Specifications of \textit{insilicoML} 1.0: A Multilevel Biophysical Model Description Language

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Abstract: An extensible markup language format, \textit{insilicoML} (ISML), version 0.1, describing multilevel biophysical models has been developed and is available in the public domain. ISML is fully compatible with CellML 1.0, a model description standard developed by the IUPS Physiome Project, for enhancing knowledge integration and model sharing. This article illustrates the new specifications of ISML 1.0 that largely extend the capability of ISML 0.1. ISML 1.0 can describe various types of mathematical models, including ordinary/partial differential/difference equations representing the dynamics of physiological functions and the geometry of living organisms underlying the functions. ISML 1.0 describes a model using a set of functional elements (modules), each of which can specify mathematical expressions of the module functions. Structural and logical relationships between any two modules are specified by edges, which allow modular, hierarchical, and/or network representations of the model. The role of edge relationships is enriched by key words in order for use in constructing a physiological ontology. The ontology is further improved by the traceability of history of the model’s development and by linking between different ISML models stored in the model’s database using meta-information. ISML 1.0 is designed to operate with a model database and integrated environments for model development and simulations for knowledge integration and discovery.

Key words: model descriptive language, multiscale modeling, model sharing, physiome.

Physiome and systems biology are emerging research fields aimed at integrating vast stores of knowledge on human physiological and pathological functions at multiple levels and scales in time and space, from molecules and cells to individual organisms \cite{1,2}. Mathematical models, in particular dynamic system models of physiological functions, play a key role for integration, since they are capable of describing time evolution of biological system states quantitatively based on physical and chemical principles or phenomenological logic governing system behavior. However, the number of mathematical models of biological functions published in peer-reviewed journals and the complexity of each of these models rapidly increase as computational performance increases. This raises difficulties in reproducing simulated behaviors of the published models and in reuse of the models by third parties, thus hindering the promotion of sciences and knowledge integration.

Systems biology markup language (SBML) \cite{3} and CellML \cite{4} have promoted pioneering efforts to overcome this problem. These are extensible markup language (XML)-based formats aimed at describing mathematical models of biological functions, such as gene expression/regulation and electrical activities of cell membranes. Models written in SBML and CellML formats can be downloaded from their model repositories. They include all information necessary to reproduce simulated results described in the corresponding publication about each model. In parallel with these efforts, we have developed \textit{insilicoML} (ISML) 0.1 to describe multilevel biophysical models. ISML 0.1 is available in the public domain through a model database at \url{www.physiome.jp} \cite{5}. The format of ISML is defined according to W3C XML specifications \cite{6}. ISML is fully compatible with CellML 1.0 and possesses several features complementary to CellML for enhancing knowledge integration and model sharing.

This article illustrates the new specifications of ISML 1.0, which largely extend the capabilities of ISML 0.1.
ISML 1.0 can describe various types of mathematical models, including ordinary differential equations (ODEs), partial differential equations (PDEs), difference equations, and agent-based simulation models that utilize IF-THEN rules, among others. These models represent the dynamics of physiological functions and the geometry, i.e., morphometrics, of living organisms underlying the functions. In ISML, a model is described by a set of functional elements (modules), each of which specifies mathematical expressions of the module functions. In this article we focus on several types of specifications that characterize ISML 1.0 and offer examples of each. They include (1) specifications that allow the user to construct biophysical models with modular, hierarchical, and/or network representations; (2) specifications of how the morphology or geometry of biological entities are incorporated with dynamic system models such as PDEs and agent-based models; (3) specifications to provide a basis for model reuse and physiological model-ontology that support easy model construction, large-scale model construction and simulation, and knowledge discovery from the multilevel complex networks of the physiological functions.

First we illustrate the overall picture of ISML and related tools, applications, and databases before getting into the details of ISML 1.0. As in ISML 0.1, ISML 1.0 is designed to cooperate with model databases and integrated environments for model development and simulations (Fig. 1). The insilicoIDE (Integrated Development Environment; ISIDE) provided by our physiome project is such an environment, which is currently the most promising tool for the use of ISML. As shown in Fig. 1, a model written in ISML 1.0 format can be constructed using a model development and simulation environment provided by ISIDE. As later described in detail, ISML 1.0 describes a model using a set of functional elements referred to as “modules,” each of which can specify mathematical expressions of the module functions. Structural and logical relationships between modules are specified by the “edges,” which allow modular, hierarchical, and network representations of the model. Models in ISML are registered in our model database in the public domain for reuse. Another way to obtain ISML models is to convert models from the CellML 1.0 model repository by using ISIDE or other APIs. ISML models that require morphological and/or time series data such as PDEs and agent-based models cooperate with the databases of morphological models and time series data. Since ISML models are written in an XML-based format, they can be easily parsed and converted to other formats, such as CellML, SBML, C++, and MPI-C++, for simulations, i.e., numerical integration with respect to time and space.

For bifurcation analysis of nonlinear dynamical system models, ISML models can be converted to model description formats of existing application software such as xppaut [7] and BUNKI [8]. Several use-cases of ISML 1.0 and future plans for our platform with the use of ISML 1.0 are described and discussed for knowledge integration and discovery at the end of this article.

1. Overview of model representation in ISML

We consider a target biological system as an aggregate of elements referred to as “modules.” These modules are characterized by a name, physical quantities representing dynamic or static states with their mathematical implementations. These implementations specify the dynamics of how the states evolve in time and space and also the model geometry, such as morphological model (shape), posture, and position, among others. As illustrated in Fig. 2, modules can functionally affect each other by transmitting values of their physical quantities. The value of a module’s physical quantity goes out through an output “port” of the module and is transmitted to an input “port” of other modules. This functional relationship can be defined between any two modules with its direction, and it is specified by its departure (head) and destination (tail) of a functional “edge” linking two modules. The
edges can also represent structural and logical relationships between two modules. The edges with functional, structural, and logical (also forwarding and capsular) types allow the user to construct biophysical models with modular, hierarchical, and network representations. The modules can also import morphological data (numerical and analytical) and time-series data acquired by experiments in the real world and/or model simulations and can utilize them in the model. With this feature, ISML 1.0 allows the user to combine experimental, theoretical, and model-based research.

Another important idea employed in ISML 1.0 is called “capsulation.” A set of modules can be tightly packed by declaring capsulation and contained by a capsule module as a representative of the modules within the capsule (Fig. 2). The capsule module can have input and output ports that provide the sole interfaces for access to and from the modules inside the capsule. Input ports of the capsule module are associated with input ports of encapsulated modules by another type of edge: “forwarding,” designated so because of similarity to the concept of port forwarding used in computer networks. Similarly, output ports of modules inside the capsule module may be connected to output ports of the capsule module by “forwarding” edges. A capsule module can be one of the modules encapsulated by another capsule module, which means that the capsulation can nest, leading to a hierarchical representation of a model. A module encapsulation covers all modules within the capsule module at the most interior (lowest) level. In many cases, a capsule module simply symbolizes a certain physiological function and can be easily reused as an element of other models.

ISML 1.0 is compatible with CellML. Let us briefly summarize the similarities between them. Both languages describe physical units for values used in models. A basic element comprising a model is called a “component” in CellML, and its counterpart is the “module” in ISML 1.0. Physical variables called “variables” in CellML can be read as “physical-quantity” in ISML 1.0. A physical-quantity for ISML 1.0 can be specified as one of several types, such as “state,” “dynamic-parameter,” “static-parameter,” and “func-expression,” among others. In particular, a major difference from CellML is that the physical-quantity of ISML 1.0 has a type of “morphology” and “time series” to deal with, numerically and analytically defined morphology and time series data acquired by real world biological experiments and/or numerical simulations. The morphology-type physical-quantity in ISML 1.0 is required, for example, in a problem with PDEs that uses morphology to define the shape of domain and boundary of the problem, or in a problem with ODEs that has constraint conditions defining a non-trivial manifold (e.g., equations of motion of a mass constrained on a surface with a complicated shape).

Ports to export/import values to/from modules outside the module under consideration specified in ISML 1.0 have no clear correspondence in CellML. This concept is indirectly expressed in CellML as public-/private-interfaces of variables. The “connections” in CellML are used to express equivalence between different variables and components, and this roughly corresponds to the edges defined in ISML. Relationships indicated by edges and connections contribute to the construction of physiological ontology based on a large set of models. There are several differences between CellML connections and ISML edges. The edges in ISML represent directional relationships among modules, whereas CellML connections are nondirectional. That is, each edge in ISML possesses “head” and “tail” modules to be linked. Moreover, each edge can have an operation-type “meaning” to which several words are assigned to specify how two modules are affected by each other.

In the following sections, we focus mainly on typical ISML 1.0 features that are not found in CellML or others.
See our Web site http://www.physiome.jp/documents/HTMLs/isml_ver1.html for more details and other specifications that are not described in this article.

2. Expressions used in ISML 1.0

In this text, the term “insilico model,” “ISML model,” or simply “model” refers to a single model described by an entire ISML document, which includes metainformation, definition of physical units, morphological data, and time series data as well as mathematical expressions representing dynamics of the physical functions under consideration.

2.1. Modularity and linkage among modules

Here we describe the Beeler-Reuter (BR) model [9] using ISML 1.0 to illustrate the modularity and linkage among modules. The BR model can simulate electrical excitation of a cardiac ventricular cell membrane. The dynamics of the membrane potential $V$ are determined by four ionic channel currents, i.e., time-independent potassium outward current, fast inward sodium current, and slow inward calcium current, and another current as an external stimulus. The dynamics of each ionic current are determined by the current, and another current as an external stimulus. The fast inward sodium current, and slow inward calcium sodium outward current, time-activated outward current, four ionic channel currents, i.e., time-independent potassium dynamics depend on the membrane potential.

$V_{m}$ is the membrane potential and $E_{r}$ is the reversal potential of $i_{r}$, $d$ and $f$ are the gate variables whose dynamics depend on the membrane potential.

Let us consider a model representing the slow calcium channel current $i_{s}$, designed as in Fig. 3. The module $M_{s}$ defining the current $i_{s}$ is designed with three submodules, which are graphically represented by the three ball-like modules connected by solid lines (representing the structural edges) in Fig. 3. In this case, the variables $d$, $f$, and $E_{r}$ on the right-hand side of Eq. 1 are specified as physical-quantities in the modules $M_{d}$, $M_{f}$, and $M_{r}$, respectively. To define $i_{s}$ in the module $M_{s}$, these three physical quantities are imported through the functional edges indicated by the dashed curves in Fig. 3. $V_{m}$ is obtained externally via the input port of this model and is used by the modules $M_{d}$, $M_{f}$ as indicated by the chain curves (representing the forwarding edges).

2.1.1. Edge. The edges shown in Fig. 3 are listed in an ISML document in the edge-set section. This section specifies the type of each edge, its departure (head) and destination (tail), and operation describing the meaning of its functional relationship with a verb or a verb phrase.

The head and tail of each edge of functional and forwarding types specify both module-id and port-id to uniquely identify a port within the model. In the case of structural, logical, and capsular edges, port-id = 0 is used as a dummy port since these edges define the relationship between modules and do not use ports.

2.1.2. Module. A description of the module $M_{s}$ starts with <module module-id="A6CD8604-2F61-...-DBD7C92378E5" type="functional"/>

where the module-id is specified as a 16-byte universally unique identifier (UUID), which is unique across all insilico models. The type of the module...
**ISML** is specified as **functional-unit**, meaning that this module possesses particular features such as **physical-quantities** necessary for modeling physiological functions.

**ISML 1.0** offers several types of modules other than **functional-unit**. These include **container**, **capsule**, and **template**. A module of **container** type does not possess **physical-quantities** and is supposed to represent a conceptual box to put several modules together, e.g., a **capsule**-type module with a name “ganglion” may be composed of many modules representing nerve cells. A **capsule**-type module acts as a symbol of a physiological function that is modeled by a set of modules. A capsule module itself thus does not possess any **physical-quantities**, but may have only input/output ports as its interfaces. A **template**-type module is similar to the concept of a class in C++, which can realize an object instantiation. Dynamic construction and destruction of instances of a template module can be managed during the run time of numerical simulations, and rules necessary for instance management can be described in ISML 1.0. This is particularly important for constructing and simulating agent-base models. A **module** is also characterized by the following four children tags: **property**, **port-set**, **physical-quantity-set** and **morphology-set**. The **property** describes basic properties of a module, such as name, keywords, track, and so on. The tags **keywords** and **track** are described in the next section.

**2.1.3. Port.** Returning to the module **MI**, example: it has four input ports and one output port. An ISML document enumerates them in the **port-set** subsection of the **module** section.

```xml
<port-set>
  <port direction="out" port-id="1">
    <name> I_s </name>
    <reference physical-quantity-id="1"/>
    <description> calcium channel current </description>
  </port>
  <port direction="in" port-id="2">
    <name> V_m </name>
    <description> potential </description>
  </port>
  ...
</port-set>
```

In this example only two of the five ports in module **MI**, are described for brevity. Each port has a direction (in or out) and is given a port-id as a sequential number unique within a module. A port with direction = “in” is called an input port, and “out” an output port. A **physical-quantity** that goes out through this output port must be specified by its ID (physical-quantity-id) in a child tag **reference** of the port. A value of a **physical-quantity** delivered from an external module through the input port is set as a **physical-quantity** of the module and used in the module.

In this case, **physical-quantity-id** is not specified. Instead, a **physical-quantity** that receives a value from the input port specifies its port-id.

**2.1.4. Physical-quantity.** In the ISML paragraph described above, the **physical-quantity** with the output port of physical-quantity-id = “1” corresponds to the slow calcium current and can be defined as a **variable-parameter** in the **physical-quantity-set** section of the module as follows.

```xml
<physical-quantity type="variable-parameter" physical-quantity-id="1">
  <name> I_s </name>
  <precision> double </precision>
  <unit unit-id="1"/>
  <dimension type="scalar"/>
  <implementation> ... </implementation>
</physical-quantity>
```

A type and **physical-quantity-id** must be given first when a **physical-quantity** is defined. In this example, **I_s** is declared as the **variable-parameter**, which expresses values that vary during simulation, such as a computed value of a static (nondynamic) mathematical function of dynamic variables (state). The values obtained by this module are defined in the **implementation** with mathematical formulae. ISML 1.0 defines other types of **physical-quantity**, i.e. **state**, **variable-parameter**, **func-expression**, **nominal**, **morphology**, and **timeseries**. In the example above, the **precision** is set to **double** (other possible candidates are **int**, **char**, and **bool**). The **unit** specifies a unit-id as a user-defined combination of the fundamental seven base units (i.e., meter, kilogram, second, ampere, kelvin, candela, mole), together with prefixes such as kilo-, milli-, and micro-. The **dimension** is set to **scalar** in this example. ISML 1.0 supports vector and matrix values for a **physical-quantity** to use in equations.

The **implementation** section is used to describe concrete contents of the **physical-quantity**, such as values for **variable-** and **static-parameters** and mathematical formulae for **states**. Specific definition is described in a child tag **definition**. For example, the **variable-parameter** **I_s** is defined as

```xml
<implementation>
<definition type="ae" format="mathml">
  <math>
    <apply><eq>
      <ci> I_s </ci>
      <apply><times>
        <ci> I_s </ci> ...
      </apply>
    </eq>
  </math>
</definition>
</implementation>
```

Here is the MathML expression for \( i_s = \rho \cdot d \cdot f \cdot (V_m - E) \).
2.1.5. Capsulation. The module \( M_I \), and its submodules are encapsulated in the example above. The capsule itself is represented as a module with the capsule type depicted as a square frame in Fig. 3. A module at the top level (referred to as the root) of a local tree structure can be encapsulated by a capsule module if all edges (except forwarding edges) are self-contained. The module \( M_I \) satisfies this condition and can be encapsulated. The capsule module is linked by a capsular edge (which is not illustrated in the figure). Once the root module is encapsulated, input and output ports of the modules under the root module are linked to the input and output ports of the capsule module by edges of the forwarding type. Whether or not the module is encapsulated is specified in the property tag of the module as

\[
\begin{align*}
\text{capsulation} & \text{ state="true">} \\
\text{capsulated-by} & \text{ module-id="CDEA5AE5-16DD--8A5848ACF217"/}
\end{align*}
\]

in which the module-id of the capsule module is specified.

Figure 4 shows the whole BR model composed of the module \( M_l \) modeling the membrane and four encapsulated modules modeling the channel currents (one of them is the slow calcium current module shown in Fig. 3). The BR model is encapsulated by the capsule module named “Beeler Reuter 1997 model,” which has an input port to receive the external stimulus current and an output to export the membrane potential. Using this model, the user can create, for example, coupled BR models by linking several individual BR modules.

2.2. Morphological data and mathematical expressions

Let us consider another example, a reaction-diffusion dynamics with the FitzHugh-Nagumo model. This model represents the excitation conduction on a two-dimensional medium as a model of cardiac tissue analyzed by Hall and Glass (1999) [10]. They investigated spatio-temporal changes in the propagating action potential on a two-dimensional sheet paved with excitable cell models. The reaction diffusion equations that they used are as follows:

\[
\frac{\partial v}{\partial t} = \frac{1}{\varepsilon} \left( v - \frac{1}{3} v^3 - w \right) + D \nabla^2 v + I_{loc} + I_{stim}(t)
\]

(2)

\[
\frac{\partial w}{\partial t} = \alpha (v - \beta - \gamma w) g(v)
\]

(3)

where \( g(v) \) is a sigmoidal function controlling the rate of the pacemaker

\[
g(v) = \frac{w_h - w_l}{1 + e^{\frac{-kv}{w}}} + w_l
\]

(4)

\( v \) and \( w \) are the excitation and recovery variables, respectively. \( I_{loc} \) is a constant current applied to a localized region at the center of the sheet. \( I_{stim}(t) \) is a pulsatile stimulation current used for resetting applied to selected points at selected timings.

We begin by designing the corresponding insilico model using ISML 1.0 (Fig. 5). The PDE with FitzHugh-Nagumo is modeled here by two modules representing excitation variable \( v \) and recovery variable \( w \), respectively, which are encapsulated to consolidate model independence. The modules \( M_v \) and \( M_w \) are mutually linked by functional edges. The module \( M_v \) receives two external electric current stimuli \( I_{stim} \) and \( I_{loc} \) via the input ports of the capsule module. The module \( M_{area} \) provides a morphology of the tissue on which the reaction diffusion problem is solved. Here the module \( M_{area} \) is linked by a structural (constituent) edge to the capsule module, which contains the modules representing the current stimulations as well as the PDE of the FitzHugh-Nagumo model, since all of them are solved on the morphology provided by module \( M_{area} \).

2.2.1 Morphology. The module \( M_{area} \) is implemented as a functional-unit type that has several physical-quantities, one of which is the morphology type. ISML 1.0 can handle numerically defined morphology, i.e., using morphological data that is an assembly of points (a set of two or three dimensional coordinates), and also morphology...
Specifications of \textit{insilicoML} 1.0

\begin{itemize}
\item Defined analytically (mathematically). In the former case, the data can be kept as an independent file either on a hard disk at a local machine or in a remote database. For the latter case, mathematical expressions are written in the \textit{insilico} model. Morphological models used in a module are described in the \texttt{morphology-set} section of the module. The simple two-dimensional square sheet used in this example (shown in Fig. 6a) can be described using MathML as follows:

\begin{verbatim}
<morphology-set>
  <morphology type="internal" morphology-id="1">
    <definition format="mathml">
      <math>
        <apply>
          <and/>
          <apply>
            <geq/>
            <ci>x</ci>
            <cn>10</cn>
          </apply>
          <apply>
            <leq/>
            <ci>y</ci>
            <cn>10</cn>
          </apply>
        </apply>
      </math>
    </definition>
  </morphology>
</morphology-set>
\end{verbatim}

A sequential unique ID number within the module is assigned to each fragment defining the morphology listed here. A type of the \texttt{morphology} tag specifies whether the morphology is defined within the \textit{insilico} model file (type = “internal”) or outside as an independent file (“external”). The morphology defined in the module $M_{area}$ can be more complicated, such as shown in Fig. 6b, defined numerically as morphological data. In that case, the description of ISML is as follows:

\begin{verbatim}
<definition format="numeric" iref="./data/morphology/sample1.wrl"/>
\end{verbatim}

\begin{verbatim}
<implementation>
  <definition type="assign" format="morphology">
    <reference morphology-id="1"/>
  </definition>
</implementation>
\end{verbatim}

Note that if the resource file defining the morphology is at a remote machine, the attribute \texttt{xref} for cross-reference must specify the remote address.

Morphological data used in an \textit{insilico} model should be cataloged in the \texttt{morphology-set} section. Each data set must be bound to a \texttt{physical-quantity} with the \texttt{morphology} type. The \texttt{physical-quantity} with the \texttt{morphology} type can be assigned morphological data (numerical data or mathematically defined morphology), and the data may be used to solve PDEs numerically. An example of \texttt{implementation} for a \texttt{morphology} type \texttt{physical-quantity} is as follows:

\begin{verbatim}
<implementation>
  <definition type="assign" format="morphology">
    <reference morphology-id="1"/>
  </definition>
</implementation>
\end{verbatim}

\begin{verbatim}
<definition type="assign" format="morphology">
  <reference morphology-id="1"/>
</definition>
</implementation>
\end{verbatim}

where the morphology-id is specified at \texttt{reference} regardless of the type of morphology, i.e., numerical data or mathematical expressions.

ISML 1.0 defines a special type of \texttt{physical-quantity} named \texttt{shape}, which is one of the ISML-predefined \texttt{physical-quantities}, i.e., all modules possess this \texttt{physical-quantity} by default. It is a special case of the \texttt{morphology}
type and can be used for binding morphological data. The physical-quantity shape represents the shape of a module. The value of this physical-quantity can be referenced by any children modules that are linked by structural or logical edges to the module possessing this physical-quantity shape. In the example (Fig. 5), the modules $M_r$, $M_e$, $M_{I_{loc}}$, and $M_{I_{stim}}$, which are encapsulated by the outer capsule module, can use the morphological data defined in the module $M_{area}$ through its shape. For example, assume that the module $M_r$ has a morphological type physical-quantity with ID = 4. In its definition in the implementation, the physical-quantity is assigned morphological data by inheriting the physical-quantity shape of the parent module $M_{area}$. This is done by specifying the module-id of the parent module as follows:

```xml
<implementation>
  <definition type="assign" format="morphology">
    <reference module-id="A6CD8604-2F61-...-DBD7C92378E5"/>
  </definition>
</implementation>
```

Notice that the parent module-id is specified instead of the morphology-id at the reference tag.

**2.2.2. Partial differential equations.** The module $M_r$ includes a PDE as the governing equation of the dynamics of $v$, which is a state-type physical-quantity. For the state-type physical-quantity, unlike the variable-parameter type, argument-set, domain, and problem-condition must be defined as well. By declaring the argument variables in argument-set, it becomes clear that the state-type physical-quantity $v$ of the FitzHugh-Nagumo model PDE is a function of time and space $(x$ and $y)$. The arguments are also used for the func-expression type physical-quantity, as shown later. The domain specifies the domain on which the function or differential equations are evaluated or solved. The following is a section of ISML 1.0 describing argument-set and domain for our PDE example:

```xml
<argument-set>
  <argument physical-quantity-id="2"/>
  <argument physical-quantity-id="3"/>
</argument-set>
<domain>
  <definition format="morphology">
    <reference physical-quantity-id="4"/>
  </definition>
</domain>
```

In the example, we are dealing with the reaction-diffusion equations defined on a two-dimensional sheet. The two arguments declared in this ISML fragment correspond to the spatial coordinate variables $x$ and $y$, which are independently defined elsewhere as the nominal-type physical-quantities with ID = 2 and 3. The nominal-type physical-quantity does not have its implementation in the definition, only the specifications of name, unit, precision, and dimension. The domain is defined by specifying a morphological physical-quantity (ID = 4) assigning morphological data from the parent module as mentioned above. The domain can also be defined mathematically in MathML, in which the argument variables define functions.

In the implementation of the state type physical-quantity $v$ for example, the PDE is described in MathML format. All terms of the equation are written on the left-hand side of the equation, and the right-hand side is always 0. Only for PDE definition, the definition tag has a subtype attribute to specify the type of PDE as one of elliptic, parabolic, hyperbolic, and others, and a form attribute specifying if the PDE is represented as weak, using variational formulation, or strong, using PDE with derivatives.

```xml
<implementation>
  <definition type="pde" sub-type="parabolic" format="mathml">
    <math>
      <apply:eq>
        <apply><partialdiff/>
          <bvar>$v$</bvar>
          <bvar>$t$</bvar>
        </apply>
      </apply>
      <apply><partialdiff/>
        <bvar>$v$</bvar>
        <bvar>$x$</bvar>
      </apply>
      <apply><partialdiff/>
        <bvar>$v$</bvar>
        <bvar>$y$</bvar>
      </apply>
    </math>
  </definition>
</implementation>
```

Here are MathML expressions for
\[
\frac{\partial v}{\partial t} - D \frac{\partial^2 v}{\partial x^2} - D \frac{\partial^2 v}{\partial y^2} - \frac{1}{\epsilon} \left( v - \frac{1}{3} v^3 - w \right) - I_{loc} - I_{stim}(t) = 0
\]

In the example of the FitzHugh-Nagumo model, zero-flux boundary conditions are applied to the four borders of the square area, which is equivalent to the zero Neumann condition at each boundary. These conditions must be described in the model. In general, ISML 1.0 specifies conditions accompanying the equations to be solved, such as boundary conditions, in the problem-condition section. The constraint, material, and initial conditions, as well as the boundary conditions, can be defined. For definitions
of the boundary conditions, the user must specify a type of either neumann, dirichlet, robin, or other. If necessary (for example, in the cases of force and flow), the direction can be defined in the direction tag by specifying argument variables, such as $x$ or $y$, or explicitly by a word, either perpendicular or parallel, indicating the direction with respect to the tangential plane (or line) at every point on the boundary. Notice that perpendicular is therefore the direction normal to the boundary. The notation Neumann($v$) is used to describe the Neumann condition. For example, if the Neumann condition of state $v$ is 0 on a segment of the boundary (specified by a segment-id described later), it is written as Neumann($v$) = 0, as shown in the example ISML document below:

```xml
<problem-condition-set>
  <problem-condition type="boundary">
    <configured-at segment-id="2">perpendicular</configured-at>
    <direction>perpendicular</direction>
    <definition type="neumann" format="mathml">
      \math \begin{align}
      \text{Neumann} & \text{ conditions are specified as a function of the boundaries.}
      \end{align}
    </definition>
  </problem-condition>
</problem-condition-set>
```

Similarly Dirichlet($) is for defining the Dirichlet condition. The user can define other types of conditions using general mathematical formulae.

For constraint conditions, a space (usually a manifold) $M$, in which the target physical-quantity $x$ is constrained, is defined in the definition section. $M$ can be described either in a mathematical format, such as $\{x, f(x) = 0\}$ with format = “mathml,” or by a set of points represented by morphological data for digitized, numerically defined cases with format = “morphology.” To specify material conditions, the values of physical material properties (such as Young’s modulus, density, and viscosity) are set in an equation format.

For those conditions to be complete, it is necessary to specify spatial positions where each condition is applied. Particular segments constituting the morphology, i.e., subsets of points of numerical morphological data or pieces of geometric objects such as an arc or a portion of a rectangle, are defined in the segment-set section in the definition of the morphology and are labeled by a sequential ID (segment-id) to be specified uniquely within the module. Each segment is also categorized either by point, line, area, or volume. An appropriate condition is then applied to each segment specified by its segment-id in the configured-at tag. Here is an example of a definition of the segments.

```xml
<morphology type="external" morphology-id="1">
  <definition ref="./data/morphology/sample1.wrl"/>
  <segment-set>
    <segment segment-id="1" type="line">
      <point no="1"/>
      <point no="3"/>
      <point no="5"/>
    </segment>
  </segment-set>
</morphology>
```

A dummy variable is used to mathematically define segments of a given morphology. For example, geometric figures can be given as $x = \cos(2\pi s)$, $y = \sin(2\pi s)$ where $s \in [0:0.6]$ is a dummy variable. For a morphology defined by numerical data, this can be done by listing points specified by their number.

In the problem-condition, conditions such as mesh, solver, and post can be defined. They are related to numerical methods and processing performed during and after simulations of the model. The mesh condition specifies the number of nodal points involved in the specified structure, and it is used by a mesh-generator. Mesh-node information is described in a separate file with a certain format. Note that instead of specifying the number of nodes in the structure, it is also possible to import external node files. The solver condition specifies a solver used for the finite element method to solve PDE with morphology. ISML 1.0 offers at least six choices: LU, Cholesky, Crout, CG, GMRES, and UMFPACK, Convergence criteria must be specified. The post condition is used to define the postprocessors. The postprocessor type determines the file format to which simulation results are saved, such as gnuplot and medit.

In our example of the FitzHugh-Nagumo model, the module $M_{inh}$ generates the constant current applied to a localized region at the center of the sheet (indicated as a gray disk in Fig. 6, a and b) either to make the media in the corresponding region oscillate or to depress excitability. The current can be described mathematically as a function of spatial coordinates $x$ and $y$: $I_{inh}(x, y) = C_{inh}$ when $x^2 + y^2 \leq r^2$; otherwise $I_{inh}(x, y) = C_{high}$, where $C_{low}$ and $C_{high}$ are constants. Thus the physical-quantity transmitted from $M_{inh}$ to the module $M$ is not a scalar value, but a set of values as the function of the space. To describe this sort of physical-quantity, the func-expression type physical-quantity is available. This type does not provide values, but instead a set of mathematical formulae of the function as it is (as text descriptions). The func-expression type physical-quantity also requires specification of the argument variables for use in the definition of the imple-
m entation of the func-expression. The definition of the func-expression type physical-quantity begins with <physical-quantity type="func-expression"-quantity-id="1">. It is characterized by name, precision, dimension, unit, argument, domain, and implementation. Note that to define the dimension, in this case the row and/or col must be specified as discretization-dependent (i.e., <dimension type="matrix"> <col> discretization-dependent <row> discretization-dependent <row> <dimension>), since the dimension of such a variable cannot be specified until a solver discretizes the domain of the function defined in the func-expression for numerical computations. The example code of the implementation for module MIloc is as follows:

```xml
<implementation>
  <definition type="conditional">
    <case-set case-set-id="1">
      <case case-id="1">
        <condition format="mathml">
          \[
          x^2 + y^2 \leq 4
          \]
        </condition>
      </case>
      <definition type="func-expression" format="mathml">
        <math>
          \ci\times
        </math>
      </definition>
    </case-set>
    <case-set case-set-id="2">
      <case case-id="2">
        <definition type="func-expression" format="mathml">
          <math>
            \ci x + \ci y
          </math>
        </definition>
      </case>
    </case-set>
  </definition>
</implementation>
```

In the implementation, the current I_{loc}, which is the function of spatial coordinates x and y, is defined using conditional branches, since the value of I_{loc} depends on the position, for example, if x^2 + y^2 \leq r^2, THEN I_{loc} = C_{low}, ELSE I_{loc} = C_{high}. The conditional branches are described by the definition tag with type = “conditional”, which takes case-set and case tags as children. The condition is described in MathML format, and the definition is evaluated if the condition is satisfied. These are described in the case section labeled by case-id. The cases in the same case-set indicate that the cases are in the same sequel, which is similar to the “IF - ELSE: IF - ELSE” syntax in computer languages. The last case in the example code above, which has no condition section, corresponds to the “ELSE” phrase.

3. Databasing, tracking, and relating models

Models created by the user are stored in a model database (ISML database) if the models satisfy certain criteria, such as publication in peer-reviewed journals. Models in the database are uniquely identified by a database ID (db-id), which is assigned by the database server when the model is registered in it. Users can download models from the database and reuse them to develop new models.

ISML 1.0 is capable of tracking any module in terms of the history of its reuse if the module is registered in the database. This is done using the property section, where an ISML document of each module records the db-id of the models that involve (reuse) the module. The reuse history of modules can provide another basis for the model-based ontology of physiological functions as well as the metainformation assigned to the modules and the edges of operation-type with “meaning” as mentioned above.

Let us take the BR model again as an example for illustrating how the reuse history is recorded in each module. When creating the BR model, we first create the model of the slow inward calcium channel current (Fig. 3) as an element of the BR model. During development of the calcium channel model until it is completed and registered in the ISML model database, the model ISML document has no database ID (db-id). In this period, when a module such as the module MI is created in the calcium channel model, module-tracking information, such as “module MI, is involved in this calcium channel model,” is written as a property of the module MI, as follows:

```xml
<track>
  <involved db-id="this" date="2007-06-06"/>
</track>
```

where “this” corresponds to the calcium channel model. This indicates that the module MI, is used as an element of the calcium channel model. When the development of the calcium channel model is completed and it is uploaded to the ISML database, the database server assigns a unique db-id to the calcium channel model. More precisely, the db-id is given to the capsule module encapsulating
the model of the calcium channel. Let us assume that the server has given database ID 41 to the calcium current model. Then the value “this” in the track tag for the module $M_I$, is replaced by the newly assigned db-id at the server, and the tracking record is rewritten as <involved db-id="41" date="2008-06-06">.

In the subsequent development process of the entire BR model (Fig. 4), let us assume that the slow calcium current model has already been registered in the database and that we can download it from the database for reuse as one of the modules composing the BR model. When the calcium channel model is reused in the BR model, a new track record is added to the module $M_I$, which has been a submodule of the calcium channel model as follows:

```xml
<track>
  <involved db-id="this" date="2008-03-22"/>
  <involved db-id="41" date="2007-06-06"/>
</track>
```

In this way, the reuse history of the module $M_I$ is accumulated and logged.

The Luo-Rudy (LR) model [11] is another representative model describing the action potential generation of the ventricular cardiac myocyte. The LR model can be considered as an extended or modified version of the BR model. Luo and Rudy modified the model by adding several new modules, including representations of extracellular ion concentrations, a novel potassium channel, and a plateau potassium current. The slow calcium current model was used in the LR model the same way as formulated in the BR model. Let us consider ISML model creation according to this actual history of the model development. That is, we reuse the slow calcium channel model of the BR model for the creation of the LR model using ISML 1.0. Upon registration of the newly created LR model, tracking for module $M_I$, is revised as

```xml
<track>
  <involved db-id="131" date="2008-08-14"/>
  <involved db-id="75" date="2008-03-22"/>
  <involved db-id="41" date="2007-06-06"/>
</track>
```

where ISML models with bd-id = 41, 75, and 131 correspond to the slow calcium channel model, the BR model, and the LR model, respectively. Once the module is created and the model including that module is registered, this history track is maintained as long as the module exists even if the module is modified (deletion or edit of some part of the module). Because of this, modifications made to the module cannot be identified clearly from the track records, though one can try to compare the module after repeated modifications with the original module, which is kept unchanged in the database. Nevertheless, because it is expected that the essential nature of the module will remain even after modification, it is worthwhile to provide the history-tracking capability.

Based on the tracking information recorded in each module, the origin of each module involved in a model can be revealed, and, with appropriate analytical tools for ISML documents on the database, the user can obtain a tree diagram of the module (model) development for an analysis of module phylogeny. In our simple example, the user can discover that the BR model and the LR model use the same or similar slow calcium channel model. Moreover, the track information enables the user to define distance or similarity between two different models. The larger the number of common modules shared in the two models, the smaller the distance between the models, leading to a model-based construction of the physiological ontology.

4. Discussion

This article illustrated the progress of the development in the description language insilicoML (ISML) 1.0 for multiscale and multilevel models of physiological functions. ISML 1.0 and integrated development environment tools, such as the ISML editor, model viewers, and simulators, are expected to be a pillar of promotion of physiology and systems biology by which researchers in the interdisciplinary fields among physiology, engineering, medicine, and informatics can make concerted efforts toward quantitative and integrated understanding of the human physiology [1]. The key concept here is to describe the physiological functions using common languages that can be shared worldwide.

The development of ISML 1.0 aims to enhance model sharing among researchers, both experimental and theoretical, by cooperating with other major model description languages (including CellML and SBML). Individual researchers constructing and simulating physiological models often use different languages and different environments, making it difficult to exchange models and ideas. ISML 1.0 and its tools will provide a common platform on which many researchers can cooperate in modeling, simulating, and analyzing their own models of particular physiological functions.

ISML 1.0 is designed to describe in a consistent manner various physiological phenomena at any spatiotemporal scale, such as the dynamics of ion movement, proteins, cells, tissues, organs, and the whole body along with relevant morphology. If unrelated researchers develop models for different targets or at different levels using a common framework such as ISML 1.0, those models can easily be combined to construct models with multiple scales and levels. Moreover, users can easily examine changes in the model dynamics when particular parts of the model are replaced by other models with the aid of the...
encapsulation used in ISML.

An important aspect provided by the development of ISML 1.0 and its database is a novel basis necessary for construction of an ontology of the physiological functions. Our idea proposed in this article for the physiological ontology has its basis in several types of relationships between mathematical models of the physiological functions. There have been several developed ontologies, such as the gene ontology and the protein ontology. However, construction of the ontology for physiological functions may not be straightforward, since the subjects have much wider varieties of nomenclature than for genomes or proteomes. If a physiological function is described using the ISML model, it can be decomposed into fundamental elements based on the modules included in the model. Each module can represent a noun by a name of the specific physiological function modeled by the module. Functional relationships between modules are represented by the edges in ISML, and each of them can describe a meaning of the relationship between any related modules in the ISML model. The functional relationship can correspond to a verb, i.e., how the state of one module can affect the state of another module through the functional relationship. A large number of nouns and verbs stored in the ISML database contribute to constructing the ontology. The history of module reuse is used to define similarities and differences between models, and can also contribute to the construction of the ontology. By quantifying those relationships, the ontology proposed in this study can determine what physiological functions exist as the elements of the ontology, what relationships are formed among them, and how these elements can be categorized according to the similarities and differences between models. Such ontology can support research activities in various ways, including (1) model search from a large number of models stored in the database, and (2) semi-automated construction of new models and large-scale models by combining individual models and modules using the ontology, among others.

For the model sharing and constructing ontology to be productive, a sufficient number of ISML models are required in a model database (ISML database) that provides storage, search, view, and distribution (download) functions. There is already a database (as of August 21, 2008) for ISML 0.1 (http://www.physiome.jp) that stores more than 100 models written in ISML 0.1. ISML 1.0 has complete backward compatibility with ISML 0.1; thus the models in the current database will function with ISML 1.0. The ISML database will be able to work in cooperation with other databases, such as gene and protein databases and SBML and CellML model repositories. Collaboration between the ISML database and other databases will also be supported by the ontology.

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