A \not\models l$  for some literal  $l \in b^-(r)$
- and remove the negative body of all other rules.

Furthermore, consider how the Evolog-Reduct of  $P$  w.r.t.  $A$  and (any arbitrary) Frame  $F$ ,  $P_F^A$ , is constructed (see Definition 3.1.6):

- Remove all rules  $r$  from  $P$  that are "blocked", i.e.  $A \not\models l$  for some negative body literal  $l \in b^-(r)$ .
- Remove all action application rules from  $P$  which are not supported by  $F$ .
- Remove the negative body from all other rules.

Since  $P$  is a "regular", i.e. non-Evolog, ASP Program it contains no action application rules by definition. It is therefore clear that for any set answer set candidate  $A$  of  $P$  and any Frame  $F$ , the GL-Reduct  $P^A$  and Evolog-Reduct  $P_F^A$  coincide. Any minimal model of the GL-reduct is therefore also a minimal model of the Evolog-Reduct (and vice-versa) and therefore the Evolog Models of  $P$  are identical with its stable models, i.e.  $EM(P) = AS(P)$ .  $\square$





# Evolog Reference Implementation

While Chapter 3 gives a complete formal specification of the Evolog language extension, this chapter describes an implementation of said specification based on the Alpha ASP solver. All code referenced in this chapter is available on the official Github repository for Alpha [Lan].

## 4.1 Architectural overview of Alpha

Alpha is a lazy-grounding ASP solver implemented in Java. In a nutshell, Alpha calculates Computation Sequences (see Definition 2.2.1) for an input program using a CDNL-inspired solving algorithm. Starting from the set of facts contained in the program, an initial truth assignment is constructed. Based on this assignment, ground instances are calculated for all rules that *could potentially fire* based on the assignment. These ground instances are converted into *noGoods* and passed to the solver component which, using an adapted CDNL approach, guesses a new assignment based on the last set of noGoods. This process is repeated until no more guesses are possible, at which point the current assignment is either returned as an answer set, or some conflict is detected, in which case a new noGood is learned, and the solver backtracks. However, the actual ground-solve-loop is - while arguably at the heart of Alpha - just a small part of the process through which programs get evaluated. Figure 4.1 gives an overview of the building blocks making up the Alpha [WBB<sup>+</sup>19] system.

### 4.1.1 Parsing and Compilation

The core ground-and-solve component of Alpha supports only a subset of the input language described in Section 2.1.1. All language features supported by the parser, but not the solver itself, get compiled into equivalent constructs in the solver's internal representation. The following transformations are applied:

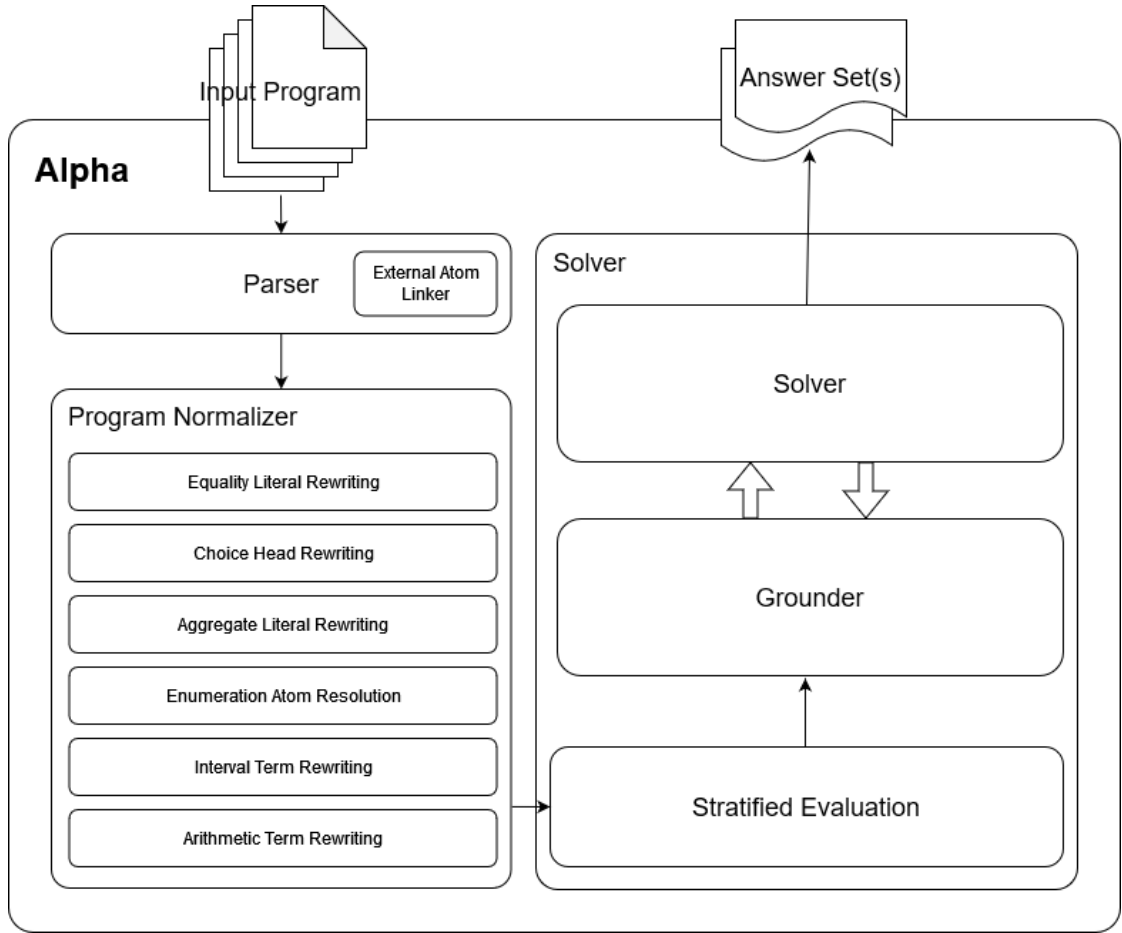


Figure 4.1: Alpha System Architecture

- *External Atom Linking*: Programs may only use external atoms whose implementations are known to the parser prior to parsing. Alpha provides a set of frequently used built-in external atoms out of the box, user-supplied code must be scanned through Alpha's API. Example A.2.1 demonstrates the use of user-supplied atom definitions. All external atom implementations are linked during parsing, i.e. every external atom in the parsed program holds a reference to the implementing Java Method.
- *Equality Literal Rewriting*: Literals like  $A = B$  in rule bodies that establish an equality between variables are removed by replacing one variable with the other (e.g.  $B$  with  $A$  in the example).
- *Choice Head Rewriting*: Rules with a choice head get replaced by a set of rules and constraints that is semantically equivalent to the choice rule.
- *Aggregate Literal Rewriting*: Aggregate literals like  $N = \#count\{ X : interesting(X) \}$

are replaced by a regular literal and a set of rules deriving instances of the replacement literal equivalent to the original aggregate literal. Subsection 4.1.1 goes into more detail on the rewriting process.

- *Enumeration Atom Resolution*: Alpha provides a feature where terms can be enumerated, i.e. the solver maps user-supplied terms to integers. This is used internally for Aggregate Literal Rewriting. Section 4.1.1 describes how the atoms in question are resolved.
- *Interval Term Rewriting*: Interval terms, i.e. terms of form  $A..B$  are transformed into regular variables that are bound through special internal literals which supply all values of the interval as ground instances of the variable.
- *Arithmetic Term Rewriting*: In order to simplify grounding later, terms constituting arithmetic expressions such as  $p(f(X*3, Y-4))$  are rewritten such that no arithmetic expressions occur in nested terms, i.e. the atom from before would be rewritten to  $p(f(R1, R2)), R1 = X * 3, R2 = Y - 4$ .

The result of the above list of transformations is what is called a *normalized* program in Alpha, which can directly be passed to the evaluation component and solved. Rewriting of Aggregate Literals and Resolution of Enumeration Atoms are of interest in the context of the Evolog reference implementation, and shall therefore be described in more detail.

### Enumeration Atoms

Alpha permits use of an "enumeration" solver directive, which allows programs to associate terms with consecutive integer keys. Listing 4.1 demonstrates using an enumeration to assign integer ordinals to a set of colors.

Listing 4.1: Using the Enumeration Directive to enumerate color symbols.

```

1 #enumeration_predicate_is ordinal.
2
3 color(white). color(red). color(magenta).
4 color(yellow). color(green). color(cyan).
5 color(blue). color(black).
6
7 numbered_color(COL, NUM) :-
8     color(COL), ordinal(colors, COL, NUM).
```

The directive `#enumeration_predicate_is` designates the predicate `ordinal/3` as an *Enumeration Predicate*. All occurrences of the an enumeration predicate get replaced with a special internal predicate `_Enumeration/3`. Listing 4.2 shows the program from before after transformation.

Listing 4.2: Transformed color numbering.

```

1 color(white). color(red). color(magenta).
2 color(yellow). color(green). color(cyan).
3 color(blue). color(black).
4
5 numbered_color(COL, NUM) :-
6     color(COL), _Enumeration(colors, COL, NUM).

```

Alpha's grounding component calculates valid ground substitutions for enumeration atoms as follows:

Given an enumeration atom  $a_e$  with terms  $t_{enum}$ ,  $t_{value}$  and  $t_{ord}$  and a partial substitution  $\sigma$ , assigning ground values to  $t_{enum}$  and  $t_{value}$ ,

- If the value  $\sigma t_{enum}$  is encountered for the first time, initialize a new empty map (i.e. set of pairs with unique first elements), and associate it with term  $\sigma t_{enum}$ .
- If the map for  $\sigma t_{enum}$  does not contain a mapping for key  $\sigma t_{value}$ , extend  $\sigma$  by the mapping  $t_{ord} \mapsto o$ , where  $o = s + 1$  and  $s$  denotes the current map size for  $\sigma t_{enum}$ . Add the mapping  $(\sigma t_{ord}, o)$  to the map and return the extended version of  $\sigma$ .
- If a mapping for  $\sigma t_{enum}$  and  $\sigma t_{value}$  exists, read the associated ordinal  $o$ , add it to  $\sigma$  and return the extended substitution.

From a semantics point of view, enumeration literals can intuitively be seen as "lazily assigned fixed-interpretation literals" (see Definition 2.1.17) in the way that every enumeration atom is true for exactly the ground substitution generated upon grounding it for the first time. The program from Listing 4.2 yields the following answer set:

$$A = \{color(black), color(blue), color(cyan), color(green), color(magenta), color(red), \\ color(white), color(yellow), numbered\_color(black, 1), numbered\_color(blue, 2), \\ numbered\_color(cyan, 3), numbered\_color(green, 4), numbered\_color(magenta, 6), \\ numbered\_color(red, 7), numbered\_color(white, 8), numbered\_color(yellow, 5)\}$$

### Aggregate Atoms

Alpha supports *Aggregate Literals* as defined in [CFG<sup>+</sup>20, p. 3] by rewriting programs containing aggregate literals into semantically equivalent aggregate-free programs. A detailed description of Alpha's implementation of Aggregate Rewriting is available at [Cona], but would exceed the scope of this Thesis. We will therefore focus on general concepts applying to all aggregate functions that are rewritten, and outline the rewriting procedure for aggregate literals where the minimum or maximum over a set of terms is calculated, which is the starting point for compilation of the newly introduced *#list* aggregate described in Section ??.

In the context of this section, we consider literals of form  $X \odot \#func\{t_1, \dots, t_n : l_1, \dots, l_m\}$ , where  $X$  is a term,  $\odot \in \{=, \leq\}$  and  $func \in \{min, max\}$ ,  $t_1, \dots, t_n$  are terms and  $l_1, \dots, l_m$  literals, respectively.

**Definition 4.1.1** (Aggregate Terms, Elements, Local and Global Variables, Dependencies). In the following, we use the notation  $var(l_1, \dots, l_n)$  for literals  $l_1, \dots, l_n$  to denote the set of variable terms occurring in said literals. Consider a rule  $r$  containing aggregate literal  $l_{agg} = X \odot \#func\{t_1, \dots, t_n : l_1, \dots, l_m\}$ :

$$H \leftarrow l_{agg}, b_1, \dots, b_k.$$

Then, the set  $var(l_1, \dots, l_n) \cap var(b_1, \dots, b_k)$  is called *global variables* of  $l_{agg}$ , denoted  $glob(l_{agg})$ . Roughly speaking, global variables of an aggregate literal are all variables occurring within the aggregate literal as well as other body literals of  $r$ . Given a set  $V = glob(l_{agg})$ , the set of literals  $dep(l_{agg})$  is the minimal set of literals that, given a substitution  $\sigma$ , must be ground after application of  $\sigma$ , in order for  $\sigma$  to also ground  $l_{agg} \cup dep(l_{agg})$ . Intuitively, global variables of an aggregate literals are all variables for which Alpha's lazy grounding component needs a ground value, in order to be able to calculate a ground instance of the aggregate literal itself. Dependencies of an aggregate literal  $l_{agg}$  are all literals of which the grounder needs ground instances in order to calculate a grounding of all global variables of  $l_{agg}$ .

In order to translate a rule  $r$  containing an aggregate literal  $l_{agg} = X \odot \#func\{t_1, \dots, t_n : l_1, \dots, l_m\}$  with global variables  $glob(l_{agg})$ , dependencies  $dep(l_{agg})$  into a semantically equivalent set of rules, the following steps are taken:

- Generate a unique identifier  $id(l_{agg})$  (typically some integer) for  $l_{agg}$
- Construct rule  $rt$  in which  $l_{agg}$  is replaced by a literal  $aggregate\_result(id(l_{agg})\_args, X)$
- Generate an *element rule* which derives one atom per element that is being aggregated over. Given the aggregate element  $t_1, \dots, t_n : l_1, \dots, l_m$ , the corresponding element rule is  $id(l_{agg})\_element\_tuple(id(l_{agg})\_args, t_1, \dots, t_n) \leftarrow l_1, \dots, l_m, dep(l_{agg})$ , i. e. the element rule body consists of all literals of the aggregate element together with all dependencies of the aggregate literal.
- Generate a set of *encoding rules* that encode the actual aggregate function over all elements as derived by the element rule and derives instances of the  $aggregate\_result/2$  predicate.

Example 4.1.1 demonstrates how an aggregate literal for the minimum function gets rewritten by Alpha.

**Example 4.1.1.** Consider the program from Listing 4.3. Based on some facts of type *employee/3* which assert that an employee works in some department and earns a given salary, we use a *#min*-aggregate to find the employee with the lowest salary in each department.

Listing 4.3: ASP program to find the worst paid employee per department.

```

1 employee(bob, sales, 2000).
2 employee(alice, development, 6000).
3 employee(dilbert, development, 4500).
4 employee(jane, sales, 3500).
5 employee(carl, controlling, 5000).
6 employee(bill, controlling, 4000).
7 employee(claire, development, 5000).
8 employee(mary, sales, 3000).
9 employee(joe, controlling, 5500).
10
11 department(DEP) :- employee(_, DEP, _).
12
13 min_salary(SAL, DEP) :-
14     SAL = #min{S : employee(_, DEP, S)},
15     department(DEP).
16 worst_paid(DEP, EMP) :-
17     min_salary(S, DEP), employee(EMP, DEP, S).

```

Listing 4.4 shows a rewritten version of the original program.

Listing 4.4: The program from Listing 4.3 in its rewritten version.

```

1 employee(bob, sales, 2000).
2 employee(alice, development, 6000).
3 employee(dilbert, development, 4500).
4 employee(jane, sales, 3500).
5 employee(carl, controlling, 5000).
6 employee(bill, controlling, 4000).
7 employee(claire, development, 5000).
8 employee(mary, sales, 3000).
9 employee(joe, controlling, 5500).
10
11 department(DEP) :- employee(_0, DEP, _1).
12 worst_paid(DEP, EMP) :-
13     min_salary(S, DEP), employee(EMP, DEP, S).
14 min_salary(SAL, DEP) :-
15     min_1_result(min_1_args(DEP), SAL), department(DEP).
16 min_1_element_tuple_less_than(ARGS, LESS, THAN) :-
17     min_1_element_tuple(ARGS, LESS),
18     min_1_element_tuple(ARGS, THAN), LESS < THAN.
19 min_1_element_tuple_has_smaller(ARGS, TPL) :-
20     min_1_element_tuple_less_than(ARGS, _3, TPL).
21 min_1_min_element_tuple(ARGS, MIN) :-

```

```

22     min_1_element_tuple(ARGS, MIN),
23     not min_1_element_tuple_has_smaller(ARGS, MIN).
24 min_1_result(ARGS, M) :-
25     min_1_min_element_tuple(ARGS, M).
26 min_1_element_tuple(min_1_args(DEP), S) :-
27     employee(_2, DEP, S), department(DEP).

```

In the rewritten version, the rule in line 14, which derives *min\_salary/2* has the aggregate literal replaced with a regular literal, instances for which are derived by newly added rules that together encode the aggregate function. The individual elements over which a minimum is being calculated are derived as instances of the *min\_1\_element\_tuple/2* predicate using the rule in line 26. In order to find the minimal instance of *min\_1\_element\_tuple/2*, we first establish a "less than"-relation (i.e. a partial order based on numeric comparison of the second term of the element tuple instances) using the rule in line 16. The minimum element is then the one for which we cannot find a "smaller" element. This element is derived by the rule in line 21. The single answer set of the rewritten program (filtered for instances of *worst\_paid/2*) is  $A = \{worst\_paid(controlling, bill), worst\_paid(development, dilbert), worst\_paid(sales, bob)\}$ .

#### 4.1.2 Evaluation

Once a program has been parsed and compiled using the steps described in 4.1.1, the compiled program can be evaluated (i.e. solved). Evaluation consists of two steps - first, the stratified bottom ("*common base program, CBP*", see 2.2.6) is evaluated using a bottom-up evaluation algorithm based on iterative fixpoint calculation as described in Definition 2.2.3. The remainder of the program, i.e. the part which cannot be handled using stratified evaluation is then solved using Alpha's CDNL-based answer set search described in Section 2.2.2. Since the stratified evaluation component is where Evolog Actions are handled, Section 4.1.2 gives an overview of how stratified evaluation is implemented in Alpha.

#### Stratified Evaluation in Alpha [Lan19]

Alpha's stratified evaluation component takes a normalized program as described in Section 4.1.1 as its input. A dependency graph for the program in question is calculated based on predicate dependencies based on Definition 2.2.5 where each dependency is labelled either "+" or "-" to distinguish dependencies through positive and negative body literals, respectively. Calculating the component graph (i.e. the graph resulting from condensing the dependency graph into its strongly connected components), one ends up with a directed acyclic graph, which is labelled such that all component nodes containing cycles through negative (i.e. labelled "-") dependencies get labelled as "unstratifiable". The CBP is then the program consisting of all rules from components that are not reachable by any path containing an unstratifiable component node. Based on the resulting subset of the component graph, a partition satisfying the criteria for a stratification

according to Definition 2.2.2 is calculated.

The stratified part of the input program is then evaluated in order of ascending stratum using Algorithm 4.1. For each stratum, we first calculate applicable ground rules based on all facts in the program as well as those derived when evaluating lower strata. As long as rule application yields new atoms, additional ground rules are computed and evaluated, until no new information can be derived, in which case a fixpoint has been reached on the current stratum. The result of the evaluation procedure is a program consisting of the rules of the non-stratified part of the input program, plus original facts as well as all new facts derived during stratified evaluation.

---

**Algorithm 4.1:** Stratified up-front evaluation of CBP in Alpha

---

**Input** : Stratification  $S = \{S_0, \dots, S_n\}$   
**Input** : Input Program  $P = (F_{in}, R_{in})$  consisting of facts and rules  $F_{in}, R_{in}$   
**Output** : partially evaluated program  $P_{eval}$

```

1 Facts  $F_{out} = facts(F_{in})$ 
2 foreach stratum  $S_i$  in  $S_0 \dots S_n$  do
3   initialize derived atoms in stratum  $A = F_{out}$ 
4   initialize derived atoms in individual run  $A_{new} = A$ 
5   do
6      $A \leftarrow A \cup A_{new}$ 
7      $A_{new} = \emptyset$ 
8     calculate applicable ground rules  $R_{app}$  in  $P_{S_i}$  based on  $A$ 
9     foreach rule  $r \in R_{app}$  do
10       $A_{new} \leftarrow A_{new} \cup fireRule(r)$ 
11    end
12  while new atoms derived in last run, i.e.  $A_{new} \neq A$ ;
13  add newly derived atoms to output facts,  $F_{out} \leftarrow F_{out} \cup A$ 
14 end
15 return  $P_{eval} = (F_{out}, (P \setminus S))$ 

```

---

## 4.2 Implementing the Evolog extension in Alpha

After the technical overview of Alpha in Section 4.1, we now turn to how Alpha has been adapted to implement the Evolog language extension specified in Chapter 3. Specifically, additions were made in the following areas:

- *Parsing* - Alpha's program parser has been extended to be able to handle rules with action heads, module definitions, and references to modules from rule bodies.
- *Aggregate Rewriting* - In order to accomodate *list aggregation literals*, a concise construction syntax for list terms, additional logic to compile this new type of



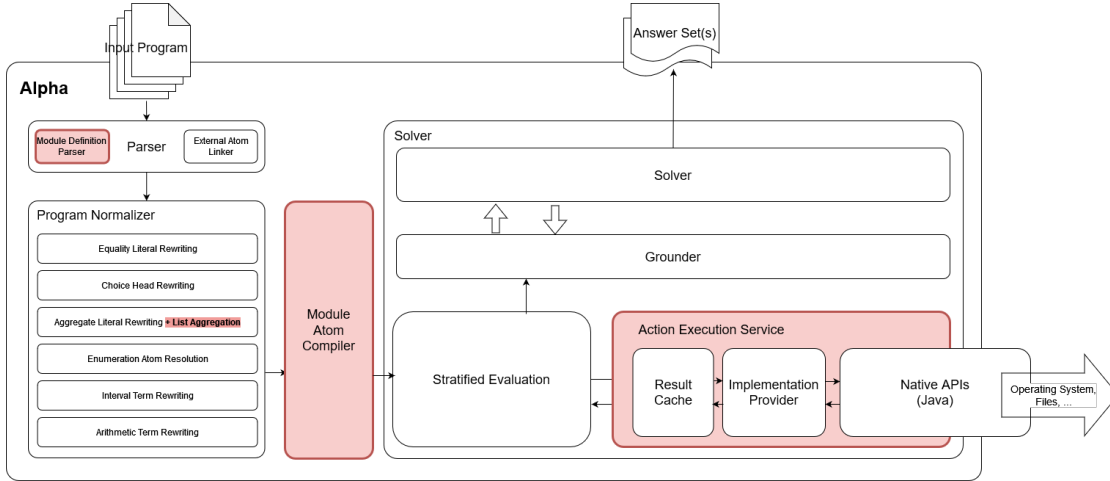


Figure 4.2: Alpha for Evolog System Architecture

aggregate literal has been added.

- *Action Execution Service* - A new component responsible for executing actions has been added. It ensures idempotency of action rule application and serves as an abstraction layer between actual action implementations and Alpha's stratified evaluation component.
- *Module Atom Compilation* - A new program preprocessing component has been added in order to compile module atoms, i.e. link name-based references to modules with the actual implementation code.

Figure 4.2 shows an updated overview of Alpha's architecture, with new components highlighted.

#### 4.2.1 Implementing Action Support

In this section, we give a detailed description of how Action support, conforming to definitions from Section 3.1 is implemented in Alpha.

**Implemented Actions** The actual actions provided in the reference implementation are intended to be sufficient for basic file (i.e. stream-)based input- and output operations. Specifically, Alpha offers actions to

- open a file handle for reading, i.e. obtaining an input stream,
- read from an input stream,
- close an input stream

- open a file handle for writing, i.e. obtaining an output stream,
- write to an output stream
- close an output stream

The idea of this selection of actions is that these can be used as the basic building blocks from which many more complex IO-Tasks, such as parsing specific file formats, waiting for user input, implementing command shells, etc. can be composed.

### Supported Programs

As stated in Section 3.1.3, Evolog programs are required to be *transparent* in terms of action execution, i.e. for every action that gets executed, there must be a corresponding witnessing atom in the respective answer set. This requirement is formalized in Definition 3.1.8. While the formal definition only states that if an action rule fires (i.e. an action is executed), the corresponding head must be contained in an answer set - which, aside from the effect that is applied on the outside world, is not any different from the general meaning of a rule that, if a the body is true, the head is as well - this leads directly to the practical problem of how to enforce this in an implementation. Since actions taken in the "outside world" (such as sending data over a network interface, writing into a file, ending a process, etc.) can not be retracted, an Evolog interpreter must be able to establish action rule transparency *prior to actual execution*. In general, any answer set search algorithm based on guesses and propagation through nogoods as described in Section 2.2.2 does not know up-front whether an atom that gets assigned as true at some point in search, will be true in a resulting answer set - every later propagation step could lead to a violation of one or more nogoods and subsequent backtracking. With that in mind, we establish that it is not feasible to have any form of guesses depend on the result of an action, since whenever a solver makes a guess, there is a chance the program might be unsatisfiable, in which case the action would not be any transparent anymore. While it may certainly possible to identify classes of programs with guesses where action transparency can be assured through clever static analysis, our implementation takes a simple approach. In the reference implementation based on Alpha, an Evolog program  $P$  is supported, if and only if  $P = cbp(e)$ , using the notion of the common base program as given in Definition 2.2.6. Since the CBP of a program is by definition stratified, we can be sure there will be an answer set, and therefore have a guarantee of action transparency as well. At the same time, programs using the guess-and-check pattern can still be executed under this restriction, by encapsulating the guess-and-check part in a module. Consider the program from Example 3.2.2, which has a single answer set, in which individual atoms correspond to answer sets of the used modules. In a program structured like that, one can use modules to encapsulate arbitrarily complicated calculations with any number of guesses while maintaining stratifiability of the top-level program. Example 4.2.1 demonstrates an unsupported program as well as a supported version of it based on a short interactive application.

**Example 4.2.1.** The program in Listing 4.5 is intended to be a very short "greeter" application. The user is asked to enter their name, which is then prepended with "Hello" and printed to the terminal. However, the constraints `:- usr_input_res(success(line(""))).` and `:- usr_input_res(error(_)).` render the program *unsupported*. In fact, the program is unsatisfiable with respect to any Frame (see Definition 3.1.3) where the user input is either empty or reading from `stdin` results in an error. Since, however, in order to evaluate the constraints in question, two actions with side-effects have to be performed first - namely one line needs to be written to `stdout` and one read from `stdin` - we would inevitably end up in a state where some side-effects have been applied to the outside world, but no corresponding action witness in an answer set exists.

Listing 4.5: An unsupported "greeter" application.

```

1  prompt_text("Hello user, enter your name: ").
2
3  write_prompt_res(R) : @streamWrite[STDOUT, PROMPT] = R :-
4      prompt_text(PROMPT), &stdout(STDOUT).
5  usr_input_res(INP) : @streamReadLine[STDIN] = INP :-
6      write_prompt_res(success(_)), &stdin(STDIN).
7
8  :- usr_input_res(success(line(""))).
9  :- usr_input_res(error(_)).
10
11 write_greeting_res(R) : @streamWrite[STDOUT, GREETING] = R
12     :-
13     usr_input_res(success(line(TEXT)), _),
14     &stdlib_string_concat["Hello ", TEXT](GREETING).
15     &stdout(STDOUT).
```

Listing 4.6 shows a supported version of the program from Listing 4.5. Here, rather than constraints, regular rules are used for error checking, and in case of an error, a corresponding message is written to `stdout`. In this version, the program might not run in the "intended" fashion, but there will always be an answer set, i.e. effects the program applies on the outside world will always be made transparent.

Listing 4.6: A supported version of the "greeter" application.

```

1  prompt_text("Hello user, enter your name: ").
2
3  write_prompt_res(R) : @streamWrite[STDOUT, PROMPT] = R :-
4      prompt_text(PROMPT), &stdout(STDOUT).
5  usr_input_res(INP) : @streamReadLine[STDIN] = INP :-
6      write_prompt_res(success(_)), &stdin(STDIN).
7
8  error("Input String empty") :-
9      usr_input_res(success(line(""))).
```

```
9      error(MSG):- usr_input_res(error(MSG)).
10     error_occurred :- error(_).
11
12     write_greeting_res(R) : @streamWrite[STDOUT, GREETING] = R
13     :-
14         usr_input_res(success(line(TEXT)), _), not
15         error_occurred.
16         &stdlib_string_concat["Hello ", TEXT](GREETING),
17         &stdout(STDOUT).
18
19     write_errmsg_res(R) : @streamWrite[STDOUT, ERRMSG] = R :-
20         error(ERR),
21         &stdlib_string_concat["An error occurred: ",
22                                ERR](ERRMSG),
23         &stdout(STDOUT).
```

### Action Implementation

Action execution in Alpha is tied into the stratified evaluation component described in Section 4.1.2. Specifically, whenever a ground instance of an action rule is fired, the corresponding action function is evaluated. The expansion of action rules described in Definition 3.1.2 happens implicitly: Instead of actually splitting a rule with an action head into an application- and projection-rule as described in the formal definition, the implementation constructs an internal representation of the ground rule in question, which is equivalent to an *action application term*  $f_{act}(S, I)$  (see Definition 3.1.2) in that it uniquely identifies a ground instance of an action rule within a program. After calculating the application for the ground rule to be fired, Alpha's action execution component queries an in-memory table called *action record* for existing entries where the key is equal to the calculated action application term. If such a mapping exists, the cached action result is returned, otherwise, the java method associated with the action function is executed, and a new mapping is added to the action record. The action record, i.e. action result cache, is used to make sure actions only get executed once, regardless of how often a certain ground instance of a rule is "fired" during program evaluation. Listing 4.7 shows an (abbreviated) version of Alpha's action execution component.

Listing 4.7: Alpha's action execution logic

```
public class ActionExecutionServiceImpl implements
    ActionExecutionService {

    private final ActionImplementationProvider
        actionProvider;
    private final Map<ActionInput, ActionWitness>
        actionRecord = new HashMap<>();
```

```
// [Constructors and utility methods omitted for
brevity]

@Override
public ActionWitness execute(String actionName, int
    sourceRuleId, Substitution sourceRuleInstance,
    List<Term> inputTerms) {
    ActionInput actInput = new
        ActionInput(actionName, sourceRuleId,
            sourceRuleInstance, inputTerms);
    return actionRecord.computeIfAbsent(actInput,
        this::execute);
}

private ActionWitness execute(ActionInput input) {
    Action action =
        actionProvider.getSupportedActions().get(input.name);
    ActionResultTerm<?> result =
        action.execute(input.inputTerms);
    return new ActionWitness(input.sourceRule,
        input.instance, input.name,
        input.inputTerms, result);
}

private static class ActionInput {

    private final String name;
    private final int sourceRule;
    private final Substitution instance;
    private final List<Term> inputTerms;

    // [Constructors and utility methods omitted for
    brevity]

}

}
```

Example 4.2.2 demonstrates a program where actions are used in recursive rules where ground instances potentially have to be evaluated multiple times and the aforementioned caching of action results comes into play.

**Example 4.2.2.** Listing 4.8 shows the "echo" program. It asks the user to enter text

on the command line, and "echoes", i.e. prints, the previous input until the user enters EXIT, at which point the program terminates. The "loop-like" behavior is realized through a positive recursive dependency cycle over predicates `write_prompt_res/2`, `usr_input_res/2` and `write_echo_res/2`.

Listing 4.8: An "echo" application which echoes user input written using Evolog actions.

```

1 prompt_text("Hello user, tell me something: ").
2 cancel_cmd("EXIT").
3
4 write_prompt_res(R, 0) : @streamWrite[STDOUT, PROMPT] = R :-
5     prompt_text(PROMPT), &stdout(STDOUT).
6 write_prompt_res(R, N) : @streamWrite[STDOUT, PROMPT] = R :-
7     prompt_text(PROMPT), write_echo_res(success(_), K),
8     N = K + 1, &stdout(STDOUT).
9 usr_input_res(INP, N) : @streamReadLine[STDIN] = INP :-
10    write_prompt_res(success(_), N), &stdin(STDIN).
11 write_echo_res(R, N) : @streamWrite[STDOUT, ECHO] = R :-
12    usr_input_res(success(line(TEXT)), N), TEXT != CANCEL_CMD,
13    cancel_cmd(CANCEL_CMD), &stdout(STDOUT),
14    &stdlib_string_concat["You said: ", TEXT](ECHO).
15
16 write_goodbye_res(R) : @streamWrite[STDOUT, MSG] = R :-
17    usr_input_res(success(line(TEXT)), _), cancel_cmd(CMD),
18    TEXT = CMD, MSG = "Goodbye, user! Sad to see you leave",
19    &stdout(STDOUT).

```

**Action Result Terms** The way action results are represented in Alpha takes its inspiration from the `Either` type present in many functional programming languages or libraries such as `Vavr` [Conb]. For example, an instance of type `Either<InputStream, Exception>` can either hold an instance of the "left" type, i.e. `InputStream` or an instance of the "right" type, i.e. `Exception`. Similarly, *action result terms* as defined in Definition 3.1.1 are function terms of form  $r_t(r_v)$ , where  $r_t \in \{success, error\}$  is the *result type* and  $r_v$  the *result value*. Result type *success* indicates the action completed normally, and in this case,  $r_v$  holds the result of the successful action, e.g. a term representing an input stream. If the result type is *error*, for example when trying to obtain a read handle on a non-existing file,  $r_v$  typically holds an error message. Listing 4.9 shows the Java Interface specifying an Action result term in Alpha.

Listing 4.9: Java definition of an action result term.

```

public interface ActionResultTerm<T extends Term> extends
    FunctionTerm {
    public static final String SUCCESS_SYMBOL = "success";
    public static final String ERROR_SYMBOL = "error";
}

```

```
/**
 * True if the action that generated this
 * result was successful (i.e. executed normally).
 */
boolean isSuccess();
/**
 * True if the action that generated this
 * result failed (i.e. threw an error in execution).
 */
boolean isError();
/**
 * Gets the actual value wrapped in this result.
 * Either a term representing the action return
 * value or a string term representing an error
 * message.
 */
T getValue();
}
```

### 4.2.2 Implementing Program Modularization

Since, formally, module atoms are just a special type of external atoms, implementing support for Modularization as described in 3.2 mainly has to deal with how to parse module definitions and how to construct external atoms according to Alpha's internal representation from a set of parsed module definition for atoms which reference these definitions. While not strictly necessary, this section also discusses the newly added `#list{...}` aggregate, which has been added as syntactic sugar for combining a set of terms into a single list term according to Definition 3.2.4.

#### List aggregation

External atoms are defined as having input- and output-*terms* (but not multisets of terms). At the same time, it is self-evident that - in most cases - one has to pass multiple input facts to a module program - for instance, a graph has sets of edges and vertices, all of which need to be passed to a module dealing with graph problems. This is where lists come in. By establishing the convention that the *single* input fact to a module program always has one or more *list terms* as arguments, we are able to encode all information that needs to be passed to the module into a single atom, while still staying true to external atom semantics as supported by Alpha. The practical problem arising from this is that constructing list terms in ASP is rather cumbersome. One has to write rules to

- specify *what* goes into the list, i.e. one or more rules encoding eligible *list elements*,

- specify *in what sequence* elements go into the list, i.e. rules which establish a total order between list elements,
- actually construct the list, i.e. starting from the end, recursively construct the term holding all list elements.

Listing 4.10 shows an example of this approach, where a list term is constructed for vertices and edges of a graph, respectively.

Listing 4.10: ASP code to create vertex and edge lists for a given graph.

```
1 %% pack vertices into a vertex list
2 vertex_element(E) :- vertex(E).
3 % First, establish ordering of elements
4 vertex_element_less(N, K) :-
5     vertex_element(N), vertex_element(K), N < K.
6 vertex_element_not_predecessor(N, K) :-
7     vertex_element_less(N, I), vertex_element_less(I, K).
8 vertex_element_predecessor(N, K) :-
9     vertex_element_less(N, K),
10    not vertex_element_not_predecessor(N, K).
11 vertex_element_has_predecessor(N) :-
12    vertex_element_predecessor(_, N).
13 % Now build the list as a recursively nested function term
14 vertex_lst_element(IDX, list(N, list_empty)) :-
15     vertex_element(N),
16     not vertex_element_has_predecessor(N),
17     IDX = 0.
18 vertex_lst_element(IDX, list(N, list(K, TAIL))) :-
19     vertex_element(N),
20     vertex_element_predecessor(K, N),
21     vertex_lst_element(PREV_IDX, list(K, TAIL)),
22     IDX = PREV_IDX + 1.
23 has_next_vertex_element(IDX) :-
24     vertex_lst_element(IDX, _),
25     NEXT_IDX = IDX + 1,
26     vertex_lst_element(NEXT_IDX, _).
27 vertex_lst(LIST) :-
28     vertex_lst_element(IDX, LIST),
29     not has_next_vertex_element(IDX).
30
31 %% pack edges into an edge list
32 edge_element(edge(V1, V2)) :- edge(V1, V2).
33 % First, establish ordering of elements
34 edge_element_less(N, K) :-
```



```

35     edge_element(N), edge_element(K), N < K.
36 edge_element_not_predecessor(N, K) :-
37     edge_element_less(N, I), edge_element_less(I, K).
38 edge_element_predecessor(N, K) :-
39     edge_element_less(N, K),
40     not edge_element_not_predecessor(N, K).
41 edge_element_has_predecessor(N) :-
42     edge_element_predecessor(_, N).
43 % Now build the list as a recursively nested function term
44 edge_lst_element(IDX, list(N, list_empty)) :-
45     edge_element(N),
46     not edge_element_has_predecessor(N),
47     IDX = 0.
48 edge_lst_element(IDX, list(N, list(K, TAIL))) :-
49     edge_element(N),
50     edge_element_predecessor(K, N),
51     edge_lst_element(PREV_IDX, list(K, TAIL)),
52     IDX = PREV_IDX + 1.
53 has_next_edge_element(IDX) :-
54     edge_lst_element(IDX, _),
55     NEXT_IDX = IDX + 1,
56     edge_lst_element(NEXT_IDX, _).
57 edge_lst(LIST) :-
58     edge_lst_element(IDX, LIST),
59     not has_next_edge_element(IDX).

```

Given facts  $vertex(a)$ ,  $vertex(b)$ ,  $vertex(c)$ ,  $edge(a, b)$ ,  $edge(b, c)$ ,  $edge(c, a)$ , we get the following answer set (filtered for predicates  $edge\_lst/1$  and  $vertex\_lst/1$ ):

$$A = \{edge\_lst(list(edge(c, a), list(edge(b, c), list(edge(a, b), list\_empty))))), vertex\_lst(list(c, list(b, list(a, list\_empty))))\}$$

Considering Listing 4.10, it is easy to see that construction of a list term looks always the same, regardless of the actual elements going into the list. Furthermore, we can observe a distinct similarity to the code generated to rewrite a  $\#min$ -aggregate outlined in Section 4.1.1. In the following, we define a new aggregate function,  $\#list$ , which Alpha rewrites into a generalized version of the list encoding from Listing 4.10.

**Definition 4.2.1** (List Aggregate). A *list aggregate* is an aggregate atom of the following form

$$t_{res} = \#list\{t_{elem} : l_1, \dots, l_n\}$$

where  $t_{res}$  and  $t_{elem}$  are terms called result- and element-term, respectively, and  $l_1, \dots, l_n$  are literals.

Definition 4.2.1 formally defines the syntax of a list aggregate. Note that - in contrast to general aggregate atoms, only equality comparisons, i.e.  $X = \#list\{\dots\}$ , are allowed, and we permit only a single element term rather than arbitrary tuples. Listing 4.11 shows a much shorter version of the program from Listing 4.10, in which list aggregation is used instead of step-by-step construction of list terms.

Listing 4.11: Creating vertex- and edge-lists using list aggregates.

```

1 %% pack vertices into a vertex list
2 vertex_lst(LIST) :-
3     LIST = #list{V : vertex(V)}.
4
5 %% pack edges into an edge list
6 edge_lst(LIST) :-
7     LIST = #list{edge(V1, V2) : edge(V1, V2)}.
```

In order to compile list-aggregates into their semantically equivalent aggregate-free versions, the following handling of list aggregates has been added to ALpha's aggregate rewriting logic:

- Substitute list aggregates in rules according to the general rules for aggregates outlined in 4.1.1.
- Element rules for list aggregates get constructed the same as for other aggregates, but permit only one element term rather than a tuple.
- In order to encode list construction, rules are added to construct a total order over all elements that should go into the list.
- List construction starts with the last element, which is defined to be the maximum element of the constructed order, successor elements are recursively added to the front of the list.
- The list is complete when a list has been constructed, such that there is no element smaller than the current head of the list.

Listing 4.12 shows the rewritten version of the program from Listing 4.11 (Note that in the hand-crafted example from Listing 4.10, lists are constructed in descending element order, rather than ascending as is the case for the aggregate-based version).

Listing 4.12: Rewritten list aggregates from Listing 4.11

```

1 vertex_lst(LIST) :-
2     list_1_result(list_1_no_args, LIST).
3 edge_lst(LIST) :-
4     list_2_result(list_2_no_args, LIST).
5 list_1_element_greater(ARGS, N, K) :-
```

```

6   list_1_element(ARGS, N),
7   list_1_element(ARGS, K),
8   N > K.
9 list_1_element_not_successor(ARGS, N, K) :-
10  list_1_element_greater(ARGS, N, I),
11  list_1_element_greater(ARGS, I, K).
12 list_1_element_successor(ARGS, N, K) :-
13  list_1_element_greater(ARGS, N, K),
14  not list_1_element_not_successor(ARGS, N, K).
15 list_1_element_has_successor(ARGS, N) :-
16  list_1_element_successor(ARGS, _0, N).
17 list_1_lst_element(ARGS, IDX, lst(N, lst_empty)) :-
18  list_1_element(ARGS, N),
19  IDX = 0,
20  not list_1_element_has_successor(ARGS, N).
21 list_1_lst_element(ARGS, IDX, lst(N, lst(K, TAIL))) :-
22  list_1_element(ARGS, N),
23  list_1_element_successor(ARGS, K, N),
24  list_1_lst_element(ARGS, PREV_IDX, lst(K, TAIL)),
25  IDX = PREV_IDX + 1.
26 list_1_has_next_element(ARGS, IDX) :-
27  list_1_lst_element(ARGS, IDX, _1),
28  NEXT_IDX = IDX + 1,
29  list_1_lst_element(ARGS, NEXT_IDX, _2).
30 list_1_result(ARGS, LIST) :-
31  list_1_lst_element(ARGS, IDX, LIST),
32  not list_1_has_next_element(ARGS, IDX).
33 list_1_element(list_1_no_args, V) :-
34  vertex(V).
35 list_2_element_greater(ARGS, N, K) :-
36  list_2_element(ARGS, N),
37  list_2_element(ARGS, K),
38  N > K.
39 list_2_element_not_successor(ARGS, N, K) :-
40  list_2_element_greater(ARGS, N, I),
41  list_2_element_greater(ARGS, I, K).
42 list_2_element_successor(ARGS, N, K) :-
43  list_2_element_greater(ARGS, N, K),
44  not list_2_element_not_successor(ARGS, N, K).
45 list_2_element_has_successor(ARGS, N) :-
46  list_2_element_successor(ARGS, _3, N).
47 list_2_lst_element(ARGS, IDX, lst(N, lst_empty)) :-
48  list_2_element(ARGS, N),

```

```
49     IDX = 0,
50     not list_2_element_has_successor(ARGS, N).
51 list_2_lst_element(ARGS, IDX, lst(N, lst(K, TAIL))) :-
52     list_2_element(ARGS, N),
53     list_2_element_successor(ARGS, K, N),
54     list_2_lst_element(ARGS, PREV_IDX, lst(K, TAIL)),
55     IDX = PREV_IDX + 1.
56 list_2_has_next_element(ARGS, IDX) :-
57     list_2_lst_element(ARGS, IDX, _4),
58     NEXT_IDX = IDX + 1,
59     list_2_lst_element(ARGS, NEXT_IDX, _5).
60 list_2_result(ARGS, LIST) :-
61     list_2_lst_element(ARGS, IDX, LIST),
62     not list_2_has_next_element(ARGS, IDX).
63 list_2_element(list_2_no_args, edge(V1, V2)) :-
64     edge(V1, V2).
```

### Parsing Module Definitions

Module definitions are parsed together with regular ASP code. Given a file containing a set of facts and rules and an arbitrary number of module definitions, Alpha's parser will group all rules and facts outside any module definition into one ASP program (i.e. the "main program"), and emit a set containing all encountered module definitions alongside with the parsed main program. Definition 4.2.2 details the technical representation of a module definition (which is basically a data structure storing all elements specified in the syntactical definition of a module, see 3.2.1). The actual code used in Alpha for parsing and storing module definitions can be found in the Appendix, see Example A.2.2.

**Definition 4.2.2** (Internal module representation). A parsed module definition gets stored as a tuple  $M = (name, p_{in}, OUT, PROG)$  where

- *name* is a string representing the name of the module. This must be unique, i.e. Alpha currently has no concept of namespaces or similar means of distinguishing equally named modules,
- $p_{in}$  is a predicate (represented as a string of form *name/arity*) called *input specification*,
- *OUT* denotes a (possibly empty) set of predicates called *output specification*,
- and *PROG* is the ASP program holding the actual implementation code for the module.

### Module Literal Compilation

Once a program and all contained module definitions have been parsed, every module atom (i.e. call to a module from a rule body) needs to be linked to its implementation. In Alpha, external atoms are evaluated during grounding. Definition 4.2.3 formally defines the notion of an *external predicate interpretation*, i.e. an interpretation function for ground instances of external atoms.

**Definition 4.2.3** (External Predicate Interpretation). Given an external atom  $\&ext[i_1, \dots, i_n](o_1, \dots, o_m)$  with input terms  $i_1, \dots, i_n$  and output terms  $o_1, \dots, o_m$ , and a substitution  $\sigma$  assigning ground values to at least all input terms of  $ex$ , the *predicate interpretation function*  $f : HU^n \mapsto 2^{HU^m}$  is a function which, for a given  $n$ -tuple over the Herbrand Universe  $HU$ , returns all  $m$ -tuples over  $HU$  for which the ground atom  $\&ext[\sigma(i_1), \dots, \sigma(i_n)](r_1, \dots, r_m)$  where  $(r_1 \dots, r_m) \in f(\sigma(i_1), \dots, \sigma(i_n))$  is true.

Implementation-wise, predicate interpretations are Java methods with a specific signature. Listing 4.13 shows the corresponding interface definition. Note that, for an implementation of `PredicateInterpretation` to be valid, it must be pure functional, i.e. not have any side-effects.

Listing 4.13: Java Interface for Predicate Interpretations

```
@FunctionalInterface
public interface PredicateInterpretation {

    Set<List<Term>> evaluate(List<Term> terms);

}
```

**Constructing interpretations for Module Atoms** Listing 4.14 shows the (abbreviated) Java code used in Alpha to construct a `PredicateInterpretation` for a module atom. Boundary checks and similar validations as well as auxiliary method calls are left out for easier readability. The interpretation itself is constructed using the Lambda-expression `PredicateInterpretation interpretation = terms -> { ... }`, where `terms` is the list of input terms, which gets wrapped into an atom of the predicate defined in `inputSpec` (i.e. the module input specification). Answer sets get filtered based on predicate names according to the module's output specification. Finally, each answer set is converted to a list of list terms using function `answerSetToTerms`, and the resulting set of term lists is returned.

Listing 4.14: Constructing module interpretations

```
private ExternalAtom translateModuleAtom(ModuleAtom atom,
    Map<String, Module> moduleTable) {
    ...
    Predicate inputSpec = definition.getInputSpec();
```

```
...
Set<Predicate> outputSpec = definition.getOutputSpec();
Set<Predicate> expectedOutputPredicates;
if (outputSpec.isEmpty()) {
    expectedOutputPredicates =
        calculateOutputPredicates(normalizedImplementation);
} else {
    expectedOutputPredicates = outputSpec;
}
...
PredicateInterpretation interpretation = terms -> {
    BasicAtom inputAtom = Atoms.newBasicAtom(inputSpec,
        terms);
    NormalProgram program = Programs.newNormalProgram(
        normalizedImplementation.getRules(),
        ListUtils.union(List.of(inputAtom),
            normalizedImplementation.getFacts()),
        normalizedImplementation.getInlineDirectives(),
        Collections.emptyList());
    java.util.function.Predicate<Predicate> filter =
        outputSpec.isEmpty() ? p -> true :
        outputSpec::contains;
    Stream<AnswerSet> answerSets =
        moduleRunner.solve(program, filter);
    if (atom.getInstantiationMode()
        .requestedAnswerSets().isPresent()) {
        answerSets = answerSets.limit(
            atom.getInstantiationMode()
                .requestedAnswerSets().get());
    }
    return answerSets.map(as -> answerSetToTerms(as,
        expectedOutputPredicates))
        .collect(Collectors.toSet());
};
return Atoms.newExternalAtom(atom.getPredicate(),
    interpretation, atom.getInput(), atom.getOutput());
}

private static List<Term> answerSetToTerms(AnswerSet
answerSet, Set<Predicate> moduleOutputSpec) {
    List<Term> terms = new ArrayList<>();
    for (Predicate predicate : moduleOutputSpec) {
        if (!answerSet.getPredicates().contains(predicate)) {
```

```
        terms.add(Terms.EMPTY_LIST);
    } else {
        terms.add(Terms.asListTerm(
            answerSet.getPredicateInstances(predicate).stream()
                .map(Atoms::toFunctionTerm)
                .collect(Collectors.toList())));
    }
}
return terms;
}
```





# Experimental Application

The goal of this chapter is to evaluate the implementation of Evolog described in Chapter 4 as a potential tool for application programming as described in Section 1.3. To that end, a complete application including user interaction, reading from and writing to files, as well as solving NP-hard search problems, is implemented purely in the Evolog language as introduced in the previous chapters. Specifically, the application in question, which we will refer to as *XML-graph-coloring* in the following sections, calculates 3-colorings for graphs, which are read from XML files. Calculated 3-colorings are again written to XML files. The file locations for both input and output files are interactively entered by the user on the commandline during runtime. More specifically, the application implements the following overall workflow:

- Upon startup, users are prompted to enter a path to a file containing an XML-encoded graph.
- The given file is read using *Action Rules* as described in Section 3.1.
- The XML data read from the input file is parsed into a function-term based representation using a dedicated xml-parsing *Module* (see Section 3.2.)
- Based on the graph representation from the previous step, another module is used to calculate 3-colorings of the parsed graph.
- Calculated 3-colorings are encoded as XML strings using a dedicated Evolog module.
- The user is prompted to enter a file path to which the XML-encoded graph colorings should be written.
- Generated colorings are written to the file indicated by the user.

The following section describes the XML-graph-coloring application in detail.

## 5.1 Interactive XML-based Graph Coloring

From an application architecture point of view, the XML-graph-coloring application consists of the following main blocks:

- Overall control flow, including user interaction and input/output actions.
- XML parsing logic, encapsulated in Module `xml_dom`.
- Graph 3-coloring calculation, implemented in Module `threecol`.
- Encoding of graph colorings as XML strings, which is realized in Module `threecol_to_xml`.

Since both the `xml_dom` as well as the `threecol_to_xml` module are relatively verbose and basically only deal with ordering elements, these are referred to as black boxes here. A detailed discussion including source code is given in Appendix A.3 and A.4. The implementation of the three-coloring module is the same as shown in Example and is therefore not discussed again here.

### 5.1.1 Reading files from user-supplied paths

Following the control flow of the application as a user perceives it, first, the user is asked to enter a file path. This file's content is then read line-by-line and aggregated into a list term. Listing 5.1 shows this part of the program.

Listing 5.1: Reading a file based on user input.

```
1 enter_input_prompt("Please enter a path to read graphs from:
   ").
2
3 write_input_prompt_res(R) : @streamWrite[STDOUT, PROMPT] = R :-
4     enter_input_prompt(PROMPT), &stdout(STDOUT).
5 usr_input_res(INP) : @streamReadLine[STDIN] = INP :-
6     write_input_prompt_res(success(_)), &stdin(STDIN).
7
8 infile(PATH) :- usr_input_res(success(line(PATH))).
9 infile_open(PATH, HD) : @fileInputStream[PATH] = HD :-
10    infile(PATH).
11
12 % Handle file opening error
13 error(io, MSG) :- infile_open(_, error(MSG)).
14
15 % Read all lines from infile
16 readline_result(PATH, 0, RES) :
17     @streamReadLine[STREAM] = RES :-
18     infile(PATH), infile_open(PATH, success(stream(STREAM))).
```

```

19 readline_result(PATH, LINE_NO, RES) :
20     @streamReadLine[STREAM] = RES :-
21     infile(PATH),
22     infile_open(PATH, success(stream(STREAM))),
23     readline_result(PATH, PREV_LINE_NO, PREV_LINE_RES),
24     PREV_LINE_RES != success(line eof),
25     LINE_NO = PREV_LINE_NO + 1.
26
27 % close stream after getting eof
28 infile_closed(PATH, RES) :
29     @inputStreamClose[STREAM] = RES :-
30     infile(PATH),
31     infile_open(PATH, success(stream(STREAM))),
32     readline_result(PATH, _, ok eof).
33
34 % Now create a list of content lines
35 file_lines(PATH, LST) :-
36     LST = #list{
37         line(LINE_NO, LINE) :
38             readline_result(PATH, LINE_NO,
39                             success(line(LINE))),
40             LINE != eof },
41     infile(PATH).

```

The rule on line 3 derives the predicate `write_input_prompt_res/1`, which wraps the result of a stream output operation - which in this case writes a prompt text to stdout. Note that the *standard output stream* STDOUT does not need to be opened using an Evolog action, but is always available to an application. A reference to STDOUT can be obtained through the external atom `stdout/1`.

If writing the prompt text succeeds, i.e. in case the result term of the write action has structure `success(_)`, user input is read from the *standard input stream* STDIN. The result of the read operation is derived on line 5 using predicate `usr_input_res/1`. Note that, as with stdout, STDIN does not have to be opened, but is obtained using the external atom `stdin/1`.

In case the read operation succeeded, we attempt to open an input stream on the file entered by the user. The rule on line 9 derives the result of this operation in the `infile_open/2` predicate. The first term is the path of the respective file, the second is an action result term which either takes the form `error(MSG)` or holds a reference to a file input stream `STREAM` wrapped in a function term `success(stream(STREAM))`.

If opening succeeds, the opened file is read line-wise, using the rules on lines 16 and 19. Lines are numbered in order to keep positional information of file content. Reading

stops once the result from the last `@streamReadLine` action is `success(line eof)` where `eof` denotes *end-of-file*.

After all lines of the input file have been read successfully an `@inputStreamClose` action is used to close the input file (see line 28). Finally, all lines from a file are aggregated into a single list term using a `#list` aggregate in the rule on line 35.

### 5.1.2 Parsing XML data

Based on the list of lines in an input file, the contained XML data is parsed using Module `xml_dom`, which is described in detail in Appendix A.3. Input files are expected to conform to the Document Type Definition (DTD) in Listing 5.2.

Listing 5.2: DTD for graph XML files.

```
1 <!ELEMENT graph (vertices, edges)>
2 <!ATTLIST graph directed (true | false) #REQUIRED>
3
4 <!ELEMENT vertices (vertex+)>
5 <!ELEMENT vertex (#PCDATA)>
6
7 <!ELEMENT edges (edge+)>
8 <!ELEMENT edge (source, target)>
9 <!ELEMENT source (#PCDATA)>
10 <!ELEMENT target (#PCDATA)>
```

Listing 5.3 shows an example input file conforming to the DTD which describes the complete graph  $K_3$ .

Listing 5.3: The complete graph  $K_3$  represented according to the DTD from Listing 5.2.

```
1 <graph directed="false">
2   <vertices>
3     <vertex>a</vertex>
4     <vertex>b</vertex>
5     <vertex>c</vertex>
6   </vertices>
7   <edges>
8     <edge>
9       <source>a</source>
10      <target>b</target>
11    </edge>
12    <edge>
13      <source>b</source>
14      <target>c</target>
15    </edge>
```

```

16         <edge>
17             <source>c</source>
18             <target>a</target>
19         </edge>
20     </edges>
21 </graph>

```

The XML-parsing module has 4 output terms which represent lists of Document Object Model (DOM) elements, parent-child relationships between DOM elements, text content of elements as well as attributes, respectively. Listing 5.4 shows the Evolog code to parse the XML content of an input file. Since each of the four result terms of the `xml_dom` module is a list term, in order to have one atom per list entry, each list needs to be "unwrapped". List terms are unwrapped by first deriving one "entry"-atom for every element of the list which holds the element itself along with all successor elements in the list (i.e. the "tail list"). Finally, the tail list of each entry atom is projected away using an additional rule, such that there is one atom for each element of the original list. In Listing 5.4, this unwrapping is performed for each output list of the XML parsing module from line 8 onward.

Listing 5.4: Parsing and consuming XML data in an Evolog program.

```

1  % Parse the string content of individual lines into a
2  % DOM-like representation
3  file_dom(PATH, dom(ELEMENTS, CHILDREN, TEXTS, ATTRIBUTES)) :-
4      file_lines(PATH, LINES),
5      #xml_dom[LINES](ELEMENTS, CHILDREN, TEXTS, ATTRIBUTES).
6
7  % Unwrap list elements from module output
8  dom_element_entry(NAME, ID, TAIL) :-
9      file_dom(_, dom(lst(dom_element(NAME, ID), TAIL), _, _),
10         _)).
11 dom_element_entry(NAME, ID, TAIL) :-
12     dom_element_entry(_, _, lst(dom_element(NAME, ID), TAIL)).
13 dom_element(NAME, ID) :-
14     dom_element_entry(NAME, ID, _).
15
16 dom_element_child_entry(NAME, ID, CHILD_ID, TAIL) :-
17     file_dom(_, dom(_, lst(
18         dom_element_child(NAME, ID, CHILD_ID), TAIL), _, _)).
19 dom_element_child_entry(NAME, ID, CHILD_ID, TAIL) :-
20     dom_element_child_entry(_, _, _, lst(
21         dom_element_child(NAME, ID, CHILD_ID), TAIL)).
22 dom_element_child(NAME, ID, CHILD_ID) :-
23     dom_element_child_entry(NAME, ID, CHILD_ID, _).

```

```
24 dom_element_text_entry(NAME, ID, TEXT, TAIL) :-
25     file_dom(_, dom(_, _, lst
26         (dom_element_text(NAME, ID, TEXT), TAIL), _)).
27 dom_element_text_entry(NAME, ID, TEXT, TAIL) :-
28     dom_element_text_entry(_, _, _, lst(
29         dom_element_text(NAME, ID, TEXT), TAIL)).
30 dom_element_text(NAME, ID, TEXT) :-
31     dom_element_text_entry(NAME, ID, TEXT, _).
32
33 dom_element_attribute_entry(
34     NAME, ID, ATTR_NAME, ATTR_VALUE, TAIL) :-
35     file_dom(_, dom(_, _, _, lst(
36         dom_element_attribute(
37             NAME, ID, ATTR_NAME, ATTR_VALUE), TAIL))).
38 dom_element_attribute_entry(
39     NAME, ID, ATTR_NAME, ATTR_VALUE, TAIL) :-
40     dom_element_attribute_entry(_, _, _, lst(
41         dom_element_attribute(
42             NAME, ID, ATTR_NAME, ATTR_VALUE), TAIL)).
43 dom_element_attribute(NAME, ID, ATTR_NAME, ATTR_VALUE) :-
44     dom_element_attribute_entry(
45         NAME, ID, ATTR_NAME, ATTR_VALUE, _).
```

### 5.1.3 Extracting a graph representation from parsed XML data

Based on the DOM-elements unwrapped in Listing 5.4, a representation of a graph using predicates `graph/1`, `graph_vertex/2`, `graph_directedness/2` and `graph_edge/2` is constructed using the code from Listing 5.5. DOM elements are translated into vertices and edges based on their names. Whether the graph in question is directed or undirected can be extracted from the corresponding attribute of the graph element.

Listing 5.5: Constructing a graph representation from parsed XML data.

```
1 % Translate generic DOM-elements into graphs
2 graph(ID) :- dom_element("graph", ID).
3 graph_vertex(ID, VERTEX) :-
4     graph(ID), dom_element("graph", ID),
5         dom_element_child("graph", ID, VERTEX_LIST_ID),
6         dom_element("vertices", VERTEX_LIST_ID),
7         dom_element("vertex", VERTEX_ID),
8         dom_element_child("vertices", VERTEX_LIST_ID,
9             VERTEX_ID),
10        dom_element_text("vertex", VERTEX_ID, VERTEX).
11 graph_directedness(ID, DIRECTED) :-
```

```

8      graph(ID), dom_element("graph", ID),
        dom_element_attribute("graph", ID, "directed",
        DIRECTED).
9 graph_undirected(ID) :- graph_directedness(ID, "false").
10 graph_directed(ID) :- graph_directedness(ID, "true").
11 graph_edge(ID, EDGE) :-
12     graph(ID), graph_vertex(ID, SRC_ID), graph_vertex(ID,
        TARGET_ID), dom_element("graph", ID),
        dom_element_child("graph", ID, EDGE_LIST_ID),
13     dom_element("edges", EDGE_LIST_ID), dom_element("edge",
        EDGE_ID), dom_element_child("edges", EDGE_LIST_ID,
        EDGE_ID),
14     dom_element("source", SRC_ELEM_ID),
        dom_element_child("edge", EDGE_ID, SRC_ELEM_ID),
        dom_element_text("source", SRC_ELEM_ID, SRC_ID),
15     dom_element("target", TARGET_ELEM_ID),
        dom_element_child("edge", EDGE_ID, TARGET_ELEM_ID),
        dom_element_text("target", TARGET_ELEM_ID, TARGET_ID),
16     EDGE = e(SRC_ID, TARGET_ID).
17 graph_edge(ID, e(TARGET, SRC)) :- graph_edge(ID, e(SRC,
        TARGET)), graph_undirected(ID).

```

#### 5.1.4 Calculating and storing graph colorings

Colorings for the graph parsed in the previous sections can be calculated using module `threecol` introduced in Example 3.2.1. For later processing (i.e. serialization to XML content), each calculated coloring is assigned a unique ID using the enumeration predicate `ids/3`.

Listing 5.6: Calculating graph colorings.

```

1 graph_threecol(ID, COL) :-
2     graph(ID),
3     VERTEX_LIST = #list{V : graph_vertex(ID, V)},
4     EDGE_LIST = #list{edge(V1, V2) : graph_edge(ID, e(V1,
        V2))},
5     #threecol[VERTEX_LIST, EDGE_LIST](COL).
6
7 graph_coloring(GRAPH, COLORING, ID) :- ids(coloring_ids,
        gcol(GRAPH, COLORING), ID), graph_threecol(GRAPH, COLORING).

```

In order to write calculated colorings to an XML file, each coloring is first converted to an XML string using module `threecol_to_xml`, which is shown in Listing ???. The module takes as input an atom of predicate `coloring/1` which is assumed to have

one argument term, where the argument is a list of terms representing vertices along with an assigned color, e.g. `coloring(lst(col(1, red), lst(col(2, blue), lst_empty)))`. In the listing, the rules on lines 2 through 8 are used to first "unwrap" the input list into one `coloring_entry/2` atom per list entry, and then establish a total ordering of said atoms using predicates `coloring_entry_pred/2`, `coloring_entry_first/1` and `coloring_entry_last/1`. Based on the derived ordering, an XML string representing the input coloring is built recursively starting from the "lowest" element. Every step of the construction process yields an instance of `coloring_xmlstr_upto/2` where the second argument term is the identifier of the "highest" vertex encoded in the XML string constituting the first argument. The output predicate of the module `coloring_xmlstr/1` is derived from the "maximal" instance of `coloring_xmlstr_upto/2`, i.e. the one corresponding to the last coloring entry according to `coloring_entry_last/1`.

Listing 5.7 shows how module `threecol_to_xml` is used in the application to obtain an XML string for each calculated graph coloring.

Listing 5.7: Generating XML strings for graph colorings.

```
1 % Translate colorings into strings
2 stringified_coloring_lst(GRAPH, COL_ID, COLSTR_LST) :-
3     graph_coloring(GRAPH, COLS, COL_ID),
4     #threecol_to_xml[COLS](COLSTR_LST).
5 stringified_coloring_entry(GRAPH, COL_ID, COL_STR, TAIL) :-
6     stringified_coloring_lst(GRAPH, COL_ID,
7         lst(coloring_xmlstr(COL_STR), TAIL)).
8 stringified_coloring_entry(GRAPH, COL_ID, COL_STR, TAIL) :-
9     stringified_coloring_entry(GRAPH, COL_ID, _,
10         lst(coloring_xmlstr(COL_STR), TAIL)).
11 stringified_coloring(GRAPH, COL_ID, COL_STR) :-
12     stringified_coloring_entry(GRAPH, COL_ID, COL_STR, _).
```

Concatenating the strings for each coloring of a respective graph into one XML string encoding all colorings found for the graph in question is done in much the same way as the string representations of individual colorings are constructed. Listing 5.8 illustrates this.

Listing 5.8: Concatenating XML strings of individual colorings.

```
1 % Stitch individual coloring strings into one string
2 stringified_coloring_before(GRAPH, COL_ID, COL_ID_AFTER) :-
3     stringified_coloring(GRAPH, COL_ID, _),
4     stringified_coloring(GRAPH, COL_ID_AFTER, _),
5     COL_ID_AFTER > COL_ID.
6 stringified_coloring_not_predecessor(GRAPH, COL_ID,
7     COL_ID_AFTER) :-
```



```

7   stringified_coloring_before(GRAPH, COL_ID, COL_ID_INTM),
8   stringified_coloring_before(GRAPH, COL_ID_INTM,
   COL_ID_AFTER).
9 stringified_coloring_predecessor(GRAPH, COL_ID, COL_ID_AFTER)
   :-
10  stringified_coloring_before(GRAPH, COL_ID, COL_ID_AFTER),
11  not stringified_coloring_not_predecessor(GRAPH, COL_ID,
   COL_ID_AFTER).
12 stringified_coloring_has_predecessor(GRAPH, COL_ID) :-
13  stringified_coloring(GRAPH, COL_ID, _),
14  stringified_coloring_predecessor(GRAPH, _, COL_ID).
15 stringified_coloring_first(GRAPH, COL_ID) :-
16  stringified_coloring(GRAPH, COL_ID, _),
17  not stringified_coloring_has_predecessor(GRAPH,
   COL_ID).
18 stringified_coloring_has_successor(GRAPH, COL_ID) :-
19  stringified_coloring(GRAPH, COL_ID, _),
20  stringified_coloring_before(GRAPH, COL_ID, _).
21 stringified_coloring_last(GRAPH, COL_ID) :-
22  stringified_coloring(GRAPH, COL_ID, _),
23  not stringified_coloring_has_successor(GRAPH, COL_ID).
24
25 xmlstring_upto(GRAPH, STR, COL_ID_FIRST) :-
26  &stdlib_string_concat("<colorings>", COLSTR) (STR),
27  stringified_coloring_first(GRAPH, COL_ID_FIRST),
28  stringified_coloring(GRAPH, COL_ID_FIRST, COLSTR).
29
30 xmlstring_upto(GRAPH, STR, COL_ID) :-
31  stringified_coloring_predecessor(GRAPH, COL_ID_PRED,
   COL_ID),
32  xmlstring_upto(GRAPH, PREV_STR, COL_ID_PRED),
33  stringified_coloring(GRAPH, COL_ID, COLSTR),
34  &stdlib_string_concat(PREV_STR, COLSTR) (STR).
35
36 colorings_xmlstring(GRAPH, RESULT_STR) :-
37  stringified_coloring_last(GRAPH, COL_ID),
38  xmlstring_upto(GRAPH, STR, COL_ID),
39  &stdlib_string_concat(STR, "</colorings>") (RESULT_STR).

```

### 5.1.5 Writing calculated graph colorings to XML files

Once all colorings for the graph(s) from the input file have been calculated and transformed into XML representation, the last remaining step is to actually write those XML strings

to files. The code used for this is shown in Listing 5.9. The way the user interaction is implemented is similar to how the input file path is obtained in Listing 5.1.

Listing 5.9: Writing Graph colorings to XML files.

```
1 % Prompt user for a path to which to write the calculated
  colorings for each graph
2 write_output_prompt_res(G, R) : @streamWrite[STDOUT, P] = R :-
3     graph(G), enter_output_prompt(PROMPT),
4     &stdlib_string_concat[PROMPT, G](S),
5     &stdlib_string_concat[S, ": "](P),
6     &stdout(STDOUT).
7 usr_output_res(G, INP) : @streamReadLine[STDIN] = INP :-
  write_output_prompt_res(G, success(_)), &stdin(STDIN).
8
9 % Open the output file
10 outfile(G, PATH) :- usr_output_res(G, success(line(PATH))).
11 outfile_open(G, PATH, HD) : @fileOutputStream[PATH] = HD :-
  outfile(G, PATH).
12
13 % Write colorings to output file
14 write_colorings_result(G, R) : @streamWrite[HD, STR] = R :-
15     colorings_xmlstring(G, STR),
16     outfile(G, PATH),
17     outfile_open(G, PATH, success(stream(HD))).
18
19 % Close the output file
20 outfile_closed(G, RES) : @outputStreamClose[STREAM] = RES :-
21     outfile(G, PATH), write_colorings_result(G,
22         success(_)),
23     outfile_open(G, PATH, success(stream(STREAM))).
```

## 5.2 Usage of the graph coloring application

In this Section, illustrate the graph coloring application's operation based on a simple input file.

**Example 5.2.1** (Trivial Graph). The simplest case on which to test our graph coloring application is a trivial graph consisting of one vertex. Listing 5.10 shows the corresponding input XML file.

Listing 5.10: One-vertex graph in XML encoding.

```
1 <graph directed="false">
2   <vertices>
```

```

3      <vertex>a</vertex>
4    </vertices>
5 </graph>

```

Listing 5.11 shows the solver call to run the XML-Graph-Coloring application for the input as well as all inputs and outputs during runtime.

Listing 5.11: Running the application for a one-vertex graph.

```

1 C:\...\evolog-thesis\evolog-samples\xml-graphcol> alpha-solver
  -i .\file-based-3col.evl -i .\xml_dom.mod.evl -i
  .\3col-module-with-lists.evl -i .\3col_to_xml.mod.evl
2 Please enter a path to read graphs from:
3 inputs/trivial.xml
4 Please enter a path to write calculated colorings for graph 1:
5 outputs/trivial.out.xml
6 Answer set 1:
7 { ... }

```

The (single) answer set  $A$  of the program is rather lengthy in its unfiltered form and is therefore given as the union of sets  $A_{input}$ ,  $A_{dom}$ ,  $A_{col}$ ,  $A_{string}$ ,  $A_{output}$  in which we group atoms by functional area they relate to. We denote as  $A_{input}$  the set of atoms in  $A$  which are derived by rules dealing with obtaining user input (i.e. the path from which to read graph specifications):

$$\begin{aligned}
 A_{input} = \{ & \text{enter\_input\_prompt}(\text{"Please enter a path to read graphs from : "}), \\
 & \text{write\_input\_prompt\_res}(\text{success}(\text{ok})), \\
 & \text{usr\_input\_res}(\text{success}(\text{line}(\text{"inputs/trivial.xml"}))), \text{infile}(\text{"inputs/trivial.xml"}), \\
 & \text{infile\_open}(\text{"inputs/trivial.xml"}, \text{success}(\text{stream}(\text{inputStream\_2}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 0, \text{success}(\text{line}(\text{"<graph directed = \"false\">"}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 1, \text{success}(\text{line}(\text{"<vertices>"}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 2, \text{success}(\text{line}(\text{"<vertex>a</vertex>"}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 3, \text{success}(\text{line}(\text{"</vertices>"}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 4, \text{success}(\text{line}(\text{"</graph>"}))), \\
 & \text{readline\_result}(\text{"inputs/trivial.xml"}, 5, \text{success}(\text{line}(\text{eof}))), \\
 & \text{infile\_closed}(\text{"inputs/trivial.xml"}, \text{success}(\text{closeResult}(\text{ok}))), \\
 & \text{file\_lines}(\text{"inputs/trivial.xml"}, \\
 & \quad \text{lst}(\text{line}(0, \text{"<graph directed = \"false\">"}), \\
 & \quad \text{lst}(\text{line}(1, \text{"<vertices>"}), \text{lst}(\text{line}(2, \text{"<vertex>a</vertex>"}), \\
 & \quad \text{lst}(\text{line}(3, \text{"</vertices>"}), \text{lst}(\text{line}(4, \text{"</graph>"}), \\
 & \quad \text{lst\_empty})))))) \}
 \end{aligned}$$

Similarly,  $A_{dom}$  contains all atoms in the answer set which model the DOM representation of the input XML file. For brevity, we do not list every individual list term of the *file\_dom* instance (since the atoms resulting from unwrapping the list are part of  $A_{dom}$  anyway). Also, helper atoms for partial list entries needed for unwrapping (such as *dom\_element\_entry*) are not included:

$$A_{dom} = \{dom\_element("graph", 1), \\ dom\_element("vertex", 3), \\ dom\_element("vertices", 2), \\ dom\_element\_attribute("graph", 1, "directed", "false"), \\ dom\_element\_child("graph", 1, 2), \\ dom\_element\_child("vertices", 2, 3), \\ dom\_element\_text("vertex", 3, "a")\}$$

The subset  $A_{col}$  holds atoms relating to the actual model of the graph as well as calculated colorings:

$$A_{col} = \{graph(1), graph\_vertex(1, "a"), graph\_undirected(1), \\ graph\_vertices(1, lst("a", lst\_empty)), graph\_edges(1, lst\_empty), \\ graph\_compact(1, lst("a", lst\_empty), lst\_empty), \\ graph\_coloring(1, lst(col("a", blue), lst\_empty), 2), \\ graph\_coloring(1, lst(col("a", green), lst\_empty), 3), \\ graph\_coloring(1, lst(col("a", red), lst\_empty), 1)\}$$

$A_{string}$  holds all atoms derived by rules dealing with serializing our calculated colorings into a single XML string. Since there is a relatively high number of pure "helper" atoms

that are only used to impose orderings etc., this set is given in a much abbreviated form:

```

A_string = {stringified_coloring(1, 1,
    "<coloring><vertex>
        <id>a</id><color>red</color></vertex></coloring>"),
stringified_coloring(1, 2,
    "<coloring><vertex>
        <id>a</id><color>blue</color></vertex></coloring>"),
stringified_coloring(1, 3,
    "<coloring><vertex>
        <id>a</id><color>green</color></vertex></coloring>"),
colorings_xmlstring(1,
    "<colorings>
        <coloring>
            <vertex><id>a</id><color>red</color></vertex>
        </coloring>
        <coloring>
            <vertex><id>a</id><color>blue</color></vertex>
        </coloring>
        <coloring>
            <vertex><id>a</id><color>green</color></vertex>
        </coloring>
    </colorings>")}
```

Last but not least,  $A_{output}$  groups together all atoms derived while prompting users for output paths to write colorings to, as well as actually writing the output file(s):

```

A_output = {write_output_prompt_res(1, success(ok)),
    usr_output_res(1, success(line("outputs/trivial.out.xml"))),
    outfile(1, "outputs/trivial.out.xml"),
    outfile_open(1, "outputs/trivial.out.xml", success(stream(outputStream_3))),
    write_colorings_result(1, success(ok)),
    outfile_closed(1, success(ok))}
```

The output file written to `trivial.out.xml` is given in Listing 5.12.

Listing 5.12: XML encoding of calculated colorings for the input file from Listing 5.10

```
1 <colorings>
```

```
2      <coloring>
3          <vertex>
4              <id>a</id>
5              <color>red</color>
6          </vertex>
7      </coloring>
8      <coloring>
9          <vertex>
10             <id>a</id>
11             <color>blue</color>
12         </vertex>
13     </coloring>
14     <coloring>
15         <vertex>
16             <id>a</id>
17             <color>green</color>
18         </vertex>
19     </coloring>
20 </colorings>
```

### 5.3 Performance of solving Evolog programs

In order to gain some idea of Alpha's solving performance for Evolog programs, a slightly modified version of the graph coloring program has been used where all rules that read user input interactively (and therefore are blocking) were removed and instead facts were added for the `infile/1` and `outfile/2` predicates.

#### 5.3.1 Test setup and inputs

Alpha's solving performance for this modified version of the XML-based-3-coloring application has been benchmarked against calculating colorings for the same graphs represented as ASP facts and without writing calculated colorings to XML files. Listing 5.13 shows the "classic ASP" representation used for the complete graph with 3 vertices  $K_3$ .

Listing 5.13: Pure-ASP version of a 3-coloring encoding for the graph  $K_3$ .

```
1 vertex(1) .
2 vertex(2) .
3 vertex(3) .
4
5 edge(1,2) .
6 edge(2,3) .
7 edge(3,1) .
8
```

```

9  % Make sure edges are symmetric
10 edge(V2, V1) :- edge(V1, V2).
11
12 % Guess colors
13 red(V) :- vertex(V), not green(V), not blue(V).
14 green(V) :- vertex(V), not red(V), not blue(V).
15 blue(V) :- vertex(V), not red(V), not green(V).
16
17 % Filter invalid guesses
18 :- vertex(V1), vertex(V2), edge(V1, V2), red(V1), red(V2).
19 :- vertex(V1), vertex(V2), edge(V1, V2), green(V1), green(V2).
20 :- vertex(V1), vertex(V2), edge(V1, V2), blue(V1), blue(V2).
21
22 col(V, red) :- red(V).
23 col(V, blue) :- blue(V).
24 col(V, green) :- green(V).

```

In addition to  $K_3$  as the simplest input graph, the application was tested against the following randomly generated graphs:

- G8-05-4: A connected graph with 8 vertices and a "density factor" of 0.5, i.e. a likelihood of 50% of there being an edge between any two vertices.
- G13-03-4: A connected graph with 13 vertices and a density factor of 0.3.
- G13-05-4: A connected graph with 13 vertices and a density factor of 0.5.
- G21-03-4: A connected graph with 21 vertices and a density factor of 0.3.
- G34-02-3: A connected graph with 34 vertices and a density factor of 0.2.
- G55-01-4: A connected graph with 34 vertices and a density factor of 0.1.

Ascending Fibonacci numbers were chosen for vertex counts in order to have an exponentially growing graph size, but avoid unintentional regularities in the input graphs, such as only having graphs with even vertex count, vertex count always being a multiple of some factor, etc.

### 5.3.2 Performance Test Results

The results of running the above described performance tests are summarized in Table ??.

Table 5.1: All execution times are given in seconds.

Instance	Classic-3col	XML-3col
K3	0.684	1.202
G8-05-4	0.731	1.520
G13-03-4	0.726	1.816
G13-05-4	0.694	1.871
G21-03-4	0.775	3.359
G34-02-3	0.805	10.318
G55-01-4	0.830	28.830

We can observe that solving performance of the basic 3-coloring program stays relatively stable across all graphs. While runtime does grow with increasing vertex count, the absolute difference is very small and would be of no concern in most real-world applications. However, in the Evolog version, where graphs are read from and written to XML files, we can observe a clearly exponential increase in runtime in step with the number of input vertices. Together with the fact that the pure 3-coloring part of the application (i.e. calculating colorings based on a graph represented as ASP atoms) does not seem to suffer from this kind of performance penalty, it seems clear that the XML processing part of the application exponentially scales with input size. This does not come as a complete surprise given that the atom counts for some predicates derived during parsing are (upper-)bounded by the square of the number of vertices in the input graph (e.g. all intermediate predicates needed to establish orderings of XML tags, list elements, etc.). See Appendix A.3 for a detailed discussion of the parser implementation.



# Discussion

In this chapter, we discuss observations from the experiments conducted in Chapter 5. It is split into two main parts: In the first part, we discuss recurring patterns in code written in Evolog as observed on the model application written in Chapter 5. In the second part of this chapter, we take a look at the performance metrics gained from the testing described in Section 5.3.

## 6.1 Observations from writing Evolog code

Considering the example application discussed in Section 5.1, this section aims to highlight potential areas for improvement of the language.

### 6.1.1 Unwrapping list terms

As the examples in Section 5.1 clearly show, there is a certain amount of boilerplate code involved whenever one has to "unpack" a list term obtained as an output term of a module literal. Since in most cases where list terms come into play, one actually wants to work on individual values on the list, we end up with many iterations of similar-looking code that is needed to transform one atom with a single list term into a set of "element atoms", each referring to one element of the list. An example of this is the subset of the 3-Coloring module (see Example 3.2.1) where the module input (an atom referencing a list of vertices and a list of edges) is deconstructed in order to represent a graph using instances of predicates `vertex/1` and `edge/2`. The relevant part of the program is shown in Listing 6.1.

Listing 6.1: Unwrapping list terms representing a graph

```

1 vertex_element(V, TAIL) :- graph(lst(V, TAIL), _).
2 vertex_element(V, TAIL) :- vertex_element(_, lst(V,
    TAIL)).

```

```

3      vertex(V) :- vertex_element(V, _).
4      edge_element(E, TAIL) :- graph(_, lst(E, TAIL)).
5      edge_element(E, TAIL) :- edge_element(_, lst(E, TAIL)).
6      edge(V1, V2) :- edge_element(edge(V1, V2), _).

```

The six lines of code shown in Listing 6.1 deal with deconstructing two list terms, a vertex list and an edge list, respectively, into individual atoms such that each atom corresponds to one element of the list. The transformation always follows the same pattern:

- One rule derives an instance of an "element atom" from the head of the list, which is obtained by deconstructing the list term into the form `lst(V, TAIL)`, where `TAIL` is the rest of the list without the head value represented as variable `V`.
- A second rule recursively derives further list elements from element atoms by further deconstructing the `TAIL` terms of the individual element atoms. Repeated evaluation of this rule eventually leads to one element atom for every element of the original list term.
- Finally, a third rule derives the actual domain atoms which are of interest by projecting away the partial lists from the individual element atoms.

Example 6.1.1 demonstrates this method of unwrapping list terms based on a list representation of the complete graph with 3 vertices  $K_3$ .

**Example 6.1.1** (List Unwrapping). Listing 6.2 shows the "graph unwrapping" program from Listing 6.1 including a fact representing the complete graph  $K_3$  encoded as two list terms.

Listing 6.2: Unwrapping the list representation of  $K_3$ .

```

1      graph(
2          lst(a, lst(b, lst(c, lst_empty))),
3          lst(edge(a, b), lst(edge(b, c), lst(edge(c,
4              a), lst_empty)))).
5      vertex_element(V, TAIL) :- graph(lst(V, TAIL), _).
6      vertex_element(V, TAIL) :- vertex_element(_, lst(V,
7          TAIL)).
8      vertex(V) :- vertex_element(V, _).
9      edge_element(E, TAIL) :- graph(_, lst(E, TAIL)).
10     edge_element(E, TAIL) :- edge_element(_, lst(E, TAIL)).
11     edge(V1, V2) :- edge_element(edge(V1, V2), _).

```

Listing 6.3 shows the (single) answer set of the program given in Listing 6.1

Listing 6.3: The unwrapped list representation of  $K_3$ .

```

1 Answer set 1:
2     { edge(a, b)
3       edge(b, c)
4       edge(c, a)
5       edge_element(edge(a, b), lst(edge(b, c), lst(edge(c,
6         a), lst_empty)))
7       edge_element(edge(b, c), lst(edge(c, a), lst_empty))
8       edge_element(edge(c, a), lst_empty)
9       graph(
10         lst(a, lst(b, lst(c, lst_empty))),
11         lst(edge(a, b), lst(edge(b, c), lst(edge(c,
12           a), lst_empty))))
13       vertex(a)
14       vertex(b)
15       vertex(c)
16       vertex_element(a, lst(b, lst(c, lst_empty)))
17       vertex_element(b, lst(c, lst_empty))
18       vertex_element(c, lst_empty) }

```

**Generalizing list-unwrapping** As Example 6.1.1 demonstrates, unwrapping a list term always takes 3 rules that only differ in predicate names. Definition 6.1.1 introduces a potential "syntactic sugar" solution we call an *unwrap rule*.

**Definition 6.1.1** (List-Unwrapping-Rule). Given an Atom  $a$  with terms  $t_1, \dots, t_k, \dots, t_n$ , where  $t_k$  is a list term, we define the following rule to be the *unwrap rule* for  $t_k$ :

$$e(V) \leftarrow \#unwrap\{V \mid t_k\}, a(t_1, \dots, t_k, \dots, t_n).$$

In the unwrap rule,  $e(V)$  is called an *element atom*, where  $V$  refers to one value from the list  $t_k$ . We define the semantics of the above rule to be equivalent to the following program:

$$\begin{aligned}
list\_element(V, TAIL) &\leftarrow a(t_1, \dots, lst(V, TAIL), \dots, t_n). \\
list\_element(V, TAIL) &\leftarrow list\_element(\_, lst(V, TAIL)). \\
e(V) &\leftarrow list\_element(V, \_).
\end{aligned}$$

Listing 6.4 shows a potential implementation of the 3-Coloring Module from Example 3.2.1 using the shorthand notation for list unwrapping proposed in Definition 6.1.1.

Listing 6.4: 3-Coloring Module with shorthand list unwrapping rules.

```

1 #module threecol(graph/2 => {col/2}) {
2     % Unwrap input

```

```
3      vertex(V) :- #unwrap{V | VLST}, graph(VLST, _).
4      edge(V1, V2) :- #unwrap{edge(V1, V2) | ELST}, graph(_,
5                          ELST).
6
7      % Make sure edges are symmetric
8      edge(V2, V1) :- edge(V1, V2).
9
10     % Guess colors
11     red(V) :- vertex(V), not green(V), not blue(V).
12     green(V) :- vertex(V), not red(V), not blue(V).
13     blue(V) :- vertex(V), not red(V), not green(V).
14
15     % Filter invalid guesses
16     :- vertex(V1), vertex(V2), edge(V1, V2), red(V1),
17         red(V2).
18     :- vertex(V1), vertex(V2), edge(V1, V2), green(V1),
19         green(V2).
20     :- vertex(V1), vertex(V2), edge(V1, V2), blue(V1),
21         blue(V2).
22
23     col(V, red) :- red(V).
24     col(V, blue) :- blue(V).
25     col(V, green) :- green(V).
26 }
```

### 6.1.2 Repetitive File-IO Actions

As we can observe in Listings 5.1 and 5.9, respectively, reading the content of a file, as well as writing to a file require a set of rules that looks the same regardless of the specific file or content in question. Listing 6.5 shows the "standard program" to read all lines from a file:

- A file path (i.e. location in storage), represented using the fact `infile(PATH)` is opened.
- Assuming opening the file produced no error, the first line of content is read using the rule on line 11.
- Based on the first line read (line number 0), we recursively continue reading as long as reading the last line was successful and did not yield `eof` (i.e. a pseudo-term denoting the end of content) as a result.
- If `eof` has been read, the file is closed, and lines are represented as atoms of form `line(LINE_NO, LINE)`.

Listing 6.5: Reading all lines from a file.

```

1  infile_open(PATH, HD) : @fileInputStream[PATH] = HD :-
2      infile(PATH).
3
4  % Handle file opening error
5  error(io, MSG) :- infile_open(_, error(MSG)).
6
7  % Read all lines from infile
8  readline_result(PATH, 0, RES) :
9      @streamReadLine[STREAM] = RES :-
10     infile(PATH), infile_open(PATH,
11         success(stream(STREAM))).
12 readline_result(PATH, LINE_NO, RES) :
13     @streamReadLine[STREAM] = RES :-
14     infile(PATH),
15     infile_open(PATH, success(stream(STREAM))),
16     readline_result(PATH, PREV_LINE_NO, PREV_LINE_RES),
17     PREV_LINE_RES != success(line eof)),
18     LINE_NO = PREV_LINE_NO + 1.
19
20 % close stream after getting eof
21 infile_closed(PATH, RES) :
22     @inputStreamClose[STREAM] = RES :-
23     infile(PATH),
24     infile_open(PATH, success(stream(STREAM))),
25     readline_result(PATH, _, ok eof)).
26
27 % Extract actual lines and numbers
28 line(LINE_NO, LINE) :-
29     readline_result(PATH, LINE_NO, success(line(LINE))),
30     LINE != eof.

```

The code to write a set of lines to a file looks similar to how a file is read. Listing 6.6 demonstrates the typical implementation. The following steps are needed:

- First, the file to which content is to be written is opened.
- Assuming lines to be written are consecutively numbered and available as atoms of form `line(LINE_NO, LINE)`, the first line (assumed to have an index of 0) is written.
- If the first line has been successfully written, all other lines are written using a recursive rule.

- We derive that all lines have been written when the line with maximum index has been written.
- Once `all_lines_written` has been derived, the file is closed.
- The file is also closed in case an error occurs during writing.

Listing 6.6: Writing lines to a file.

```
1 % Open the output file
2 open_result(PATH, RES) : @fileOutputStream[PATH] = RES :-
    outfile(PATH).
3
4 % Write lines in order of ascending line number
5 write_result(0, RES) : @streamWrite[STREAM, LINE] = RES :-
    open_result(PATH, success(stream(STREAM))), outfile(PATH),
    line(0, LINE).
6 write_result(LINE_NO, RES) : @streamWrite[STREAM, LINE] = RES
    :-
7     open_result(PATH, success(stream(STREAM))),
8     write_result(LINE_NO - 1, success(ok)),
9     outfile(PATH), line(LINE_NO, LINE).
10 all_lines_written :- write_result(LINE_NO, success(ok)),
    LINE_NO = #max{NUM : line(NUM, _)}.
11
12 % The file should be closed once all lines were successfully
    written
13 should_close(PATH, STREAM) :- all_lines_written,
    open_result(PATH, success(stream(STREAM))), outfile(PATH).
14
15 % The file should also be closed when an IO error occurs
    during writing
16 should_close(PATH, STREAM) :-
17     write_result(_, error(_)),
18     open_result(PATH, success(stream(STREAM))),
19     outfile(PATH).
20
21 % Close the file
22 close_result(PATH, RES) : @outputStreamClose[STREAM] = RES :-
23     should_close(PATH, STREAM),
24     open_result(PATH, success(stream(STREAM))),
25     outfile(PATH).
```

**Module-based abstraction of actions** As we’ve demonstrated in Listings 6.5 and 6.6, frequently used file operations like reading all lines from and writing a set of lines to a file, form repetitive code patterns that would greatly benefit from some shortened abstraction. The obvious solution seems to be to simply permit actions in Evolog Modules. However, this approach presents some challenges with regards to transparent semantics.

Recalling the demand for *action transparency* (see Definition 3.1.8), i.e. that the result of every firing (ground) action rule must be part of an answer set, modules containing actions can not simply be used in module literals as they’re described in Section 3.2. Instead, we’d have to apply the definitions of action functions in rule heads to an Evolog Module and have a module function as an action interpretation function. Example 6.1.2 demonstrates this approach.

**Example 6.1.2** (Action Modules). In this example, we show a potential implementation for a module that takes a list of numbered strings and a string describing a file location as input, and writes all lines to the referenced file. Listing 6.7 gives an example of how a module definition for a module containing actions might look.

Listing 6.7: Prototypical definition of a module containing Evolog actions.

```

1 #action write_lines(lines/1, path/1 => (success:
   lines_written/1 | error: io_error/1)) {
2     % Unwrap line list
3     line(IDX, LINE) :- #unwrap{line(IDX, LINE) | LST},
       lines(LST).
4     % Unwrap 1-element-list path/1
5     outfile(PATH) :- #unwrap{P | PLST}, path(PLST).
6
7     % Open the output file
8     open_result(PATH, RES) : @fileOutputStream[PATH] = RES
       :- outfile(PATH).
9
10    % Write lines in order of ascending line number
11    write_result(0, RES) : @streamWrite[STREAM, LINE] =
       RES :- open_result(PATH, success(stream(STREAM))),
       outfile(PATH), line(0, LINE).
12    write_result(LINE_NO, RES) : @streamWrite[STREAM,
       LINE] = RES :-
13        open_result(PATH, success(stream(STREAM))),
14        write_result(LINE_NO - 1, success(ok)),
15        outfile(PATH), line(LINE_NO, LINE).
16    all_lines_written :- write_result(LINE_NO,
       success(ok)), LINE_NO = #max{NUM : line(NUM, _)}.
17
18    % The file should be closed once all lines were
       successfully written

```

```

19      should_close(PATH, STREAM) :- all_lines_written,
      open_result(PATH, success(stream(STREAM))),
      outfile(PATH).
20
21      % The file should also be closed when an IO error
      occurs during writing
22      should_close(PATH, STREAM) :-
23          write_result(_, error(_)),
24          open_result(PATH, success(stream(STREAM))),
          outfile(PATH).
25
26      % Close the file
27      close_result(PATH, RES) : @outputStreamClose[STREAM] =
      RES :-
28          should_close(PATH, STREAM),
29          open_result(PATH, success(stream(STREAM))),
30          outfile(PATH).
31
32      io_error("Failed opening file: " + MSG) :-
33          open_result(_, error(MSG)).
34      io_error("Failed writing line " + IDX + ": " + MSG) :-
35          write_result(IDX, error(MSG)).
36      io_error("Failed closing file: " + MSG) :-
37          close_result(_, error(MSG)).
38
39      lines_written(CNT) :- CNT = #count{N, R :
      write_result(N, R)}, not io_error(_).
40 }

```

The action module proposed in Listing 6.7 differs from a "regular" module in the following ways:

- The module output definition (`success: lines_written/1 | error: io_error/1`) distinguishes between a success and an error case.
- Since we assume action result terms to always be unary function terms with symbol either *success* or *error*, the atoms of the given predicates would then be arguments of the respective (success or error) term in their termified form according to Definition 3.2.5.
- The above items imply that an action module result can only be success *or* error, but not both. In an actual implementation of this concept, ways of enforcing this rule would need to be addressed.



Listing 6.8 shows a prototypical example of how an action module could be used.

Listing 6.8: Prototypical usage of an action module providing an abstraction to write content to a file.

```

1 line(0, "lorem ipsum dolor").
2 line(1, "sit amet consectetur").
3 line(2, "adipiscing elit")
4
5 write_result(R) : @write_lines[LINES, "/tmp/out.txt"] = R
6               :- LINES = #list{line(I, L) : line(I, L)}.

```

While the approach to writing composite actions using a special form of modules sketched in Example 6.1.2 looks promising at first glance, we note that it also, potentially, projects away information about side-effects: Consider two distinct Frames  $F_1$  and  $F_2$ , where in  $F_1$  the `stream_write` action in module `write_lines` gives an error for line 1, and in  $F_2$ , the same action yields an error for line 2. In both frames, the result of the action rule in the top-level program is going to be an error, specifically `write_result(io_error("Failed writing line 1: error"))` for  $F_1$ , and `write_result(io_error("Failed writing line 2: error"))` (note that wrapping list terms mandated by module output translation were ignored since there is only one answer set holding one instance of `io_error` anyway). While in this case the line on which writing failed is made transparent through the error message, in general, it would be possible to project away the information how many lines were written, i.e. two frames might yield different outcomes for the actions inside the module while yielding the exact same result for the overall program. Potential implementations of this concept would have to make allowances for this.

## 6.2 Potential improvements to solving performance

The performance test results from Section 5.3.2 show a strong increase in runtime of the XML-3-Coloring application with increasing vertex count of the input graph. Intuitively, this seems easily explained - the XML-parsing module described in Section A.3 has a number of predicates whose (ground) instance count in an answer set is  $O(n^2)$  with regards to the number  $n$  of instances of the predicates they depend on. A typical example are rules like in Listing A.11 where, in order to find pairs of corresponding opening and closing XML tags, we construct a partial ordering of tokens, resulting in  $\frac{n^2-n}{2} = \binom{n}{2}$  atoms for the relation predicate. Similar constructs occur multiple times in the XML parsing module. While information about the position of tokens in the input text relative to each other is necessary for the program, it would not have to be kept in memory at all times - the program in the XML parsing module is stratified, and the output predicates which are actually reported back to the calling program all reside in the uppermost stratum.

Based on the observations above, we could reasonably adapt Alpha’s stratified evaluation algorithm (see Section 4.1.2) to discard predicates from working memory that are no longer needed for evaluation (i.e. all instances of all directly dependent predicates have been calculated) and are not part of a module’s output predicates, thereby reducing memory footprint and in turn reduce the amount of time spent on (fruitless) garbage collection cycles in the JVM running Alpha.

# Conclusion

The idea of this work was to extend traditional answer set programming with a means to apply "side-effects", i.e. influence the outside world, from a program, while still preserving fully declarative semantics.

To that end, we introduced the language extension *Evolog* in Chapter 3, which is a conservative extension to the ASP language that is understood by the Alpha ASP solver. Evolog adds a formal representation of actions, which is defined in Section 3.1, to ASP that is inspired by the Monad concept from functional programming in that it treats the state of the world on which side-effects are applied as input and output, respectively, of an action function. In Evolog, this is captured in the notion of a *Frame*, which describes the interactions of a program with the outside world by means of an interpretation function for action rules. In order to enable an implementation of the action semantics which allows for programs with actions to be run on a physical computer with reasonable results, we defined necessary restrictions on the structure of those programs. The most important such restriction is that, whenever an action rule fires, the resulting atom must be part of an answer set, thereby ensuring that for every effect applied to the outside world, there is a witnessing atom in an answer set. In our reference implementation of said action semantics, in order to get to a condition that can actually be statically verified by an interpreter, we narrowed this to only allow actions in stratified programs. While this may seem like a rather large limitation, it neatly lines up with the fact that - even though ASP has multiple-model-semantics - we only run a given program on one computer at a time, so attempting to apply contradicting actions from multiple models would not be practical.

The second part of the Evolog extension is the program modularization concept defined in Section 3.2. Modularization in Evolog treats modules which encapsulate sub-programs as a special kind of external atoms. By defining each answer set of a module program to constitute one ground substitution of the corresponding module atom, Evolog modules

also provide a means to use program parts with multiple models (and guesses through e.g. the guess-and-check pattern) alongside Actions.

In Chapter 4 we described our implementation of the Evolog extension, which is based on the lazy-grounding ASP solver Alpha, which we then tested by implementing a "real world" application, combining user interaction, file in- and output, parsing, as well as solving an NP-complete decision problem, in Chapter 5. We described a pure-Evolog implementation of an application that parses XML files representing graphs, on which we the search for three-colorings, and write the found colorings to an XML file. Chapter 5 also contains some short performance test results from a variation of the XML-based 3-coloring application.

Last but not least, Chapter 6 made an attempt to draw lessons for further improvement of the Evolog language extension as well as implementation performance from the code described in Chapter 5 - we identified code constructs that are often repeated and discussed potential optimizations. Furthermore, we also discussed solving performance for Evolog programs, and sketched a possible approach to reducing the memory footprint of an Evolog program.

## 7.1 Outlook

While the work described here delivers a possible start on action semantics and a compatible modularization concept for (lazy-grounding) ASP solvers, it does not go beyond a first step, and there are a number of topics worth exploring further.

An intuitive first step to further improving the Evolog extension would be actually implementing the language constructs sketched in Section 6.1, namely syntactic shorthands for working with lists as well as composite actions. Furthermore, the current implementation does not support nested module calls, i.e. a module program cannot use another module. Since there is no semantic reason prohibiting this, it would suggest itself as a major improvement in the area of developer comfort.

Apart from developing language features, implementing additional applications using actions could lead to further insights. The actions provided for the reference implementation shown here are very centered around file input and output. While means to write to and read from file descriptors already enable a vast number of different programs, further actions, for example to interact with network sockets, or even build graphical user interfaces, would greatly increase the usability of Evolog as a potential general purpose programming language.

# Additional Material

## A.1 Installing Alpha

**Using a pre-built binary file** The Alpha ASP solver is written in Java. In order to run it, a Java Runtime Environment (JRE) compatible with Java 11 (i.e. Version 11 or newer) is required. Alpha builds can be obtained from the Github page<sup>1</sup>.

The artifact required to run Alpha is the bundled jar file containing the commandline application, i.e. `alpha-cli-app-${version}-bundled.jar`.

In order to solve an ASP program stored in file `program.asp`, run the jar file as demonstrated in Listing A.1.

Listing A.1: Solving an ASP program using Alpha.

```
$>java -jar /path/to/alpha/
    alpha-cli-app-x.y.z-bundled.jar -i program.asp
```

In order to run the Alpha jar from any location in the file system without always having to enter the full path to the jar, one can create a launcher script for Alpha. This launcher script then needs to be placed in a directory that is included in the system's PATH environment variable (e.g. `/usr/local/bin` on most Linux systems). Listing A.2 shows a launcher script suitable for a Linux system (i.e. in the Bash scripting language) which assumes the Alpha jar for Alpha version 0.8.0 to be stored in `/usr/local/bin`.

Listing A.2: Bash launcher script for Alpha.

```
1 #!/bin/bash
2
3 java -jar /usr/local/bin/alpha-cli-app-0.8.0-bundled.jar $@
```

<sup>1</sup><https://github.com/alpha-asp/Alpha/releases>

Listing A.3 shows a Windows-compatible script which assumes that the Alpha jar is stored in `C:/Program Files/alpha-asp`.

Listing A.3: Windows-Batch launcher script for Alpha.

```
1 java -jar "C:\Program
   Files\alpha-asp\alpha-cli-app-0.8.0-SNAPSHOT-bundled.jar" %*
```

Using a central launcher script, Alpha can be called through the name of the launcher script from any working directory. This is demonstrated in Listing A.4, where `alpha-solver` is assumed as name of the launcher script.

Listing A.4: Solving an ASP program using Alpha through a launcher script.

```
$>alpha-solver -i program.asp
```

**Building Alpha from source code** As an alternative to using a pre-built binary package, Alpha can also be built from sources. For this, a Java Development Kit (JDK) compatible with Java 11 is required. In order to build Alpha, clone the git repository<sup>2</sup> and start the build through the Gradle Wrapper script as demonstrated in Listing ?? . The actual executable jar is built using the gradle task *bundledJar*, but to make sure all automated tests succeed as expected, is it recommended to run a full build (using task *build*) first. After the *bundledJar* task has completed, the executable file is located in the build directory of the `alpha-cli-app` module.

Listing A.5: Building Alpha from source code.

```
$>git clone https://github.com/alpha-asp/Alpha.git
$>cd Alpha
$>./gradlew build
$>./gradlew bundledJar
$>java -jar alpha-cli-app/build/libs/
      alpha-cli-app-0.8.0-bundled.jar -i program.asp
```

## A.2 Examples

**Example A.2.1** (Fibonacci-Numbers using external atoms). The following code snippet demonstrates how to run a program which includes user-supplied external atoms using Alpha. Since the Alpha Commandline-App currently does not support loading Atom Definitions from jar files, the solver is directly called from Java code in this example.

In this example, we use an external predicate definition `fibonacci_number/2` to efficiently calculate Fibonacci numbers. The actual ASP program generates all Fibonacci

---

<sup>2</sup><https://github.com/alpha-asp/Alpha.git>

numbers up to index 40 that are even. The ASP program is shown in Listing A.6, while Listing A.7 shows the Java implementation of the external predicate.

Listing A.6: ASP program to find even Fibonacci numbers.

```

1 %% Find even fibonacci numbers up to F(40).
2 fib(N, FN) :- &fibonacci_number[N](FN), N = 0..40.
3 even_fib(N, FN) :- fib(N, FN), FN \ 2 = 0.

```

The Java implementation of the `fibonacci_number/2` predicate makes use of *Binet's Formula*<sup>3</sup> to efficiently compute Fibonacci numbers using a closed form expression of the sequence.

Listing A.7: Fibonacci number computation in Java.

```

public class CustomExternals {

    private static final double GOLDEN_RATIO =
        1.618033988749894;
    private static final double SQRT_5 = Math.sqrt(5.0);

    /**
     * Calculates the n-th Fibonacci number using a
     * variant of Binet's Formula
     */
    @Predicate(name="fibonacci_number")
    public static Set<List<ConstantTerm<Integer>>>
        fibonacciNumber(int n) {
        return
            Set.of(List.of(Terms.newConstant(binetrRounding(n))));
    }

    public static int binetrRounding(int n) {
        return (int) Math.round(Math.pow(GOLDEN_RATIO,
            n) / SQRT_5);
    }

}

```

The Alpha solver is invoked from a simple Java method, which uses Alpha's API to first compile the external atom definition, and then run the solving process for the parsed ASP program. This is illustrated in Listing A.2.1

```

public class CustomExternalsApp {

```

<sup>3</sup>[https://en.wikipedia.org/wiki/Fibonacci\\_sequence#Binet's\\_formula](https://en.wikipedia.org/wiki/Fibonacci_sequence#Binet's_formula)

```
public static void main(String[] args) throws
    IOException {
    Alpha alpha = AlphaFactory.newAlpha();
    Map<String, PredicateInterpretation>
        customExternals =
            Externals.scan(CustomExternals.class);
    String aspCode =
        Files.readString(
            Paths.get("src/main/resources/
                      customExternals.asp"));
    InputProgram program =
        alpha.readProgramString(aspCode,
            customExternals);
    alpha.solve(program).forEach(as -> {
        System.out.println("Answer set:\n" +
            as);
    });
}
```

**Example A.2.2** (Module Parsing). Listing A.8 shows the actual ANTLR-grammar used in Alpha to parse module definitions. Parsed Module Definitions are stored in instances of the Module type. Listing A.9 shows the corresponding Java interface definition.

Listing A.8: ANTLR grammar for module definitions

```
1 directive_module:
2     SHARP DIRECTIVE_MODULE id
3         PAREN_OPEN module_signature PAREN_CLOSE
4         CURLY_OPEN statements CURLY_CLOSE;
5
6 module_signature :
7     predicate_spec ARROW CURLY_OPEN
8         ('*' | predicate_specs) CURLY_CLOSE;
9
10 predicate_specs:
11     predicate_spec (COMMA predicate_specs)?;
12
13 predicate_spec: id '/' NUMBER;
```

Note how the `module_signature` rule permits the shorthand `*` instead of an output predicate list. This is used as a shorthand to express the list of all predicates derived by any rule in the module (i.e. emit full, unfiltered answer sets).



Listing A.9: Java interface specifying Alpha's internal representation of a module definition.

```
public interface Module {

    String getName();

    Predicate getInputSpec();

    Set<Predicate> getOutputSpec();

    InputProgram getImplementation();

}
```

### A.3 An XML-DOM parser in ASP

The module `xml_dom` which is used in Listing 5.4 implements a generic XML-parser that generates a DOM-representation based on external atoms for regex evaluation.

Input is encoded as a list of strings, each one representing a line of input. At first, all input is tokenized in order to find opening and closing XML tags. Listing A.10 shows the Evolog code used for tokenization in the `xml_dom` module.

Listing A.10: Tokenizing XML input.

```
1 token_regex(tag_open, "(<(\w+) ( \w+=\\\" \w+\\\" ) *>) ").
2 token_regex(tag_close, "</\w+>").
3 opening_tag_name_regex("<(\w+) ( \w+=\\\" \w+\\\" ) *>").
4 closing_tag_name_regex("</(\w+)>").
5 attribute_regex("( \w+=\\\" \w+\\\" )").
6 attribute_name_regex("( \w+)=\\\" \w+\\\" ").
7 attribute_value_regex(" \w+=\\\" (\w+)\\\" ").
8
9 % Unwrap lines
10 line_element(E, TAIL) :- input_lines(lst(E, TAIL)).
11 line_element(E, TAIL) :- line_element(_, lst(E, TAIL)).
12 line(LINE_NO, LINE) :- line_element(line(LINE_NO, LINE), _).
13
14 % Match basic tokens
15 token(t(TOK, VALUE), LINE_NO, FROM, TO) :-
16     line(LINE_NO, LINE), token_regex(TOK, REGEX),
17     &regex_matches[REGEX, LINE](VALUE, FROM, TO).
```

Regular expressions to match opening and closing XML tags are represented as facts the predicate `token_regex/2`. The external atom `regex_matches/5` is used to determine whether a line matches a token regex, as well as the respective parts of the string where the match occurs.

Once all tokens denoting opening and closing tags - up until that point ignoring actual tag names and relative positions of tags to each other - the next step is extracting actual tag names and pairs of opening and closing tags of the same name. This part of the parsing process is demonstrated in Listing A.11. First, a relative ordering of tokens is established using the `token_before/8` predicate, which is derived by the rules on lines 2 through 11. Furthermore, we extract the name of each tag by using a variant of the tag open/close regexes with capture groups such that the first group is only the actual name of the tag, see lines 14 to 29. A matching pair of opening and closing tags in a well-formed XML document is characterized as an opening and a closing tag of the same name, with no opening tag of that name in between. In order to find these pairs, we first derive all matching pairs which do have another opening tag between them as instances of predicate `tag_opening_between/7`. The actual tag pairs making up well-formed XML elements are the derived using the rule on line 54 as those pairs of matching tags for which no corresponding instance of `tag_opening_between/7` exists.

Listing A.11: Finding pairs of matching opening and closing XML tags.

```
1 % Establish a token ordering
2 token_before(TOK1, TOK1_LINE_NO, TOK1_FROM, TOK1_TO,
3             TOK2, TOK2_LINE_NO, TOK2_FROM, TOK2_TO) :-
4     token(TOK1, TOK1_LINE_NO, TOK1_FROM, TOK1_TO),
5     token(TOK2, TOK2_LINE_NO, TOK2_FROM, TOK2_TO),
6     TOK1_LINE_NO = TOK2_LINE_NO, TOK1_TO < TOK2_FROM.
7 token_before(TOK1, TOK1_LINE_NO, TOK1_FROM, TOK1_TO,
8             TOK2, TOK2_LINE_NO, TOK2_FROM, TOK2_TO) :-
9     token(TOK1, TOK1_LINE_NO, TOK1_FROM, TOK1_TO),
10    token(TOK2, TOK2_LINE_NO, TOK2_FROM, TOK2_TO),
11    TOK1_LINE_NO < TOK2_LINE_NO.
12
13 % Extract tag names
14 tag_opening(TAG_NAME, t(tag_open, TOK_VALUE),
15            LINE_NO, TAG_FROM, TAG_TO) :-
16    token(
17        t(tag_open, TOK_VALUE),
18        LINE_NO, TAG_FROM, TAG_TO),
19    &regex_matches
20    [TAG_NAME_REGEX, TOK_VALUE](TAG_NAME, _, _),
21    opening_tag_name_regex(TAG_NAME_REGEX).
22 tag_closing(TAG_NAME, t(tag_close, TOK_VALUE),
23            LINE_NO, TAG_FROM, TAG_TO) :-
```

```
24     token(  
25         t(tag_close, TOK_VALUE),  
26         LINE_NO, TAG_FROM, TAG_TO),  
27     &regex_matches  
28     [TAG_NAME_REGEX, TOK_VALUE](TAG_NAME, _, _),  
29     closing_tag_name_regex(TAG_NAME_REGEX).  
30  
31 % We have an opening/closing tag pair if tag name is the same,  
32 % opening tag is before closing tag,  
33 % and no opening tag of same name between.  
34 tag_opening_between(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,  
35     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-  
36     tag_opening(TAG_NAME, OPEN_TOK,  
37     OPEN_LINE_NO, OPEN_FROM, OPEN_TO),  
38     tag_closing(TAG_NAME, CLOSE_TOK,  
39     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
40     token(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM, OPEN_TO),  
41     token(CLOSE_TOK, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
42     token(INTM_OPEN_TOK, INTM_OPEN_LINE_NO,  
43     INTM_OPEN_FROM, INTM_OPEN_TO),  
44     tag_opening(TAG_NAME, INTM_OPEN_TOK,  
45     INTM_OPEN_LINE_NO, INTM_OPEN_FROM, INTM_OPEN_TO),  
46     token_before(  
47     OPEN_TOK, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,  
48     INTM_OPEN_TOK, INTM_OPEN_LINE_NO,  
49     INTM_OPEN_FROM, INTM_OPEN_TO),  
50     token_before(  
51     INTM_OPEN_TOK, INTM_OPEN_LINE_NO, INTM_OPEN_FROM,  
52     INTM_OPEN_TO, CLOSE_TOK, CLOSE_LINE_NO,  
53     CLOSE_FROM, CLOSE_TO).  
54 element(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,  
55     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-  
56     tag_opening(TAG_NAME, OPEN_TOK,  
57     OPEN_LINE_NO, OPEN_FROM, OPEN_TO),  
58     tag_closing(TAG_NAME, CLOSE_TOK,  
59     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
60     token(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM, OPEN_TO),  
61     token(CLOSE_TOK, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
62     token_before(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM,  
63         OPEN_TO,  
64         CLOSE_TOK, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
65     not tag_opening_between(  
66     TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
```

```
66 | CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) .
```

Having parsed all XML elements, the next step is to determine which of these are terminal elements in the sense that they only contain text, as opposed to elements which consist of further XML elements. This is achieved using the code form Listing A.12.

Listing A.12: Parsing details of XML elements

```
1 % Any string between two tags with no other opening
2 % or closing tags between is content
3 any_tag_opening_between(TAG_NAME, OPEN_LINE_NO, OPEN_FROM,
4   OPEN_TO, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-
5     element(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
6       CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),
7     token(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM, OPEN_TO),
8     token(CLOSE_TOK, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),
9     token_before(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM,
10      OPEN_TO, t(tag_open, INTM_TOK), INTM_LINE_NO,
11      INTM_FROM, INTM_TO),
12     token_before(t(tag_open, INTM_TOK), INTM_LINE_NO,
13      INTM_FROM, INTM_TO, CLOSE_TOK, CLOSE_LINE_NO,
14      CLOSE_FROM, CLOSE_TO) .
15 any_tag_closing_between(TAG_NAME, OPEN_LINE_NO, OPEN_FROM,
16   OPEN_TO, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-
17     element(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
18       CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),
19     token(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM, OPEN_TO),
20     token(CLOSE_TOK, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),
21     token_before(OPEN_TOK, OPEN_LINE_NO, OPEN_FROM,
22      OPEN_TO, t(tag_close, INTM_TOK), INTM_LINE_NO,
23      INTM_FROM, INTM_TO),
24     token_before(t(tag_close, INTM_TOK), INTM_LINE_NO,
25      INTM_FROM, INTM_TO, CLOSE_TOK, CLOSE_LINE_NO,
26      CLOSE_FROM, CLOSE_TO) .
27 any_tag_between(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
28   CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-
29     any_tag_opening_between(TAG_NAME, OPEN_LINE_NO,
30      OPEN_FROM, OPEN_TO, CLOSE_LINE_NO, CLOSE_FROM,
31      CLOSE_TO),
32     any_tag_closing_between(TAG_NAME, OPEN_LINE_NO,
33      OPEN_FROM, OPEN_TO, CLOSE_LINE_NO, CLOSE_FROM,
34      CLOSE_TO) .
35 terminal(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
36   CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO) :-
```

```

19     element(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,
20             CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),
21     not any_tag_between(TAG_NAME, OPEN_LINE_NO, OPEN_FROM,
22                         OPEN_TO, CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO).
23 % Extract content between tags (currently assumes start and
24   end tags to be on the same line)
25 element_text(ELEMENT_NAME, FROM_POS, TO_POS, TEXT) :-
26     terminal(ELEMENT_NAME, LINE_NO, OPEN_FROM, OPEN_TO,
27             LINE_NO, CLOSE_FROM, CLOSE_TO),
28     line(LINE_NO, FULL_LINE),
29     &string_substring[FULL_LINE, OPEN_TO,
30                     CLOSE_FROM](TEXT),
31     FROM_POS = pos(LINE_NO, OPEN_FROM),
32     TO_POS = pos(LINE_NO, CLOSE_TO).

```

In Listing A.12, the rule on line 18 derives whether an XML element is a *terminal* element in that it does not enclose any additional elements other than plain text. In order to determine this, we check if there are any opening or closing tags between the opening and closing tags of the element in question, see lines 3 through 15. For all terminal elements, we parse the text content of each individual element by taking a substring of the line containing the opening and closing tags ranging from after the last character of the opening tag to before the first character of the closing tag. (Note that this implementation assumes that content within a tag does not contain newline characters in order to keep the code a bit shorter.)

Every XML tag can also have any number of attributes. The parsing logic for attributes is shown in Listing A.13. First, attributes are detected by checking every string that has been matched to be an opening XML tag against regular expressions matching attribute name and values, see the rule on line 2. Attributes derived using this approach are then associated with the corresponding element/7 instance through the predicate `element_attribute/8`.

Listing A.13: Parsing XML attributes.

```

1 %% Extract attributes
2 tag_opening_attribute(TAG_NAME, t(tag_open, TOK_VALUE),
3                       LINE_NO, FROM, TO, attribute(NAME, VALUE)) :-
4     tag_opening(TAG_NAME, t(tag_open, TOK_VALUE), LINE_NO,
5                 FROM, TO),
6     &regex_matches[ATTRIBUTE_REGEX,
7                 TOK_VALUE](ATTRIBUTE_STR, _, _),
8     attribute_regex(ATTRIBUTE_REGEX),
9     &regex_matches[ATTRIBUTE_VALUE_REGEX,
10                 ATTRIBUTE_STR](VALUE, _, _),
11     attribute_value_regex(ATTRIBUTE_VALUE_REGEX),

```

```
8      &regex_matches[ATTRIBUTE_NAME_REGEX,  
9      ATTRIBUTE_STR](NAME, _, _),  
10     attribute_name_regex(ATTRIBUTE_NAME_REGEX) .  
11 element_attribute(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,  
12     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO, attribute(NAME,  
13     VALUE)) :-  
14     element(TAG_NAME, OPEN_LINE_NO, OPEN_FROM, OPEN_TO,  
15     CLOSE_LINE_NO, CLOSE_FROM, CLOSE_TO),  
16     tag_opening_attribute(TAG_NAME, t(tag_open, _),  
17     OPEN_LINE_NO, OPEN_FROM, OPEN_TO, attribute(NAME,  
18     VALUE)) .
```

The final step in actual parsing of the XML content is to derive the complete hierarchy of all XML tags, i.e. determine which tags are direct children of another. This is done using the code from Listing A.14. First, we establish elements that enclose other elements (i.e. opening and closing tag of the enclosed element between opening and closing tags of the enclosing element) using the rule on line 4. An element  $e_1$  is not the parent element of an element  $e_2$  enclosed by  $e_1$  if there is a third element  $e_i$  which is enclosed by  $e_1$ , but also encloses  $e_2$ . If, for any element enclosing another element such an intermediate element exists, the rule from line 18 derives this as an instance of `element_not_parent/6`. An element  $e_1$  is the parent element of  $e_2$  if it encloses  $e_2$  and is not known to be not the parent. Element pairs fulfilling this definition are derived by the rule on line 22 and expressed as instances of `element_parent/6`.

Listing A.14: Parsing XML attributes.

```
1 % Derive parent elements  
2 % An element encloses another if the opening and closing tag  
3   of the enclosed  
4 % element lie fully between opening and closing tags of the  
5   enclosing element  
6 element_encloses(ENC, ENC_POS_FROM, ENC_POS_TO, LOWER,  
7   LOWER_POS_FROM, LOWER_POS_TO) :-  
8   element(LOWER, LOWER_OPEN_LINE_NO, LOWER_OPEN_FROM,  
9   LOWER_OPEN_TO, LOWER_CLOSE_LINE_NO,  
10  LOWER_CLOSE_FROM, LOWER_CLOSE_TO),  
11  element(ENC, ENC_OPEN_LINE_NO, ENC_OPEN_FROM,  
12  ENC_OPEN_TO, ENC_CLOSE_LINE_NO, ENC_CLOSE_FROM,  
13  ENC_CLOSE_TO),  
14  token(LOWER_OPEN_TOK, LOWER_OPEN_LINE_NO,  
15  LOWER_OPEN_FROM, LOWER_OPEN_TO),  
16  token(LOWER_CLOSE_TOK, LOWER_CLOSE_LINE_NO,  
17  LOWER_CLOSE_FROM, LOWER_CLOSE_TO),
```

```

9      token(ENC_OPEN_TOK, ENC_OPEN_LINE_NO, ENC_OPEN_FROM,
10         ENC_OPEN_TO),
11      token(ENC_CLOSE_TOK, ENC_CLOSE_LINE_NO,
12         ENC_CLOSE_FROM, ENC_CLOSE_TO),
13      token_before(ENC_OPEN_TOK, ENC_OPEN_LINE_NO,
14         ENC_OPEN_FROM, ENC_OPEN_TO, LOWER_OPEN_TOK,
15         LOWER_OPEN_LINE_NO, LOWER_OPEN_FROM, LOWER_OPEN_TO),
16      token_before(LOWER_CLOSE_TOK, LOWER_CLOSE_LINE_NO,
17         LOWER_CLOSE_FROM, LOWER_CLOSE_TO, ENC_CLOSE_TOK,
18         ENC_CLOSE_LINE_NO, ENC_CLOSE_FROM, ENC_CLOSE_TO),
19      ENC_POS_FROM = pos(ENC_OPEN_LINE_NO, ENC_OPEN_FROM),
20      ENC_POS_TO = pos(ENC_CLOSE_LINE_NO, ENC_CLOSE_TO),
21      LOWER_POS_FROM = pos(LOWER_OPEN_LINE_NO,
22         LOWER_OPEN_FROM),
23      LOWER_POS_TO = pos(LOWER_CLOSE_LINE_NO,
24         LOWER_CLOSE_TO).
25
26 % An element E1 is parent of element E2 if E1 encloses E2 and
27   there is no other element that encloses E2 and is enclosed
28   by E1.
29
30 element_not_parent(E, E_POS_FROM, E_POS_TO, P, P_POS_FROM,
31   P_POS_TO) :-
32     element_encloses(P, P_POS_FROM, P_POS_TO, E,
33       E_POS_FROM, E_POS_TO),
34     element_encloses(P, P_POS_FROM, P_POS_TO, I,
35       I_POS_FROM, I_POS_TO),
36     element_encloses(I, I_POS_FROM, I_POS_TO, E,
37       E_POS_FROM, E_POS_TO).
38
39 element_parent(E, E_POS_FROM, E_POS_TO, P, P_POS_FROM,
40   P_POS_TO) :-
41     element_encloses(P, P_POS_FROM, P_POS_TO, E,
42       E_POS_FROM, E_POS_TO),
43     not element_not_parent(E, E_POS_FROM, E_POS_TO, P,
44       P_POS_FROM, P_POS_TO).

```

Last but not least, the `xml_dom` module has a number of rules to project away positional information of tags which are not deemed to be of interest to the calling component. Listing A.15 shows all rules used to achieve this. First, each parsed XML element is assigned a unique identifier represented using the predicate `dom_element_id/2` by the rule on line 1. Identifier terms are obtained from the *enumeration literal* (see Section 4.1.1) `id_enum/3`. Based on the thereby established relation, versions of the predicates `element/7`, `element_parent/6`, `element_text/4`, `element_attribute/8` in which the individual terms making up the respective element's position within the document are replaced with the corresponding identifier.

Listing A.15: Deriving simplified representations of parsed XML content.

```
1 dom_element_id(e(NAME, POS_FROM, POS_TO), ID) :-
2     element(NAME, START_LINE_NO, START_FROM, START_TO,
3         END_LINE_NO, END_FROM, END_TO),
4     POS_FROM = pos(START_LINE_NO, START_FROM),
5     POS_TO = pos(END_LINE_NO, END_TO),
6     id_enum(element_ids, e(NAME, POS_FROM, POS_TO), ID).
7
8 dom_element(NAME, ID) :-
9     dom_element_id(e(NAME, _, _), ID).
10
11 dom_element_child(NAME, ID, CHILD_ID) :-
12     dom_element(NAME, ID),
13     dom_element_id(e(NAME, POS_FROM, POS_TO), ID),
14     dom_element_id(e(CHILD_NAME, CHILD_POS_FROM,
15         CHILD_POS_TO), CHILD_ID),
16     element_parent(CHILD_NAME, CHILD_POS_FROM,
17         CHILD_POS_TO, NAME, POS_FROM, POS_TO).
18
19 dom_element_text(NAME, ID, TEXT) :-
20     dom_element(NAME, ID),
21     dom_element_id(e(NAME, POS_FROM, POS_TO), ID),
22     element_text(NAME, POS_FROM, POS_TO, TEXT).
23
24 dom_element_attribute(NAME, ID, ATTR_NAME, ATTR_VALUE) :-
25     dom_element(NAME, ID),
26     dom_element_id(e(NAME, pos(START_LINE_NO, START_FROM),
27         pos(END_LINE_NO, END_TO)), ID),
28     element_attribute(NAME, START_LINE_NO, START_FROM, _,
29         END_LINE_NO, _, END_TO, attribute(ATTR_NAME,
30         ATTR_VALUE)).
```

## A.4 Serializing graph-colorings to XML

Graph colorings calculated in the application from Chapter 5 are converted to XML strings using the rules shown in Listing A.16

Listing A.16: Generating XML strings for graph colorings.

```
1 #module threecol_to_xml (coloring/1 => {coloring_xmlstr/1}) {
2     coloring_entry(vertex(ID, COL), TAIL) :-
3         coloring(lst(col(ID, COL), TAIL)).
```



```

3      coloring_entry(vertex(ID, COL), TAIL) :- coloring_entry(_,
4          lst(col(ID, COL), TAIL)).
5      coloring_entry_pred(ID, ID_PRED) :-
6          coloring_entry(vertex(ID, _), lst(col(ID_PRED, _), _)).
7      coloring_entry_has_pred(ID) :- coloring_entry_pred(ID, _).
8      coloring_entry_is_pred(ID) :- coloring_entry_pred(_, ID).
9      coloring_entry_first(ID) :- coloring_entry(vertex(ID, _),
10          _, not coloring_entry_has_pred(ID)).
11      coloring_entry_last(ID) :- coloring_entry(vertex(ID, _),
12          _, not coloring_entry_is_pred(ID)).
13
14      coloring_tag_open("<coloring>").
15      coloring_tag_close("</coloring>").
16
17      coloring_part(ID, STR) :-
18          coloring_entry(vertex(ID, COL), _),
19          &stdlib_string_concat["<vertex><id>", ID] (S1),
20          &stdlib_string_concat[S1, "</id><color>"] (S2),
21          &stdlib_string_concat[S2, COL] (S3),
22          &stdlib_string_concat[S3, "</color></vertex>"] (STR).
23
24      coloring_xmlstr_upto(STR, ID) :-
25          coloring_entry_first(ID),
26          coloring_part(ID, FIRST_PARTSTR),
27          coloring_tag_open(OPENSTR),
28          &stdlib_string_concat[OPENSTR, FIRST_PARTSTR] (STR).
29
30      coloring_xmlstr_upto(STR, ID) :-
31          coloring_entry_pred(ID, ID_PRED),
32          coloring_xmlstr_upto(PREV_STR, ID_PRED),
33          coloring_part(ID, PARTSTR),
34          &stdlib_string_concat[PREV_STR, PARTSTR] (STR).
35
36      coloring_xmlstr(STR) :-
37          coloring_entry_last(ID),
38          coloring_xmlstr_upto(PREV_STR, ID),
39          coloring_tag_close(CLOSESTR),
40          &stdlib_string_concat[PREV_STR, CLOSESTR] (STR).
41  }

```



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# Acronyms

**ASP** Answer Set Programming. 1–6, 8, 9, 11, 13, 30, 38, 41

**CBP** Common Base Program. 21, 22, 47, 48, 50, 111

**CDNL** Conflict-driven Nogood Learning. 21, 41, 47

**DOM** Document Object Model. 69, 76

**DTD** Document Type Definition. 68



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