Abstract

Background: In addition to known protein-coding genes, large amounts of apparently non-coding sequence are conserved between the human and mouse genomes. It seems reasonable to assume that these conserved regions are more likely to contain functional elements than less-conserved portions of the genome.

Methods: Here we used a motif-oriented machine learning method based on the Relevance Vector Machine algorithm to extract the strongest signal from a set of non-coding conserved sequences.

Results: We successfully fitted models to reflect the non-coding sequences, and showed that the results were quite consistent for repeated training runs. Using the learned models to scan genomic sequence, we found that they often made predictions close to the start of annotated genes. We compared this method with other published promoter-prediction systems, and showed that the set of promoters which are detected by this method is substantially similar to that detected by existing methods.

Conclusions: The results presented here indicate that the promoter signal is the strongest single motif-based signal in the non-coding functional fraction of the genome. They also lend support to the belief that there exists a substantial subset of promoter regions which share several common features including, but not restricted to, a relative abundance of CpG dinucleotides. This subset is detectable by a variety of distinct computational methods.

Background

Since the publication of draft sequences for the human [1] and mouse [2] genomes, several groups have run large-scale comparisons of the sequences to detect regions of conserved sequence. An initial survey of these was published along with the draft mouse genome [2], with additional comparisons appearing since then [3]. Briefly, protein coding genes are – as we might expect – among the most strongly conserved regions, but homologous sequences can be found throughout the genome. In total, it is possible to align up to 40% of the mouse genome to human sequence [4], but it seems likely that at least some of this is just random "comparative noise" – regions of sequence which serve no particular purpose but which, purely by chance, have not yet accumulated enough mutations to make their evolutionary relationship unrecognizable. However, it is widely accepted that some of the noncoding-but-similar regions, especially those with the...
highest levels of sequence identity between the two species, are preferentially conserved because they perform some important function. It has been estimated that around 5% of the genome is under purifying selection [2], indicating that mutations in these regions have deleterious effects: a strong suggestion of some important function.

Here, we apply the Eponine Windowed Sequence (EWS) sequence analysis method which uses a Relevance Vector Machine (RVM) [5] to extract a minimal set of short motifs which are able to discriminate between two sets of sequences: in this case, a positive set of conserved non-coding sequences and a negative set of randomly picked non-coding sequences. The EWS model is an adaptation of the Eponine Anchored Sequence (EAS) model, first applied for transcription start site prediction in [6] and subsequently used to predict a range of additional biological features including translation start sites and transcription termination sites [A. Ramadass, unpublished]. While EAS is designed to classify individual points in a sequence – a feature which allows the model to predict precise locations for features such as transcription start sites – EWS classifies complete blocks (windows) of sequence. The basis functions (inputs) of the RVM are sums of position-weight matrix scores [7] across the whole window.

**Results**

We considered a set of alignments made by the blastz program [4] between release NCBI33 of the human genome and release NCBI30 of the mouse genome. Since unprocessed blastz aligns around 40% of human sequence to the mouse genome, we chose to focus on the 'tight' alignments. These are a subset of alignments which are rescored and thresholded using a set of parameters given in [4], and cover only around 5.6% of the human genome – a proportion much closer to the fraction of non-repetitive portions of chromosome 6, using the same criteria to define repeats and coding sequence. While it is probable that this set also included some functional sequences, we would expect them to be represented at a substantially lower level than in the conserved set.

In total, the tight blastz set contained 787173 blocks of sequence with high-scoring alignments between the two genomes. We considered only those blocks assigned to human chromosome 6, a 170 Mb chromosome which has recently undergone manual annotation of gene structures and other features [8]. This chromosome included 44105 (5.6%) of the total alignments. These varied in length from 34 to 9382 bases, with a length distribution skewed towards relatively short alignments, as shown in figure 1.

![Histogram showing number of alignments covering human sequences of various lengths.](image)

**Figure 1**

Blastz alignments between human chromosome 6 and the mouse genome. Histogram showing number of alignments covering human sequences of various lengths.

Since we were interested in non-coding features of the genome, we ignored all regions where an alignment overlapped an annotated gene structure. This removed 20.8% of aligned bases. It is possible that some genes, and especially pseudogenes, have been missed by the annotation process, so we also removed portions covered by ab initio gene predictions from the Genscan program [9]. This eliminated an additional 4.3% of aligned bases. Finally, repetitive sequence elements annotated by the programs RepeatMasker [10] and trf [11] (5.9%) were removed from the working set. The remainder of the aligned regions were split into non-overlapping 200 base windows, ignoring any portions less than 200 bases. This gave a set of 13925 sequences which are well-conserved between human and mouse – and therefore likely to be functional – but which are very unlikely to be part of the protein-coding repertoire. These formed the positive training set for our machine learning strategy.

A negative training set of equal size was prepared by picking 200-base windows at random from the non-coding, non-repetitive portions of chromosome 6, using the same criteria to define repeats and coding sequence. While it is probable that this set also included some functional sequences, we would expect them to be represented at a substantially lower level than in the conserved set.

These two sets of sequence were presented to the Eponine Windowed Sequence machine learning system, as described in the methods section. Randomly chosen 5-base words were used as seed motifs, and three independent training runs were performed, each for 2000 cycles. The set of motifs used in model 1 is shown in table 1.

While the exact set of motifs used in the model varied somewhat from run to run, testing pairs of models on non-overlapping windows from a 1 Mb region of human chromosome 22 and plotting the scores showed that the model outputs were highly correlated (e.g. figure 2). We calculated the Pearson correlation coefficient for all pairs, and in all cases this was greater than 0.96. From this...
strong correlation, we concluded that any variations in the model were simply the result of the trainer picking one representative from a group of motifs which provide similar information.

We scanned genomic sequences using these models at a range of thresholds, and examined the results on the Ensembl genome browser [12] using a Distributed Annotation System [13] server. Visual inspection showed that many of the highest-scoring regions were localised near the start of genes. This prompted us to look at the distribution of high-scoring sequences with respect to the starts of a set of well-annotated genes. We considered the GD_mRNA genes from version 2.3 of the human chromosome 22 annotation. These are confidently annotated genes with experimental evidence as described in [14], which confirms at least the approximate location of the ends of the transcripts, and are independent from the chromosome 6 training data. Figure 3 shows the density of predictions with EWS scores $\geq 0.90$ relative to the annotated 5' ends of these genes. This shows a strong peak of predictions close to the annotated starts, demonstrating that the model is predicting some sequences commonly located around the transcription start site of genes. Combining this observation with the fact that the model was trained from conserved (and therefore presumed functional) sequences, we believe that it is detecting signals found in the promoter regions of genes.

Table 1: Motifs used in EWS homology model 1. The entries in this table show consensus sequences of the weight matrices used in the model (note that it is possible for two distinct weight matrices to have the same consensus sequence). Motifs are listed in both forwards and reverse-complement orientation, and the two sections of the table indicate whether that motif is given a positive or negative weight in the learned linear model.

| Postive | Reverse | Negative | Reverse |
|---------|---------|----------|---------|
| Forward | Reverse | Forward | Reverse |
| gcca    | tgac    | taccgt   | accgt   |
| tatgt   | caata   | gggca    | tggcc   |
| tgcca   | tggca   | gcca     | tggac   |
| ggca    | tgcct   | acaat    | acgt    |
| tacgt   | acgta   | gggc     | gcgcc   |
| tatac   | agtac   | tact     | agta    |
| taac    | gtaa    | cctcc    | gppag   |
| tttt    | aaaa    | gcca     | tggc    |
| acaat   | atgt    | tattg    | catta   |
| caatt   | aattg   | tattg    | catta   |
| caggc   | gcggc   | aatttt   | aatttt  |
| cag     | cctg    | caatt    | atgt    |
| cggct   | atcgg   | gatt     | ataccg  |
| aaatt   | aatt   | cctgg    | cctgg   |
| gctcg   | ctpgc   | cattg    | cttg    |
| gcc     | gcc     | act      | agt     |
| aaaaa   | tttt    | caatt    | cttta   |

**Figure 2**
Correlation of model scores. Scatter plot showing the scores of EWS models 1 and 2 on a set of human sequences.

**Figure 3**
Localisation of predictions. Density of predictions from one of the homology models around known gene starts on human chromosome 22.
precise location at which transcription begins. In [6], we developed a pseudochromosome, derived from release 2.3 of the chromosome 22 annotation. As described above, this includes a subset of 284 experimentally verified gene structures. The pseudochromosome was constructed to include these genes while omitting all other annotated genes (which could be substantially truncated). We considered predictions (groups of one or more overlapping windows which all have scores greater than some chosen threshold) to be correct if they lie withing 2 kb of an annotated gene start, and false otherwise. Plotting accuracy (proportions of predictions which are correct) against coverage (proportion of transcript starts which are detected by one of the correct predictions) gives a Receiver Operating Characteristic (ROC) curve. Using this criterion, a totally random set of predictions would be given an accuracy of around 0.07. ROC curves are plotted for the three independently trained models in figure 4. Firstly, this shows that predictive performance for all three models is rather similar. It also shows that they can function as accurate promoter predictors, with accuracy rising to a plateau of around 0.7, much higher than expected for random predictions.

We picked model 1 for further study. Using a score threshold of 0.91, this gives an accuracy of 0.68 and a coverage of 0.31. We compared the set of genes correctly detected by this model to two other methods: firstly, the EponineTSS predictor described in [6], and secondly, the published results from the PromoterInspector program [15]. PromoterInspector results were mapped to pseudochromosome coordinates using the procedure described in [6]. Figure 5 shows how the set of promoters detected by these three distinct methods overlaps. There are clearly strong correlations between all three methods. In particular, at this threshold the EWS homology model detects 98 promoters which were found by at least one of the other methods, but only 4 novel promoters.

We investigated the robustness of the signal learned by this process by retraining models with a variety of seed word sizes, from 2 to 6 bases. During training, motifs can be trimmed to lengths shorter than that of the seed words (down to a minimum of 2 bases) but can never grow longer than the seed word size. When evaluated on the pseudochromosome, the resulting models always showed a preference for regions around gene starts, regardless of word length, as shown in figure 6. However, the accuracy was reduced when using short seed words – particularly words of length of 2. The best accuracy was seen for a seed word length of 5, and decreased somewhat for words of length 6.

This suggests that a large fraction (but not all) of the information learned by these models can be encoded in dinucleotide frequencies. It is well known that many transcription start sites are close to regions of relatively high CpG dinucleotide composition (CpG islands) [16]. To investigate the contribution that CpG dinucleotides make to our models, we deleted all CpG dinucleotides from the

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**Figure 4**

Accuracy and coverage of TSS prediction. Plots of Accuracy vs. coverage at a range of score thresholds (ROC curves) for three independently trained homology models.

**Figure 5**

Comparison of TSS prediction methods. Sets of pseudochromosome promoters correctly predicted by three different prediction methods: EponineTSS [6] with a score threshold of 0.999, PromoterInspector (labelled “Prom’spectr”), and the homology-EWS model 1 with a score threshold of 0.91 (“Homol_1”).
training data, then re-evaluated the resulting models on the pseudochromosome (also with CpG dinucleotides removed), as shown in figure 7. Perhaps not surprisingly, dinucleotide models now show very little tendency to detect gene starts. However, as the word size increases, the preference for gene starts gradually increases, until a seed size of 6 gives an accuracy comparable to that see when CpG dinucleotides are included, although the maximum coverage before accuracy begins to drop rapidly is somewhat lower. Broadly similar results are seen if CpG dinucleotides are randomly replaced with other dinucleotides.

**Conclusions**

We have shown here that, when presented with a set of non-coding sequences which are strongly conserved between human and mouse, a simple motif-oriented machine learning system consistently builds models which are able to detect a substantial fraction of human promoter regions with good accuracy. This strongly suggests that this promoter signal represents the most widely used motif-based signal in functional non-coding sequence. While the model learned here can clearly be applied for the purpose of genome-wide promoter annotation, in practise existing methods offer better coverage and (in the case of the EponineTSS predictor) predictions for the precise location of the transcription start site.

It is interesting that the promoter model learned by this technique detected substantially the same set of promoters as found by the EponineTSS and PromoterInspector methods. It has previously been remarked that these two methods detect similar sets [6], but this could perhaps be explained by the fact that both methods were initially derived from similar sets of known promoter sequences (in both cases, training data was extracted from the EPD database [17]. In the case of the homology models described here, there is no connection with EPD, or any similar set of known promoters: the training data was picked purely on the basis of its high similarity to corresponding portions of the mouse genome. These results therefore support the alternate view that there is a particular 'easily detected' subclass of promoter sequences.

One distinct group of promoters, which previous results show may correspond to this easily detected family, is the set of promoters associated with CpG islands [16]. However, while a number of the motifs listed in table 1 are G/C rich and/or contain the CpG dinucleotide, by no means all of the motifs match this description, and indeed one motif containing CpG has a negative weight in the linear model – its presence in a sequence will reduce the model’s output score – while some A/T rich motifs have positive weights. We therefore believe that the signals detected here are significantly more complex than a simple over-representation of CpG dinucleotides. Experiments with smaller seed-word sizes support this assumption: while dinucleotide-based models were also able to predict promoter regions, the accuracy was lower than for models including longer motifs. Finally, we show that while the predictive capacity of dinucleotide models is largely eliminated once CpG dinucleotides are removed from the sequence, models including longer words are still able to make correct promoter predictions in many cases. So
while CpG dinucleotides are an important contribution to
the promoter signal, they are clearly not the only
component.

Methods
Genomic sequence and annotation
Human genome sequence release NCBI33 and mouse
genome release NCBIM30 were extracted from Ensembl
databases [12], which also contained gene predictions
from Genscan [9] and repeat data from RepeatMasker [10]
and trf [11]. Curated annotation of gene structures on
human chromosome 6 was obtained from the Vega data-
bases [18]. Vega and Ensembl data was extracted directly
from the SQL databases using the BioJava toolkit with bio-
java-ensembl extensions [19].

Genome alignments
Human-mouse genome alignments were generated by the
blastz alignment program. These were subsequently re-
scored and filtered to give a 'tight' set of high-confidence
alignments, as described in [4]. We downloaded the tight
alignment set from the UCSC genome website [20].

Pseudochromosome for testing promoter-finding methods
A 16.3 Mb pseudochromosome sequence was produced
based on version 2.3 of the curated annotation for human
chromosome 22. This includes all the experimentally-val-
iddated gene structures and their upstream regions, while
omitting regions containing genes that are predicted but
not fully verified. In the case of a pair of divergent genes
where one has been verified and the second has not, their
shared upstream region was cut at the midpoint. More
information about pseudochromosome construction is
given in [6].

Eponine Windowed Sequence learning
The Eponine Windowed Sequence (EWS) model is
designed by analogy to the Eponine Anchored Sequence
model first described in [6], but rather than targeting indi-
vidual points in the sequence, it is designed to classify
small regions or windows of a sequence, based purely on
their own sequence content.

The EWS model uses the Relevance Vector Machine [5]
algorithm to drive the training process. Relevance Vector
Machines solve classification and regression problems by
building Generalised Linear Models (GLMs) as weighted
sums of a "working set" of basis functions. During the
training process, those basis functions which are not
informative are given weights close to zero and eventually
discarded from the working set. To explore very large sets
of possible basis functions, it is possible to add extra basis
functions during the course of the training process [6].

The "sensors" of the EWS model are DNA position-weight
matrices [7], which make convenient models of short
sequence motifs. When using weight matrices to analyse
sequence windows, we sum the weight matrix probability
scores for all possible positions within the sequence.
Normalising for the length of the sequence being
inspected and the size of the PWM, the basis functions of
the model take the form:

$$\phi(S) = \frac{4|W|}{|S| - |W| + 1} \sum_{i=1}^{S-|W|+1} W(S_i^j W_{i-j}^j)$$

where $W(s)$ is the probability that sequence $s$ was emitted
by weight matrix $W$, $|S|$ is the sequence length, $|W|$ is the
weight matrix length, and $S_i^j$ denotes a subsequence from
$i$ to $j$.

An initial set of basis functions is proposed by taking all
possible DNA motifs of a specified length (typically 5)
and generating weight matrices which preferentially rec-
ognise these motifs. As the relevance vector machine
trainer removes non-informative basis functions from the
working set, they are replaced by applying one of the fol-
lowing sampling strategies to a basis function picked ran-
domly from the working set:

- Generate a new weight matrix in which each column is
  a sample from a Dirichlet distribution with its mode equal
to the weights in the corresponding column of the parent
  weight matrix.

- Generate a new weight matrix one column shorter than
  the parent by removing either the first or the last column.

By using these sampling rules, the trainer is able to explore
motif space. The process of generating candidate motifs
using these rules then selecting the most informative
using the RVM can be seen as a form of genetic algorithm.

Authors’ contributions
TD and TH conceived and designed this study, and ana-
lysed results. TD implemented the Eponine machine
learning system and drafted the manuscript. All authors
read and approved the final manuscript.

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