



Anatomy of Escherichia coli Ribosome Binding Sites

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During translational initiation in prokaryotes, the 3' end of the 16S rRNA binds to a region just upstream of the initiation codon. The relationship between this Shine-Dalgarno (SD) region and the binding of ribosomes to translation start-points has been well studied, but a unified mathematical connection between the SD, the initiation codon and the spacing between them has been lacking. Using information theory, we constructed a model that treats these three components uniformly by assigning to the SD and the initiation region (IR) conservations in bits of information, and by assigning to the spacing an uncertainty, also in bits. To build the model, we first aligned the SD region by maximizing the information content there. The ease of this process confirmed the existence of the SD pattern within a set of 4122 reviewed and revised Escherichia coli gene starts. This large data set allowed us to show graphically, by sequence logos, that the spacing between the SD and the initiation region affects both the SD site conservation and its pattern. We used the aligned SD, the spacing, and the initiation region to model ribosome binding and to identify gene starts that do not conform to the ribosome binding site model. A total of 569 experimentally proven starts are more conserved (have higher information content) than the full set of revised starts, which probably reflects an experimental bias against the detection of gene products that have inefficient ribosome binding sites. Models were refined cyclically by removing non-conforming weak sites. After this procedure, models derived from either the original or the revised gene start annotation were similar. Therefore, this information theory-based technique provides a method for easily constructing biologically sensible ribosome binding site models. Such models should be useful for refining gene-start predictions of any sequenced bacterial genome.

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Introduction

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Ribosomes play a central role in cells by reading mRNA and synthesizing proteins.¹ The entire high-resolution atomic structure of the 50S² and 30S³ ribosomal subunits have been determined recently, but a full understanding of translation will also require quantitative mathematical descriptions. Because codons are three bases long, translational initiation must be directed to within one base on the mRNA. This requires a pattern in the

mRNA known as a ribosome binding site, which includes the initiation codon. The completion of entire genome sequences, and the identification of likely genes within them, now allows for the inspection of most ribosome binding sites and allows for the statistics of the patterns to be determined in greater detail than was possible previously.^{4–7}

In eukaryotes, ribosomes recognize the 7-methyl guanine cap to help identify the translation initiation codon.⁸ Prokaryotes, however, lack this marker and instead have a contact between the 3' end of the 16S rRNA in the 30S ribosomal subunit and a region upstream of the initiation codon, referred to as the Shine-Dalgarno region (SD).^{9,10} The ribosome protects RNA further downstream than just the initiation codon;¹¹ therefore, the

Abbreviations used: SD, Shine-Dalgarno; IR, initiation region; GS, gap surprisal.

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downstream region should be accounted for when modeling ribosome binding. The region around the initiation codon is referred to as the initiation region (IR).

The SD has strong effects on translation,^{10,12,13} and one of its most intriguing features is the variable spacing between it and the initiation region. Preferential binding of the 16S rRNA at certain spacings has been shown.^{11,14–17} We investigated how this spacing affects the sequence conservation of the SD and IR, and the patterns being bound for the majority of ribosome binding sites in *Escherichia coli*.

Nucleic acid and protein sequences can be analyzed by information theory, an approach that was originally applied to quantify the movement of data in communication systems.^{18,19} Unlike statistical measures of significance, information measured in bits defines the minimum number of binary choices needed to represent some data. The advantage of this measure, over all other measures, is that information from independent sources can be added together, and bits provide a universal scale. In molecular biology, the amount of information indicates the degree of sequence conservation among a set of aligned sequences. It is a quantitative measure that has proven to be more useful than consensus sequences for understanding a variety of genetic systems.^{6,20-23} The average information computed from a set of related sequences⁶ describes the overall conservation at each position in the alignment, and this can be shown with a sequence logo graphic (e.g. see Figure 1(a)).²⁴ The individual information present in a single sequence²⁵ measures how that sequence contributes to the average sequence conservation of the sequence family, and this can be shown with sequence walker graphics (e.g. see Figure 4).²⁶

Individual information is calculated as the sum of the conservation at each base position. Some bases are not favored and can have a negative value.²⁵ A site with overall negative information content should, according to the second law of thermodynamics, have a positive ΔG , and therefore should not be bound.²⁵ The theory, therefore, naturally provides a way to detect anomalous sites. Such sites can be removed to refine the dataset and thereby produce a more consistent model. Anomalous sites can be investigated to determine whether they represent sequencing errors, database errors, or novel biological phenomenon.

Although the spacing between the SD contact and the IR is variable, a "rigid" ribosome model functioned reasonably well.^{6,7,27} However, as more sites were added to the model, the information content of the SD region dropped, suggesting that the model was not sufficient to explain the variation among sites. We therefore investigated a ribosome binding model where the spacing between the SD and the IR was allowed to vary. This flexible model provides a better representation than a rigid model does.

We describe four main results. First, multiple alignment of the regions upstream of E. coli genes by maximizing the information content identified the SD pattern without reference to the 16S rRNA sequence. Secondly, ribosome binding could be modelled using a unified mathematical representation for the aligned SD, the IR, and the distribution of spacings. Thirdly, the second law of thermodynamics sets zero as the theoretical lower bound for the information of binding sites,^{25,28} so we could iteratively remove sites with negative information to heighten the model's predictive capabilities. Finally, further characterization of the Shine-Dalgarno model allowed us to observe how the SD pattern varies with distance from the initiation region.

Theory

Since early ribosome binding site models did not account for variable spacing between binding components,^{7,27} a new method for analyzing flexible sites was developed. First, the individual information of a binding site is computed from a rigid weight matrix defined as:²⁵

$$R_{iw}(b, l) = 2 - (-\log_2 f(b, l) + e(n(l))) \text{ (bits per base)}$$
(1)

where f(b,l) is the frequency of each base (*b*) at position (*l*) in the aligned binding site sequences and e(n(l)) is a sample size correction factor for the (*n*) sequences at position (*l*) used to create f(b,l).⁶ Then, to evaluate the individual information of a ribosome binding site using a flexible model, we calculated three values:

Flexible Site Information
=
$$R_i(SD) + R_i(IR) - GS(d)$$
 (bits/site). (2)

 R_i (SD) is the individual information of the aligned SD, R_i (IR) is the individual information of the aligned IR, and GS(d) is the gap surprisal, which accounts for the variable spacing *d*.

The SD was aligned in two steps. First, the sequences upstream of the initiation codon were embedded into random sequence so as not to trigger alignment by the well-conserved initiation codon. Second, the sequences were shuffled to maximize the information content.²⁹ By aligning the SD, we obtained the distribution of distances from the IR. Any probability distribution has an uncertainty measured in bits:

$$H = -\sum_{d} p_{d} \log_{2} p_{d}$$

= $\sum_{d} p_{d}(-\log_{2} p_{d})$ (bits) (3)

where p_d is the probability of the distance d.^{18,19} Rewriting the uncertainty as shown on the right-hand side shows that it can be expressed as an



Figure 1. Rigid and flexible ribosome binding site sequence logos. (a) The rigid model of the entire EcoGene12 set;³⁴ (b) the EcoGene12 unrefined flexible model; (c) the EcoGene12 refined flexible model; (d) the Verified refined flexible model. For all logos, the height of each stack of letters corresponds to the total sequence conservation at that position, measured in bits.⁶ The height of each letter corresponds to the relative frequency of that base at that position.²⁴ The sine wave represents the 11 base twist of *A*-form RNA.⁵³ The histogram between each pair of flexible logos represents the distribution of distances between the Shine-Dalgarno and the initiation region zero coordinates. A Gaussian distribution with the same mean and standard deviation is shown for comparison. All logos on the left represent the Shine-Dalgarno alignment and all logos on the right represent the initiation region alignment. The sequence shown under each SD logo is the anti-Shine-Dalgarno sequence found on the 3' end of the 16S rRNA.

average of the surprisal function:³⁰

$$u_d = -\log_2 p_d$$
 (bits/spacing) (4)

n(d), the number of sites with a binding distance d, is divided by the total number of sites, n, to obtain the frequency of binding at each distance. The *GS* equation is therefore:

$$GS(d) = -\log_2 \frac{n(d)}{n} + e(n) \quad (bits/spacing) \quad (5)$$

where e(n) is a small-sample correction for *GS*, required because we have substituted a frequency for the probability p_d .^{6,25}

GS(d) is positive when there is more than one spacing possibility and it has the same units (bits) as R_i (SD) and R_i (IR). We assume that the spacing is independent of the SD and IR,¹⁶ so we subtracted GS(d) from the SD and IR individual information to obtain equation (2). Other similar methods^{31,32} cannot be used to compare models from different datasets, because they use consensus sequences, which are sensitive to small changes in the sequences. In contrast, the individual information method allows comparison between matricies from different recognizers (SD and IR in this case) and evaluations converge to a single value as the data set size increases.²⁵

Results

Characteristics of flexible ribosome binding site models

Several different E. coli ribosome binding site models were used for various purposes. Models may be rigid, in which case all parts are fixed relative to a zero coordinate, or flexible, in which case the model contains two rigid parts (SD and IR) separated by a variable distance. These models are further characterized as being either unrefined or refined. Refinement refers to a cyclic process in which an individual information model is made from the current set of binding sites and then sites that have negative information content are removed from the set. This process is repeated until only positive sites remain (see Materials and Methods). In order for the information to be calculated for a flexible model, one must take into account the statistics of the spacing between binding components. The effect of the spacing, called the gap surprisal (GS) (Theory, equation (5)), is given in bits and is subtracted from the sum of the information present in the SD and IR binding components, also measured in bits (Theory, equation (2)).

We used three databases in this work. The protein-coding feature locations in the completegenome GenBank entry U00096 have not been updated since the original publication,³³ so our first database was the alternative set of gene intervals present in EcoGene12.34 This revised database, which contains 4122 known and putative translation start sites in *E. coli*, is the result of an intense and continuous effort to improve the annotation and prediction of E. coli genes. Second, we used the Verified subset of this database, which is composed of protein start sites confirmed by N-terminal protein sequencing. The third database is the original E. coli annotation from Blattner (Blattner et al.,33 GenBank U00096). To create a reliable baseline model, we refined the Verified set. In contrast, a ribosome model built from the refined Eco-Gene12 database is probably the most representative of all genes. We also refined the Blattner database to determine if we could automatically derive a model comparable to EcoGene12.

The Verified model is derived from ribosome binding sites for proteins that have been well studied and/or detected as spots on 2D gels and it probably lacks many sites that show lower binding affinity. Despite this bias, the Verified model is useful, since it is composed only of sites proven to be actual ribosome binding sites. For example the range of SD to IR spacing from -18 to -4 was established by observing spacings utilized within the Verified set. The EcoGene12 model is based on the full set of proven and predicted gene starts, and thus is representative of all ribosome binding sites, including weak sites responsible for low-level protein expression. Although EcoGene12 may contain a few predicted sites that turn out to be incor-

rect, we consider it to be the most accurate model, and therefore we used it as our benchmark model.

The rigid model sequence logo made from all EcoGene12 translation start sites (Figure 1(a)) shows the expected strong conservation for the initiation region at bases 0, +1 and +2 and a low region of conservation from bases -12 to -6 for the SD. When the SD was re-aligned to maximize the information,²⁹ its information present rose from $1.53(\pm 0.03)$ to $4.96(\pm 0.04)$ bits. (We report here the mean R_{sequence} and standard error of this mean from the individual information distribution.²⁵) The range of re-alignment (-18 to -4) was selected to allow for all spacings observed in the Verified model. The SD was realigned using only sequences from translation start sites, and this was done independently of the 16S rRNA sequence, yet the sequence logo closely complements the 16S rRNA 3' end. This flexible model has an SD-IR spacing of -18 to -4 bases, with a peak of occurrence at -9(Figure 1(b)). When the model was tightened by the exclusionary refinement using process (Figure 1(c)), there was again an increase in the information present in the SD logo to $5.23(\pm 0.04)$ bits. In contrast, the refined Verified SD has 5.77(\pm 0.10) bits with an SD-IR spacing of -18 to -4 bases, with a peak of occurrence at -10(Figure 1(d)).

Logos were made in the same fashion as Figure 1(a)-(c) for the Blattner sites. Since the logos looked similar to the EcoGene12 logos, they are not shown. When the SD region was re-aligned to maximize information in the Blattner model, the information rose from $0.91(\pm 0.03)$ SD to $3.87(\pm 0.05)$ bits. This model has an SD-IR spacing of -16 to -2 bases, with a peak of occurrence at -8. Refinement of the Blattner model also showed a further increase in the SD information to $5.01(\pm 0.04)$ bits. The most noticeable difference between the unrefined and the refined Blattner model is at the zero position of the aligned SD. This position went from a partially conserved G at \sim 1.5 bits to a fully conserved G at 2 bits, while the rest of the positions increased proportionally. Interestingly, upon refinement the SD-IR spacing shifted to -18 to -4 and the peak shifted to -9bases. This is the same range as that seen in the well-characterized Verified model (Figure 1(d)).

For all models, a Gaussian distribution with the same mean and standard deviation as the respective SD-IR spacing distributions was plotted along with the spacing histogram. In all cases, the histogram did not match the Gaussian plot.

Using the individual information method,²⁵ all sites in the Verified set were evaluated by the rigid and flexible EcoGene12 refined models over the range of 30 bases upstream to 14 bases downstream of the first base of the initiation codon. This is the range required to identify a site with the maximum SD-IR spacing of 18 bp. Previous information theory-based ribosome evaluations with a rigid model have been reasonably accurate,⁷ but since that model does not take into account vari-

able spacing, it is limited in its analysis of ribosome binding. The rigid EcoGene12 model (range -21 to +14) picked up about the same number of upstream non-sites (sites with more than zero bits of information other than those annotated in the data set) as the flexible model (92 *versus* 89, respectively). The two models identified nearly the same number of Verified start points (565 *versus* 567). The average site strength assessed by the rigid model was $9.50(\pm 0.13)$ bits, and with the flexible model it was $10.17(\pm 0.14)$ bits, indicating that the flexible model generally assessed the Verified sites more strongly.

Shine-Dalgarno as a function of spacing

To understand better the function of the Shine-Dalgarno, we examined SD sequence logos at every SD-IR spacing in the EcoGene12 set (Figure 2). The shape and pattern of the SD remained fairly constant, but the information present fluctuated. There was a constant increase in the information as the spacing was increased from -4 to -9, and a decrease in information for -9 to -18. This is reflected in the change of the size of the bases surrounding the central G. The information present in the SD at each alignment relative to the IR is only weakly related to the conservation of information in the IR (r = -0.17) (Figure 3). When the total flexible site information, as calculated from equation (2) (see Theory), was examined for all positions, an increase and decrease in information was observed similar to that observed with the SD region alone. When the refined Blattner sites were split into spacing classes, similar results were obtained (data not shown).

For each spacing class, the program diana³⁵ was used to determine if there was any correlation between bases in the SD and IR. None was observed at any spacing (data not shown), suggesting independence between the SD and the IR. In addition, no correlation is observed between parts of the refined EcoGene12 SD (Figure 1(c)) when all classes are combined.

For all spacings of -4 to -11 there is an A with low conservation at position -3,^{5,6} and it is present also from -16 to -18, indicating that conservation at this position is an effect of the initiation contact and not the SD (Figure 2).

The minimum SD-IR spacing of -4 has been observed in *nadB*³⁶ but appeared infrequently in EcoGene12. Binding of regions with spacings more than 18 bases is known, but these are rare and due to RNA structural effects such as hairpins that bring the SD closer to the IR.¹²



Figure 2. The Shine-Dalgarno as a function of spacing. Sequence logos were constructed for all distances between the SD and IR zero coordinates observed in the Eco-Gene12 refined set. The black circle falls under the central G of the Shine-Dalgarno, which is the zero coordinate of the SD in the variable model.



Figure 3. Quantification of ribosome binding site components as a function of spacing. The information present in the Shine-Dalgarno regions of Figure 2 (shown in green boxes) were plotted at their respective distances. The information content was measured over the region 12 bases prior to and 4 bases after the central G of the Shine-Dalgarno, except for the spacing of -4, whose information is measured over the range of -12to +3, because of interference with the IR at position 0. The information present in the IR for the range of -3 to +14 at each distance is shown in black (with small filled circles). The gap surprisal GS computed by equation (5) from the distance distribution in Figure 1(c), is plotted as open blue circles. The red curve with no symbols shows the total flexible information at each spacing, as calculated by equation (2). For all cases, error bars are plotted with black I symbols (the error for GS is smaller than the circle).

Correlation between the refined Blattner and EcoGene12 models

To test whether the refined EcoGene12 model is accurate and can be used to correct sequence annotations, we scanned the model across several proven ribosome binding sites and across several sites predicted by Blattner that have been corrected in EcoGene12 (Figure 4). When we applied our model to the well-studied lacZ and lacI initiation regions, it concurred with Blattner's locations (Figure 4(a) and (b)). In the case of the 8.1 bit lacl ribosome binding site, which starts at a GTG in the context atGTGa, a second weaker 5.5 bit site is seen at the out-of-frame ATG just upstream. Interestingly, ribosomes binding to this site should terminate immediately at the TGA. Using two of Blattner's sites that have been corrected based on N-terminal protein sequencing, we tested whether our model locates the correct binding sites. In mhpD (Figure 4(c)), we saw a 12.8 bit site at the correct location six bases downstream from Blattner's prediction. Our model did not predict any site ($R_i > 0$) at the Blattner location. In *yhbL* (Figure 4(d)) there is a predicted weak 4.5 bit site at the incorrect position, but experimentally the start site was proven to be nine bases downstream and our model favored this location (13.7 bits). As expected, in both cases the correct site was found in the same reading frame as the predicted site. When the refined Blattner model was scanned over these same sites, the same predictions were made, indicating that the refined Blattner model is comparable to the refined EcoGene12 model.

To investigate further the effect of refinement, we scanned both the Blattner unrefined and refined models over all of the EcoGene12 sites for regions 100 bases upstream and 100 bases downstream of each of the 3900 refined EcoGene12 start points. The unrefined Blattner model found 21,464 non-sites and the refined model found considerably fewer non-sites (12,018). This large number of sites detected may represent weak ribosome binding sites, untranscribed regions or may be false positive artifacts of this model. Alternatively, some of these sites may be occluded by RNA secondary structure. Since the unrefined Blattner model contains many non-sites, it has a lower information content and therefore picks up more non-sites than the refined Blattner model. Both the unrefined and the refined Blattner models identified all of the EcoGene12 sites.

To generalize Figure 4(c) and (d), we scanned both the refined Blattner model and the refined EcoGene12 model over the 26 sites in the Blattner annotation that have been corrected in the Eco-Gene12 dataset based on experimental verification. The Blattner model identified the experimentally reported start site as the strongest site in 21 of the 26 cases. In four of the five other cases, the model assessed the Blattner annotation more strongly, but also found a site at the confirmed start point. For one gene (gntK), the model did not match either Blattner's annotation or the experimentally proven TTG start. When this same analysis was done using the refined EcoGene12 model, the correct site was predicted in 22 of the 26 sites and three of Blattner's annotations were favored. As with the refined Blattner model, no site was predicted for the *gntK* gene at either the Blattner or EcoGene12 locations. As exemplified by Figure 4(d), in approximately half of the 26 corrected sites both models predicted strong sites at the verified locations and these were accompanied by weaker sites at Blattner's locations. In the three (Eco-Gene12) or four (Blattner) cases where the verified start was weaker than the Blattner site for either model, the difference in site strength between the site at the Blattner location and the site at the Verified location was generally only 1 to 2 bits (except for one case, where the difference was around 5 bits). These results show that refined flexible



Figure 4. Lister maps with sequence walkers for four ribosome binding sites. Blattner's sequence, GenBank accession number U00096, is annotated with 4290 gene starts,³³ four of which are illustrated; (a) lacZ, (b) lacI, (c) mhpD and (d) yhbL. The EcoGene12 flexible model (Figure 1(c)) was scanned across each sequence. The sites that are found $(R_i > 0)$ are shown by two part sequence walkers. A walker is a graphic consisting of several adjacent letters with varying heights.²⁶ Vertical green rectangles indicate the zero coordinate of each sequence walker and provide a scale from -3 to +2 bits. Braces ({ and }) connected by a broken line are used to link SD and IR walkers. This feature, created by the program biscan, also reports the distance of separation, the coordinate of the IR and the flexible site information value according to equation (2). All correct translation start points, based on experimental data, are identified by a filled black arrow starting at the initiation start point. The broken-line arrow shows Blattner's predicted gene start. The color bar above the sequence cycles through three colors to illustrate the reading frames. In cases (c) and (d), it is obvious that the predicted (boxed and broken-line arrow) and corrected sites (boxed and full arrow) fall in the same reading frame because the adenine bases lie under the same color. The sine waves represent the 11 base twist of A-form RNA.53 The asterisks and numbers above the sequence indicate positions on the Escherichia coli genome.33

information models can be used to improve ribosome binding site predictions.

Can we create a valid ribosome model from the large lists of gene start-points determined from open reading frames that are presented as annotations for complete genome sequences? To test for relatedness, we compared various models using the Euclidean distance between $R_{iw}(b,l)$ matrices (Materials and Methods, equation (6)). The distance between the unrefined Blattner SD matrix and the refined EcoGene12 SD matrix was 13.0 bits and the distance between the corresponding IR matrices was 2.2 bits. In contrast, when the refined Blattner was compared to the refined EcoGene12 model, there was a much smaller difference: for the SD matrix there was a distance of 0.9 bit. Refining

the Blattner model brought it closer to the refined EcoGene12 model, which is representative of the bulk of *E. coli* ribosome binding sites.

When the individual information distributions for all models were compared, there was a general increase in the strength of sites from the unrefined to the refined to the Verified model (Figure 5 and Table 1). This effect may occur not only because the refinement process removes negative sites, but also because the well-characterized sites in the Verified model may tend to neglect weaker sites, as these may often be harder to characterize biochemically. The sets overlap reasonably well, since 507 of the 569 Verified sites are found in the refined Blattner set, and all but six of the Verified sites are found in the refined EcoGene12 model.



Figure 5. Individual information distributions for five ribosome binding site models. The ordinate is the individual information and the abscissa is the frequency of occurrence. (a) The information distributions for the Eco-Gene12 unrefined (red circles), the EcoGene12 refined (green boxes) and the Verified model (black, with no symbols). (b) The information distributions for the Blattner unrefined (red circles), the Blattner refined (green boxes) and the Verified model (black, with no symbols).

Discussion

Unlike gene-finding programs,^{37,38} ribosomes do not use open reading frames or other global factors to recognize translational start points, so our philosophy is to model the ribosome explicitly. A pure model has the advantage that it can identify ribosome binding sites in the center of genes, such as the out-of-frame one at *E. coli* coordinate 3919396 in *atpB* (formerly *uncB*)³⁹ and those of short polypeptides as in transcription attenuation.⁴⁰

To create a flexible ribosome model, we first removed the initiation codon and downstream open reading frame by embedding the SD region into random sequence. This allowed us to use multiple alignment to focus the SD region by maximizing its information content.²⁹ The SD emerged easily (Figure 1(b)), demonstrating mathematically the existence of this feature in the majority of *E. coli* ribosome binding sites. Furthermore, the general pattern matches the 3' end of the 16S rRNA well, confirming independently that these are complementary to each other.³⁷

In contrast with the notion of an SD consensus sequence, sequence logos show that the SD is variable and its pattern depends on how far the sequence is from the IR (Figure 2). Despite this variability, the information content of the SD is relatively constant at various spacings, smoothly increasing and decreasing in a range of only 2 bits from -18 to -5 (Figure 3). Surprisingly, the SD-IR spacing contributes more variation to the total information than either the SD or the IR. Furthermore, the variation of the SD information works in the same direction as the gap surprisal; they do not compensate for each other but instead work together. This sets up a maximal range of variation for efficiency of translational initiation. These observations are consistent with the SD-anti-SD helix formed between the rRNA and the mRNA as being a reasonably consistent "object", whose placement relative to the IR is important.

In all cases (Figure 1) the spacing distribution between the SD and the initiation region was similar to but differed from a Gaussian distribution. There is a predominance of -9 and -8 spacings (and -10 for the refined Verified set). A spring (simple harmonic oscillator⁴¹) moving under the influence of random thermal noise should produce a Gaussian spacing distribution.⁴² Since there is a non-Gaussian distribution, the SD-mRNA helix appears to have more constraints than a freely oscillating spring. What these bounds are may become apparent only when crystal structures of initiating ribosomes have been determined, but a clue that the meaning is related to the placement of the SD-anti-SD helix comes from the shape of the SD sequence logos.

Unlike the rectangular block that a consensus sequence would make on an information graph, the SD sequence logo rises smoothly and declines with position (Figure 1(c)). This is consistent with the idea that mismatches at the center of an RNA-

Table 1. Comparing individual information distribution values

1 0					
Model	Mean (bits)	Std Dev. (bits)	SEM (bits)	п	
Blattner unrefined	6.83	4.84	0.07	4290	
Blattner refined	8.82	3.63	0.06	3509	
EcoGene12 unrefined	8.81	3.99	0.06	4122	
EcoGene12 refined	9.28	3.58	0.06	3900	
Verified	10.35	3.73	0.16	569	

We report the mean, standard deviation (Std Dev.), standard error of the mean and number of sites for each model. These values correspond to the distributions in Figure 5.

RNA hybrid should be more disruptive than mismatches towards the ends. However, the situation may be more complicated. Sequence logos for duplex DNA binding proteins also rise and decline with position.^{21,43–45} One intriguing explanation is that the formation of the mRNA-rRNA hybrid is followed by binding of a ribosomal protein or RNA^{3,46} into the resulting major or minor groove as a step during translational initiation. Such a model accounts for the shape of the SD sequence logo because proteins tend to evolve contacts on one face of a helix, and such contacts become progressively more difficult to form when they are close to the back face.44 The proximity of protein S1 to the SD^{11,47,48} suggests it as a candidate for this process, but other proteins such as S7, S18 and S21,⁴⁹ and various 16S rRNA positions^{50,51} that crosslink to the mRNA⁵² could be involved. To allow us to judge the validity of this model, we added a dashed sine wave to the sequence logos. The peaks of this wave are separated by 11 bp, which is the distance between two major grooves of A-form RNA.53 Preferred spacings of the SD (Figure 2) are consistent with this model, but there is clearly a greater degree of flexibility than in DNA-protein interactions. However, tight packing is observed throughout the 70S subunit^{2,54} and there is close packing in the 30S^{,3} so it is likely that the fully assembled initiation region is also tightly packed. This suggests a mechanism for initiation in which the binding of the 16S 3' end to the mRNA SD allows the resulting helix to be smaller than unpaired single strands would be. The smaller helix could pack against other components of the ribosome, reducing the volume further and completing initiation, perhaps by creating sufficient space in the A site for the next tRNA. Even a nonspecific RNA phosphate backbone binding into the minor groove between the SD and the mRNA^{3,46} could account for the sinusoidal shape of the aligned SD sequence logo. IF3 appears to recognize codon-anticodon complementarity at the initiation codon rather than direct recognition of the codon itself.55 Because complementarity usually creates a more compact structure than a mismatch, this effect is also consistent with a packing model for initiation. Finally, this tight-packing model may account for why the SD-IR spacing is narrower than a Gaussian distribution: the simple harmonic oscillator is confined in a box.

The concept of individual information²⁵ allows us to consistently apply an information measure not only to the SD and IR but also to the gap distance between individual sequences, thereby creating a flexible search tool. The problem of how to compute the information content of flexible binding sites was recognized previously.⁶ If two sequence elements have a variable distance between them, then the uncertainty in position decreases the overall information content. For example, GC, with an information content of 2 bases × 2 bits per base = 4 bits, is found every 16 bases in equiprobable DNA, while GNC is found at the same frequency. A shorthand notation for the set containing both of these is G1EC, in which 1E means to search for G followed by C with an extendable spacing of 1 or 0 bases.⁵ Because it contains the search for both GC and GNC, G1EC occurs approximately every 8 bases in equiprobable random DNA. So, although the G and C contribute 4 bits of information, the variable spacing removes 1 bit and the site is therefore effectively only 3 bits. With G3EC there are four possible search patterns, GC, GNC, GNNC and GNNNC; this removes $\log_2 4 = 2$ bits. Interestingly, G15EC has 16 search patterns and this removes 4 bits giving the, at first sight, odd result that the information content is 0 bits. However, in a sequence M bases long, G15EC will occur roughly M times because of overlapping cases, so the result is consistent. It is interesting to note that there can be sites with negative information by this method: in a sequence of length M, G31EC will occur roughly 2M times, giving an apparent information of -1 bit. The reason for this odd effect is that there are many overlapping sites. We interpret zero or negative information to mean that the two components are independent.

We have extended these computations by using Shannon's uncertainty measure to consistently assess the contribution when different spacings occur with different frequencies. Because frequencies are not probabilities, a small sample correction was applied.⁶

Fortunately, the negative information effect does not occur for ribosome binding sites. In the refined EcoGene12 model (Figure 1(c)), the SD contains $5.80(\pm 0.04)$ bits, the IR contains $6.72(\pm 0.04)$ bits, and the uncertainty of the distance between them (gap uncertainty, H_{gap}) is 3.25(±0.02) bits, giving information content⁶ of $R_s(SD) +$ total а $R_s(\text{IR}) - H_{\text{gap}} = 9.28(\pm 0.06)$ bits. This is similar to the refined rigid EcoGene12 model, which has $8.92(\pm 0.05)$ bits, but quite different from the flexible Verified model with $10.35(\pm 0.16)$ bits. We suggest that the difference occurs because strong sites tend to be experimentally identified first and some non-functional sites may still contaminate the refined EcoGene12 model. The latter effect can be observed in Figure 5, where the unrefined Eco-Gene12 has examples of sites below zero, while the refined EcoGene12 set does not have any sites below zero bits, by definition. While there are no sites below zero in the refined Verified set (because we removed the 13 that we found), the lower end of the distribution curve is smaller than that for the refined EcoGene12. Further, the shape of the Verified distribution is a more Gaussian-like curve, trailing smoothly down to nearly zero at zero bits,25 while the refined EcoGene12 distribution still has members near zero bits and is therefore discontinuous. It is not known if these very weak sites are functional.

The Verified sites that we removed during refinement (gene at U00096 coordinates and orientations: *uppS* 194903+, *gsk* 499349+, *fes* 612038+,

dbpA 1407535+, topB 1844984-, guaB 2632090-, xseA 2632252+, trmD 2743359-, pcm 2867542-, cysI 2888122-, dnaN 3879949-, aceK 4216175+ and arcA 4637875-) presumably initiate differently from the majority of sites. Surprisingly, this set does not contain *infC*, which codes for IF3. In the absence of this initiation factor the ribosome can use the AUU start of *infC* (1798662–) for initiation, forming a regulatory feedback loop.⁵⁵ By the Verified model, the AUU-containing IR is -4.8 bits but this is compensated by a GS of 2.3 bits at the optimal spacing of -9 bases and an SD of 9.6 bits to give a total of 2.5 bits. This anomalous site was removed automatically during refinement of Eco-Gene12, because the G at the third base of start codons is otherwise invariant and rare bases are weighted against more heavily in larger datasets.²⁵ By the EcoGene12 model, the *infC* IR is -8.4 bits with an SD at a -9 spacing of 8.7 bits for a total of -2.0 bits. This model predicts that mutating the start codon from AUU to AUG should bring the IR up to 5.5 bits to give a strong 11.9 bit site.

Other mechanisms may be needed to explain the anomalous Verified sites. Only two excluded cases in the Verified set have GTG starts, which are known to be weaker than ATG starts.¹³ With fewer examples in the data set, marginal GTG starts could have been removed because of statistical noise.

Another way to explain the Verified site anomalies is that RNA secondary structures might bring an SD closer to the IR¹² and so influence translation.^{11,56} As shown in Figure 6, this mechanism might be involved in *fes* (612038) in which a four base helix (-4.5 kcal/mol)⁵⁷ may bring a 3.4 bit SD to position -11 with respect to the IR and *pcm* (2867542) in which a five base helix (-7.8 kcal/mol) brings a 3.5 bit SD to position -9with respect to the IR. The other sites do not appear to use this mechanism.

The relatively large number of initiation regions that do not conform to the majority model (i.e. the rejected Verified sites) suggests the possibility that there are even more alternative mode(s) of translation initiation. We are left with a number of likely and proven genes that fail to have ribosome binding sites that conform to our model. A combination of computational and experimental approaches will be needed to identify alternative models among the rejected sites. Of course, one simple possibility is that apparent anomalies can be caused by sequencing errors.²⁶

An empirical observation for human splice junctions is that, in addition to the thermodynamic bound at zero bits,²⁵ sites with less than 2.4 bits are non-functional.⁵⁸ We suspect that such a fuzzy non-zero bound may also apply to the majority of ribosome binding sites. However, unlike the case with splice junctions, experimental data are not available to suggest what a natural bound may be that delineates a functional from a non-functional ribosome site. For this reason we used the zero bound, which is based on the second law of thermodynamics, for cyclic refining.

The process of subtracting the gap uncertainty is similar to the accounting of gaps provided by hidden Markov models (HMM),^{38,59} except that the gap size we use is variable and the frequencies of different gap distances are accounted for.³⁷ It may be possible to extend the model given here to a full information-theory based HMM, but this was not attempted.

The information theory approach allowed us to build models that represent the vast majority of ribosome sites without having to assume that some sequences were not sites. In contrast, training with a neural net²⁷ requires this assumption, because data on where ribosomes do not bind are sparse. Because the data set is so large, it could be split into spacing classes (Figure 2), effectively dissecting the ribosome binding sites. The resulting models revealed that the weakly conserved "A at $-3^{\prime\prime 5,6}$ correlates with the IR and not with the SD. This is consistent with the presence of an A at -3relative to the translational start in eukaryotic mRNAs, which do not have an SD.8 The function of this conservation is unknown but crosslink experiments place it close to U1381 on the 16S rRNA⁵¹ and to the S7 protein.^{3,49}

The effort required to generate a data set as exemplified by EcoGene12 is enormous. The refining process described here gives results comparable to EcoGene12, so we believe it will be useful for gene analysis in other species. Ideally, all organisms would have models consisting of biochemically supported sites, rather than sites that were chosen by computer algorithms that do not model the SD. However, as shown here, it is possible to use information theory methods to help produce a reasonably clean identification of gene starts. This technique will be useful to



Figure 6. mRNA folding may rescue *fes* and *pcm* translation. Structure base-pairings are indicated by parentheses. Start sites were predicted by sequence walkers as in Figure 4.

better characterize medically important disease organisms.

Materials and Methods

Databases

To create our models, we drew from three databases. One database that we used was the 4122 sites in the Eco-Gene12 database, which represent the majority of *E. coli* genes.³⁴

The second database was a carefully compiled list of 569 experimentally supported sites, referred to as the Verified database.³⁴ This database is a subset of the Eco-Gene12 database and provided us with an initial comparison model that was used to determine the allowed SD-IR spacings and the general pattern of the Shine-Dal-garno. Rudd³⁴ has catalogued from the biomedical literature 717 *E. coli* proteins whose N termini have been determined directly by protein sequencing. The Verified proteins that have cleaved signal peptides were omitted, since these N-terminal protein sequences do not verify the translation start codons as definitively as the 569 Verified proteins that are uncleaved or have only the initiator methionine residue cleaved. This dataset can be obtained from the internet at: http://www.lecb.ncifcrf.gov/~toms/papers/flexrbs/

The third database was the 4290 gene starts presented by Blattner *et al*³⁷ and extracted from their complete *E. coli* GenBank entry, U00096 (version M52, September 02, 1997); this is referred to as the Blattner database. The refining method described below was performed on these databases.

Creating a ribosome model

Our ribosome models have two rigid binding elements connected by a flexible bond that allows the spacing between the elements to vary. If both elements do not find a suitable contact at an appropriate spacing, then the model will not bind. The first binding element is represented by a sequence logo made from translation start codons, which we refer to as the initiation region (IR) (Figure 1(a)). The range of this model is from -3 to +14, representing the predominantly A conservation at -3 through the downstream mRNA protected by the ribosome.^{6,11,17} To create this logo, standard Delila tools were used.^{24,44}

For the SD there is only a low sequence conservation over the range of -11 to -7 in the rigid logo (Figure 1(a)), so we used multiple alignment to build on the rigid model to create a flexible ribosome model. Random sequences were generated by the markov program and the -20 to -4 range of the ribosome binding sites was embedded into the random sequence using the embed program. By replacing the IR with random sequence we avoided alignment by the IR in the next step. This step was to use the malign program to realign the SD region to maximize the information present.²⁹ The resulting alignment was then represented by a logo using the previously described method. This realigned set displays the complement of the anti-Shine-Dalgarno found on the 16S rRNA (Figure 1(b)) and was used as our Shine-Dalgarno model. The zero coordinate of this model was shifted to the coordinate of the large central G using instshift. Since this base position contains the maximum information, presumably it is the most stable position to use and it will appear as the most significant base in a sequence walker (Figure 4). By this definition, our +4 base corresponds to the SD_{ref} reference point defined by Chen *et al.*,¹⁴ and our spacing measures are "aligned spacing" according to these authors. This measures the distance from a fixed point on the 16S rRNA to the initiation codon, as they advocate.

Once we had both an SD and an IR model, we made a histogram of the distances between their zero coordinates for all sites in the database. This was done using the diffinst program, which calculates the distance between coordinates in a pair of Delila instruction sets. These distances were then presented in a histogram using the genhis program and graphed in postscript with genpic.

The program used to compute equation (1) is ri,²⁵ which generated the $R_{iw}(b,l)$ weight matrix for both the SD and IR for further analysis of the individual information conserved in ribosome binding sites (see Theory). The SD sites were assessed for the region -12 to +4 and the IR sites were assessed for the region -3 to +14 because this is the region covered by footprints.⁷ The program used to compute equation (2) is biscan (see Theory). Biscan finds pairs of SD and IR that fall within the range of the spacing histogram, and then the flexible site information is calculated for each coupling using the distribution of distances from the genhis histogram.

Further information about the programs is available at http://www.lecb.ncifcrf.gov/~toms/and a web-based server with guest-access is available at http://www.lecb.ncifcrf.gov/~toms/delilaserver.html

Cyclic refinement

The 5' ends of genes are often placed incorrectly in sequence database feature tables. To obtain a reliable ribosome model containing a minimum number of misplaced sites, a cyclic refinement method was used. To do this, we computed the flexible site information for all sites in the set. We removed all sites whose information content was less than zero (this is the theoretical boundary for binding because of the second law of thermodynamics²⁵) and we rebuilt the model with the corrected set. Following every round of refining, the SD was realigned by malign as previously described.²⁹ Refinement used 1000 realignments, maximized the information in a window from -20 to -4 and allowed the sequences to shift from -8 to +6. This range was chosen to match the known binding range of the Verified model. If more than one Shine-Dalgarno site was found upstream of an initiation start site, then the SD that gave the strongest flexible site information was used in the model. The alignment with the highest information content was chosen from the 1000 realignments. This process was repeated until no site remained in the data set with a negative flexible site information. The EcoGene12 set required ten rounds of refining and lost 222 sites; the Verified set required two rounds and we dropped 13 sites, and Blattner's set required 20 rounds of refining and we dropped 781 sites. Each round took approximately two hours on a 450 MHz Sun Ultra60 Sparc workstation.

Dissecting the SD

To generate the SD as a function of spacing (Figure 2), a logo was made for each of the 15 observed spacing groups. For example, six sites were observed to have an SD-IR spacing of -4 bases, so a

logo was made for those sites (upper left corner of the Figure). This was repeated for the range of -18 to -4 for the refined EcoGene12 model. The graph of information present in the SD *versus* spacing (Figure 3), is the R_{sequence} for the range of -12 to +4 around the central G in the SD portion of the logo. The range -12 to +3 was used for the spacing of -4, because of interference with the zero position of the initiation region. The range for the IR information curve was -3 to +14. To examine relatedness between nucleotides for an SD-IR spacing, correlations between nucleotides were computed using the program diana.³⁵

Sequence walkers

To make flexible sequence walkers²⁶ (Figure 4), biscan generated features that were then mapped by the programs live, mergemarks and lister. Live created a color bar that changed hue every three bases to mark reading frames.²² Mergemarks combined marks from various sources and lister generated the sequence walker graphics.²⁶ The refined EcoGene12 model was used for this analysis. For Figure 6, mfold 3.1⁵⁷ was used to fold RNA sequences and the structures were displayed along with the walkers using mfoldseq, which generates sequence files for mfold, and mfoldfea, which uses the output from mfold to create features for lister.

Comparing matrices

To compare two weight matrices, we calculated the Euclidean distance between them using the following equation:

Distance =
$$\sqrt{\sum_{l} \sum_{b} (Ri_1(b, l) - Ri_2(b, l))^2}$$
 (bits) (6)

Here, the difference is taken between the individual information (R_i) of each base (b) at each position (l) between matrices 1 and 2. The difference is then squared and summed for all positions and the square-root of this value is taken, giving the distance in bits. The program used to do this was diffribl.

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