



NRC Publications Archive Archives des publications du CNRC

Positional Weight Matrix as a Sequence Motif Detector Pan, Youlian; Phan, Sieu

Publisher's version / Version de l'éditeur:

Oligonucleotide Array Sequence Analysis, pp. 421-440, 2008

NRC Publications Record / Notice d'Archives des publications de CNRC:

<https://nrc-publications.canada.ca/eng/view/object/?id=ea9161ea-ede6-4c8c-8794-afca7eac4710>

<https://publications-cnrc.canada.ca/fra/voir/objet/?id=ea9161ea-ede6-4c8c-8794-afca7eac4710>

Access and use of this website and the material on it are subject to the Terms and Conditions set forth at

<https://nrc-publications.canada.ca/eng/copyright>

READ THESE TERMS AND CONDITIONS CAREFULLY BEFORE USING THIS WEBSITE.

L'accès à ce site Web et l'utilisation de son contenu sont assujettis aux conditions présentées dans le site

<https://publications-cnrc.canada.ca/fra/droits>

LISEZ CES CONDITIONS ATTENTIVEMENT AVANT D'UTILISER CE SITE WEB.

Questions? Contact the NRC Publications Archive team at

PublicationsArchive-ArchivesPublications@nrc-cnrc.gc.ca. If you wish to email the authors directly, please see the first page of the publication for their contact information.

Vous avez des questions? Nous pouvons vous aider. Pour communiquer directement avec un auteur, consultez la première page de la revue dans laquelle son article a été publié afin de trouver ses coordonnées. Si vous n'arrivez pas à les repérer, communiquez avec nous à PublicationsArchive-ArchivesPublications@nrc-cnrc.gc.ca.





National Research
Council Canada

Conseil national
de recherches Canada

Institute for
Information Technology

Institut de technologie
de l'information

NRC - CNRC

Positional weight matrix as a sequence motif detector *

Pan, Y., Phan, S.
October 2008

* published in Nova Science Publishers, Inc., October 2008. pp. 421-440.
ISBN: 978-1-60456-542-3. Editors: M.K. Morettie and L.J. Rizzo. NRC
50391.

Copyright 2008 by
National Research Council of Canada

Permission is granted to quote short excerpts and to reproduce figures and tables
from this report, provided that the source of such material is fully acknowledged.

Chapter 14

POSITIONAL WEIGHT MATRIX AS A SEQUENCE MOTIF DETECTOR*

Youlian Pan and Sieu Phan

Institute for Information Technology, National Research Council Canada
1200 Montreal Road, Ottawa, Ontario, Canada K1A 0R6

ABSTRACT

In biological sequence research, the positional weight matrix (PWM) is often used for motif signal detection. A set of experimentally verified oligonucleotides known to be functional subsequences, which can be bound by a transcription factor (TF), as translational initiation sites or pre-mRNA splicing sites, are collected and aligned. The frequency of each nucleotide A, C, G, or T at each column of the alignment is calculated in the matrix. Once a PWM is constructed, it can be used to search from a nucleotide sequence for the subsequences that possibly perform the same function. The match between a subsequence and a PWM is usually described by a score function, which measures the closeness of the subsequence to the PWM as compared with the given background. However, selection of threshold scores that legitimately qualify a functional subsequence has been a great challenge. Many laboratories have attempted tackling this problem; but there is no significant breakthrough so far. In this chapter, we discuss the characteristics of a PWM and factors that affect motif predictions and propose a new score function that is tied into information content and statistical expectation of a PWM. We also apply this score function in the PWMs from public databases and compare it favorably with the broadly used Match method.

1. INTRODUCTION

The term motif is broadly used to describe a conserved pattern, which include a conserved domain (Pérez-Arellano et al., 2007), a functional structure (Schreiter and Drennan, 2007; Kiedziarska et al., 2007), a specific architecture in biological networks (Zhu et al., 2007) and a string pattern in protein or nucleotide sequences (Pan, 2006). A nucleotide sequence motif often refers to a short functional pattern in the DNA or RNA sequence and is

* This is publication NRC XXXXX of the National Research Council Canada

often used to characterize the interaction between a DNA and a protein, such as the binding sites of nucleases and transcription factors. Others at RNA levels are involved in translation, such as ribosome binding, and mRNA processing, such as splicing between exons and introns. Some motifs are involved in transcription termination, such as microRNAs binding. These motifs are usually conserved in the nucleotide sequence pattern.

Many proteins are able to bind to a DNA subsequence with alternative nucleotides at one or more positions in a motif. Among a set of functional subsequences that are known to have the same function, the most dominant ones are called *consensus* for that function. The term of *consensus* has later been relaxed to include degenerate, yet functional subsequences that match all example sites closely, but not necessarily exactly. A consensus describes statically the interaction between a protein and a DNA, but is unable to describe precisely the significance of one nucleotide over the others at certain position of a motif. In this regard, a positional weight matrix (PWM) is often more powerful to express a motif (Stormo, 2000). In the literature, a PWM is also referred as frequency matrix, position specific weight matrix, and positional probability matrix (see Pan, 2006 and refs there in).

A PWM is a probabilistic description of a binding motif for a given protein. Due to its nature of pattern conservation, many similar subsequences could be discovered through computational pattern recognitions. A significant number of such subsequences have been experimentally validated in the biology laboratories across the world. Through the multiple alignments of a set of short nucleotide sequences that are known to perform the same function, probability of each nucleotide appears at certain position of the motif is derived simply by counting its occurrences and can be expressed either in the form of motif logo (Schneider and Stephens, 1990) or a probability matrix (Stormo, 2000).

Over the past two decades, many computational approaches are developed to discover conserved motifs with certain degree of success. Computational motif discovery process can be considered in two categories, unsupervised *de novo* motif discovery and supervised known motif prediction (Pan, 2006). In the unsupervised *de novo* motif discovery processes, numerous DNA subsequences of a given length w are locally aligned. The idea behind these methods is that each sequence in the input dataset contains one or more subsequences that could be instances of a conserved motif, but there is no prior knowledge with regard to their binding agent, motif pattern structure, or probability of each nucleotide at certain position. The conserved motifs are iteratively evolved through various optimization algorithms.

The optimization algorithms in *de novo* motif discovery can be subdivided into three distinct approaches: *deterministic optimization*, *stochastic optimization*, and *enumeration* (Pan, 2006, D'haeseleer, 2006). Information content is one of the objective functions applied to measure the quality of a local alignment (see Section 2). Optimization of the information content of a matrix for a fixed number of occurrences has been widely used in matrix-based motif discovery.

While performing local alignment, *deterministic optimization* methods simultaneously optimize a positional weight matrix describing a motif and probabilities of its associated sites. Expectation maximization (EM, Lawrence and Reilly, 1990) is one of such methods. The EM algorithms are named for their two iterative steps, expectation and maximization, which are repeated until a convergence criterion is satisfied. The expectation step evaluates the likelihood of each subsequence of length w to fit the PWM profile with respect to the background of the input sequence. A likelihood value Z_{ij} is computed for each position i of input sequence j . The maximization step puts the best subsequence (with maximum likelihood

value) from each input sequence j and builds a new alignment PWM profile. A popular implementation of these algorithms is in MEME software (Bailey and Elkan, 1994). However, the result of the EM algorithm is sensitive to the initial subsequence selection from each input sequence and, therefore, it could be trapped by local maxima.

Stochastic optimization applies stochastic sampling instead of exhaustive survey of all subsequence of length w . Initially, one such subsequence is selected randomly from each of the k input sequences and a $4 \times w$ PWM profile is built on subsequences from $k-1$ input sequences. Similar to the EM algorithm, the likelihood value of each possible w -mer (Z_i) from the remaining sequence is evaluated to represent how well the w -mer fits to the model. A w -mer is sampled stochastically according to the relative likelihood values ($Z_i / \sum_i Z_i$,

Lawrence *et al* 1993). This relative likelihood value indicates that a w -mer that fits well (with high Z_i value) is more likely sampled. This procedure is repeated for each sequence until all input sequences have been sampled. The whole procedure is repeated until a convergence criterion is satisfied. Theoretically, the relative information is maximized only after an infinite number of cycles. Practically, this algorithm often converges very quickly (Zhang, 2002). Similar to the EM approach, stochastic sampling could be locked to a local maximum. Inserting another step into the algorithm solves this problem. After a specific number of cycles, one can automatically shift (called a phase shift) all aligned subsequences to the left or right by a certain number of nucleotides and a $4 \times w$ PWM profile is built after the shift. The iterative sampling is repeated. This approach has been implemented in the Gibbs Sampling family of tools that are very popular in motif discovery (Liu *et al* 1995; Hughes *et al*, 2000; Liu *et al.*, 2001; Thijs *et al*, 2001; Thompson *et al.*, 2003, 2007; Favorov *et al*, 2005).

Enumerative algorithms search exhaustively for all possible combinations of nucleotides and select statistically the top few overrepresented motifs from a set of input sequences. Subsequently, motif discovery is treated as a feature selection problem; a motif is treated as a feature of input sequences that discriminate the potential motifs from background sequences (Sinha, 2003). Instead of counting the frequency of overall occurrences of the motifs in a set of sequences, information about motif distribution in individual sequences is used to evaluate motif overrepresentation. This kind of methods has been considered too slow due to the fact that their time complexity grows exponentially with the length of the motif. Given a motif length w , these methods need to evaluate 4^w candidate patterns before an optimal solution is found. However, by indexing the input sequences with a suffix tree, the execution time becomes exponential with respect to the number of allowed substitutions, which is usually small, instead of motif length (Pavesi *et al*, 2004). These methods are guaranteed to find the globally most overrepresented motifs (Sinha and Tompa, 2002). Some of these methods compare favorably with others with regard to accuracy (Tompa *et al.*, 2005).

Biological methods for motif discovery also become significant and more reliable. These include “photofootprinting” (Becker and Wang, 1984), chromatin immunoprecipitation (ChIP, Solomon and Varshavsky, 1985), and a combination of a modified version of the ChIP procedure (Orlando, 2000) with DNA microarray technology (chip) called ChIP-chip (Horak and Snyder, 2002), protein binding microarray (Bulyk *et al*, 2001; Mukherjee *et al.*, 2004) and others reviewed in Pan (2006). These methods brought the motif discovery to a new stage and many computationally discovered motifs are validated.

Additionally, motif discovery are being advanced through phylogenetic footprinting. Mainly, two phylogenomic approaches have been considered. The first is to find motifs that

are common from multiple orthologous sequences. Footprinter (Blanchette and Tompa, 2002) and OrthoMEME (Prakash et al., 2004) are two examples in this category. The second approach is to globally align the promoter sequences of the orthologous genes and then identify conserved windows. For example, EMnEM (Moses et al, 2004), CompareProspector (Liu et al., 2004), PhyME (Sinha et al, 2004), and PhyloGibbs (Siddharthan et al, 2005), Phylo-HMMs (Siepel and Haussler, 2005), phylogenetic Gibbs sampler (Newberg et al, 2007) and others reviewed in Pan (2006) falls into this category.

Through both computational and biological approaches, or through a combination of both approaches, many motif profiles in the form of PWM became known and validated. Many (trained) PWMs are available in public databases, such as TRANSFAC (Matys et al, 2006) and JASPAR (Sandelin et al, 2004). Such PWMs are extensively used for *supervised* prediction of putative motifs. Many PWM-based motif prediction tools have been developed over the past decades. These include Match (Kel et al., 2003), P-Match (Chekmenev, 2005), MatInspector (Cartharius et al, 2005) and GAPWM (Li et al, 2007).

The main challenge in *supervised* PWM-based motif prediction methods is the objective score function and determination of a threshold score. The score functions usually depend on PWM parameters such as its length and information content. Therefore, a threshold scores that legitimately qualify a functional motif is very hard to select without subjectivity. This chapter is to discuss the characteristics of a PWM, the existing score functions, the factors that affect motif predictions and finally propose a new score function that is tied into information content and statistical expectation of a PWM. We also apply this score function in the PWMs from public databases and compare it with the broadly used Match method.

In the following sections, we describe the existing objective functions, their strengths and weaknesses in Section 2, and ways to improve the score functions based on the existing ones in Section 3. We apply the proposed score function in two datasets in Section 4, discuss the result in Section 5, and finally conclude in Section 6.

2. EXISTING OBJECTIVE (SCORE) FUNCTIONS

Before describing a score function, it is necessary to review the information theory (Schneider et al, 1986) that is the backbone of most score functions. Information theory describes the uncertainty of a discrete random variable $X = \{x_1, x_2, \dots, x_k\}$ as

$$H(X) = \sum_{b=1}^k p_b \log_2(1/p_b) = -\sum_{b=1}^k p_b \log_2(p_b) \text{ (bit)} \quad (1),$$

where p_b is the probability of x_b and $\sum_{b=1}^k p_b = 1$. One bit of information resolves the uncertainty of choice between 2 equally likely symbols. For a nucleotide sequence, there are 4 bases ($X = \{A, C, G, T\}$, $k = 4$). The maximum uncertainty corresponds to the case that all 4 bases have equal probability, $p_b = 0.25$; thus the maximum uncertainty is 2 bits. If only one base, say G, appears at certain position of a PWM, we have $p_A = 0$, $p_C = 0$, $p_G = 1$, $p_T = 0$. The uncertainty of that column is 0 bit ($0 \log 0 = 0$ and $1 \log 1 = 0$). Therefore the uncertainty value for each column of a PWM is between 0 and 2.

The information content at column i of a PWM can be measured by a Kullback-Leibler (Kullback and Leibler, 1951) distance (or relative entropy), which is defined as

$$I_i = \sum_{b=1}^4 p_{i,b} \log_2 \frac{p_{i,b}}{f_b} \quad (2)$$

or

$$I_i = -\sum_{b=1}^4 p_{i,b} \log_2 f_b + \sum_{b=1}^4 p_{i,b} \log_2 p_{i,b} \quad (3),$$

where $p_{i,b}$ is the probability of nucleotide b found at position i of the site, and f_b is the probability of nucleotide b on sequences other than the sites (i.e. background). With the assumption of equally probable background model in a DNA sequence, Equation (3) becomes

$$I_i = -\log_2 f_b \left(\sum_{b=1}^4 p_{i,b} \right) + \sum_{b=1}^4 p_{i,b} \log_2 p_{i,b} = -(-2)(1) + \sum_{b=1}^4 p_{i,b} \log_2 p_{i,b}.$$

This special version of Equation (3) is the traditional Shannon distance (Shannon, 1948)

$$I_i = 2 + \sum_{b=1}^4 p_{i,b} \log_2 p_{i,b} \quad (4).$$

Information content of a PWM is the summation of information on each column:

$$I_{matrix} = \sum_{i=1}^w I_i \quad (5).$$

Based on information content in each column of a PWM, the PWM can also be expressed graphically as sequence logo (Schneider and Stephens, 1990). An objective function of each PWM-based motif discovery method is to discriminate a subsequence between a background model and the PWM model. There are mainly six categories of the objective functions, *log-odd score (also called log-likelihood ratio)*, *MAP score*, *relative information*, *Z score*, *specificity score*, and *positional bias scores*.

The *log-odd score* is to measure the relative likelihood of a subsequence s being an instance of the PWM against the given background model. It is the summation of the score on each base and can be expressed as:

$$V_s = \log_2 \left(\prod_{i=1}^w \frac{p(b_i)}{f(b_i)} \right) = \sum_{i=1}^w \log_2 \left(\frac{p(b_i)}{f(b_i)} \right) \quad (6).$$

The log-odd score has been extensively used in various domains. It is the core of Viterbi algorithm that is extensively used in sequence alignment, hidden Markov model (Krogh et al,

1994) and many motif finding tools. However, the log-odd score is dependent on length of the motifs and PWM models.

The *MAP* (*maximum a posteriori probability*) score combines the information content of the PWM and the rareness of PWM with regard to the background model (Liu et al, 2002):

$$MAP = \frac{\log_2(n)}{w} \left[\sum_{i=1}^w \sum_{b=1}^k p_{i,b} \log_2(p_{i,b}) - \frac{1}{n} \sum_{j=1}^n \log_2 f(s_j) \right] \quad (7),$$

where n is the number of sites used to construct the PWM and s_j is an individual site

The *relative information* measures the relative information of a site S based on PWM

$$Score(S) = \sum_{i=1}^w I_i p_{i,b_i} \quad (8),$$

where b_i is nucleotide i in the site. The *relative information* score does not consider background information and applied by MatInspector (Cartharius et al., 2005), Match (Kel et al., 2003) and P-Match (Chekmenov et al., 2005). These applications take a max score and then take a ratio between a site score and the max score:

$$R = \frac{Score(S)}{Score(max)} \quad (9),$$

but Match (Kel et al., 2003) scales between the maximum score and minimum score and then takes the ratio

$$R(Match) = \frac{Score(S) - Score(min)}{Score(max) - Score(min)} \quad (10).$$

The *Z-score* measures the significance of a prediction based on its over-representation and is expressed as

$$Z(s) = \frac{obs(s) - E(s)}{\sigma(s)} \quad (11),$$

where $obs(s)$ is the predicted number of PWM instances s , $E(s)$ is the expected number of s in the background model and $\sigma(s)$ is the standard deviation. However, the statistical over-representation as measured by *Z-score* does not necessarily mean binding preference (Li and Tompa, 2006).

The *group specificity score* measures the goodness of a given motif that is localized to the set of input sequences as compared to the background. The rationale is that the true motif is a subsequence that is clearly more frequent in the selected sequence set than in all sequences. This score is used in AlignACE (Hughes et al, 2000).

The *sequence specificity score* measures the likelihood of the set of input sequence potentially contains a functional motif. A prediction having multiple motifs in one sequence is less significant than a prediction having a balanced number of motifs in each sequence (Li and Tompa, 2006). This score is applied in ANN-Spec (Workman and Stormo, 2000) and Weeder (Pavesi et al, 2004).

Due to the uneven distribution of certain motifs (Pan et al, 2004, Smith et al, 2007), *positional bias* (Hughes et al, 2000) and *local positional bias* (Friberg et al, 2005) scores are being used to measure the concentration of the motifs within certain local regions of the input sequence set.

Most of the above described objective functions are exclusively used for finding a PWM representing a motif, such as *MAP score* and *group specificity*. Great effort has been placed in discovering PWMs through both biological and computational approaches and hundreds of known PWMs became available. The question is how to make use of these resources in predicting new instances of known PWMs. There is always an issue of a threshold selection for each PWM in predicting the new instance of the PWM. In Match and P-Match, three threshold values (minimum false positive value (minFP), minimum false negative value (minFN), and minimum of FP+FN) are generated for each PWM to satisfy users with different perspectives based on the sequences used to construct the PWM. This chapter investigates this problem by incorporating various sources of available information.

3. PROPOSAL OF AN INTEGRATED APPROACH THAT HAS THREE COMPONENTS

3.1. The Log Odd Score

The log odd score as described by Equation (6) is motif length dependent. For a consensus motif c of the PWM, the log-odd score

$$V_c = \sum_{i=1}^w \log_2 \left(\frac{p(b_{c,i})}{f(b_{c,i})} \right) \quad (12).$$

V_c is the best possible log-odd score of the PWM. For a subsequence s , $V_s = V_c$ means s is a best fit with the PWM and most likely a TFBS that the PWM specifies, while a $V_s = 0$ means s is identical to the background and very unlikely to be a TFBS that the PWM specifies. Thus, we defined the goodness (G , Pan and Phan, 2008) of fit between s and the PWM as:

$$G = \begin{cases} V_s / V_c, & \text{while } V_s \geq 0 \\ 0, & \text{while } V_s < 0 \end{cases} \quad (13).$$

The value of G is between 0.00 and 1.00 since $V_s \leq V_c$ and is independent of pattern length. From the G value, one could easily tell the confidence level of the putative TFBS that is found by a PWM regardless of the length of the subsequence.

3.2. Statistical Expectation

For a given PWM of length w , the theoretical number of motif variants is 4^w based on background model. Each motif variant has a probability of $4^{-w} \times (L - w + 1)$ to appear in a sequence S of length L . We can enumerate each motif variants in the PWM space and calculate its log odd-score and G value. At a given threshold, assume n motif variants satisfy the threshold, thus the probability of the PWM appearing on S is

$$p = 4^{-w} \times n(L - w + 1) \quad (14).$$

3.3. Information Content of the PWM

MAP score measures relative information content as compared with background and has been widely used and maintains a good balance between PWM specificity and diversity (Liu et al., 2002). However, the MAP score, as we can see from Equation (7), takes average information of all columns in the PWM. This simplistic averaging effect implicitly penalizes longer PWMs that usually have higher proportion of degraded columns that have lower information content. Overall information content on the other hand, as described by Equation (5), captures all information in the PWM.

3.4. Proof of Concept

For proof of concept, we retrieved 107 PWMs from JASPAR core database (Sandelin et al., 2004) on September 19, 2007. Since these PWMs are trained from different organisms ranging from yeast, insects to human and other mammals, we set the background probability as default ($f_b = 0.25$, $b = \{A, C, G, T\}$) and calculated best log-odd score, statistical expectation at different thresholds of G , MAP score, and information content of the PWM.

For a given G threshold g , we are interested to know the statistical expectation of finding a motif variant having a G value higher than g . This probability is called p_value associated with the given g . Theoretically, we calculate the probability density function, $e(g)$, over the entire g span (from 0 to 1). It is established by counting the number of motif variants found in a given interval g . The probability distribution function, $E(g)$, is then determined by

$$E(g) = \int_0^g e(\tau) d\tau \quad (15).$$

And the p_value is determined by

$$p_value(g) = 1 - E(g) \quad (16).$$

Figure 1 shows the p -value function of the human PWM PAX2_01 (from TRANSFAC). With knowledge of p _value, we can now select a proper threshold with a desired level of confidence. This in turn suggests that for every PWM, we establish an accompanying p _value table. The G threshold can be selected from the table according to the desired confidence level.

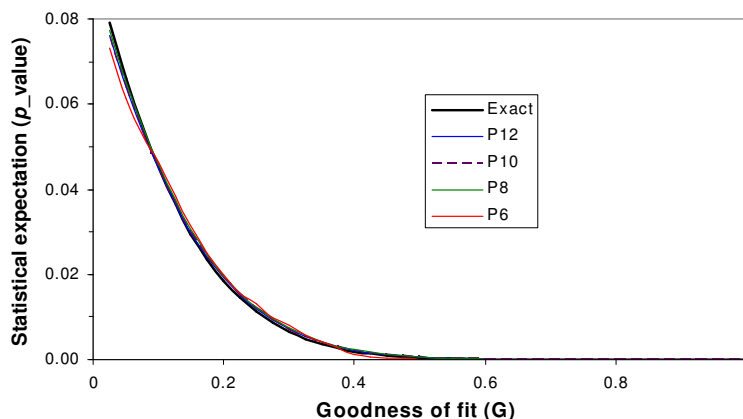


Figure. 1. p _value function of human PWM PAX2_01. Exact: all instances in the PWM space are evaluated, Px: 4^x instances from the PWM space are sampled.

The establishment of the p -value function for a lengthy PWM is computationally expensive. For example, a PWM of 24 columns would take more than 100 days to complete the calculation under the current state of technology. To alleviate this problem, we devised a random sampling scheme to perform the calculation. Intuitively,

$$e(g) = \lim_{n \rightarrow N} e_n(g) \quad (17),$$

where n is the number in the sampling subspace from the entire PWM space of $N = 4^w$ (w is the length of PWM) and $e_n(g)$ is the density function obtained by evaluating the G value of Equation (13) over the selected n variants. We verified that our random sampler and ensured distinct motif variants with no single repeat within the PWM space before it was used for random sampling. The p _value functions based on random sampling were verified to be technically identical as if the entire PWM space is sampled as long as sampling size was over 4^{10} (Fig. 1). It is feasible to perform exact calculation for a PWM of $w \leq 18$. For a PWM of $w > 18$, we applied the random sampling scheme to perform the computation.

The result indicates that the statistical expectation is highly correlated ($p < 0.0001$) with PWM length (w), information content, and the best log-odd score (Table 1). It appears that MAP score is not able to reveal the statistical expectation of a PWM since MAP score implicitly penalized longer PWM as described above. In this regard, we conducted the correlation analysis for the 19 PWMs that have equal length of 10. The correlation (MAP for $L=10$) appears to increase as result of this selection, but still not comparable with others such as best log-odd score, information content and PWM length. Additionally, in order to eliminate the effect of PWM length, we also calculated the cumulative information of 5 columns in the PWM that have the highest information content and named it as *core*

information and analyzed its correlation with p -value regardless of the length of PWM. This correlation appears to be significant at lower G threshold and is not helpful in selection of a threshold value. Thus, the core information is not considered further. At a given length of PWM, the p -values significantly correlated with threshold G values (Fig. 2).

Table 1. Correlation between the statistical expectations values and various properties of the 107 PWMs from JASPAR core database at a given threshold of G value.

$G >$	0	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9	0.95
Length	<u>0.57</u>	<u>0.56</u>	<u>0.65</u>	<u>0.71</u>	<u>0.77</u>	<u>0.83</u>	<u>0.88</u>	<u>0.91</u>	<u>0.95</u>	<u>0.97</u>	<u>0.98</u>
Information	<u>0.93</u>	<u>0.95</u>	<u>0.97</u>	<u>0.98</u>	<u>0.99</u>	<u>0.98</u>	<u>0.96</u>	<u>0.93</u>	<u>0.88</u>	<u>0.83</u>	<u>0.80</u>
Core Info	<u>0.73</u>	<u>0.72</u>	<u>0.70</u>	<u>0.66</u>	<u>0.61</u>	<u>0.53</u>	<i>0.46</i>	<i>0.39</i>	<i>0.30</i>	<i>0.22</i>	<i>0.19</i>
Best Log-odd	<u>0.78</u>	<u>0.81</u>	<u>0.84</u>	<u>0.89</u>	<u>0.93</u>	<u>0.97</u>	<u>0.99</u>	<u>1.00</u>	<u>0.99</u>	<u>0.98</u>	<u>0.96</u>
MAP score	0.10	0.09	0.08	0.05	0.00	0.12	0.12	0.19	0.28	0.39	0.44
MAP(L=10)	0.23	0.29	0.36	0.43	0.52	0.58	0.64	0.68	0.61	0.45	0.06

Note: The underlined correlation coefficients are at $p < 0.0001$. The italicized correlation coefficients are at $p < 0.05$. The row of "MAP (L=10)" has $n = 19$. All other rows have $n = 107$.

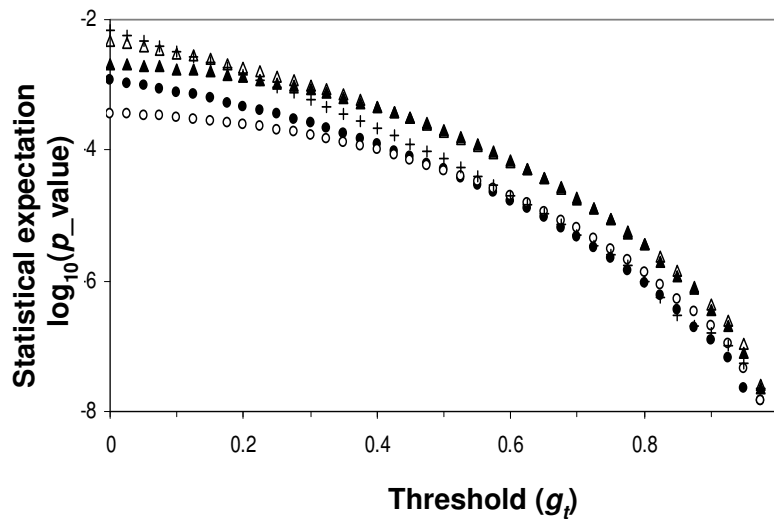


Figure 2. Relationship between probability and threshold G value of the 5 PWMs (MA0010, MA0017, MA0046, MA0069, MA0082). $w = 14$.

3.5. Integration

It is obvious that a selection of threshold should be based on a selection of G value in combination of either of PWM length (w), information content, or the best log-odd score. Since length factor implicitly incorporated in both information content and the log-odd score, we select the better one between information content and the best log-odd score. Figure 3

indicates that a combination of G values with the best log-odd scores performs better in representing the statistical expectation than the combination of G values with information contents. Thus, a combination of G values with the best log-odd scores is used as a guide to selecting a G threshold. The combination simply revert G values back to log-odd score.

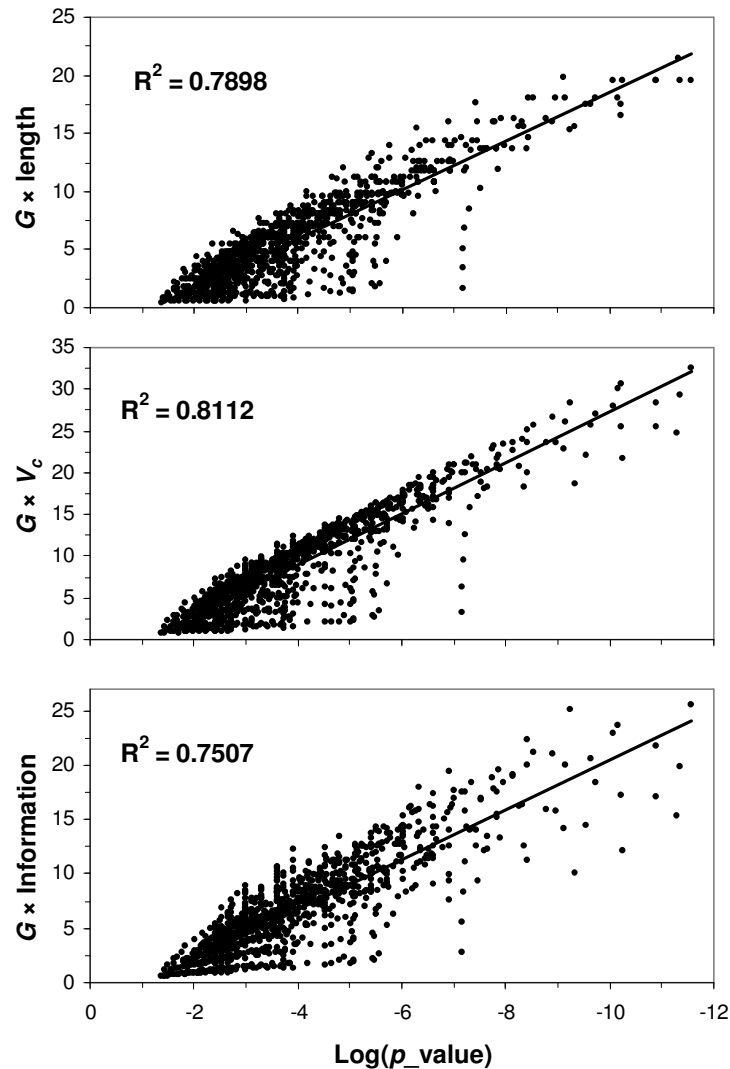


Figure 3. Correlation between statistical expectation and the combination of G value with various properties of the PWM.

4. APPLICATIONS

We applied the above method to generate the statistical expectation function (over G) for each of the 585 vertebrate PWMs and 56 yeast PWMs from TRANSFAC database (Matys et al., 2006). The background probabilities are generated from enumeration of the nucleotides in

all known genes' promoters of each species. We also searched for the putative binding sites of yeast transcription factors GCN4 and ROX1 using the TRANSFAC yeast PWMs, GCN4_C and ROX1_Q6, respectively. Finally, we searched for putative binding sites of the human tumor suppressor HIC-1 using the two vertebrate PWMs, HIC1_02 and HIC1_03, from TRANSFAC. In order to keep simplicity in calculation, no pseudo-count is applied in these applications. Therefore, all subsequences that contain a nucleotide with a 0.00 probability in the PWM are not considered in this study.

4.1. Vertebrate and Yeast PWMs

We used both species specific background and default background for comparison. The results indicate there is no perceivable difference between the two with regard to correlation between a combination of G value with PWM length, best log-odd score, or information content on one hand and the statistical expectation on the other (Table 2). Both vertebrate and yeast data confirms that a combination of G value with best log-odd score is the best in representing the statistical expectation.

Table 2. Correlation between the statistical expectations with various properties of PWMs from three different databases

PWM data Source	JASPAR	TRANSFAC Vertebrate		TRANSFAC Fungi	
	default	human	default	yeast	default
G x best log-odd	0.9007	0.8726	0.8752	0.8617	0.8637
G x Length	0.8887	0.8352	0.8359	0.8611	0.8637
G x Information content	0.8664	0.8403	0.8416	0.8120	0.8096

* Background: default = {0.25, 0.25, 0.25, 0.25};
 human = {0.2259, 0.2716, 0.2767, 0.2258};
 yeast = {0.3096, 0.1932, 0.1893, 0.3079}.

4.2. Yeast Genes

Among the available data, the yeast genome (*Saccharomyces cerevisiae*) is best studied. From the TRANSFAC database (Version 10.4), we retrieve two PWMs (GCN4_C and ROX1_Q6), which model the binding sites for transcription factors GCN4 and ROX1, respectively. The promoter sequences of all 5769 genes from SGDgene table were retrieved from the UCSC Genome Browser (<http://genome.ucsc.edu/>). The promoter sequences contain 600 bp upstream of the transcription start site (TSS). In order to validate the result, we retrieved the known (documented) associations between the transcription factors and their respective target genes from YEASTRACT database (Teixeira et al., 2006).

We searched the promoter sequences by using the two PWMs and setting different thresholds of G values ranging from 0.40 to 1.00. At each threshold, we considered a putative association between a TF and a target gene if a putative binding site of the TF is found in the

promoter sequence of this gene. The result of such putative association is validated by known associations obtained from the YEASTRACT database.

For the purpose of comparison, we apply the terminology of *Sensitivity* (S_n) and *Positive Predictive Value* (PPV) as defined in (Altman and Bland, 1994a, 1994b). *Sensitivity* is the proportion of all known associations ($TP+FN$) that are accurately predicted (TP); and the *Positive Predictive Value* is the proportion of predicted associations ($TP+FP$) that are true (TP). Traditionally, *Specificity* (proportion of negatives that are predicted false) is used in the evaluation of a method. In genomic sequences, true negative (TN) is predominately higher than any of FP , TP or FN . *Specificity* as defined in Altman and Bland (1994a) would not be able to reveal signals effectively as its value would be very close to 1.00 in almost all instances (Bursset and Guigo, 1996). Therefore, we adopted the *Positive Predictive Value* instead of *Specificity* in this study so that the signals can be comparable.

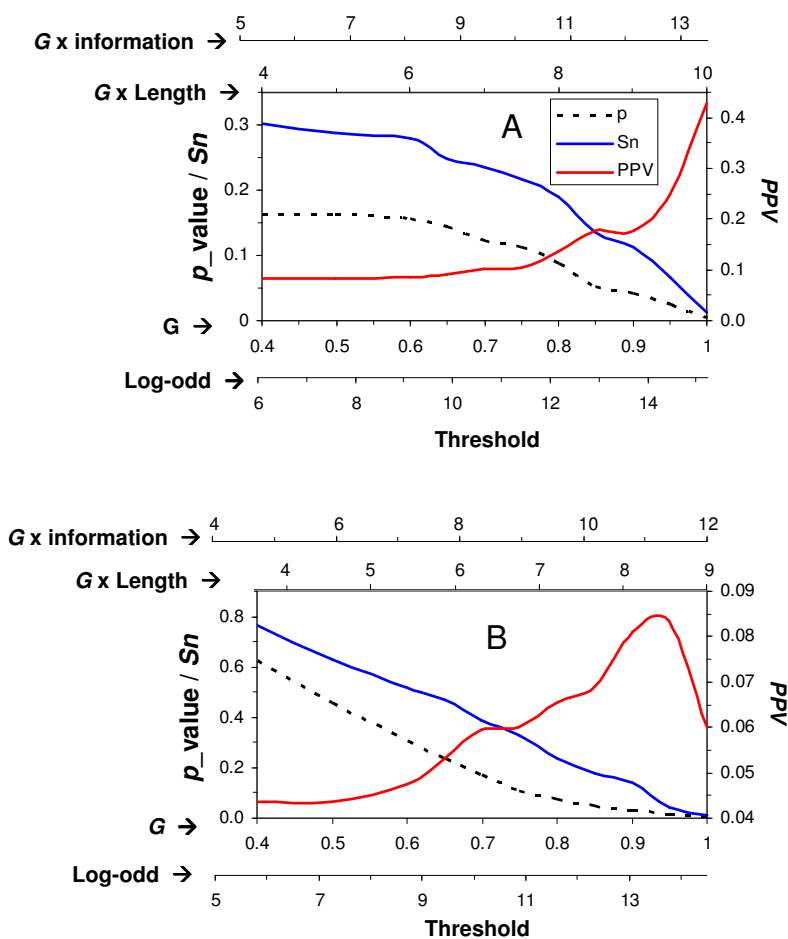


Figure 4. Performance evaluation on yeast data application case. The plot legends in panel A (GCN4_C) also apply to panel B (ROX1_Q6). S_n : sensitivity, PPV : positive predictive value, p : probability.

By decreasing the threshold, more known associations between TFs and their targets are found by the corresponding PWM as reflected by the *Sensitivity* values (Fig. 4). However, the number of false positives increases as reflected by the *Positive Predictive Values*. Based on the *p_value*, users will be able to find a corresponding threshold. For example, the log-odd threshold ($p \leq 0.05$) should be 13 for GCN4_C and 12.25 for ROX1_Q6.

4.3. Human Cancer Genes

We retrieved 406 cancer gene entries from the CancerGenes Resequencing Resource (Higgins et al., 2006). These 406 entries represent 385 distinct genes. We retrieved promoter sequences of these cancer genes from the UCSC Genome Browser (<http://genome.ucsc.edu/>). The promoter sequences cover the range of 1000 bp upstream and 200 bp downstream of TSS with a total length of 1200 bp. We included the 200bp downstream sequences in this study to account for potential alternative TSS annotations that may appear in some genes. We retrieved two PWMs (HIC1-02, HIC1_03) for a tumor suppressor gene HIC-1 from the TRANSFAC database and searched the promoter sequences for putative TFBSs that fit the two PWMs over the *G* threshold between 0.40 and 0.90. There is no similar database to YEASTRACT for known associations between the transcription suppressor and its target genes in the human genome.

We searched promoters of human cancer genes and predicted about 20% of these cancer gene have potential association with the transcription suppressor HIC-1 (Fig. 5, $p \leq 0.05$). Since our promoter sequences are from a normal human individual (not a cancer patient), it is not unreasonable that 20% of the cancer genes are associated with the transcription suppressor.

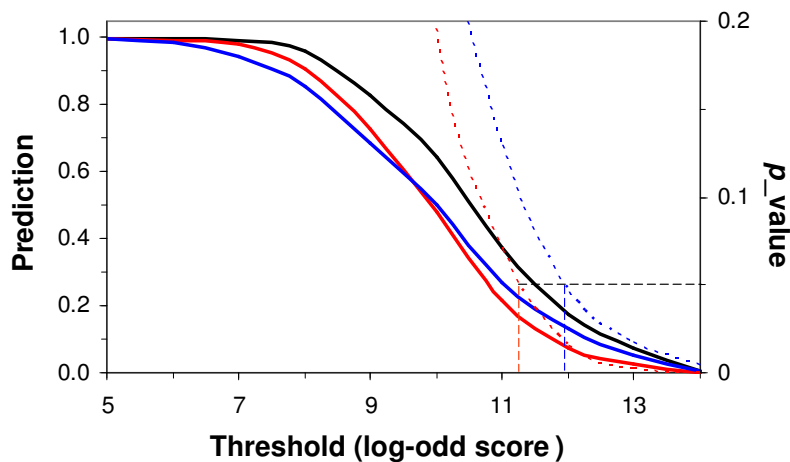


Figure 5. Prediction of association of transcription factor HIC1 with target human cancer genes. Red: predicted by HIC1_02, Blue: predicted by HIC1_03, Black: union prediction of HIC1_02 and HIC1_03. Solid curves: predicted percentage, Dotted curves: variation of *p*-values. Broken lines: marks of significant levels ($p < 0.05$).

4.4. Comparison with Match

We identified 16 yeast genes (Table 3) that are known to be associated with both transcription factors GCN4 and ROX1 (YEASTRACT database, Teixeira, et al., 2006) and used them to validate our method and compared it with Match (Kel et al., 2003). While using the default similarity threshold, Match could only find one of the 32 known associations. All associations are found by Match when decrease the threshold because it considers pseudo-count, which would guarantee to find all instance at a low threshold and to have very high false positive predictions. Without knowing their statistical expectation, we did not include those misleading predictions in Table 3. Our method found 16 of the 32 associations, of which 3 are significant ($p < 0.05$).

Table 3. Validation and comparison with Match (Kel et al., 2003). The values in the current method columns indicate that the associations are found at threshold of G / p_value.

Gene	Match		Current method	
	ROX1_Q6	GCN4_C	ROX1_Q6	GCN_C
ADH1			0.71 / 0.16	
ADH5			0.69 / 0.18	
BOP2		+	0.71 / 0.16	1.00 / 0.005
CWP1			0.81 / 0.07	
CWP2				
GAT2			0.64 / 0.24	
GID8			0.72 / 0.15	
HSP12			0.47 / 0.50	
HSP26			0.88 / 0.04	
HXT5				0.63 / 0.15
IDH1			0.77 / 0.09	
LYS1			0.55 / 0.39	
LYS9			0.47 / 0.50	
MUC1			0.47 / 0.50	
RAD16			0.53 / 0.41	
RPI1			0.91 / 0.03	

5. DISCUSSIONS

Positional weight matrix has been extensively used in discovery of sequence motifs, such as transcription factor binding sites. The main challenge in using PWM is to find a threshold

for various objective functions. There is no substantial breakthrough so far in solving this problem. Match (Kel et al, 2003) appears more appealing than the others and widely used. Match takes the maximum and minimum score and scales them between 0.00 and 1.00 and it also takes into consideration five consecutive nucleotides whose maximum score is the best in any region of the PWM space. P-Match (Chekmenev et al., 2005) combines pattern matching and weight matrix approaches and claimed to be more accurate. We tried to search for the GCN4_C and ROX1_Q1 motifs using P-Match, but unable to find any by the threshold setting defined by those authors. It is expected to find all associations at a lower threshold same as those predicted by Match. However, those potential associations would be buried in a large amount of false positive predictions. Without prior knowledge of these associations, it would be nearly impossible to distinguish them from false positives. For the methods such as Match and P-Match, without an indication of statistical expectation of the predicted motifs, users would be skeptical on the result.

Nucleotide frequency varies across different genomes, for example, the frequencies of A, C, G, T in human promoter sequences are 0.23, 0.27, 0.27 and 0.23, respectively, which are not too much different from the default frequency (0.25 for each nucleotide). But in the yeast promoter sequences, they are 0.31, 0.19, 0.19 and 0.31, respectively. For this reason, we use genome specific nucleotide frequency. Additionally, the nucleotide frequencies vary over various regions of genomics sequences (Pan et al., 2004, Smith et al., 2007). For more precise prediction, regional nucleotide frequencies should be applied.

Occasionally, the log-odd score of a motif could be dominated by one or a few positions because of their extremely high or low frequency values for certain nucleotide(s). Probably, one could argue that a log-odd value of 0.00 for a subsequence might not represent its identity with the background model because the influence of one or more high value(s) of the high frequency nucleotide(s) at certain position(s) is neutralized by the influence of one extremely low value derived from a low frequency nucleotide at another position. Nevertheless, because the overall log-odd score is close to 0.00, no matter whether it is caused by the neutralization of frequencies across various positions or by overall values close to 0.00, this subsequence is not likely a true TFBS. Therefore, a 0.00 log-odd score indicates the subsequence is most unlikely a TFBS.

It is important to consider the statistical expectation of a predicted motif instance. With the density function that we generated for each PWM, it is convenient to find the statistical expectation of each predicted motif instance base on its G value. This G value can easily be reverted back to log-odd score or relative information as can be seen in Fig. 4. Since log-odd value is closely related with statistical expectation as indicated in this study, people would question the necessity of going through the step of G value. The necessity of G value step is demonstrated through the generation of p -value distribution and through its value in measuring the distance between the generated log-odd value and the PWM as compared to the background.

A transcription factor usually binds on a DNA sequence through several positions. Numerous previous studies indicated that the positions inside a motif are somewhat interdependent. For this reason, PWMs are converted to high order hidden Markov models (Krogh et al, 1994). The log-odd score of a motif instance can be calculated based on the state transition probability (e.g. Viterbi score) of the high order hidden Markov model. The same calculation proposed in Equation (13) can be applied to scale the log-odd scores derived from hidden Markov models and statistical expectations can be calculated accordingly.

Many biological problems can not be easily revealed by simply measuring statistical expectation. For example, in our application of yeast genome, too stringent threshold would exclude many potential candidates, such as the threshold set in Match for the two yeast PWMs (Table 3) and our work in ROX1_Q6 (Fig. 4B). With varying threshold incorporating both the G values and the distribution function of statistical expectation, we are able to find more motifs using the method proposed in this paper.

Generally, a less stringent threshold would incur higher false positive prediction (Fig. 4). This can be complemented by incorporating other information such as microarray gene expression data (Hertzberg et al., 2007) or through comparative genomic approaches (e.g. Newberg et al., 2007). Incorporating gene expression data is certainly a boost in motif finding. However, such data are not always available. With comparative genomics approach, it is arguable even though some successes were shown. Our recent study indicates that promoters of most human genes are significantly different from their orthologues in mice or rats. Similarly, the divergence in the binding sites of the pseudohyphal regulators Ste12 and Tec1 among three closely related yeasts, *Saccharomyces cerevisiae*, *S. mikatae*, and *S. bayanus*, is significant (Borneman et al., 2007). In that case, substantial number of functional motifs in one organism may not appear in the promoter of orthologous genes in another organism, even if both are closely related. Such motifs could be excluded during the prediction process.

Based on the application cases, we suggest taking consideration of both objective function scores and statistical expectation in selecting a threshold. In choosing a threshold, log-odd score appears to be a good objective function as compared with other functions. It is highly correlated with statistical expectation.

In the application of cancer genes, we predicted 20% of the 385 cancer genes are subjected to transcription suppression of HIC-1. Since the promoter sequences are from normal human genome, one would expect all cancer genes are suppressed one way or another. We need to realize that many other cancer gene suppressors are not included in this study. For example, the popular cancer suppressor gene P53 is one among many others.

6. CONCLUSIONS

In prediction of a sequence motif using positional weight matrix, it is important to find a statistically meaningful threshold. In this chapter, we described an approach to find a threshold that incorporates the goodness-of-fit of a subsequence to a PWM, statistical expectation, and information content. We demonstrated the strength of this approach through various application cases and compared the method favorably with the broadly used Match method using 16 yeast genes of known association with two transcription factors. We also predicted that 20% of the known cancer genes have association with the tumor suppressor gene HIC-1.

REFERENCES

- Altman DG, Bland JM, 1994a. Statistics Notes: Diagnostic Tests 1 - sensitivity and specificity. *British Medical Journal* **308**: 1552.

- Altman DG, Bland JM, 1994b. Statistics Notes: Diagnostic Tests 2 - Predictive Values. *British Medical Journal* **309**: 102.
- Bailey EL, Elkan C, 1994. Fitting a mixture model by expectation maximization to discovery motifs in biopolymers. In: *Proceeding of the Second International Conference on intelligent System for Molecular Biology*. AAAI Press, pp. 28-38.
- Becker MM, Wang JC, 1984. Use of light for footprinting DNA *in vivo*. *Nature* **309**: 682-687.
- Blanchette M, Tompa M, 2002. Discovery of regulatory elements by comparative method for phylogenetic footprinting. *Genome Res* **12**: 739-748.
- Borneman AR, Gianoulis TA, Zhang ZD, et al., 2007. Divergence of transcription factor binding sites across related yeast species. *Science* **317**: 815-819.
- Bulyk ML, Huang X, Choo Y, Church GM. Exploring the DNA-binding specificities of zinc fingers with DNA microarrays. *PNAS* **98**: 7158-7163.
- Burset M, Guigo R, 1996. Evaluation of gene structure prediction programs. *Genomics* **34**: 353-367.
- Cartharius K, Frech K, Grote K, et al, 2005. MatInspector and beyond: promoter analysis based on transcription factor binding sites. *Bioinformatics* **21**: 2933-2942.
- Chekmenev DS, Haid C, Kel AE, 2005. P-Match: transcription factor binding site search by combining patterns and weight matrices. *Nucleic Acids Res* **33**: W432-437.
- D'haeseleer P, 2006. How does DNA sequence motif discovery work? *Nature Biotechnology* **24**: 959-961.
- Favorov AV, Gelfand MS, Gerasimova AV, Ravcheev DA, Mironov AA, Makeev VJ, 2005. A Gibbs sampler for identification of symmetrically structured, spaced DNA motifs with improved estimation of the signal length. *Bioinformatics* **21**: 2240-2245.
- Friberg M, von Rohr P, Gonnet G, 2005. Scoring functions for transcription factor binding site prediction. *BMC Bioinformatics* **6**: 84.
- Hertzberg L, Izraeli S, Domany E, 2007. STOP: searching for transcription factor motifs using gene expression. *Bioinformatics* **23**: 1737-1743.
- Higgins ME, Claremont M, Major JE, Sander C, Lash AE, 2006. CancerGenes: a gene selection resource for cancer genome projects. *Nucleic Acids Res* **35**: D721-726.
- Horak CE, Snyder M, 2002. ChIP-chip: A genomic approach for identifying transcription factor binding sites. *Methods In Enzymology* **350**: 469-483.
- Hughes JD, Estep PW, Tavazoie S, Church GM, 2000. Computational identification of cis-regulatory elements associated with groups of functionally related genes in *Saccharomyces cerevisiae*. *J Mol Biol* **296**: 1205-1214.
- Kel AE, Gossling E, Reuter I, et al, 2003. MATCH: A tool for searching transcription factor binding sites in DNA sequences. *Nucleic Acids Res* **31**: 3576-3579.
- Kiedziarska A, Smietana K, Czepczynska H, Otlewski J, 2007. Structural similarities and functional diversity of eukaryotic discoidin-like domains. *Biochim Biophys Acta* **1774**:1069-1078.
- Krogh A, Brown M, Mian IS, Sjolander K, Haussler D, 1994. Hidden Markov models in computational biology: application to protein modeling. *J Mol Biol* **235**: 1501-1531.
- Kullback S, Leibler RA, 1951. On information and sufficiency. *Ann Math Stat* **22**: 79-86.
- Lawrence CE, Reilly AA, 1990. An expectation maximization (EM) algorithm for the identification and characterization of common sites in unaligned biopolymer sequences. *Proteins Struct funct Genet* **7**: 41-51

- Lawrence CE, Altschul SF, Boguski MS, Liu JS, Neuwald AF, Wootton JC, 1993. Detecting subtle sequence signals: a Gibbs sampling strategy for multiple alignments. *Science* **262**: 208-214.
- Li L, Liang Y, Bass RL, 2007. GAPWM: a genetic algorithm method for optimizing a position weight matrix. *Bioinformatics* **23**: 1188-1194.
- Li N and Tompa M, 2006. Analysis of computational approaches for motif discovery. *Algorithms for Molecular Biology* **1**:8.
- Liu JS, Lawrence CE, Neuwald A, 1995. Bayesian models for multiple local sequence alignment and its Gibbs sampling strategies. *J Amer Statist Assoc* **90**: 1156-1170.
- Liu X, Brutlag DL, Liu JS, 2001. BioProspector: discovering conserved DNA motifs in upstream regulatory regions of co-expressed genes. *Pac Symp Biocomput* 2001: 127-138.
- Liu X, Brutlag DL, Liu JS, 2002. An algorithm for finding protein-DNA binding sites with applications to chromatin-immunoprecipitation microarray experiments. *Nature Biotech* **20**: 835-839.
- Liu Y, Liu XS, Wei L, Altman RB, Batzoglu S, 2004. Eukaryotic regulatory element conservation analysis and identification using comparative genomics. *Genome Res* **14**: 451-458.
- Matys V, Kel-Margoulis OV, Fricke E, et al, 2006. TRANSFAC and its module TRANSCompel: transcriptional gene regulation in eukaryotes. *Nucleic Acids Res* **34**: D108-110.
- Moses AM, Chiang DY, Eisen MB, 2004. Phylogenetic motif detection by expectation-maximization on evolutionary mixtures. *Pac Symp Biocomput* 2004: 324-35.
- Mukherjee S, Berger MF, Jona G, et al, 2004. Rapid analysis of the DNA-binding specificities of transcription factors with DNA microarrays. *Nat Genet* **36**: 1331-1339.
- Newberg LA, Thompson WA, Conlan S, Smith TM, McCue LA, Lawrence CE 2007. A phylogenetic Gibbs sampler that yields centroid solutions for cis-regulatory site prediction. *Bioinformatics* **23**:1718-1727
- Orlando V, 2000. Mapping chromosomal proteins *in vivo* by formaldehyde-crosslinked-chromatin immunoprecipitation. *Trends Biochem Sci* **25**: 99-104.
- Pan Y, 2006. Advances in the Discovery of cis-Regulatory Elements. *Curr Bioinfo* **1**:321-336
- Pan Y, Phan S, 2008. Guide to Threshold Selection for Motif Prediction Using Positional Weight Matrix. *Proceedings of the IAENG International Conference in Bioinformatics* (ISBN: 978-988-98671-8-8), March 19-21, 2008, Hong Kong; pp 151-156.
- Pan Y, Smith B, Fang H, Famili FA, Sikorska M, Walker PR, 2004. Selection of putative cis-regulatory motifs through regional and global conservation. In: *Proceedings of the 2004 IEEE Computational System Bioinformatics Conference (CSB2004)*, 16-19 August 2004, Stanford, CA, USA 2004; pp 684-685.
- Pavesi G, Mereghetti P, Mauri G, Pesole G, 2004. Weeder web: discovery of transcription factor binding sites in a set of sequences from co-regulated genes. *Nucleic Acids Res* **32**: W199-203.
- Prakash A, Blanchette M, Sinha S, Tompa M, 2004. Motif discovery in heterogeneous sequence data. *Pac Symp Biocomput* 2004; 248-259.
- Sandelin A, Alkema W, Engstrom P, Wasserman WW, Lenhard B, 2004. JASPAR: an open-access database for eukaryotic transcription factor binding profiles. *Nucleic Acids Res* **32**: D91-94.

- Schneider TD and Stephens RM, 1990. Sequence logos: a new way to display consensus sequences. *Nucleic Acids Res.* **18**: 6097-6100.
- Schneider TD, Stormo GD, and Gold L, 1986. Information content of binding sites on nucleotide sequences. *J Mol Biol* **198**: 415-431.
- Schreiter ER, Drennan CL, 2007. Ribbon-helix-helix transcription factors: variations on a theme. *Nat Rev Microbiol.* **5**:710-720.
- Shannon CE, 1948. A mathematical theory of communication. *Bell System Tech J* **27**: 379-423, 623-656.
- Siddharthan R, Siggia ED, van Nimwegen E, 2005. PhyloGibbs: a Gibbs sampling motif finder that incorporates phylogeny. *PLoS Comput Biol* 2005; 1: 534-56.
- Siepel A, Haussler D, 2005. Phylogenetic hidden Markov models. In: Nielsen R Ed, *Statistical Methods in Molecular Evolution*. Springer, New York 2005; pp 325–351.
- Sinha S. Discriminative motifs, 2003. *J Comput Biol* **10**: 599-615.
- Sinha S, Blanchette M, Tompa M, 2004. PhyME: a probabilistic algorithm for finding motifs in sets of orthologous sequences. *BMC Bioinformatics* **5**: 170.
- Sinha S, Tompa M, 2002. Discovery of novel transcription factor binding sites by statistical overrepresentation. *Nucleic Acids Res* **30**: 5549-5560.
- Smith B, Fang H, Pan Y, Walker PR, Famili AF, Sikorska M, 2007. Evolution of motif variants and positional bias of the cyclic-AMP response element. *BMC Evol Biol.* **7**: S15.
- Solomon MJ, Varshavsky A, 1985. Formaldehyde-mediated DNA-protein crosslinking: a probe for *in vivo* chromatin structures. *PNAS* **82**: 6470-6474.
- Stormo GD, 2000. DNA binding sites: representation and discovery. *Bioinformatics* **16**: 16-23.
- Teixeira MC, Monteiro P, Jain P, *et al.* 2006. The YEASTRACT database: a tool for the analysis of transcription regulatory associations in *Saccharomyces cerevisiae*. *Nucleic Acids Res.* **34**: D446-451.
- Thijs G, Lescot M, Marchal K, *et al*, 2001. A higher order background model improves the detection of regulatory elements by Gibbs Sampling. *Bioinformatics* **17**: 1113-1122.
- Thompson W, Rouchka EC, Lawrence CE, 2003. Gibbs recursive sampler: finding transcription factor binding sites. *Nucleic Acids Res* **31**: 3580-3585.
- Thompson WA, Newberg LA, Conlan S, McCue LA, and Lawrence CE 2007. The Gibbs Centroid Sampler. *Nucleic Acids Res* **35**: W232-237.
- Tompa M, Li N, Bailey TL, *et al*, 2005. Assessing computational tools for the discovery of transcription factor binding sites. *Nat Biotech* **23**: 137-144.
- Workman CT and Stormo GD, 2000. ANN-Spec: a method for discovering transcription factor binding sites with improved specificity. *Pac Symp Biocomput* 2000: 467-478.
- Zhang MQ, 2002. Computational methods for promoter recognition. In: Jiang T, Xu Y, Zhang MQ Eds, *Current Topics in Computational Molecular Biology*. MIT Press, Cambridge, Massachusetts. pp 249-268.
- Zhu X, Gerstein M, Snyder M, 2007. Getting connected: analysis and principles of biological networks. *Genes & Dev* **21**: 1010-1024.