Interpolated Markov chains for eukaryotic promoter recognition

Uwe Ohler^{1,2}, Stefan Harbeck¹, Heinrich Niemann¹, Elmar Nöth¹ and Martin G. Reese²

¹Chair for Pattern Recognition (Computer Science V), University of Erlangen-Nuremberg, Martensstraße 3, D-91058 Erlangen, Germany and ²Department of Molecular and Cell Biology, University of California at Berkeley, 539 Life Sciences Addition, Berkeley, CA 94720-3200, USA

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Abstract

Motivation: We describe a new content-based approach for the detection of promoter regions of eukaryotic protein encoding genes. Our system is based on three interpolated Markov chains (IMCs) of different order which are trained on coding, non-coding and promoter sequences. It was recently shown that the interpolation of Markov chains leads to stable parameters and improves on the results in microbial gene finding (Salzberg et al., Nucleic Acids Res., 26, 544-548, 1998). Here, we present new methods for an automated estimation of optimal interpolation parameters and show how the IMCs can be applied to detect promoters in contiguous DNA sequences. Our interpolation approach can also be employed to obtain a reliable scoring function for human coding DNA regions, and the trained models can easily be incorporated in the general framework for gene recognition systems.

Results: A 5-fold cross-validation evaluation of our IMC approach on a representative sequence set yielded a mean correlation coefficient of 0.84 (promoter versus coding sequences) and 0.53 (promoter versus non-coding sequences). Applied to the task of eukaryotic promoter region identification in genomic DNA sequences, our classifier identifies 50% of the promoter regions in the sequences used in the most recent review and comparison by Fickett and Hatzigeorgiou (Genome Res., 7, 861–878, 1997), while having a false-positive rate of 1/849 bp.

Contact: ohler@informatik.uni-erlangen.de

Introduction

Today's state-of-the-art eukaryotic gene-finding algorithms (such as Kulp *et al.*, 1996; Krogh, 1997; Burge and Karlin, 1998) are based on a statistical framework which is, in many cases, a generalization of a hidden Markov model (HMM), also called hidden semi-Markov model. Within this framework, several scoring functions for signals such as splice sites and for regions such as exons, introns or promoters are

combined. After the search for possible signals and the judgement of the segments in between, the standard HMM decoding algorithm then provides the best path through the graph of all possible segmentations of the whole sequence. Although much progress has been made with this approach, there is still a considerable need for robust algorithms to classify the individual signals and segments, as the accuracy of the system output depends on the accuracy of its components. In the following, we will present new models for the classification of individual DNA segments, and will mainly focus on the recognition of eukaryotic promoter regions.

Popular content-based measures for primary DNA sequences make use of Markov chains (MCs) of a fixed order (closely related to oligomer measures) and have been employed, for example, in the widespread GeneMark and GeneMark.hmm prokaryotic gene finders (Lukashin and Borodovsky, 1998). Recently, the linear interpolation of MCs of different order has been described for microbial gene recognition (Salzberg *et al.*, 1998). An interpolation provides a better parameter estimation, as, with increasing order of the Markov chain, the training algorithms lack a suitable amount of data because the number of model parameters increases exponentially.

Here we present a new interpolation scheme which has been successfully applied by our group for various speech recognition tasks (see Schukat-Talamazzini *et al.*, 1997). In the context of speech recognition, interpolated Markov chains (IMCs) to judge the likelihood of symbol sequences are commonly referred to as stochastic language models. In contrast to the method described by Salzberg *et al.* (1998), who provided a function including a χ^2 test on statistical significance to calculate parameters for a linear interpolation, we use a disjoint part of the training sample to estimate automatically optimal interpolation parameters with respect to a statistical objective function.

We will show how this kind of IMC can improve the detection of eukaryotic promoter sequences in unknown genomic DNA. Recent progress in the understanding of the structure and function of these polymerase II promoters is reviewed in detail by Kornberg (1996; and other articles in the same issue) or Nikolov and Burley (1997).

The survey of Fickett and Hatzigeorgiou (1997) provides an excellent introduction to the topic of automated recognition of eukaryotic promoters and a comparison of the available systems for general-purpose Pol II promoter prediction. Among these are linear discriminative (Solovyev and Salamov, 1997) as well as neural network (Reese and Eeckman, 1998) or content-based (Hutchinson, 1996; Audic and Claverie, 1997) methods. Content-based measures were up to now either plagued by too large a number of false positives, or imposed restrictions on the number of predictions. The results obtained by our IMCs will demonstrate the improvement of the recognition rate compared to the best methods available. Our goal was to build a general-purpose promoter recognition system that can be applied to the general task of promoter recognition; computer models constructed for specific tissue types as in Frech et al. (1998) have a much lower false-positive recognition rate. On the other hand, there is an apparent need to add a general promoter recognition module to a gene recognition system. This should help to split contiguous stretches of DNA into the right number of genes and detect the correct transcription start site, which might be far upstream from the translated region.

1. Algorithm

Let us assume that we have *K* classes $\Omega_1 \dots \Omega_k$ and wish to classify a sequence $w = w_1 \dots w_T$ with symbols w_i , taken from a finite vocabulary *V*, into one class. In the case of molecular genetics, the alphabet might consist of amino acids or nucleotides. We can make use of the chain rule to compute the likelihood of a particular sequence for each class:

$$P_k := P(\boldsymbol{w}|\boldsymbol{\Omega}_k) = \prod_{i=1}^{T} P(w_i|w_1...w_{i-1}, \boldsymbol{\Omega}_k)$$
(1)
context

This equation shows that one symbol in a sequence is dependent on all its predecessors, i.e. on the context of preceding symbols. Using Bayes' rule, we are able to classify the sequence into sequence class \hat{k} according to the largest a posteriori probability:

$$\hat{k} = \operatorname{argmax}_{k} P(\Omega_{k} | \boldsymbol{w}) = \operatorname{argmax}_{k} (p_{k} \cdot P_{k})$$

$$k \qquad (2)$$

If we have no exact knowledge about the a priori probabilities p_k of our sequence classes, the values p_k are assumed to be uniformly distributed and can be neglected. We therefore need to assign a likelihood to the symbol sequence w. If we can establish a model which computes this probability, we have the means to determine how likely a sequence will occur in a specific class.

1.1 Maximum Likelihood parameter estimation

In the following, we will drop the condition on class Ω_k for simplicity. The right-hand side of equation (1) contains a context of arbitrary length which cannot be handled; therefore, an approximation is made by imposing a restriction. A possible approximation of the probability P(w) is thus made by limiting the context length to N - 1:

$$P(\boldsymbol{w}) \approx \prod_{i=1}^{T} P(w_i | w_{i-N+1} \dots w_{i-1})$$
(3)

The resulting model is called a Markov chain of order N-1. Our goal is to obtain parameters—in our case values for the conditional probabilities $P(w_i|w_{i-N+1} \dots w_{i-1})$ —which lead to the best possible recognition rate on the *K* classes under consideration. As we cannot optimize the recognition rate directly, we have to use objective functions which show the desired behaviour and for which a solution can be found. One well-known objective function is Maximum Likelihood (ML). If Λ_k denotes the set of the parameters of model M_k for class Ω_k , we optimize the following function $R(\Lambda_k)$:

$$R(\Lambda_k) = \prod_{i=1}^{n_k} P(\boldsymbol{w}_{ki} | \boldsymbol{M}_k)$$
(4)

where n_k is the number of training sequences for class k. Each class is regarded as independent of the others, and ML estimation tries to maximize the probability that the given training sample was generated, knowing to which class each sequence belongs.

Using a training sample, the ML estimation of the conditional probabilities $\tilde{P}(w_i|w_{i-N+1}...w_{i-1})$ can be performed simply by counting the oligomers of length *N* and *N* – 1 in a set of training sequences:

$$\tilde{P}(w_i|w_{i-N+1}^{i-1}) = \frac{\#(w_{i-N+1}^i)}{\#(w_{i-N+1}^{i-1})}$$
(5)

where w_x^y is an abbreviation for the partial sequence from position *x* to position *y*, and # denotes the frequency of its argument in the training sample. Here, we have to meet two problems.

- 1. The approximation by a large context gets closer to the real probability as denoted in equation (1). Unfortunately, the number of parameters which have to be estimated increases exponentially with the number of N, and thus the ML estimates become far from being reliable because of the limited training sample size.
- 2. With increasing length, some *N*-mers might not occur at all in the training sample. This has the consequence that the likelihood of the whole sequence *w* is set to zero if it contains any unseen *N*-mer. This might be justified if it really is not a part of the considered class. On the other hand, the sample size might simply be too small to contain every single *N*-mer. As we do not know

which case is true, we must not set any likelihood to zero.

A solution to these problems—the trade-off between the model context and the training sample size, and the problem of unseen *N*-mers—can be found by introducing a weighted interpolation scheme.

1.2 Interpolation techniques

The basic idea of applying interpolation methods is to fall back on the probability estimation of subsequences shorter than *N* if the frequencies of an *N*-mer $v = v_1 \dots v_N$ cannot be reliably estimated. In principle, interpolation leads us to a re-estimation of the initial parameter values [equation (5)]. Here, we will consider two different interpolation techniques. The first one is the linear interpolation between all conditional probabilities with increasing context length up to N - 1:

$$\hat{P}(v_{N}|v_{1}^{N-1}) := \rho_{0}\frac{1}{L}$$

$$+ \rho_{1}\tilde{P}(v_{N})$$

$$+ \rho_{2}\tilde{P}(v_{N}|v_{N-1})$$

$$\vdots$$

$$+ \rho_{N}\tilde{P}(v_{N}|v_{1}^{N-1})$$
(6)

The fraction (1/L) accounts for unseen events and ensures that no probability is set to zero. The coefficients are nonnegative values which sum up to one to guarantee that the new parameter values $\hat{P}(\cdot | \cdot)$ again form a probability distribution.

Setting all the weights $\rho_0 \dots \rho_{N-1}$ to zero and ρ_N to one is very similar to the well-known oligomer approach, with the only difference that in a Markov chain the parameters are normalized with respect to the context [see equation (5)]. The models with linear interpolation are thus a straightforward generalization combining oligomers of different length. The advantage of interpolation is that the model can take into account statistics of a higher order without running into the danger of overfitting the model to the training data.

Equation (6) contains only one vector of interpolation coefficients, whether all the subsequences up to length *N* really occurred in the training data or not. Additionally, all parameters are treated equally, whereas the interpolation coefficient assigned to a parameter with a frequently occurring context should be larger than the coefficient for a rare event. By introducing an additional function $g_i(\mathbf{v}')$ which scores the reliability of the context $\mathbf{v}' = v_1^{N-1}$ monotonically, the linear interpolation can be extended to handle this problem accurately:

$$\hat{P}(v_N|\boldsymbol{v}') := \frac{\sum_{i=0}^N \rho_i \cdot g_i(\boldsymbol{v}') \cdot \tilde{P}_i(v_N|\boldsymbol{v}')}{\sum_{i=0}^N \rho_i \cdot g_i(\boldsymbol{v}')}$$
(7)

where $\tilde{P}(v_N|v')$ serves as an abbreviation for the estimates of different context lengths *i*, as was shown in detail in equation (6). This interpolation scheme is called rational interpolation. It overcomes the problems of linear interpolation by using the function $g_i(v')$, which we chose to be a sigmoid function dependent on the frequency of the last *i* symbols of v':

$$g_i(v') = \frac{\#_i(v')}{\#_i(v') + C}$$
(8)

The shape of the sigmoid function is dependent on the constant bias C. In the case of C = 0, the function g_i is always equal to one and equation (7) becomes equivalent to linear interpolation. Also, with an increasing amount of training data, the bias C becomes less and less important; the rational interpolation thus has the largest impact if the training sample size is small.

1.3 Maximum Likelihood estimation of interpolation coefficients

We still lack the means to specify appropriate coefficients ρ_i for both linear and rational interpolation. In our approach, optimal coefficients according to the ML objective function are calculated using a second disjoint part of the training sample. This step is called validation and is carried out after the initial estimation of the conditional probabilities (Section 1.1). There is no closed solution for a maximum of the ML objective function in the case of IMCs, but for the coefficients used in linear interpolation, a local optimum can be found with the iterative Expectation Maximization (EM) algorithm (Dempster et al., 1977): we regard the coefficients as hidden variables in a double stochastic process. Afterwards, a large weight will be assigned to those contexts for which we can obtain reliable estimations; if only sparse data are at hand, the weights belonging to short contexts will be increased.

For rational interpolation, the EM algorithm cannot be applied and the computation of locally optimal interpolation weights is carried out with a gradient descent algorithm instead. The detailed re-estimation formulas are omitted at this point and can be found in Schukat-Talamazzini *et al.* (1997). This automated estimation of optimal parameters is the main difference of our interpolation methods to those described for parsing microbial sequences (Salzberg *et al.*, 1998), where the coefficients are calculated using a predefined function based on the χ^2 statistical test.

1.4 Sequence classification using interpolated Markov chains

After an IMC has been trained according to Sections 1.1–1.3 for each of the considered sequence classes, the IMCs can be used in parallel to classify a sequence using equation (2). Sometimes, though, the focus is put on the right classification



Fig. 1. Content-based classification with interpolated Markov chains (IMC). The output is the difference between scores of the best background model and the model of interest; this score is then classified with a suitable threshold.

of only one class. In this case, we have one class of interest and one or more 'background' classes, and it is not important which particular class a sequence from the background is assigned to, as long as it is not classified into the class of interest. This situation occurs in promoter recognition, where we want to distinguish promoters ('class of interest') from non-promoters (the 'background' consists of several models for exonic, intronic and intergenic sequences). We can tune the IMCs with respect to sensitivity and specificity for the class of interest using the following approach. First, we compute the likelihood P_k for each class Ω_k , and then we determine the difference between the score for the model of interest P_M and the best of the background models P_B . Including a length normalization, we obtain the following equation for the total score S:

$$S(w) = \frac{P_B(w) - P_M(w)}{len(w)}$$
(9)

In practice, the logarithms of the probabilities are used because of the more efficient computation and the prevention of numerically unstable values when regarding long sequences. In Figure 1, an overview of the resulting algorithm is given.

Choosing a suitable threshold value for the total score *S*, we can select any percentage of false positives (i.e. patterns

out of one of the background classes which were classified into the class of interest). The curve of false-positive rate versus recognition rate over the whole range is called receiver operating characteristics (ROC) and will be used to compare the performance of different classifiers. Additionally, we will provide the correlation coefficient (*CC*), which is defined as follows:

$$CC = \frac{(TP \cdot TN) - (FN \cdot FP)}{\sqrt{(TP + FN) \cdot (TN + FP) \cdot (TP + FP) \cdot (TN + FN)}} (10)$$

Herein, TP stands for true positives, TN for true negatives, FP for false positives and FN for false negatives; these numbers denote the absolute numbers of correctly and wrongly classified sequences.

1.5 Application of IMCs to search for regulatory regions

We will now briefly describe our system for the detection of eukaryotic polymerase II promoters in contiguous DNA sequences. The system consists of one IMC model for promoter sequences and two background IMC models for coding and non-coding sequences. To search for promoters in contiguous sequences, we use a sliding window of 300 bases (motivated by the size of the training sequences; see Section 2). Every 10 bases, the current sequence in the window is classified as promoter or non-promoter using a scoring threshold that has previously been selected empirically on the training data (see Figure 1). Because a whole promoter region is very likely to cause multiple predictions of several overlapping windows, a prediction is only made for each local minimum of the difference between background and promoter score which lies below the chosen threshold. The transcription start site is then assumed to be located at position 250 within the window. To eliminate single false predictions, a post-processing operation is applied on the graph of the score function S. By a smoothing algorithm, single false promoter predictions as well as single non-promoter predictions within a promoter region are filtered out. We chose to apply the hysteresis threshold algorithm, where a smoothing cursor of a chosen height is shifted over the curve from left to right. As the local minima within the smoothed graph usually comprise several positions with the same value, the prediction is then made at the position with the lowest value in the original graph. More detailed information can be found in Ohler and Reese (1998).

2. Data sets

We have built strongly needed representative training and test sets for eukaryotic promoter recognition which allow for a thorough comparison of different methods. These data sets are suited for algorithms aiming at human and Drosophila mela*nogaster* promoter prediction. The data do not contain only promoter sequences, which can be retrieved quite easily from the Eukaryotic Promoter Database EPD, but also carefully chosen coding and non-coding sequences. For the human promoter set, we extracted all non-related vertebrate sequences, except retroviruses, from EPD Release 50 (Perier et al., 1998). Retrieving only human promoter sequences would result in a too small data set to fit the parameters of our models; EPD Release 50 contained only 181 independent human sequences. Sequences with less than 40 bases upstream or 5 bases downstream from the annotated transcription start site were discarded to ensure that at least the possible TATA box and the initiator site were contained in each entry. This resulted in 565 entries, from which sequences of 300 bases (250 upstream and 50 downstream) were extracted.

For the coding and non-coding sequences, we used the exon and intron sequences of human genes contained in the data set of 1998 for the GENIE genefinding system (Kulp *et al.*, 1996; Reese *et al.*, 1997). The exons were concatenated to form long coding sequences. Then, 300-base-long non-overlapping sequences were extracted. Owing to the still limited amount of data, we divided the human data into five sets containing 113 promoter, 180 coding and 869 non-coding sequences each. On these sets, reliable results can now be

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obtained by carrying out a 5-fold cross-validation. In each experiment, the model is trained on four parts of the sequence data, leaving one part out at a time and testing the performance on the part not used for training. Then the average over all five experiments is computed and used as a result for comparison. All the data sets and more detailed information are publicly available, and can be retrieved via the URL http://www-hgc.lbl.gov/inf/human.html; this site also contains a link to the similar set of *D.melanogaster* data. We encourage researchers working in the field of promoter recognition to compare their algorithms on these representative sets.

To evaluate the performance of the system on long contiguous sequences, we made use of the data set in Fickett and Hatzigeorgiou (1997). Using these data, we evaluated our IMC-based system on a more realistic problem of recognizing transcription start sites and the corresponding promoters in DNA stretches of genomic DNA, and were able to compare our results with other programs. The set consists of 18 vertebrate sequences containing 24 annotated and experimentally proven promoters with a total of 33 120 bp. The evaluation on the contiguous sequences was carried out on both strands; recognition results are therefore given in base pairs instead of single bases.

3. Results and discussion

To get a first impression, we compared different context lengths (4–6 bases) and interpolation methods (none, linear and rational) on the classification of human promoters and coding sequences from the fixed length sequence set (see Section 2). Figure 2 shows a part of the ROC using IMCs of sixth order and pure simple hexamer frequencies, for which the best results could be obtained. The figure shows clearly that rational interpolation outperforms drastically the oligomer approach without interpolation; it is also superior to the simpler linear approach, thus confirming that interpolation helps us to avoid the effect of overfitting the models to the sparse training data.

As a second step, we applied careful 5-fold cross-validation experiments on the complete fixed-length sequence set (promoters, introns, coding sequences), using IMCs with a context length of six and rational interpolation. To get a better insight, we tested the promoter model not only against both non-promoter models at once, but also individually against one non-promoter class. Table 1 therefore contains the average of the five experiments for three discrimination tasks: promoter versus coding sequences, promoter versus intron sequences, and promoters versus both coding and non-coding sequences. Choosing a threshold for more than 5% of false positives here does not lead to a practically useful number of predictions.



Fig. 2. Comparison of the performance of IMC models with oligomer statistics without interpolation for eukaryotic promoter recognition. The results (which are the best for each considered method) were achieved for hexamers and IMCs based on 7-mers. The ROC curve for promoter/coding sequence classification in the range of 0–15% of false positives is shown. The models were trained on a set of 452 promoters and 720 coding sequences of 300 bases length, and evaluated on a disjoint test set of 113 promoters and 180 coding sequences.

Table 1. Promoter classification on vertebrate sequences with Markov chain models using rational interpolation and an order of six. For a certain percentage of false positives, the corresponding cross-validated recognition rate and the correlation coefficient are given. The recognition rate with the highest correlation coefficient is in bold (CDS is coding sequence)

False	Recognized promoters (%)		
positives (%)	Promoter versus CDS	Promoter versus intron	Promoter versus CDS/intron
0.0	58.6 (0.68)	12.9 (0.33)	3.9 (0.16)
1.0	69.4 (0.74)	32.2 (0.46)	29.9 (0.45)
2.0	78.8 (0.80)	42.5 (0.51)	41.8 (0.50)
3.0	80.5 (0.81)	49.7 (0.53)	48.7 (0.52)
4.0	85.7 (0.83)	51.9 (0.52)	53.6 (0.52)
5.0	88.9 (0.84)	54.7 (0.51)	56.6 (0.51)

The discrimination performance between promoters and coding regions is stunning: at a false-positive rate of 5%, almost 89% of the promoter sequences were classified correctly (correlation coefficient 0.84). Nevertheless, it is also very clear that a classification between promoter and introns is much more difficult: the best CC value obtained was 0.53, at a false-positive rate of 3% and a recognition rate of 49.7%. Most probably, this stems from the much weaker information contained in the introns compared to the strong coding information of the exons. On applying models on the three-part set of promoters, non-coding and coding sequences, the results are comparable to the two-class problem of promoters

and non-coding sequences, resulting from the much larger sample size of intronic sequences. Corresponding results were obtained for the *D.melanogaster* set (Ohler and Reese, 1998).

We applied one model trained on promoters, coding and non-coding sequences to the task of finding promoter regions in longer vertebrate DNA sequences, following the principles described in Section 1.5 and using the set of contiguous sequences from the promoter prediction program survey of Fickett and Hatzigeorgiou (1997). In this survey, a prediction is judged as correct if an annotated transcription start site lies within 200 bases downstream and 100 bases upstream from the predicted site. Using this criterion, and a threshold set at a rate of 4% false positives (highest CC value), we could detect 12 out of the 24 promoters (50%) while having one false prediction on average every 849 bp. The two programs which achieved the best performance in the survey could detect 54 and 42 % of the promoters with a false-positive rate of 1/460 and 1/789 bp, respectively (Reese and Eeckman, 1998; Solovyev and Salamov, 1997). These numbers show that the performance of the IMCs is slightly better than the best available tools for promoter prediction, but the number of test sequences is too small to make a general statement possible.

An example of the performance on the longest test sequence, the human phenol sulphotransferase gene (5663 bases, forward strand of GenBank accession code HSU54701), is shown in Figure 3. Following the approach described in Section 1.5, two predictions are made within this sequence, one of which is located close to one of the two annotated transcription start sites. A complete graph describing the regulatory potential over the sequence positions is calculated. Even if no clear decision is possible at the default threshold, a manual inspection of the graph may still reveal where a sudden change from regulatory (low values) to nonregulatory (high values) takes place.

A closer look at the contiguous sequences in the Fickett and Hatzigeorgion (1997) data set and the behaviour of the system concludes this section and helps to reveal some advantages and shortcomings of the current approach:

- The overall results are certainly influenced by the fact that our system was established as a promoter predictor for human sequences, whereas seven of the 18 sequences were of non-human origin.
- One start site missed was located only a few bases downstream of the sequence start. As we score a window which is assumed to contain 250 bases upstream and 50 bases downstream, no predictions are made before position 250.
- In sequence MMG67PRO, three annotated start sites were located within 300 bp, and our program made only one detection. This is not unexpected since a whole 300 bp region is scored at once, and the post-pro-



Fig. 3. Output of the system on a large contiguous sequence (GenBank accession HSU54701), before and after applying the automated smoothing step. The original graph depicts the difference of the best non-promoter and the promoter model score on the forward strand of the sequence. Two predictions are made, one for each local minimum below a predefined threshold on the smoothed output. Here, the predictions are located at positions 980 and 3580; the identified annotated transcription start site is located at position 935. Another start site located at position 2002 is not revealed.

cessing smoothes out the small local maxima that might help to separate the individual start sites.

- The prediction accuracy of the TSS location was quite good despite the fact that MC do not use location-specific information: seven of the 12 correct predictions were made within 30 bases from the annotated start site.
- Only two of the missed promoters were also not detected by any of the nine programs evaluated in Fickett and Hatzigeorgiou (1997). On the other hand, one promoter detected by our MC could not be identified by any other program. This means that much improvement could be achieved by a combination of several systems.

At the moment, we do not have a non-promoter model for intergenic sequences; if a reliable training sample for this sequence class can be obtained, the performance is likely to improve because of the more accurate sequence modelling. Obtaining such a sample though is difficult; most database entries contain only single genes, and for large sequences generated in the genome projects, the genes and especially the transcription start site annotations are mostly computational and not experimentally verified, and therefore not reliable.

The probably most widespread application of MCs so far is found in gene recognition systems, where they serve as a classifier for coding versus non-coding parts of a DNA sequence. Thus, we also compared the performance of our interpolated models to the standard MC, following the guidelines of the coding measure survey of Fickett and Tung (1992). On the GENIE data set of human exons and introns, the average recognition rate on 108-bp-long sequences is 85%, which is an improvement of 2.2 per cent points (frame-independent classification) compared to the best reviewed method, a non-interpolated Markov chain. Detailed results will be presented elsewhere.

4. Conclusions

In this paper, we describe the application of IMCs to contentbased DNA classification problems. The performance of our models on two different applications, the recognition of promoter regions and the discrimination of coding and non-coding sequences, is consistently better than that of oligomer models which realize MCs of a fixed order. We therefore recommend the use of interpolated models in any case, even if enough data are at hand—due to the estimation of optimal interpolation parameters, the interpolated model will in the 'worst' case again result in a conventional non-interpolated Markov chain.

For the classification of promoter regions, we could demonstrate on the test set of Fickett and Hatzigeorgiou (1997) that our method performs equally or better than any signalor content-based method in the survey. Signal-based approaches rely on the application of position-specific models, e.g. neural networks or weight matrices trained on a frequently occurring pattern such as the TATA box or the initiator site. In the case of general purpose promoter prediction where no certain combination of transcription factor binding sites is expected in advance, the judgement of the overall sequence proves to be equally suitable. Further research towards the integration of content- and signal-based approaches therefore seems appropriate; a first step in this direction was described by Solovyev and Salamov (1997).

In our opinion, another important factor for the success of our promoter recognizer is the competition of several models. Promoter predictors which only consist of a model for promoter sequences and rely on a certain fixed threshold have to meet the problem that it often depends not only on the sequence itself, but also on the particular context whether a region is functionally active. Because we use several models and judge the difference of the particular likelihoods, this is implicitly captured.

The integration of a promoter recognition module into gene parsers like GENIE (Kulp *et al.*, 1996) or GenScan (Burge and Karlin, 1997), where the different sensors are trained separately and can be easily exchanged, is in principle straightforward. However, up to now, the only system incorporating a promoter module is GenScan, and this is a fairly simple model incorporating weight matrices for the TATA and the initiator region, coupled with a null model to cope with promoters with a weakly conserved core region. According to Burge and Karlin (1997), this approach is due to the lack of sensitivity of current predictors. The performance of promoter prediction algorithms is still much worse than those for coding regions or signals involved in the transcription process, such as splice sites, and therefore a cautionless employment of a promoter module may lead to an overall deterioration of the system. Nevertheless, the good classification results for promoters versus exons, especially, lead us to the expectation that a future integration of our promoter recognizer into a gene parsing framework will be successful.

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