Information theory as a model of genomic sequences

Chengpeng Bi and Peter K. Rogan
Laboratory of Human Molecular Genetics, Children’s Mercy Hospital,
Schools of Medicine and Computer Science and Engineering, University of
Missouri-Kansas City, US

Keywords: Information theory, entropy, thermodynamics, surprisal, weight matrices,
binding sites, sequence logo, sequence walker, model refinement, evolution

Correspondence:
Peter K. Rogan, Ph.D.
Children’s Mercy Hospital and Clinics
2401 Gillham Rd.
Kansas City MO  64108
816-983-6511
Fax 816-983-6515
progan@cmh.edu
Abstract

Shannon information theory can be used to quantify overall sequence conservation among sets of related sequences. Variation in nucleic acid sequences recognized by proteins can be comprehensively modeled with information weight matrices that permit each member sequence to be rank ordered according to its respective individual information contents. These rankings can be used to compute the affinities of recognition sites by proteins and to predict the effects of nucleotide substitutions in the sequences of these sites. The distribution of information across a set of protein binding sites in DNA is related to the pattern of intermolecular contacts that stabilize the protein-nucleic acid complex (i.e. the corresponding helical structure of double stranded DNA).

A. Theory

Shannon and Weaver (1949) developed their theory of information in order to understand the transmission of electronic signals and model the communication system. Gatlin (1972) first described its extension to biology. Information theory is an obvious tool to use in looking for patterns in DNA and protein sequences (Schneider, 1995). Information theory has been applied to the analysis of DNA and protein sequences in several ways: (1) by analyzing sequence complexity from the Shannon-Weaver indices of smaller DNA windows contained in a long sequence; (2) by comparing homologous sites in a set of aligned sequences by means of their information content; and (3) by examining the pattern of information content of a sequence divided into successively longer words (symbols) consisting of a single base, base pairs, base triplets and so forth.

Some of the most useful applications of molecular information theory have come from studies of binding sites (typically protein recognition sites) in DNA or RNA recognized by the same macromolecule, which typically contain similar but non-identical sequences. Because average information measures the choices made by the system, the theory can comprehensively model the range of sequence variation present in nucleic sequences that are recognized by individual proteins or multi-subunit complexes.
Treating a discrete information source (i.e. telegraphy or DNA sequences) as a Markov process, Shannon defined entropy ($H$) to measure how much information is generated by such a process. The information source generates a series of symbols belonging to an alphabet with size $J$ (e.g. 26 English letters or 4 nucleotides). If symbols are generated according to a known probability distribution $p$, the entropy function $H(p_1, p_2, \ldots, p_J)$ can be evaluated. The units of $H$ are in bits, where one bit is the amount of information necessary to select one of two possible states or choices. In this section we describe several important concepts regarding the use of entropy in genomic sequence analysis.

**Entropy** is a measure of the average uncertainty of symbols or outcomes. Given a random variable $X$ with a set of possible symbols or outcomes $A_X = \{a_1, a_2, \ldots, a_J\}$, having probabilities $\{p_1, p_2, \ldots, p_J\}$, with $P(x = a_i) = p_i$, $p_i \geq 0$ and $\sum_{x \in A_X} P(x) = 1$, the Shannon entropy of $X$ is defined by

$$H(X) = \sum_{x \in A_X} P(x) \log_2 \frac{1}{P(x)} \tag{1}$$

Two important properties of the entropy function are: (a) $H(X) \geq 0$ with equality for one $x$, $P(x) = 1$; and (b) Entropy is maximized if $P(x)$ follows the uniform distribution. Here the uncertainty or surprisal, $h(x)$, of an outcome $(x)$ is defined by

$$h(x) = \log_2 \frac{1}{P(x)} \text{ (bits)} \tag{2}$$

For example, given a DNA sequence, we say each position corresponds to a random variable $X$ with values $A_X = \{A, C, G, T\}$, having probabilities $\{p_A, p_C, p_G, p_T\}$, with $P(x = A) = p_A$, $P(x = C) = p_C$, and so forth. Suppose the probability distribution $P(x)$ at a position of DNA sequence is

$$P(x = A) = 1/2; P(x = C) = 1/4; P(x = G) = 1/8; P(x = T) = 1/8.$$ 

The uncertainties (surprisals) in this case are $h(A) = 1$, $h(C) = 2$, $h(G) = h(T) = 3$ (bits). The entropy is the average of the uncertainties: $H(X) = \mathbb{E}[h(x)] = 1/2(1) + 1/4(2) + 1/8(3) + 1/8(3) = 1.75$ bits. In a study of genomic DNA sequences, Schmitt and Herzel (1997) found that genomic DNA sequences are closer to completely random sequences than to written text, suggesting that higher-order interdependencies between neighboring or adjacent sequence positions make little contributions to the block entropy.
The entropy (average uncertainty), $H$, is 2 bits if each of the four bases is equally probable (uniform distribution) before the site is decoded. The information content ($IC$) is a measure of a reduction in average uncertainty after the binding site is decoded. $IC(X) = H_{\text{before}} - H_{\text{after}} = \log_2|A_x| - H(X)$, provided the background probability distribution $P(\text{before})$ is uniform (Schneider, 1997a). If the background distribution is not uniform, the Kullback-Leibler distance (relative entropy) can be used (Stormo, 2000). The information content calculation needs to be corrected for the fact that a finite number of sequences were used to estimate the information content of the ideal binding site, resulting in the corrected $IC, R_{\text{sequence}}$ (Schneider et al., 1986). This measures the decrease in uncertainty before vs. after the macromolecule is bound to a set of target sequences. Positions within a binding site with high information are conserved between binding sites, whereas low information content positions exhibit greater variability. The $R_{\text{sequence}}$ values obtained precisely describe how different the sequences are from all possible sequences in the genome of the organism, in a manner that clearly delineates the conserved features of the site.

Relative entropy For two probability distributions $P(x)$ and $Q(x)$ that are defined over the same alphabet the relative entropy (also known as the Kullback-Leibler divergence or KL-distance) is defined by

$$D_{KL}(P \parallel Q) = \sum_{x \in A_x} P(x) \log \frac{P(x)}{Q(x)}$$

Note that the relative entropy is not symmetric: $D_{KL}(P || Q) \neq D_{KL}(Q || P)$; and although it is sometimes called the KL-distance, it is not strictly a distance (Koski, 2001; Lin 1991). Relative entropy is an important statistic for finding unusual motifs/patterns in genomic sequences (Durbin et al., 1998; Lawrence et al., 1993; Bailey and Elkan, 1994; Hertz and Stormo, 1999; Liu et al., 2002).

$R_{\text{sequence}}$ vs. $R_{\text{frequency}}$ The fact that proteins can find their required binding sites among a huge excess of non-sites (Lin and Riggs, 1975; von Hippel, 1979) indicates that more information is required to identify an infrequent site than a common binding site in the same genome. The amount of information required for these sites to be distinguished from all sites in the genome, $R_{\text{frequency}}$, is derived independently from the size and
frequency of sites in the genome. $R_{\text{frequency}}$, like $R_{\text{sequence}}$, is expressed in bits per site. $R_{\text{sequence}}$ cannot be less than the information needed to find sites in the genome. With few exceptions, it has been found that $R_{\text{sequence}}$ and $R_{\text{frequency}}$ are similar (Schneider et al. 1986). This empirical relationship is strongly constrained by the fact that all DNA-binding proteins operating on the genome are encoded in the genome itself (Kim et al. 2003).

**Molecular machines** are characterized by stable interactions between distinct components, for example, the binding of a recognizer protein to a specific genomic sequence. The behavior of a molecular machine can be described with information theory. The properties of molecular machine theory may be depicted on multiple levels: on one level, sequence logos, which describe interactions between the molecules (see Figure 1), are equivalent to transmission of information by the recognizer as a set of binary decisions; on another level, the information capacity of the machine, which represents the maximum number of binary decisions (or bits) that can be made for the amount of energy dissipated by the binding event; and finally, the relationship between information content and the energy cost of performing molecular operations (Schneider 1991, Schneider 1994). The molecular machine capacity is derived from Shannon’s channel capacity (Shannon, 1949). The error rate of the machine can be specified to be as low as necessary to ensure the survival of the organism, so long as the molecular machine capacity is not exceeded. Entropy decreases as the machine makes choices, which corresponds to an increase in information.

The second law of thermodynamics can be expressed by the equation: $dS \geq dQ/T$. The equation states that for a given increment of heat $dQ$ entering a volume at some temperature $T$, the entropy will increase $dS$ at least by $dQ/T$. If we relate entropy to Shannon’s uncertainty, we can rewrite the second law in the following form:

$$\varepsilon_{\text{min}} = \kappa_B T \ln(2) \leq -\frac{q}{IC} \quad \text{(joules per bit)}$$

(4)

where $\kappa_B$ is Boltzmann constant and $q$ is the heat. This equation states that there is a minimum amount of heat energy that must be dissipated (negative $q$) by a molecular machine in order for it to gain $IC = 1$ bit of information.
**Individual information** The information contained in a set of binding sites is an average of the individual contributions of each of the sites [Shannon 1948; Pierce 1980; Sloan and Wyner 1993; Schneider 1995]. The information content to each individual binding site sequence can be determined by a weight matrix so that the average of these values over the entire set of sites is the average information content [Schneider 1997].

The individual information weight matrix is:

\[
R_{iw}(b, l) = 2.0 - (-\log_2 (f(b, l) + e(n(l)))) \text{ (bits per base)}
\]  

(5)

in which \(f(b, l)\) is the frequency of each base \(b\) at position \(l\) in the binding site sequences; \(e(n(l))\) is a correction of \(f(b, l)\) for the finite sample size (\(n\) sequences at position \(l\)) [Schneider et al 1986]. The \(j^{th}\) sequence of a set of binding sites is represented by a matrix \(s(b, l, j)\), which contains 1’s in cells from base \(b\) at position \(l\) of a binding site and zeros at all other matrix locations. The individual information of a binding site sequence is the dot product between the sequence and the weight matrix:

\[
R_i(j) = \sum_l \sum_{b=a}^t s(b, j, j)R_{iw}(b, l) \text{ (bits per site)}
\]  

(6)

**B. Applications**

**Displaying sequence conservation** Sequence logos, which display information about both consensus and non-consensus nucleotides, are visual representations of the information found in a binding site (an example is shown in Figure 1). This is the information that the decoder (i.e. a binding protein) uses to evaluate potential sites in order to recognize actual sites. The calculation of sequence logos uses the assumption that each site is evaluated independently, i.e. that there is no correlation between a change in nucleotide at one position with another position, which is reasonable for most genomic sequences (Schmitt and Herzel 1997). An advantage of the information approach is that the sequence conservation can be interpreted quantitatively. \(R_{\text{sequence}}\), which is the total area under the sequence logo and measures the average information in a set of binding
site sequences, is related to the specific binding interaction between the recognizer and the site. $R_{sequence}$ is an additive measure of sequence conservation; thus it is feasible to quantitatively compare the relative contributions of different portions within the same binding site.

Structural features of the protein-DNA complex can be inferred from sequence logos. When positions with high information content are separated by a single helical turn (10.4 base pairs), this suggests that the protein makes contacts across the same face of the double helix. Sequence conservation in the major groove can range anywhere between 0 and 2 bits depending on the strength of the contacts involved, and usually correlates with the highest information content positions (Papp et al. 1993). Minor groove contacts of B-form DNA allow both orientations of each kind of base pair so that rotations about the dyad axis cannot easily be distinguished, hence a single bit is the information content in native B-form DNA (Schneider 1996). Higher levels of conservation for bases within the minor groove indicate that these positions are accessed protein distortion of the helix, i.e. bending accompanied by base pair opening and flipping (Schneider, 2001).

**Visualizing individual binding site information**

Because sequence logos display the average information content in a set of binding sites, they may not accurately convey protein-DNA interactions with individual DNA sequences, especially at highly variable positions within a binding site. The walker method (Schneider 1997b) graphically depicts the nucleotide conservation of a known or suspected site compared to other valid binding sites defined by the individual information weight matrix (Schneider 1997a). Walkers apply to a single sequence (rather than a set of binding sites); only a single letter is visualized at each position of the binding site (Fig. 2). The height of the letter, which is in units of bits, represents the contribution of that nucleotide at each position in the binding site by the information weight matrix, $R_{iw}(b,l)$. Evaluation of the $R_i$ value at each position in a genomic DNA sequence is equivalent to moving the walker along that sequence. Walkers are displayed for sequences with
positive $R_i$ values, since these are more likely to be valid binding sites (see equation 4 and discussion below). Sequence walkers facilitate visualization and interpretation of the structures and strengths of binding sites in complex genetic intervals and can be used to understand the effects of sequence changes (see below), and engineer overlapping or novel binding sites.

**Mutation and polymorphism analysis** Because the relationship between information and energy can be used to predict the effects of natural sequence variation at these sites, phenotypes can be predicted from corresponding changes in the individual information contents ($R_i$, in bits) of the natural and variant DNA binding sites (Rogan et al., 1998; see g206101). For splice site variants, mutations have lower $R_i$ values than the corresponding natural sites, with null alleles having values at or below zero bits (Equation 4; Kodolitsch et al., 1999). The decreased $R_i$ values of mutated splice sites indicate that such sites are either not recognized or are bound with lower affinity, usually resulting in an untranslatable mRNA. Decreases in $R_i$ are more moderate for partially functional (or leaky) mutations that reduce but do not abolish splice site recognition and have been associated with milder phenotypes (Rogan et al 1998). The minimum change in binding affinity for leaky mutations is $\geq 2^{\Delta R_i}$ lower fold than cognate wild type sites. Mutations that activate cryptic splicing may decrease the $R_i$ value of the natural site, increase the strength of the cryptic site, or concomitantly affect the strengths of both types of sites (see Figure 2). Non-deleterious changes do not alter the $R_i$ value of splice sites significantly (Rogan and Schneider, 1995). Increases in $R_i$ indicate stronger interactions between protein and cognate binding sites.
Information evolution  How do genetic systems gain information in a binding site of genomic DNA sequence by evolutionary processes? Schneider (2000) proposed an answer to this question. Given a binding site for an artificial protein, his simulation begins with zero information and, as in naturally occurring genetic systems, the information measured in the fully evolved binding sites ($R_{\text{sequence}}$) is close to that needed to locate the sites in the genome ($R_{\text{frequency}}$).

Model refinement  Information models based on small numbers of proven binding sites may fail to detect valid binding sites and tend to predict $R_i$ inaccurately. Iterative selection of functional binding sites has been used to optimize (Lund et al., 2000) and to introduce bias (Shultzaberger and Schneider, 1999) into the frequencies of each nucleotide in computing the information theory-based weight matrices of binding sites. Significant differences between information weight matrices have been determined from their respective evolutionary distance metrics (for example, see Shultzaberger et al., 2001). The effects of model refinements can be monitored by comparing the genome scan results for pairs of successive information weight matrices based on additional binding sites (Gadiraju et al., 2003) Other potential applications include determining the locations of overlapping binding sites recognized by different proteins and comparisons of binding sites detected with information models of orthologous proteins from different species.

Regulatory binding sites  Information theory-based models have been used in searching for regulatory sites in genomic DNA or RNA sequences of prokaryotes (Hengen et al., 1997) and eukaryotes (see g402310). The binding sites in prokaryotes include the ribosome binding sites (Shultzaberger et al., 2001), T7 promoters, plasmid replication initiator protein binding sites (Papp et al., 1993) and binding sites for repressors and polymerases (Schneider et al., 1986) and cyclic AMP receptor protein (Stormo and Fields, 1989) in E. coli.

A bipartite pattern is an independent functional unit on the upstream or downstream side of a regulated gene that is recognized by a protein binding complex such as a
heterodimer. A model built to simulate a bipartite pattern in genomic sequences has
left and right motif sub-models, plus an associated gap penalty function, \( g(d) \) defined as
\(-\log(n(d)/n)\), where \( n(d) \) is the number of sites with gap size \( d \). Shannon’s entropy can
be used to evaluate such sites by calculating the total information content, \( IC \), given as
\[
IC = IC(\text{left} \mid d) + IC(\text{right} \mid d) - g(d)
\]
where
\[
IC(m \mid d) = \sum_{i=1}^{J_m} \left( \log_2 |A_{x_i}| - H_{ml}(X) \right), \quad m \in \{\text{left, right}\}
\]
\[
H_{ml}(X) = \sum_{x \in A_X} P_{ml}(x) \log_2 \frac{1}{P_{ml}(x)}, \quad A_X = \{A,C,G,T\}
\]
Here \( J_m \) is the width of motif \( m \), and \( P_{ml}(x) \) is the probability of \( x \) at position \( l \) given
motif \( m \). The left and right motifs are not allowed to be overlapping and the gap size \( d \)
is set to a limited range \([d_{\text{min}}, d_{\text{max}}]\) based on empirical observations. The goal is to
maximize the total information content which can be reduced to minimize the total
Shannon’s entropy. We used a Monte Carlo strategy to greedily search the multiple
alignment space and find an optimal solution to the bipartite pattern search problem (Bi et al., 2004).

We developed a new method for the bipartite cis-regulatory pattern discovery based on
minimizing entropy, and applied the method to a set of known PXR/RXR\( \alpha \) binding sites
in the human genome. The PXR/RXR\( \alpha \) heterodimer binding controls the expression of
coo-regulated genes such as \( CYP3A4 \), which is involved in detoxification of drugs and
xenobiotics (see g206211). This work is an extension of Shultzaberger et al., 2001.
Using the assumption that two proteins (i.e. PXR and RXR\( \alpha \)) cooperatively bind to the
bipartite site with constrained spacers, we built models for different motif widths and
validated them based on the relative binding strength of the a series of test sequences.
The results supported our hypothesis that PXR and RXR\( \alpha \) transcription factors
cooperatively bind to two adjacent motifs with variable spacing (Bi et al 2004).

Genome-wide analyses Information weight matrices of binding sites can be developed
directly from validated sets of binding sites extracted from genome sequences provided
that the locations of sequence features are accurately annotated. As this is not always the
case, we built a genome-wide human splice junction database by initially extracting the coordinates and sequences of donor and acceptor splice regions from both strands of the human genome reference sequence (Rogan et al., 2003). After redundant sites were eliminated, the splice site database consisted of 170,144 acceptor and 170,450 donor sites. The information weight matrix was recomputed after each of iteration and scanning of the resultant set of sites. Successive models iteratively utilized the modified matrix by excluding sites with negative $R_i$ values. After 8 cycles of refinement, the final models were then defined by 108,079 unique acceptor sites and 111,772 donor sites (Sequence logos of model sites are shown in Figure 1).

The average information contents of the acceptor and donor sites are respectively, 9.8 bits/site and 8.1 bits/site. These values are similar to those previously reported by Schneider and Stephens (1992), i.e. 9.35 bits for acceptors and 7.92 bits for donors, which were based on about 65-fold fewer splice sites. Individual splice site strengths in the genome have an approximately Gaussian distribution.

C. Prospects for information theory based analyses of genomic sequences

As the functions for regulatory and expressed nucleic acid sequences are elucidated, it is becoming evident that multiple protein components catalyze these processes. Modeling such molecular machines by information theory will require the development of procedures that account for cooperative and interdependent binding events between two or more recognizers. Frameworks for building multipartite information models will therefore have to incorporate corrections for overlapping sites and mutual information.

There are opportunities to enhance currently available genomic applications by scaling currently available software for information theory analyses (Gadiraju et al. 2003) to investigate genome annotation. For example, changes in IC contents due to mutation may be of assistance in prioritizing single nucleotide polymorphisms for functional analyses. It is also possible that comparative genomic analyses of binding sites with orthologous DNA recognition domains from multiple
species may reveal possible identities of functionally-analogous regulatory sequences in these systems.

Medical genetic applications of information theory based binding site models are a promising avenue to improve diagnosis and prognosis of disease causing mutations (see g106404). Accurate models will be required for use of information theory in a clinical setting. To calibrate individual information measures of protein-nucleic acid binding with the thermodynamic properties of these complexes will require more comprehensive models, ie. based on larger numbers of binding sites spanning a wide range of binding affinities.
Acknowledgements
This work was supported by NIEHS (Grant ES10855). We thank Dr. Thomas Schneider for valuable suggestions and comments.

Related Articles

References


Figure Captions

Figure 1. Examples of sequence logos. Models of human (A) 108,079 acceptor and (B) 111,772 donor splice sites derived from both strands of the human genome reference sequence (April, 2003) are shown. A sequence logo visually represents the sequence conservation among a common set of recognition sites, with the height of each nucleotide stack corresponding to the the average information content at that position. The height of each nucleotide is proportional to its frequency. Sampling error is indicated by error bar at the top of each stack. The zero coordinate represents the intronic position immediately adjacent to the splice junction. The average information contents ($R_{\text{sequence}}$) of the acceptor and donor sites are respectively, 9.8118 ± 0.0001 bits/site and 8.12140 ± 0.00009 bits/site.

Figure 2. Examples of sequence walkers. A synonymous C>T substitution at codon 608 activates a cryptic donor splice site in exon 11 the LMNA gene in patients with Hutchinson-Gifford progeria (Ericksson et al. 2003). The walker, shown below the sequence, indicates a pre-existing 8.7 bit cryptic site that is strengthened by the mutation to 10.2 bits ($\geq 2.8$ fold). The height and orientation of each nucleotide correspond to contribution that nucleotide makes to the overall information content in the binding site. The green rectangle indicates the location of valid splice site ($R_i > 0$) and delineates the scale displayed; the lower and upper limits shown are, respectively, -4 bits and +2 bits. Sequence coordinates are from GenBank accession L12401 (4277C>T).
Figure 1

(A) 

(B)
Figure 2

Normal sequence of LMNA

Donor 8.7 bits @4272

G608G (GGC>GCT)

Donor 10.2 bits @4272