

Ontological integration of data models for cell signaling pathways

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Abstract

Databases have collected masses of information concerning cell signaling pathways that include information on pathways, molecular interactions as well as molecular complexes. However we have no general data model to represent comprehensive properties of cell signaling pathways, so that this type of information has been represented by two different data models that we call 'binary relation' and 'state transition'. The disagreement between the existing models derives from lack of consensus about a factor of causality in reactions in cell signaling pathways, which is often called 'signal'. We developed an ontology named CSNO (Cell Signaling Networks Ontology) based on device ontology. As device ontology is a research product of knowledge engineering, CSNO is the first application of it to biological knowledge. CSNO defines the factor of causality called 'signal', offers an integrative viewpoint for the two different data models, explicates intrinsic distinctions between signaling and metabolic pathways, and eliminates ambiguity from representation of complex molecules.

Keywords: device ontology, cell signaling pathway, database integration, role concept, metabolic pathway

1. Introduction

A cell signaling pathway (signaling pathway) represents a series of chemical reactions ending up in achieving cellular functions. Information on signaling pathways is crucial in functional genomics because it gives a clue on how to systematically connect genes or gene products to cellular functions. Databases have accumulated masses of data concerning signaling pathways that include pathways, molecular interactions as well as molecular complexes [1,2,6,9,11,13,15,16,20,21]. The data are collected either from literature or from large scale high throughput experiments. Speed of data accumulation has been further accelerated because of computerization of the data collecting procedures including information extraction and information retrieval [3,6,9,15].

As all the data collected in databases are descriptions of signaling phenomena at molecular level, we then have to proceed with systematical construction of functional information on the data, so as to reconstruct cellular phenomena in the computer. However, the data collected so far have been represented in two different models called 'binary relation' [1,11,13,16,20] and 'state transition' [2,5,21]. Binary relation model regards a signaling pathway as a series of molecular interactions, such as interactions between an enzyme and a substrate, a receptor and a ligand, an ion channel and an ion, or a transcription factor and DNA. The state transition model regards a signaling pathway as a series of state changes of molecules, such as change of chemical modification state, localization state, and state of complex represented by composition of the complex. Each model represents a different aspect of signaling pathways. The molecular interaction represents a requirement for starting a reaction. The state transition represents a result of a reaction on a molecule. However the problem how to integrate them into a general model is yet unsolved. It remains to be elucidated how the two aspects contribute to entire nature of signaling pathways. Such a general model will facilitate not only reusability of all the collected information but also monotonous increase of further collection of information on signaling pathways in an integrated database schema.

The disagreement between the data models arises from a lack of consensus about a factor of causality in reactions of signaling pathways. The factor is what is called 'signal'. While the two different models virtually represent concepts at 'biochemical level', an integrative model of them should be at a more abstract level, which provides us with a viewpoint for capturing cellular functions that a signaling pathway ends up in achieving, e. g., cell cycle, gene expression, and cytoskeletal rearrangement. Device ontology provides us with a framework to systematize concepts at such the abstract level [8]. Device ontology tells us that a function should be defined as an interpretation of 'B1 behavior' [8]. B1 behavior is a particular type of

behavior called 'B1' that yields the output from input on demand according to the specific context. The output in device ontology is something that becomes an input of another B1 behavior in succession. Thus B1 behavior represents an intrinsic property of causality in a sequence that finally achieves a purpose at its end. Definition of B1 behavior may enable us to define a factor of causality in reactions of signaling pathways and to assign an individual molecule a function having a meaning of how the individual molecule contributes to achievement of a cellular function.

We have developed an ontology for signaling pathways named CSNO (Cell Signaling Networks Ontology) based on device ontology [19]. CSNO provides us with a clear structured framework for systematic, consistent, and shareable description of signaling pathways. As device ontology is a research product of knowledge engineering, CSNO is the first application of it to biological knowledge. CSNO defines a factor of causality in reactions of signaling pathways, offers a module that integrates two existing different data models, explicates intrinsic distinctions between signaling and metabolic pathways, and eliminates ambiguity from representation of complex molecules.

2. Methods

Device ontology has been developed aiming at systematization of functional knowledge for design of artifacts in engineering [8]. Device ontology provides us with a consistent viewpoint that every constituents of the target phenomena founds on, so that we can systematically assign functions to the constituents. Device ontology specifies 'role concept' of the individual constituent. Role concept is a concept that is defined only in a certain context of the world. In comparison, 'basic concept' is a concept that is defined without premising any contexts. An instance of a basic concept can play a role represented by a role concept. A definition of role concept consists of inherited attributes from a basic concept and intrinsic attributes of the role. We take a role concept 'teacher role' as an example [10]. 'Teacher role' has 'name' attribute inherited from a basic concept 'human'. 'The subject that the teacher teaches' is an intrinsic attribute of 'teacher role'. Any role can be eliminated from the particular individual without causing any change on its being a basic concept that the role depends on. For example, even when Mr. Smith resigns a 'teacher role', he is still a 'human'. Thus definition of role concept specifies intrinsic properties of a certain context.

Device ontology specifies a physical phenomenon caused by device (agent), operand, medium, and B1 behavior and function of the device (Figure 1). It decomposes a target phenomenon into actions of devices on demand according to the specific context. Operand is defined as something that is processed by the device. Medium is defined as something that carries an operand.



Figure 1: Schematic representation of device ontology.

B1 behavior is defined as a change of attribute values of an operand when it is processed by a device. Function of device is defined as teleological interpretation of B1 behavior. Let us take a heat exchanger as an example [17] whose B1 behavior is 'to transfer' heat carried by the warmer stream (medium) to the colder stream (medium). Device ontology specifies the physical phenomenon as 'to warm' (function) is an interpretation of B1 behavior of the heat exchanger (device) in a context in which someone focuses on the colder stream and 'to cool' in a context in which someone focuses on the warmer stream. Although device ontology has been applied to artifact modeling in engineering, it affords a general framework for systematization of any physical phenomena in nature. This study shows our successful application of device ontology to knowledge of cellular biology.

Concerning the modeling of physical phenomena, there exist two major viewpoints: device-centered and process-centered views [14]. The device ontology specifies the former and the process ontology specifies the latter. The major difference between the two is that while device ontology has an agent (device) which is considered as something which plays the main actor role in obtaining the output, process ontology does not have such an agent but has participants which only participate in the phenomena being occurring. It is natural to apply process ontology to model chemical reactions. However we decided to apply device ontology to model the reactions in order to have a viewpoint in which a molecule is not a participant but an agent of a reaction, so that we can specify a function as an interpretation of B1 behavior of a molecule.

We refer to [18] for knowledge of 'TGF-beta pathway' and [12] for knowledge of 'B cell activation pathway'.

3. Results

3.1 Is-a hierarchy of CSNO

The skeleton of an ontology is an is-a hierarchy, which consists of is-a relations that represent super-sub (generalized-specialized) relations between concepts. Figure 2 shows is-a hierarchy of CSNO.

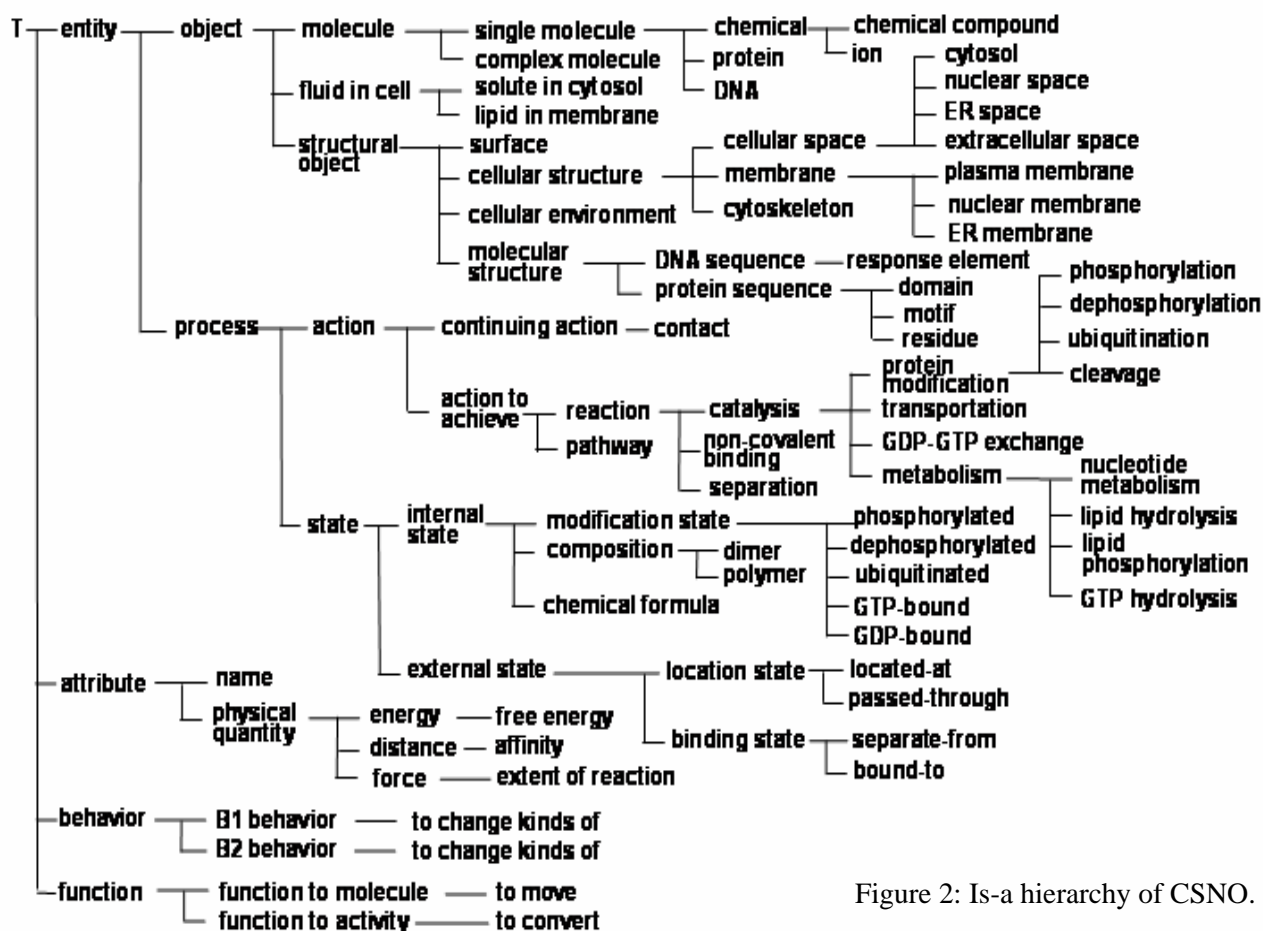


Figure 2: Is-a hierarchy of CSNO.

3.2 Definition of a factor of causality in reactions of signaling pathways

3.2.1 Definition of activity that is converted by a molecule

Any biological pathway is a causal sequence of chemical reactions. The causality is an intrinsic property of any pathway either in engineering or in natural science. In signaling pathways biologists assume 'signal' to be a factor of the causality. The causality in reactions yields a pathway that ends up in achieving a function. In case of a signaling pathway, a pathway ends up in achieving cellular functions, e.g. cell cycle, gene expression, and cytoskeletal rearrangement.

We firstly investigated a factor of causality in reactions without premising any biological contexts. Physical chemistry tells of two theories that 'free energy' of a reaction determines whether the reaction occurs spontaneously or not and that 'reaction kinetics' of reactions determines which reaction is most probable to occur in a certain condition [4]. A pathway in physical chemistry is a sequence of reactions most probably to occur in individual conditions changing progressively. Physical chemistry tells that not 'free energy' but 'reaction kinetics' determines a factor of causality. 'Affinity' is a physical quantity of reaction kinetics that determines the causality of a molecule of a certain state. 'Affinity' is obtained by differentiating 'free energy' with respect to 'extent of reaction' that is a kind of 'distance' in physical quantity. Because differentiation of 'energy' with respect to 'distance' yields 'force', 'affinity' is a kind of 'force'. Thus 'affinity' is a concept that represents 'a factor of causality in reactions of a pathway' in physical chemistry.

Then we investigate interpretation of 'affinity' in a context of signaling pathways. We found 'activity' is an interpretation of 'affinity'. While 'affinity' is a 'force' between molecules, 'activity' is a 'force' not only between molecules but also between a molecule and a reaction or a function. Device ontology tells

us that 'activity' is a role concept. It also tells us how to systematize underlying relations between the role and its basic concept 'affinity'. Figure 3 shows conceptual hierarchy of 'activity'. While there are various roles like a role dependent on a task or a role dependent on an action, 'activity' is a 'part-role concept' that depends on a part-whole relation [10]. Is-a hierarchy of part-role concepts can be defined only in an is-a hierarchy of whole concepts that the part-role concepts are parts of. In case of definition of 'activity', we firstly defined its whole concept 'reaction' and then defined 'activity' in the definition of 'reaction' (Figure 3). 'Activity' is specialized into 'bind activity', 'phosphorylate activity', 'be-phosphorylated activity', and so on.

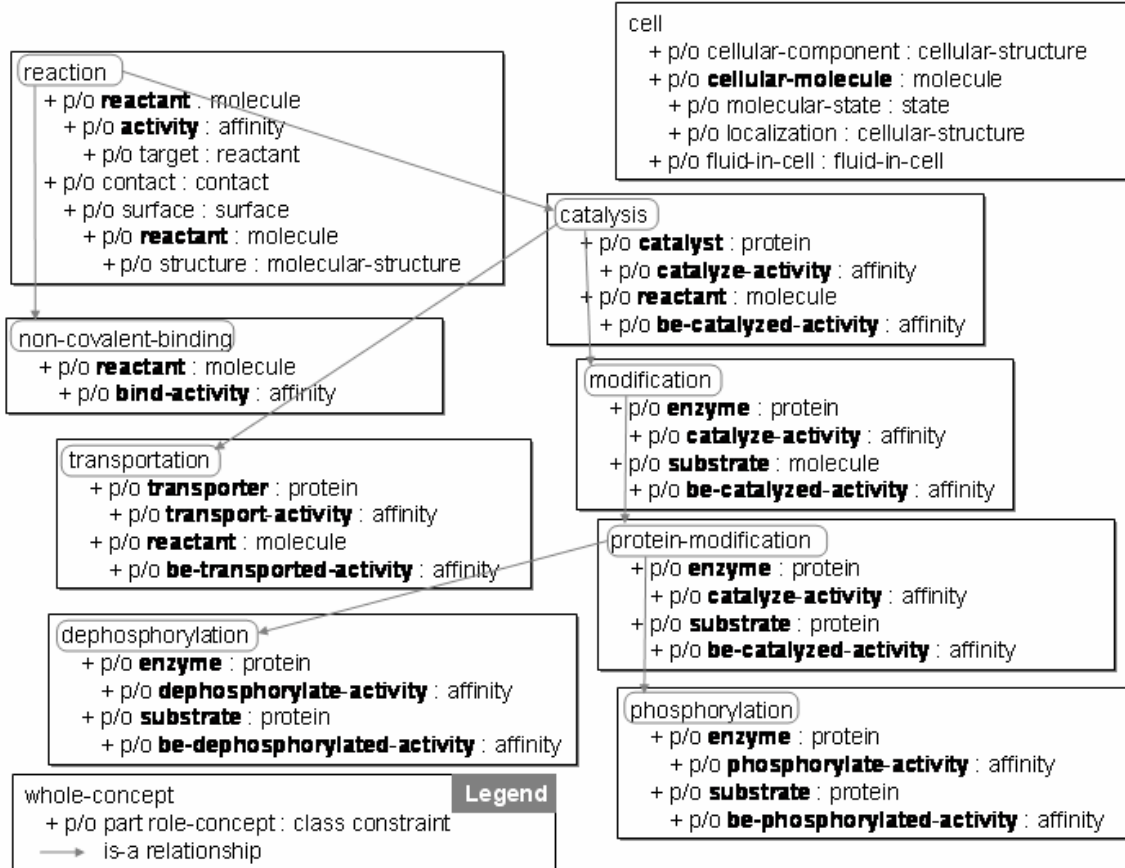


Figure 3: Conceptual hierarchy of 'activity' (portion) including part-whole relations between 'activity' and 'reactant' (part-role concepts) and 'reaction' (whole concept). This figure also shows part-whole relation between 'cellular-molecule' (part-role concept) and 'cell' (whole concept). Role concepts of interest are in bold.

'Activity' is an abstracted feature from all the reactions occurred in signaling pathways in a context of causality. Based on 'activity' we can represent all the reactions uniformly. Table 1 shows a definition of 'activity' based on device ontology. 'Activity' is defined as operand. We define 'reactant' as device. 'Reactant' is also a role concept that is an interpretation of basic concept 'molecule' in a context of 'being a participant of a reaction', which we will give detailed explanation in section 3.3. We define 'contact surface' between reactants as medium, according to a precedent study of device ontology by Kitamura and Mizoguchi [8]. The study found that contact surface of device can be medium when 'force' is operand. Because 'force' is transmitted by no other than a contact surface between devices, the contact surface is regarded as medium which carries the 'force'. 'Activity' is processed by device so as to be changed like that 'bind activity is changed into phosphorylate activity', which is observed in allosteric activation of a phosphorylating enzyme. Such the action 'to change kinds of activity' is occurred in every reactant yielding a pathway, so that we define 'to change kinds of' as B1 behavior (Figure 2). At last we define 'to convert' as function of 'reactant' whose behavior is 'to change kinds of activity'. This is the

Table 1: Definitions of 'to move' and 'to convert' functions in device ontology.

	Function 'to move'	Function 'to convert'
Device	Cellular environment	Reactant
Operand	Cellular-molecule	Activity
Medium	Fluid in cell	Contact Surface

which carries the 'force'. 'Activity' is processed by device so as to be changed like that 'bind activity is changed into phosphorylate activity', which is observed in allosteric activation of a phosphorylating enzyme. Such the action 'to change kinds of activity' is occurred in every reactant yielding a pathway, so that we define 'to change kinds of' as B1 behavior (Figure 2). At last we define 'to convert' as function of 'reactant' whose behavior is 'to change kinds of activity'. This is the

function to be assigned to individual molecules that cause a sequence ends up in achieving a cellular function.

3.2.2 Definition of movement of a molecule carrying activity

Although 'activity' is a factor of causality in reactions, it fails to explain localization changes of the reactions in a cell. Localizations of reactions are progressively changed from plasma membrane to mostly nucleus or to other organelles. A signaling pathway is then considered as orthogonal combination of a sequence of 'to convert' functions and a sequence of localization changes. Purpose of the former sequence is to finally achieve a certain cellular function. Purpose of the latter sequence is to bring the final reaction to a certain place in a cell where the target cellular function should occur. While the former sequence is considered as a main stream that is essential in finally achieving the target function, the latter sequence is considered as an auxiliary stream that assists the main stream to be in succession. The assistance of the auxiliary stream is indispensable because cells are compartmentalized into functional areas by organelles where only certain reactions can occur.

We define the localization change as B1 behavior based on device ontology (Table 1). We define 'cellular-molecule' as operand. 'Cellular-molecule' is a role concept that is an interpretation of a basic concept 'molecule' in a context of 'being a constituent of a cell', which we will give detailed explanation in section 3.3. 'Cellular environment' and 'fluid in cell' are defined as device and medium, respectively. Then we define 'to move' as function of 'cellular-molecule' whose behavior is 'to change kinds of localization'.

The two functions 'to convert' and 'to move' are orthogonal because each function has distinct agent and operand. Thus we finally define that **'a factor of causality in reactions in signaling pathways' is activity and cellular-molecule carrying the activity.**

We define that an active transportation across a membrane is not a 'to move' but a 'to convert' function, although the transportation contributes to a localization change. Since device of 'to move' function is 'cellular environment', the function allows only diffusions in an organelle. However the active transportation is a process that a device called 'channel' or 'transporter' lets a 'cellular-molecule' traverse between organelles. 'Transport activity' is a subsumption of 'catalyze activity' in CSNO (Figure 3).

3.3 A consistent viewpoint for capturing a complex molecule

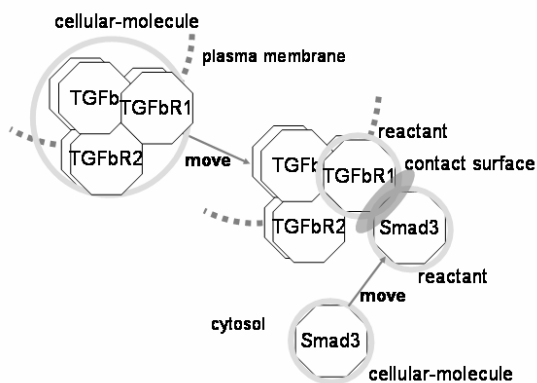


Figure 4: Schematic representation of 'cellular molecule' and 'reactant' roles held by a complex molecule (TGFb dimer, TGFbR1 dimer, and TGFbR2 dimer) and a single molecule (Smad3). This figure represents the same reaction explained in Figure 6. Please refer to Figure 6 for abbreviations.

In contrast to metabolic pathways composed of only modification reactions, signaling pathways include non-covalent binding as well as modification reactions. Because of that, complex molecules often participate in signaling pathways. It has been controversial among biologists whether a whole complex molecule or a component of the complex should be regarded as an agent. Either binary relation or state transition model has not answered this question.

CSNO answers the question as follows. CSNO explains us that a molecule has two roles: operand of 'to move' function and device of 'to convert' function. In case that operand of 'to move' function is a complex molecule, the complex molecule as a whole is regarded as the operand, since all the members of a complex molecule actually move at a time (Figure 4). This view emphasizes existence of the complex molecule as a constituent of a cell. Since this view represents a role of a molecule, we define 'cellular-molecule' as a role concept that is an interpretation

of a basic concept 'molecule' in the view (Figure 3). Cellular-molecule has an attribute 'localizations' intrinsic to the role and an attribute 'molecular state' inherited from the basic concept (Figure 3).

On the other hand, in case that device of 'to convert' function is a complex molecule, a component of the complex is regarded as the device, since the component actually converts 'activity' and also makes 'contact surface' (Figure 4). This view emphasizes participation of a molecule in a reaction. Since this view represents another role of a molecule, we define 'reactant' as a role concept that is an interpretation of a basic concept 'molecule' in the view (Figure 3). Reactant has an attribute 'activity' intrinsic to the role and an

attribute 'structure', like domain and motif, inherited from the basic concept (Figure 3).

In contrast to a complex molecule, a single molecule plays both roles of 'cellular-molecule' and 'reactant' as intact (Figure 4). Thus CSNO provides us with a consistent viewpoint for capturing complex molecules as well as single molecules so as to eliminate ambiguity in representation of them. CSNO also explains us that a complex molecule affords several reactants in it. Such a complex can accumulate several reactions in it so as to integrate signaling effects at a certain point in the pathway.

3.4 Definition of a module that integrates existing data models

When two molecules move and meet at a certain place in cells, a reaction occurs and an 'activity' is converted by an individual reactant (Figure 5A). A combination of orthogonal functions of 'to move' and 'to convert' determines a module for signaling pathways, which is a standardized and independent unit used in construction of the pathway. We name the module 'CSNO-module'. CSNO-module is defined as a simple combination of definitions of 'to move' and 'to convert' functions in device ontology (Figure 5B). CSNO-module includes B2 behavior which is usually disregarded in definitions of device ontology. B2 behavior represents an internal change of device, but it is not a factor related to causality because it does not yield any output [8]. In the context of signaling pathways, B2 behavior of reactant (device) is defined as a 'state' including 'modification state' and 'binding state' (Figure 2). In signaling pathways, instances of B2 behavior is commonly observed and described in biological literatures.

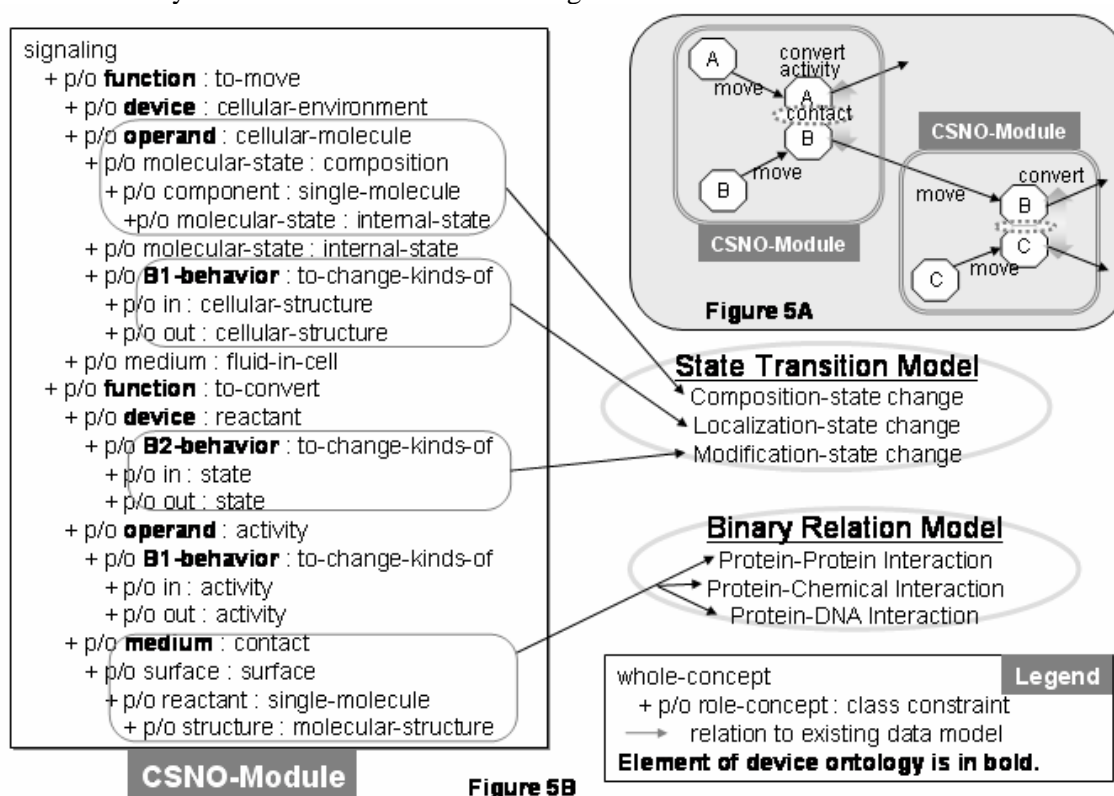


Figure 5: CSNO-module. Schematic representation of CSNO-module (Figure 5A). Definition of CSNO-module and relations between its attributes and exiting data models (Figure 5B).

CSNO-module integrates two existing models for signaling pathway; binary relation and state transition (Figure 5B). Operand of 'to move' function represents information equivalent to what state transition model represents as 'composition-state change'. B1 behavior of 'to move' function represents information equivalent to what state transition model represents as 'localization-state change'. B2 behavior of 'to convert' function represents information equivalent to what state transition model represents as 'modification-state change'. Medium of 'to convert' function represents information equivalent to what binary relation model represents as 'molecular interaction' that contains protein-protein, protein-chemical, and protein-DNA interactions. Thus CSNO-module represents comprehensive aspects of signaling pathways including not only aspects that have been modeled so far but also a new aspect introduced in this study which is an intrinsic factor for causality yielding a sequence to achieve a function.

3.5 Instances of CSNO-module

We take 'TGF-beta pathway' and 'B cell activation' as examples to show instances of CSNO-module. Figure 6 shows an instance that represents complex formation between TGF-beta receptor 1 (TGFbR1) and Smad3 occurred in 'TGF-beta pathway'. Operand of 'to move' function indicates that TGFbR1 constitutes a complex with TGF-beta and TGF-beta receptor 2 (TGFbR2). B1 behavior of 'to move' function indicates that the complex is located at plasma membrane. B1 behavior of 'to convert' function indicates that Smad3 produces a new 'activity' to 'be-phosphorylated' against TGFbR1. Medium of 'to convert' function indicates domain and motif structures at a contact surface between TGFbR1 and Smad3. B2 behavior of 'to convert' function indicates that state of TGFbR1 changes into 'bound-to state' targeting Smad3 as well as that state of Smad3 changes into 'bound-to state' targeting TGFbR1, which represents construction of a complex at the contact surface between TGFbR1 and Smad3.

Figure 7A shows an instance that represents a metabolic reaction in a signaling pathway 'B cell activation'. Even if it is a metabolic reaction, it is regarded as a 'to convert' function when it occurs in signaling pathways. As figure 7A shows, the reaction not only produces IP3 but also causes 'activity' to bind to a protein having 'IP3 binding' motif. Change of chemical formula from PIP(4,5)P2 to IP3 is indicated as B2 behavior of device 'phospholipid'.

A complete set of instances for 'TGF-beta pathway' and 'B cell activation' is not shown in this article but available from <http://athos.is.s.u-tokyo.ac.jp/CSNO/>.

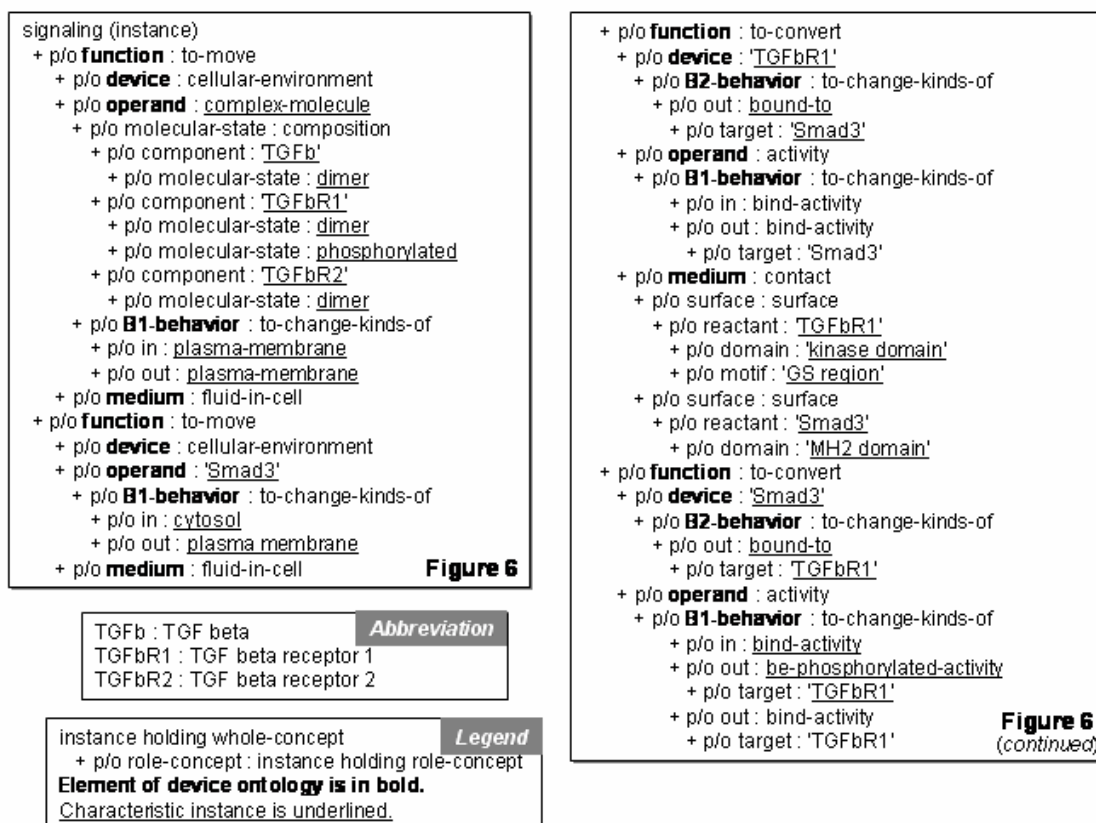


Figure 6: An instance of CSNO-module that represents a complex formation between TGF-beta receptor 1 and Smad3.

3.6 Intrinsic distinctions between signaling and metabolic pathways

When we represent metabolic pathways in device ontology, we find that metabolic pathways consist of sequences of 'to convert' functions without any interventions of 'to move' functions. Table 2 shows a definition of 'to convert' function in metabolic pathways based on device ontology. We define 'enzyme' as device. 'Enzyme' is a role concept that is an interpretation of a basic concept 'molecule' in a context of 'metabolism'. 'Chemical structure' and 'chemical compound' are defined as operand and medium, respectively. Intrinsic distinction between signaling and metabolic pathways is in their definitions of operand. In a signaling pathway, 'activity' and 'cellular-molecule' carrying the 'activity' are operands, which enable both progressive activations of reactions and progressive localization changes of the reactions. In contrast, in

Table 2: Definitions of 'to convert' functions in cell signaling and metabolic pathways in device ontology

	Cell signaling pathway	Metabolic pathway
Device	Reactant	Enzyme
Object	Activity	Chemical Structure
Medium	Contact Surface	Chemical Compound

point out that a signaling pathway may also control a metabolic pathway as another end point.

The same metabolic reactions producing IP3 from PIP(4,5)P2 occur both in signaling and metabolic pathways. When it occurs in a metabolic pathway, e.g. a pathway of further phosphorylation of IP3 in order to produce IP6P [7], change of chemical structure from PIP(4,5)P2 to IP3 is an intrinsic property (B1 behavior) of the reaction (Figure 7B). When it occurs in a signaling pathway, acquisition of 'bind activity' by phospholipid is an intrinsic property (B1 behavior) of the reaction (Figure 7A).

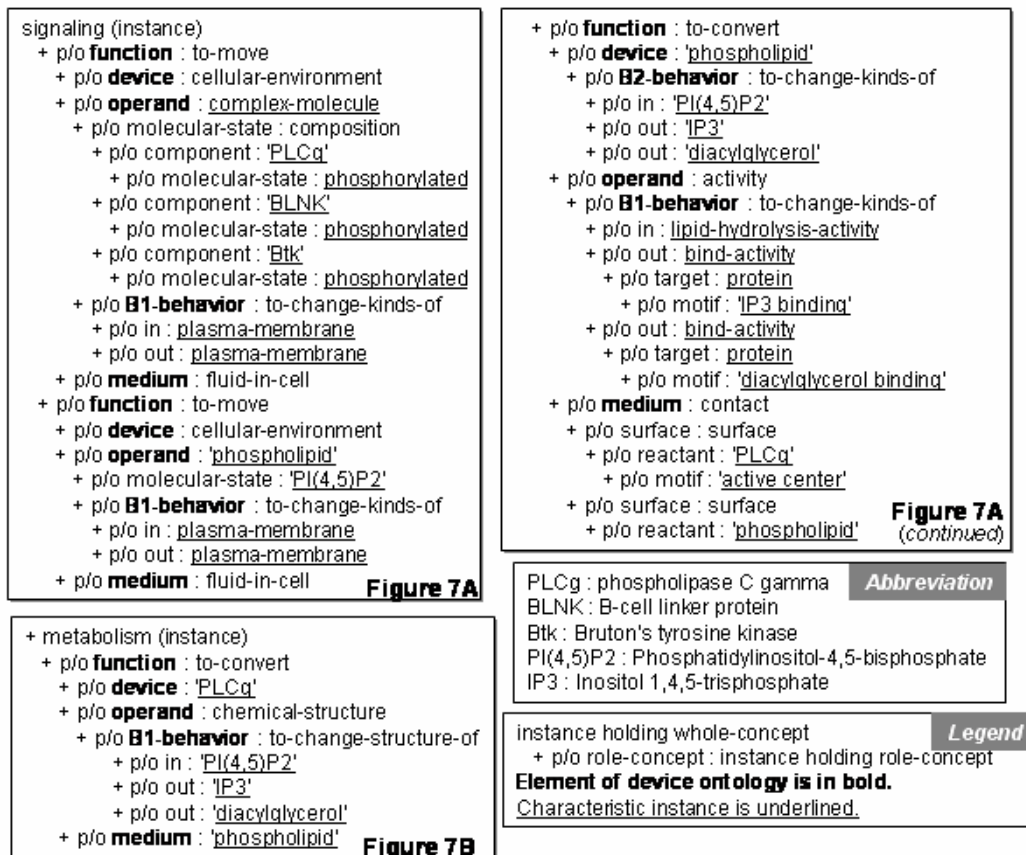


Figure 7: An instance of CSNO-module that represents production of IP3 from PI(4,5)P2 occurred in a signaling pathway (Figure 7A) as well as occurred in a metabolic pathway (Figure 7B).

4. Conclusions and discussion

Device ontology provides us with a viewpoint that a physical process is an action of an agent. Based on device ontology we define a signaling pathway as an orthogonal combination of a sequence of 'to move' functions and a sequence of 'to convert' functions. 'To move' function is an action of 'cellular environment' that moves a molecule named 'cellular-molecule' in cells. 'To convert' function is an action of a molecule named 'reactant' that converts 'activity' that is a factor of causality in reactions. In comparison, we define a metabolic pathway as a sequence which consists of only 'to convert' functions that convert chemical structures. The intrinsic difference between the two pathways depends on biological functions that the individual pathways finally achieve. A signaling pathway ends up in achieving cellular functions required in a cell. A metabolic pathway ends up in producing chemical compounds required in a cell. All the definitions

described in this article comprise CSNO. CSNO also provides us with a consistent viewpoint for capturing complex molecules so as to eliminate ambiguity in representation of them.

We define a module for signaling pathways based on CSNO. CSNO-module indicates how to integrate two existing data models; binary relation and state transition. We offer CSNO-module to the biological community as a general model to describe signaling pathways. To facilitate that, we will soon open an extensive collection of representative instances of CSNO-module to the public.

This study offers a comprehensive model to describe signaling pathways at molecular level. Next we have to proceed with systematization of functional information based on CSNO, so that we can finally reconstruct cellular phenomena in computers. Functional decomposition based on device ontology sheds light on the systematization of the functions, as our previous study has shown [19].

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