KEGG: Kyoto Encyclopedia of Genes and Genomes

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ABSTRACT

Kyoto Encyclopedia of Genes and Genomes (KEGG) is a knowledge base for systematic analysis of gene functions in terms of the networks of genes and molecules. The major component of KEGG is the PATHWAY database that consists of graphical diagrams of biochemical pathways including most of the known metabolic pathways and some of the known regulatory pathways. The pathway information is also represented by the ortholog group tables summarizing orthologous and paralogous gene groups among different organisms. KEGG maintains the GENES database for the gene catalogs of all organisms with complete genomes and selected organisms with partial genomes, which are continuously re-annotated, as well as the LIGAND database for chemical compounds and enzymes. Each gene catalog is associated with the graphical genome map for chromosomal locations that is represented by Java applet. In addition to the data collection efforts, KEGG develops and provides various computational tools, such as for reconstructing biochemical pathways from the complete genome sequence and for predicting gene regulatory networks from the gene expression profiles. The KEGG databases are daily updated and made freely available (http://www.genome.ad.jp/kegg/).

INTRODUCTION

The progress in structural genomics will soon uncover the complete genome sequences for hundreds and thousands of organisms. However, roughly a half of the genes identified in every genome thus far sequenced still remain unknown in terms of their biological functions. New experimental and informatics technologies in functional genomics are urgently required for systematic identification of gene functions. Kyoto Encyclopedia of Genes and Genomes (KEGG) is an effort to computerize the current knowledge of biochemical pathways and other types of molecular interactions that can be used as reference for systematic interpretation of sequence data (1). At the same time, KEGG attempts to standardize the functional annotation of genes and proteins, and maintain gene catalogs for all complete genomes and some partial genomes including mouse and human.

The basic concepts of KEGG (1) and underlying informatics technologies (2,3) have already been published. KEGG is tightly integrated with the LIGAND chemical database for enzyme reactions (4,5) as well as with most of the major molecular biology databases by the DBGET/LinkDB system (6) under the Japanese GenomeNet service (7). The database organization efforts require extensive analyses of completely sequenced genomes, as exemplified by the analyses of metabolic pathways (8) and ABC transport systems (9). In this article, we describe the current status of the KEGG databases and discuss the use of KEGG for functional genomics.

OBJECTIVES OF KEGG

In May 1995, we initiated the KEGG project under the Human Genome Program of the Ministry of Education, Science, Sports and Culture in Japan. We wish to automate human reasoning steps for interpreting biological meaning encoded in the sequence data. We consider the problem of predicting gene functions as a process of reconstructing a functioning biological system from the complete set of genes and gene products. Thus, it is critical to understand how genes and molecules are networked to form a biological system. Specifically, the objectives of KEGG are summarized in the following four points.

First, KEGG aims at computerizing the current knowledge of genetics, biochemistry, and molecular and cellular biology in terms of the pathway of interacting molecules or genes. The KEGG pathway database contains the information of how molecules or genes are networked, which is complementary to most of the existing molecular biology databases that contain the information of individual molecules or individual genes. During the first two years of the KEGG project we focused on the metabolic pathways, but since July 1997 we have also been collecting a large body of knowledge in the regulatory aspects of cellular functions.

Second, KEGG maintains the gene catalogs for all organisms with completely sequenced genomes and selected organisms with partial genomes. Because the criteria of interpreting sequence similarity are different for different authors, the quality of gene function annotations varies significantly in GenBank (10). KEGG’s gene catalogs are intended to provide consistent and standardized annotations by linking individual genes to components of the KEGG biochemical pathways.

Third, KEGG maintains the catalog of chemical elements, compounds, and other substances in living cells as the LIGAND

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Table 1. The summary of KEGG release 8.0 (October 1998)

<table>
<thead>
<tr>
<th>Database / auxiliary data</th>
<th>Number of entries</th>
<th>Contents</th>
<th>Direct links</th>
<th>Additional links by DIGEST/LinkDB*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PATHWAY</td>
<td>2,085 entries</td>
<td>GIF image map showing biochemical pathways</td>
<td>GENES, ENZYME, COMPOUND, other PATHWAY</td>
<td>Goodrich, EMBL, SWISS-PROT, PIR, PRF, PDB, PROSITE, EBI, TRANSFAC, PROSITE, GMM, PMID, AAindex, LITDB, Medline, similar sequences</td>
</tr>
<tr>
<td>GENES</td>
<td>76,891 entries in 17 complete genomes and 12 partial genomes</td>
<td>Nomenclature, classification, codon frequency, amino acid sequence, foldilf sequence</td>
<td>PATHWAY, ENZYME, GENOME, map, ENZYME, SWISS-PROT</td>
<td></td>
</tr>
<tr>
<td>LIGAND/GROUP</td>
<td>68,566 entries</td>
<td>Nomenclature, chemical formula, molecular weight, CAS number</td>
<td>PATHWAY, ENZYME, GENOME, map, ENZYME, SWISS-PROT</td>
<td></td>
</tr>
<tr>
<td>LIGAND/ENZYME</td>
<td>3,391 entries</td>
<td>EC number, nomenclature, classification, reaction formula, substrates, product, inhibitor, effects, targets</td>
<td>PATHWAY, ENZYME, GENOME, map, ENZYME, SWISS-PROT</td>
<td></td>
</tr>
<tr>
<td>Genome map</td>
<td>17 complete genomes and 1 partial genome</td>
<td>Java applet for searching and browsing genes in a genome map</td>
<td>GENES, PATHWAY</td>
<td></td>
</tr>
<tr>
<td>Comparative genome map</td>
<td>17 x 37 complete genome comparisons</td>
<td>Java applet for comparing genes in two genome maps</td>
<td>GENES</td>
<td></td>
</tr>
<tr>
<td>Ortholog group table</td>
<td>4,772 genes in 51 tables</td>
<td>HTML table for functionally related genes, orthologs and paralogs, and operon structure</td>
<td>PATHWAY, ENZYME, SWISS-PROT</td>
<td></td>
</tr>
<tr>
<td>Gene catalog</td>
<td>43 catalogs</td>
<td>Classification of genes (hierarchical taxon)</td>
<td>PATHWAY, GENES, ENZYME, SWISS-PROT</td>
<td></td>
</tr>
<tr>
<td>Molecular catalog</td>
<td>8 catalogs</td>
<td>Classification of proteins, RNAs, and compounds</td>
<td>GENES, ENZYME, COMPOUND, SWISS-PROT</td>
<td></td>
</tr>
</tbody>
</table>

*Additional links can be obtained by clicking on the entry name of each entry; in the case of a graphical pathway map, by clicking on the title box.

database (4,5) and they are again linked to pathway components. This is based on our view that both the genetic information encoded in the genome and the chemical information encoded in the cell are required for understanding cellular functions.

Fourth, in addition to the database efforts, KEGG aims at providing new informatics technologies that combine genomic and functional information towards predicting biological systems and designing further experiments. For example, the KEGG reference pathways can be used to uncover molecular interactions and pathways that underlie gene expression profiles obtained by microarray experiments (11).

THE KEGG DATABASES

All the data in KEGG can be accessed from the KEGG table of contents page (http://www.genome.ad.jp/kegg/kegg2.html ), which is divided into two broad categories: the pathway (functional) information and the genomic (structural) information. The user may enter the KEGG system top-down starting from the pathway information or bottom-up starting from the genomic information. In either case it should be easy to navigate through KEGG since the information is highly integrated. The KEGG data are also linked with many other molecular biology databases by the DBGET/LinkDB system. A summary of KEGG data contents is shown in Table 1.

PATHWAY database

The KEGG/PATHWAY database is a collection of graphical diagrams (pathway maps) for the biochemical pathways. Table 2 shows the current list of the KEGG biochemical pathways, which is highly biased toward the metabolism. There are about 90 reference maps for the metabolic pathways that are manually drawn and continuously updated according to biochemical evidence. The organism-specific pathway maps are then automatically generated by matching the EC numbers in the gene catalogs and in the reference maps. The maps for the regulatory pathways are drawn separately for each organism, since they are too divergent to be represented in a single reference map.

Figure 1 shows an example of the KEGG metabolic pathway map for citrate (TCA) cycle for Haemophilus influenzae and Helicobacter pylori. A box is an enzyme (gene product) with the EC number inside and a circle is a metabolic compound. They are both clickable objects to retrieve detailed molecular information. The enzymes (boxes) whose genes are identified in the genome are colored green (shaded in Fig. 1) by the process of matching the gene catalog and the reference pathway according to the EC numbers. It is interesting to note that the two organisms seem to

Table 2. The KEGG biochemical pathways (release 8.0, October 1998)

| Metabolism | D-Alanine metabolism | Glutathione metabolism | Alkaloid biosynthesis I | Alkaloid biosynthesis II | Methionine of Complex Carbohydrates | Starch and sucrose metabolism | Peptidoglycan biosynthesis | Methionine of Complex Lipids | Glyceraldehyde metabolism | Isoprenoid metabolism | Glycine and aspartate metabolism | Serine metabolism | Methionine metabolism | Riboflavin metabolism | Vitamin B1 metabolism | Nicotinate and oxidoreductase metabolism | Fumarate and CoA biosynthesis | Biotin metabolism | Folate biosynthesis | One carbon pool from folate | Retinol metabolism | Pyrophosphate and chlorophyll metabolism | Terpenoid biosynthesis | Xenobiotics metabolism | Uroquinase metabolism | Flavonoids, alkaloids, and lignin biosynthesis | Methionine of Macromolecules | mRNA Biosynthesis (bacteria) | RNAI and RNAII Biosynthesis (bacteria) | RNAI Bioynthesis (bacteria) | RNAII Biosynthesis (bacteria) | Proline Biosynthesis (bacteria) | Proline Biosynthesis (yeast) | Amylase-I RNAI biosynthesis | Cell Processes | Membrane Transport | ATP-binding cassette (ABC) transport system | Phosphatase system (PTS) | Signal Transduction | Two-component system | Bacterial chemotaxis | MAPK signaling pathway | Ligand-activated receptor signaling | G-protein coupled receptors | Cytokine receptors | Molecular Assembly | Ribosome assembly | Flagellar assembly | FHF ATP synthase |
Figure 1. The KEGG pathway map of citrate (TCA) cycle for (a) *Haemophilus influenzae* and (b) *Helicobacter pylori*. A rectangle and a circle represent, respectively, an enzyme and a compound. The enzymes whose genes are identified in the genome are shown by colored (shaded) rectangles.

have only the lower and upper half of the TCA cycle, respectively, although a missing enzyme in *H. pylori* still needs be identified to make the pathway continuous.

The PATHWAY database can be retrieved most conveniently from the top menu of the KEGG table of contents page, metabolic pathways and regulatory pathways that are categorized by the
Figure 2. The correlation of a physical unit and a functional unit of genes. (a) Starting from the KEGG genome map for *Escherichia coli*, (b) a chromosomal region is selected in the zoom-up window. (c) By clicking on the 'Pathway' button, the user can examine if the biochemical pathway can be formed from a set of genes in the zoom-up window.

The GENES database is maintained as follows. First the information of all genes in an organism is automatically generated from the complete genomes section of the GenBank database (10). Then the EC number assignment is performed by GFIT (8) and other programs with manual verification efforts. The gene function annotations are continuously re-evaluated according to the KEGG/PATHWAY database and by comparing with SWISS-PROT (12) and other databases.

**Genome maps and comparative genome maps**

The genome map represents a one-dimensional network of genes that are physically located in a circular or linear genome. The gene order turns out to be extremely valuable information in functional annotation, especially for bacteria and archaea. As shown in Figure 2, the genome map is linked to the pathway map by the Pathway button. The user can check if a physical unit of closely located genes (e.g., genes in an operon) would form a functional unit of related proteins that appear at close positions in the hierarchical classification of Table 2. Alternatively, the PATHWAY database can be searched by the DBGET/LinkDB system.
Figure 3. The comparative genome map in KEGG. (a) An overview of comparing Mycoplasma genitalium (horizontal) and Mycoplasma pneumoniae (vertical) genomes is shown where each dot represents significant amino acid sequence similarity of two genes. (b) The zoom-up window for the boxed area in (a) is shown where each gene name can be identified and pre-calculated homologous gene strings can optionally be displayed in color.

By clicking on the List button in the genome map, the user can also invoke a sequence similarity search to see if a stretch of genes would match a functional unit in the pathway.

The co-linearity of genes between two genomes is quite useful for identification of clusters of orthologous genes. KEGG provides the comparative genome map for identification of such clusters and for functional annotation of newly sequenced genomes (Fig. 3). The comparative genome map is analogous to the dot matrix for comparing two nucleotide or amino acid sequences. Here, a dot represents significant sequence similarity between two genes at the amino acid sequence level. To present this map, the sequence similarity scores are pre-computed by SSEARCH for all pairs of organisms and homologous gene clusters are pre-identified so that they can be optionally highlighted in the map.

**Ortholog group tables**

The ortholog group table is a summary table that represents functional correlations in the pathway, physical (positional) correlations in the genome, and evolutionary (sequence similarity) correlations among species. Currently there are about 4800 genes that belong to about 140 groups of functional units. The ortholog group table is most useful as a reference data set for functional annotations. KEGG provides a computational tool to search against the ortholog group tables for sequence similarity of a set of query sequences (see below).

A case in point is the annotation of ABC transporters. A typical ABC transporter consists of a substrate-binding protein, two membrane proteins, and two ATP-binding proteins. The genes for these molecular components usually form an operon, and there are many paralogous genes that are responsible for the transport of different substances. If a set of query sequences that are located at physically close positions in the genome match against all components, it is a good indication that the transporter unit is correctly reconstructed. Since the KEGG ortholog group table for ABC transporters is functionally categorized (9), the pattern of matches and their similarity scores can be used to deduce substrate specificity.

**Molecular catalogs and LIGAND database**

The KEGG molecular catalogs are intended to provide functional and structural classifications of proteins, RNAs, other biological macromolecules, chemical substances and molecular assemblies. However, the organization is still rudimentary except for enzymes. The information of chemical compounds, enzyme molecules, and enzymatic and non-enzymatic reactions is stored in the LIGAND database as described in the accompanying paper (5). Based on this database KEGG provides several molecular catalogs for classifications of enzymes and a preliminary classification of chemical compounds.

**FUNCTIONAL GENOMICS**

**Pathway reconstruction with reference maps**

The KEGG reference maps for metabolic pathways represent biochemical knowledge containing all chemically identified reaction pathways. The constraint of the genome, i.e., a list of enzymes encoded in the genome, will reconstruct organism-specific pathways, which are represented by coloring of boxes in the KEGG pathway maps (Fig. 1). Furthermore, the constraint of operons will often predict functional units or conserved pathway motifs, which are represented by additional coloring in the KEGG pathway maps (Fig. 2). The operon information probably reflects a regulatory unit of transcription. Thus, it is easy to see how the information of gene expression profiles can be used as still another constraint against the KEGG reference pathway maps. In fact, KEGG provides a tool to color the pathway maps in order to visualize, for example, the microarray patterns of gene expression profiles.

Table 3 shows the list of currently available tools for search and analysis of KEGG pathway maps and genome maps. The user-interfaces for these tools can be accessed from the KEGG network.
Pathway reconstruction from binary relations

It is often the case that no homology can be found when searching against the KEGG reference pathways, which suggests that the current biochemical knowledge is not sufficient to predict a pathway. If there is a missing portion in the reconstructed metabolic pathway, KEGG provides a tool to predict alternative paths with alternative enzymes. The tool actually computes all possible reaction pathways between two compounds from a set of substrate–product relations, i.e., from a set of enzymes (5): http://www.genome.ad.jp/kegg-bin/mk_pathcomp_html

This tool also has a feature called query relaxation to incorporate grouping or hierarchy of relations. Whenever any member of the group is identified in the genome by sequence similarity (e.g., an enzyme in the same hierarchy of EC numbers), the entire group is incorporated for computation (e.g., to represent wider substrate specificity). This effectively increases the number of substrate–product relations and expands possible reaction pathways.

This type of computation, which we call pathway reconstruction from binary relations, can be performed in a more general way. A binary relation can be a substrate–product relation in metabolic pathways, a gene–gene interaction observed in gene expression profiles, or a protein–protein interaction observed by yeast two hybrid system experiments. In practice, the reconstruction from binary relations works better when the problem size is not large. For example, if most of the pathway is reconstructed from the reference but there still remain a few missing enzymes, then the computation with binary relations may fill the missing links of fragmented pathways. In addition to the substrate–product binary relations, KEGG will provide tools to integrate and compute different types of binary relations. Perhaps this is the most challenging area of computational problems that have become accessible by the KEGG project.

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