## Timeline of Deliverables

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March 14, 2016

**Proposed title:** Locating Transcription Factor Binding Sites: Methods in search of *moralistic* motifs

Expected length: 8-10 pages

**References:** See References section. In places where required I have mentioned relevant review articles that I would be keeping as my main reference.

Key reference: Assessing computational tools for the discovery of transcription factor binding sites [1]

**Expected duration**: 6 weeks(14<sup>th</sup> March to 30<sup>th</sup> April)

Section Organization: See table 2

Tool	Original Findings	Principle	Reference	$de ext{-}novo$
AlignACE	Discovery of over represented motifs in unaligned sequences, typically in the upstream region of corregulated genes	Gibbs sampling	[2]	1
ANN-Spec	finding low-complexity patterns present in high frequency. Suit- able for locating TFBS given a background sequence	Artificial Neural Network for parameter fitting to maximize posterior probability and then Gibbs sampling for	[3]	/
Consensus	Statistically significant alignments of DNA or protein sequences to determine evolutionary/functional perspective	Greedy algorithm that searches for motifs maximizing information iteratively(Fixed width)	[4]	<b>✗</b> (Check, constraint: Fixed width)
Consite	TFBS prediction using phylogenetic footprinting	Accounts for evolutionary constraints by aligning regulatory sequence from orthologous pairs of genes	[5]	х
GLAM	Locating functional sites by MSA	Simulated Annealing for automatically determining the width (An improvement over Stormo's Gibbs sampling approach [4]	[6]	1
The Improbizer	Identifying cis-regulatory elements that activate gene- expression within pharyngeal gene clusters	EM algorithm, zero, first or second order markov model for background sequences	[7]	/
MEME	Over-represented motifs in DNA, protein sequences	EM algorithm to search for optimum motif	[8]	<b>✓</b>
MONKEY	Identifying conserved transcription factor binding sites in MSA	Probabilistic model of binding site-specificity accounting for evolutionary tree by modeling one as background	[9]	х
CENTIPEDE	Used to predict genome wide map of 800K TFBS of 200+ TFs	Integrates histone modification, gene annotation, DNAse I cleavage pattern to predict TFBS	[10]	<b>✓</b>
$\operatorname{SeqGL}$	TFBS prediction using ChIP, DNAse or ATAC-seq data	Group lasso regularization to extract most important <i>k-mer</i> groups distinguishing peaks from flanking sequences followed by motif finding across regions that have non zero weight	[11]	<b>/</b>
YMF	Statistically significant motifs	Given regulatory regions of <i>related</i> genes, find motifs with greater Z-score	[12]	х
Weeder	Predicting Regulatory motifs	Models the significant occurrence of motifs over a seventh order markov chain expected background	[13]	<b>/</b>
TFEM	TFBS prediction	Position specific priors based on phylogenetic conservation, pe- nalization based on deviation from conserved profiles	[14]	<b>√</b>

Kellis et al.	Prediction of regulator target for drosophila	BLS: Branch Length score based cutoffs for finding most significant motifs	[15]	<b>✓</b>
PhyloGibbs	Regulatory motif finding in multiple local sequence alignment of orthologous sequences	MCMC, simulated annealing based approach that treats alignments as the sites for binding and intergenic DNA as 'background', taking into account evolutionary distances	[16]	×
REDUCE	Discovering cis-regulatory sequences using expression data without the need of gene clustering	Models the log fold change expression as a linear with the number of occurrences(or Information content) of motif as covariates. Motif lengths are determined by user	[17]	х
GMEP	Modeling Sequence to expression(S2E) profiles. Hierarchical clustering of GMEP identified clusters of motifs with known TFs	Enumerate motifs for different length to find the weight contri- bution to gene expression, sim- ilar to the REDUCE algorithm discussed above	[18]	×
EMnEM/PhyME	Identify motifs in orthologous sequences	Phylogenetic EM based approach	[19, 20]	Х
RCADE	Motif discovery in C2H2-ZF ChIP-seq data	Use a previously established recognition code for C2H2 to identify motifs in target sequences, which are then tested for enrichment using sequences from endogenous retroelements(ERE) and non-ERE regions	[21]	×
DME	Identifying tissue specific TFBS	Identifies motifs over- represented in one set of sequences over the back- ground(promoters of deferen- tially expressed genes)	[22]	<b>✓</b>
INSIGHT	Not for motif discovery, but to gauge impact of mutations on TFBS	Probabilistic model to measure impact of natural selection on TFBS	[23]	х
rVISTA	Finding cis-regulatory sequences	Clustering of TFBS and interspecies sequence conservation	[24]	Х
TRAWLER	Regulatory motif discovery pipeline	Uses a suffix-tree implementation and a Z-score approximation	[25]	<b>√</b>
MEME-prior	TFBS prediction	Prior probabilities based on phylogenetic or other background information assigned to bases	[26]	/
FootPrinter	Regulatory motif prediction in homologous sequences	Uses MSA and evolutionary conservation to determine motifs	[27]	Х

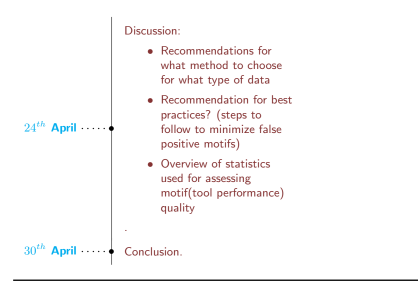
 ${\bf Table\ 1:\ Tools,\ purpose,\ methods}$ 

## References

[1] Martin Tompa, Nan Li, Timothy L Bailey, George M Church, Bart De Moor, Eleazar Eskin, Alexander V Favorov, Martin C Frith, Yutao Fu, W James Kent, Vsevolod J Makeev, Andrei A Mironov, William Stafford Noble, Giulio Pavesi, Graziano

Table 2: Timeline of deliverables

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$14^{th}$ March · · · · · •	Deliverables proposed.
$21^{st}$ March $\cdots \bullet$	Page 1-2: Introduction:  • Why TFBSs matter?: Biological motivation  • TFBS prediction: Computational Challenges(Hints from review article [28])
$2^{nd}$ April $\cdots \bullet$	Page 3-5:TFBS finding: A two step approach:  1. Representation of Motif(Consensus Based or Profile Based) [Comprehensive Review: [29]]  • Position Weight Matrices (PWMs) [30], with gaps and correlation [31, 4]  • Dinucleotide Weight Matrices(DWMs) [32]  • Other Variable width permitting models [33, 34, 35]  • Regression Based models [36, 37]  • Energy Modeling [38]  2. Identifying binding sites given a motif representation
$17^{th}$ April $\cdots \bullet$	Page: 6-10 Motif Discovery:  • Motif discovery  - Comparative     Approaches , See     table 1  - de-novo approaches  • Motif discovery and     Phylogenetic footprinting     - See table 1



Pesole, Mireille Rgnier, Nicolas Simonis, Saurabh Sinha, Gert Thijs, Jacques van Helden, Mathias Vandenbogaert, Zhiping Weng, Christopher Workman, Chun Ye, and Zhou Zhu. Assessing computational tools for the discovery of transcription factor binding sites. *Nature Biotechnology*, 23(1):137–144, January 2005.

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