Pathway Specification and Comparative Queries: A High Level Language with Petri Net Semantics

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Abstract

Understanding biological pathways is an important activity in the biological domain for drug development. Due to the parallelism and complexity inherent in pathways, computer models that can answer queries about pathways are needed. A researcher may ask 'what-if' questions comparing alternate scenarios, that require deeper understanding of the underlying model. In this paper, we present overview of such a system we developed and an English-like high level language to express pathways and queries. Our language is inspired by high level action and query languages and it uses Petri Net execution semantics.

Introduction

Biological pathways are highly interconnected networks of biochemical processes. These processes execute autonomously in parallel, driven by natural constraints of ingredient supply and demand, i.e., a process can execute as soon as its preconditions are satisfied. For example, a metabolic reaction can occur as soon as sufficient quantities of its ingredients become available. Many processes are also governed by additional preconditions, including availability of additional substances that are not ingredients, substance gradients, inhibition, and stimulation. Collectively, these preconditions regulate the reactions through feedback loops and feed-forwards in the network. Reactions execute at different speeds, generating products on completion, which become ingredients for the down-stream reactions. The state of a pathway is defined by the available substance quantities, and the chain of reactions between a starting state and an ending state of a pathway define a trajectory of pathway's state evolution. The interplay between limited quantity of ingredients and other preconditions can present multiple choices for alternate trajectory evolutions at each pathway state.

Understanding these pathways is of fundamental importance in biological research for disease diagnosis and drug development. However, the aforementioned complexities make it difficult for one person to retain all aspects of the pathway. As a result, computer based systems are needed to

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represent these pathways, allowing biologists to pose queries against them. An important class of questions in this regard are the so called 'what-if' questions, which compare alternate scenarios of a pathway. We find such questions in college level books that a professor may ask his students to gauge their understanding of a pathway. For example the following question from (Reece et al. 2010) appeared in a recent knowledge representation and reasoning challenge ¹:

Question 1. "At one point in the process of glycolysis, both DHAP and G3P are produced. Isomerase catalyzes the reversible conversion between these two isomers. The conversion of DHAP to G3P never reaches equilibrium and G3P is used in the next step of glycolysis. What would happen to the rate of glycolysis if DHAP were removed from the process of glycolysis as quickly as it was produced?"

Answering such questions require simulating different scenarios and reasoning with the results. For example, question 1 asks for comparison of the *rate of glycolysis* between the nominal pathway and an alternate pathway in which *DHAP is removed as quickly as it is produced.*

In this paper we describe an English-like high level language to express pathways and queries. We highlight important features of its syntax and semantics and give an overview of an implementation that understands this language and answer questions about them. Our language is inspired by high level action and query languages such as (Gelfond and Lifschitz 1993; Giunchiglia and Lifschitz 1998; Lee, Lifschitz, and Yang 2013). However, compared to most existing action languages, which describe transition systems (Gelfond and Lifschitz 1998) ², our language describes trajectories. Our language is geared towards modeling natural systems, in which actions occur autonomously (Reiter 1996) when their pre-conditions are satisfied; and substance quantities do not become negative. Compared to most other action languages, substance quantities produced and consumed are additive ³. Our system also supports a richer query component, which is missing from most query languages that accompany action languages.

¹https://sites.google.com/site/2nddeepkrchallenge/

²Some languages like C+ (Giunchiglia et al. 2004) allow autonomous actions, but their query languages lack expressiveness.

 $^{^3}$ Although some languages like C+ have been extended to allow additive fluents (Lee and Lifschitz 2003).

Some aspects of our system are similar to (Baral et al. 2004), but their work is limited to signaling pathways, does not allow numeric quantities or has provision of loops inherent in biological pathways.

We use Petri Nets (Peterson 1977) for representing pathways as they mimic biological pathway diagrams and can represent parallel systems. For ease of reasoning and extensibility, we use Answer Set Programming (ASP) (Baral 2003) for simulating the pathway using the approach by (Anwar, Baral, and Inoue 2013a; 2013b).

The rest of the paper is organized as follows: we present important aspects of our high-level language syntax and semantics. Then we illustrate the use of our language with an example. After that, we briefly describe implementation of our system that understands this language and conclude with main contributions.

Description of our system

We present the key elements and discriminating factors of our high level language below.

Pathway Specification Language

The alphabet of pathway specification language \mathcal{P} consists of disjoint nonempty domain-dependent sets A, F, representing actions, and fluents, respectively; a fixed set S of firing styles; a fixed set K of keywords as syntactic sugar (shown in bold face); a fixed set of punctuations $\{`,'\}$; and a fixed set of special constants $\{'1', '*, 'max'\}$; and integers.

Each fluent $f \in F$ has a domain dom(f) which is either integer or binary and specifies the values f can take. A *state* s is an interpretation of F that maps fluents to their values. We write s(f) = v to represent "f has the value v in state s". States are indexed, such that consecutive states s_i and s_{i+1} represent an evolution over one time step from i to i+1 due to firing of an action set T_i in s_i . We illustrate the role of various constructs using a hypothetical example from the biological domain s_i :

domain of
$$sug$$
 is $integer$, fac is $integer$, $acoa$ is $integer$, $h2o$ is $integer$ (1)

 box may execute causing fac change value by -1,

 $acoa$ change value by +1 (2)

if $h2o$ has value 1 or higher (3)

inhibit box if sug has value 1 or higher (4)

initially sug has value 3, fac has value 4,

 $acoa$ has value 0, $h2o$ has value 0 (5)

Above query is about a process called beta-oxidation represented by 'box', the effect of which is captured in lines (2)-(3). Line (1) declares fluents for the substances used in the pathway and their domain, i.e. sugar ('sug'), fatty-acids ('fac'), acetyl-CoA ('acoa'), and water ('h2o') are represented by numeric fluents. Line (2) describes the effect of beta oxidation on its inputs and outputs, i.e. when beta-oxidation occurs, it consumes 1 unit of fatty-acids and produces 1 unit of acetyl-CoA. It implicitly defines a precondition that beta-oxidation cannot occur unless there is at least

1 unit of fatty-acids available. Line (3) describes an explicit precondition (or *guard*) of beta oxidation, i.e. it cannot occur unless there is at least 1 unit of water available. The water is, however, not consumed during beta oxidation. Line (4) explicitly inhibits beta-oxidation when there is any sugar available; and line (5) sets up the initial conditions of the pathway, s.t. initial quantities of sugar, fatty-acids, acetyl-CoA, and water are 3,4,0,0, respectively.

Now we introduce various statements and clauses, give their intuitive definitions, and show how they are combined to construct a pathway specification (or domain description). In the following description, f is a fluent, a is an action, $w \in \mathbb{N}^+ \cup \{0\}, d \in \mathbb{N}^+, S \in \{1, *, max\}, e \in (\mathbb{N} \setminus \{0\}) \cup \{*\}$ for integer or $e \in \{1, -1, *\}$ for binary fluents.

An effect clause has the form:

$$f$$
 change value by e (6)

A *guard_cond* clause can take the forms:

$$f$$
 has value w or higher (7)

$$f$$
 has value lower than w (8)

The domain description contains statements of the forms:

$$\begin{array}{ll} \textbf{domain of } f \textbf{ is '} integer' | \text{'} binary' & (9) \\ a \textbf{ may execute causing } effect_1, \dots, effect_m \\ \textbf{ if } guard_cond_1, \dots, guard_cond_n & (10) \\ \textbf{inhibit } a \textbf{ if } guard_cond_1, \dots, guard_cond_n & (11) \end{array}$$

initially
$$f$$
 has value w (12)

$$a$$
 executes in d time units (13)

firing style
$$S$$
 (14)

where m > 0 and $n \ge 0$.

A fluent domain declaration statement (9) declares the values a fluent f can take. While binary domain is commonly used for representing substances in a signaling pathway, metabolic pathways use positive numeric values. Since the domain is for a physical entity, we disallow fluents with negative values. A may-execute statement (10) captures the pre-conditions of an action a and its impact. A single mayexecute statement must not have effect, effect, with $e_i < 0$ and $e_i < 0$; or $e_i > 0$ and $e_i > 0$ for the same fluent. An *inhibit* statement (11) captures explicit inhibition conditions for an action a. An *initial condition* statement (12) captures the initial state of pathway (e.g. as substance distribution). A *duration* statement (13) represents the duration d an action a takes to execute. A firing style statement (14) specifies how many actions may execute in parallel, where, \hat{S} is either "1", "*", or "max" for serial execution, arbitrary amount of parallelism, and maximum parallelism. Actions execute automatically when fireable, subject to the available fluent quantities.

Definition 1 (Pathway Specification). A pathway specification *is composed of one or more* domain, may-execute, inhibit, initially, duration *statements, and one* firing style *statement.*

A pathway specification is consistent if (i) there is at most one firing style statement (ii) at most one duration statement for an action a; (iii) the $guard_cond_1, \ldots, guard_cond_n$ from a may-execute are disjoint from any other may-execute

⁴Multiple **domain** and **initially** statements have been written in a comma-separated compact form.

for the same action ⁵; (iv) domain of fluents, effects, conditions and numeric values are consistent, i.e., effects and conditions on binary fluents must be binary; and (v) the pathway specification does not cause it to violate fluent domains by producing non-binary values for binary fluents.

When missing, duration of an action is assumed to be 1; and initial fluent quantity for a fluent is assumed to be 0.

Intuitively, a pathway specification $\mathbf D$ represents a set of trajectories of the form: $\sigma=s_0,T_0,s_1,\ldots,s_{k-1},T_{k-1},s_k$. Each trajectory encodes an evolution of the pathway. Starting from an *initial state* s_0 , each s_i,s_{i+1} pair represents the state evolution in one time step due to execution of the action set T_i in state s_i producing s_{i+1} . An action set T_i is only executable in state s_i , if the total decrease of fluent values due to $e_i<0$ and $e_i=*$ will not result in any of the fluents becoming negative. Changes to fluents due to $e_i>0$ for the action set T_i occur over subsequent time-steps depending upon the durations of actions involved. Thus, the state $s_i(f_i)$ is the sum of $e_i>0$ for actions of duration d that occurred d time steps before (current time step) i, i.e. $a\in T_{i-d}$.

Query Specification Language

The alphabet of query language \mathcal{Q} consists of the same sets A, F from \mathcal{P} representing actions, and fluents, respectively; a fixed set of reserved keywords K shown in bold in syntax below; a fixed set $\{`:`,`;`,`,`,`,''\}$ of punctuations; a fixed set of $\{<,>,=\}$ of directions; and constants. Our query language asks questions about biological entities and processes in a biological pathway described by a pathway specification (or domain description). A query statement is composed of a query description (the quantity, or property being sought by the question), interventions (changes to the pathway), observations (about states and actions of the pathway), and initial setup conditions. For example, the following query indirectly determines the direction of change in the rate of glycolysis by comparing the rate of production of 'bpg13' w.r.t. the glycolysis pathway given in (Reece et al. 2010, Figure 9.9); and corresponds to question (1).

 ${\bf direction\ of\ change\ in}\ average$

rate of production of bpg13 is d

when observed between time step 0 and time step k; comparing nominal with modified pathway obtained due to interventions:

remove dhap as soon as produced;

using initial setup:

continuously supply
$$f16bp$$
 in quantity 1; (15)

Intuitively, a query statement is evaluated against the trajectories of a pathway (or domain description). The pathway is first modified by first applying the initial setup conditions, and the interventions. The modified pathway is them simulated and its trajectories are filtered to retain only those which satisfy the observations specified in the query statement. Next we define the syntax of the query language and

its elements 6 , give their intuitive meaning, and how these components fit together to form a query statement. In the following description, a's are actions, f's are fluents, n's are numbers, q's are positive integer numbers, d is one of the directions from $\{<,>,=\}$.

C / / J	
$\langle point \rangle ::= \mathbf{time\ step}\ ts$	(16)
$\langle interval \rangle ::= \langle point \rangle \text{ and } \langle point \rangle$	(17)
$\langle aggop \rangle ::= \mathbf{minimum} \mid \mathbf{maximum} \mid \mathbf{average}$	(18)
$\langle quant\ intf \rangle \ ::= \ {f rate}\ {f of}\ {f production}\ {f of}\ f\ {f is}\ n$	(19)
rate of firing of a is n	(20)
$\langle quant \ ptf \rangle \ ::= \ \mathbf{value \ of} \ f \ \mathbf{is} \ \mathbf{higher \ than} \ n$	(21)
value of f is n	(22)
$\langle qual\ intf \rangle \ ::= \ f \ {f is \ accumulating}$	(23)
$\langle qual\; ptf \rangle \; ::= \; a \; \mathbf{occurs}$	(24)
a does not occur	(25)
$ a_1 $ switches to a_2	(26)
$\langle quant \ agg \ intf \rangle \ ::= \ \langle aggop \rangle \ {f rate} \ {f of} \ {f firing} \ {f of} \ a \ {f is} \ n$	(27)
$ \langle aggop \rangle$ rate of production of f is n	(28)
$\langle quant \ agg \ ptf \rangle ::= \langle aggop \rangle \ \mathbf{value \ of} \ f \ \mathbf{is} \ n$	(29)
$\langle quant \ cagg \ intf \rangle ::= \ \mathbf{direction} \ \mathbf{of} \ \mathbf{change} \ \mathbf{in}$	
$\langle aggop \rangle$ rate of production of f is d	(30)
direction of change in	
$\langle aggop \rangle$ rate of firing of a is d	(31)
$\langle quant \ cagg \ ptf \rangle ::= $ direction of change in	
$\langle aggop \rangle$ value of f is d	(32)
$\langle simp\ intf \rangle \ ::= \ \langle quant\ intf \rangle \ \ \langle qual\ intf \rangle$	(33)
$\langle simp\ ptf \rangle \ ::= \ \langle quant\ ptf \rangle \ \ \langle qual\ ptf \rangle$	(34)
$\langle int\ obs \rangle$	
$::= \langle simp \; ptf \rangle \; \; \langle simp \; ptf \rangle \; \mathbf{at} \; \langle point \rangle \; \; \langle simp \; intf \rangle$	(35)
$ \langle \text{simp intf} \rangle $ when observed between $\langle interval \rangle$	(36)
$\langle qdesc \rangle ::= \langle \text{int obs} \rangle$	(37)
$ \langle \text{int obs} \rangle $ in all trajectories	(38)
\(\langle \quant agg \text{intf} \rangle	
when observed between $\langle interval \rangle$	(39)
$ $ \langle quant agg ptf \rangle when observed at \langle point \rangle	(40)
$\langle cqdesc \rangle ::= \langle quant cagg intf \rangle$	
when observed between $\langle interval \rangle$	(41)
$ $ \langle quant cagg ptf \rangle when observed at \langle point \rangle	(42)
$\langle interv \rangle ::= $ remove f as soon as produced	(43)
disable a	(44)
continuously supply f in quantity q	(45)
add delay of q time units in availability of f	(46)
set value of f to q	(47)
$\langle icond \rangle ::= (45) \mid (47)$	(48)
$\langle qstmt \rangle ::= \langle qdesc \rangle;$	
due to interventions : $\langle interv \rangle_1, \dots, \langle interv \rangle_{N1};$	
due to observations : $\langle \text{int obs} \rangle_1, \dots, \langle \text{int obs} \rangle_{N2};$	
using initial setup : $\langle icond \rangle_1, \dots, \langle icond \rangle_{N3}$;	(49)

⁶Although some of our single-trajectory queries can be represented as LTL formulas, we have chosen to keep the current representation as it is more intuitive for our biological domain.

 $^{^5}$ Note that 'f has value 5 or higher' overlaps with 'f has value 7 or higher'.

| \(\text{cqdesc} \); comparing nominal pathway with modified pathway obtained

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due to interventions :\langle interv \rangle_1, \dots, \langle interv \rangle_{N1};
due to observations :\langle int obs \rangle_1, \dots, \langle int obs \rangle_{N2};
using initial setup :\langle icond \rangle_1, \dots, \langle icond \rangle_{N3}; (50)
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Given a pathway specification (domain description) P with trajectories of the form $\sigma = s_0, T_0, s_1, \dots, s_{k-1}, T_{k-1}, s_k$, where each s_i, T_i, s_{i+1} is an evolution from state s_i to state s_{i+1} due to firing of T_i . Intuitively, a *point* is a time-step and interval defines a continuous range of time-steps on a trajectory, then *point* formulas (represented by ptf) are evaluated w.r.t. a point on the trajectory, while interval formulas (represented by int f) are evaluated w.r.t. an interval on a trajectory, e.g. rate of production of f over interval [i, j] is given by $(s_i(f) - s_i(f))/(j-i)$. Quantitative formulas (represented by quant) are evaluated for some quantity n, while qualitative formulas (represented by qual) are evaluated for some qualitative attribute of a state / trajectory. Aggregate quantitative formulas (represented by $quant \ aqq$) are evaluated for some quantity n, which is an aggregate of quantities n_1, \ldots, n_m for trajectories $\sigma_1, \ldots, \sigma_m$, e.g. the average aggregate is computed as $n = (n_1 + \cdots + n_m)/m$. Comparative quantitative formulas (represented by $quant \ cagg$) are evaluated for some direction of change between aggregate quantities n (over trajectories $\sigma_1, \ldots, \sigma_m$) and n' (over trajectories $\sigma'_1, \ldots, \sigma'_{m'}$), e.g. the change direction is '>' if n' > n.

Intuitively, an internal observation (represented by *int obs*) is a simple point formula that holds at any point on a trajectory, a simple point formula that holds at a specific point, a simple interval formula that holds over any interval on a trajectory, or a simple interval formula that holds over a specific interval. Intuitively, an internal observation filters the set of trajectories produced by a pathway specification.

Intuitively, a query description (represented by qdesc) is one of the possible point formulas that holds at a specific point, an interval formula that holds over a specific range, an internal observation, an internal observation over all trajectories in a set of given trajectories. A comparative query description (represented by cqdesc) is one of the quantitative comparative aggregate point formula at a specific point on two sets of trajectories, or a quantitative comparative aggregate interval formula over a specific interval on two sets of trajectories. Intuitively, a query description or a comparative query description specifies the property that we want to have hold true over the trajectories of the domain pathway.

Intuitively, an intervention (represented by *interv*) specifies a modification to the pathway specification, e.g. intervention (43) modifies the pathway such that all quantity of f is removed as soon as it is produced.

Intuitively, a query statement (represented by *qstmt*) is a *comparative query statement* if it contains a comparative query description, and non-comparative otherwise. A query statement is composed of a query statement, interventions to the pathway, internal observations to filter the trajectories, and initial conditions. Intuitively, a *query statement* asks whether a *query description* holds in a pathway, after modifying it with initial setup, interventions and observations.

While, a *comparative query statement* asks whether a *query description* holds when a nominal pathway is compared to a modified pathway, subject to same initial setup, but interventions and observations only applied to the modified pathway.

Pathway Semantics

The semantics of our pathway are given by a Guarded-Arc Petri Net, which allows choice between different effects (arc-sets) of a transition, such that only one effect (arc-set) is active for a transition in a given state determined by its arc-guard ⁷.

Definition 2 (Guard). A guard condition *takes one of the following forms*: $(f < v), (f \le v), (f > v), (f \ge v), or (f = v)$, where f is a fluent; and v is a fluent or a numeric constant. A guard is a propositional formula of guard conditions, with each guard condition treated as a proposition.

An interpretation of a guard G is a possible assignment of a value to each fleuent $f \in G$ from the domain of f. A guard G is satisfied w.r.t. a state s, written $s \models G$ iff G has an interpretation in which each of its fluents f has the value s(f) and G is true.

Definition 3 (Guarded-Arc Petri Net). A Guarded-Arc Petri Net is a tuple $PN^G = (P, T, G, E, R, W, D, TG, L)$:

P is a finite set of places

T is a finite set of transitions

G is a set of guards as defined in definition (2)

 $TG: T \rightarrow G$ are the transition guards

 $E \subseteq (T \times P \times G) \cup (P \times T \times G)$ are the guarded arcs

 $R \subseteq P \times T \times G$ are the guarded reset arcs

 $W: E \to \mathbb{N}^+$ are arc weights

 $D: T \to \mathbb{N}^+$ are the transition durations

 $L: P \to \mathbb{N}^+$ specifies maximum tokens for each place

subject to constraints: (i) $P \cap T = \emptyset$ (ii) $R \cap E = \emptyset$ (iii) Let $t \in T$ be a transition, and $gg_t = \{g : (t, p, g) \in E\} \cup \{g : (p, t, g) \in E\} \cup \{g : (p, t, g) \in R\}$ be the set of arcguards for normal and reset arcs connected to it, then any two distinct guards $g_1 \in gg_t, g_2 \in gg_t$ must not have an interpretation that makes both g_1 and g_2 true.

We make a simplifying assumption that all places are readable by using their place names. Execution of the PN^G occurs in discrete time steps. The *marking (or state)* of a Guarded-Arc Petri Net PN^G is the token assignment of each place $p_i \in P$. Initial marking is given by $M_0: P \to \mathbb{N}^0$, while the token assignment at step k is written as M_k .

Next we define the execution semantics of PN^G . We start with terminology used below. Let (i) $s_0 = M_0$ represent the the initial marking (or state), $s_k = M_k$ represent the marking (or state) at time step k, (ii) $s_k(p)$ represent the marking of place p at time step k, such that $s_k = [s_k(p_0), \ldots, s_k(p_n)]$, where $P = \{p_0, \ldots, p_n\}$ (iii) T_k

⁷Our model is similar to the model in (Jensen, Kristensen, and Wells 2007) in many aspects with differences in certain key semantics related to biological modeling, such as reset arts.

be the firing-set that fired in step k, (iv) en_k be the set of enabled transitions in state s_k , (v) $del_k(p,\{t_1,\ldots,t_n\})$ be the sum of tokens that will be consumed from place p if transitions t_1,\ldots,t_n fired in state s_k , (vi) $overc_k(\{t_1,\ldots,t_n\})$ be the set of places that will have over-consumption of tokens if transitions t_1,\ldots,t_n were to fire simultaneously in state s_k , (vii) $sel_k(fs)$ be the set of possible firing-set choices in state s_k using fs firing style (viii) $add_k(p)$ be the total production of tokens in place p (in state s_k) due to actions terminating in state s_k , (ix) s_{k+1} be the next state computed from state s_k due to firing transition-set T_k . Then, the execution semantics of the Guarded-Arc Petri Net PN^G starting from state s_0 using firing-style fs is given as follows:

$$\begin{split} en_k &= \{t: t \in T, s_k \models TG(t), \forall (p,t,g) \in E, \\ &(s_k \models g, s_k(p) \geq W(p,t,g))\} \\ del_k(p, \{t_1, \dots, t_n\}) &= \\ &\sum_{i=1,\dots,n} W(p,t_i,g) : (p,t_i,g) \in E, s_k \models g \\ &+ \sum_{i=1,\dots,n} s_k(p) : (p,t_i,g) \in R, s_k \models g \\ overc_k(\{t_1, \dots, t_n\}) &= \{p: p \in P, \\ s_k(p) < del_k(p, \{t_1, \dots, t_n\})\} \\ sel_k(1) &= \{\{ss\} : ss \in en_k, overc_k(\{ss\}) = \emptyset\} \\ sel_k(*) &= \{ss : ss \in 2^{en_k}, overc_k(ss) = \emptyset\} \\ sel_k(max) &= \{ss : ss \in 2^{en_k}, overc_k(p, ss) = \emptyset, \\ (\nexists ss' \in 2^{en_k} : ss \subset ss', overc_k(ss') = \emptyset)\} \\ T_k &\in sel_k(fs) \\ add_k(p) &= \sum_{j=0,\dots,k} W(t_i, p, g) \\ &: (t_i, p, g) \in E, t_i \in T_j, D(t_i) + j = k + 1 \\ s_{k+1}(p) &= min(s_k(p) - del_k(p, T_k) + add_k(p), L(p)) \end{cases} \tag{51} \end{split}$$

Definition 4 (Trajectory). $\sigma = s_0, T_0, s_1, \ldots, s_{k-1}, T_{k-1}, s_k$ is a trajectory of PN^G iff given $s_0 = M_0$, each T_i is a possible firing-set in s_i whose firing produces s_{i+1} per PN^G 's execution semantics in (51).

Query Semantics

First we give the semantics of domain modification due to an intervention by examples. Consider intervention (43), we create the modified domain description $\mathbf{D}' = \mathbf{D} \diamond (\mathbf{remove} \ f \ \mathbf{as} \ \mathbf{soon} \ \mathbf{as} \ \mathbf{produced})$ as follows:

 $\mathbf{D}' = \mathbf{D} + \{tr \text{ may execute causing } f \text{ change value by } *\}$

Applying intervention (45) $\mathbf{D}' = \mathbf{D} \diamond (\mathbf{continuously} \ \mathbf{supply} \ f \ \mathbf{in} \ \mathbf{quantity} \ q)$ results in the following changes:

$$D' = D + \{t_f \text{ may execute causing } f \text{ change value by } + q\}$$

Next we define the semantics of some common formulas and observation using LTL-style. Let $\sigma = s_0, T_0, s_1, \ldots, T_{k-1}, s_k$ represent a trajectory of domain **D** with initial marking s_0 . Let actions T_i firing in state s_i be observable in s_i such that $T_i \subseteq s_i$.

 $\begin{array}{l} s_i \text{ such that } T_i \subseteq s_i. \\ \text{ Let } \langle s_i, \sigma \rangle \models F \text{ represent } F \text{ holds at point } i \text{ in } \sigma; \\ \{\langle s_i^1, \sigma_1 \rangle, \ldots, \langle s_i^m, \sigma_m \rangle\} \models F \text{ represent } F \text{ holds at point } \end{array}$

 $\begin{array}{l} i \ \ \text{in trajectories} \ \sigma_1, \ldots, \sigma_m; \ \left\{\left\langle s_i^1, \sigma_1\right\rangle, \ldots, \left\langle s_i^m, \sigma_m\right\rangle\right\}, \\ \left\{\left\langle \bar{s}_i^1, \bar{\sigma}_1\right\rangle, \ldots, \left\langle \bar{s}_i^{\bar{m}}, \bar{\sigma}_{\bar{m}}\right\rangle\right\}\right\} \ \ \models \ F \ \ \text{represent} \ F \ \ \text{holds} \ \ \text{at} \\ \text{point} \ \ i \ \ \text{in both sets} \ \left\{\sigma_1, \ldots, \sigma_m\right\} \ \ \text{and} \ \left\{\bar{\sigma}_1, \ldots, \bar{\sigma}_{\bar{m}}\right\}; \\ \left(\left\langle s_i, \sigma\right\rangle, j\right) \ \ \models \ F \ \ \text{represent} \ F \ \ \text{holds} \ \ \text{over interval} \ \ [i,j] \ \ \text{in} \\ \sigma_i, \left(\left\langle s_i^1, \sigma_1\right\rangle, \ldots, \left\langle s_i^m, \sigma_m\right\rangle\right\}, j\right) \ \ \models \ F \ \ \text{represent} \ F \ \ \text{holds} \ \ \text{over interval} \ \ \ [i,j] \ \ \text{in} \\ \sigma_1, \ldots, \sigma_m; \ \ \text{and} \ \ \left(\left\{\left\langle s_i^1, \sigma_1\right\rangle, \ldots, \left\langle s_i^m, \sigma_{\bar{m}}\right\rangle\right\}\right\}, j\right) \ \ \ F \ \ \text{represent} \\ F \ \ \text{holds} \ \ \ \text{over interval} \ \ \ [i,j] \ \ \ \text{over two sets} \ \ \left\{\sigma_1, \ldots, \sigma_m\right\} \ \ \text{and} \\ \left\{\bar{\sigma}_1, \ldots, \bar{\sigma}_{\bar{m}}\right\}. \ \ \text{Then the semantics of a rate query are given as follows:} \end{array}$

$$(\langle s_i,\sigma\rangle,j) \models \text{ rate of production of } f \text{ is } n \\ \text{ if } n = (s_j(f) - s_i(f))/(j-i) \\ (\{\langle s_i^1,\sigma_1\rangle,\ldots,\langle s_i^m,\sigma_m\rangle\},j) \models \\ average \text{ rate of production of } f \text{ is } r \\ \text{ if } \exists [r_1,\ldots,r_m]: (\langle s_i^1,\sigma_1\rangle,j) \models \text{ rate of production } \\ \text{ of } f \text{ is } r_1\ldots(\langle s_i^m,\sigma_m\rangle,j) \models \text{ rate of production } \\ \text{ of } f \text{ is } r_m \text{ and } r = (r_1+\cdots+r_m)/m \\ (53) \\ (\{\langle s_i^1,\sigma_1\rangle,\ldots,\langle s_i^m,\sigma_m\rangle\},\{\langle \bar{s}_i^1,\bar{\sigma}_1\rangle,\ldots,\langle \bar{s}_i^{\bar{m}},\bar{\sigma}_{\bar{m}}\rangle\}\},j) \\ \models \text{ direction of change in } average \text{ rate of } \\ \text{ production of } f \text{ is } d \\ \text{ if } \exists n_1: (\{\langle s_i^1,\sigma_1\rangle,\ldots,\langle s_i^m,\sigma_m\rangle\},j) \models average \text{ rate } \\ \text{ of production of } f \text{ is } n_1 \text{ and } \\ \exists n_2: (\{\langle \bar{s}_i^1,\bar{\sigma}_1\rangle,\ldots,\langle \bar{s}_i^{\bar{m}},\bar{\sigma}_{\bar{m}}\rangle\},j) \models average \text{ rate } \\ \text{ of production of } f \text{ is } n_2 \text{ and } n_2 \ d \ n_1 \\ \\ \{\langle s_0^1,\sigma_1\rangle,\ldots,\langle s_0^m,\sigma_m\rangle\},\{\langle \bar{s}_0^1,\bar{\sigma}_1\rangle,\ldots,\langle \bar{s}_0^{\bar{m}},\bar{\sigma}_{\bar{m}}\rangle\}\} \\ \models \text{ direction of change in } average \text{ rate of } \\ \text{ production of } f \text{ is } d \text{ when observed between } \\ \text{ time step } i \text{ and time step } j \\ \text{ if } \Big(\Big\{\{\langle s_i^1,\sigma_1\rangle,\ldots,\langle s_i^m,\sigma_m\rangle\},\{\langle \bar{s}_i^1,\bar{\sigma}_1\rangle,\ldots,\langle \bar{s}_i^{\bar{m}},\bar{\sigma}_{\bar{m}}\rangle\}\Big\},j\Big) \\ \models \text{ direction of change in } \langle average \text{ rate of } \\ \text{ direction of change in } \langle average \text{ rate of } \\ \end{pmatrix}$$

Definition 5. Let \mathbf{D} be a domain description and \mathbf{Q} be a query statement (49) with query description U, interventions $V_1,\ldots,V_{|V|}$, internal observations $O_1,\ldots,O_{|O|}$, and initial conditions $I_1,\ldots,I_{|I|}$. Let $\mathbf{D_1} \equiv \mathbf{D} \diamond I_1 \diamond \cdots \diamond I_{|I|} \diamond V_1 \diamond \cdots \diamond V_{|V|}$ be the modified domain description constructed by applying the initial conditions and interventions from \mathbf{Q} and σ_1,\ldots,σ_m be its trajectories that satisfy $O_1,\ldots,O_{|O|}$. Then, \mathbf{D} satisfies \mathbf{Q} (written $\mathbf{D} \models \mathbf{Q}$) if $\{\sigma_1,\ldots,\sigma_m\} \models U$.

(55)

production of f is d

Definition 6. Let \mathbf{D} be a domain description and \mathbf{Q} be a comparative query statement (50) with query description U, interventions $V_1, \ldots, V_{|V|}$, internal observations $O_1, \ldots, O_{|O|}$, and initial conditions $I_1, \ldots, I_{|I|}$. Let $\mathbf{D_0} \equiv \mathbf{D} \diamond I_1 \diamond \cdots \diamond I_{|I|}$ be the nominal domain description constructed by applying the initial conditions from \mathbf{Q} and $\sigma_1, \ldots, \sigma_m$ be its trajectories. Let $\mathbf{D_1} \equiv \mathbf{D} \diamond I_1 \diamond \cdots \diamond I_{|I|} \diamond V_1 \diamond \cdots \diamond V_{|V|}$ be the alternate domain description constructed by applying the initial conditions and interventions from \mathbf{Q} and $\bar{\sigma}_1, \ldots, \bar{\sigma}_{\bar{m}}$ be its trajectories that sat-

isfy $O_1, \ldots, O_{|O|}$. Then, \mathbf{D} satisfies \mathbf{Q} (written $\mathbf{D} \models \mathbf{Q}$) if $(\{\sigma_1, \ldots, \sigma_m\}, \{\bar{\sigma}_1, \ldots, \bar{\sigma}_{\bar{m}}\}) \models U$.

The following propositions follow from the above description.

Proposition 1. Let \mathbf{D} be a domain description and \mathbf{Q} be a non-comparative query statement. Then, $\mathbf{D} \models \mathbf{Q}$ iff $\mathbf{D}' \models \mathbf{Q}'$ where $\mathbf{D}' \equiv \mathbf{D} \diamond I_1 \cdots \diamond I_{|I|} \diamond V_1 \cdots \diamond V_{|V|}$ and $\mathbf{Q}' \equiv \mathbf{Q}$ with $I_1, \ldots, I_{|I|}, V_1, \ldots, V_{|V|}$ removed, where $I_1, \ldots, I_{|I|}$ are the initial conditions in \mathbf{Q} and $V_1, \ldots, V_{|V|}$ are the interventions in \mathbf{Q} .

Proposition 2. Let \mathbf{D} be a domain description and \mathbf{Q} be a non-comparative query statement. Then, $\mathbf{D} \models \mathbf{Q}$ iff $\{\sigma_1,\ldots,\sigma_n\} \models \mathbf{Q}'$, where σ_1,\ldots,σ_n are trajectories of \mathbf{D}' that satisfy $O_1,\ldots,O_{|O|}$, $\mathbf{D}'\equiv\mathbf{D}\diamond I_1\cdots\diamond I_{|I|}\diamond V_1\cdots\diamond V_{|V|}$ and $\mathbf{Q}'\equiv\mathbf{Q}$ with $I_1,\ldots,I_{|I|},V_1,\ldots,V_{|V|},O_1,\ldots,O_{|O|}$ removed; $I_1,\ldots,I_{|I|}$ are the initial conditions in \mathbf{Q} , $V_1,\ldots,V_{|V|}$ are the interventions in \mathbf{Q} , and $O_1,\ldots,O_{|O|}$ are internal observations in \mathbf{Q} .

Proposition 3. Let \mathbf{D} be a domain description and \mathbf{Q} be a comparative query statement. Then, $\mathbf{D} \models \mathbf{Q}$ iff $(\{\sigma_1,\ldots,\sigma_n\},\{\sigma'_1,\ldots,\sigma'_{n'}\}) \models \mathbf{Q}'$, where σ_1,\ldots,σ_n are trajectories of \mathbf{D}' , and $\sigma'_1,\ldots,\sigma'_{n'}$ are trajectories of \mathbf{D}'' that satisfy $O_1,\ldots,O_{|O|}$, $\mathbf{D}' \equiv \mathbf{D} \diamond I_1 \cdots \diamond I_{|I|}$, $\mathbf{D}'' \equiv \mathbf{D} \diamond I_1 \cdots \diamond I_{|I|} \diamond V_1 \cdots \diamond V_{|V|}$, $\mathbf{Q}' \equiv \mathbf{Q}$ with $I_1,\ldots,I_{|I|},V_1,\ldots,V_{|V|},O_1,\ldots,O_{|O|}$ removed; $I_1,\ldots,I_{|I|}$ are the initial conditions in \mathbf{Q} , $V_1,\ldots,V_{|V|}$ are the interventions in \mathbf{Q} , and $O_1,\ldots,O_{|O|}$ are internal observations in \mathbf{Q} .

Intuitively, proposition 1 states that a domain description D satisfies a (non-comparative) query statement Q whenever the modified domain description D' constructed by applying interventions and initial conditions from Q to D satisfies the modified query statement Q' constructed by removing interventions and initial conditions from Q. Proposition 2 states that a domain description D satisfies a (noncomparative) query statement Q whenever the trajectories of the modified domain description D' constructed by applying interventions and initial conditions from Q to D that satisfy observations in Q, satisfy the modified query statement \mathbf{Q}' constructed by removing interventions, initial conditions, and observations from Q. Proposition 3 states that a domain description D satisfies a comparative query statement Q whenever the comparison between trajectories of the nominal domain description D' constructed by applying initial conditions from Q and the trajectories of the alternate domain description D'' constructed by applying interventions and initial conditions from Q that satisfy the observations in Q satisfy the query statement Q' constructed by removing interventions, initial conditions and observations from **Q**.

Proof of the above propositions are based on the definitions 5,6 of satisfiability of a query statement \mathbf{Q} by domain \mathbf{D} ; the semantics of the query description U in the query statement; and the structure of the query statements. The forward direction is proven by showing that the trajectories of the modified domain descriptions (constructed by apply-

ing initial conditions and interventions) filtered by the internal observations (as appropriate) satisfy the query description U in the query statement \mathbf{Q} . The reverse direction is proven by showing that one can select a domain description \mathbf{D} that, when modified through selected initial setup conditions and interventions represents the same trajectories as \mathbf{D}' , and one can select observations that filter these trajectories to only those satisfied by the query description U. The selected initial setup conditions, interventions, and the observations when added to \mathbf{Q}' give us \mathbf{Q} .

Illustrative Example

We illustrate our high level language by applying it to question (1) and the relevant glycolysis pathway.

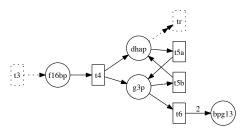


Figure 1: Petri Net for question 1.

The following pathway specification encodes the domain description \mathbf{D} for question (1) and produces the PN in Fig. 1 minus the tr, t3 transitions:

domain of f16bp is integer, dhap is integer, g3p is integer, bpg13 is integer t4 may execute causing f16bp change value by -1, dhap change value by +1, g3p change value by +1 t5a may execute causing dhap change value by -1, g3p change value by +1 t5b may execute causing g3p change value by -1, dhap change value by +1 t6 may execute causing g3p change value by -1, bpg13 change value by +2
initially f16bp has value 0, dhap has value 0, g3p has value 0, bpg13 has value 0
firing style max

The question is asking for the direction of change in the *rate* of glycolysis when the nominal pathway is compared against a modified pathway in which DHAP is removed as soon as it is produced. Since this rate can vary with the trajectory of the world evolution, we consider the average change in rate. Using domain knowledge (Reece et al. 2010, Figure 9.9), we measure the rate of glycolysis indirectly by measuring the rate of production of bpg13, a downstream product, which is converted into equivalent quantity of the end product of glycolysis. We also add continuous supply of source ingredient f16bp at the rate of 1 unit per time step to prevent starvation. This results in a query statement (15) $\bf Q$ for some simulation length k. We evaluate it for the direction d using

our deep reasoning system, which gives us d='<', suggesting that the rate of glycolysis will be lowered when DHAP is removed as soon as it is produced.

Implementation

We implemented a system ⁸ that understands a subset of our high level language using Python as the driver as well as the high level language parser; and Clingo (Gebser et al. 2011) as the ASP solver. The system takes a pathway specification, a query specification, and simulation parameters, such as simulation length and maximum possible tokens at any place for query evaluation. Our system uses Petri Net semantics for modeling the biological pathway and ASP for simulating it. Interventions in questions are modeled as Petri Net extensions and translated into ASP using the approach in (Anwar, Baral, and Inoue 2013a; 2013b). This gave us the translations of extensions such as reset arcs, inhibit arcs (Peterson 1977), read arcs (Christensen and Hansen 1993), colored tokens (Peterson and others 1980), and timed transitions (Ramchandani 1974) etc. We augmented the encoding further to include additional extensions, such as conditional effects of actions, and more generalized inhibitions supported by our language.

Next, we briefly summarize how our system processes a non-comparative aggregate quantitative query statement, then describe how the comparative quantitative query from previous section is evaluated.

To evaluate a non-comparative query, the system builds a Petri Net model from the pathway specification. It then applies the initial conditions and interventions from the query statement to this model. The model is then translated into ASP. The resulting ASP code is augmented with constraints for (internal) observations. Answer sets of the augmented code provide the filtered trajectories. Atoms needed to compute the quantity specified in the query (e.g. rate of production) are extracted from the answer sets and quantity values aggregated across answer sets. The aggregated value is compared against any aggregate value provided for query statement satisfaction or the computed value is returned.

To evaluate comparative quantitative query statement from the previous section is decomposed into two subqueries, \mathbf{Q}_0 for nominal case and \mathbf{Q}_1 for the modified case:

 $\mathbf{Q_0} \equiv average \text{ rate of production of } bpg13 \text{ is } n_{avg}$ when observed between time step 0 and time step k; using initial setup:

continuously supply f16bp in quantity 1;

 $\mathbf{Q_1} \equiv average \text{ rate of production of } bpg13 \text{ is } n'_{avg}$ when observed between time step 0 and time step k; due to interventions:

remove dhap as soon as produced; using initial setup :

continuously supply f16bp in quantity 1;

The queries are evaluated w.r.t. the same initial conditions for average rates n_{avg} , n'_{avg} that satisfy direction d in the original query statement. The Petri Net model in Figure 1 shows nominal case ($\mathbf{D_0}$) in solid lines and the interventions

added for the alternate case $(\mathbf{D_1})$ as dotted lines. The average results are compared using d to determine if they specify the direction given in the comparative query statement.

Conclusion

We have presented a high level (English like) language for specifying pathways and asking queries against them. Our pathway specification language uses Petri Net semantics as the simulation model and our query language is inspired by action languages. Our query language is one of the main contributions of this paper. It allows aggregate queries over a set of trajectories, comparative aggregate queries over two sets of trajectories, and interventions that are more general than actions which can be used to modify the pathway as specified in a query statement. Our approach improves on the query languages associated with action languages, by implementing comparative queries.

Though some aspects of our language may appear cumbersome to a person with background in action languages, we retained the syntax to allow coherent description of a variety of complex interventions and queries.

We illustrated how an example question comparing alternate scenarios of a biological pathway is encoded in our high level language; and summarized how our implementation performs query evaluation. We will elaborate on these aspects in companion papers.

Our approach fits into a larger problem of representing and reasoning about biological pathways gaining interest lately, geared towards developing an end-to-end system that extracts biological knowledge from texts, assembles it into pathways, determines the right level of abstraction eliminating irrelevant details, and answers such questions about them that require understanding of the inner workings of these pathways; with the ability to pose questions in natural language and present results in a natural language or a visual representation such as graphs. These aspects serve as guidelines for future extension to our work, and indeed there is existing work on many of these, including from our group, e.g., see (Tari et al. 2009; 2010) for pathway construction from text extraction and translating natural language questions to formal queries that we will extend.

The importance of determining the right level of abstraction by eliminating irrelevant details is two fold. On one hand it can improve performance by reducing problem size and computational resources needed, while on the other, it presents cleaner results that are easier to interpret by eliminating extraneous information. In this regard, techniques from multi-scale modeling (see (Dada and Mendes 2011; Heiner and Gilbert 2013)) could be applicable.

Additional areas of improvements include encapsulating time information in the queries such that it is hidden from the user, as well as the ability to apply interventions at arbitrary points during the simulation. Our current implementation uses Clingo, which handles discrete quantities only. We intend to extend this work to include continuous real numbers based on work by (Lee and Meng 2013).

⁸https://sites.google.com/site/deepqa2014/

References

- Anwar, S.; Baral, C.; and Inoue, K. 2013a. Encoding higher level extensions of Petri nets in answer set programming. In Cabalar, P., and Son, T., eds., *Logic Programming and Nonmonotonic Reasoning*, volume 8148 of *Lecture Notes in Computer Science*. Springer Berlin Heidelberg. 116–121.
- Anwar, S.; Baral, C.; and Inoue, K. 2013b. Encoding Petri nets in answer set programming for simulation based reasoning. *TPLP* 13(4-5-Online-Supplement).
- Baral, C.; Chancellor, K.; Tran, N.; Tran, N.; Joy, A.; and Berens, M. 2004. A knowledge based approach for representing and reasoning about signaling networks. *Bioinformatics* 20(suppl 1):i15–i22.
- Baral, C. 2003. *Knowledge Representation, Reasoning and Declarative Problem Solving*. Cambridge University Press.
- Christensen, S., and Hansen, N. D. 1993. *Coloured Petri nets extended with place capacities, test arcs and inhibitor arcs*. Springer.
- Dada, J. O., and Mendes, P. 2011. Multi-scale modelling and simulation in systems biology. *Integrative Biology* 3(2):86–96.
- Gebser, M.; Kaminski, R.; Kaufmann, B.; Ostrowski, M.; Schaub, T.; and Schneider, M. 2011. Potassco: The Potsdam answer set solving collection. *AICom* 24(2):105–124.
- Gelfond, M., and Lifschitz, V. 1993. Representing action and change by logic programs. *The Journal of Logic Programming* 17(2):301–321.
- Gelfond, M., and Lifschitz, V. 1998. Action languages. *Electronic Transactions on AI* 3(16).
- Giunchiglia, E., and Lifschitz, V. 1998. An action language based on causal explanation: Preliminary report. In *AAAI/IAAI*, 623–630. Citeseer.
- Giunchiglia, E.; Lee, J.; Lifschitz, V.; McCain, N.; and Turner, H. 2004. Nonmonotonic causal theories. *Artificial Intelligence* 153(1):49–104.
- Heiner, M., and Gilbert, D. 2013. Biomodel engineering for multiscale systems biology. *Progress in biophysics and molecular biology* 111(2):119–128.
- Jensen, K.; Kristensen, L. M.; and Wells, L. 2007. Coloured Petri nets and CPN Tools for modelling and validation of concurrent systems. *International Journal on Software Tools for Technology Transfer* 9(3-4):213–254.
- Lee, J., and Lifschitz, V. 2003. Describing additive fluents in action language $\mathcal{C}+$. In *Proc. of IJCAI 2003*.
- Lee, J., and Meng, Y. 2013. Answer set programming modulo theories and reasoning about continuous changes. *IJ-CAI'13*.
- Lee, J.; Lifschitz, V.; and Yang, F. 2013. Action language BC: Preliminary report. In *Proceedings of International Joint Conference on Artificial Intelligence (IJCAI)*.
- Peterson, J., et al. 1980. A note on colored Petri nets. *Information Processing Letters* 11(1):40–43.
- Peterson, J. L. 1977. Petri nets. *Computing Surveys* 9(3):223–252.

- Ramchandani, C. 1974. Analysis of asynchronous concurrent systems by Petri nets. Technical report, DTIC Document
- Reece, J.; Cain, M.; Urry, L.; Minorsky, P.; and Wasserman, S. 2010. *Campbell Biology*. Pearson Benjamin Cummings.
- Reiter, R. 1996. Natural actions, concurrency and continuous time in the situation calculus. *KR* 96:2–13.
- Tari, L.; Hakenberg, J.; Gonzalez, G.; and Baral, C. 2009. Querying a parse tree database of medline text to synthesize user-specific biomolecular networks. *Proc Pac Symp Biocomput* 14:87–98.
- Tari, L.; Baral, C.; Anwar, S.; Liang, S.; and Hakenberg, J. 2010. Synthesis of pharmacokinetic pathways through knowledge acquisition and automated reasoning. *Pacific Symposium on Biocomputing* 15:465–476.