**Structural bioinformatics**

**ballaxy: web services for structural bioinformatics**

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Associate Editor: Alfonso Valencia

**ABSTRACT**

**Motivation:** Web-based workflow systems have gained considerable momentum in sequence-oriented bioinformatics. In structural bioinformatics, however, such systems are still relatively rare; while commercial stand-alone workflow applications are common in the pharmaceutical industry, academic researchers often still rely on command-line scripting to glue individual tools together.

**Results:** In this work, we address the problem of building a web-based system for workflows in structural bioinformatics. For the underlying molecular modelling engine, we opted for the BALL framework because of its extensive and well-tested functionality in the field of structural bioinformatics. The large number of molecular data structures and algorithms implemented in BALL allows for elegant and sophisticated development of new approaches in the field. We hence connected the versatile BALL library and its visualization and editing front end BALLView with the Galaxy workflow framework. The result, which we call ballaxy, enables the user to simply and intuitively create sophisticated pipelines for applications in structure-based computational biology, integrated into a standard tool for molecular modelling.

**Availability and implementation:** ballaxy consists of three parts: some minor modifications to the BALL library and an integration into the BALL framework and the BALLView application for molecular modelling. Modifications to Galaxy will be submitted to the Galaxy project, and the BALL and BALLView integrations will be integrated in the next major BALL release. After acceptance of the modifications into the Galaxy project, we will publish all ballaxy tools via the Galaxy toolshed. In the meantime, all three components are available from http://www.ball-project.org/ballaxy. Also, docker images for ballaxy are available at https://registry.hub.docker.com/u/anih/ballaxy/dockerfile/. ballaxy is licensed under the terms of the GPL.

**Supplementary information:** Supplementary data are available at Bioinformatics online.

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Received on January 29, 2014; revised on August 7, 2014; accepted on August 20, 2014

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**1 INTRODUCTION**

Workflow systems have become increasingly popular in many areas of computational biology. In addition to the creation and sharing of powerful workflows that can be seen as a modern alternative to shell-based scripting, such systems foster the reproducibility of results in computational science and offer benefits with respect to the deployment and configuration of software packages. This latter part is particularly clear for systems that offer a web-based user interface that can be accessed using a standard web browser with its commonly accepted set of interface metaphors. Installation and maintenance of such systems can be centralized, although parameter settings and special combinations of tools into processing chains can be stored per user and manipulated through the browser. One important example is the popular Galaxy system (Goecks et al., 2010).

In structural bioinformatics, the number of workflow-enabled portals appears to be smaller than in computational genomics, for instance. One prominent example of such a system for computational chemistry is the MoSGrid (Molecular Simulation Grid) portal (Herres-Pawlis et al., 2012), which offers predefined workflows for quantum calculations, molecular dynamics and docking, based on UNICORE and WS-PGrade/gUSE technologies (Gesing et al., 2012). Although many typical use cases are well covered, it remains difficult for novice users to modify or extend the existing MoSGrid workflow.

One reason for the relative scarcity of easily extensible structure-based workflow systems is the difficulty of supporting much of the functionality of an inherently 3D discipline through the confines of a typically 2D workspace of a workflow toolkit, such as the one offered by the browser connected to a Galaxy instance. While sequence-based tools mainly require text-based visualization and 2D plots, sophisticated molecular structure manipulation demands versatile tools that interact with a 3D molecular system.

Hence, a web-based workflow system for structural computational biology would, at present, require constant switching between tools: from the commonly locally installed molecular modelling application to the web service, and vice versa. Recurrent data transfer causes repeated up- and downloads, which are cumbersome and can be time-consuming.

In this work, we present an approach to combine web-based workflow management with a classical powerful molecular modelling interface.
2 IMPLEMENTATION

The first step of our approach consists in an extension of the BALL project (Hildebrandt et al., 2010), where common functionality has been encapsulated in command-line tools. These have then been supplemented with a mechanism for the generation of description files for workflow systems—Galaxy in particular—which then allows for the integration of the tools into a workflow. In the process, we also extended the Galaxy system to support molecular file formats and automatically load the BALL tools.

To solve the problem of providing workflow functionality in combination with a versatile modelling environment, we then extended BALLView (Moll et al., 2005, 2006) by a plugin that tightly integrates communication with a ballaxy server (local or remote) into the modelling toolkit. ballaxy is based on the Galaxy workflow engine and uses its powerful user management, tool handling and workflow environment.

To this end, we extend Galaxy to understand molecular file formats such as PDB or MOL2, and by tools for structural data in the context of molecular modelling and computer-aided drug design. Automated file format detection inside Galaxy uses the python interface of BALL. Structures can be downloaded, visualized and manipulated in BALLView and, through an entry in their context menu, directly uploaded to the server. A browser window embedded into BALLView is then used to interact with ballaxy in the usual fashion to create or run tools and workflows. The results of these workflows can then be directly downloaded into the BALLView instance through the click of a button, where they are displayed in 3D and can be further manipulated or stored.

To provide a collection of useful tools, we further extended the BALL library. These tools will be integrated into BALL version 1.5, which is in development at the time of writing. Besides convenience tools like molecular file converters or a connected component splitter, we currently account for four main application areas: NMR shift prediction (Dehof et al., 2013, 2011a), ligand optimal bond order assignment (Dehof et al., 2011b), pose clustering (Hildebrandt et al., 2013) and docking (Kohlbacher, 2012).

Furthermore, various tools for computer-aided drug design provide functionality to set up ligand-based and structure-based virtual screening workflows. The ligand-based part comprises quantitative structure-activity relationship (QSAR) tools for reading and preprocessing of QSAR datasets, model generation and validation, feature selection and activity prediction. The structure-based part offers tools to read, check and prepare protein structures and virtual compound libraries, for pocket detection, receptor grid generation and protein-ligand docking. Tools for target and antitarget rescoring of initial docking poses allow customized improvement of the docking outcome. Post-docking analysis tools like a ScoreAnalyzer or an RMSDCalculator enable the examination of final docking results and their clustering. All the tools support a standard set of parameters, e.g. for file input, output and help texts. All the tools also support different execution environments: the tool can be used for standard command line usage as well as for integration into a workflow toolkit. Each tool is additionally able to export its own configuration file used to tell the workflow package about its existence and mode of operation.

3 EXAMPLE

Workflow systems such as Galaxy offer great benefits for many fields of science. Using a system such as ballaxy, these benefits also apply to applications in structural-based drug discovery. A typical docking workflow using CADDSuite (Kohlbacher, 2012) tools is shown in Supplementary Figure S1, where it is described in detail.

Finally, addition of new tools using BALL is simple and is described in our documentation (http://ball-trac.bioinf.uni-sb.de/wiki/ballaxy).

ACKNOWLEDGEMENTS

The authors wish to thank the BALL development team. They gratefully acknowledge the use of software from OpenEye Scientific Software, Inc.

Funding: A.H. acknowledges financial support from the Intel Visual Computing Institute (IVCI) of Saarland University and the ‘Schwerpunkt Rechnergestützte Forschungsmethoden in den Naturwissenschaften’ of Johannes-Gutenberg University Mainz, A.H. and H.P.L. financial support from DFG (BIZ4:1-4). O.K. acknowledges financial support from DFG core facilities (grant KO 2313/6-1), the European Commissions Seventh Framework Programme (FP7/2007-2013) under grant agreement no. 283481 (SCI-BUS). J.K. and O.K. would like to thank the German Federal Ministry of Education and Research (BMBF) for the opportunity to do research in the MoSGrid project (reference 01IG09006).

Conflict of interest: none declared.

REFERENCES