Systems biology

GetBonNie for building, analyzing and sharing rule-based models

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ABSTRACT

Summary: GetBonNie is a web-based application for building, analyzing and sharing rule-based models encoded in the BioNetGen language (BNGL). Tools accessible within the GetBonNie environment include (i) an applet for drawing graphs that correspond to BNGL code; (ii) a network-generation engine for translating a set of rules into a chemical reaction network; (iii) simulation engines that implement generate-first, on-the-fly and network-free methods for simulating rule-based models; and (iv) a database for sharing models, parameter values, annotations, simulation tasks and results.

Availability: GetBonNie is free at http://getbonnie.org.

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1 INTRODUCTION

The systems-level dynamics of molecular interactions in cellular regulatory systems are difficult to model using approaches that rely on explicit specification of a chemical reaction network. The reason is combinatorial complexity (Hlavacek et al., 2006), the potential of molecular interactions to generate large numbers of chemical species and reactions. Rule-based modeling approaches have been developed to address this problem (Hlavacek et al., 2006). In these approaches, molecules and molecular complexes are typically represented using graphs, or the equivalent, and molecular interactions are represented using (graph-rewriting) rules. A rule implicitly defines the reactions that can be generated by the molecular interaction that it represents.

A variety of software tools have been developed to facilitate rule-based modeling (Hlavacek et al., 2006). These tools, which provide complementary capabilities, are based on similar formal foundations. However, interoperability of these tools is limited because a standard format, such as the Systems Biology Markup Language (SBML) (Hucka et al., 2003), has yet to emerge for the exchange of rule-based models. Moreover, a public repository, such as the BioModels Database (Le Novère et al., 2006) in which models are stored in SBML, is unavailable for rule-based models. SBML is notable because it allows chemical reaction networks to be specified in a form that can be processed by (numerous) SBML-compliant software tools (http://sbml.org). Although a set of rules can be translated into a chemical reaction network, a rule set often implies a vast reaction network, and as a result, the current version of SBML, Level 2, is not suited for efficient encoding of rule-based models. This deficiency has been recognized and a Level 3 extension of SBML appropriate for representation of rule-based models is under discussion (http://sbml.org/Community/Wiki).

In part, to provide a means to exchange rule-based models in terms of rules (rather than in terms of the networks implied by rules) and, in addition, to provide additional computational infrastructure for rule-based modeling of biochemical systems, we developed GetBonNie. GetBonNie provides a web-based toolkit for building, viewing, simulating and sharing rule-based models encoded in the BioNetGen language (BNGL) (Faeder et al., 2009), which is closely related to K-calculus (Danos et al., 2007a), another language for encoding rule-based models. In GetBonNie, models are stored in an XML-based format, which is defined by an XML schema documented at the GetBonNie web site. The name GetBonNie is an anagram of BioNetGen.

2 FEATURES

GetBonNie combines the capabilities of several stand-alone programs in one freely accessible web-based application. Screenshots are shown in Figure 1. GetBonNie allows users to create rule-based models and share them within a private group or with the public. Public models are each assigned a unique accession number for reference purposes. Model specifications can be either uploaded to or built de novo within the GetBonNie environment, which provides interactive forms to guide model building and simulations (Fig. 1A). Model specifications are automatically checked to ensure that they are well formed. Online help is available.

2.1 Model building and sharing

GetBonNie represents and displays the elements of a model specification as a tree, which can be navigated by a user. A model is built by using web forms to add/delete and edit BNGL code (and associated annotations) at each leaf of a model tree. BNGL
Fig. 1. Screenshots. (A) The Build tool illustrated here provides a form-based interface for specifying a rule-based model using BNGL. The tree view at left allows a model specification to be easily navigated. In this screenshot, the BNGL Textbox Editor displays one of the seven rules that comprise the model specification outlined in the tree view. This specification is available as a BioNetGen input file (simple_egfr.bngl) at the BioNetGen web site (http://bionetgen.org), and it is discussed at length in the tutorial of Faeder et al. (2009). This model is also available in the public library of models of GetBonNie (Model #000002). The Action Assistant allows a user to add actions to a model, such as simulation directives. A user can view a model specification in BNGL format by clicking on the BNGL tab at top center. By clicking on the Build Help tab, a user can view online help. Help tabs are available throughout GetBonNie. (B) The RuleBuilder Lite tool illustrated here can be used to visualize BNGL code entered in the box at bottom. It can also be used to generate BNGL code using the drawing tools at top. In the center of this screenshot, the BNGL code that appears in the BNGL Textbox Editor of the Build tool window shown in panel (A) is displayed according to the graphical conventions of Faeder (2005b). As can be seen, RuleBuilder Lite has a toolbar, a drawing area, a current status indicator (located above the box used to enter BNGL code) and an input box for BNGL code. (C) A contact map (Danos et al., 2007a) produced by the Viewer tool is shown here. This contact map illustrates the molecules, molecular components and molecular interactions defined in Model #000002. The contact map is generated by analyzing a model specification. A gray box represents a molecule, whereas a blue box inside a gray box represents a component of a molecule. The possible internal states of a component are also drawn in blue boxes that appear below component boxes. Lines connect binding partners. The contact map shown here indicates that a tyrosine of EGFR (Y1068) has two states: unphosphorylated (U) and phosphorylated (P). Note that, according to this contact map, the SH2 domain of Grb2 binds tyrosine 1068 of EGFR only if this tyrosine is phosphorylated.
A PHP script parses XML-encoded BNGL to generate drawing of rules into a chemical reaction network, and the resulting network server-side network generation engine can be used to translate a set online and/or download the raw simulation data. Simulation settings systems. When a simulation is finished, a user can plot the results

2009; Danos et al

Brent, 2005) and stochastic network-free methods (Colvin et al

et al 2004), stochastic on-the-fly methods (Faeder et al, 2009). Simulation results are plotted online using the Google™ visualization API. A web forum for discussion of GetBonNie is provided using the phpBB® forum solution (http://www.phpbb.com).

GetBonNie uses the software tools BioNetGen and DYNSTOC (Colvin et al., 2009) as engines for simulation, network generation and network export. BioNetGen provides generate-first and on-the-fly simulation functions and network-generation and export functions. DYNSTOC provides a network-free simulation function.

4 FUTURE DIRECTIONS

The formalism of rule-based modeling enables one to make precise statements about the molecules and site-specific details of the molecular interactions in a system, and the formal elements of a model specification can be associated with annotation for the corresponding molecules and interactions. In the future, we plan to extend GetBonNie to allow biological knowledge, such as protein sequences, to be linked to formal elements of models. This type of annotation capability, which was one of the motivating factors for developing GetBonNie as a web-based application, should enable users to make models more understandable and easier to refine and extend.

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Conflict of Interest: none declared.

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