PriorsEditor: a tool for the creation and use of positional priors in motif discovery

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ABSTRACT

Summary: Computational methods designed to discover transcription factor binding sites in DNA sequences often have a tendency to make a lot of false predictions. One way to improve accuracy in motif discovery is to rely on positional priors to focus the search to parts of a sequence that are considered more likely to contain functional binding sites. We present here a program called PriorsEditor that can be used to create such positional priors tracks based on a combination of several features, including phylogenetic conservation, nucleosome occupancy, histone modifications, physical properties of the DNA helix and many more.

Availability: PriorsEditor is available as a web start application and downloadable archive from http://hare.medisin.ntnu.no/priorseditor requires Java 1.6). The web site also provides tutorials, screenshots and example protocol scripts.

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1 INTRODUCTION

Computational discovery of transcription factor binding sites in DNA sequences is a challenging problem that has attracted a lot of research in the bioinformatics community. So far more than a hundred methods have been proposed to target this problem (Sandve and Drabløs, 2006) and the number of publications on the topic is steadily increasing.

There are two general approaches for discovering potential transcription factor binding sites with computational tools. One is to examine regulatory regions associated with a group of genes that are believed to be regulated by the same factors and search for patterns that occur in all or most of these sequences. This approach, often referred to as de novo motif discovery, can be used when we have no prior expectations as to what the binding motifs might look like. One concern with this approach, however, is that it might be necessary to consider rather long sequence regions to ensure that the target sites are indeed covered. Since binding motifs for transcription factors are usually short and often allow for some degeneracy, the resulting signal-to-noise ratio can be quite low, making it difficult to properly discriminate motifs from background. Another problematic issue is that DNA sequences inherently contain a lot of repeating patterns, such as tandem repeats and transposable elements, which can draw focus away from the target binding motifs when searching for similarities between sequences.

One general motif discovery approach, called motif scanning, searches for sequence matches to previously defined models of binding motifs, for instance in the form of position weight matrices (PWMs; Stormo, 2000). The main drawback with motif scanning is that it tends to result in an overwhelming number of false positive predictions. According to the ‘futility theorem’ put forward by Wasserman and Sandelin (2004), a genome-wide scan with a typical PWM could incur in the order of 1000 false hits per functional binding site, which would make such an approach practically infeasible for accurate determination of binding sites. The problem here lies not so much in the predicted binding patterns themselves, since many of these would readily be bound by transcription factors in vitro. In vivo, however, most such binding sites would be non-functional, perhaps because the chromatin conformation around the sites precludes access to the DNA (Segal et al., 2006) or because the target factors require the cooperative binding of additional factors nearby to properly exert their regulatory function (Ravasi et al., 2010).

One way to improve accuracy in motif discovery is to try to narrow down the sequence search space as much as possible beforehand, for instance, by masking out portions of the sequences that resemble known repeats or considering only sequence regions that are conserved between related species (Duret and Bucher, 1997). Kolbe et al. (2004) introduced a measure they called ‘Regulatory Potential’ which combines phylogenetic conservation with distinctive hexamer frequency profiles to identify possible regulatory regions. This measure calculates a score for each position along the sequence, and regions receiving higher scores are deemed more likely to have a regulatory role. Regulatory Potential can be considered as an example of a ‘positional prior’ since each position is associated with an a priori probability of possessing some specific property. Positional priors can be used as an aid in motif discovery by assigning high prior values to regions that we consider more likely to contain functional binding sites and then focus the search on these regions. Besides conservation and oligonucleotide frequencies, other features that can be relevant for assigning prior values include: localized physical properties of the DNA double helix, distance from transcription start site or other binding sites, ChIP-chip and ChIP-seq data, and potentially tissue-specific epigenetic factors such as the presence of nucleosomes and associated histone modifications. Many of the aforementioned features have previously been applied and shown to improve the performance of motif discovery by themselves (see e.g. Bellora...
Fig. 1. The top left panel in this screenshot shows examples of some of the features that can be used as a basis to create positional priors. These features are visualized as data tracks in the main panel for a selected set of sequences. The bottom-most track contains predicted matches to TRANSFAC and JASPAR motifs in regions with non-zero RegulatoryPotential7X scores.

We present here a program called PriorsEditor, which allows users to easily construct positional priors tracks by combining various types of information and utilize these priors to potentially improve the motif discovery process (Fig. 1).

2 SOFTWARE DESCRIPTION

The first step in constructing a priors track with PriorsEditor is to specify the genomic coordinates for a set of sequences one wishes to analyze. Next, data for various features can be imported to annotate these genomic segments. PriorsEditor supports three types of feature data. The first type, numeric data, associates a numeric value with each position in the sequence and can be used to represent features such as phylogenetic conservation scores, DNA melting temperatures and nucleosome-positioning preferences. Numeric data tracks are also used to hold the final positional priors. The second feature type, region data, can be used to refer to continuous stretches of the DNA sequence that share some unifying properties which distinguish them from the surrounding sequence. Different regions are allowed to overlap, and regions can also be assigned values for various attributes, including type designations, score values and strand orientations. Features best represented as regions include genes, exons, repeat regions, CpG-islands and transcription factor binding sites. The last feature type, DNA sequence data, represents the DNA sequence itself in single-letter code. DNA sequence data can be passed on to motif discovery programs for further analysis, and it can also be used to estimate various physical properties of the DNA double helix, such as GC content, bendability and duplex-free energy. Additional feature data can be obtained from web servers such as the UCSC Genome Browser (Rhead et al., 2010) or be loaded from local files.

Once the data for the desired features have been loaded, the data tracks can be manipulated, compared and combined to create a priors track using a selection of available operations. These include operations to extend regions by a number of bases upstream and/or downstream, merge overlapping regions or regions within close proximity, filter out regions, normalize data tracks, smooth numeric data with sliding window functions, interpolate sparsely sampled
of operations can be automatically applied to new sequence sets while in recording mode. With protocol scripts these same series of operations can be programmed to repeat the same procedure to analyze other sequence sets as and requires several steps to complete. If a user discovers a good set of starting up the normal graphical interface. This allows the user to run PriorsEditor from a command-line interface instead of starting up the normal graphical interface. This allows the construction and use of positional priors to be incorporated into a batch-processing pipeline.

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