

The EcoCyc and MetaCyc Databases

Peter D. Karp*, Monica Riley**, Milton Saier***, Ian T. Paulsen***, Suzanne M. Paley*
and Alida Pellegrini-Toole**

October 8, 1999

*Pangea Systems Inc, 4040 Campbell Ave., Menlo Park, CA 94025; pkarp@PangeaSystems.com.

**Marine Biological Laboratory, Woods Hole, MA 02543; mriley@mbi.edu.

***Department of Biology, University of California at San Diego, La Jolla, CA 92093-0116; msaier@ucsd.edu.

Abstract

EcoCyc is an organism-specific Pathway/Genome Database that describes the metabolic and signal-transduction pathways of *E. coli*, its enzymes, and — a new addition — its transport proteins. MetaCyc is a new metabolic-pathway database that describes pathways and enzymes of many different organisms, with a microbial focus. Both databases are queried using the Pathway Tools graphical user interface, which provides a wide variety of query operations and visualization tools. EcoCyc and MetaCyc are available at URL <http://ecocyc.PangeaSystems.com/ecocyc/>.

1 Introduction

The EcoCyc and MetaCyc databases (DBs) are online reference sources for metabolic data. They are similar in describing metabolic pathways, reactions, enzymes and substrate compounds. Both DBs use the same DB schema. Both are accessed using the same software environment, called the Pathway Tools. Both are review-level DBs in that a given entry in either DB often integrates information from multiple literature citations. There is also overlap in the content of the DBs — both contain all pathways of *E. coli* small-molecule metabolism.

The DBs do differ significantly in content. EcoCyc aims to describe the full biochemical network of *E. coli*. As well as describing the metabolism of *E. coli*, EcoCyc describes its signal-transduction pathways, its transporters, and all *E. coli* genes. MetaCyc does not focus on a single organism as EcoCyc does. It describes pathways from a wide variety of species, with a focus on microorganisms, but includes some human and other mammalian pathways as well.

Intended uses of the DBs include the following.

- MetaCyc is a general reference source on metabolic pathways for the scientific community. It also serves as a reference pathway DB for prediction of the pathway complement of an organism from the annotated genome of the organism [1].
- EcoCyc is a resource for analysis of microbial genomes at the level of individual genes. Because the *E. coli* genome has a high fraction of genes whose functions were determined experimentally, it is an accurate reference for inferring gene function by sequence similarity.
- EcoCyc and MetaCyc describe the subunit structures of many enzymes, and therefore could be used as training or validation datasets for algorithms that detect protein–protein interactions.
- Because of its links to sequence DBs such as Swiss-Prot, EcoCyc can be used to perform function-based retrieval of DNA or protein sequences, for example to prepare datasets for studies of protein structure–function relationships.
- Both EcoCyc and MetaCyc are used as an aid in teaching biochemistry.

This article describes recent enhancements to EcoCyc and MetaCyc, and how to access them. We request that users of these DBs cite this article in publications related to their use.

Version 5.0 of EcoCyc was released in June, 1999; version 5.0 of MetaCyc was released in September 1999. In the future the same version number will be used for both DBs, and their releases will be synchronized.

2 Pathway Tools Software and Database Environment

The Pathway Tools software that underlies EcoCyc and MetaCyc provides query, editing, and visualization operations for Pathway/Genome DBs [2, 3, 1]. The Pathway Tools is an environment for *functional bioinformatics* — for managing, curating, and computing with a functional genome annotation. The Pathway Tools utilize a frame knowledge representation system (FRS) called Ocelot [3, 4]. FRSs use an object-oriented data model that organizes information within classes: collections of objects that share similar properties and attributes.

Recent software enhancements now allow users to issue structured queries to EcoCyc and MetaCyc using a new HTML query form that is available at URL <http://ecocyc.pangeasystems.com:1555/query.html>. This query interface allows the user to specify the class of objects they wish to query, plus a set of constraints that select the objects of interest, plus a set of attributes to be returned for each object. For example, the user might query the class of all genes to select all genes whose chromosomal location is within a specific location, and ask that the results include the unique identifier of each gene, its name, the function of its product, and its map position.

The EcoCyc/MetaCyc web site was upgraded and reorganized for the 5.0 release. Although the location of the EcoCyc home page has not changed, the URL of

the page used to formulate most queries to EcoCyc and MetaCyc has changed to <http://ecocyc.pangeasystems.com:1555/server.html>.

3 EcoCyc

Table 1 shows the current size of the principal EcoCyc and MetaCyc classes.

One of the recent enhancements to Ecocyc is the inclusion of membrane transport systems. Metabolic pathways are dependent on membrane transport processes for the uptake of exogenous substrates and for the export of metabolic end products. Membrane transporters also play a role in other important cellular processes such as energy interconversion, ion homeostasis, and resistance to drugs and toxic compounds. Therefore, we believe that membrane transport systems represent a significant addition to Ecocyc and will help provide an understanding of the interplay between metabolic pathways and their corresponding transport systems.

A complete list of known and putative *E. coli* cytoplasmic membrane transporters has previously been compiled, and each of these transporters was classified on the basis of amino acid sequence similarities and transport function [5] (see also <http://www-biology.ucsd.edu/~ipaulsen/transport/> and (<http://www-biology.ucsd.edu/~msaier/transport/>). This compilation of over 300 transporters was imported into Ecocyc, and each transporter is now being annotated using the same detailed, literature-based approach that EcoCyc uses for *E. coli* enzymes and pathways. For each transporter, a comprehensive literature search of Medline and Current Contents is performed for relevant keywords. Citations in relevant journal papers are used to identify further pertinent articles. Additionally, the background knowledge of transport systems of the two transport editors proved invaluable, and in some instances experts working on particular transport systems are contacted to clarify their findings.

For each transporter, where possible, the annotation includes: (a) Functional information such as the energy-coupling mechanism, substrate specificity and relative affinity of each transporter, (b) A brief description of the experimental evidence for function, e.g., cloning and expression data, knockout mutants, protein purification and functional reconstitution, vesicle or whole cell transport assays, (c) Sequence similarity to other known transporters, (d) The inferred physiological role of the transporter and/or the role of its substrate in metabolism, (e) In some cases additional relevant information is provided, such as structure/function studies, domain structure, details of transcriptional regulation.

Version 5.0 of EcoCyc contains all of the *E. coli* phosphotransferase system (PTS) transporters. Subsequent versions will include detailed descriptions of all of the other transporters present in *E. coli* (primary and secondary transporters, and channels). The PTS functions by transporting and concomitantly phosphorylating its sugar substrates using phosphoenolpyruvate as a phosphate donor. Each PTS transporter characteristically consists of a sugar-specific multidomain Enzyme II complex with between one and four protein subunits and two general energy coupling proteins, Enzyme I and HPr. The Version 5.0 release describes each of the 20 Enzyme II complexes and the two general energy coupling proteins of the PTS.

A representative example of an Ecocyc transporter entry is available at WWW URL <http://ecocyc.PangeaSystems.com:1555/ECOLI/new-image?type=ENZYME&object=CPLX-166>. This entry describes the mannitol PTS transporter Enzyme II^{Mtl}, encoded by the *mtlA* gene. Mannitol is imported and concomitantly phosphorylated by Enzyme II^{Mtl} to yield mannitol-1-phosphate. Phosphoenolpyruvate is used as energy source and phosphate donor. The phosphoryl transfer reaction involves Enzyme I and HPr in addition to Enzyme II^{Mtl}.

The inclusion of transport systems in the Ecocyc KB not only provides a valuable resource for researchers, but also provides the opportunity for probing the interrelationships between transport and metabolism at a global level. For instance, in preliminary analyses we have compared the complete inventory of compounds transported into the cell with the starting reactants of all of the metabolic pathways of *E. coli*, as well as with the complete lists of metabolites and cofactors present in *E. coli* (Paulsen and Karp, unpublished data). This analysis provides an idea of the transporters which may be present in *E. coli*, but have not yet been experimentally identified, as well as highlighting the physiological role of the transported substrates.

Our schema for transport data builds upon our representations of enzyme function. Each transporter is represented by one or more DB objects that encode the transporter and its monomer subunits, if any. The transporter is linked to one or more (in the case of multifunctional transporters) objects that describe its function as a biochemical reaction. We have extended our schema for reactions to allow each substrate to be tagged with a cellular compartment, which if omitted defaults to the cytoplasm. For example, the representation of the mannitol PTS transporter tags the substrate mannitol with the compartment periplasm; the other substrates default to the cytoplasm. The PTS provides a strong example of why the reaction-based representation is a suitable one, since these transport systems both translocate and chemically alter their substrates.

4 MetaCyc

MetaCyc is a meta-metabolic pathway database that contains pathways from a variety of different organisms, which are listed in Table 2. MetaCyc describes metabolic pathways, reactions, enzymes, and substrate compounds. The MetaCyc data were gathered from a variety of literature and on-line sources. “MetaCyc” is pronounced “met-a-sike”. It sounds like “encyclopedia”.

MetaCyc has two major goals: to provide a reference source on metabolic pathways that is accessible to scientists through the WWW, and to serve as a reference for computational prediction of metabolic pathways for sequenced genomes [6]. The philosophy of MetaCyc is to encode pathways that have been reported in the experimental literature, and to label each pathway with the organism(s) in which it is known to occur, based on wet-lab experiments. Thus, unlike EcoCyc, which aims to describe the complete metabolic map of a single organism, MetaCyc aims to provide a smorgasbord of pathways from many organisms, none of whose metabolic maps have been studied in as much detail in the laboratory as has that of *E. coli*.

MetaCyc employs the same database schema as does EcoCyc. It aims to provide the same rich literature-based annotation for each pathway as does EcoCyc, although a minority of pathways currently lack the extensive commentary and literature citations that we plan to provide. Each

MetaCyc pathway contains a citation to the source from which it was obtained. Unlike EcoCyc, MetaCyc does not provide genomic data such as genomic maps or sequences.

MetaCyc was initialized to contain all metabolic pathways of EcoCyc. Many additional pathways were then added to the database. The majority of pathways have been added by Dr. Riley's group at the Marine Biological Laboratory, but some pathways have added by Dr. Karp's group at Pangea Systems. The pathways have been gathered from many different phyla, including micro-organisms, plants, and humans.

Each MetaCyc pathway has a slot (attribute) called "species distribution" that lists the one or more species in which the experimental literature reports that this pathway has been observed. The fact that a given species is not listed in the species distribution of a pathway does not necessarily imply that the pathway is not present in that species — experiments may not yet have been performed, or the relevant literature may not have been located. Table 2 lists the species for which MetaCyc 5.0 contains pathways. Figure 1 shows the length distributions of MetaCyc pathways.

Each distinct pathway (where a pathway is defined as a set of reaction steps that are connected in a particular topology) is encoded only one time in MetaCyc — separate observations of the same pathway in new organisms do expand the species distribution recorded for that pathway, but those new observations do not result in new pathway records in MetaCyc unless the pathway differs in its component reactions. Different variants of a pathway that are observed in different organisms are recorded as distinct pathway objects in the DB. Different pathway variants might differ by the addition or removal of one or more reactions, or by substitution of one reaction for a similar reaction — such as substituting a reaction in which NAD is the hydride acceptor for a reaction in which NADP is the hydride acceptor.

The pathways within MetaCyc are annotated at different levels of detail. For all pathways, the pathway DB object is linked to objects for each reaction in the pathway. For some pathways, additional DB objects are provided for each enzyme and for each enzyme subunit in the pathway, and the pathway and enzyme include extensive commentary and literature citations.

MetaCyc contains all reactions in the enzyme-classification system devised by the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology (NC-IUBMB). It also contains thousands of objects representing individual metabolites.

The MetaCyc data reside within the same software environment as used for EcoCyc: the Pathway Tools. The Pathway Tools run in both a WWW mode and an X-windows mode. Both EcoCyc and MetaCyc are accessed through a query form at the URL <http://ecocyc.PangeaSystems.com:1555/server.html>. The user selects the DB to query using the dataset selector button near the top of the form. All query options in the remainder of the form can be applied to either DB. Put another way, all of the visualization and query tools available for EcoCyc are also available for MetaCyc (the exceptions being the genomic-map browser, which is not available for MetaCyc because MetaCyc contains no genomic maps).

5 Distribution

EcoCyc and MetaCyc are available under license from Pangea Systems in two forms:

- EcoCyc and MetaCyc are accessible online through the WWW at URL <http://ecocyc.PangeaSystems.com/ecocyc/> (this version supports a subset of the GUI functionality of the X-windows version).
- An X-windows version of EcoCyc and MetaCyc for the Sun workstation bundles together the Pathway/Genome Navigator software with the EcoCyc and MetaCyc DBs.

Both of the preceding forms of access are free to academic institutions for research use (contact pkarp@PangeaSystems.com); a fee applies to other forms of use. The EcoCyc/MetaCyc WWW site provides background information about the DBs and software, and access to the publications produced by the EcoCyc project.

Acknowledgments

This work was supported by grant 1-R01-RR07861-01 from the Comparative Medicine Program at the National Center for Research Resources. The contents of this article are solely the responsibility of the authors and do not necessarily represent the official views of the National Institutes of Health.

References

- [1] Karp, P.D. (1999) *Trends in Biotechnology*, 17(7):275–281.
- [2] Karp, P.D. (1999) In *EcoCyc: The Resource and the Lessons Learned*, 47–62. Kluwer Academic Publishers.
- [3] Karp, P.D., and Paley, S. (1996) *J. Comp. Biol.*, 3(1):191–212.
- [4] Karp, P.D., Chaudhri, V.K., and Paley, S.M. (in press) *Journal of Intelligent Information Systems*.
- [5] Sliwinski, M.K., Paulsen, I.T., and Saier, M.H. (1998) *J. Mol. Biol.*, 277:573–592.
- [6] Karp, P.D., Ouzounis, C., and Paley, S.M. (1996) In States, D.J., Agarwal, P., Gaasterland, T., Hunter, L., and Smith, R., editors, *Proceedings of the Fourth International Conference on Intelligent Systems for Molecular Biology*, 116–124, Menlo Park, CA, . AAAI Press.

	EcoCyc	MetaCyc
Metabolic Pathways	139	305
Signaling Pathways	20	0
Reactions	946	3786
Enzymes	629	71
Genes	4390	0
tRNAs	79	0
Compounds	1868	1963
Citations	1944	315

Table 1: The number of objects in version 5.0 of EcoCyc and MetaCyc.

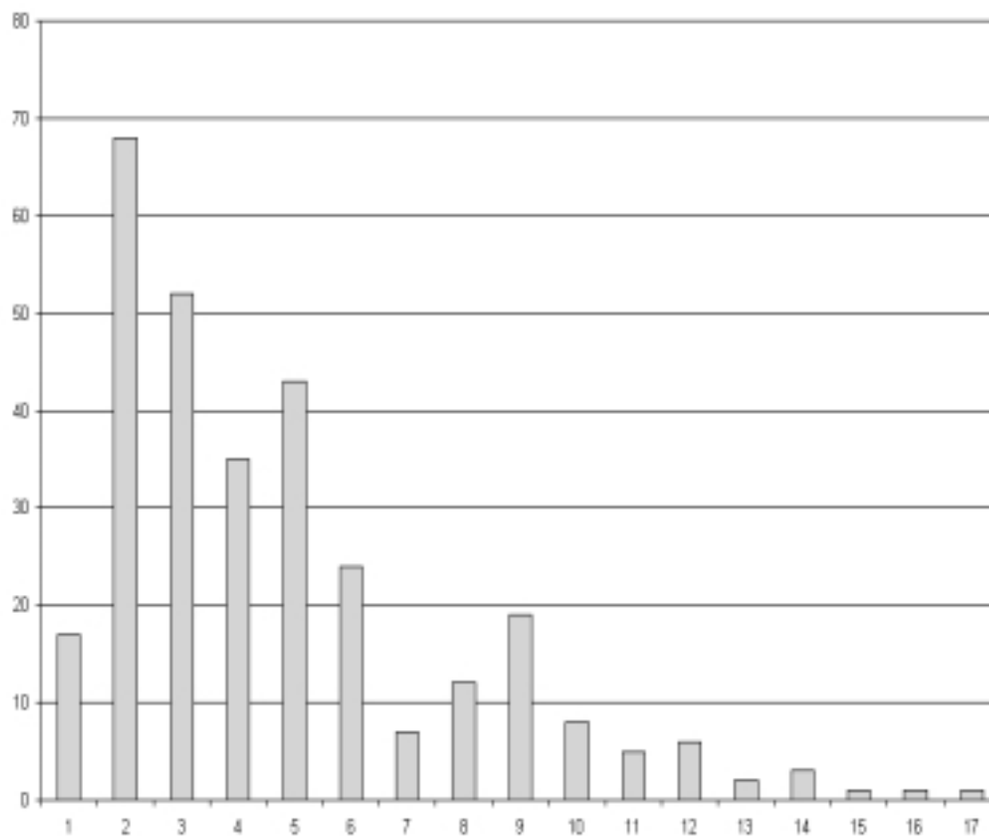


Figure 1: The length distribution of MetaCyc pathways. We graph the number of MetaCyc pathways having a given length in reaction steps.

<i>Acinetobacter</i>	<i>Methanosarcina thermophila</i>
<i>Actinomycetes</i>	<i>Methanospirillum spp.</i>
<i>Aerobacter aerogenes</i>	<i>Methanothermus spp.</i>
<i>Archaeobacteria</i>	<i>Micrococcaceae</i>
<i>Arthrobacter aurescens TW17</i>	<i>Moraxella sp</i>
<i>Arthrobacter globiformis</i>	<i>Mycoplasma arthritidis</i>
<i>Arthrobacter simplex</i>	<i>Mycoplasma capricolum</i>
<i>Ascomycotina</i>	<i>Mycoplasma genitalium</i>
<i>Aves</i>	<i>Neisseriaceae</i>
<i>Azotobacter beijerinckii</i>	<i>Nocardia sp. B-1 and TW2</i>
<i>Brevibacterium</i>	<i>Oryctolagus cuniculus</i>
<i>Brevibacterium helvolum</i>	<i>Ovis aries</i>
<i>Bacillus subtilis</i>	<i>Petunia sp</i>
<i>Burkholderia (Pseudomonas) cepacia</i>	<i>Physarum polydephalum</i>
<i>Citobacter</i>	<i>Protozoa</i>
<i>Clostridium acetobutylicum</i>	<i>Pseudomonadacea</i>
<i>Clostridium pasteurianum</i>	<i>Pseudomonas acidovorans</i>
<i>Clostridium propionicum</i>	<i>Pseudomonas aeruginosa</i>
<i>Clostridium tetanomorphum</i>	<i>Pseudomonas alcaligenes NCIB 9867</i>
<i>Corynebacterium</i>	<i>Pseudomonas aureofaciens</i>
<i>Desulfovibrio gigas</i>	<i>Pseudomonas diminuta MG</i>
<i>E. coli</i>	<i>Pseudomonas fluorescens</i>
<i>Embryophyta</i>	<i>Pseudomonas mendocina (pWWO) or (pTOL)</i>
<i>Enterobacter aerigenes</i>	<i>Pseudomonas picketti</i>
<i>Enterobacter cloacae</i>	<i>Pseudomonas putida</i>
<i>Flavobacterium</i>	<i>Pseudomonas putida F1</i>
<i>Fungi imperfecti</i>	<i>Pseudomonas putida mt-2</i>
<i>Haemophilus influenzae</i>	<i>Rattus norvegicus</i>
<i>Homo sapiens</i>	<i>Rhizobiaceae</i>
<i>Klebsiella aerogenes</i>	<i>Rhodobacteriaceae</i>
<i>Klebsiella pneumoniae</i>	<i>Rhodococacea</i>
<i>Lactobacillaceae</i>	<i>Rhodococcus maris</i>
<i>Lactococcus lactis</i>	<i>Rodentia</i>
<i>Mammalia</i>	<i>Ruminantia</i>
<i>Methanobacterium thermoautotrophicum</i>	<i>Saccharomyces cerevisiae</i>
<i>Methanobrevibacter spp.</i>	<i>Salmonella typhimurium</i>
<i>Methanococcus spp.</i>	<i>Sporosarcina</i>
<i>Methanocorpusculum spp.</i>	<i>Streptomyces parvulus</i>
<i>Methanoculleus spp.</i>	<i>Sulfolobus solfataricus</i>
<i>Methanogenium spp.</i>	<i>Sus scrofa</i>
<i>Methanomicrobium spp.</i>	<i>Synechocystis sp. strain PCC 6803</i>
<i>Methanoplanus spp.</i>	<i>Synechocystis spp. PCC 6803</i>
<i>Methanopyrus spp.</i>	<i>Teleostei</i>
<i>Methanosarcina barkeri</i>	
<i>Methanosarcina spp.</i>	

Table 2: MetaCyc contains pathways from these species.