



ON THE FORMALIZATION OF ZSYNTAX WITH APPLICATIONS IN MOLECULAR BIOLOGY

SOHAIB AHMAD, OSMAN HASAN, UMAIR SIDDIQUE*

Abstract. Recent progress in nanotechnology and optical imaging offers promising features to develop effective drugs to cure chronic diseases, such as cancer and malaria. However, qualitative characterization of biological organisms (such as molecules) is the foremost requirement to identify key drug targets. One of the most widely used approaches in this domain is molecular pathways, which offers a systematic way to represent and analyse complex biological systems. Traditionally, such pathways are analysed using paper-and-pencil based proofs and simulations. However, these methods cannot ascertain accurate analysis, which is a serious drawback given the safety-critical nature of the applications of molecular pathways. To overcome these limitations, we recently proposed to formally reason about molecular pathways within the sound core of a theorem prover. As a first step towards this direction, we formally expressed three logical operators and four inference rules of Zsyntax, which is a deduction language for molecular pathways. In the current paper, we extend this formalization by verifying a couple of behavioural properties of Zsyntax based deduction using the HOL4 theorem prover and developing a biologist friendly graphical user interface. Moreover, we demonstrate the utilization of our work by presenting the formal analysis of cancer related molecular pathway, i.e., TP53 degradation and metabolic pathway known as Glycolysis.

Key words: Molecular biology, logical deduction

AMS subject classifications. 68Q25, 92C40, 03D05

1. Introduction. Molecular biology is extensively used to construct models of biological processes in the form of networks or pathways, such as protein-protein interaction networks and signalling pathways. The analysis of these biological networks, usually referred to as biological regulatory networks (BRNs) or gene regulatory networks (GRNs) [10], is based on the principles of molecular biology to understand the dynamics of complex living organisms. Moreover, the analysis of molecular pathways plays a vital role in investigating the treatment of various human infectious diseases and future drug design targets. For example, the analysis of BRNs has been recently used to predict treatment decisions for sepsis patients [15].

Traditionally, the molecular biology based analysis is carried out by biologists in the form of wet-lab experiments (e.g. [7, 13]). These experiments, despite being very slow and expensive, do not ensure accurate results due to the inability to accurately characterize the complex biological processes in an experimental setting. Other alternatives for deducing molecular reactions include paper-and-pencil proof methods (e.g. using Boolean modelling [27] or kinetic logic [28]) or computer-based techniques (e.g. [29]) for analysing molecular biology problems. The manual proofs become quite tedious for large systems, where the calculation of unknown parameters takes several hundred proof steps, and are thus prone to human errors. The computer-based methods consist of graph-theoretic techniques [21], Petri nets [11] and model checking [3]. These approaches have shown very promising results in many applications of molecular biology (e.g. [8, 14]). However, these methods are not generic and hence have been used to describe some specific areas of molecular biology only [4]. Moreover, the inherent state-space explosion problem of model checking [20] limits the scope of this success only to systems where the biological entities can acquire a small set of possible levels.

Theorem proving [12] can be considered as the second most widely used formal method. It does not suffer from the state-space explosion problem of model checking and has also been advocated for conducting molecular biology based analysis [30]. The main idea behind theorem proving is to construct a computer based mathematical model of the given system and then verify the properties of interest using deductive reasoning. The foremost requirement for conducting the theorem proving based analysis of any system is to formalize the mathematical or logical foundations required to model and analyse that system in an appropriate logic. There have been several attempts to formalize the foundations of molecular biology. For example, the earliest axiomatization even dates back to 1937 [31] and other efforts related to the formalization of biology are presented in [32, 25]. Recent formalizations, based on K -Calculus [6] and π -Calculus [22, 23, 24], also include some formal

*School of Electrical Engineering and Computer Science (SEECS), NUST, Islamabad, Pakistan ({11mseesahmad, osman.hasan, umair.siddique}@seecs.nust.edu.pk).

reasoning support for biological systems. But the understanding and utilization of these techniques is very cumbersome for a working biologist as highlighted by Fontana in [9].

In order to develop a biologist friendly formal deduction framework for reasoning about molecular reactions, we propose to formalize the Zsyntax [4] language in higher-order logic. Zsyntax is a formal language that supports modelling and logical deductions about any biological process. The main strength of Zsyntax is its biologist-centred nature as its operators and inference rules have been designed in such a way that they are understandable by the biologists. Traditionally, logical deductions about biological processes, expressed in Zsyntax, were done manually based on the paper-and-pencil based approach. This limits the usage of Zsyntax to smaller problems and also makes the deduction process error-prone due to the human involvement. As a first step towards overcoming this limitation, we formalized the logical operators and inference rules of Zsyntax in higher-order logic [2]. In the current paper, we build upon these formal definitions to verify a couple of key behavioural properties of Zsyntax based molecular pathways using the HOL4 theorem prover. The formal verification of these properties raises the confidence level in our definitions of Zsyntax operators and inference rules, which have complex interrelationships. Moreover, these formally verified properties can be used to facilitate the formal reasoning about chemical reactions at the molecular level. In order to illustrate the usefulness and effectiveness of our formalization for analysing real-world problems in molecular biology, we present the formal analysis of a cancer related molecular pathway involving TP53 degradation and a metabolic pathway known as Glycolysis. As a part of this work, we have also developed a user-friendly graphical user interface (GUI) for conducting the Zsyntax based formal analysis of molecular pathways. The distinguishing features of this tool includes its biologist friendliness and dedicated proof tactic that allows the users to perform automated reasoning about molecular pathways.

Our current framework handles static reactions but it can be further extended to study the reaction kinetics [4] due to the flexibility of Zsyntax. The main motivation behind using higher-order-logic theorem proving in our work is to be able to leverage upon the high expressiveness of higher-order logic and thus reason about differential equations and probabilistic properties, which form an integral part of reaction kinetics. However, the scope of the current paper is on the formalization of Zsyntax based deduction calculus for molecular pathways but this formalization can later be extended to support reaction kinetics as well because it is done in a higher-order-logic theorem prover.

The rest of the paper is organized as follows: Section 2 provides an introduction to Zsyntax and the HOL4 theorem prover. The higher-order-logic formalization of Zsyntax operators and inference rules using HOL4 are described in Sect. 3. This is followed by the descriptions of the behavioural properties of Zsyntax along with their formal proof sketches in Sect. 4 and 5, respectively. We provide a brief overview of the GUI developed as a part of this work in Sect. 6. In order to demonstrate the use of our formalization, we present two case studies on the TP53 degradation and glycolytic pathway in Sect. 7. We conclude the paper in Sect. 8 while highlighting some interesting potential applications of our work.

2. Preliminaries.

2.1. Zsyntax. Zsyntax [4] exploits the analogy between biological processes and logical deduction. Some of the key features of Zsyntax are: 1) the ability to express molecular reactions in a mathematical way; 2) heuristic nature, i.e., if the conclusion of a reaction is known, then one can deduce the missing data from the initialization data; 3) computer implementable semantics. Zsyntax consists of the following three operators:

Z-Interaction: The interaction of two molecules is expressed by the Z-Interaction ($*$) operator. In biological reactions, Z-interaction is not associative.

Z-Conjunction: The aggregate of same or different molecules (not necessarily interacting with each other) is formed using the Z-Conjunction ($\&$) operator. Z-Conjunction is fully associative.

Z-Conditional: A path from A to B under the condition C is expressed using the Z-Conditional (\rightarrow) operator as: $A \rightarrow B$ if there is a C that allows it.

Zsyntax supports four inference rules, given in Table 2.1, that play a vital role in deducing the outcomes of biological reactions:

Besides the regular formulas that can be derived based on the above mentioned operators and inference rules, Zsyntax also makes use of *Empirically Valid Formulae* (EVF). These EVFs basically represent the non-logical axioms of molecular biology and are assumed to be validated empirically in the lab.

TABLE 2.1
Zsyntax Inference Rules

Inference Rules	Definition
Elimination of Z-conditional(\rightarrow E)	if $C \vdash (A \rightarrow B)$ and $(D \vdash A)$ then $(C \& D \vdash B)$
Introduction of Z-conditional(\rightarrow I)	$C \& A \vdash B$ then $C \vdash (A \rightarrow B)$
Elimination of Z-conjunction($\&$ E)	$C \vdash (A \& B)$ then $(C \vdash A)$ and $(C \vdash B)$
Introduction of Z-conjunction($\&$ I)	$(C \vdash A)$ and $(D \vdash B)$ then $(C \& D) \vdash (A \& B)$

It has been shown that any biological reaction and its final outcome can be derived by using the above mentioned three operators and four inference rules [4]. For example, consider a scenario in which three molecules A, B and C react with each other to yield another molecule Z. This can be represented as a Zsyntax theorem as follows:

$$A \ \& \ B \ \& \ C \ \vdash \ Z$$

The Z-Conjunction operator $\&$ is used to represent the given aggregate of molecules and then the inference rules from Table 1 are applied on these molecules along with some EVFs (chemical reactions verified in laboratories) to obtain the final product Z. For the above example, these EVFs could be:

$$A \ * \ B \ \rightarrow \ X \ \text{and} \ X \ * \ C \ \rightarrow \ Z$$

meaning that A will react with B to yield X and X in return will react with C to yield the final product Z.

The main contribution of our paper is the formal verification of the Zsyntax based deduction method based on the higher-order-logic formalization of the above-mentioned operators and inference rules using the HOL4 theorem prover. This work will in turn facilitate the derivation of biological reactions within the sound core of HOL4.

2.2. HOL4 Theorem Prover. HOL4 is an interactive theorem prover developed at the University of Cambridge, UK, for conducting proofs in higher-order logic. It utilizes the simple type theory of Church [5] along with Hindley-Milner polymorphism [17] to implement higher-order logic. HOL4 has been successfully used as a verification framework for both software and hardware as well as a platform for the formalization of pure mathematics.

In order to ensure secure theorem proving, the logic in the HOL4 system is represented in the strongly-typed functional programming language ML [19]. An ML abstract data type is used to represent higher-order logic theorems and the only way to interact with the theorem prover is by executing ML procedures that operate on values of these data types. The HOL4 core consists of only 5 basic axioms and 8 primitive inference rules, which are implemented as ML functions. Soundness is assured as every new theorem must be verified by applying these basic axioms and primitive inference rules or any other previously verified theorems/inference rules.

A HOL4 theory is a collection of valid HOL4 types, constants, axioms and theorems, and is usually stored as a file in computers. Users can reload a HOL4 theory in the HOL4 system and utilize the corresponding definitions and theorems right away. Various mathematical concepts have been formalized and saved as HOL4 theories by the HOL4 users. We utilize the HOL4 theories of Booleans, arithmetics and lists extensively in our work. Table 2.2 provides the mathematical interpretations of some HOL4 symbols and functions frequently used in this paper.

3. Formalization of Zsyntax. We modelled the molecules as variables of arbitrary data types (α) in our formalization of Zsyntax [2]. A list of molecules (α list) represents the Z-Interaction or a molecular reaction among the elements of the list. The Z-Conjunction operator forms a collection of non-reacting molecules and can now be formalized as a list of list of molecules (α list list). This data type allows us to apply the Z-Conjunction operator between individual molecules (a list with a single element) or multiple interacting molecules (a list with multiple elements). The Z-Conditional operator is used to update the status of molecules, i.e., generate a new set of molecules based on the available EVFs (wet-lab verified reactions). Each EVF is modelled in our formalization as a pair (α list # α list list) where the first element is a list of molecules (α list) indicating the reacting molecules and the second element is a list of list of molecules (α list list) indicating the resulting set of

TABLE 2.2
HOL4 Symbols and Functions

HOL4 Symbol	Standard Symbol	Meaning
\wedge	<i>and</i>	Logical <i>and</i>
\vee	<i>or</i>	Logical <i>or</i>
\neg	<i>not</i>	Logical <i>negation</i>
$::$	<i>cons</i>	Adds a new element to a list
$++$	<i>append</i>	Joins two lists together
HD L	<i>head</i>	Head element of list <i>L</i>
TL L	<i>tail</i>	Tail of list <i>L</i>
EL n L	<i>element</i>	n^{th} element of list <i>L</i>
MEM a L	<i>member</i>	True if <i>a</i> is a member of list <i>L</i>
LENGTH L	<i>length</i>	Length of list <i>L</i>
FST	$\text{fst}(a, b) = a$	First component of a pair
SND	$\text{snd}(a, b) = b$	Second component of a pair
SUC n	$n + 1$	Successor of a <i>num</i>

molecules after the reaction between the molecules of the first element of the pair has taken place. A collection of EVFs is represented as a list of EVFs ($(\alpha \text{ list } \# \alpha \text{ list } \text{list})\text{list}$) in our formalization.

The elimination of Z-Conditional rule is the same as the elimination of implication rule (Modus Ponens) in propositional logic and thus it can be directly handled by the HOL4 simplification and rewriting rules. Similarly, the introduction of Z-Conditional rule can also be inferred from the rules of propositional logic and can be handled by the HOL4 system without the introduction of a new inference rule. The elimination of the Z-Conjunction rule allows us to infer the presence of a single molecule from an aggregate of inferred molecules. This rule is usually applied at the end of the reaction to check if the desired molecule has been obtained. Based on our data types, described above, this rule can be formalized in HOL4 by returning a particular molecule from a list of molecules:

DEFINITION 3.1. Elimination of Z-Conjunction Rule

$\vdash \forall L m. \text{z_conj_elim } L m = \text{if MEM } m L \text{ then } [m] \text{ else } L$

The function `z_conj_elim` has the data type $(\alpha \text{ list} \rightarrow \alpha \rightarrow \alpha \text{ list})$. The above function returns the given element as a single element in a list if it is a member of the given list. Otherwise, it returns the argument list as it is.

The introduction of Z-Conjunction rule along with Z-Interaction allows us to perform a reaction between any of the available molecules during the experiment. Based on our data types, this rule is equivalent to the append operation of lists.

DEFINITION 3.2. Intro of Z-Conjunction and Z-Interaction

$\vdash \forall L m n. \text{z_conj_int } L m n = \text{FLAT } [\text{EL } m L; \text{EL } n L] :: L$

The above definition has the data type $(\alpha \text{ list } \text{list} \rightarrow \text{num} \rightarrow \text{num} \rightarrow \alpha \text{ list } \text{list})$. The HOL4 functions `FLAT` and `EL` are used to flatten a list of list to a single list and return a particular element of a list, respectively. Thus, the function `z_conj_int` takes a list *L* and appends the list of two of its elements *m* and *n* on its head.

Based on the laws of stoichiometry [4], the reacting molecules using Z-Conjunction operator have to be deleted from the aggregate of molecules. The following function represents this behaviour in our formalization:

DEFINITION 3.3. Reactants Deletion

$\vdash \forall L m n. \text{z_del } L m n = \text{if } m > n$
 $\quad \text{then del (del } L m) n$
 $\quad \text{else del (del } L n) m$

Here the function `del L m` deletes the element at index *m* of the list *L* and returns the updated list as follows:

DEFINITION 3.4. Element Deletion

$$\vdash \forall L. \text{del } L \ 0 = \text{TL } L \wedge$$

$$\forall L \ n. \text{del } L \ (n + 1) = \text{HD } L :: \text{del } (\text{TL } L) \ n$$

Thus, the function $\text{z_del } L \ m \ n$ deletes the m^{th} and n^{th} elements of the given list L . We delete the higher indexed element before the lower one in order to make sure that the first element deletion does not effect the index of the second element that is required to be deleted. The above data types and definitions can be used to formalize any molecular pathway (which is expressible using Zsyntax) and reason about its correctness within the sound core of the HOL4 theorem prover.

4. Formal Reasoning support for Zsyntax. Our main objective is to develop a framework that accepts a list of initial molecules and possible EVFs and allows the user to formally deduce the final outcomes of the corresponding biological experiment within the sound core of HOL4.

In this regard, we first develop a function that compares a particular combination of molecules with all the EVFs and upon finding a match introduces the newly formed molecule in the initial list and deletes the consumed instances.

DEFINITION 4.1. EVF Matching

$$\vdash \forall L \ E \ m \ n. \text{z_EVF } L \ E \ 0 \ m \ n =$$

$$\text{if FST } (\text{EL } 0 \ E) = \text{HD } L$$

$$\text{then } (\text{T}, \text{z_del } (\text{TL } L \ ++ \ \text{SND } (\text{EL } 0 \ E)) \ m \ n)$$

$$\text{else } (\text{F}, \text{TL } L) \wedge$$

$$\forall L \ E \ p \ m \ n.$$

$$\text{z_EVF } L \ E \ (p + 1) \ m \ n =$$

$$\text{if FST } (\text{EL } (p + 1) \ E) = \text{HD } L$$

$$\text{then } (\text{T}, \text{z_del } (\text{TL } \ ++ \ \text{SND } (\text{EL } (p + 1) \ E)) \ m \ n)$$

$$\text{else } \text{z_EVF } L \ E \ p \ m \ n$$

Here, HD and TL returns head and tail of a list respectively. Similarly, FST and SND returns first and second element of a pair respectively. The data type of the function z_EVF is: $(\alpha \text{ list list} \rightarrow (\alpha \text{ list} \# \alpha \text{ list list}) \text{ list} \rightarrow \text{num} \rightarrow \text{num} \rightarrow \text{num} \rightarrow \text{bool} \# \alpha \text{ list list})$. The function LENGTH returns the length of a list. The function z_EVF takes a list of molecules L and recursively checks its head, or the top most element, against all elements of the EVF list E . If there is no match then the function returns a pair with its first element being false (F), indicating that no match occurred, and the second element equal to the tail of the input list L . Otherwise, if a match is found then the function replaces the head of list L with the second element of the EVF pair and deletes the matched elements from the initial list as these elements have already been consumed. This modified list is then returned along with a true (T) value, which acts as a flag to indicate an element replacement.

Next, in order to deduce the final outcome of the experiment, we have to call the function z_EVF recursively by placing all the possible combinations of the given molecules at the head of list L one by one. This can be done as follows:

DEFINITION 4.2. Recursive Function to model the argument n in Def. 4.1

$$\vdash \forall L \ E \ m. \text{z_recur1 } L \ E \ m \ 0 =$$

$$\text{z_EVF } (\text{z_conj_int } L \ m \ 0) \ E \ (\text{LENGTH } E - 1) \ m \ 0 \wedge$$

$$\forall L \ E \ m \ n.$$

$$\text{z_recur1 } L \ E \ m \ (n + 1) =$$

$$\text{if FST } (\text{z_EVF } (\text{z_conj_int } L \ m \ (n + 1)) \ E \ (\text{LENGTH } E - 1) \ m \ (n + 1)) \Leftrightarrow \text{T}$$

$$\text{then } \text{z_EVF } (\text{z_conj_int } L \ m \ (n + 1)) \ E \ (\text{LENGTH } E - 1) \ m \ (n + 1)$$

$$\text{else } \text{z_recur1 } L \ E \ m \ n$$

DEFINITION 4.3. Recursive Function to model the augment m in Def. 4.1

$$\vdash \forall L \ E \ n. \text{z_recur2 } L \ E \ 0 \ n =$$

$$\text{if FST } (\text{z_recur1 } L \ E \ 0 \ n) \Leftrightarrow \text{T}$$

$$\text{then } (\text{T}, \text{SND } (\text{z_recur1 } L \ E \ 0 \ n))$$

```

    else (F,SND (z_recur1 L E 0 n)) ∧
  ∀ L E m n.
z_recur2 L E (m + 1) n =
  if FST (z_recur1 L E (m + 1) n) ⇔ T
  then (T,SND (z_recur1 L E (m + 1) n))
  else z_recur2 L E m (LENGTH L - 1)

```

Both the above functions have the same data type, i.e., $(\alpha \text{ list list} \rightarrow (\alpha \text{ list} \# \alpha \text{ list list}) \text{ list} \rightarrow \text{num} \rightarrow \text{num} \rightarrow \text{num} \rightarrow \text{bool} \# \alpha \text{ list list})$. The function `z_recur1` calls the function `z_EVF` after appending the combination of molecules indexed by variables `m` and `n` using the introduction of Z-Conjunction rule. Thus, the new combination is always placed on the top of the list, which is passed as the variable `L` to the function `z_EVF`. Moreover, we provide the length of the EVF list (`LENGTH E - 1`) as the variable `p` of the function `z_EVF` so that the new combination is compared to all the entries of the EVF list. It is also important to note that the function `z_recur1` terminates as soon as a match in the EVF list is found. This function also returns a flag indicating the status of this match as the first element of its output pair. The second function `z_recur2` checks this flag and if it is found to be true (meaning that a match in the EVF list is found) then it terminates by returning the output list of the function `z_recur1`. Otherwise, it continues with all the remaining values of the variable `m` recursively. In the first case, i.e., when a match is found, the above two functions have to be called all over again with the new list. These iterations will continue until there is no match found in the execution of functions `z_recur1` and `z_recur2`. This overall behaviour can be modelled by the following recursive function:

DEFINITION 4.4. Final Recursion Function for Zsyntax

```

⊢ ∀ L E m n.
z_deduction_recur L E m n 0 = (T,L) ∧
  ∀ L E m n q.
z_deduction_recur L E m n (q + 1) =
  if FST (z_recur2 L E m n) ⇔ T
  then z_deduction_recur
    (SND (z_recur2 L E m n)) E (LENGTH (SND (z_recur2 L E m n)) - 1)
    (LENGTH (SND (z_recur2 L E m n)) - 1) q
  else (T,SND (z_recur2 L E (LENGTH L - 1) (LENGTH L - 1)))

```

The function `z_deduction_rec` also has the same data type as the above mentioned recursion functions. The variable of recursion in this function should be assigned a value that is greater than the total number of EVFs so that the application of none of the EVF is missed to guarantee correct operation. Similarly, the variables `m` and `n` above should be assigned the values of (`LENGTH L - 1`) to ensure that all the combinations of the list `L` are checked against the elements of the list of EVFs. Thus, the final deduction function for Zsyntax can be expressed in HOL4 as follows:

DEFINITION 4.5. Final Deduction Function for Zsyntax

```

⊢ ∀ L E. z_deduction L E =
  SND (z_deduction_recur L E (LENGTH L - 1) (LENGTH L - 1) LENGTH E)

```

The type of the function `z_deduction` is $(\alpha \text{ list list} \rightarrow (\alpha \text{ list} \# \alpha \text{ list list}) \text{ list} \rightarrow \alpha \text{ list list})$. It accepts the initial list of molecules and the list of valid EVFs and returns a list of final outcomes of the experiment under the given conditions. Next, the output of the function `z_deduction` has to be checked if the desired molecule is present in the list. This can be done by using the elimination of the Z-Conjunction rule given in Definition 1.

The formal definitions, presented in this section, allow us to recursively check all the possible combinations of the initial molecules against the first elements of given EVFs. In case of a match, the corresponding EVF can be applied and the process can restart again to find other possible matches from the new list of molecules. This process terminates when no more molecules are found to be reacting with each other and at this point we will have the list of post-reaction molecules. The desired result can then be obtained from these molecules using the elimination of Z-Conjunction rule. The main benefit of the development, presented in this section, is that it facilitates automated reasoning about the molecular biological experiments within the sound core of a theorem, which will be demonstrated later.

5. Formal Verification of Zsyntax Properties. In order to ensure the correctness and soundness of our definitions, we use them to verify a couple of properties representing the most important characteristics of molecular reactions. The first property deals with the case when there is no combination of reacting molecules in the list of molecules and in this case we verify that after the Zsyntax based experiment execution both the pre and post-experiment lists of molecules are the same. The second property captures the behaviour of the scenario when the given list of molecules contains only one set of reacting molecules and in this case we verify that after the Zsyntax based experiment execution the post-experiment list of molecules contains the product of the reacting molecules minus its reactants along with the remaining molecules provided initially. We represent these scenarios as formally specified properties in higher-order logic using our formal definitions, given in the previous section. These properties are then formally verified in HOL4.

5.1. Scenario 1: No Reaction. We verify the following theorem for the first scenario:

Theorem 1.

$$\begin{aligned} &\vdash \forall E L. \\ &\quad \sim(\text{NULL } E) \wedge \sim(\text{NULL } L) \wedge \\ &\quad (\forall a m n. \text{MEM } a E \wedge m < \text{LENGTH } L \wedge n < \text{LENGTH } L \\ &\quad \quad \Rightarrow \sim\text{MEM } (\text{FST } a) [\text{HD } (\text{z_conj_int } L m n)]) \\ &\quad \Rightarrow \text{z_deduction } L E = L \end{aligned}$$

The variables E and L represent the lists of EVFs and molecules, respectively. The first two assumptions ensure that both of these lists have to be non-empty, which are the pre-conditions for a molecular reaction to take place. The next conjunct in the assumption list of Theorem 1 represents the formalization of the no-reaction-possibility condition as according to this condition no first element of any pair in the list of EVFs E is a member of the head of the list formed by the function `z_conj_int`, which picks the elements corresponding to the two given indices (that range over the complete length of the list of molecules L) and appends them as a flattened single element on the given list L . This constraint is quantified for all variables a , m and n and thus ensures that no combination of molecules in the list L matches any one of the first elements of the EVF list E . Thus, under this constraint, no reaction can take place for the given lists L and E . The conclusion of Theorem 1 represents the scenario that the output of our formalization of Zsyntax based reaction would not make any change in the given molecule list L and thus verifies that under the no-reaction-possibility condition our formalization also did not update the molecule list.

The verification of this theorem is interactively done by ensuring the no-update scenario for all molecule manipulation functions, i.e., `z_EVF`, `z_recur1`, `z_recur2` and `z_deduction_recur`, under the no-reaction-possibility condition [1]. For example, the corresponding theorem for `z_EVF` function is as follows:

Theorem 2.

$$\begin{aligned} &\vdash \forall E L m n P. \\ &\quad \sim(\text{NULL } E) \wedge \sim(\text{NULL } L) \wedge m < \text{LENGTH } L \wedge n < \text{LENGTH } L \wedge \\ &\quad P < \text{LENGTH } E \wedge (\forall a. \text{MEM } a E \Rightarrow \sim\text{MEM } (\text{FST } a) [\text{HD } L]) \\ &\quad \Rightarrow \text{z_EVF } L E P m n = (\text{F}, \text{TL } L) \end{aligned}$$

The assumptions of above theorem ensure that both lists L and E are not empty and the arguments of the function `z_EVF` are bounded by the `LENGTH` of L and E . The last conjunct in the assumption list models the no-reaction-possibility condition in the context of the function `z_EVF`. The conclusion of the theorem states that no update takes place under the given conditions by ensuring that the function `z_EVF` returns a pair with the first element being `F` (False), representing no match, and the second element being equal to `TL L`, which is actually equal to the original list L since an element was appended on head of L by the parent function.

5.2. Scenario 2: Single Reaction. The second scenario complements the first scenario and, caters for the case when a reaction is possible and we verify that the molecules list is indeed updated based on the outcomes of that reaction. In order to be able to track the reaction and the corresponding update, we limit ourselves to only one reaction in this scenario but since we verify a generic theorem (universally quantified) for all possibilities our result can be extended to cater for multiple reactions as well. The theorem corresponding to this scenario 2 is as follows:

Theorem 3.

$$\begin{aligned} & \vdash \forall E L z m' n'. \\ & \sim\text{NULL } E \wedge \sim\text{NULL } (\text{SND } (EL z E)) \wedge 1 < \text{LENGTH } L \wedge m' < \text{LENGTH } L \wedge \\ & n' < \text{LENGTH } L \wedge z < \text{LENGTH } E \wedge \text{ALL_DISTINCT } (L ++ \text{SND } (EL z E)) \wedge \\ & (\forall a b. a \neq b \Rightarrow \text{FST } (EL a E) \neq \text{FST } (EL b E)) \wedge m' \neq n' \wedge \\ & (\forall K m n. m < \text{LENGTH } K \wedge n < \text{LENGTH } K \wedge \\ & (\forall j. \text{MEM } j K \Rightarrow \text{MEM } j L \vee \exists q. \text{MEM } q E \wedge \text{MEM } j (\text{SND } q)) \Rightarrow \\ & \quad \text{if } (EL m K = EL m' L) \wedge (EL n K = EL n' L) \\ & \quad \text{then HD } (z_conj_int K m n) = \text{FST } (EL z E) \\ & \quad \text{else } \forall a. \text{MEM } a E \Rightarrow \text{FST } a \neq \text{HD } (z_conj_int K m n)) \\ & \Rightarrow z_deduction L E = z_del (L ++ \text{SND } (EL z E)) m' n' \end{aligned}$$

The first two assumptions ensure that neither the list E , i.e., the list of EVFs, nor the second element of the pair at index z of the list E is empty. Similarly, the third assumption ensures that the list L , i.e., the list of initial molecules, contains at least two elements. These constraints ensure that we can have at least one reaction with the resultant being available at index z of the EVF list. The next four assumptions ensure that the indices m' and n' are distinct and these along with the index z fall within the range of elements of their respective lists of molecules L or EVFs E . According to the next assumption, i.e., $\text{ALL_DISTINCT } (L ++ \text{SND } (EL z E))$, all elements of the list L and the resulting molecules of the EVF at index z are distinct, i.e., no molecule can be found two or more times in the initial list L or the post-reaction list E . The next assumption, i.e., $(\forall a b. a \neq b \Rightarrow \text{FST } (EL a E) \neq \text{FST } (EL b E))$, guarantees that all first elements of the pairs in list E are also distinct. Note that this is different from the previous condition since the list E contains pairs as elements and the uniqueness of the pairs does not ensure the uniqueness of its first elements. The final condition models the presence of only one pair of reactants scenario. According to the assumptions of this implication condition, the variable K is used to represent a list that only has elements from list L or the second elements of the pairs in list E . Thus, it models the molecules list in a live experiment. Moreover, the variables m and n represent the indices of the list K and thus they must have a value less than the total elements in the list K (since the first element is indexed 0 in the HOL4 formalization of lists). Now, if the indices m and n become equal to m' and n' , respectively, then the head element of the $z_conj_int K m n$ would be equal to FST of $EL z E$. Otherwise, for all other values of indices m and n , no combination of molecules obtained by $\text{HD}(z_conj_int K m n)$ would be equal to the first element of any pair of the list E . Thus, the if case ensures that the variables m' and n' point to the reacting molecules in the list of molecules L and the variable z points to their corresponding resultant molecule in the EVF list. Moreover, the else case ensures that there is only one set of reacting molecules in the list L . The conclusion of the theorem formally describes the scenario when the resulting element, available at the location z of the EVF list, is appended to the list of molecules while the elements available at the indices m' and n' of L are removed during the execution of the function $z_deduction$ on the given lists L and E .

The proof of Theorem 3 is based on verifying sub-goals corresponding to this scenario for all the sub-functions, i.e., z_EVF , z_recur1 , z_recur2 and $z_deduction_recur$. The formal reasoning for all of these proofs involved various properties of the del function for a list element and some of the key theorems developed for this purpose in our development are given in Table 5.1 and more details can be found in [1].

The formalization described in this section consumed about 500 man hours and approximately 2000 lines of HOL4 code, mainly due to the undecidable nature of higher-order logic. However, this effort raises the confidence level on the correctness of our formalization of Zsyntax. This fact distinguishes our work from all the other formal methods based techniques used in the context of BRNs, where the deduction rules are applied without being formally checked. Moreover, our formally verified theorems can also be used in the formal analysis of molecular pathways. The assumptions of these theorems provide very useful insights about the constraints under which a reaction or no reaction would take place. To the best of our knowledge, this is the first time that properties, like Theorems 1 and 3, about a molecular pathway experiment have been formally verified. Thus, the identification of these properties and their formal verification both constitute contributions of this paper.

As part of this work, we also developed a simplifier Z_SYNTAX_SIMP [1] that simplifies the proof with a single iteration of the function $z_deduction_recur$ and works very efficiently with the proofs involving our functions. The proof steps can be completely automated and the proof can be done in one step as well. However, we have kept the reasoning process manual purposefully as this way users can observe the status of the reaction

TABLE 5.1
Formally verified Properties of the *del* Function

Signature	Theorem
<code>del_ASSOC_THM</code>	$\vdash \forall L E m. m < \text{LENGTH } L$ $\Rightarrow \text{del } (L ++ E) m = \text{del } L m ++ E$
<code>del_LENGTH_THM</code>	$\vdash \forall L E m. m < \text{LENGTH } L$ $\Rightarrow \text{LENGTH } (\text{del } L m) = \text{LENGTH } L - 1$
<code>del_EL_THM</code>	$\vdash \forall L m n. m < n \wedge n < \text{LENGTH } L \wedge 1 < \text{LENGTH } L$ $\Rightarrow \text{EL } m L = \text{EL } m (\text{del } L n)$
<code>del_DISTINCT_THM</code>	$\vdash \forall L n. n < \text{LENGTH } L \wedge \text{ALL_DISTINCT } L$ $\Rightarrow \text{ALL_DISTINCT } (\text{del } L n)$
<code>del_MEM_THM</code>	$\vdash \forall L a m. m < \text{LENGTH } L \wedge \text{MEM } a (\text{del } L m)$ $\Rightarrow \text{MEM } a L$
<code>del_NOT_MEM_THM</code>	$\vdash \forall L m. \text{ALL_DISTINCT } L \wedge m < \text{LENGTH } L$ $\Rightarrow \sim \text{MEM } (\text{EL } m L) (\text{del } L m)$

at every iteration, which is a very useful feature to get an insight of what is happening inside a reaction. Each application of `Z_SYNTAX_SIMP` on the reaction, depicted in Fig. 7.1, would result in moving from a state n to $n + 1$.

6. Graphical User Interface. The interest of using formal methods in the domains of bio-informatics and system biology has been rapidly growing since the last decade. However, there is a notable gap between biologists, who prefer wet lab experiments, and computer scientists, who can provide efficient tools to perform computer experiments with a unified framework. This fact has somewhat limited the usage of formal methods by the bio-informatics and pharmaceutical industry in research and development. We believe that one of the major reasons for the above-mentioned gap is the unavailability of biologists friendly formal tools. The idea on which our GUI functions is that the user needs to provide the initial list of molecules and possible EVFs. Consequently, the verification relating the two inputs can be done automatically. The GUI, developed in C#, is capable of initiating a HOL4 session in the command line and automatically converting user provided data into the language supported by HOL4. These translated texts are then utilized to form a HOL 4 theorem, which is then passed to HOL4 via the command line and the verification process automatically starts using the formal definitions and tactics developed in our work. Figure 6.1 provides a snapshot of our GUI while handling the example of Glycolysis pathway, which will be described in more detail in the next section. Many real-world

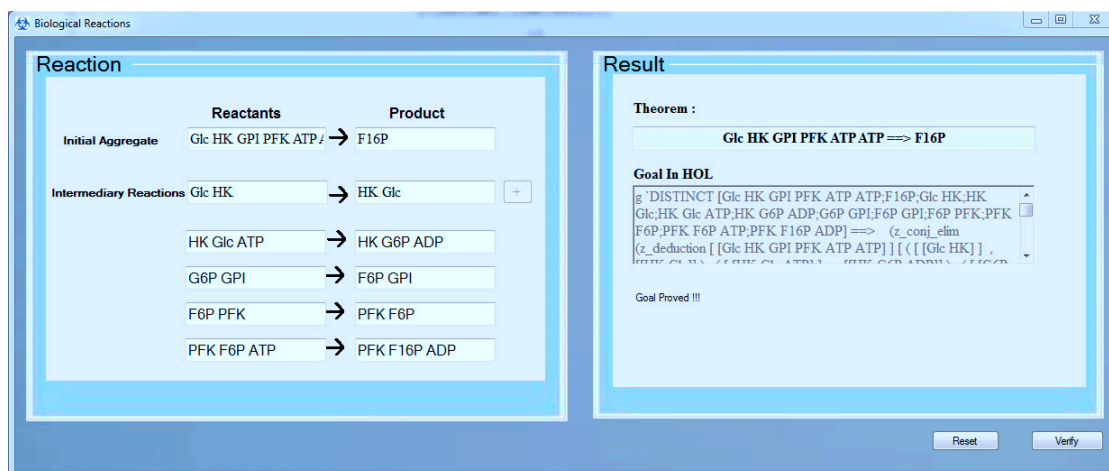


FIG. 6.1. Graphical User Interface

case studies have been automatically verified, including the examples that are described in the next section, by our GUI. Upon the successful verification of a theorem the GUI returns a message that the given pathway has been successfully deduced. The current version of our GUI is available for download at [1] and we are working to further enhance its usability by providing graphical insights about the verification.

7. Case Studies. In the section, we present the utilization of our Zsyntax formalization and its corresponding properties by two real-world cases studies: 1) Reaction involving TP53 degradation. 2) Molecular pathway of Glycolysis which is responsible for the production of Pyruvate and also provides energy for cell growth.

7.1. TP53 Degradation. TP53 gene encodes p53 protein, which plays a crucial role in regulating the cell cycle of multicellular organisms and, thus, functions as a tumour suppressor for preventing cancer [4]. We utilize the proposed framework to analyse the TP53 degradation reaction [4]. The regulatory loop leading to the TP53 degradation is illustrated in Fig. 7.1, where the dark coloured circles denote the list of molecules given by biologists and the hollow circles depict the molecules that do not interact with any other molecule in a particular step of the reaction.

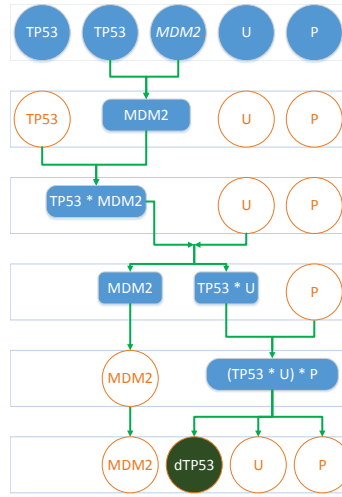


FIG. 7.1. Reaction Representing Degradation of TP53

A theorem representing the regulatory loop involving MDM2, $MDM2$ and TP53 and leading to TP53 degradation [4], can be described as follows:

$$TP53 \ \& \ TP53 \ \& \ MDM2 \ \& \ U \ \& \ P \vdash d(TP53)$$

In HOL4, this biological reaction can be written as the following theorem using our reported framework:

```

⊢ DISTINCT [TP53;dTP53;P;U;iMDM2;MDM2] ⇒
z_conj_elim (z_deduction [[TP53];[TP53];[iMDM2];[U];[P]]
  ([[TP53;MDM2],[[MDM2]]];
  ([MDM2;TP53],[[TP53;MDM2]]);
  ([TP53;MDM2;U],[[TP53;U];[MDM2]]);
  ([TP53;U;P],[[dTP53];[U];[P]])) [dTP53]
= [[dTP53]]

```

The DISTINCT function used in the assumption of the above theorem ensures that all the molecules are distinct. The first list argument of the function `z_deduction` is the initial aggregate (IA) of molecules that are available for reaction and the second list argument of the function `z_deduction` represents the valid EVFs for this reaction. Thus, the function `z_deduction` would deduce the final list of molecules under these particular conditions. The function `z_conj_elim` will return the molecule `dTP53` if it is present in the post-reaction list of molecules, as previously described.

Figure 7.1 shows the step-wise reaction leading to $d(\text{TP53})$. The blue-coloured circles show the chemical interactions and green colour represents the desired product in the pathway, whereas each rectangle shows the total number of molecules in the reaction at a given time. It is obvious from the figure that whenever a reaction yields a product, the reactants get consumed (no longer remain in the list) hence satisfying the stoichiometry of a reaction.

The script of TP53 Degradation Theorem is given below:

```
e(RW_TAC std_ss [DISTINCT, UNIQ_MEM, MEM]);
e(Z_SYNTAX_SIMP z_EVF_def );;
e(Z_SYNTAX_SIMP z_EVF_def );;
e(Z_SYNTAX_SIMP z_EVF_def );;
e(Z_SYNTAX_SIMP z_EVF_def );;
e(Z_SYNTAX_SIMP z_EVF_def );;
e(Z_SYNTAX_SIMP z_EVF_def );;
e(REWRITE_TAC [MEM, z_conj_elim_def]);
```

It is quite evident from the proof script of Theorem 1 that its proof steps can be completely automated and the proof can be done in one step as well. However, we have kept the reasoning process manual purposefully as this way the users can observe the status of the reaction at every iteration. For example, each application of `Z_SYNTAX_SIMP` on the reaction, depicted in Fig. 7.1, would result in moving from a state n to $n + 1$.

Note that we can also record the time required to complete each iteration. For example, in case of above examples, the time for each step in the above case-study is given in Table 7.1.

TABLE 7.1
Runtime per Iteration

Iteration	Duration (Seconds)
1 \rightarrow 2	95.969
2 \rightarrow 3	51.829
3 \rightarrow 4	27.127
4 \rightarrow 5	28.062
5 \rightarrow 6	0.2130

7.2. Molecular Pathway of Glycolysis. Glycolysis is a metabolic pathway which converts glucose into pyruvate and energy released in this process is used for the cell growth [18]. In molecular biology literature, Glycolysis is a determined sequence of ten enzyme-catalyzed reactions. Formation of Glucose-6-phosphate, Fructose-1,6-bisphosphate (F1,6P), etc. are intermediate steps in glycolysis [18]. In order to deduce the complete Glycolysis pathway, it is convenient to divide the pathway in three blocks as shown in Fig. 7.2. All the abbreviations used in Fig. 7.2 are described in Table 7.5. The dark coloured circles represent enzymes and each of them participates in the reaction at a certain point and only helps in the formation of new molecules. It is important to note that enzymes do not get consumed, which means that they retain their state after the completion of the reaction.

The first phase of the pathway leads to formation of Fructose-1,6-bisphosphate (F1,6P). The theorem representing the reaction of the glycolytic pathway leading from D-Glucose to F1,6P [4] can be described in classical Zsyntax format as follows:

$$\text{Glc} \ \& \ \text{HK} \ \& \ \text{GPI} \ \& \ \text{PFK} \ \& \ \text{ATP} \ \& \ \text{ATP} \vdash \text{F1,6P}$$

We can formally encode this theorem in HOL4 as follows:

```
⊢ DISTINCT [Glc; HK; GPI; PFK; ATP; ADP; G6P; F6P; F16P] ⇒
(z_conj_elim (z_deduction [[Glc]; [HK]; [GPI]; [PFK]; [ATP]; [ATP]]
  [[Glc; HK], [[HK; Glc]];
   ([HK; Glc; ATP], [[HK]; [G6P]; [ADP]]);
   ([G6P; GPI], [[F6P]; [GPI]]);
   ([F6P; PFK], [[PFK; F6P]]);
   ([PFK; F6P; ATP], [[PFK]; [F16P]; [ADP]])]) [F16P]
= [[F16P]]
```

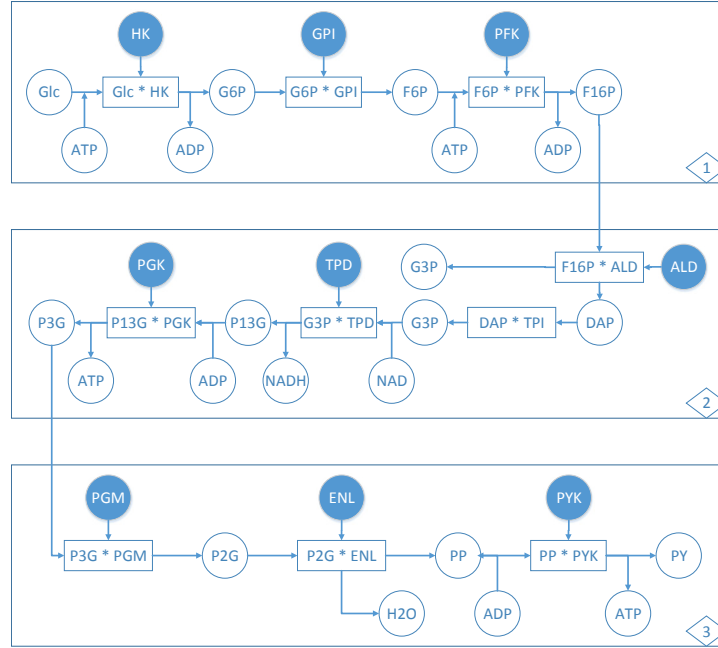


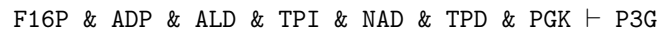
FIG. 7.2. Formation of Pyruvate through Glycolysis

where the first list argument of the function `z_deduction` is the initial aggregate (IA) of molecules that are available for reaction and the second list represents the valid EVFs for this reaction. The EVFs mentioned in the form of pairs and involving the molecules (G6P, F6P, etc.) are obtained from wet lab experiments, as reported in [4]. The `DISTINCT` function used above makes sure that all molecules (from initial aggregate and EVFs) used in this theorem are indeed distinct. Thus, the function `z_deduction` would deduce the final list of molecules under these particular conditions. The function `z_conj_elim` returns the molecule F1,6P if it is present in the post-reaction list of molecules. The verification time required for each iteration step is given in Table 7.2.

TABLE 7.2
Runtime per Iteration

Iteration	Duration (Seconds)
1 → 2	11.996
2 → 3	7.376
3 → 4	12.964
4 → 5	12.756
5 → 6	9.240
6 → 7	0.048

Fructose-1,6-bisphosphate formed in the above reaction further reacts with different enzymes to form 3-Phosphoglycerate, which completes the second phase of Glycolysis pathway. Note that during this phase two molecules of G3P are formed and each molecule follows the same path. For the sake of simplicity, we only provided one NAD molecule so that only one molecule can complete the pathway to form one Pyruvate molecule. After the successful deduction of first phase of Glycolysis, we can model the second phase as follow:



which can be formalized in HOL4 as follows:

```

⊢ DISTINCT [F16P; ALD; TPI; NAD; TPD; PGK; G3P;
            DAP; P13G; P3G; NADH; ADP; ATP] ⇒
(z_conj_elim (z_deduction [[F16P]; [ADP]; [ALD]; [TPI]; [NAD]; [TPD]; [PGK]]
              [( [F16P; ALD], [G3P]; [DAP] ] );
             ([DAP; TPI], [G3P]; [TPI]));
            ([G3P; TPD], [TPD; G3P]);
            ([TPD; G3P; NAD], [P13G]; [TPD]; [NADH]);
            ([P13G; PGK], [PGK; P13G]);
            ([PGK; P13G; ADP], [PGK]; [P3G]; [ATP])) ) [P3G]
= [[P3G]]

```

The time required for each iteration of the second phase of Glycolysis is given in Table 7.3.

TABLE 7.3
Runtime per Iteration

Iteration	Duration (Seconds)
1 → 2	24.940
2 → 3	20.532
3 → 4	22.676
4 → 5	16.804
5 → 6	20.780
6 → 7	17.120
7 → 8	0.188

3-Phosphoglycerate (P3G) formed in the above step further reacts with other enzymes to yield the final product of Glycolysis pathway, i.e., Pyruvate. A theorem representing the conversion of 3-Phosphoglycerate (P3G) to Pyruvate (PY) can be described as follows:

P3G & ADP & PGM & ENL & PYK ⊢ PY

Consequently, the corresponding HOL4 expression is given as follows:

```

⊢ DISTINCT [PGM; ENL; PYK; P3G; P2G; H2O; PP; PY; ADP] ⇒
(z_conj_elim (z_deduction [P3G]; [ADP]; [PGM]; [ENL]; [PYK]]
              [( [P3G; PGM], [P2G]; [PGM] ] );
             ([P2G; ENL], [H2O]; [PP]; [ENL]);
             ([PP; PYK], [PYK; PP]);
             ([PYK; PP; ADP], [PYK]; [PY]; [ATP])) ) [PY]
= [[PY]]

```

Similar to the above steps, the timing statistics for the above theorem are given in Table 7.4.

TABLE 7.4
Runtime per Iteration

Iteration	Duration (Seconds)
1 → 2	5.860
2 → 3	5.676
3 → 4	9.384
4 → 5	7.784
5 → 6	0.044

This completes the description of the utilization of our work to formally reason about two real-world case studies, i.e., TP53 degradation and molecular pathway of Glycolysis.

Our HOL4 proof script is available for download [1], and thus can be used for further developments and analysis of different molecular pathways. It is important to note that formalizing Zsyntax and then verifying its

TABLE 7.5
Abbreviations of molecules used in Glycolysis

Abbr.	Biological Entity	Abbr.	Biological Entity
Glc	Glucose	PYK	Pyruvate Kinase
HK	Hexokinase	G6P	Glucose-6-phosphate
ATP	Energy	F6P	Fructose-6-phosphate
GPI	Phosphoglucomutase	F16P	Fructose-1,6-diphosphate
PFK	Phosphofruktokinase	G3P	Glyceraldehyde-3-Phosphate
ALD	Aldolase	DAP	Dihydroxyacetonephosphate
TPI	Triosephosphateisomerase	P13G	1,3-biphosphoglycerate
TPD	Triosephosphate Dehydrogenase	P2G	2-phosphoglycerate
PGK	Phosphoglycerokinase	PP	Phosphoenolpyruvate
PGM	Phosphoglyceromutase	PY	Pyruvate
ENL	Enolase	P3G	3-Phosphoglycerate

properties was a very tedious effort. However, it took around 10 lines of code to formally define and verify each theorem related to the above case studies in HOL4, which clearly illustrates the usefulness of our foundational work. We have shown that our formalization is capable of modelling molecular reactions using Zsyntax inference rules, i.e., given a set of possible EVFs, our formalism can derive a final aggregate **B** from an initial aggregate **A** automatically. In case of a failure to deduce **B**, the proposed method still provides the biologist with all the intermediate steps so that one can examine the reaction in detail and figure out the possible cause of failure. The evident benefit of our reasoning approach is its automatic nature as the user does not need to think about the proof steps and which EVFs to apply where. However, the most useful benefit of the proposed approach is its accuracy as the theorems are being verified in a formal way using a sound theorem prover. Thus, there is no risk of human error or wrong application of EVFs. Finally, due to the computer-based analysis, the proposed approach is much more scalable than the paper-and-pencil based analysis presented in [4].

8. Conclusions. Most of the existing formal verification research related to molecular biology has been focussed on using model checking. As a complementary approach, the primary focus of the current paper is on using a theorem prover for reasoning about molecular pathways. The main strength of this approach, compared to existing model checking related work, is that the underlying methods and deduction rules can also be formally verified besides the verification of a particular molecular pathway case. Leveraging upon this strength, we formally verified two key behavioural properties of molecular pathways based on the Zsyntax language, which presents a deduction style formalism for molecular biology in the most biologist-centred way. Besides ensuring the correctness of our formalization of the Zsyntax operators and inference rules, the formally verified properties also play a vital role in reasoning about molecular pathways in the sound core of a theorem prover. The practical utilization and effectiveness of the proposed development has been shown by presenting the automatic analysis of reaction involving TP53 degradation and metabolic pathway known as Glycolysis.

The proposed work opens the doors to many new directions of research. Firstly, we are enhancing our GUI to add more biologist friendly features in it. Moreover, we are also targeting some larger case studies, such as Dysregulation of the cell cycle pathway during tumour progression [16] and Fanconi Anemia/Breast Cancer (FA/BRCA) pathway [26]. Another interesting future direction is to leverage upon the high expressiveness of higher-order-logic and utilize calculus and differential theoretic reasoning to add reaction kinetics support in our formalism.

REFERENCES

- [1] S. AHMAD. Formal Reasoning about Molecular Pathways - HOL Proof Script. <http://save.seecs.nust.edu.pk/projects/holsyntax/holzsyntax.html>, 2014.
- [2] S. AHMAD, O. HASAN, AND U. SIDDIQUE. Towards Formal Reasoning about Molecular Pathways in HOL. In *International Conference on Enabling Technologies: Infrastructure for Collaborative Enterprises*, pages 378–383. IEEE, 2014.

- [3] C. BAIER AND J. KATOEN. *Principles of Model Checking*. MIT Press, 2008.
- [4] G. BONIOLO, M. D'AGOSTINO, AND P. DI FIORE. Zsyntax: a Formal Language for Molecular Biology with Projected Applications in Text Mining and Biological Prediction. *PloS ONE*, 5(3):e9511–1–e9511–12, 2010.
- [5] A. CHURCH. A Formulation of the Simple Theory of Types. *Journal of Symbolic Logic*, 5(2):56–68, 1940.
- [6] V. DANOS AND C. LANEVE. Formal Molecular Biology. *Theoretical Computer Science*, 325(1):69–110, 2004.
- [7] N.H. HUNT ET AL. Immunopathogenesis of Cerebral Malaria. *International Journal for Parasitology*, 36(5):569–582, 2006.
- [8] L. TRILLING FAB. CORBLIN, E. FANCHON. Applications of a Formal Approach to Decipher Discrete Genetic Networks. *BMC Bioinformatics*, 11(1):385, 2010.
- [9] W. FONTANA. Systems Biology, Models, and Concurrency. *SIGPLAN Notices*, 43(1):1–2, January 2008.
- [10] F. CASSEZ ET AL. G.BERNOT. Semantics of Biological Regulatory Networks. *Electronic Notes Theoretical Computer Science*, 180(3):3–14, 2007.
- [11] PETER J. E. GOSS AND J. PECCOUD. Quantitative Modeling of Stochastic Systems in Molecular Biology by using Stochastic Petri Nets. *Proceedings of the National Academy of Sciences*, 95(12):6750–6755, 1998.
- [12] J. HARRISON. *Handbook of Practical Logic and Automated Reasoning*. Cambridge University Press, 2009.
- [13] K. HIRAYAMA. Genetic Factors Associated with Development of Cerebral Malaria and Fibrotic Schistosomiasis. *Korean J. Parasitol*, 40(4):165–172, Dec 2002.
- [14] M. MAGNIN L. PAULEVÉ AND O. ROUX. Abstract Interpretation of Dynamics of Biological Regulatory Networks. *Electronic Notes Theoretical Computer Science*, 272(0):43–56, 2011.
- [15] C.J. LANGMEAD. Generalized Queries and Bayesian Statistical Model Checking in Dynamic Bayesian Networks: Application to Personalized Medicine. In *Proc. International Conference on Computational Systems Bioinformatics*, pages 201–212, 2009.
- [16] R. MAGLIETTA, V. LIUZZI, E. CATTANEO, E. LACZKO, A. PIEPOLI, A. PANZA, M. CARELLA, O. PALUMBO, T. STAIANO, F. BUFFOLI, A. ANDRIULLI, G. MARRA, AND N. ANCONA. Molecular Pathways Undergoing Dramatic Transcriptomic Changes During Tumor Development in the Human Colon. *BMC Cancer*, 12(1):608, 2012.
- [17] R. MILNER. A Theory of Type Polymorphism in Programming. *Journal of Computer and System Sciences*, 17(3):348–375, 1977.
- [18] D. NELSON. *Lehninger Principles of Biochemistry*. W.H. Freeman, New York, 2008.
- [19] L.C. PAULSON. *ML for the Working Programmer*. Cambridge University Press, 1996.
- [20] R. PELÁNEK. Fighting State Space Explosion: Review and Evaluation. In *Formal Methods for Industrial Critical Systems*, volume 5596 of *Lecture Notes in Computer Science*, pages 37–52. Springer, 2008.
- [21] J. POSPCHAL AND V. KVASNIKA. Reaction Graphs and a Construction of Reaction Networks. *Theoretica Chimica Acta*, 76(6):423–435, 1990.
- [22] A. REGEV AND E. SHAPIRO. Cells as Computation. *Nature*, 419:343, 2002.
- [23] A. REGEV AND E. SHAPIRO. The π -Calculus as an Abstraction for Biomolecular Systems. In *Modelling in Molecular Biology*, Natural Computing Series, pages 219–266. Springer, 2004.
- [24] A. REGEV, W. SILVERMAN, AND E. Y. SHAPIRO. Representation and Simulation of Biochemical Processes Using the pi-Calculus Process Algebra. In *Pacific Symposium on Biocomputing*, pages 459–470, 2001.
- [25] M. RIZZOTTI AND A. ZANARDO. Axiomatization of Genetics. 1. Biological Meaning. *Journal of Theoretical Biology*, 118(1):61–71, 1986.
- [26] A. RODRÍGUEZ, D. SOSA, L. TORRES, B. MOLINA, S. FRÍAS, AND L. MENDOZA. A Boolean Network Model of the FA/BRCA Pathway. *Bioinformatics*, 28(6):858–866, 2012.
- [27] L. THOMAS AND R. D'ARI. *Biological Feedback*. CRC Press, USA, 1990.
- [28] R. THOMAS. *Kinetic Logic: A Boolean Approach to the Analysis of Complex Regulatory Systems*, volume 29 *Lecture Notes in Biomathematics*. Springer-Verlag, 1979.
- [29] J.J TYSON, C.A. NAGY, AND B. NOVAK. The Dynamics of Cell Cycle Regulation. *Bioessays*, 24(12):1095–1109, 2002.
- [30] O. WOLKENHAUER, D. SHIBATA, AND M.D. MESAROVIC. The Role of Theorem Proving in Systems Biology. *Journal of Theoretical Biology*, 300(0):57–61, 2012.
- [31] J.H. WOODGER, A. TARSKI, AND W.F. FLOYD. *The Axiomatic Method in Biology*. The University Press, 1937.
- [32] A. ZANARDO AND M. RIZZOTTI. Axiomatization of Genetics 2. Formal Development. *Journal of Theoretical Biology*, 118(2):145–152, 1986.

Edited by: Jesus Carretero

Received: September 14, 2014

Accepted: October 24, 2014