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The Use of Weighted Graphs for Large-Scale Genome Analysis

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Introduction

Biology is undergoing a revolution due to the remarkable increase in the availability of DNA sequence data. This data is replete with biological knowledge. An increasingly important scientific challenge is therefore how best to extract this knowledge. This depends on the development of better data-structures and algorithms.

One area that has been transformed by the sequencing revolution is the area of prokaryotic metabolism. The complete genome sequences of over a thousand prokaryotic species are now known. These have been used to infer the complement of enzymes: taxonomic (these summarize phylogenetic importance), isoenzymatic (these summarize enzymatic variety/redundancy), and sequence-similarity (these summarize sequence conservation); and we applied these types of weighted graph to survey prokaryotic metabolism. To demonstrate the utility of this approach we have compared and contrasted the large-scale evolution of metabolism in Archea and Eubacteria. Our results provide evidence for limits to the contingency of evolution.

Abstract

There is an acute need for better tools to extract knowledge from the growing flood of sequence data. For example, thousands of complete genomes have been sequenced, and their metabolic networks inferred. Such data should enable a better understanding of evolution. However, most existing network analysis methods are based on pair-wise comparisons, and these do not scale to thousands of genomes. Here we propose the use of weighted graphs as a data structure to enable large-scale phylogenetic analysis of networks. We have developed three types of weighted graph for enzymes: taxonomic (these summarize phylogenetic importance), isoenzymatic (these summarize enzymatic variety/redundancy), and sequence-similarity (these summarize sequence conservation); and we applied these types of weighted graph to survey prokaryotic metabolism. To demonstrate the utility of this approach we have compared and contrasted the large-scale evolution of metabolism in Archea and Eubacteria. Our results provide evidence for limits to the contingency of evolution.

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Surveying prokaryotic metabolism

A number of previous studies have surveyed the large-scale evolution of metabolism. Yamada & Bork [5] surveyed metabolic (and protein-protein) interactions using a graph-theory framework. In our view the most interesting surveys are those of Peregrin-Alvarez et al. [6] and Freilich et al. [7]. Peregrin-Alvarez et al. compared the metabolic graph of E. Coli with the genomic evidence known at the time. The main emphasis of the Freilich...
et al. paper is on comparisons with mammalian enzymes, compared to our work they also analysed an order of magnitude fewer genomes, and did not attempt to sample uniformly. More recently Kreimer et al. [8] examined \( \approx 300 \) prokaryotic genomes using graph theory to estimate the modularity of metabolism. In an interesting application of large-scale analysis Borenstein et al. [9] estimated the environmental requirements of prokaryotic species.

**Three types of weighted graphs for surveying metabolism**

We propose three different ways to use the instantiations of the super-metabolic graph to add weights to the nodes (enzymes) to form weighted graphs for the large-scale analysis of metabolic pathways.

The first approach is to define the enzyme weights to be the proportion of genome instantiations where a gene for the enzyme is found. We call this type of weighted graph taxonomic. Such graphs summarize the phylogenetic importance of enzymes: enzymes with high weights occur in many species, and those with low weights rarely occur.

The second approach is to define the enzyme weights to be the average number of protein sequences of that enzyme in a genome. We call these weighted graphs isoenzymatic. Such graphs summarize the enzymatic variety/redundancy of metabolism: a high number indicates more isoenzymes.

The third approach is to define the enzyme weights to be the average sequence similarity of that enzyme in the genome instantiations. These are termed sequence similarity weighted graphs. Such graphs summarize the sequence conservation of metabolism: a high number indicates high sequence conservation.

To the best of our knowledge weighted graphs have not been used in any of these ways before.

**Comparing the evolution of metabolism in Eubacteria and Archaea**

We wished to test the utility of our weighted graphs for large-scale metabolic analysis. For this we selected the problem of investigating the diversity of prokaryotic metabolic pathways. Specifically we investigated the oldest and most fundamental split in the evolution of life, that between the two prokaryotic domains: Archaea and the Eubacteria [10]. Although recognition of the importance of this division came late [10], it is now clear that it is the deepest known phylogenetic division, and probably occurred 2–3 Billion years ago. Our idea is to use the newly available data on the biodiversity of metabolic pathways to investigate how different pathways have evolved since the divergence of the Archaea and the Eubacteria.

We recognize that because of gene transfer prokaryotic species/enzyme evolution does not have a pure tree topology (it is a directed acyclic graph). Genes have jumped between species, and across the Archaea Eubacteria divide. However, we hypothesize that this effect is insufficient to obscure the main signal from evolutionary descent.

**Sampling genomes**

Any conclusions that we draw regarding prokaryotic metabolism should be generally true for prokaryotic genomes. However, only a limited number of genomes have been sequenced. If these sequenced genomes were an unbiased sample from all existing sequenced and non-sequenced prokaryotic genomes, then one could argue that the sample is representative of the whole. But unfortunately this is not the case. (1) The sequenced genomes are very biased towards prokaryotic groups that are of special interest to humans (e.g., pathogens), and also towards groups that are easy to cultivate in the lab. (2) When comparing Eubacteria and Archaea there is also the problem that Eubacteria genomes outnumber Archaea by an order of magnitude. It is unclear how much this imbalance is due to the fact that Eubacteria have been studied more (for example, because Eubacteria cause diseases while Archaea do not), and how much it is due to there being more species of Eubacteria. These biases mean that we had to be careful how we used the sequenced genomes. We therefore decided to sample genomes uniformly across evolutionary space. By this we mean that if you envisage prokaryotic species as leaves of the tree of life with the branches evolutionary distance, then we will sample leaves uniformly distant from each other.
Materials and Methods

Data preparation

To generate our super-metabolic graph we selected the 192 pathways that occur in prokaryota from Kyoto Encyclopedia of Genes and Genomes (KEGG) [11] (http://www.genome.jp/kegg/ http://www.genome.jp/kegg/) (Release 59.0, July 1, 2011). This produced a graph with 2,365 enzymes with complete EC codes, and 43,627 metabolite edges (N.B. this is not the cardinality of the set of metabolites.).

We selected all prokaryotic (Archaea and Eubacteria) species with complete genomes from this release of KEGG. This gave 108 Archaea species and 1,287 Eubacteria species.

The assignment of protein function to these genes was taken directly from KEGG. It is clear that the functions of most genes from most genomes is not based on direct experimental evidence, but rather on inferred conservation of function with homology — a form of abductive reasoning [12]. Such inferences, like all abductions, are prone to error and must be treated with caution. However, these functional assignments are generally based on reasonably close homology, and are generally trusted. If these predictions were systematically wrong this could lead to bias in our results.

To sample genomes we first applied CD-HIT (http://weizhong-lab.ucsd.edu/cd-hit/) to cluster species based on their 16S ribosomal RNA sequences similarities at 0.8 level in each domain. We obtained 15 clusters of Archaea and 114 clusters of Eubacteria species. The different number of clusters reflects the difference in sampling (and possibly a difference in genomic diversity). To fairly compare sampling from the two domains, we sampled the same number of genomes from both domains. To generate the sampling datasets: for Archaea we randomly chose one species from each cluster; for Eubacteria we first randomly chose 15 clusters, then from each cluster randomly chose one species. We repeated the sampling process 100 times uniformly from both domains to provide 200 datasets each containing 15 genomes. We argue that this procedure produces datasets sampled uniformly across evolutionary space.

Our data is available free online at: http://www.cs.helsinki.fi/research/discovery/data/plosone2014/

Weighted metabolic network construction

In a metabolic network $G=(V,E)$, nodes correspond to enzymes: $V = \{ec_1, ec_2, \ldots, ec_n\}$. Two nodes (enzymes) $ec_i, ec_j \in V$ are connected, that is $(ec_i, ec_j) \in E$, if the reactions which they catalyse share compounds. For example, consider the following two reactions where $C_i$ is a compound: (1) $C_i + C_j \leftrightarrow C_k$, and (2) $C_d + C_e \Rightarrow C_d + C_e$. If the reactions share at least one compound, that is if $\{C_i, C_j, C_k\} \cap \{C_d, C_e\} \neq \emptyset$, then $ec_i$ and $ec_j$ are connected by an edge. We here only take into account compounds whose entries begin with “C” in KEGG, and also remove uncommon-cofactor-S1.pdf common cofactors (Table S1 (common cofactors)) taken from the article [13].

For taxonomic graphs let $H$ represent a set of selected genomes from one domain, $h_i$ represent a genome in $H$, and $c(h_i, ec)$ be a function showing whether $h_i$ contains enzyme $ec_i$. If $h_i$ contains $ec_i$, then $c(h_i, ec_i) = 1$, otherwise $c(h_i, ec_i) = 0$. The taxonomic weight of an enzyme $ec_i$ is $I_{tax}(ec_i) = \sum_{h_i \in H} c(h_i, ec_i)$. The range of $I_{tax}(ec_i)$ is $[0,1]$. A high taxonomic weight implies the enzyme ubiquitously exists in the domain.

For isoenzymatic graphs, the average isoenzymatic weight, denoted by $I_{iso}$, illustrates the average number of different protein sequences it may present. Let $n(h_i, ec_i)$ represent the number of different forms of protein sequences of an enzyme $ec_i$ in the genome $h_i$. The isoenzymatic weight of an enzyme $ec_i$ is

$$I_{iso}(ec_i) = \frac{\sum_{h_i \in H} n(h_i, ec_i)}{|H|},$$

The range of $I_{iso}(ec_i)$ is $[0, \infty]$. A high weight indicates a high redundancy of the enzyme.

For sequence similarity graphs we computed the average similarity of the enzyme’s protein sequences, represented as $S_{seq}(ec_i)$, measuring how conserved an enzyme’s protein sequence is during evolution. For each enzyme we first randomly picked one protein sequence of the enzyme from each selected genome, on the condition that the genome contains such enzyme. Let seq$_h$ be the protein sequence of enzyme $ec_i$ in the genome $h_i$. We then used the Laign [14] program of the FASTA package [15] with default parameters to compute the similarities of any pair of the selected protein sequences, denoted by $S(seq_{h_i}, seq_{h_k})$. We next calculated the mean value of the pairwise similarities. The sequence similarity weight of an enzyme $ec_i$ is

$$I_{seq}(ec_i) = \frac{\sum_{h_i \in H} \sum_{h_k \in H} S(seq_{h_i}, seq_{h_k})}{(\sum_{h_i \in H} c(h_i, ec_i))(\sum_{h_k \in H} c(h_k, ec_i) - 1)}$$

The range of $I_{seq}(ec_i)$ is $[0, 1]$. A high sequence-similarity weight shows the enzyme’s protein sequence conserved well.

For notational convenience, we write $I()$ to represent either kind of weighting when we are not discussing a specific weighting.

Weighted Graphs for Large-Scale Genome Analysis

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Inter and Intra domain correlation of weighted graphs

We wished to investigate the correlation between weights in the metabolic graphs. We first deleted non-informative regions of the graph: 1,112 enzymes have zero taxonomic weight in all 200 sampled datasets. (702 enzymes do not have taxonomy information in KEGG, and 347 enzymes only exist in Eubacteria kingdom, and the rest 63 enzymes exist in a tiny number of prokaryotic species.) For intra-domain correlations we included all enzymes that exist in at least two genomes in the selected data set. For the inter-domain correlations we examined all enzymes that have weights in either kingdom. We used Spearman’s rank correlation to calculate correlations as it is robust. The correlations between datasets were calculated by using cor.test() function in the R program, and values of $p < 0.05$ were considered to be significant.

We performed permutation tests to systematically examine the statistical significance of correlation between the weighted graphs we formed. In our setting the topology of the super-metabolic graphs in both domains is the same, and weights are only associated with nodes. Thus a rearrangement of the weights on the nodes produces a graph with the same topology and distribution of weights, but with no biological information — a random graph. We used these random graphs for the permutation tests. Let $N$ be the number of weighted enzymes in the graph. We randomly generated $10^5$ random graphs as a sample of the $N!$ possible permutations. For illustration, let $X$ and $Y$ be two groups of enzyme weights. We first applied sample() function in the R program to rearrange the values of one group of data $X$ without replacement, denoted by $X^*$, and then recomputed Spearman’s
rank correlation coefficient between the permuted group $X'$ and the other group $Y$. We repeated this step $10^5$ times, and totally obtained $10^5$ random correlation coefficient values. For the one-tailed test we calculated the ratio of the values in the permutational distribution of the statistic that are equal to or larger than the original correlation coefficient between $X$ and $Y$. For ratios smaller than 0.05 we considered the null hypothesis to not be consistent with the observations.

Specific or Ubiquitous enzymes across domains

We further analyzed the enzyme weights to identify whether the enzyme is specific or ubiquitous to a domain. An enzyme is regarded to be specific to a domain if its mean weight is above a specified high threshold in one domain, and is below a specified low threshold in the other domain. Conversely an enzyme is regarded as ubiquitous (to both domains) if its mean weights are above a specified high threshold in both domains.

Let $\tau$ be the high threshold. For taxonomic weights and sequence-similarity weights, since both ranges are between 0 and 1, let $1 - \tau$ be the low threshold. When $\tau$ gets close to 1, then an enzyme is more specific to one domain. We set $\tau$ to be 0.667. For isoenzymatic weights, as the range is $[0, +\infty)$, $\tau$ was set to be 1.

Metabolic pathways

Traditionally sub-parts of metabolism have been classified into different pathways, and these pathways have proven useful in analysis. We therefore used pathway information from KEGG. Our construction of a weighted metabolic pathway $P_i = (V_i,E_i)$ is analogous to the construction of the metabolic network. Node set $V_i$ is the set of enzymes that are needed in the pathway $P_i$. Two nodes (enzymes) are connected if the reactions which they catalyze are in the pathway $P_i$, and also share compounds.

Because of limited availability of enzyme taxonomic data, nearly half of enzymes have zero weight in all sampled datasets. For analysis, we selected pathways that averagely contain at least 10 weighted enzymes in both domains. To examine the importance of each selected pathway to a domain, we calculated the ratio of specific (or ubiquitous) enzymes the pathway contains to the other selected pathway to a domain, we calculated the ratio of the values in the permutational distribution of the statistic that are equal to or larger than the original correlation coefficient between $X$ and $Y$. For ratios smaller than 0.05 we considered the null hypothesis not to be consistent with the observations.

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Weighted graph compression

We further analyzed the weighted metabolic graphs using graph compression techniques. The model we applied is the extended version of the work, proposed by Toivonen et al [16], to take node weights into account during compression. Nodes with similar neighbors are grouped into super-nodes, and their edges grouped into super-edges. The idea is to compress an enzyme weighted metabolic network into a smaller one where the information related to enzymes with high weights is retained. We hypothesized that comparing compression across and between pathways would be informative for understanding the biodiversity and evolution of prokaryote metabolisms.

A graph is compressed iteratively through executing a series of operations. There are two basic operations: individual edge removal and node-pair merge. In the edge deletion operation, a single edge is removed, and if the removal leaves an edge endpoint isolated then the endpoint is removed as well. In the merge operation, a pair of nodes are merged into a new super-node, and the new super-node links with the neighbors of the merged nodes, and the weights on the super-edges re-assigned. Whether the new super-node links with their neighbors or not depends on whether the omission of the super-edge produces smaller error with respect to the extra saved space. In each iteration the effect of two operations are computed, and the one that produces a more compressed graph with smaller error is executed. An illustrated example is in Figure 2. Nodes $EC_i$ and $EC_j$ are merged into a new super-node. The new super-node only links with two previous neighbors $EC_k$ and $EC_m$, and the node $EC_k$ is deleted.

Compression may cause four types of error: a) a superedge may represent edges that do not exist in the original graph; b) edges in the original graph may not be represented by any superedge; c) nodes in the original graph may not be preserved in the compressed graph; and d) edge weights may be changed. The quality of the compressed graph is measured by computing the distance between the original graph and the decompressed graph of the compressed graph with respect to node weights, that is,

$$\text{dist}(G, \text{dec}(S)) = \sqrt{\sum_{[u,v] \in E_G \times V_G} \left( I(u)I(v)(w_G(u,v) - w_{\text{dec}}(S)(u,v))^2 \right)}.$$  (1)

Here $\text{dec}(S)$ is the decompressed graph of the compressed graph $S$, $I(u)$ represents the weight on the node $u$, and $w(u,v)$ represents the weight on the edge between $u$ and $v$. The decompressed graph $\text{dec}(S)$ is a graph where nodes are all node identities that are inside the super-nodes in the compressed graph $S$, and edges link nodes if there are super-edges between the corresponding super-nodes. The weight of an edge equals the weight of the corresponding super-edge.

The weights on the superedges are set to minimize the distance between the original and the compressed graphs. However, the minimization problem is computational hard. Since the distance metric (Equation 1) satisfies the triangle inequality, the distance between the original and the compressed graphs is upper bounded by the sum of the distance between the increasingly compressed graphs. We then propose a solution to minimize the upper bound of the distance. The weights on the supernodes therefore are set to minimize the distance between two sequential compressed graphs. Assume that (super)nodes $u$ and $v$ of graph $S_{i-1}$ are compressed into supernode $z$ in the resulting graph $S_i$, and $x_j$ is one of $u$’s and $v$’s neighbors. The weight of the superedge $\{z,x_j\}$ then is

$$w_z(z,x_j) = \frac{I(u)w_{S_{i-1}}(u,x_j) + I(v)w_{S_{i-1}}(v,x_j)}{I(u) + I(v)}.$$  (2)

![Figure 2](https://example.com/figure2.png)

**Figure 2.** A example of weighted graph compression based on node weights. Node weights are in the parenthesis.
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The compression ratio $cr$ measures how much smaller the compressed graph is with respect to the original graph, which is defined as the cardinalities of nodes and edges, that is,

$$cr(S) = \frac{|V_S| + |E_S|}{|V_G| + |E_G|}$$

(3)

where $V_S$ and $E_S$ represent the number of super-nodes and super-edges in the compressed graph $S$, and $V_G$ and $E_G$ represent the number of nodes and edges in the original graph $G$. The values of $cr$ are in the range from 0 to 1. Specially, when $cr$ gets close to 0, the compressed graph becomes more abstract.

The goal of weighted graph compression is to produce a compressed graph $S$ of a given weighted graph $G$ at a specified compression ratio $cr$, such that the distance between the original and compressed graph with respect to the node weights $\text{dist}(G, \text{dec}(S))$ is minimized.

To investigate the conservation of evolution, we computed the average enzyme weights in the compressed graphs with respect to different compression ratios, and the average enzyme weights in the compressed part that exist in both Archaea and Eubacteria domains. We also calculated the similarity between the original and the compressed pathways. Suppose for the pathway $P_i$, after compression, the number of removed nodes are $|V'_i|$, and the number of changed edges are $|E'_i|$ (including the number of added and deleted edges). The similarity between the original and compressed pathway is $1 - \frac{|V'_i| + |E'_i|}{|V_i| + |E_i|}$. A large value indicates a high similarity between the original and the compressed pathway.

**Results**

**The example of glycolysis**

To illustrate the result of processing of the sampled metabolic networks into weighted graphs, Figures 3–5 show the result of forming the three types of weighted graph for a section of the glycolysis pathway for Eubacteria and Archaea. In order to better illustrate the differences of weights intuitively we have adjusted the thickness of nodes according to their node weights. We selected glycolysis because it is the proto-typical pathway, and because it is found in most prokaryota. Yet even in glycolysis large differences in weight are observed.

![Figure 3. Taxonomic weight in a section of glycolysis.](image1)

The weights for Archaea are in red, the weights for Eubacteria are in blue.

![Figure 4. Sequence similarity weight in a section of glycolysis.](image2)

The weights for Archaea are in red, the weights for Eubacteria are in blue.

The taxonomic weights are in Figure 3. From the figure it is clear that some enzymes are ubiquitous, i.e. found in most species of both Eubacteria and Archaea, e.g. pyruvate kinase (EC 2.7.1.40); while other enzymes occur more frequently in one domain or the other, e.g. pyruvate synthase (EC 1.2.7.1) occurs much more frequently in Archaea than Eubacteria; other enzymes are uncommon in both domains, e.g. PEP carboxylase (EC 4.1.1.32).

The sequence similarity weights are in Figure 4. Here again there are large differences in weights across the pathway and between Eubacteria and Archaea. Enolase (EC 4.2.1.11) has relatively high sequence similarity in both Eubacteria and Archaea, while Pyruvate synthase (EC 1.2.7.1) has relatively low sequence similarity. In contrast PEP carboxylase (EC 4.1.1.32) has a high sequence similarity in Archaea and low similarity in Eubacteria. Such differences do not seem to be explainable by technical problems with alignment or sampling.

The isoenzymatic weights are in Figure 5. There are sizable differences in this type of weights across the pathway and between...
**Eubacteria and Archaea.** PEP carboxylase (EC 4.1.1.32) has the lowest isoenzymatic weights in both **Eubacteria** and **Archaea**, while Pyruvate synthase (EC 1.2.7.1) has the highest weights. The most notable difference between **Eubacteria** and **Archaea** is Acetate—CoA ligase (EC 6.2.1.1).

**The relationship between the different types of enzyme weights**

We investigated the correlation relationship between the different types of enzyme weights. In both **Archaea** and **Eubacteria** there is a statistically significant and moderate negative correlation between sequence-similarity and isoenzymatic weights (Figure 6). The average correlation coefficient values are $-0.41$ and $-0.39$ respectively. The average P values (of 100 tests) for these significance values are $4.18\times10^{-16}$ and $4.31\times10^{-15}$. The average permutation-based P values (Figure 7) in both **Archaea** and **Eubacteria** are 0, which show that there is a highly significant relationship between sequence-similarity and isoenzymatic weights in both domains. This means that enzymes where the sequences have low sequence similarities are more likely to have a greater number of isoenzymes, this seems intuitively reasonable as isoenzymes could enable greater sequence divergence.

In **Archaea** there is a weak but statistically significant positive correlation (Correlation coefficient $= 0.2$, P value $= 0.0013$) between taxonomic and isoenzymatic weights (Figure 8). This suggests that more common enzymes are slightly more likely to have isoenzymes. This also seems reasonable, as the more common enzymes are likely to be more important and need greater control, but it is unclear why the correlation is so low. In **Eubacteria** the correlation between taxonomic and isoenzymatic weights is negligible (Correlation coefficient $= 0.12$, P value $= 0.035$). Their significance test results (Figure 9) indicate that a clear relationship (average permutation-based P value $= 0.0007$) between taxonomic and isoenzymatic weights exists in **Archaea**, whereas the relationship (average permutation-based P value $= 0.017$) is not quite obvious in **Eubacteria**. It is not clear why there is a difference between domains.

**Figure 6.** Correlation between sequence similarity and isoenzymatic weights.

**Figure 7.** The strength of relationship between sequence similarity and isoenzymatic weights as estimated by permutation tests.

There is a weak but statistically significant negative correlation (Correlation coefficient $= -0.16$, P value $= 0.008$) between taxonomic and sequence-similarity weights in **Archaea** (Figure 10). This means that more common enzymes are slightly more likely to have less diverged sequences. This is also reasonable, as they are likely to be under more constraints. The corresponding negligible negative correlation (Correlation coefficient $= -0.07$, P value $= 0.18$) in **Eubacteria** is not significant. The significance tests (Figure 11, average permutation-based P values of $0.0038$ and $0.09$ respectively) give the same outcomes. Again it is not clear why there is a difference between domains.

**Figure 8.** Correlation between taxonomic and isoenzymatic weights.

**Figure 9.** Correlation coefficient between sequence–similarity and isoenzymatic weights.
It is noteworthy that in all cases the correlation is stronger in Archaea than Eubacteria, the reasons for this are unclear.

Comparison of the Archaea and Eubacteria domains

Summary statistics. We calculated the mean weights for all enzymes, and for all enzymes with non-zero weights for the three weightings (Table 1). The mean weights for Archaea are set in roman, and mean weights for Eubacteria are in italics. The mean weights for a specific type of weighting are similar across domains.

We then used Spearman’s rank correlation to compute correlation of the weights as this gives a broad indication how similar the metabolic graphs are across domains (Figure 12). The correlation of 0.48 in taxonomic weights between domains (i.e. Archaea and Eubacteria) is highly significant (P value = 1.468e-28). The correlation of 0.19 in isoenzymatic weights between domains is also significant but at a much lower level (P value = 0.0055). There is however no significant correlation (Correlation coefficient = −0.08) in sequence-similarity weights between domains in Archaea and Eubacteria (P value = 0.124). The significance test results (Figure 13) provide evidence that average permutation-based P values are 0 and 0.0027 respectively, so a clear relationship exists between domains in both taxonomic weights and isoenzymatic weights, but the relationship (average permutation-based P value = 0.067) between domains is not quite obvious in sequence-similarity weights.

Individual enzymes. In order to compare the similarities and differences between Archaea and Eubacteria in more detail we identified specific and ubiquitous enzymes across domains.

In the case of taxonomic weights, for the 1,253 enzymes that have weights in either kingdom, 73 enzymes are classed as domain specific, and 120 as ubiquitous. For example valine-tRNA ligase (EC 6.1.1.9) occurs in every genome examined. (Valine is in some sense the most average of amino acids.) The commonest enzyme that isn’t associated directly with macromolecule processing is ribose-phosphate diphosphokinase (EC 2.7.6.1). Other extremely ubiquitous enzymes are phosphoglycerate kinase (EC 2.7.2.3) and glycine hydroxymethyl-transferase (EC 2.1.2.1).

Some enzymes occur much more frequently in one domain than the other. Some of these enzymes are expected, for example UDP-N-acetylmuramate dehydrogenase (EC 1.1.1.158) is involved in peptoglycan metabolism and is much more common in Eubacteria. However, others are more surprising, such as the comparatively frequent occurrence of L-lactate dehydrogenase (EC 1.1.1.27) in Eubacteria, but not Archaea. Examples of enzymes that occur much more frequently in Archaea are mevalonate kinase (EC 2.7.1.36), and nicotinamide-nucleotide adenylyltransferase (EC 2.7.7.1).

Pathways. To examine the importance of each pathway to the domain we calculated the ratio of special enzymes (e.g. specific or ubiquitous enzymes) to the size of enzymes with non-zero weights in each pathway (Table 2). The pathways that contain the
Table 1. Mean enzyme weights in different types of weights. The weights for Archaea are set in roman, the weights for Eubacteria are in italics.

<table>
<thead>
<tr>
<th>Weight Type</th>
<th>All enzymes</th>
<th>Non-zero enzymes</th>
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<tbody>
<tr>
<td></td>
<td>Archaea</td>
<td>Eubacteria</td>
</tr>
<tr>
<td>Taxonomic weight $I_{tax}$</td>
<td>0.104</td>
<td>0.14</td>
</tr>
<tr>
<td>Isoenzymatic weight $I_{iso}$</td>
<td>0.257</td>
<td>0.367</td>
</tr>
<tr>
<td>Sequence-similarity weight $I_{seq}$</td>
<td>0.168</td>
<td>0.247</td>
</tr>
</tbody>
</table>

Figure 12. Correlation of weights between domains.
doi:10.1371/journal.pone.0089618.g012

Figure 13. The strength of relationship of weights between domains as estimated by permutation tests.
doi:10.1371/journal.pone.0089618.g013
highest ratio of ubiquitous enzymes are: aminoacyl-tRNA biosynthesis; valine, leucine, and isoleucine biosynthesis; and phenylalanine, tyrosine, and tryptophan biosynthesis. The origin of these pathways clearly predates the 
Eubacteria Archaea divide, and we may speculate they are perhaps the oldest of all pathways. The pathways that contain the highest ratio of specific enzymes are: terpenoid backbone biosynthesis, and pantothenate and Co-A biosynthesis. It is also noteworthy that both glycolysis/gluconeogenesis has quite a high ratio of specific enzymes.

Compression results on average enzyme weights. To further investigate the conservation of evolution we applied weighted graph compression to abstract the metabolic networks into smaller ones utilizing taxonomic enzyme weights. The idea of using compression is that with increased compression more and more nodes and edges with lower weights are removed; therefore the compressed graphs may also be informative about the common ancestor of 
Eubacteria and 
Archaea.

We calculated the average taxonomic enzyme weight in the compressed graphs as a function of compression ratio, and plotted the results of 
Archaea in the dashed-line in Figure 14, and the results of 
Eubacteria in Figure 15. As expected, the average enzyme weight increases when there is more compression, i.e., in a smaller compression ratio. As enzymes with relatively lower weights are removed in the compression process, the enzymes left in the compression graphs become more important to the kingdom. We next computed the average taxonomic weight of enzymes in
compressed part shared by both kingdoms. The results, plotted with solid lines in Figures 14 and 15, show that the average taxonomic weight of enzymes in the shared compressed part also becomes higher when there is more compression. More interestingly, the average weight in the shared part for a specific kingdom (e.g., Figure 15) is much higher than the weight in the whole compressed graphs.

**Compression results on pathways.** We further investigated how similar the compressed pathways are to the original ones. The correlations of compression results of pathways across domains are significant and very strong positive in three types of weights (Table 3).

We ranked pathways based on the descending similarity between the original and the compressed graphs (Table 4). The pathways in the top rank have high similar original structure in the compressed results. Since the average enzyme weight becomes higher when there is more compression (Figures 14 and 15), the pathways in the top rank probably contain more enzymes with high weights, and it also implies that parts of these pathways possibly exist in the ancestor of the domain.

**Discussion**

**Summary of prokaryotic metabolism**

The weighted graphs, in Figures 3–5, concisely and intuitively illustrate how different enzymes exist between the prokaryotic domains. According to Figure 12, for taxonomic weights, there is moderate correlation (0.48) between two domains. This implies that if an enzyme is common in species in one domain, it is likely to also be common in the other domain. In contrast the correlation (0.19) between the number of isoenzymes for a given enzyme is weak, and there is no correlation (0.08) in sequence-similarity weights.

By comparing different types of weighted graphs we can also get some useful understandings of the correlation among sequence-similarity weights, isoenzymatic weights and taxonomic weights. For example, when an enzyme is common in a domain it is more likely to have a higher number of isoenzymes (Figure 6).

Another useful analysis is about how many specific or ubiquitous enzymes each pathway contains (Table 2). If a pathway contains a high ratio of ubiquitous enzymes it is more likely that the pathway exists in the core metabolism of both domains. In contrast, a pathway contains a high ratio of specific enzymes implies that the pathway is more common in one domain.

**The utility of weighted graphs in analyzing the contingency of evolution**

One of the most fundamental questions in evolutionary biology is to what extent the paths that evolution has taken are stochastic, and to what extent they are determined by constraints imposed by the environment and biochemistry. Eminent evolutionary biologists have taken radically different views on this question of stochasticity. Stephen Jay Gould in many essays, and most notably in his book *Wonderful Life* [17], argued for contingency in evolution. For him evolutionary biology, in seeking to explain the past was a historical science, so if the process could somehow be run again then one would expect a radically different result. In contrast, Simon Conway Morris [18] has argued that the constraints on living organisms are such that it is likely that evolution would take broadly the same path and intelligent organisms such as humans are likely to evolve.

The central problem with investigating this question is that it is generally impossible to repeat the experiment — evolution. However, it is possible to get some understanding of the stochasticity of the problem by looking at cases where evolution started from the same starting points, i.e. the *Archaia* and *Eubacteria*.
Our most relevant results regarding this question are Figure 12 and Table 4. In Figure 12 the correlation of 0.48 between taxonomic weights in *Archaea* and *Eubacteria* is highly significant ($P$ value = $1.468 \times 10^{-28}$). This means that knowing the phylogenetic importance of an enzyme in one domain is informative about its importance in another domain. Our interpretation of this result is that as the process of evolution proceeded and new species of prokaryota were formed, evolution was constrained to use enzymes similarly in both domains. We argue that these results are evidence for limits to the contingency of evolution, however it is difficult to quantify the extent of contingency.

A similar but weaker pattern is seen for isoenzymatic weights. However, there is no correlation in the case of sequence-similarity weights. Our interpretation of this is that most sequence changes are neutral and not selective [19]. If this is the case then we would expect to observe little correlation between domains for this type of weight.

Figures 14 and 15 show, for a specific domain, the average taxonomic enzyme weight in the compressed part shared by both domains is higher than the weight in the whole compressed graph. This means the enzymes in the shared part have high weights in both domains. Our interpretation of this is that the shared enzymes are more common than average in both kingdoms. These

### Table 4. The rankings of pathways with respect to the descending similarity between the original and the compressed pathways (compression ratio is 0.09).

<table>
<thead>
<tr>
<th>Pathway Name</th>
<th>Taxonomy</th>
<th>Isoenzyme</th>
<th>Sequence-similarity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Archaea</td>
<td>Eubacteria</td>
<td>Archaea</td>
</tr>
<tr>
<td>Aminoacyl-tRNA biosynthesis</td>
<td>1</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Selenocompound metabolism</td>
<td>2</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>Citrate cycle (TCA cycle)</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Valine, leucine and isoleucine biosynthesis</td>
<td>4</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>Carbon fixation in photosynthetic organisms</td>
<td>5</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>One carbon pool by folate</td>
<td>6</td>
<td>8</td>
<td>9</td>
</tr>
<tr>
<td>Pentose phosphate pathway</td>
<td>7</td>
<td>5</td>
<td>15</td>
</tr>
<tr>
<td>Porphyrin and chlorophyll metabolism</td>
<td>8</td>
<td>9</td>
<td>6</td>
</tr>
<tr>
<td>Carbon fixation pathways in prokaryotes</td>
<td>9</td>
<td>3</td>
<td>11</td>
</tr>
<tr>
<td>Glycolysis/Gluconeogenesis</td>
<td>10</td>
<td>15</td>
<td>8</td>
</tr>
<tr>
<td>Alanine, aspartate and glutamate metabolism</td>
<td>11</td>
<td>11</td>
<td>13</td>
</tr>
<tr>
<td>Lysine biosynthesis</td>
<td>12</td>
<td>6</td>
<td>10</td>
</tr>
<tr>
<td>Methane metabolism</td>
<td>13</td>
<td>24</td>
<td>12</td>
</tr>
<tr>
<td>Phenylalanine, tyrosine and tryptophan biosynthesis</td>
<td>14</td>
<td>14</td>
<td>7</td>
</tr>
<tr>
<td>Pyrimidine metabolism</td>
<td>15</td>
<td>16</td>
<td>13</td>
</tr>
<tr>
<td>Glycine, serine and threonine metabolism</td>
<td>16</td>
<td>21</td>
<td>18</td>
</tr>
<tr>
<td>Pantothenate and CoA biosynthesis</td>
<td>17</td>
<td>17</td>
<td>18</td>
</tr>
<tr>
<td>Pyruvate metabolism</td>
<td>18</td>
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<td>28</td>
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<tr>
<td>Nitrogen metabolism</td>
<td>19</td>
<td>12</td>
<td>22</td>
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<tr>
<td>Terpenoid backbone biosynthesis</td>
<td>20</td>
<td>13</td>
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<tr>
<td>Butanoate metabolism</td>
<td>21</td>
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<td>Histidine metabolism</td>
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<td>Cysteine and methionine metabolism</td>
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<td>Arginine and proline metabolism</td>
<td>24</td>
<td>22</td>
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<tr>
<td>Amino sugar and nucleotide sugar metabolism</td>
<td>25</td>
<td>23</td>
<td>25</td>
</tr>
<tr>
<td>Propanoate metabolism</td>
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<td>Nicotinate and nicotinamide metabolism</td>
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<td>20</td>
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<td>Purine metabolism</td>
<td>28</td>
<td>18</td>
<td>19</td>
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<tr>
<td>Fructose and mannose metabolism</td>
<td>29</td>
<td>27</td>
<td>31</td>
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<tr>
<td>Valine, leucine and isoleucine degradation</td>
<td>30</td>
<td>29</td>
<td>30</td>
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<tr>
<td>Galactose metabolism</td>
<td>31</td>
<td>30</td>
<td>29</td>
</tr>
<tr>
<td>Glyoxylate and dicarboxylate metabolism</td>
<td>32</td>
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<td>32</td>
</tr>
<tr>
<td>Starch and sucrose metabolism</td>
<td>33</td>
<td>34</td>
<td>35</td>
</tr>
<tr>
<td>Pentose and glucuronate interconversions</td>
<td>34</td>
<td>33</td>
<td>34</td>
</tr>
<tr>
<td>Glutathione metabolism</td>
<td>35</td>
<td>32</td>
<td>35</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0089618.t004

Weighted Graphs for Large-Scale Genome Analysis
results provide the evidence of the conservation of evolution. Likewise, Table 4 lists the ordered pathways that are well preserved in the compressed graphs, which indicates that parts of these pathways are more likely to have been present in their common ancestral organism. The correlation of compression results of pathways between domains (Table 4) is statistically significant and very strong positive (Correlation coefficient = 0.91, P value = 3.91e−14) for taxonomic weights. This means the pathways that are important in one domain are also important in another domain. Again this is evidence of limits to the contingency of evolution.

Application of weighted graphs to other biological networks

We have demonstrated the use of weighted graphs as a way of efficiently analyzing large amounts of genomic information about metabolic networks. Similar weighted graphs could also be applied to other types of network: regulatory genetic, protein interaction, etc. The definitions of taxonomic and sequence similarity weighted graphs are directly applicable to other regulatory genetic and protein interaction graphs, and the definition of isoenzymatic weighted graphs could be adapted to be isofunctional. This would open up an interesting range of types of analysis for application to large numbers of genomes.

Supporting Information

Table S1 The list of removed compounds taken from the article “Using a Logical Model to Predict the Growth of Yeast,” authored by K. E. Whelan and R. D. King, published in BMC Bioinformatics in 2008.

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Author Contributions

Conceived and designed the experiments: RDK. Performed the experiments: FZ. Analyzed the data: FZ. Contributed reagents/materials/analysis tools: FZ HT. Wrote the paper: FZ RDK.

References