WEB SITE REVIEW

Review of the Enzymes and Metabolic Pathways (EMP) Database

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INTRODUCTION

The Enzymes and Metabolic Pathways (EMP) database is an electronic compilation of enzymatic and metabolic data from over 15,000 literature sources covering over 1800 different organisms (Selkov et al., 1996). The Metabolic Pathways Database (MPW) is a derivative of the larger EMP database containing over 3000 pathway diagrams of primary and secondary metabolism, membrane transport, signal transduction, intracellular traffic, translation, and transcription (Selkov et al., 1998). The assembly of the encoded data and of the pictorial pathway diagrams is an ongoing project centered at the Laboratory of Mathematical Simulation of Multi-enzyme Systems at the Institute of Theoretical and Experimental Biophysics of the Russian Academy of Sciences, in Pushchino, Russia. These resources can be accessed free of charge by academic and nonprofit users at http://www.empproject.com.

DESCRIPTION OF EMP AND MPW

Each record in the EMP database translates the entire factual content of a journal publication into a structured, indexed, and searchable form. Altogether over 300 subject fields are employed to describe the various categories of biological importance shown in Table 1. A comprehensive description of the subject fields is available at the EMP Web site (2001). The database can be queried by means of a whole text search, taxonomy search, by accession number (i.e., a unique key that identifies a record), or by enzyme commission (EC) number. In addition to the type of information shown in Table 1, each entry provides search links to the MEDLINE (Wheeler et al., 2001), ENZYME (Bairoch, 2000), BREND A (2001), and LIGAND (Goto et al., 2000) databases for complementary information as well as the metabolic pathway name for easy retrieval from MPW.

TABLE 1

The Number of Subject Fields in each EMP Informational Category

<table>
<thead>
<tr>
<th>Subject Fields</th>
<th>Total Subject Fields</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bibliography</td>
<td>16</td>
</tr>
<tr>
<td>Biochemical Genetics</td>
<td>18</td>
</tr>
<tr>
<td>Biological Source</td>
<td>19</td>
</tr>
<tr>
<td>Cell Cultivation</td>
<td>19</td>
</tr>
<tr>
<td>Enzyme Kinetics and</td>
<td>21</td>
</tr>
<tr>
<td>Reaction Mechanisms</td>
<td>51</td>
</tr>
<tr>
<td>Enzyme Regulation</td>
<td>15</td>
</tr>
<tr>
<td>Enzyme Structure</td>
<td>30</td>
</tr>
<tr>
<td>Enzyme and Reaction</td>
<td>28</td>
</tr>
<tr>
<td>Equilibrium and</td>
<td>14</td>
</tr>
<tr>
<td>Thermodynamics</td>
<td>19</td>
</tr>
<tr>
<td>Host</td>
<td>31</td>
</tr>
</tbody>
</table>

The collection of pathways found in the MPW database originated as the “working notes” of Evgeni Selkov (Selkov et al., 1997) and is currently updated with approximately 1000 pathways a year by the Pushchino team (EMP Web site, 2001). Pathway maps include not only the stoichiometry of the metabolic pathways, but also the substrate and coenzyme specificity of their enzymes, their subcellular locations, required prosthetic groups and cofactors, and the taxonomic occurrence of the pathways. All pathway names are generated in the format substrates-products-function(coenzymes)–(locations)–[comment]. Figure 1 shows the MPW-generated pathway diagram for glutathione biosynthesis named L-glutamate, L-cysteine–glutathione anabolism (ATP) (cytosol). This pathway was also used as an example in the last issue's Web site review (Maranas and Burgard, 2001) of the EcoCyc and MetaCyc databases (Karp et al., 2000). As is the case with the EcoCyc and MetaCyc pathways, each compound or reaction number in the MPW pathways is linked to additional information. On the other hand, unlike the EcoCyc pathways which focus solely on the metabolism of Escherichia coli, species-specific regulatory structure information is not available through MPW.

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Cytosol

FIG. 1. Glutathione biosynthesis pathway obtained from the EMP-MPW Web site.

STRENGTHS AND WEAKNESSES

A key strength of the EMP database is the depth of enzyme-specific biochemical data for an extensive list of organisms. This information finds applications ranging from the analysis and mathematical simulation of metabolic networks (Goryanin et al., 1999) to metabolic modification (Klinke et al., 2000) and drug development (Black and Hare, 2000; Selzer et al., 2000). Although the high degree of detail for each reference summary may be overwhelming for novice users, a quite useful "help" program and tutorial are available at the EMP Web site. The major strength of the MPW part of the EMP database is its usefulness in metabolic reconstructions. Once the functional assignment of a sequenced genome is complete, genes can be attached to metabolic reconstructions. Nevertheless, information along these lines can be obtained by querying the EMP database literature sources. In summary, the wealth of biochemical data accessible through the EMP/MPW database makes it a very good starting point for identifying detailed information on enzymes, compounds, or pathways.

REFERENCES


