SPINE 2: a system for collaborative structural proteomics within a federated database framework

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Received February 15, 2003; Revised and Accepted April 9, 2003

ABSTRACT

We present version 2 of the SPINE system for structural proteomics. SPINE is available over the web at http://nesg.org. It serves as the central hub for the Northeast Structural Genomics Consortium, allowing collaborative structural proteomics to be carried out in a distributed fashion. The core of SPINE is a laboratory information management system (LIMS) for key bits of information related to the progress of the consortium in cloning, expressing and purifying proteins and then solving their structure by NMR or X-ray crystallography. Originally, SPINE focused on tracking constructs, but, in its current form, it is able to track target sample tubes and store detailed sample histories. The core database comprises a set of standard relational tables and a data dictionary that form an initial ontology for proteomic properties and provide a framework for large-scale data mining. Moreover, SPINE sits at the center of a federation of interoperable information resources. These can be divided into (i) local resources closely coupled with SPINE that enable it to handle less standardized information (e.g. integrated mailing and publication lists), (ii) other information resources in the NESG consortium that are interlinked with SPINE (e.g. crystallization LIMS local to particular laboratories) and (iii) international archival resources that SPINE links to and passes on information to (e.g. TargetDB at the PDB).

INTRODUCTION

The structural genomics effort is generating a vast amount of data, underscoring the need for database systems and servers that can organize this information (1–4). Structural genomics consortia have been formed to consolidate these efforts. These consortia, including the Northeast Structural Genomics Consortium (NESG), are composed of numerous researchers in disparate locations working cooperatively at each step of the structural determination process, from selection of targets through to analysis of the results. The SPINE (Structural Proteomics In the NorthEast) database (5) was designed to coordinate the efforts of these researchers in the NESG as an information management and data analysis resource.

SPINE was created in 1999 as a data repository with associated data mining tools. Through many revisions, its tracking functionality has expanded to accommodate detailed histories for individual samples, thereby presenting a more complete framework for transmitting information through all stages in high-throughput protein production and structure determination. In the original publication, Bertone et al. (5) described the system architecture and how it was interlinked with specific tools to enable data mining. There have been previous discussions regarding a role of proteomics ontologies in structural genomics (6) and, at this stage, SPINE can be described as the beginning of an ontology of standardized protein properties. Here we present version 2 of SPINE and describe its overall development over the last 3 years.

Our system encompasses several aspects. Firstly, the core of the SPINE system (core SPINE) is a centralized information management system, which tracks protein targets through the structure determination process, from the cloning of expression constructs to final biophysical and structural characterization and submission of PDB coordinate sets, providing histories for particular samples. Secondly, SPINE sits at the center of a federation of computational resources; there are a number of SPINE-integrated web tools, both local and remote, that allow members of the consortium to post further information related to protein targets. Finally, the database

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The authors wish it to be known that, in their opinion, the first three authors should be regarded as joint First Authors
can be analyzed retrospectively to identify factors that contribute to the ease with which individual proteins may be studied. Although the system is tailored to the needs and goals of the NESG, we expect that many of its features could be readily adapted to similar projects.

IMPLEMENTATION OF CORE SPINE SYSTEM

The core database is implemented in MySQL on a Unix platform, with its user interface written entirely in the Perl programming language and integrated with the Apache web server. (Previous versions of the database used PHP extensively; the current implementation provides a more consistent platform.) This approach offers a considerable speed advantage and allows sharing of libraries with offline programs used in the development of future releases and associated tools. The suitability of Perl for systems programming also allows a wide variety of other modules to be used in the server with minimal set-up and administrative overheads, such as the BLAST package (7), Java and Lisp code for data analysis. The core data elements are stored in a number of tables that record the experimental progress of individual targets (Fig. 1). Auxiliary tables control access to the database and record a history of individual changes.

Initially, the basic unit tracked by SPINE was the expression construct. However, with the evolution by the NESG consortium of a systematic protein target selection process (8), these protein targets, each of which may have multiple associated constructs, have been made the focus of the database. All derivative records from the target onwards comprise one-to-many relationships. At the level of protein purification, we have introduced parent–child relationships for individual protein samples, broadening the data structure instead of compressing multiple purification records into a single instance. Most records include some form of unstructured data, often in the form of analytical images that have been uploaded to the server. This has even been extended to encompass Email (see below) related to specific targets.

A key feature of the database is the ability of any registered member of the consortium to add and modify entries via an intuitive web interface. This is regulated by journaling all changes to data (and the user responsible) and by restricting
access to certain entries. More flexible and customized methods of data entry are also possible. The use of direct SQL and the Open Database Connectivity (ODBC) protocols enable a variety of remote interfaces to the server (such as Excel spreadsheets or Java programs), and data interchange uses standard XML or table formats. These features enable bulk uploading of local datasets into SPINE. In the future, development will focus mainly on the schema, the SPINE data dictionary and display functions, leaving data entry as the prerogative of individual users.

INTEGRATION OF CORE SPINE WITH A FEDERATION OF OTHER RESOURCES

The NESG consortium is comprised of various database management systems used to store and search critical data. SPINE provides federation technologies to provide a common unified interface for these diverse systems. The core SPINE database system sits at the center of a federation of information resources diagrammed in Figure 2. The core of SPINE is a relational database handling highly standardized information that interoperates with a set of local resources designed to handle more heterogeneous data that is not readily stored in tables. Some of these features include the incorporation of free text fields, the ability to upload data files and the utility of its data mining tools and servers.

SPINE is also associated with external resources that are coupled together in a loose federation associated with the NESG. These resources can be categorized into three tiers.

Local integrated resources

SPINE is integrated with a number of ‘local’ resources, resident on the same machine and tightly coupled to it. These include the following.

The NESG website. This is built around a wiki (http://wiki.org/) platform that lets users edit or create web pages by using the web browser. This platform allows for easy remote editing of such things as links to related projects and is useful for web-based collaboration.

A structure gallery. This is used for displaying completed 3-dimensional structures of protein targets.

A publication page. This is built on elaborating the NCBI PubMed XML dump to incorporate such things as targets and web sites. It allows the direct cross-referencing of targets, URLs and MEDLINE identifiers.

The target info bulletin board. The idea behind this is that there is a lot of information that people would like to track about a particular target that does not fit into standardized tables. This can be easily sent in the form of simple Email messages that are copied to this bulletin board. These messages are automatically parsed for specific target identifiers and each instance of a target identifier in the archive is linked to its corresponding record in SPINE, and vice versa.

Other NESG resources

There are a number of other computational resources that are part of the NESG project which are connected to SPINE (Table 1). In particular, the diverse needs of experimentalists have led to the creation of several specialized databases within the consortium, dedicated to aspects of the project such as NMR data collection or crystal screening that are not well-served by a single central resource. SPINE is currently being extended to facilitate storage of summary information from these satellite databases and even perform remote queries. Some of the resources that SPINE interoperates with are the PEP (9) cluster viewer at Columbia University and ZebaView target list at Rutgers University, where new target entries are
Table 1. Description of other NESG resources

<table>
<thead>
<tr>
<th>NESG resources</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target resources</td>
<td></td>
</tr>
<tr>
<td>Dynamic target prioritization website</td>
<td>Web site that prioritizes targets that can provide useful information for constructing structural models of other proteins. <a href="http://maat.med.cornell.edu/nseg.html">http://maat.med.cornell.edu/nseg.html</a></td>
</tr>
<tr>
<td>PEP: database for prediction of entire</td>
<td>Used mainly for target selection, PEP provides clustering sequence identity information of potential targets. It also predicts structural and functional features of the targets to aid in experimental analysis. <a href="http://cubic.bioc.columbia.edu/db/PEP/">http://cubic.bioc.columbia.edu/db/PEP/</a></td>
</tr>
<tr>
<td>proteomes</td>
<td></td>
</tr>
<tr>
<td>Zebaview: the official NESG target list</td>
<td>Organizes and reports on the progress of the NESG consortium protein targets. <a href="http://www-nmr.cabm.rutgers.edu/bioinformatics/ZebaView/">http://www-nmr.cabm.rutgers.edu/bioinformatics/ZebaView/</a></td>
</tr>
<tr>
<td>Annotation resources</td>
<td></td>
</tr>
<tr>
<td>PartsList/Gene Census.org</td>
<td>Database web tools that focus on comparing genomes globally. <a href="http://bioinfo.mbb.yale.edu/genome/">http://bioinfo.mbb.yale.edu/genome/</a></td>
</tr>
<tr>
<td>Structure/functional annotation website</td>
<td>This provides detailed information on solved structures. <a href="http://trantor.bioc.columbia.edu/sharon/Target_list_1.html">http://trantor.bioc.columbia.edu/sharon/Target_list_1.html</a></td>
</tr>
<tr>
<td>Structure validation website</td>
<td>A site that summarizes information related to the accuracy and validity of each structure determined. <a href="http://www.cabm.rutgers.edu/~aneerban/NESS_sample_reports/">http://www.cabm.rutgers.edu/~aneerban/NESS_sample_reports/</a></td>
</tr>
<tr>
<td>LIMS systems</td>
<td></td>
</tr>
<tr>
<td>HWI crystallization database</td>
<td>Database for information repository and tracking of crystallization data for robotic crystallization generated at the Hauptman Woodward Medical Research Institute, Buffalo, NY.</td>
</tr>
<tr>
<td>PNNL</td>
<td>Database for information repository associated with Pacific Northwest National Laboratories, principally related to the progress of NMR structures.</td>
</tr>
<tr>
<td>Proteus crystallization database</td>
<td>Database for information repository and tracking for crystallization data generated at Columbia University.</td>
</tr>
<tr>
<td>Rutgers protein production</td>
<td>Laboratory information management system for Rutgers University.</td>
</tr>
<tr>
<td>LIMS</td>
<td></td>
</tr>
<tr>
<td>Toronto protein production LIMS</td>
<td>Laboratory information management system for Toronto Structural Proteomics Initiative at the University of Toronto.</td>
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</tbody>
</table>

Automatically downloaded nightly and inserted into SPINE. Other web resources SPINE is linked with include the SPINS database at Rutgers (17), the PartsList and Gene Census databases at Yale University (18), the Proteus crystallization database at Columbia University and a laboratory information management system (LIMS) system at the University of Toronto.

External archival resources

SPINE is also connected to resources outside the NESG through an evolving portal called SmartLink. This system handles much of the difficulty of translating ORF and structure identifiers and dealing with missing or dangling links. Most of the information pertaining to 3-dimensional structure determinations is transferred to the PDB (10). Other resources that SPINE is connected to include SwissProt (11), PIR (12), BMRB (13), TargetDB (10), the RCSB registry for structural genomics projects and Wormbase (14).

Detailed Sample Tracking and History

The requirements of tracking the progress of a target across multiple institutions include the ability to maintain a list of individual samples and their locations. For example, a protein may be purified at one site, shipped to another for crystallization screening and then sent to a third site for structural characterization. Protein production requires greater flexibility, since not only protein samples but also construct stocks and fermentation batches must be stored and tracked. The current system handles all sample types via tube records, whose contents are determined from their `parent' record (construct, expression or purification). Each `sample tube’ generated in the pipeline of sample production is assigned a unique tube identifier, which eventually will be mapped into a bar coding system. Therefore, a collection of protein samples may be assigned for biophysical analysis without regard for their specific target, since the database automatically determines their history based on tube identifier. This concept has been extended to handle sample plates, which behave as an aggregation of tube records identified by their well number. An example screenshot can be viewed in Figure 3C. With this approach, information pertaining to shipments as well as physical location can be easily associated with sample records. This provides accounting of material transfer between institutions and a more accurate picture of the progress of individual targets.

Discussion

In this paper we have described SPINE version 2, a relational database system that serves as the LIMS for the NESG consortium. This new version serves as the center of a federation of interoperable resources to allow for distributed functionality across the entire consortium. The major goals of the database system include the following.

A distributed LIMS

Constructing a distributed LIMS enables information to be shared among all the members of the consortia. From a vast number of local and external resources, SPINE can incorporate and process data, which can then be made accessible through a simple user browser interface at any location.
Figure 3. Assorted screenshots from the NESG/SPINE web server. (A) The NESG home page, providing links to all the institutions involved and to the gallery of structures produced by them. (B) The SPINE target summary page showing the data recorded in SPINE at each stage of the protein production and structure determination process (blue shading indicates data is present) for every target in the project. (C) One of the new additions in version 2 of SPINE is the ability to track in detail a given sample tube down to a well, plate and physical location. Any changes can be made by clicking on the desired sample entry (blue circle). (D) The NESG structure gallery showing all the structures that have been determined by the NESG consortium. The structure gallery also contains links to the PDB, the BMRB, the Structure Validation Website and the structure/functional annotation website.
Standardized proteomics ontology

An improved representation of protein properties that is standardized will allow for more efficient retrospective analysis of structural and functional information, which can be mined using an incorporated generalized data mining web tool.

Data mining

The schema for SPINE has been designed to facilitate analysis of the collected data to further optimize target selection criteria. Bertone et al. (5) demonstrated the potential data mining capabilities of the SPINE database by developing a decision tree algorithm that was used to infer whether a protein was soluble, and which biochemical properties contributed most to solubility, from a dataset of 562 M. thermoautotrophicum protein expression constructs. The decision tree analysis indicated that protein characteristics such as protein length, hydrophobicity and percent composition of charged residues were the strongest determinants in inferring the solubility of a protein. At present, the number of targets in the NESG is 7866, roughly 15 times the size of the original dataset, and presents a wealth of new information for mining (C.-S. Goh et al., in preparation).

Future directions in interoperation

We envision enhancing the integration of other databases by establishing a tighter interoperation with external databases, particularly those in the second tier.

To this end, we are currently investigating automatic data integration tools to span our heterogeneous federation of databases into what appears as a single (but virtual) database. Such an approach seems like a logical progression since not only does it remove implementation dependencies for each site, but it also manages the interconnections between sites. Users at different project locations and people from the outside could then interface, query and interact with the uniform interface at a single site, rather than considering the details of all the different laboratory databases. This will allow us to adopt a hub and spoke topology of database connections rather than a fully interconnected network, simplifying the interface as compared to an all-with-all federation. It also provides a graceful transition from a very loose federation to a single unified database.

A number of technologies that may be useful in this include the SDSC Storage Resource Broker (SRB), http://www.npaci.edu/DICE/SRB, a piece of client–server middleware that provides a uniform interface for connecting to heterogeneous data resources over a network and accessing replicated data sets), Bio-Kleisli (15) and the Object-Protocol Model (16).

OBTAINING AND USING THE SOFTWARE

The complete code for SPINE (2.0), including table creation commands and the associated data dictionary, may be downloaded at http://spine.nesg.org/download/. Currently a number of the targets in SPINE are publicly available. These correspond to any protein targets whose structure has been solved. A demo version of the SPINE database including this public data generated by the NESG consortium can be accessed at http://spine-dev.nesg.org/. In the future, it is planned that all of SPINE and its data will be made publicly available.

ACKNOWLEDGEMENTS

We thank Michael Baran, Natalia Denissenova and Chi Kent Ho for helpful discussions. All of the authors belong to the Northeast Structural Genomics Consortium.

REFERENCES