Hyperlink Management System and ID Converter System: enabling maintenance-free hyperlinks among major biological databases

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ABSTRACT

Hyperlink Management System (HMS) is a system for automatically updating and maintaining hyperlinks among major public databases in the field of life science. We daily create corresponding tables of data IDs of major databases for human genes and proteins, and provide a CGI-program that returns correct and up-to-date URLs for showing data of various databases that correspond to user-specified IDs. The HMS can deal with various IDs: accession numbers of International Nucleotide Sequence Databases, HUGO Gene Symbols and IDs of UniProt, PDB, H-InvDB and others, and it can return URLs of various databases: H-InvDB, HUGO Gene Nomenclature Committee Database, NCBI Entrez Gene, UniProt, PDB and others. For example, 23,297 pages of Locus view of H-InvDB are reachable by using HUGO Gene Symbols through the HMS. Not only the CGI-program, the HMS provides a Web page for finding and opening URLs of these databases. Although hyperlinking is an effective way of relating biological data among different databases, setting hyperlinks among them is a laborious work. The HMS fully automates the job, enabling maintenance-free hyperlinks. We also developed the ID Converter System (ICS) for simply converting data IDs by using corresponding tables in the HMS. The HMS and ICS are freely available at http://biodb.jp/.

INTRODUCTION

There are currently 1170 biological databases in the world (1) that are being utilized for various researches and developments. It is now possible for biologists to try to make discoveries by combining and linking data of different types obtained from multiple databases. For example, by combining protein sequences and protein structure data that are stored separately in major databases, biologists can study the basic rules of protein 3D structure such as how amino-acid sequences organize themselves into specific and stable 3D structures. In this way, linking specific data stored in multiple databases will lead to the organization of biological knowledge and will provide useful resources for novel researches.

Finding corresponding data in different databases and setting hyperlinks among them is an effective way of integrating information that are stored in independently operated databases in different organizations. The maintenance of the hyperlinks among many databases, however, requires a huge cost, because the number of pairs of databases is generally very large. For example, possible ways of hyperlinks among \( N \) related databases will be \( \binom{N}{1}^N \), if we consider the directions of hyperlinks. Suppose if one of the \( N \) databases is updated, \( 2(N-1) \) ways of hyperlinks need to be updated. If each of \( N \) databases is periodically updated in every 3 months, at most \( 8(N-1)N \) ways of hyperlinks need to be updated in a year, which will be a laborious work. We thus need a novel system to automate the maintenance of hyperlinks.

Here, we developed a Web server for automatically managing and updating hyperlinks that can connect...
molecular information in multiple databases about human genes and proteins. Furthermore, we developed a Web server for converting data IDs from one type to another. These Web servers will be useful for both database developers and biologists. The reason for it is that, for database developers, these systems can realize automated, work-saving and efficient management of databases, and can reduce the large cost of maintaining biological databases. Also, for biologists, especially those who are unfamiliar with the data IDs of major biological databases, our newly developed Web servers will be of help in carrying out efficient analyses of ‘big data’ that are being produced massively in laboratories (2). We hope that many database managers and researchers will find our Web servers useful.

WEB SERVER FEATURES

Usage of the Hyperlink Management System (HMS) CGI-program

HMS is a tool for automatically updating and maintaining hyperlinks between databases in the field of life science. The HMS automatically downloads the lists of data IDs for human genes and proteins and their relations from major biological databases everyday, and produces large corresponding tables of all data IDs of these databases (Tables 1 and 2). Then, using the corresponding tables of data IDs, the HMS shows the most up-to-date URLs for data in a database that are corresponding to a user-specified ID of another database. The URL is easily obtained by running a CGI-program of the HMS.

Suppose that the manager of database A wishes to create hyperlinks from database A to database B. Usually in this case, the manager uses data IDs of database B in the URLs of hyperlinks. However, if the manager uses the HMS, he can use data IDs of database A to create hyperlinks to database B. This is very useful because we can easily create hyperlinks without knowing data IDs of target databases. We can therefore drastically reduce the cost of creating and maintaining hyperlinks among databases.

Currently, the HMS collects 17 types of data IDs for human genes and proteins (Table 2). Among the 17 types of data IDs, we defined accession numbers of International Nucleotide Sequence Database (INSD; DDBJ/EMBL/GenBank), HUGO Gene Symbols and UniProt IDs as ‘common IDs’ that are used as key IDs in creating corresponding tables. We collect corresponding tables between a common ID and an other ID from each of source databases (Figure 1). We also obtain a corresponding table between accession numbers and HUGO Gene Symbols from NCBI and a corresponding table between accession numbers and UniProt IDs from UniProt. Using these tables, we create a large corresponding table that shows the relationships among 17 types of data IDs. The common IDs define the route of ID conversion (as shown in Figure 1). For example, when converting HIX of H-InvDB into PDB IDs, we first convert HIX into UniProt IDs and then convert it into PDB IDs.

The HMS directly opens a Web page of the target database if there is only one data ID corresponding to the source ID. However, in converting data IDs, an ID does not always correspond to one ID of another type. If there

| Database name | Viewer name | Format |
|---------------|-------------|--------|
| H-InvDB (3,4) | Transcript view LOCUSVIEW | TRANSCRIPTVIEW |
| H-Inv cluster ID (HIX) | HIX_ID | HIT_ID |
| H-Inv protein ID (HIP) | HIP_ID | 219765 |
| INSD Accession Number | ACC_ID | 4159 |
| NCBI OMIM ID | OMIM_ID | 133629 |
| RefSeq ID | Ref_SEQ | 222136 |
| Entrez Gene ID | GENE_ID | 48671 |
| GeMDBG dbSNP rs# | JSNP_ID | 40235 |
| HUGO Gene Symbol | HUGO SYMBOL | 76024 |
| H-GOLD Marker Name | MS_ID | 26421 |
| PDB PDB ID | PDB | 494 |
| UniProt UniProt ID | UNIPROT | 50958 |
| Ensembl Ensembl Transcript ID | ENST | 84422 |
| Ensembl Gene ID | ENSG | 63280 |
| FRNAdb FRNAdb ID | FR_ID | 37435 |
| HGPD cDNA clone ID | PDB | 41501 |
| FLJ cDNA clones | FLJ | 21340 |

*Parameters of database names used in the HMS CGI-program, ICS and Web service.

Table 2. Data IDs used in the HMS

| Source database | Data ID | Format | Count |
|-----------------|---------|--------|-------|
| H-InvDB | H-Inv transcript ID (HIT) | HIT_ID | 219765 |
| H-Inv cluster ID (HIX) | HIX_ID | 4159 |
| H-Inv protein ID (HIP) | HIP_ID | 133629 |
| INSD | Accession Number | ACC_ID | 222136 |
| NCBI OMIM ID | OMIM_ID | 16253 |
| RefSeq ID | Ref_SEQ | 48671 |
| Entrez Gene ID | GENE_ID | 40235 |
| GeMDBG dbSNP rs# | JSNP_ID | 76024 |
| HUGO Gene Symbol | HUGO SYMBOL | 26421 |
| H-GOLD Marker Name | MS_ID | 494 |
| PDB PDB ID | PDB | 50958 |
| UniProt UniProt ID | UNIPROT | 84422 |
| Ensembl Ensembl Transcript ID | ENST | 63280 |
| Ensembl Gene ID | ENSG | 37435 |
| FRNAdb FRNAdb ID | FR_ID | 41501 |
| HGPD cDNA clone ID | PDB | 21340 |
| FLJ cDNA clones | FLJ | 21340 |

*Parameters of ID types used in the HMS CGI-program, ICS and Web service.

*bNumber of data as of 23 January 2009.
are more than one IDs corresponding to the source ID, the HMS shows the list of all corresponding data IDs with appropriate hyperlinks, from which users can choose and open the data of interest.

In order to create hyperlinks using the HMS CGI-program, use the following URL as a target URL.  

http://biodb.jp/hfs.cgi?id=[ID]&type=[ID Type]&db=[Database Name]

Here, [ID] is a data ID of the database from which hyperlink is set, [ID Type] is a kind of the data ID and [Database Name] is the name of database to which hyperlink is set. Details of parameter settings are shown in Tables 1 and 2 and also in the online HELP pages of the HMS. For example, using BC053 657, an accession number of INSD, you can create a hyperlink toward the corresponding ‘Transcript View’ of H-InvDB in the following way.

  &lt;a href = "http://biodb.jp/hfs.cgi?id=BC053657&type=ACC_ID&db=TRANSCRIPTVIEW"&gt;BC053 657&lt;/a&gt;

This CGI-program is freely available to all users in the world. For database managers, if their databases have any one of those data IDs shown in Table 2, it is possible to create hyperlinks from the database to all the databases shown in Table 1 through the HMS. Because the hyperlinks of the HMS are automatically updated everyday, the connection to the latest data is always guaranteed.

Usage of the HMS Web Server

The HMS CGI-program is supposed to be used for hyperlinking among databases. However, there are situations that we want to query with a small number of data IDs. We thus developed a Web server of the same function as the CGI-program (Figure 2). Here, by utilizing the same corresponding tables of data IDs as those of the CGI version, users can open Web pages of a database corresponding to a specified data ID of another database. The usage of the HMS Web server is quite simple and requires no complicated explanations. For example, if you enter an accession number of INSD in the ‘Source ID’ and push the ‘Search’ button, you can open the ‘Locus view’ of H-InvDB. In the textbox, users can enter multiple data IDs delimited by a space, tab or comma. In this case, the result page shows a list of multiple URLs with hyperlinks corresponding to each data.

Usage of the ID Converter System (ICS)

ICS is a tool for converting data IDs used in a database into other, corresponding IDs used in other databases (Figure 3). All the data IDs used in the HMS can be dealt with. Multiple IDs can be converted at a time, if users enter multiple data IDs delimited by a space, tab or comma. When a data ID can be converted into multiple corresponding IDs, the ICS shows all the corresponding IDs. Users can also specify a file on their client PC, and convert a list of data IDs in it. Furthermore, the ICS is accessible by computer programs using the Web service, as will be described later.

Web service and other functions

For those who wish to analyze massive data by computers, the HMS and ICS accept queries through the Web service, which makes it possible to call functions of these systems through computer programs. The details of commands for the Web service will be shown if you click the ‘Web service’ button on the upper right corner of the Web pages (Figures 2 and 3). On these pages, there are some sample programs for the Web service. Furthermore, we prepared statistics pages of the HSM and ICS. If you click the ‘Update Info’ button on the upper right corner of the top pages (Figures 2 and 3), the number of data IDs downloaded from each database everyday will be shown. If you click the ‘Release Info’ button, the number of pages of each database that are hyperlinked through various data IDs will be shown. Finally, there are help pages that explain how to use the HMS and ICS.

DISCUSSION

As has been stated earlier, there are currently 1170 databases in the field of life sciences that are listed in the online Molecular Biology Database Collection of Nucleic Acids...
Research (1). The number of databases is increasing every year. Because biologists usually wish to combine a certain type of data with another, integration of these various and huge databases will be of great significance. In principle, there are two possible ways of integrating databases. One is to collect all the databases at one place and merge them, while the other is to combine the databases virtually by hyperlinking corresponding data among them. The former method requires an enormous amount of disk space of computers and a large number of people to maintain and update the databases. In other words, the cost of the former method is too large to implement. In fact, there was an attempt to construct a huge integrated database of many biological data by the former method, which failed in a few years probably due to the high running cost. The latter method, on the other hand, can be realized relatively easily if we adopt the automated hyperlinking that we introduced in this article. The HMS will be a fundamental platform to virtually integrate databases that are operated independently by different organizations. Furthermore, the HMS will make it possible for us to construct integrated databases in a sustainable way.

The ICS is also a quite useful tool. Because major databases such as INSD and UniProt update their 'live lists' of...
data IDs on a daily basis, the role of ICS to provide users with the up-to-date information of ID conversion is very important. The usage of the ICS is very simple, and anyone can make full use of it easily. For example, even when one has a list of HUGO gene symbols for 100 genes, he/she can easily find corresponding data in H-InvDB, UniProt, PDB and other databases through the ICS. Advantages of the ICS are not limited to this. The most important feature of the ICS is that ID conversion can be done between two types of data IDs that are not directly related. The ICS can convert from and to all data IDs shown in Figure 1, including those that are not directly connected. For example, the ICS can directly convert accession numbers of ISND into PDB IDs. We thus think that the ICS will be a useful Web server for many researchers.

We plan to add more and more databases into the HMS and ICS in the future. It is easy to add new databases as destination of HMS hyperlinks if the database uses either accession numbers of INSD, UniProt IDs or PDB IDs. We welcome any proposals about incorporating additional databases into the HMS. Furthermore, we have been dealing with databases about human molecular data, but we plan to extend the HMS to molecular data of other species in near future. The ICS will be also extended to incorporate other IDs that are frequently used by biologists. For example, we will incorporate IDs of literature databases, which will enable combining biological knowledge in literature with molecular data. We will extend the ICS according to the requests from users.

The HMS and ICS will be of considerable use as an automated tool for database managers and as a new

Figure 3. Usage of the ID Converter System on the Web. Input source IDs in the textbox, and select the type of IDs. You may also specify a text-file with a list of source IDs. Then, select the type of data ID to convert, and push the ‘Search’ button. Then, the system searches for IDs that correspond to source IDs, and shows the list of them. The list can be downloaded by pushing the ‘Download’ button. This figure shows an example of converting two accession numbers, BC053657 and Z34290, into HUGO Gene Symbols.
indispensable tool for biologists. We may be able to extend these systems to incorporate keyword searches that are frequently used in many databases. By indexing general and technical terms by assigning them some IDs, the HMS and ICS will be able to deal with keyword searches. Such an extension will further increase the value of these systems.

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