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Developing Science Gateways for Drug Discovery in a Grid Environment

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ABSTRACT

Methods for in-silico screening of large databases of molecules increasingly complement and replace experimental techniques to discover novel compounds to combat diseases. As these techniques become more complex and computationally costly we are faced with an increasing problem to provide the research community of life-sciences with a convenient tool for high-throughput virtual screening (HTVS) on distributed computing resources. To this end, we recently integrated the biophysics-based drug screening program FlexScreen into a service applicable for large-scale parallel screening and reusable in the context of scientific workflows. Our implementation, based on Pipeline Pilot and Simple Object Access Protocol (SOAP) provides an easy-to-use graphical user interface to construct complex workflows which can be executed on distributed computing resources, thus accelerating the throughput by several orders of magnitude.

Keywords: Virtual Screening, FlexScreen, Science gateways, Drug Discovery, High Performance Computing

INTRODUCTION

Drug discovery can be drastically accelerated with the use of high-throughput virtual screening (HTVS) methods (Friesner et al., 2004; Halgren et al., 2004; Meng et al., 1992; Merlitz and Wenzel, 2002, 2004), an ongoing trend in medical research taking advantage of recent developments in algorithms and computer technology. In order to identify promising candidates for novel drugs, chemical compound databases with millions of ligands (Irwin and Shoichet, 2005) need to be screened using HTVS against structurally resolved receptors and thus distributing the workload on resources such as computing grids becomes essential. On the other hand, optimization of existing methods for HTVS to utilize novel high performance computer (HPC) architectures such as GPUs (Pérez-Sánchez and Wenzel, 2011; Sánchez-Linares et al., 2011b) can significantly reduce the run time per ligand.

Currently, HPC resources are mostly accessed remotely through low-level front-end machines (user interface machines) or using grid middleware and thus require from the non-expert end users in-depth knowledge of diverse batch systems or grid middleware protocols, respectively. To acquire the knowledge to use this complex low-level infrastructure for real-life applications makes the learning curve for scientists very steep. This is why efforts have still to be made to hide the complexity embedded in the Grid and to provide productive high-level services that allow scientists to take more effectively further advantage of the distributed resources.

Science gateways are the primary solutions dedicated to bridge such knowledge gaps. A science gateway is defined as a community-developed set of tools, applications, and data that is integrated via
a portal or a suite of applications, usually in a graphical user interface, that is further customized to meet the needs of a targeted community (Catlett, 2005, 2002). With science gateways non-grid-aware users can use grid infrastructure to run shared, well-tested applications customized for their own research field. Generally these solutions contain a set of research-specific applications developed by (and for) the community, and provide services integrated in a unified user interface, usually a web portal or a stand-alone graphical user interface. In the context of HTVS, this problem is paramount because the target user community consists of pharmacists and biologists not trained or experienced in the use of HPC/grid infrastructures.

Very often, science gateways provide special higher-level services for construction and execution of scientific workflows, i.e., means to automate processing of multiple steps in parallel or in a sequence, including branching and loops. Scientific workflows are abstract logical maps of complex simulation protocols and require that each step (often a different scientific application) provides common interfaces for execution and data exchange. Diverse mature science gateways or science gateway frameworks have evolved in different projects, which additionally allow for workflow management. For example, the UNICORE workflow engine and its workbench have been used in the area of Quantitative Structure-Activity Relationship (QSAR) and Quantitative Structure-Property Relationships (QSPR) models (Sild et al., 2005), and the Gridbus workflow for brain imaging (Pandey et al., 2009). Other very widely used workflow-enabled science gateways are Pipeline Pilot (http://www.accelrys.com), with different licensing options depending on the academic or industry version, Kepler (Ludäscher et al., 2006), Galaxy (Goecks et al., 2010; Blankenberg et al., 2010; Giardine et al., 2005), WS-PGRADE (Kacsuk et al., 2012), KNIME (Berthold et al., 2008) and Taverna (Wolstencroft et al., 2013) with open source licenses. For a review on scientific workflows we refer to Deelman et al. (2009).

To get an idea about the difficulties with the direct exploitation of HPC systems using HTVS methods we will describe how this process is usually carried out by expert users without use of science gateways. There are mainly three differentiated stages involved in the process:

1. Simulation data preparation: all the necessary data for the simulation must be conveniently prepared and the HPC system set up accordingly. In a classical parallel HPC system the total simulation is divided into different simulation units. Those units belong to thousands or more configuration files that must be arranged from a single file valid for the sequential execution of the program. This is not easy to do for end users, since it requires the use of different shell scripts for preparing those input files. Besides, specific configuration files for the queuing system must be set up for each independent simulation. Therefore, advanced knowledge of different IT technologies like tasks parallelization, input file structure, etc., is required at this stage.

2. Execution of the simulation: using different methods, the different simulation units are sent to the HPC system for their execution. The user needs to take care that there are no errors, to check continuously that the system is working properly and calculations are being performed seamlessly, and when the computations are finished, that there have been no errors.

3. Processing and interpretation of the results: it is usually necessary to move all the relevant data produced in the simulation to a local machine for its posterior analysis. Advanced knowledge of how HPC filesystems work is generally required at this stage. Lastly, all data needs to be curated and processed, normally using different advanced tools.

Given all the different and complex stages of the general simulation process, it is clear that for a user to be able to run calculations in this fashion, advanced knowledge of several tools, filesystems, etc., is required. Therefore, not all specialized users would be able to run computational experiments in this environment but only the advanced ones. This is why a work environment of another quality should be provided for the end user to exploit these resources effectively.

In order to make HTVS methods accessible for the relevant community, we identified the following goals. (i) The screening method has to be made accessible via an easy-to-use graphical interface; (ii) The HTVS application has to be integrated in such a way that it becomes reusable in different scientific workflows in combination with other applications; (iii) The screening method has to provide a seamless access to large-scale computing resources to enable large screening campaigns. In this work, we present a solution for the HTVS application FlexScreen which will take into account these three aspects. In general, our research belongs to the problem of subject adaptation of existing IT technologies, its customization...
for known domain of expertise of end users. This is an expansion of our previous work (Pérez-Sánchez et al., 2011). In the next section we will consider requirements for development of such type solutions. In Methods Section, we will introduce the FlexScreen application as well as the methods we employ to integrate FlexScreen into workflows for HTVS. In Implementation Section, we will particularly describe how we adopted Pipeline Pilot and the Simple Object Access Protocol (SOAP) to implement our concept and present a study case with use of the developed machinery. Furthermore, we present our investigation of integrating the implemented methods in diverse workflow-enabled science gateways. In Conclusions and future work Section, we will conclude and give an outline of future work.

PROBLEM STATEMENT

Our research belongs to the general problem of subject adaptation of existing IT technologies, its customization for known domain of expertise of end users. For the moment there are several approaches, intended for simplification of using HTVS methods (the list of corresponding software tools can be found at Jacob et al. (2012)).

The idea of our research is to find the most effective way of customization of HTVS methods. Analysis of existing approaches allows us to formulate next preconditions for the possible solution:

1. Simple and easy development of results by users, having no specific IT knowledge and qualification;
2. Rapid development, as e.g. by following RAD (Rapid Applications Development) technology;
3. Possibility of quick redevelopment of a solution without changing IT infrastructure;
4. Flexibility of solution, i.e. it should not be hardcoded inside a corresponding tool and so allowing development of extensions by advanced users;
5. The solution should allow to a user easy and naturally express logic (properties and behavior) of a domain;
6. Generality of solution, i.e. possibility to take into account specifics of modeling different domains (as biology, chemistry, physics, geometry);
7. Possibility of sharing and reusing existing solutions;
8. The solution should allow feature analysis (best of all, by attracting visual techniques);
9. On-fly testing and verification of basic properties (incl. syntax check) before executing;
10. The implementation environment should be easy and commonly used.

Having these requirements we have analyzed existing technologies and corresponding software tools. Simple development means that most of calculation issues (effectiveness, optimization, scheduling, resources etc.) are automated, and user can concentrate on the task in terms of his domain of expertise (e.g. domain specific data types). Details of the underlying programming code also should normally be hidden. At the same time solution should take into account different qualification of users, which can include biology scientists, persons with IT background and people from pharmacy companies. Thus solution should be flexible, and allow quick redevelopment to follow possible changes in HTVS methods.

To give to users the freedom in expression of logic we decided not to use point solutions - software tools having interface to HTVS, but introduce language, allowing to users to express logic of domain in the way they want to do it.

Here the problem of development of Domain Specific Languages (DSL) can be addressed. In general, the approaches for modeling domains can be divided into two parts: 1) using a so-called General Purpose Language (GPL) or 2) developing a DSL. Although existing GPLs are good for expressing computational domains, they are not suitable for modeling biological domains. At the same time, biological modeling approaches do not allow us to express data structures and computational processes.

Thus, we need to develop an approach, that allows us to express heterogeneous semantics of interlinked biological and computational domains. The main task here is to specify protocol of calculation, can be considered as workflow of tasks. In general, Workflow Management System (WMS) can be used here,
allowing to develop and manage different protocols as a sequence of tasks. Note, workflow approach is becoming more and more popular nowadays for modeling distributed IT environments, including grids and cloud computing. They allow to manage the execution of various distributed processes.

A scientific workflow system is a special type of a WMS, allowing to build protocols for some scientific application as simplified maps of complex simulation protocols. Development of scientific workflows for using HTVS methods will be considered as an effective solution for our research problem. We choose Pipeline Pilot for implementation due to its flexibility, allowing to develop workflows for different domains. Pipeline Pilot can be defined as scientific visual and dataflow programming language, allowing construction and execution of scientific workflows. At the same time the Pipeline Pilot is simple enough to be used by people having no specific IT knowledge and skills.

Note, that using visual languages (VL) allows the manipulation of graphical objects as mathematical complexes or data structures. Using VL, e.g. visual programming languages, belongs to RAD technology. VLs are also effectively used for data analysis in quite different domains.

As most of VLs, Pipeline Pilot uses idea of drawing boxes and connecting them by arrows (pipes). It allows development in an interactive way and checking syntax on the fly. As for protocol of calculation, Pipeline Pilot implemented the idea of dataflow programming, emphasizing the movement of data throw pipes. This approach allows users to automate parallelization.

Due to popularity, protocols developed in Pipeline Pilot enable scientists to publish scientific services making them available across scientific community. Moreover, Pipeline Pilot workflows language is a standard, which allows to encapsulate and deploy the best practices. So in general the proposed solution reduces not only development time, but also costs, spent by integrating with HTVS point solution software.

METHODS

FlexScreen
In this work, HTVS calculations have been performed with the all-atom receptor–ligand docking program FlexScreen (Merlitz and Wenzel, 2002; Kokh and Wenzel, 2008), which employs a force-field based scoring function (similar to Autodock (Morris et al., 1996)) and a Monte-Carlo based search algorithm based on the stochastic tunneling method (Wenzel and Hamacher, 1999). This method has the advantage that it suffers only a comparatively small loss of efficiency when an increasing number of degrees of freedom of the receptor is considered.

A physical model is implemented, which takes implicitly into account the influence of the solvent in the interaction between ligands and proteins. The free energy of the system includes a vacuum contribution that has been previously available in FlexScreen as well as additional solvation terms for the individual species and for the complex as a linear sum of atomic parameters (Eisenberg and McLachlan, 1986). This latter model has the advantage that it is faster than other methods presently used and has still proven to be reasonably accurate. The solvent accessible surface area of the molecules must be determined, which is a computationally intensive task. The other main advantage of the method is the determination of the weight parameters for different atom and bond types deriving from experimental partition coefficients in the cases of octanol–water and gas–water.

Pipeline Pilot
Pipeline Pilot provides for applications based on SOAP standard methods to communicate with each other over the HPC resources (Yang et al., 2010), allowing very effective workflow life-cycle management, i.e. it ensures maximum reuse of already integrated modules. In this way, in addition to its built-in functionality, the architecture of Pipeline Pilot has been organized for integration and extensibility and designed to interoperate with external software objects and applications. A number of mechanisms are available to automate the execution of a remote program. Additional options are available if the screening code resides on the workflow server.

Different mechanisms are used for remote execution ranging from simple Telnet and File Transfer Protocol (FTP) up to more elaborated standards such as SOAP (Snell et al., 2002) and web services. The SOAP standard provides methods for applications to communicate with each other over the HPC resources. The Pipeline Pilot supports SOAP with Web Services Description Language (WSDL) extensions for efficient decoupling of workflow management from the internal implementation of services. The SOAP framework is independent of any particular programming model, environment, or language. It is a structured method for sharing messages between server and client, and relies on the language XML.
to store and transmit the information and adds the necessary HTTP headers to the information. Most applications do not deal directly with the underlying SOAP data structures. Instead, they use a toolkit specific to their programming language and operating system. The toolkit simplifies the process of making SOAP calls and processing the returned results.

RESTful services gain popularity, which typically work more faster comparatively with SOAP implementations. At the same time, it is more difficult to broadcast RESTful services, which is a significant point in the context of development of scientific gateways. SOAP provides an interface for WSDL, allowing to define complex protocols, which is exactly the case of using Pipeline Pilot. So REST and SOAP have their own advantages and drawbacks and both are intensively used in modern web-based systems. The decision to choose the needed protocol can depend on problem domain only.

Pipeline Pilot provides several integration methods so that several applications existing either in the workflow server, remote server or cluster can be executed automatically in a workflow. Pipeline Pilot provides also data integration tools that assist in the assembly of information from different formats and pertaining to different databases. A convenient and intuitive graphical user interface via a web browser is provided for constructing and executing the workflows. The workflows are assembled using modules that are represented as icons in the graphical user interface. The workflows are stored in an XML format and can be easily exchanged between users. The modules, called components, include a variety of data readers, manipulators, calculators, data viewers, and data writers. For example, there are convenient data reading modules for ISIS files, SD-files, and SMILES, as well as delimited text and Excel spreadsheet files. Data viewers and writers include standard applications, such as WebLabViewerPro and Spotfire. An HTML molecular table viewer provides a convenient way to view tabular results with chemical structures. Although the applicability of the pipelining provided by this software is generic, the numerous (more than 200) specific components provided by SciTegic are heavily geared toward chemoinformatics environments. For academic users there is a free version of Pipeline Pilot available.

Workflows and Data Pipelining

A workflow in Pipeline Pilot refers to the way a protocol is defined, usually in form of several disconnected pipelines, each of which is made of components joined by pipes. A component refers to an individual operation to be performed on a set of data records. The order of execution depends on the order in which the components are joined since the protocols are executed from left to right and from top to bottom.

In the specific form of a workflow called data pipelining, records are passed individually down the pipes. Data pipelining allows the automation of the HTVS process and the integration of several related modeling and database packages. Thus, in addition to orchestration of multiple workflow steps, the data pipelining provides means for seamless data exchange between the individual application modules. The end users’ work in HTVS projects can be tremendously facilitated by the exploitation of already prepared sets of commonly used collections of tasks in the form of workflows. These protocols can be later deployed on HPC resources in a simple and automated fashion. An advantage of the pipelining approach is the ability to capture and conveniently share workflows for better reuse.

IMPLEMENTATION

Pipeline Pilot Modules for FlexScreen

FlexScreen was initially designed as a standalone command line application. Therefore, most previous users are familiar with this method of running it. The GUI that the gateway provides is meant for those users who are not familiar with running command line applications. Nevertheless, the gateway is meant for both kinds of users. From one side, users that were familiar with the command line version, will not find any difficulty when using the GUI, whereas new FlexScreen users or users that have never used any command line application will be quickly able to setup docking simulations.

In the first part of our work we implemented a set of Pipeline Pilot modules that were required to run FlexScreen within Pipeline Pilot. The required executables and template configuration files were placed in the Pipeline Pilot server. The FlexScreen integration in Pipeline Pilot is depicted in Fig.1. In pipelines 1 and 2 end users need to specify receptor and ligand database files in the standard molecular PDB format. If the user works with other molecular formats (smi, sdf, etc.), the protocol can be easily modified using molecular format converters included in the standard component collection of Pipeline Pilot. Afterwards, the initial receptor and ligand files can be parameterized depending on the charge model used, hydrogen model, etc. and additional components (pH, tautomers, etc.) can also be easily
included in the pipeline. Once the molecules are ready for the HTVS calculations, the docking parameters (degree of flexibility, simulation length, physical model, etc.) and parallel calculation parameters (batch size, number of processors to use, etc.) are also specified at the beginning of the third pipeline. In any case, the protocol also provides default parameters for all the components, so that the end user only needs to select ligand, receptor and binding site parameters to run FlexScreen calculations.

Figure 1. Integration of FlexScreen into Pipeline Pilot workflows. Pipelines 1 and 2 read and format the ligand database and receptor files. In Pipeline 3 the input molecules are received and the docking simulation parameters are specified. Then the FlexScreen component performs the SOAP calls and runs the calculations on the HPC resources. Finally the results are processed and presented in an interactive table format.

Data Analysis from Virtual Screening Calculations
One of the challenges in a virtual screening experiment is to analyze and organize the returned results. Again, an expert modeler is familiar with tools available within a modeling environment to examine and filter the results. But for a non-expert end user, the analysis and presentation must be automated so that they can correctly generate the information that is needed for further decision making. Using a single PC as a server, a single user is thus able to design and run application workflows that link all available Pipeline Pilot modules with FlexScreen for HTVS.

SOAP Implementation of FlexScreen
The integration in Pipeline Pilot alone, or in other words, the use of Pipeline Pilot on just a desktop machine is, however, insufficient for really large in-silico screening campaigns. The improved accuracy of FlexScreen comes at the price of the computation cost of the underlying biophysical model. Therefore, we have implemented the FlexScreen Pipeline Pilot modules as a SOAP-based (Snell et al., 2002) client-service pair capable to operate on distributed architectures such as computing grids and clouds. We have developed a SOAP-based web service for the remote FlexScreen application using software such
Algorithm 1 Pseudocode of the FlexScreen workflow using SOAP services with Pipeline Pilot

1: \textbf{for} \ i = 1 \ \textbf{to} \ \text{numberofbatches} \ \textbf{do} \\
2: \quad \text{SendLigandData} \\
3: \quad \text{SendReceptorData} \\
4: \quad \text{SendSimulationParameters} \\
5: \quad \textbf{end for} \\
6: \quad \textbf{for} \ i = 1 \ \textbf{to} \ \text{numberofbatches} \ \textbf{do} \\
7: \quad \quad \text{ExecuteSimulationUnits} \\
8: \quad \quad \textbf{end for} \\
9: \quad \text{ReceiveResultsData} \\
10: \quad \text{ReportResultsData} \\

as Apache/Tomcat (http://tomcat.apache.org) or the Perl SOAP::Lite module (http://soaplite.com). The SOAP server contains sufficient processing functionality to perform the following tasks (cf. Fig.2):

![Architecture of the implemented FlexScreen module](image)

**Figure 2. Architecture of the implemented FlexScreen module (cf. Fig.1).** This figure represents the case of use of distributed HPC resources via a SOAP client-server pair.

1. Receive a batch of ligands and receptor file as a SOAP message and save them to a file (steps 2 and 3 of Algorithm 1). One of the advantages of using SOAP is that it allows a batch size to be specified, allowing the collation of a series of individual docking requests in a single request for efficiency.
2. Receive complementary information as SOAP messages (step 4 of Algorithm 1) and save it to files, e.g., protein active site, configuration files related to simulation parameters, etc.
3. Generate and submit jobs to execute FlexScreen on HPC resources using the files previously created (step 7 of Algorithm 1).
4. Read the resulting files (step 9 of Algorithm 1) and pass them back as a SOAP message to the calling component. A report on the results will be automatically prepared (step 10 of Algorithm 1) as an interactive HTML report, a PDF document, or a spreadsheet.
Examples of Use

Results from a HTVS calculation performed by an end user are shown in Figs.3 and 4. As it can be seen in Fig.4, the resulting data is clearly organized in tables which are directly opened in the web browser after the screening calculations. The user can control the degree of detail in the final report interacting with the “table parameters” component as well as reorganize easily and sort the final data with a few mouse clicks in the web browser. There is also the possibility of exporting the results to other standard formats, i.e., PDF, Word, Excel spreadsheets, CSV text files, etc. The end user can also obtain detailed information about the 3D structure of the docked receptor–ligand conformations as can be seen in Fig.3, very useful for compound optimization, posterior screenings, etc.

Figure 3. 3D representation of the HTVS results obtained for two different receptor-ligand pairs. Blue color denotes the experimental ligand binding mode, orange color the FlexScreen prediction without considering solvation, and the red color the prediction with the consideration of solvation.

From the perspective of users’ experience, we found that the access to well-developed and validated workflows using FlexScreen encourages the user to test and explore new ideas. Informal discussions with users who have performed HTVS calculations with FlexScreen in this way confirms that the deployment of HTVS methods does not just get the same answers faster, but that scientists end up asking many more “what-if” questions and running many more experiments than they would have done when a modeler had to be involved in each case.

Integration of the FlexScreen Services in Further Science Gateways

In the last four years a few new science gateways have been developed or existing ones have been extended to support the HTVS user community, e.g. MoSGrid (Krüger et al., 2014) developed on top of WS-PGRADE or KNIME. To allow the reusability of services in the users’ preferred virtual environment, we investigated the possibilities to integrate the FlexScreen services in the context of further science gateways.

Since the FlexScreen services are SOAP based, a crucial prerequisite is the support of such services in the science gateway. Furthermore, the science gateway needs to be workflow enabled for the different tasks accomplished by each of the services to provide the whole pipeline of analysis steps. Since users may have established preparation and post processing steps for the HTVS pipeline, another prerequisite for considering a science gateway is the possibility to configure the execution environment of tasks in a workflow or pipeline independent from each other. In our investigation we considered four workflow-enabled science gateways widely used in the biomedical community.

WS-PGRADE (Kacsuk et al., 2012) is the flexible web user interface of the workflow system gUSE, which supports the management of DAG-based workflows. The control structure is defined by data dependencies and parameter sweep mechanisms allow for emulating loops over a defined range of parameters and data. Each task in a workflow is represented by a job with input and output datasets and each job can be configured for exploiting a resource independent of the configuration of dependent jobs. Thus, a job can be configured as SOAP web service and connected with jobs defined for applying local, cluster, grid and cloud resources or another SOAP web service. Thus, users can reuse the FlexScreen services in an intuitive way.
### Docking Results of Streptavidin

| NAME     | MOLECULE | AFFINITY | RMSD  |
|----------|----------|----------|-------|
| jnk vs_1-81_1 | ![Image](image1.png) | -102.113727 | 1.803142 |
| jnk vs_1-82_1 | ![Image](image2.png) | -84.001330 | 1.882979 |
| jnk vs_1-83_1 | ![Image](image3.png) | -34.849769 | 1.812968 |
| jnk vs_1-84_1 | ![Image](image4.png) | -45.391462 | 2.092398 |

**Figure 4. Sample of the output results in HTML format directly from the web browser.** HTVS results are presented in consecutive rows for the different ligands of the database. Different columns contain information about each ligand regarding name, energy calculations, RMSD, etc. Clicking on each ligand 2D representation opens a new window with detailed information about the 3D ligand binding mode as shown in Fig. 3.

The concept behind Galaxy (Goecks et al., 2010; Blankenberg et al., 2010; Giardine et al., 2005) differs from WS-PGRADE but it also offers an intuitive web user interface with workflow management capabilities for DAG-based workflows. It is designed as a tool box for intuitively creating and invoking workflows with pre-configured tools in local, cluster and cloud environments. The administrator of a Galaxy instance can configure SOAP web services, which are then available to the users (Wang et al., 2009). Hence, users are able to integrate the FlexScreen services in their workflows.

Taverna (Wolstencroft et al., 2013) follows a different approach on the client side compared to WS-PGRADE and Galaxy and the workbench needs to be installed by the users. Despite this drawback on the users’ side, it is widely adopted in the community. It supports besides DAG-based workflows also loops as workflow constructs and is especially based on configuring each step in a workflow as SOAP-based service. It is an ideal candidate for reusing the FlexScreen services.

While KNIME (Berthold et al., 2008) is also an easy-to-use workbench, which has to be installed by the users, it supports command line tools and SOAP-based web services via its Generic Webservice Client (https://tech.knime.org/webservice-client). KNIME is especially user-friendly, has rich workflow management features and offers pre-configured packages. A user can easily integrate the FlexScreen services into the workbench.

These four examples prove that the FlexScreen services are not only applicable in the native Pipeline Pilot environment but also in other science gateways and, thus, reusable for a large user community employing diverse science gateways for their research topics. The services can be connected with other tools and services to improve the user experience on accomplishing their research in one user interface.

### Conclusions and Future Work

In this paper, we have considered the general problem of subject adaptation of existing IT technologies, its customization for known domain of expertise of end users. We have described the implementation of a HTVS methodology in a science gateway environment making use of the workflow environment provided
by Pipeline Pilot. The solution basing on SOAP and web services enables the exploitation of distributed HPC resources using a grid computing strategy.

From our point of view, the main drawback of Pipeline Pilot is that a yearly paid license is required. Therefore, not all research institutions would be able to cover these costs. It seems that open source alternatives to Pipeline Pilot exist, such as UNICORE, Kepler and Taverna, but we are not sure yet whether they offer the same or similar alternative. Thus, we will explore them in further studies.

Currently, we are also developing improved GPU-based versions of FlexScreen (Guerrero et al., 2011; Sánchez-Linares et al., 2011b,a) and planning its deployment on grid resources.

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