A modeling and simulation language for biological cells with coupled mechanical and chemical processes

Endre Somogyi\textsuperscript{1} and James A. Glazier\textsuperscript{2}

\textsuperscript{1}Department of Computer Science, Indiana University
\textsuperscript{2}Department of Intelligent Systems Engineering, Indiana University

February 28, 2017

Abstract

Biological cells are the prototypical example of active matter. Cells sense and respond to mechanical, chemical and electrical environmental stimuli with a range of behaviors, including dynamic changes in morphology and mechanical properties, chemical uptake and secretion, cell differentiation, proliferation, death, and migration.

Modeling and simulation of such dynamic phenomena poses a number of computational challenges. A modeling language describing cellular dynamics must naturally represent complex intra and extra-cellular spatial structures and coupled mechanical, chemical and electrical processes.

Domain experts will find a modeling language most useful when it is based on concepts, terms and principles native to the problem domain. A compiler must then be able to generate an executable model from this physically motivated description. Finally, an executable model must efficiently calculate the time evolution of such dynamic and inhomogeneous phenomena.

We present a spatial hybrid systems modeling language, compiler and mesh-free Lagrangian based simulation engine which will enable domain experts to define models using natural, biologically motivated constructs and to simulate time evolution of coupled cellular, mechanical and chemical processes acting on a time varying number of cells and their environment.

\textbf{Keywords:} Biological Systems Modeling, Simulation, Spatial Hybrid Systems

1 Introduction

Life is the result of a complex interplay between chemical, mechanical, electrical and other physical processes [5]. Biological cells exist in a dynamic, spatial, fluid environment where they sense stimuli in their environment such as physical forces, chemical and electrical signals, fluid pressure
and flow, or other physical properties. Cell responses include changes in morphology or mechanical properties, proliferation, death, differentiation, motion and production of signals.

Mechanistic models of natural systems do not attempt to replicate physical reality in exact detail, but describe the mechanisms which quantifiably predict specific observed behavior. Since all aspects of a model are controlled, while many aspects of an experiment are not controllable or not observable, simulations enable researchers to test and validate hypotheses. Simulations are often faster than real time and are generally cheaper, safer and sometimes more ethical than real-world experiments.

Building mechanistic models of biological phenomena can be challenging, both because of the inherent complexity of the biological system being modeled and because models are typically written in programming languages that do not easily describe physical concepts. Most programming languages either represent abstractions of the underlying Von Neumann hardware architecture, or are meant to represent models of computation (e.g., \( \lambda \) calculus) rather than natural physical phenomena.

In order to enable scientists to create mechanistic models easily, we are developing a new programming language and simulation environment, \textit{Mechanica}. Our motivating problem domain is \textit{mesoscopic} molecular and cellular biology, where atomistic simulation approaches are not computationally tractable and deterministic continuum approximations are not valid. Mechanica is a physically motivated model description language and simulation engine suitable for many kinds of mesoscopic agent-based phenomena.

Mechanica describes models in terms of objects and processes. Objects are the physical things being studied in our problem domain. Objects are state-full things that exist as snapshots in time; objects themselves are time-invariant – they do not have any intrinsic dynamics. Processes act on objects and change the state of objects over time. They define the dynamics and interactions of and between objects. Objects and processes work together to define how a system evolves over time; i.e., they describe a dynamical model.

2 Related Work

Languages designed for modeling dynamic physical systems differ from mainstream programming languages in that they incorporate notions of time and/or space \cite{2} and alleviate the need for users to think about low-level computational implementation details. Many modeling approaches can describe different aspects of biological processes. Some modeling approaches are well-suited for macro-scale chemical reactions, others at nano-scale atomistic dynamics, but few approaches exist for meso-scale dynamics that allow for a natural description of coupled chemical and mechanical processes.

Systems biology is a holistic or integrative approach that seeks to understand phenomena as a whole, and such approaches frequently represent biological phenomena like gene regulatory, chemical or metabolic networks. Systems biology focuses on how individual components interact and communicate rather than investigating the internal workings of the components. Systems biology modeling approaches tend to have constructs for representing transformation processes such as chemical reactions and discrete events, but lack concepts to represent dynamic geometry and
mechanical processes. Systems biology models are represented with languages such as Systems Biology Markup Language (SBML), Petri Nets or process algebras.

SBML (http://www.sbml.org/) can describe phenomena such as chemical reactions in a well-stirred compartment, but has limited support for spatial processes and no concept of forces, dynamic geometry or structural rearrangement. SBML itself has well-defined semantics for discrete events which we adopt in Mechanica, and individual SBML models can be connected together as part of a larger simulation environment such as discrete event simulation [3].

Petri Nets are a modeling formalism introduced by Carl A. Petri to represent chemical networks. Petri Nets are well-suited to describe concurrent communication and synchronization networks. Colored Petri Nets (CPN) [10] combine Petri Nets with a functional programming language to create a discrete event modeling language. The integration of a functional programming language simplifies model construction and increases the modeling capability of CPNs. CPNs are frequently applied in systems biology, and multiple CPNs can be coupled and arranged in a grid to simulate spatial systems such as bacterial colonies [13]. CPNs generally lack concepts of dynamic space and do not have a natural way to describe mechanical processes. Certain CPN-related process algebras (like Cardelli’s 3π language [6]) can describe dynamic spatial arrangements. Mechanica also incorporates a functional programming language similar to the way in which CPNs combine reaction networks, discrete events and a functional programming language.

Systems biology approaches in general are not ideally suited for describing bio-mechanics, as they do not have natural ways to represent mechanical constructs such as forces, stress, constitutive relations, or geometry transformations.

Particle-based approaches start at the nano rather than system scale and describe natural phenomena from an atomistic perspective. Traditional MD simulators model a fixed number of spherical, volume-excluding particles (atoms) interacting with a classical force field. MD simulators have simple particle types and are restricted to a finite set of predefined forces. Reactive MD simulators such as ReaxFF [17], ReaDDy [14], or SRSim [8] extend classical MD with reacting particles. ReaxFF evolves particles via energy or temperature constrained Newtonian dynamics, and ReaDDy and SRSim via over-damped Langevin dynamics. Reactive MD simulators can model self-assembly of complex materials out of particles. ReaDDy supports particle creation and annihilation, whereas particle count is fixed in ReaxFF and SRSim. Reactive MD alters bonded relationships between particles to form chemical complexes by by adding or removing mechanical bonds between particles. SRSim uses BioNetGen to specify reaction rules, whereas ReaxFF and ReaDDy use their own specification format.

Reactive MD excels at describing the nano-scale, however meso- and multi-scale phenomena present challenges. Reactive MD particles are simple, in they only have position, momentum and orientation, and are not typically augmented with additional state variables, thus individual particles can not model more complex things like macromolecules or cells. Model specification in Reactive MD can be challenging because users typically specify custom forces in C++ or Java instead of a model description language. Reactive MD does not have concepts of continuous chemical reactions or fields like systems biology approaches.

Stochastic particle simulators such as MCell [16], Smoldyn [1], or ML-Space [4] are well-suited for spatial biochemical simulations such as reaction-diffusion in complex geometries. These
simulators move individual rigid particles with either spatial Gillespie or Brownian dynamics as opposed to interaction potentials as with MD. MCell is restricted to point particles with no volume exclusion, ML-Space particles are volume excluding and Smoldyn particles can be either. Particles can react, diffuse, be confined by surfaces and bind to membranes. Smoldyn and MCell particles are simple, but ML-Space particles can contain other particles and support hierarchical composition. ML-Space and MCell models are specified with a natural and elegant rule-based approach similar to BioNetGen [7], and Smoldyn supports rule-based reaction network generation. However, most stochastic particle simulators only support fixed geometries, and cannot model mechanical or electrical processes or dynamic morphology. Multi-scale simulations are challenging because these simulators because do not have a way of describing continuous fields.

Transport Dissipative Particle Dynamics (tDPD) [11] combine features from Systems Biology and MD to create a very general multi-scale modeling paradigm. tDPD augments the simple MD particles with additional state variables such as chemical concentrations, and adds continuous chemical reaction networks. tDPD particles can represent either solids or fluids. tDPD is ideally suited for multi-scale models because it represents both large, coarse-grained particles which interact with small diffusing particles represented as continuum concentrations. tDPD is a relatively new modeling approach, hence most tDPD simulation appear to be hard-coded in general-purpose programming languages. To our knowledge, no model specification language exists for tDPD. No tDPD simulation appears to support reacting particles as with Reactive MD, nor do any tDPD simulations support discrete events as do the systems biology approaches.

Mechanica builds on the tDPD and incorporates reacting particles and discrete events, and defines a model declaration language.

3 Approach

The Mechanica modeling language (MML) enables construction of mechanistic models of natural phenomena. MML is based on a functional subset of Microsoft Typescript, and includes conventional constructs such as variables and functions, but adopts syntactic concepts from POVRay, a widely used spatial scene description language for geometry description, and pattern matching rules inspired by ideas from BioNetGen, OCaml and Mathematica.

MML has two basic constructs: objects and processes, to reflect the objects and processes found in naturally occurring phenomena. Objects in MML can be a conventional primitive type (e.g., scalars, integers, booleans, enums), or a composite. An object is an instance of a type. A type defines the object’s structure and layout, and a state space, or the set of permissible values that can be stored in the object. Objects have an identity, a symbol which names and refers to the object. MML introduces new types such as particles, concentrations, amounts, spatial regions, and spatial fields with special semantics to support spatial hybrid systems modeling.

MML represents materials as a collection of particle and link objects. MML particles typically represent “ parcels” of small pieces of real materials rather than atoms. Links are persistent structural relationships which connect two or more material objects, and apply a force between the connected objects. Links generalize bonded relationships from molecular dynamics, with two main differences: links can be dynamically formed and broken, and their functional form is user spec-
ified rather than fixed. This particle-based approach enables MML to create a completely unified treatment of all materials. Solids are strongly bonded particles, fluids are weakly bonded particles, and Mechanica can simulate any range in between.

Processes in MML are different from processes in the traditional computing sense in which processes typically are a sequence of instructions that define a series of discrete state changes in objects. MML processes, on the other hand, define either continuous or discrete transformations, which includes creation, deletion and state change. Processes map a set of inputs to a set of outputs, and define a transformation as either a discrete atomic operation or a continuous operation. Both kinds of processes can have \( \text{when} \) and \( \text{while} \) predicates which specify the conditions for activating and deactivating the process. Processes have \textit{body} which calculates a probability (real value, \([0, 1]\)) for discrete processes or a \textit{rate} for continuous processes. The probability for discrete processes defines the likelihood that the process will be applied if its predicates are true; the probability defaults to 1 if not present. The rate for continuous processes specifies the rate of transformation of its inputs to outputs. Process and function definitions are side effect free – they cannot directly modify non-local symbols.

Discrete processes are similar to events in Modelica or SBML. They obey the same semantics as SBML events, and we evaluate them using the SBML event handling algorithm we developed in [15]. Discrete process definitions specify a \textit{pattern} of what objects to act on, and conditions that specify when to trigger the process. Like SBML, discrete processes instantaneously change variable values. Additionally MML discrete processes can create and destroy object instances. (Note: this paper focuses on continuous processes and does not discuss discrete processes in detail.)

Continuous processes are a generalization of chemical reactions and define the time evolution of participating \textit{continuous variables} such as \textit{concentrations, amounts}, real or complex numbers. Continuous variables can be attached to any particle type following [11]. Thus, as the particles move, the attached chemical amounts are advected along with them. Continuous processes definitions specify the \textit{rate} at which the process occurs.

Particles are the basic building blocks of materials. They have a position and optional attributes such as spatial extent, orientation, mass. They can be point particles or have complex shapes. Links can be applied either explicitly or implicitly. A link can be applied explicitly to a set of material objects, and this set will remain explicitly bound until the link is removed. The link can be removed either explicitly or by defining a \textit{while} predicate in the link definition. Links can also be implicitly applied to a \textit{spatial region} by specifying a \textit{when} predicate. Here, a process will monitor the \textit{when} predicate, and apply the link to any set of objects that match the predicate. In this sense, implicitly applied forces behave similarly to long-range or non-bonded interactions in MD.

We can define a simple particle-link system with a pair of point particles, \( a \) and \( b \), located at \((0, 0, 0)\), and \((1, 0, 0)\) respectively, and explicitly connect them with a spring link as

\[
a: \text{particle}(0,0,0,\text{mass}=1); \ b: \text{particle}(1,0,0,\text{mass}=1); \\
\text{link}(a,b){\text{-k*(1-dist(a,b))}};
\]

This particular link defines a Hookean spring force, with a rest length of 1, where the \textit{dist} function calculates the distance between two objects. Link definitions however can contain arbitrary user-specified force definitions. This yields a very flexible model because users may specify the interaction force to be as complex as they like. For example, a two body force could be as simple
as a Hookean interaction, or as complex as a dipole, quadrupole or octopole interaction.

A biological cell has on the order of $10^{14}$ atoms, so it is simply not computationally feasible to model a cell at all-atom resolution. Many smaller ($< 100$nm) sized molecules are soluble and readily diffuse through solution. If we are interested in micro- rather than nano-scale phenomena, we may approximate diffusing chemicals as continuous, spatially varying concentrations. MML has concentration and amount data types which can be attached to any material object. Concentration values are automatically scaled by the volume of the material object that it is attached to. Say a particle has a concentration attribute and the particle’s volume increases, then the value of the concentration decreases. As each material point moves, the attached values move along with it. Chemical concentrations can be attached to any kind of material such as fluids (solvents), surfaces (cell membranes) or fibers, and they can also be attached to spatial regions. Concentration or amount types can be added to any object by adding them to the object’s definition. For example, we could create a new particle $p$ with attributes $A$ as a concentration, $B$ as a constant concentration (boundary value), and $C$ as an amount by:

$$p: \text{particle} \{ A: \text{conc}; B: \text{const conc}(5.0); C: \text{amount}(1.23); \}$$

We may want to fill a region with material particles. Fluid particles have a set of forces act on them which effectively models a fluid. To create a spherical fluid droplet, we first define a MaterialRegion, which is bounded by a surface, and fill it with fluid particles. The fill function determines the available space inside a region and packs it with objects of the given type.

```
MaterialRegion{
    surface: Sphere \{ radius:5, resolution:1 \};
    body: fill(type=FluidParticle\{radius=5\});
}
```

Material regions can also have homogeneous, spatially uniform interiors if we do not specify a body. Homogeneous regions can behave as a liquid bag if we attach a volume preservation link to the object. When concentrations, amounts or other attributes are attached to a homogeneous material region, their value is constant for the entire region.

Chemical reactions are fundamentally important in biology. Reactions define the transformation of a set of reactants into a set of products and can be readily described by processes. A set of continuous processes between a set of inputs (reactants) and outputs (products), such as a set of concentrations, defines a reaction network as in Fig. 1. The Mechanica compiler reads the process definitions and generates a system of ordinary differential equations (ODEs) which define the time evolution of reactants and products. Process definition syntax is similar to conventional chemical reaction notation, and is based on Microsoft Typescript type specification syntax. For example, to add a chemical reaction network to an object, one would simply write a set of process definitions in the object definition. Process definitions begin with the proc keyword, have an optional name, a set of inputs, a set of outputs and a reaction rate expression as in Fig. 1. If the object definition does not already have local definitions corresponding to the product names from a process definition, then the process definition implicitly adds concentration types in the object definition for these names. We do this as a user convenience since chemical reaction networks are among the most common modeling tasks.

Processes can define intra-particle reactions, such as a flux between values located at neigh-
boring particles. Thus, processes can define general reaction-transport problems. Spatial process definitions are written and behave identically to processes confined to one object. The spatial information is simply added in the input argument definitions. For example, to define a Fickian (passive diffusion) flux of the value of A attached to material objects, a and b of type MyMaterial, we would write:

```
proc (a: MyMaterial .A) -> (b: MyMaterial .A)
  when (dist(a,b) < 5) {k * (a - b)};
```

This process definition has a when clause that defines a cutoff distance of 5, and the dist function recognizes that the values are located at different locations. Fickian diffusion flux of continuous valued concentrations between material points is the same approach that [11] used to model advection-diffusion-reaction, \( \frac{dC_i}{dt} = \nabla (D_i \nabla C_i) + Q_i \), \( i = 1, \ldots, N \). They validated their approach against a variety of analytical solutions and demonstrated that augmenting a particle fluid model with scalar values at each material site incurs negligible computational overhead.

```
type MyCell : MaterialRegion {
  surface : Sphere {
    radius:5, resolution:1
  };
  proc (A) -> (X) {k1 * A};
  proc (X, 2 Y) -> (3 X) {
    k2 * X * Y**2
  };
  proc (B, X) -> (Y, D) {
    k3 * B * X
  };
  proc (X) -> (E) {k4 * X};
}
```

Figure 1: Mechanica source code for creating a MaterialRegion derived type which is bounded by a spherical surface composed of a set of points and triangular faces. Four local continuous transformation processes define a reaction network between symbols A, B, D, E, X and Y. The Mechanica compiler generates a system of ordinary differential equations from the continuous process definitions. The compiler also generates a set of particles to form the bounding surface of the MyCell type. The right column shows a screenshot of the MyCell object in its initial configuration, then at a later time after it moves along a surface.

**A field** is a mapping between a value at any spatial location and a set of values associated with material points. In general, any continuously valued field, \( A(\mathbf{r}) \) can be approximated as a sum over a set of appropriate kernel functions, \( A(\mathbf{r}) = \sum_i f(A_i, \mathbf{r}, \mathbf{r}_i) \), where \( A_i \) is some quantity located at the source location \( \mathbf{r}_i \), and \( \mathbf{r} \) is the field location. This definition enables us to define any continuous field as a function of the material. For example, the kernel function for an electric scalar potential would be \( q/4\pi\varepsilon_0|\mathbf{r} - \mathbf{r}_i| \). In general, if the value of some scalar quantity \( A \) is known at a finite set of material points, then the value may be approximated at any location in space \( A(\mathbf{r}) \) with a suitable interpolation function [12]. This allows us to attach a chemical concentration to a material point and evaluate that concentration at any point in space. For example, one might define a chemical compound attached to solvent points and read the chemical concentration value at some non-solvent
material point. We provide default kernels for chemical concentrations and charge, but users are free to define their own kernels.

Most traditional languages employ symbol scoping blocks. These are regions of code that define where a symbol (variable definition) is visible. Scoping blocks can contain other scoping blocks. Depending on the language, code in the child blocks can usually access variables defined in parent blocks. Lexically scoped languages resolve non-local symbols to the scoping block in which the function was defined. Dynamically scoped languages such as PERL resolve non-local symbols to the scoping block in which the function is executing.

In order to account for the spatial nature of our problem domain, we introduce a new kind of scope resolution we call spatial scoping. Spatial scoping is based on the idea that processes represent a physical process, so processes always have to execute somewhere in space. The present location of a process is implicit, much like the this pointer is implicit in Java. Non-local symbols in process bodies resolve to the present spatial environment in which the process currently runs. Spatial scoping is similar to dynamic scoping in that symbol resolution depends on the calling context of a function; however, in spatial scoping, the calling context is a containing spatial region.

Users typically never have to explicitly keep track of coordinate systems; instead the runtime manages all this. All users have to know is the name of the value they want to read. Spatial scoping treats different kinds of spatial objects uniformly. For example, consider a transformation process which converts one substance to another, where the rate (the body of the transformation process definition) depends on the concentration of some other substance,

\[
\text{proc } (A) \rightarrow (B) \{k1 * A * C\};
\]

Here, \(C\) is a non-local symbol, so the runtime will search in the containing spatial environment. This process for example could be attached to a fluid particle. This particle type may have a \(C\) attribute attached to it, in which case \(C\) resolves to that value. If the particle does not have such an attribute, then the compiler will check if the containing spatial environment defines the \(C\) symbol. The containing spatial region may define the \(C\) as a scalar, in which case the value of \(C\) is uniform for the entire region, and this symbol resolves to that value. Or \(C\) may be defined as a spatially varying field, in which case the compiler will generate code that reads from that field at the present location of the particle. If this symbol cannot be resolved in any containing environment, and there is no field definition, then the compiler will report an error.

Processes in MML are very general, being used to define not only transformations of continuous valued objects like amounts and concentrations, but also discrete objects such as particles, materials, and other composite objects. Processes acting on discrete objects have the identical syntax as processes acting on continuous objects, the only difference is the semantic interpretation of the process body. A discrete process body is expected to return a probability value, a real value between zero and one, as opposed to a reaction rate for continuous processes. For example, line 1 defines two new types, \(A\) and \(B\), derived from particle. Line 2 defines a process which consumes two discrete particles of type \(A\), and produces one particle of type \(B\). Line 3 defines a process which consumes a single object of type \(B\) and produces two objects of type \(A\) with random probability.

1 type A:particle{radius :1}; type B:particle{radius :2};  
2 proc (a:A, b:A) \rightarrow (B) \text{ when } (\text{dist}(a,b) < 5) \{\text{exp}(-\text{dist}(a,b)**2)\};  
3 proc (a:B) \rightarrow (A, A) \{\text{rand()}\};
Processes can also be used to change the values stored in one or more attributes. Process can be defined so that their inputs have to be in a certain state, so that they match a particular pattern. We borrow the record pattern matching syntax from OCaml to specify the state of an object where a process can be applied. For example, say an object of type A has an attribute named activated which is an enum, and we want to toggle this from Inactive to Active when this object is near another object of type Activator.

\[ \text{proc} \quad (a:A\{\text{activated} = \text{Inactive}\}, b: \text{Activator}) \rightarrow \\
(a\{\text{with activated} = \text{Active}\}, b) \quad \text{when} \quad (\text{dist}(a, b) < 5); \]

The with keyword is borrowed from OCaml, and specifies that the attributes in the with expression should take on new values, but all other attributes should remain unaltered. The second argument, b of type Activator, is unaffected and passes straight through. The body of this process is omitted, which defaults the probability to 1 and tells the compiler that this process should always be applied whenever these two particles match the when pattern, i.e. are within a distance of 5.

Many proteins have binding sites where other molecules can attach, altering the behavior of the protein. MML introduces a new data type called site to represent binding sites; any spatial object can have binding site attributes. We may for example add two binding sites to a particle type, and use a process to bind two objects together dynamically when certain conditions are met.

\[ \text{type} \quad A: \text{particle}\{s1: \text{site}(\text{empty}); s2: \text{site}(\text{empty})\}; \]

\[ \text{proc} \quad (a:A\{s1=\text{empty}\}, b:A\{s2=\text{empty}\}) \rightarrow \\
(\text{link}(a.\text{s1}, b.\text{s2})\{-k*(1-\text{dist}(a-b))\}) \quad \text{when} \quad (\text{dist}(a, b) < 5); \]

This process binds the s1 binding site of a to the s2 binding site of b with a new link. When a link or some other connector attaches to a binding site, the Mechanica semantics state that that binding site should become non-empty.

Mechanica combines a tDPD system with discrete events and dynamic connectivity. Any variable that is produced or consumed by a continuous process is a continuous variable; the set of continuous variables, \(C_1, C_2, \ldots, C_n\) form the continuous state vector \(C(t)\). Each process definition body can access the present continuous variable states, particle attributes, and constant parameters to calculate the process transformation rate. The set of continuous processes defines a system of ordinary differential equations (ODEs) in the form of

\[
\frac{d}{dt} C = \begin{bmatrix}
C_f \\
C_r
\end{bmatrix} = \begin{bmatrix}
N \cdot \nu(C, r, v, p) \\
f(C, r, v, p)
\end{bmatrix}.
\]

The state vector comprises two partitions: \(C_f\) is the vector of independent continuous variables (which participate in a transformation processes, i.e., a reaction network), and \(C_r\) is a vector of variables which are defined by rate processes. The vector \(p\) consists of time independent parameters. The reaction network of \(m\) continuous variables and \(n\) transformation processes defines the \(m \times n\) stoichiometry matrix \(N\). Each stoichiometric element, \(N_{i,j}\) is the net number of continuous variables \(i\) produced or consumed in transformation process \(j\). All of the transformation process definitions combine to form the transformation rate function \(\nu\). The second part of the state vector, \(C_r\), is a set of variables which form a system of conventional ODEs, i.e. each element in this vector is defined by a rate process. Only the Mechanica runtime can directly change variable values.
The net force on each particle is the sum of the linked, $F^L$, conservative, $F^C$, random, $F^R$, dissipative, $F^D$, and external, $F^{ext}$ forces. The linked force is the sum of the linked relationships between the particles. The conservative force is a soft interaction which ensures volume exclusion and keeps materials from interpenetrating. The dissipative force represents the effects of viscosity, and the random force represents the effects of thermal fluctuations. Integration time steps can be large because these inter-particle forces are soft and have short-range interactions. More details on the conservative, random and dissipative forces can be found in [11]. The time evolution of particle positions, $r$, and velocities, $v$, is defined as

$$\frac{d^2r_i}{dt^2} = \frac{dv_i}{dt} = \frac{1}{m_i} \mathbf{F}_i = \frac{1}{m_i} \left( \sum_{i \neq j} \left( F^L_{ij} + F^C_{ij} + F^D_{ij} + F^R_{ij} \right) + F^{ext}_i \right).$$

(2)

The runtime calculates the time evolution of the model via *time slicing* – time is partitioned into a series of discrete time steps. Each time step has three phases: (A) evaluate and apply the when and while predicates of the discrete processes determine if they should be triggered, and apply them. Links are attached and detached in phase 1 according to their when and while predicates. (B) integrate the continuous state vector variables according to eqn. 1 and (C) integrate the particle positions according to eqn. 2.

### 3.1 Mechanica model of chemotaxis

Chemotaxis is a biological process which describes the motion of a cell towards or away from a chemical signal. We use chemotaxis as a biological example to illustrate the coupling of chemical and mechanical processes and demonstrate how we can represent a few chemotaxis mechanisms in Mechanica. Cells undergoing chemotaxis can sense spatial gradients in chemoattractant concentrations. Chemoattractants are frequently secreted from some source, such as a bacterium at a specific location. Here, we define a rule to secrete a chemoattractant from a fixed point source in space. We first define the the model environment with an an explicit solvent. We write a rule in the top level spatial region which fills it with explicit solvent particles (line 1). This rule states that any space that is not filled with sub-objects gets filled with solvent particles. This rule creates a symbol called *solvent* which is of type water. *CXC* is a common biological chemoattractant, which we will secrete from a point source located at $(20,20,0)$ (line 2). This rule increases the concentration of CXC on the solvent particles whenever they are within a distance of 5 from the point $[20,20,0]$. Because the CXC symbol is listed as a product, the rule implicitly adds the symbol CXC as a chemical concentration attribute attached to the solvent particles. The solvent will transport the CXC along with it, but we want to allow the CXC to diffuse through the solvent. We can use the built-in diffusion rule or we can write our own. Say we want simple Fickian diffusion between solvent particles (line 3). This process defines a transformation process to represent diffusion of solute between neighboring particles. It has a flux rate of $k*(a-b)$, where $k$ is a constant, and $a,b$ are concentrations of CXC located at neighboring particles.

```markdown
1 solvent:fill(type=Water);
2 proc () -> (b:solvent.CXC) when (dist([20,20,0], b) < 5)
3 {aRateConstant};
```
Our cell model will only define a single basic cell object that sticks to a horizontal surface. In order to represent a single cell, we can define a basic MaterialRegion derived type as in Fig. 1, where the initial cell is shown in the top-right. We can place a single instance centered at \([0,0,2.5]\) via

\[
\text{mycell: MyCell \{origin: [0,0,2.5]\}}
\]

Cells frequently stick to the surface that they’re on. We can represent this adhesion with a link that causes the cell’s surface to stick to the base surface. Stickiness tends to be asymmetric, so we can write the link rule to attach at a shorter distance than when it releases. We can think of it as similar to a Velcro adhesion, where the two components must be close to activate, but the force can hold on a bit longer when it stretched. In addition to the \(\text{when}\) clause, links can also have a \(\text{while}\) clause which can define a deactivation condition that differs from the activation condition. This rule creates links which connect the particles of the cell’s surface to the floor of our simulation domain.

\[
\text{link(a: mycell.surface, b: BoundingPlanes[FLOOR])}
\]

\[
\begin{align*}
\text{when} & \ (\text{dist}(a,b) < 0.5) \quad \text{while} \quad (\text{dist}(a,b) < 2) \\
& \{-k*(\text{dist}(a,b))\}
\end{align*}
\]

In our simple model we approximate the cytoplasm as water. We fill the body of the cell with water particles by adding the following line to the cell definition.

\[
\text{mycell.body: fill(type=Water)}
\]

A biological cell’s surface often contains many chemoattractant receptors. Essentially these bind and degrade diffusing chemoattractant molecules in the fluid medium around the cell, and initiate a spatially varying signaling cascade in the cell. We observe that the receptor can be either active or inactive, and only active receptors initiate signaling inside the cell. We can model these observations with the following rules: Line 1 creates a surface bound concentration of active receptors \(\text{mycell.body.ARectpt}\), with an initial concentration of 0.0. Line 2 represents passive diffusion of solvent CXC to cell’s surface, with a diffusion rate of \(k*(a - b)\). Lines 3-4 define a reaction that consumes CXC on the surface and produces active receptor \(\text{ARectpt}\). There are a finite number of receptors, so we want this reaction to saturate, hence the form of the reaction rate.

\[
\begin{align*}
\text{mycell.surface.ARectpt: conc(0.0)}; \\
\text{proc (a: solvent.CXC) -> (b: mycell.surface.CXC)} \\
\text{when} \ (\text{dist(a,b) < 5}) \ {k * (a - b)}; \\
\text{proc (a: mycell.surface.CXC) -> (b: mycell.surface.Recept)} \{ \\
\text{k2 * a * exp(-b**2)}; \\
\}
\end{align*}
\]

Signaling cascades inside the cell usually involve many stages, but here we assume that an internal compound Rho (known to induce actin polymerization) can be activated by an active surface receptor, \(\text{ARectpt}\). Many chemotaxis models propose that actin filament polymerization in the leading edge of a migrating cell is principally responsible for lamellipodium extension. We approximate this mechanism with a set of spring-like forces that push the cell nucleus away from the cell surface. In the presence of active Rho, the springs elongate and adjust their applied force between the nucleus and the surface particles; otherwise they have no effect. We model Rho as a cytosol diffusing
compound which decays over time. Line 1 creates a new concentration called \( \text{Rho} \) attached to the body of the cell (the cytoplasm). Line 2 produces \( \text{Rho} \) on cytoplasm particles at a rate proportional to surface \( \text{ARecpt} \) concentration. This is an example of spatial scoping, where the reaction occurs at a cytoplasm fluid particle (the reaction is producing \( \text{Rho} \) on the cytoplasm particles), so symbols referring to concentrations not attached to the particle resolve to the spatial field produced by these concentrations. The \( \text{mycell.surface.ARecpt} \) symbol resolves to a value that approaches the exact surface concentration value when the cytoplasm particle is near, but falls off rapidly if the cytoplasm particle’s distance away from the surface is high. Line 3 defines a passive diffusion of \( \text{Rho} \) between neighboring cytoplasm fluid particles. Line 4 cause \( \text{Rho} \) to decay at a rate of \( k \cdot [\text{Rho}] \), or proportional to the concentration of \( \text{Rho} \). Lines 5-7 define a set of links that attaches the cell nucleus particle to the cell surface particles. Spatial scoping evaluates non-local symbols in link definitions at the midpoint of the link. Thus, when cytoplasm carrying high concentrations of \( \text{Rho} \) move near the link midpoint location, the \( \text{Rho} \) symbols evaluates to a high value, which causes the spring force to increase. If the concentration of \( \text{Rho} \) at the link location decreases, say, the link moves away or the \( \text{Rho} \) decays at this location, the spring force will contract, causing the cell membrane to retract.

```
1 mycell.body.Rho:conc(1.5);
2 proc () -> (a:mycell.body.Rho) {k * mycell.surface.ARecpt};
3 proc (a:mycell.body.Rho) -> (b:mycell.body.Rho)
   when (dist(a,b) < 5) {k * (a - b)};
4 proc (a:Rho) -> () {k * a};
5 link(a:mycell.nucleus, b:mycell.surface) {
6   -k * (dist(a,b) - restLength + k2 * Rho);
7 }
```

### 3.2 Simulator Details

The Mechanica simulator is loosely based Model-View-Controller paradigm. It is comprised of five key objects: Simulator, Model, View, Integrator, and Compiler. The Simulator is the top-level object which orchestrates the interaction of the others. It uses the Compiler to read a source MML document(s) and generate a Model. The Model contains the compiled form of the MML model and the present state. The Integrator calculates the time-evolution of the Model. The View displays the present model state in an OpenGL window. The Model notifies the view when its state changes so that the View can update the display.

The front end of the Mechanica compiler is a conventional recursive descent parser which generates an abstract syntax tree (AST). The semantic analysis phase of the compiler, however, differs from procedural programming language compilers. Unlike procedural languages where there is a reasonably close mapping between language statements and the resulting machine instructions, MML statements specify rules and relationships rather than explicit instructions. The Mechanica compiler analyzes the model processes and rules and use this information to generate the two time evolution systems of equations, eqn. (1) and eqn. (2). The Mechanica compiler performs semantic analysis of the continuous process definitions using techniques we established in [15]. The Mechanica compiler presently generates C code, and uses GCC to compile this into a shared library.
The compiler then creates a new Model object which loads this shared library. Although MML object definitions appear similar to a C struct, our in-memory storage is different. All the continuous variables are stored in a single contiguous array, and the particle position and velocity vectors are also stored in contiguous arrays.

The Mechanica integrator uses CVODE [9] to calculate the time evolution of the continuous variables. We chose the contiguous memory layout to match CVODE. At each time step, CVODE operates in-place on this shared memory block. We also use CVODE’s root finder to check if any when/while predicates should trigger discrete processes. The Mechanica Integrator uses a modified version of Pedro Gonnet’s mdcore library (http://mdcore.sourceforge.net/) to calculate the time evolution of the particle positions. The contiguous array storage of the particles and velocities also matches mdcore’s layout, thus mdcore operates in-place with no memory copying. mdcore uses the compiled MML model force function to calculate inter-particle potentials. The integrator uses the distance cut-off information from the link predicates to specify a potential cut-off distance for mdcore, which mdcore then uses to construct optimal Verlet and cell lists. Unlike traditional molecular dynamics or particle systems, the particle count in Mechanica models changes over time. The runtime initially allocates a large block, more than the initial particle count, and as particle count increases, it performs a realloc on that block. We are investigating more efficient approaches for dealing with this challenge.

4 Conclusion

We are developing the first known modeling language which can represent generalized continuous transformation processes, forces, fluid flow, and material state transition in the same language. The Mechanica language enables users to create physically motivated models of complex natural phenomena using mechanistic instead of computational building blocks. Mechanica is currently under active development, all source code and binaries will be made available on the Mechanica website, (http://www.mechanica.org) under a permissive open source license (BSD/GPL dual license).

acknowledgments

We acknowledge generous financial support the National Institutes of Health, National Institute of General Medical Sciences, National Institute of Environmental Health Sciences and National Institute of Biomedical Imaging and Bioengineering, grants U01 GM111243, R01 GM076692 and R01 GM077138. We thank Dr. Marie Gingras, Dr. Amr Sabry and Dr. Paul Macklin for their discussion and insights.

References

[1] Andrews, S. S., N. J. Addy, R. Brent, and A. P. Arkin. 2010, March. “Detailed Simulations of Cell Biology with Smoldyn 2.1”. PLoS Computational Biology vol. 6 (3), pp. e1000705.
[2] Beal, J., S. Dulman, K. Usbeck, M. Viroli, and N. Correll. 2013. “Organizing the Aggregate: Languages for Spatial Computing”. In Formal and Practical Aspects of Domain-specific Languages Recent Developments, pp. 436–501. Hershey, PA, Information Science Reference.

[3] Belloli, L., G. Wainer, and R. Najmanovich. 2016. “Parsing and model generation for biological processes”. In Proceedings of the Symposium on Theory of Modeling Simulation. ACM.

[4] Bittig, A., and A. Uhrmacher. 2016. “ML-Space: Hybrid Spatial Gillespie and Particle Simulation of Multi-level Rule-based Models in Cell Biology”. IEEE/ACM Transactions on Computational Biology and Bioinformatics vol. PP (99), pp. 1–1.

[5] Brodland, G. W. 2015. “How computational models can help unlock biological systems”. Seminars in Cell & Developmental Biology vol. 47-48, pp. 62–73.

[6] Cardelli, L., and P. Gardner. 2010. “Processes in space”. In Conference on Computability in Europe, pp. 78–87. Berlin Heidelberg, Springer, Programs.

[7] Faeder, J. R., M. L. Blinov, and W. S. Hlavacek. 2009, January. “Rule-Based Modeling of Biochemical Systems with BioNetGen”. In Systems Biology, pp. 113–167. Totowa, NJ, Humana Press.

[8] Gruenert, G., B. Ibrahim, T. Lenser, M. Lohel, T. Hinze, and P. Dittrich. 2010, June. “Rule-based spatial modeling with diffusing, geometrically constrained molecules”. BMC bioinformatics vol. 11 (1), pp. 307.

[9] Hindmarsh, A. C., P. N. Brown, K. E. Grant, S. L. Lee, R. Serban, D. E. Shumaker, and C. S. Woodward. 2005. “SUNDIALS: Suite of nonlinear and differential/algebraic equation solvers”. ACM Transactions on Mathematical Software (TOMS) vol. 31 (3), pp. 363–396.

[10] Jensen, K., L. M. Kristensen, and L. Wells. 2007. “Coloured Petri Nets and CPN Tools for modelling and validation of concurrent systems.”. International Journal on Software Tools for Technology Transfer vol. 9, pp. 213–254.

[11] Li, Z., A. Yazdani, A. Tartakovsky, and G. E. Karniadakis. 2015, July. “Transport dissipative particle dynamics model for mesoscopic advection-diffusion-reaction problems”. Journal of Chemical Physics vol. 143 (1), pp. 014101.

[12] Liu, M. B., and G. R. Liu. 2010. “Smoothed Particle Hydrodynamics (SPH): an Overview and Recent Developments”. Archives of computational methods in engineering vol. 17 (1), pp. 25–76.

[13] Párvu, O., D. Gilbert, M. Heiner, F. Liu, N. Saunders, and S. Shaw. 2015, April. “Spatial-Temporal Modelling and Analysis of Bacterial Colonies with Phase Variable Genes”. ACM Transactions on Modeling and Computer Simulation (TOMACS) vol. 25 (2), pp. 13–25.
[14] Schöneberg, J., and F. Noé. 2013, September. “ReaDDy - A Software for Particle-Based Reaction-Diffusion Dynamics in Crowded Cellular Environments”. *PLoS ONE* vol. 8 (9), pp. e74261.

[15] Somogyi, E. T. 2014, December. *Simulation of electrochemical and stochastic systems using just in time compiled declarative languages*. Ph. D. thesis, Indiana University, Bloomington.

[16] Stiles, J. R., T. M. Bartol et al. 2001. *Monte Carlo methods for simulating realistic synaptic microphysiology using MCell*. CRC Press, Boca Raton, FL.

[17] van Duin, A. C. T., S. Dasgupta, F. Lorant, and W. A. Goddard. 2001, October. “ReaxFF: A Reactive Force Field for Hydrocarbons”. *The Journal of Physical Chemistry A* vol. 105 (41), pp. 9396–9409.