YMDB: the Yeast Metabolome Database

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ABSTRACT
The Yeast Metabolome Database (YMDB, http://www.ymdb.ca) is a richly annotated ‘metabolomic’ database containing detailed information about the metabolome of Saccharomyces cerevisiae. Modeled closely after the Human Metabolome Database, the YMDB contains >2000 metabolites with links to 995 different genes/proteins, including enzymes and transporters. The information in YMDB has been gathered from hundreds of books, journal articles and electronic databases. In addition to its comprehensive literature-derived data, the YMDB also contains an extensive collection of experimental intracellular and extracellular metabolite concentration data compiled from detailed Mass Spectrometry (MS) and Nuclear Magnetic Resonance (NMR) metabolomic analyses performed in our lab. This is further supplemented with thousands of NMR and MS spectra collected on pure, reference yeast metabolites. Each metabolite entry in the YMDB contains an average of 80 separate data fields including comprehensive compound description, names and synonyms, structural information, physico-chemical data, reference NMR and MS spectra, intracellular/extracellular concentrations, growth conditions and substrates, pathway information, enzyme data, gene/protein sequence data, as well as numerous hyperlinks to images, references and other public databases. Extensive searching, relational querying and data browsing tools are also provided that support text, chemical structure, spectral, molecular weight and gene/protein sequence queries. Because of S. cerevisiae’s importance as a model organism for biologists and as a biofactory for industry, we believe this kind of database could have considerable appeal not only to metabolomics researchers, but also to yeast biologists, systems biologists, the industrial fermentation industry, as well as the beer, wine and spirit industry.

INTRODUCTION
Metabolomics is a field of ‘omics’ research that is primarily focused on the identification and characterization of small molecule metabolites in cells, organs and organisms (1). Together with genomics, transcriptomics and proteomics these four ‘omics’ disciplines form the cornerstones to systems biology. However, relative to its more mature ‘omics’ cousins, metabolomics still lags far behind in developing or formalizing its software and database infrastructure (2). This is because the needs of metabolomics researchers span a very diverse range of scientific disciplines including organic chemistry, analytical chemistry, biochemistry, molecular biology and systems biology. In other words, metabolomics requires a tight blending of the tools found in both bioinformatics and cheminformatics. To address these informatics challenges, we (and others) have been steadily developing a set of comprehensive and open access tools to lay a more solid software/database foundation for metabolomics (2–4). In particular, our group has developed several widely used organism- or discipline-specific databases including the Human Metabolome Database (HMDB) (5), DrugBank (6), the CyberCell database (CCDB) (7), the Toxin/Toxin-Target database (T3DB) (8) and the Small Molecule Pathway Database (SMPDB) (9). HMDB, T3DB, DrugBank and SMPDB were specifically developed to address the metabolomics, toxicology, pharmacology and systems biology associated with humans (i.e. Homo sapiens), whereas CCDB was specifically developed to address the metabolomics and systems biology needs for Escherichia coli.

We believe that the establishment and maintenance of organism-specific metabolomics databases is absolutely

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critical to the field of metabolomics as each organism has a unique and chemically distinct metabolome. The ‘naïve’ identification of metabolites, by simple mass matching for instance, without regard to their origin (organism or man-made) frequently leads to spurious, humoros or meaningless compound identifications (10). Therefore, as part of our ongoing effort to create species-specific metabolomic resources for other model organisms we have now turned our attention to yeast, or more specifically, Saccharomyces cerevisiae.

The metabolic byproducts of S. cerevisiae fermentation are particularly interesting from both a biochemical and an industrial point of view. Indeed, S. cerevisiae (and its various strains) is perhaps the world’s most important microbial biofactory, playing a key role in industrial chemical or biofuel production (ethanol), in the baking industry, as well as in beer, wine and spirit production. Together, these yeast-based industries are worth more than one trillion dollars per year to the global economy (11). As a model organism for molecular biologists, S. cerevisiae is certainly the most intensively studied microbe and perhaps the most well understood living thing on earth. Being one of the first organisms to be fully sequenced (12) and being particularly amenable to unique and powerful genetic manipulations (13,14) the sequence, function and interacting partner(s) of every gene/protein in S. cerevisiae is now almost completely known. This knowledge is contained in a number of excellent yeast-specific resources including SGD (15), YPD (16), CYGD (17) and FunSpec (18). This remarkably detailed molecular knowledge has also made S. cerevisiae a favorite model organism for systems biologists, leading to the development of some very useful resources aimed at modeling or describing yeast pathways and metabolism including YeastNet (19), MetaCyc (20), KEGG (21) and Reactome (22). Each of these excellent databases contains valuable information on primary yeast metabolic reactions, pathways and primary yeast metabolites.

Unfortunately, none of these systems biology databases contains information on the secondary metabolites of yeast fermentation (those compounds that give wine, beer and certain cheeses or breads their flavor or aroma), yeast-specific lipids, yeast volatiles or yeast-specific ions. These actually represent hundreds of industrially and biochemically important compounds. Furthermore, none of today’s current set of yeast systems biology databases provides detailed metabolite descriptions, intra- or extracellular concentrations, growth conditions, physicochemical properties, subcellular locations, reference Nuclear Magnetic Resonance (NMR) or Mass Spectrometry (MS) spectra or other parameters that might typically be needed by researchers interested in yeast metabolism or yeast fermentation. For metabolomics researchers, as well as industrial chemists working with yeast byproducts, these kinds of data need to be readily available, experimentally validated, fully referenced, easily searched and readily interpreted. Furthermore, they need to cover as much of the yeast metabolome as possible. In an effort to address these shortcomings with existing yeast systems biology databases and to create a database specifically targeting the needs of yeast metabolomics, we have developed the Yeast Metabolome Database (YMDB).

DATABASE DESCRIPTION

The YMDB is a combined bioinformatics–cheminformatics database with a strong focus on quantitative, analytic or molecular-scale information about yeast metabolites and their associated properties, pathways, functions, sources, enzymes or transporters. The YMDB builds upon the rich data sets already assembled by such resources as YeastNet 4.0 (19), MetaCyc (20), KEGG (21), UniProt (23), ChEBI (24) and HMDB (5). But it also brings in a large body of independently collected literature data, as well as a significant quantity of experimental data, including NMR spectra, MS spectra and validated metabolite concentrations, to compliment this electronic or literature-derived data.

The diversity of data types, the quantity of experimental data and the required breadth of domain knowledge made the assembly of the YMDB both difficult and time-consuming. To compile, confirm and validate this comprehensive collection of data, more than a dozen textbooks, several hundred journal articles, nearly 30 different electronic databases and at least 20 in-house or web-based programs were individually searched, accessed, compared, written or run over the course of the past 18 months. The team of YMDB contributors and annotators included analytical chemists, NMR spectroscopists, mass spectrosocists and bioinformaticians with dual training in computing science and molecular biology/chemistry.

The YMDB currently contains more than 2000 yeast metabolite entries that are linked to nearly 27 000 different synonyms. These metabolites are further linked to some 66 non-redundant pathways and 916 reactions involving 857 distinct enzymes and 138 transporters. More than 750 compounds are also linked to experimentally acquired ‘reference’ 1H and 13C NMR and MS/MS spectra. Concentration data (intracellular and extracellular) is also provided for a total of 627 compounds. The complete collection of data in the YMDB occupies a total of 1.1 GB. Relative to other yeast metabolite/pathway databases, YMDB is substantially larger and significantly more comprehensive. A detailed comparison of YMDB to other widely known yeast resources is provided in Table 1.

The YMDB is modeled closely after the HMDB. As a result, it has many of the features found in the HMDB including efficient, user-friendly tools for viewing, sorting and extracting metabolites, proteins, pathways or chemical taxonomy information (Figure 1). These are available through the YMDB navigation bar (located at the top of every YMDB web page) that lists seven pull-down menu tabs (‘Home’, ‘Browse’, ‘Search’, ‘About’, ‘Help’, ‘Download’ and ‘Contact Us’). To further aid in navigation and searching, nearly every viewable page in the YMDB, including the ‘Home’ page, supports simple text queries through a text search box located near the top of each YMDB web page. This text
search tool, which can be specified to search through either protein or metabolite data fields, supports text matching, accommodates misspellings and highlights the text where the word is found. A more advanced text search that supports Boolean constructs and permits more precise data field specifications is also available.

In addition to these extensive text search capabilities, the YMDB also offers general database browsing via the ‘Browse’ buttons located in the YMDB menu bar. Five different Browsing options are available including Metabolite Browse (for viewing and sorting metabolites), Protein Browse (for viewing and sorting proteins), Reaction Browse (for viewing chemical reactions), Pathway Browse (for viewing yeast-specific KEGG pathways) and Class Browse (for viewing groups of compounds by their chemical taxonomy or class). Each of the Browsing views is presented as a set of navigable/sortable synoptic summary tables. These tables are, in turn, linked to more detailed ‘MetaboCards’ and ‘ProteinCards’ similar to those found in DrugBank and HMDB. Clicking on a MetaboCard or ProteinCard button opens a web page describing the compound or protein of interest in much greater detail. Every MetaboCard entry contains 50 data fields devoted to chemical or physico-chemical data and synoptic biological data (names, sequences, accession codes). Each ProteinCard entry contains 30 data fields devoted to biochemical, nomenclature, gene ontology and sequence data for metabolically important yeast enzymes and transporters. In addition to providing comprehensive numeric, textual and factual data, each MetaboCard and ProteinCard also contains hyperlinks to many other databases (KEGG, BioCyc, PubChem, ChEBI, PubMed, PDB, UniProt, GenBank), abstracts, references, digital images and applets for viewing molecular structures.

Table 1. Comparison of the size and content of different yeast-specific or yeast-containing metabolism/metabolomics databases

| Database Content          | YMDB | YeastNet 4* | MetaCyc | KEGG |
|---------------------------|------|------------|---------|------|
| Number of metabolites     | 2007 | 792        | 688     | 720  |
| Number of data fields     | 81   | 14         | 19      | 12   |
| Number of NMR spectra     | 1540 | 0          | 0       | 0    |
| Number of MS spectra      | 1346 | 0          | 0       | 0    |
| Number of external        | 15   | 4          | 3       | 5    |
| database hyperlinks       |      |            |         |      |
| Concentrations            | Yes  | No         | No      | No   |
| Compound descriptions     | Yes  | No         | No      | No   |
| Cell locations            | Yes  | Yes        | No      | No   |
| Pathways                  | Yes  | No         | Yes     | Yes  |
| Sequence search           | Yes  | No         | Yes     | Yes  |
| Structure search          | Yes  | No         | No      | Yes  |
| Molecular weight search   | Yes  | No         | Yes     | No   |
| NMR spectral search       | Yes  | No         | No      | No   |
| MS spectral search        | Yes  | No         | No      | No   |
| Chemical taxonomy         | Yes  | No         | Yes     | No   |

*YeastNet 4.0 reports a total of 1494 metabolites but only 792 are unique.

The YMDB’s NMR and MS search utilities allow users to upload peak lists and to search for matching compounds from the database’s collection of MS and NMR spectra. The YMDB currently contains 1540 experimentally obtained 1H and 13C NMR spectra (with spectral collection conditions) for 466 different compounds (most collected in water at pH 7.0, 10 mM for 1H, 50 mM for 13C) measured in our lab or obtained from the BioMagResBank (BMRB) (26). Most of the NMR spectra are fully assigned. It also contains 951 MS/MS (Triple-Quad) spectra for 317 pure compounds analyzed by our laboratory. An additional 400 MS or MS/MS spectra were obtained from MassBank (27). The YMDB
Spectral search utilities allow both pure compounds and mixtures of compounds to be identified from their MS or NMR spectra via peak matching algorithms that were developed in-house (28,29).

Adjacent to the ‘Search’ menu, the ‘About’ pull-down menu contains information on the YMDB database, recent news or updates, links to other databases, data sources and database statistics. The ‘Help’ pull-down menu provides general documentation on database definitions, data field types and data field sources. It also contains information on experimental methods (for metabolite concentration measurements performed by our lab for the YMDB), details on how to cite YMDB, as well as a tutorial on how to use YMDB’s advanced text search utilities. Finally the ‘Download’ menu contains downloadable data for all YMDB chemical structures (as Structure Data Format (SDF) files), all enzyme/protein sequences (in FASTA format), as well as complete flat file data sets of the current YMDB release in JSON format.

Figure 1. A screenshot montage of the YMDB showing several of the YMDB’s search and data display tools describing the metabolite L-Glutamine. Not all fields are shown.
DATABASE IMPLEMENTATION

YMDB employs a Ruby on Rails (version: 3.09)-based front-end attached to a sophisticated MySQL relational database (version: 5.0.77) at its back-end. All data are entered directly through a custom-built web interface with each YMDB MetaboCard having an edit page, which allows database curators to manually make changes to YMDB entries. The public user interface and the internal database both read from the same database.

All structures in the YMDB are stored in a centralized structure hub. This hub is a RESTful web resource that automatically stores and updates chemical properties such as molecular weight, solubility and logP. Additionally, the hub renders the structure images and thumbnails visible on the public YMDB site. The centralized nature of this structure hub helps to maintain consistency for all structures stored in YMDB. Whenever a structure is changed or updated, all properties are automatically recalculated and made available on the public site at http://www.ymdb.ca.

QUALITY ASSURANCE, COMPLETENESS AND CURATION

The same quality assurance, quality control and data compilation procedures implemented during the development of HMDB, T3DB and DrugBank were used in the development of YMDB. In particular, the compounds in YMDB were identified using a combination of methods, including manual literature surveys, text mining of on-line journals or abstracts and data mining of other electronic databases. Literature sources included specialty journals on metabolomics, food composition and analysis, systems biology, analytical chemistry and textbooks on wine and beer chemistry. All primary metabolites had to have at least two databases confirm their existence and inclusion (with evidence that the necessary enzymes or pathways are present), whereas all secondary metabolites (such as those found in wine or beer) were required to have a traceable literature/experimental reference. For many secondary yeast metabolites the relevant starting compounds, reactions, pathways and catalyzing enzymes are not yet known. Hopefully, with time and improved technology, this information will become available. With many yeast secondary metabolites it is sometimes difficult to know if the compound was present in the media (wort or grape must) prior to fermentation or whether it arose as a consequence of fermentation. For those compounds where there was some ambiguity regarding their source (plant versus yeast), we attempted to cross-check our findings through multiple literature sources in order to exclude possible grape, hops or barley metabolites.

For those yeast metabolites found to match to previously existing entries from either the HMDB or CCDB, only the chemical data fields were imported into the YMDB (except the compound description which was manually edited to include or remove organism-specific references). The biological data for these HMDB/CCDB imported compounds was generated de novo since yeast biology is very different than E. coli or human biology. In order to ensure both completeness and correctness, each metabolite record entered into the YMDB was reviewed and validated by a member of the curation team after being annotated by another member. Other members of the curation group routinely performed additional spot checks on each entry. Several software packages including text-mining tools, chemical parameter calculators and protein annotation tools were developed, modified and used to aid in data entry and data validation. One particular program, BioSpider (30), was used extensively to acquire routine, machine retrievable or easily calculated/verifiable chemical data on metabolites. To facilitate and monitor the data entry process, all of YMDB’s data is entered into a centralized, password-controlled database, allowing all changes and edits to the YMDB to be monitored, time-stamped and automatically transferred.

CONCLUSIONS

To summarize, the YMDB is a richly annotated, web-accessible ‘metabolomics’ database that brings together quantitative chemical, physical and biological data about nearly 2000 S. cerevisiae metabolites. Relative to other yeast metabolism/pathway databases, YMDB has between 2–3× more metabolites and 5–10× more data. The YMDB also uniquely contains detailed information on hundreds of secondary metabolites that are critically important to the food, beverage, chemical and biofuel industry. Among the other distinguishing features of YMDB are: (i) the breadth and depth of its annotations (>80 data fields); (ii) the large number of hyperlinks and references to other resources; (iii) the availability of detailed compound descriptions; (iv) the inclusion of thousands of reference NMR and MS spectral data; (v) the inclusion of intra- and extracellular metabolite concentration data; (vi) the quantity of biological and biochemical information included in each compound entry and (vii) the support for queries by text, chemical structure, spectra, molecular weight and gene/protein sequence. Owing to these unique characteristics, we believe the YMDB fills an important niche in yeast biology as it addresses not only the specialized analytical needs of metabolomics researchers, but also the interests of molecular biologists, systems biologists, the industrial fermentation industry, as well as the beer, wine and spirit industries.

While the YMDB certainly fills an important niche for yeast metabolomics, it is also a work in progress. As with many areas in metabolomics, new compounds are constantly being discovered, new concentrations are being reported, new pathways/reactions are being elucidated and new metabolite functions are being determined. So long as our resources permit, we intend to continue to update and enhance the YMDB as this new information is published or acquired.

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