The percentage of bacterial genes on leading versus lagging strands is influenced by multiple balancing forces

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ABSTRACT

The majority of bacterial genes are located on the leading strand, and the percentage of such genes has a large variation across different bacteria. Although some explanations have been proposed, these are at most partial explanations as they cover only small percentages of the genes and do not even consider the ones biased toward the lagging strand. We have carried out a computational study on 725 bacterial genomes, aiming to elucidate other factors that may have influenced the strand location of genes in a bacterium. Our analyses suggest that (i) genes of some functional categories such as ribosome have higher preferences to be on the leading strands; (ii) genes of some functional categories such as transcription factor have higher preferences on the lagging strands; (iii) there is a balancing force that tends to keep genes from all moving to the leading and more efficient strand and (iv) the percentage of leading-strand genes in a bacterium can be accurately explained based on the numbers of genes in the functional categories outlined in (i) and (ii), genome size and gene density, indicating that these numbers implicitly contain the information about the percentage of genes on the leading versus lagging strand in a genome.

INTRODUCTION

It has been observed that the majority of bacterial genes tend to be located on the leading strand in a genome, and the percentage of such genes has a large variation across different bacteria, ranging from ~45% to ~90% (1,2). A number of studies have been carried out aiming to provide explanations for such observations. A key factor considered in these studies is the different mechanisms used by bacterial cells in replication of the leading and the lagging strands when cell replication and transcription occur simultaneously (3,4). Specifically, during chromosomal replication, deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) polymerases move in the same direction on the leading strand but in opposite directions on the lagging strand, creating the possibility of head-on collisions between the two polymerases during transcription of some genes on the lagging strand, hence making the lagging strand the less efficient one between the two (1,4). In an earlier study, Brewer (3) suggested that bacterial cells may be under a selection pressure to have highly expressed genes reside on the leading strand. Rocha and Danchin (5,6) recently argued that it is really the essentiality instead of the needed expression levels of genes that may have driven certain genes to the leading strand. Although this interpretation seems to be correct, it provides only a partial answer as essential genes account for only a small portion of the whole gene set encoded in a bacterial genome, e.g. ~10% in Escherichia coli (7,8) and ~10% in Bacillus subtilis (9). Price et al. (10) observed that longer operons tend to be on the leading strand and suggested that there may be a selection pressure to have such an arrangement to avoid interruptions during transcription of such operons. Furthermore, Rocha (6,11) observed that the presence/absence of the DNA polymerase PolC in a genome is highly correlated with bacterial genomes having at least 70% of their genes on the leading strand or not. Hu et al. (12) proposed that replication-associated purine asymmetry may also contribute to the strand bias in a genome. In addition, Lin et al. (13) found that the essential genes on the leading strand are enriched in only a few of sub-categories of clusters of orthologous groups (14). Although this analysis provided useful insights of

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functional preference of genes to the leading and lagging strand, a larger analysis involving more genes and organisms is needed to ensure the generality of the observation. More importantly, the general issues of why the majority of bacterial genes tend to be located on the leading strands and why the percentage of leading strand genes has such a large variation across different organisms remain largely unanswered.

We present in this study a computational analysis of all the sequenced bacterial genomes aiming to provide a more general explanation to the above two observations. Our key findings are (i) genes of different functional categories have different level of tendency to be on the leading strand; (ii) genes of some functional categories such as transcription factor have higher preferences to be on the lagging strands; (iii) there is at least one balancing force that keeps genes from all moving to the leading strand during evolution, i.e. a more balanced genome facilitates a higher gene density in a genome and (iv) the percentage of leading-strand genes for a bacterium can be accurately explained in terms of genes in some functional categories outlined in (i) to (ii), genome size and gene density. On the basis of these findings, we believe that the percentage of genes on the leading versus lagging strand in a genome is the result of two sets of balancing forces, one that tends to drive genes of certain functional categories to the leading strands to make the bacteria more efficient in their responses to environmental changes and one that tends to keep the genome as compact as possible to stay energetically efficient when replicating and maintaining the genome.

MATERIALS AND METHODS

Data

The 725 bacterial genome sequences along with their predicted genes and functional annotations were retrieved from the NCBI FTP site (ftp://ftp.ncbi.nih.gov/genomes/Bacteria/) as of 11 December 2010. The gene ontology (GO) annotations for these genomes were from the GOA Proteome Sets (v52) (15), and the GOslim definitions were downloaded from the Gene Ontology site (http://www.geneontology.org/GO_slims/goslim_generic.obo) (16). The microarray data for *E. coli* are downloaded from the M3D web site (http://m3d.bu.edu) (17).

High-level functional annotations of genes

GO (16) was used to define functional categories of gene products. Based on the GO annotation and GO hierarchy information, the Perl script map2slim (http://search.cpan.org/~emungall/go-perl/scripts/map2slim) was used to the bacterial genomes for assignment of GOslim-based functional categories.

Determination of genes on leading and lagging strands

To determine whether a gene is on the leading versus lagging strand of a genome, the origin and the terminus of replication are needed. The origin of replication for each of the 725 bacterial genomes was retrieved from the DoriC database (18), which has been widely used in the comparative genomics analysis (19–21) and the origin prediction for newly sequenced genomes (22–28). The terminus of replication is thus calculated as the location of origin of replication plus half of the chromosome length. With these two positions, the leading and lagging strands are determined for each half of the chromosome according to a well-known fact that the leading strand always has more genes than the lagging strand does (11). For each bacterium, only the major chromosome is considered, and plasmids are excluded in this study.

Preference of functional categories on different strands

Given a GO functional category, an index *x* is calculated using the following formula:

\[
x = \frac{n_0}{n_0 + n_1}
\]

where *n*0 is the number of leading-strand genes of this functional category, *n*1 is the number of lagging-strand genes of this category; *x* is calculated for all the GOslim functional categories, respectively, on the leading strand for all 725 genomes, so that for each category, there is a data set (A) of 725 values. In addition, the overall percentage of leading-strand genes (data set B) is obtained for each of the 725 genomes as well. For each GOslim functional category, a Wilcoxon rank sum test was performed to test whether the data sets A and B are from two distinct distributions. We also used the similar procedure to assess the preference of functional categories on the lagging strand. All the statistical analyses are conducted using the R statistical language (http://www.r-project.org).

Prediction of the percentage of leading-strand genes in a genome

Network training

A neural network model, with one hidden layer of 10 nodes, is used to predict the percentage of leading-strand genes in a genome using the total 57 inputs and then selected smaller numbers of 30, 25, 20, 15, 10 and 5 ones in this study. To reduce the possibility of the over-fitting problem, we used an early stopping technique, which divides the data into three subsets: training set used for computing gradient and updating the network weights and biases, validation set used for monitoring training process by its error rate and testing set used for assessing the neural network performance independently. When the network starts to over-fit the data, the error on the validation set begins to rise, and hence, the training process is stopped early. We used the default setting in the MATLAB Neural Network Toolbox, which arbitrarily divides the data into the three subsets, respectively: 507 (70%) for training set, 109 (15%) for validation set and 109 (15%) for testing set. The performance of a neural network is measured by mean squared error (MSE) and Pearson correlation score (*R*). The trained neural network
Variable selection
Out of the initial set of 57 input variables, we have conducted a variable selection process based on the idea of mean impact value (MIV) (29). Based on the ranks of the MIVs of the input variables, input variables with insignificant MIV will be eliminated from the neural network model. MIVs are calculated as follows: vary the value of each input variable by increasing and decreasing 10% for all samples and get two outputs. Then subtract one from the other and obtain the impact change value [impact value (IV)] of the output due to the changes of the input variable values. Then the MIV is obtained by averaging the IVs across all trained networks: \( \text{MIV} = \frac{\text{IV}}{n} \), where \( n \) is the times of network training.

RESULTS
Characteristics of genes on leading strands
We have analyzed 725 sequenced eubacterial genomes for which origins of replication and GO-based annotations are available in terms of the strand biases of their protein-encoding genes, and archaea were excluded from our analysis as they may have multiple origins of replication. Figure 1A shows the percentage distribution of leading-strand genes across all the 725 genomes, ranging from 45% to ~90%. This observation extends a previous observation made based on a few bacterial genomes. Across these 725 bacteria, the percentage of leading-strand genes does not show any correlation with genome sizes in terms of gene numbers (Supplementary Figure S1), whereas different phyla have substantially different averaged percentages of leading-strand genes (Supplementary Figure S2), which is consistent with a previous finding made on a smaller group of bacterial genomes (11,30).

We have also examined the relationship between leading strand bias and the growth rate for 104 of the 725 bacterial genomes, for which the doubling-time information is available (31). We found that bacterial genomes with high leading-strand bias (>70%) tend to have higher growth rates than those with low leading-strand bias (<70%) measured by the Wilcoxon rank sum test with \( P \) value: \( 1.9 \times 10^{-4} \), as shown in Figure 1B. This observation changes the previous conclusion that fast-growing bacteria have similar leading-strand bias to that of the slow-growing bacteria (11), which was made based on a substantially smaller number of genomes with known growth rates.

Functional categories whose genes have different preference to different strands
We have examined whether genes of different functional categories may have different level of preference to be on the leading versus the lagging strands across all bacteria. To do this, we checked 55 of the 127 GOslim functional categories (16) that have available gene assignments in at least 36 (5%) of the 725 genomes and have the number of genes with the median being between 5 and 500 (categories with >500 genes will be too general for our study) across all the genomes under consideration. For each of the 55 functional categories, we consider a functional category prefers a strand if genes in this category have a higher percentage than the average percentage of all genes on the strand across all genomes. The Wilcoxon rank sum test is used to assess the statistical significance of an observed preference measured using a \( P \) value. We found that 32 of the 55 categories prefer the leading strand with \( P \) value < 0.01, including genes related to ribosome, structural molecular activity, translation, DNA binding and cell cycle, with the detailed information presented in Table 1; and 11 categories prefer the lagging strand with \( P \) value < 0.01, including DNA-binding transcription factor activity, signal transducer activity and regulation of biological process, with details in Table 2. On average, 52% of the genes encoded in a bacterial genome are covered by the 43 (32+11) functional categories, and the detailed distribution of percentage across different bacterial genomes is given in Supplementary Table S1. Notably, transcription factor
Table 1. Preference of GO slim categories toward leading strands across 725 bacterial genomes

| GO branch | GO category                                      | Preference |
|-----------|--------------------------------------------------|------------|
| MF        | GO:0005198 structural molecule activity          | 7.28E-15   |
| MF        | GO:0003723 RNA binding                           | 1.15E-10   |
| MF        | GO:0008135 translation factor activity           | 4.67E-78   |
| MF        | GO:0005515 protein binding                       | 1.37E-32   |
| MF        | GO:0003774 motor activity                        | 2.05E-24   |
| MF        | GO:0000116 nucleotide binding                    | 5.93E-14   |
| MF        | GO:0003676 nucleic acid binding                  | 1.04E-13   |
| MF        | GO:0032354 enzyme regulator activity             | 1.97E-08   |
| MF        | GO:0016740 transferase activity                  | 9.62E-05   |
| BP        | GO:0006412 translation                           | 6.77E-118  |
| BP        | GO:0007049 cell cycle                            | 8.66E-72   |
| BP        | GO:0019538 protein metabolic process             | 7.64E-56   |
| BP        | GO:0015031 protein transport                     | 7.60E-34   |
| BP        | GO:0016043 cellular component organization       | 1.81E-30   |
| BP        | GO:0009695 response to external stimulus         | 1.89E-20   |
| BP        | GO:0007154 cell communication                     | 2.51E-18   |
| BP        | GO:0006919 generation of precursor metabolites   | 1.11E-17   |
|           | and energy                                       |            |
| BP        | GO:0005975 carbohydrate metabolic process        | 7.17E-16   |
| BP        | GO:0019748 secondary metabolic process           | 3.10E-15   |
| BP        | GO:0006259 lipid metabolic process               | 1.37E-11   |
| BP        | GO:0009565 catabolic process                     | 1.51E-06   |
| BP        | GO:0006519 cellular amino acid and derivative    | 2.41E-06   |
|           | metabolic process                                |            |
| BP        | GO:0006811 ion transport                         | 5.37E-05   |
| BP        | GO:0006850 response to stress                   | 3.26E-03   |
| CC        | GO:0005840 ribosome                             | 1.97E-13   |
| CC        | GO:0043226 organelle                            | 5.17E-11   |
| CC        | GO:0007377 cytoplasm                            | 6.29E-56   |
| CC        | GO:0005622 intracellular                        | 6.67E-41   |
| CC        | GO:0005694 chromosome                           | 1.11E-26   |
| CC        | GO:0043234 protein complex                      | 6.98E-26   |
| CC        | GO:0005618 cell wall                            | 5.31E-11   |
| CC        | GO:0008380 plasma membrane                      | 1.42E-06   |

The first column represents the three major GO categories: molecular function (MF), cellular component (CC) and biological process (BP).

Table 2. Preference of GO slim categories toward lagging strands across 725 bacterial genomes

| GO branch | GO category                                                      | Preference |
|-----------|------------------------------------------------------------------|------------|
| MF        | GO:0003700 sequence specific DNA binding                         | 3.09E-34   |
| MF        | GO:0016209 antioxidant activity                                  | 3.31E-11   |
| MF        | GO:0003677 DNA binding                                           | 3.41E-11   |
| MF        | GO:0004971 signal transducer activity                            | 2.10E-06   |
| MF        | GO:0004072 protein kinase activity                               | 9.34E-06   |
| MF        | GO:0008233 peptidase activity                                    | 1.08E-05   |
| BP        | GO:0007879 regulation of biological process                      | 3.20E-15   |
| BP        | GO:0019725 cellular homeostasis                                  | 3.08E-11   |
| BP        | GO:0006350 transcription                                         | 2.94E-09   |
| BP        | GO:0006464 protein modification process                          | 1.06E-06   |
| BP        | GO:0007165 signal transduction                                   | 6.31E-05   |

The first column represents the three major GO categories: molecular function (MF), cellular component (CC) and biological process (BP).

The analysis suggests that there might be a selection pressure for a bacterium to have a more compact genome (i.e. a shorter genome without losing genes), particularly in a complex environment. To check this hypothesis, we have examined the percentages of coding regions in the two groups of bacteria, one containing all bacteria with at least 70% of the genes on the leading strands and one containing all the other 725 bacteria and checked their relationship with the living styles of the bacteria. Our analysis revealed that (i) the bacteria in the second group (with lower strand bias) tend to have higher percentages of coding regions than those in the first group, with a P value 1.8 × 10^{-8} based on the Wilcoxon rank sum test, as shown in Figure 2A and (ii) this tendency is more significant for bacteria living in complex environments, with P values ranging from 0.25 to 2.8 × 10^{-9}, as shown in Figure 2B–F. One possible explanation is that there might be a selection pressure for bacteria living in nutrient-depleted environments to keep their genomes as compact as possible (without losing genes), and having a more balanced genome is one way to achieve this goal (a more balanced genome seems to allow a higher degree of overlap between regulatory regions of operons).

A balancing force: strand bias versus gene density

Our main hypothesis is that the percentage of leading-strand genes in a genome reflects the relationship between the key functionalities and the living environment of an organism. To check this hypothesis, we have examined the population of genes in each functional category encoded in each genome to see whether some of them can be used to predict the percentage of leading-strand genes.

We trained 10 times a neural network with 57 input nodes, one node for each of the 55 functional categories, one node for gene density and one node for the genome size; one hidden layer of 10 nodes and one output node, where gene density is calculated as the percentage of leading-strand genes.
of non-coding region length against chromosome length. We split the 725 genomes into three sets: 507 (70%) as the training set, 109 (15%) as the validation set and 109 (15%) as the testing set. At the end of the training, the neural network has the following average performance results on the three data sets: \( \text{MSE} = 0.0015 \) and \( R = 0.91 \) between the desired and predicted values on the training set; \( \text{MSE} = 0.0023 \) and \( R = 0.85 \) on the validation set and \( \text{MSE} = 0.0021 \) and \( R = 0.87 \) on the testing set. Figure 3 shows the performance of a trained neural network on the different data sets.

Using the variable selection procedure outlined in ‘Materials and Methods’ section, we have examined the IV of each input on the performance of each neural network trained by increasing or decreasing its value by 10% and used the averaged IV (MIV) for the 10 trained neural networks as a measure of the importance level of that variable. We have examined the performance of neural networks with smaller numbers of inputs with top MIV values: 30, 25, 20, 15, 10 and 5, as shown in Figure 4. Each network was trained three times. The networks with 25 inputs work best on the different data sets with the following average performance: \( \text{MSE} = 0.0018 \) and \( R = 0.89 \) on the training set; \( \text{MSE} = 0.0019 \) and \( R = 0.86 \) on the validation set and \( \text{MSE} = 0.0017 \) and \( R = 0.88 \) on the testing set. Specifically, these 25 inputs are listed in Table 3: cell cycle, iron transport, transport, response to stress, nucleobase, cellular homeostasis, translation and generation of precursor metabolites and energy under the biological process category; ribosome, cytoplasm, cell envelope and protein complex under the cellular component category; RNA binding, electron carrier activity, kinase activity, translation factor activity and structure molecule activity under the molecular function category; along with genome size and gene...
density. This prediction result is highly consistent with our above results shown in Figure 3.

Using the selected 25 variables from each genome, we have constructed a new neural network with one hidden layer to predict the overall percentage of genes on the leading strand as follows:

\[
P = \sum_{j=1}^{k_2} w_j^{(2)} f \left( \sum_{i=1}^{k_1} w_{ij}^{(1)} p_i \right),
\]

where \( P \) is the percentage of leading-strand genes in a genome, \( f \) is a hyperbolic tangent sigmoid transfer function, \( w_{ij}^{(1)} \) is the weight of the \( i \)th input to the \( j \)th node of the hidden layer and \( w_j^{(2)} \) is the weight of the \( j \)th node in the hidden layer to the output node in the neural network model, \( k_1 \) is the number of variables, \( k_2 \) is the number of nodes of the hidden layer, \( p_i \) is a scaling factor calculated as the ratio between the variable \( x_i \) and the max value \( x_{i,max} \) of this variable across all 725 bacterial genomes.

We speculate that the genes of certain functional categories need to be on the leading strands when living in certain environments to out-compete their competitors when food is limited and the competition is high; otherwise, the organism may keep the genes on the lagging strands as a more balanced genome may mean a more compact genome, which requires lower maintenance energy. A good example is that the chemotactic response of \textit{Pseudoalteromonas haloplanktis} in exploiting ephemeral microscale nutrient patches is at least 10 times faster than that of \textit{E. coli} (35), suggesting that \textit{P. haloplanktis} may be genetically optimized for this particular capability. To check whether some genes are specifically located on the leading strand of the organism, we examined the strand distribution of genes across the 55 GOslim functional categories on \textit{P. haloplanktis} and \textit{E. coli}, and we found that genes of some functional categories are significantly enriched (with \( P \) value < 0.05) on the leading strand of \textit{P. haloplanktis} than that of \textit{E. coli}, including protein transport, DNA metabolic process, ion transport and signal transduction under the biological process category; organelle and intracellular under the cellular component; antioxidant activity and motor activity (Supplementary Table S3). This clearly makes sense as...
collectively having more genes related to motor activity, transporter activity and signal transduction on the leading strand may enable the bacteria to react much faster when the nutrients become available (36,37).

**DISCUSSION**

It has been observed that bacterial genomes have a large variation in terms of the percentage of their leading-strand genes, ranging from ~45% to ~90%. We have provided an explanation for the large variation of observed strand biases across 725 bacterial genomes, which extends substantially the previous explanations. Our key contributions through this study include that (i) the genes of certain functional categories that need to be on the leading strands of genomes, to enhance the survivability of the host; (ii) genes of some functional categories such as transcription factor have higher preference to be on the lagging strands; (iii) there is at least one balancing force that keeps genes from all moving to the more efficient leading strands during evolution, particularly in nutrient-depleted environments and (iv) the percentage of leading-strand genes for a bacterial genome can be well explained using the numbers of genes in 25 functional categories outlined in (i) to (ii), genome size and gene density. We anticipate that more sophisticated analyses could possibly lead to quantitative models relating the percentage of leading-strand genes in a bacterial genome can be well explained using the numbers of genes in 25 functional categories outlined in (i) to (ii), genome size and gene density. We anticipate that more sophisticated analyses could possibly lead to quantitative models relating the percentage of leading-strand genes in a bacterial to a few parameters, which reflect the relationships between the living environments of an organism and the ‘intended’ capabilities of the organism and the needs for its survival, giving rise to improved understanding about the rules that may determine which genes will be on the leading versus the lagging strand of a genome.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online: Supplementary Tables 1–3 and Supplementary Figures 1–3.
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