Wave enables real-time collaboration between users and also integrates software tools. Various types of content are incorporated in one evolving communication document. Processes are handled in a menu driven user interface, biological data handling including DAS and an internal database communication. We demonstrate its use by implementing robots for gene–data retrieval, manipulation and display. The initial development of SynBioWave demonstrates the power of the underlying Google Wave protocol for Synthetic Biology and lays the foundation for continuous and user-friendly extensions. Specialized wave-robots with a manageable set of capabilities will divide and conquer the complex task of creating a genome in silico.

Availability: The robot is available at SynBioWave@appspot.com, the source code at http://synbiowave.sourceforge.net
Contact: kristian@biologie.uni-freiburg.de
Supplementary information: Supplementary data are available at Bioinformatics online and at http://www.synbiowave.org.

Received on April 5, 2010; revised on August 30, 2010; accepted on September 4, 2010

1 INTRODUCTION

Scientific research is a collaborative effort requiring continuous exchange of ideas and data interlinking researchers at every level of the work flow. Since geographic separation should pose no hindrance, such networking is best achieved using capabilities of the so-called Web 2.0. In the life sciences, numerous databases provide information and web services solve specific tasks. Despite these efforts, a technology to share the research working process from the decision and design phase of an experiment to the data collection and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes. SynBioWave is a software suite that enables multiple distributed users to analyze and construct genetic parts and finally genomes.

2 SOFTWARE

2.1 Usage

As SynBioWave is based on Google Wave, usage differs from typical desktop or database-front ends. A user first opens a Wave client, which is a dynamic web site within a web browser, and creates or opens a conversation object called a Wave. The user may invite further people to this Wave and start to communicate. The SynBioWave environment is initialized by adding the SynBioWave robot (synbiowave@appspot.com) to the conversation. This robot adds a user interface (UI) to the Wave and provides core biological functionality. All users participating in the Wave can collaboratively use SynBioWave in real time.

For example, one user uploads a DNA sequence and another user wishes to translate the sequence and to search for reading frames. Since the latter functions are not included in the SynBioWave core, the user adds the participant/extension ‘synbiowave-translate@appspot.com’, which instantly inserts the translate option in the pull-down menu of the main toolbar. Finally, users can choose between various ways to display biological data (Supplementary Fig. S1). A more extensive user guide is available in the Supplementary Material.

2.2 Architecture

The SynBioWave core consists of one robot, the SynBioWave robot and an associated gadget. Together, these programs generate a UI, communicate with an internal database, and import or export annotated sequence formats. Specialized bioinformatics tasks are implemented in additional robots (Supplementary Fig. S2). Such further robots can be developed based on our extended
robot application programming interface (API), which provides an abstract class for creating robots. Hence, further robots can be built without detailed Wave know-how. By selecting task-relevant robots, end users can dynamically manage the complexity of the SynBioWave front end.

The concept of the SynBioWave core robot for organizing common tasks also overcomes a shortcoming of the original Wave platform, which does not offer coordination between third-party robots that may co-exist within a single Wave.

2.3 Graphical UI

Providing a consistent user experience and organization of robot services posed a major challenge. Therefore, we extended the Wave UI with a dynamically generated toolbar that integrates the features of the robots added to the Wave. This pull-down menu furnishes the web-based SynBioWave with a look and feel similar to a desktop application.

This was achieved by developing a new protocol, named qooXWave. The protocol defines a client-to-server and server-to-client communication for building UI-elements, event reporting and data transmission. Communication is realized via JSON strings that are stored inside a gadget’s state object. Both server (robot) and client (gadget) can access the state object and both can react to changes of this state object (the latter is a feature of the Wave API). The client-side implementation of the qooXWave protocol is based on the rich-internet application (RIA) framework qooXdoo (http://qooxdoo.org). Wave gadgets import the qooXdoo Javascript application into the Wave-client. A robust store-model-view-controller architecture ensures a proper client-side implementation of the complex protocol. This technology can also be used for other scientific or non-scientific Wave extensions and provides a template for implementing new functions into SynBioWave.

2.4 Data handling

Biological information is provided in many different formats in databases, locally stored files and from user input. As Google Wave provides sharing capability and textual editing only, we improved the Google Java API to access, process and convert biological information. A very general internal Java representation of DNA, RNA and protein sequences was created allowing for easy conversion of the internal format to commonly used formats and vice versa (Supplementary Fig. S3). This representation merges the basic information of the GenBank format (http://www.ncbi.nlm.nih.gov/Sitemap/sampleresult.html) with the annotation system of the Google Wave API. Within SynBioWave sequence information is stored in a separate workspace wave, which is automatically made accessible to all users added to the main wave. Import and export to databases is handled via the Java Data Objects (JDO) protocol. Use of JDO enables the user to choose compatible external databases (Supplementary Fig. S4). In addition, we implemented a manager class for the database protocol “Distributed Annotation System” (DAS) (Dowell et al., 2001). This protocol allows users to collect sequences and annotations from DAS-enabled databases, collate the information and display them in a single view within SynBioWave.

Java Wave robots per se represent Http Servlets originally restricted to serve robot events. Therefore, we extended their capabilities to enable file up- and downloads. File accession is handled based on the apache commons-upload project and information is then parsed via BioJavaX classes (Holland et al., 2008). Due to the underlying BioJavaX API, the upload is easily extendable to sequence files in different formats. A simple function for converting directly typed or pasted sequences into the internal sequence format is also provided.

2.5 Extensibility

An important feature of the SynBioWave is extensibility by user contributed robots. The SynBioWave API provides easy access to the aforementioned features allowing a programmer to focus on specific tasks. The supplement gives an example of the API documentation (Supplementary Fig. S5) and a step-by-step developer guide.

3 DISCUSSION

SynBioWave builds upon Google Wave, a real-time online collaboration-software for molecular and synthetic biology. We added several features that are important for scientific tasks and provide a general-purpose extension framework. Wave is still undergoing significant development. A continuous co-development of SynBioWave is envisioned and we, for example, upgraded to Google’s robot API v2. At present, performance of the Google Wave server varies and the long-term availability is unclear. Users can set up their own Wave servers and clients using open source software. (Supplementary Fig. S6). Due to its open architecture, SynBioWave can be adapted to various client–server implementations. To ensure wide access and compatibility with the latest updates, the SynBioWave robot is currently hosted on the Google Wave server. The code of SynBioWave is maintained over the sourceforge platform for open-source projects enabling subversion control and feature request. Primary contact point is the project web site (http://www.synbiowave.org), which provides documentation, a screencast, and a wordpress forum.

ACKNOWLEDGEMENTS

We thank the Qooxdoo development team for technical help.

Funding: Excellence Initiative of the German Federal and State Governments [EXC 294]; Genedata AG, Basel, Switzerland; Roche, Basel, Switzerland; Human Frontier Science Program (LT fellowship to R.G.).

Conflict of Interest: none declared.

REFERENCES


