SOAP-based services provided by the European Bioinformatics Institute

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ABSTRACT

SOAP (Simple Object Access Protocol) (http://www.w3.org/TR/soap) based Web Services technology (http://www.w3.org/ws) has gained much attention as an open standard enabling interoperability among applications across heterogeneous architectures and different networks. The European Bioinformatics Institute (EBI) is using this technology to provide robust data retrieval and data analysis mechanisms to the scientific community and to enhance utilization of the biological resources it already provides [N. Harte, V. Silventoinen, E. Quevillon, S. Robinson, K. Kallio, X. Fustero, P. Patel, P. Jokinen and R. Lopez (2004) Nucleic Acids Res., 32, 3–9]. These services are available free to all users from http://www.ebi.ac.uk/Tools/webservices.

INTRODUCTION

Today, biological databases are large collections of data that are relatively difficult to maintain outside the centres and institutions that produce them. These data are traditionally accessed using browser-based World Wide Web interfaces. When large amounts of data need to be retrieved and analysed, this often proves to be tedious and impractical. Web Services technology enables scientists to access these data and analysis applications as if they were installed on their laboratory computers. Similarly, it enables programmers to build complex applications without the need to install and maintain the databases and analysis tools (1) and without having to take on the financial overheads that accompany these. Moreover, Web Services provide easier integration and interoperability between bioinformatics applications and the data they require.

THE TECHNOLOGIES

The European Bioinformatics Institute (EBI) has tried and tested standards such as CORBA (http://www.corba.org) and Web Services. CORBA is standardized and mature; it uses the Inter-ORB Protocol (IIOP) and can be tunneled through HTTP but does not natively support HTTP. It is trickier to communicate through firewalls. Web Services uses SOAP (the Simple Object Access Protocol) over HTTP. It interacts with other systems using messages based on eXtensible Markup Language (XML) (http://www.w3.org/XML). A SOAP message can be transferred using almost any application or transport protocol. SOAP uses the Web Services Description Language (WSDL) (http://www.w3.org/TR/wsdl) to describe its interface. A SOAP client can read the WSDL at runtime and dynamically select the proper data-encoding scheme and network transfer protocol. SOAP implementations are available for many programming languages, including Perl and Java, which are popular languages among bioinformaticians.

On the basis of these observations, the EBI has chosen to use the Web Services technology to expose its services in a programmatically accessible manner. All that is required by the bioinformatics programmer is a lightweight program that communicates with existing services running at the EBI. These services have several advantages. As traditional web browsers cannot be used programmatically, these services provide an easy and flexible way to deal with repetitive tasks such as bulk submission with minimal intervention from the user. Web Services clients allow the programmer as well as the service provider to integrate and build more complex analysis workflows using existing EBI services. Also, using these services effectively avoids the need to maintain many programs and databases locally.

SERVICES

In this article, we describe services currently available at the EBI via a SOAP server. These include tools for sequence and literature data retrieval, sequence similarity search services, protein function analysis and structural analysis tools that access the Macromolecular Structure Database (MSD) (2) and a set of Web Services called Soaplab (3) for the European...
InterProScan, respectively. These services are implemented
through web services such as Fasta (13), WUBlast (14) and the protein function
analysis tool InterProScan (15). These are Web Services that provide the same
functionality as the traditional analysis tool InterProScan (15). These Web Services
are available from http://www.ebi.ac.uk/Tools/webservices/WSFasta.html,
http://www.ebi.ac.uk/Tools/webservices/WSUWUBlast.html, and
http://www.ebi.ac.uk/Tools/webservices/WSInterProScan.html.

The EBI provides Web Services for sequence similarity tools
such as Fasta (13), WUBlast (14), and the protein function
analysis tool InterProScan (15). These are Web Services
that provide the same functionality as the traditional
browser-based services found at http://www.ebi.ac.uk/fasta,
http://www.ebi.ac.uk/blast2 and http://www.ebi.ac.uk/
InterProScan, respectively. These services are implemented
on a Perl-based, SOAP::Lite (http://www.soaplite.com) server.
The methods provided for using these services are listed
in Table 3. Fully functional Java and Perl client programs
are available from http://www.ebi.ac.uk/Tools/webservices/
download.html. A sample client in Perl is shown in Figure 3,
and Figure 4 illustrates how to run the client. Depending on
the input and the databases chosen for the search, jobs may
take seconds to complete or up to a few hours. Two modes of
job submission exist: synchronous and asynchronous.

**Synchronous mode.** This mode is equivalent to a user running
a command on a console or terminal and waiting for it to
complete. This requires the client to be constantly connected
to the server. This mode is suitable for database searches that
can be executed in up to 5 min (e.g., protein versus protein
searches).

**Asynchronous mode.** In this mode, the user submits a job and
receives a job identifier in return. This is the same as running
a UNIX command in the background and obtaining a job id.
The user can use the ‘jobs’ command to list processes that are
running in the background. Similarly, the user can query or
poll the status of an asynchronous mode job and receive the
following four states in response: JOB RUNNING (i.e., the
job is currently being processed), JOB PENDING (i.e., the job
is in a queue waiting processing), JOB NOT FOUND (i.e., the job
id is no longer available); job results are deleted after 24 h) and
JOB FAILED (i.e., the job failed or no results where found).

Typically, the asynchronous submission mode is recommended
when users are submitting batch jobs (e.g., many protein
sequences to analyse using InterProScan) or large
database searches (e.g., searching the whole of the EMBL
nucleotide sequence database). One advantage of this mode is
that it is impervious to system or network failure. The results of jobs are stored at the EBI for 24 h after the job has completed.

Structural Analysis
The EBI provides a Web Services interface to tools that access the MSD. This service enables software developers to query the MSD directly from their own application programs and is further described at http://www.ebi.ac.uk/msd-srv/docs/api. The available functions are described in the corresponding WSDL description at http://www.ebi.ac.uk/msd-srv/docs/api/msd_soap_service.wsdl. As well as simple extraction of data from the database, the interface also provides methods for performing complex queries on the MSD relational database remotely.

Figure 1. A sample Perl client calling the fetchData method.

```perl
#!/usr/bin/perl
use SOAP::Lite;
my $uri = 'urn:Dbfetch';
my $proxy = 'http://www.ebi.ac.uk/ws/services/Dbfetch';
my $soap = new SOAP::Lite(uri => $uri, proxy => $proxy);
my $result = $soap->fetchData("uniprot:wap_rat", "fasta", "raw");
print $result;
```

Figure 2. A sample client invocation showing the method called and result obtained.

```
% java DbfetchClient fetchData uniprot:wap_rat fasta raw
>uniprot|P01174|WAP_RAT Whey acidic protein precursor (Whey phosphoprotein) (WAP).
MRCSISLVLGLLALAEVALARNLQEHVFNSQSMCDSDSS4DTBCICQTNVECA
QNDMCCPSCSSCR6CTPVNEVQAGRCPWNPIQM1AAGPCPKDNPSCIDSDCSG
TMKCCENICSCMDPPEKSPTVISFQ
```

Figure 3. A sample Perl client for WSFasta calling the doFasta method asynchronously.

```perl
#!/usr/bin/perl
use SOAP::Lite;
my $WSDL = 'http://www.ebi.ac.uk/ws/WSFasta.wsdl';
my $soap = SOAP::Lite->service($WSDL);
my %params = (program => 'fasta3',
               database=>'uniprot',
               searchtype=>1, # indicates asynchronous mode
);
open INPUT, "input.txt"
my $content = <INPUT>;
close INPUT;
my $result = $soap->doFasta(SOAP::Data->name('params')
                         ->type(map=>%params),
                         SOAP::Data->name(content=>$content)
                         ->type("base64"));
print $result;
```

Figure 4. A sample client invocation. Note that ‘\’ means a continuous one-line command. The input file (e.g. mysequence) is a Fasta-formatted sequence.
For protein structure analysis, MSDfold, a protein secondary structure-matching tool, is available as a Web Service. An example client is described at http://www.ebi.ac.uk/msd-srv/docs/api/examples.html.

Soaplab

Soaplab (http://www.ebi.ac.uk/soaplab) is a tool that can automatically generate and deploy Web Services on top of existing command-line analysis programs. It is especially well suited for EMBoss-type applications. It allows the integration of many applications within a single programming interface. It can also interoperate with other Web Services described earlier (e.g. WSInterProScan) and it can create Web Services on top of existing web resources (e.g. extracting data from a third-party web page and providing its data as a Web Service).

Soaplab in its basic form is a tool for non-programmers who need only to create metadata describing resources (command-line applications, web pages) and let Soaplab generate the rest. The resulting Web Services are uniform and provide a good platform for integration into a workflow such as in Taverna (http://taverna.sourceforge.net). The initial metadata are available from the Soaplab Web Services interface. They make the services self-describing. Soaplab is also a reference implementation of the OMG (Object Management Group, http://www.omg.org/) standard for the Life Sciences Analysis Engine (LSAE).

CONCLUSION

We present here a set of applications that give the user more direct access to data and services from the EBI. From the user’s perspective, these are equivalent to installing and maintaining software and databases on local computers. From the programmer’s point of view, Web Services provide a robust and flexible environment in which to build applications and provide complex and novel services.

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Conflict of interest statement. None declared.

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