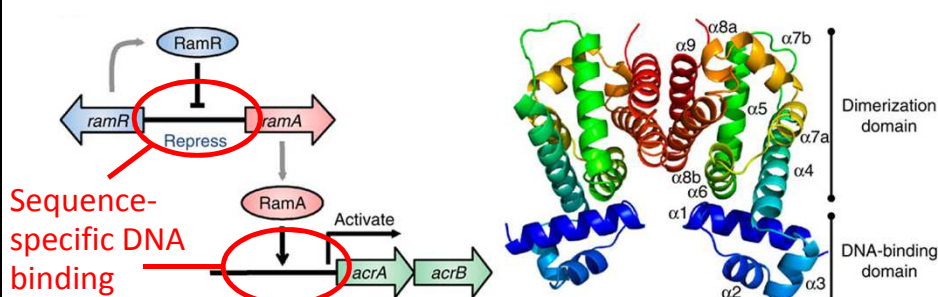


Motifs

BCH364C/391L Systems Biology / Bioinformatics – Spring 2015
Edward Marcotte, Univ of Texas at Austin

Edward Marcotte/Univ. of Texas/BCH364C-391L/Spring 2015

An example transcriptional regulatory cascade
Here, controlling *Salmonella* bacteria multidrug resistance

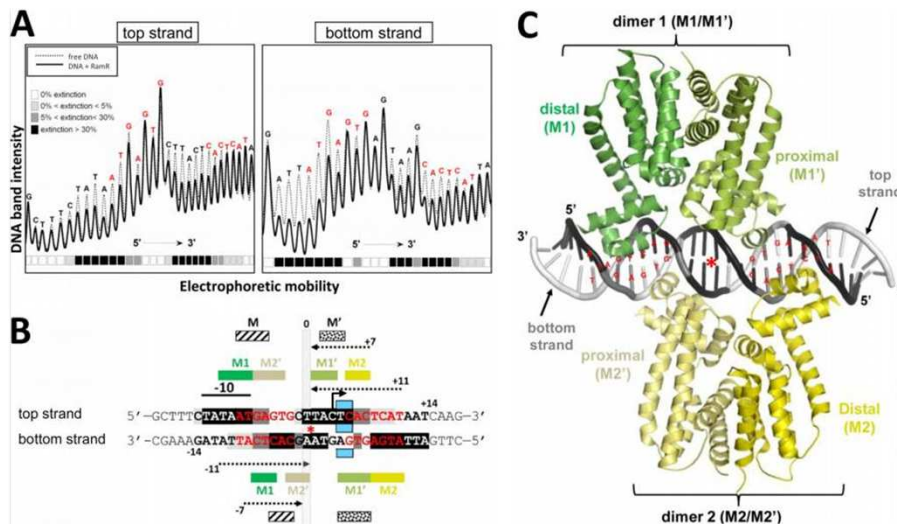


RamR represses the *ramA* gene, which encodes the activator protein for the *acrAB* drug efflux pump genes.

RamR dimer

Nature Communications 4, Article number: 2078 doi:10.1038/ncomms3078

Historically, DNA and RNA binding sites were defined biochemically (DNase footprinting, gel shift assays, etc.)



Hydroxyl radical footprinting of *ramR-ramA* intergenic region with RamR

Antimicrob Agents Chemother. Feb 2012; 56(2):942-948.

Historically, DNA and RNA binding sites were defined biochemically (DNase footprinting, gel shift assays, etc.)

Now, many binding motifs are discovered bioinformatically

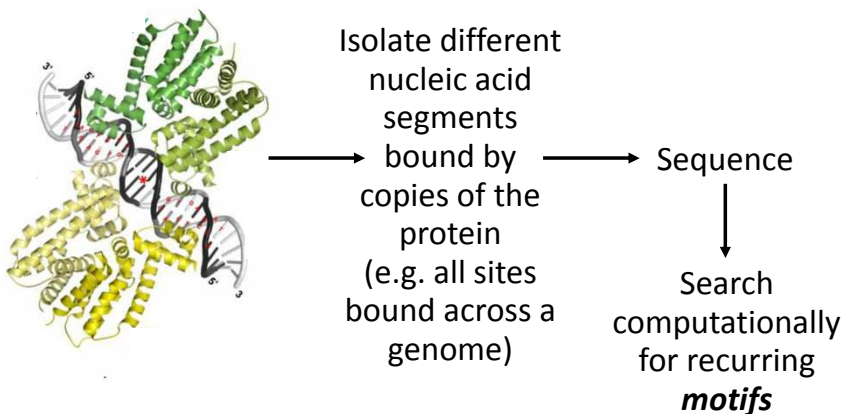
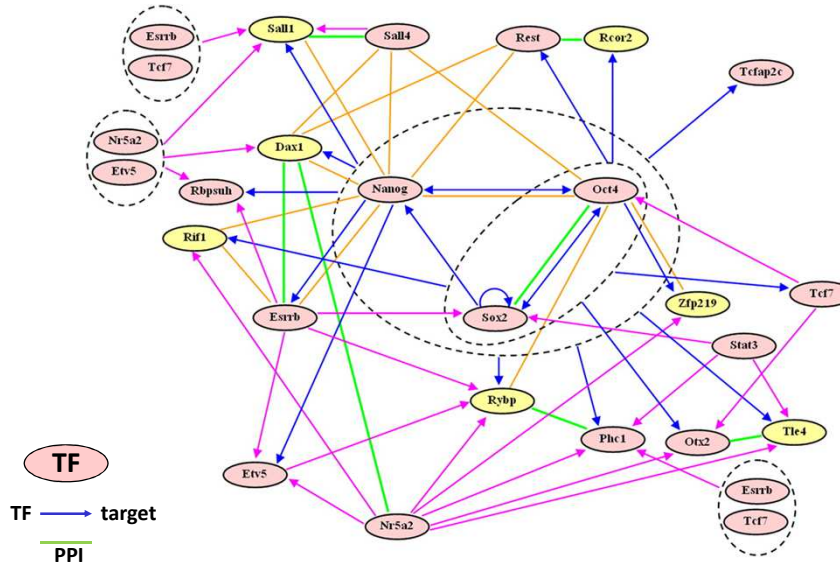


Image: Antimicrob Agents Chemother. Feb 2012; 56(2):942-948.

Transcription factor regulatory networks can be highly complex, e.g. as for embryonic stem cell regulators



<http://www.pnas.org/content/104/42/16438>

MOTIFS

HEM13	CCCATTTGTTCTC
HEM13	TTTCTGGTTCTC
HEM13	TCAATTGTTTAG
ANB1	CTCATTGTTGTC
ANB1	TCCATTGTTCTC
ANB1	CCTATTGTTCTC
ANB1	TCCATTGTTCGT
ROX1	CCAATTGTTTG

Binding sites of the transcription factor ROX1

consensus

A	002700000010
C	464100000505
G	000001800112
T	422087088261

frequencies



$$I_{\text{seq}}(i) = - \sum_b f_{b,i} \log_2 \frac{f_{b,i}}{p_b}$$

↑
freq of nuc b in genome

frequency of nuc b at position i

NATURE BIOTECHNOLOGY VOLUME 24 NUMBER 4 APRIL 2006

So, here's the challenge:

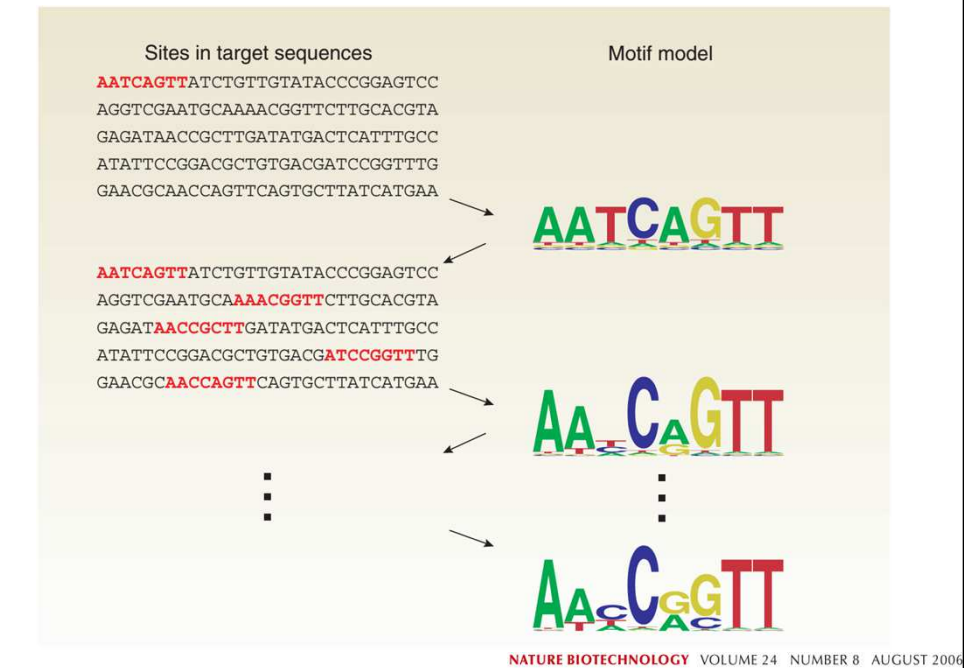
**Given a set of DNA sequences
that contain a motif (e.g.,
promoters of co-expressed
genes), how do we discover it
computationally?**

**Could we just count all
instances of each k -mer?**

Why or why not?

**→ promoters and DNA
binding sites are not well
conserved**

How does motif discovery work?

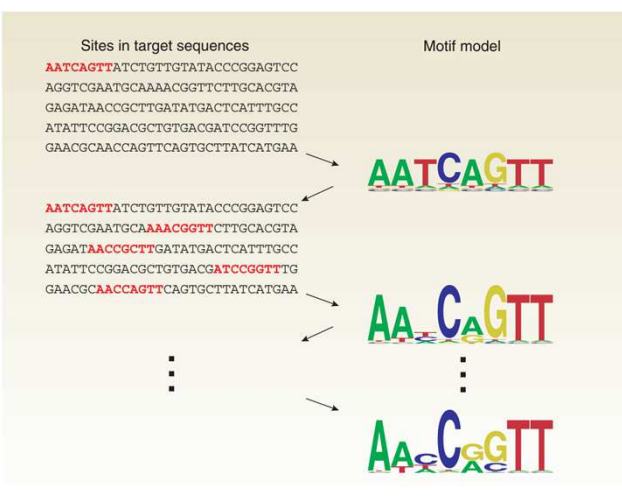


How does motif discovery work?

Assign sites to motif

Assign sites to motif

Assign sites to motif



Update the motif model

Update the motif model

Update the motif model

What does this process remind you of?

NATURE BIOTECHNOLOGY VOLUME 24 NUMBER 8 AUGUST 2006

How does motif discovery work?

Motif finding often uses expectation-maximization (like the k-means clustering we already learned about), i.e. alternating between building/updating a motif model and assigning sequences to that motif model.

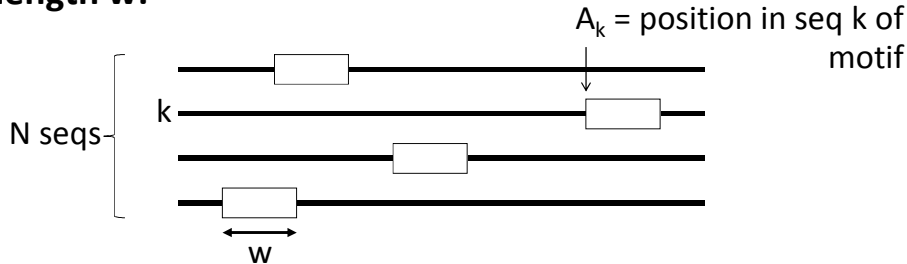
Searches the space of possible motifs for optimal solutions without testing everything.

Most common approach = *Gibbs sampling*

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski, Jun S. Liu, Andrew F. Neuwald, John C. Wootton

We will consider N sequences, each with a motif of length w:



q_{ij} = probability of finding nucleotide (or aa) j at position i in motif
i ranges from 1 to w

j ranges across the nucleotides (or aa)

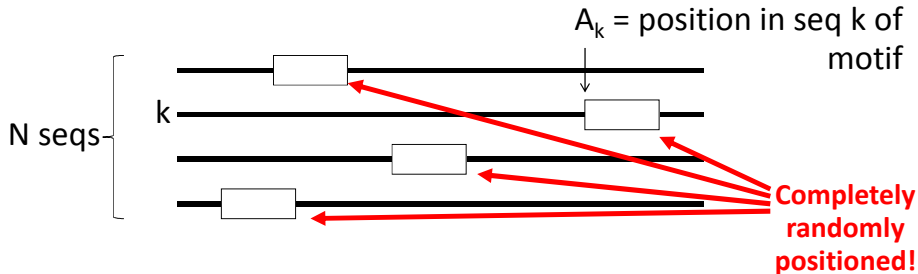
p_j = background probability of finding nucleotide (or aa) j

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski,
Jun S. Liu, Andrew F. Neuwald, John C. Wootton

NOTE: You won't give any information at all about what or where the motif should be!

Start by choosing w and randomly positioning each motif:



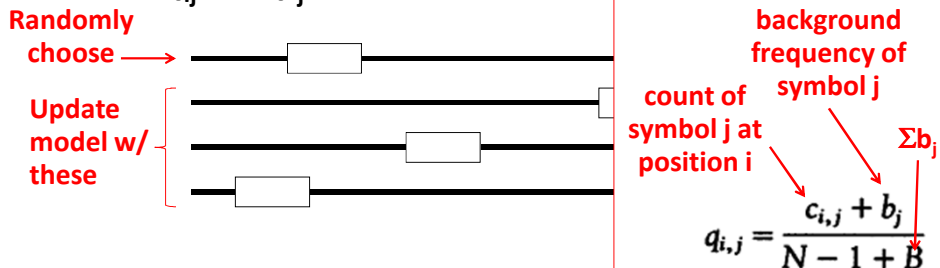
q_{ij} = probability of finding nucleotide (or aa) j at position i in motif
i ranges from 1 to w
j ranges across the nucleotides (or aa)
 p_j = background probability of finding nucleotide (or aa) j

SCIENCE • VOL. 262 • 8 OCTOBER 1993

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski,
Jun S. Liu, Andrew F. Neuwald, John C. Wootton

Predictive update step: Randomly choose one sequence, calculate q_{ij} and p_j from N-1 remaining sequences



q_{ij} = probability of finding nucleotide (or aa)
i ranges from 1 to w
j ranges across the nucleotides (or aa)
 p_j = background probability of finding nucleotide (or aa) j

p_j is calculated similarly from the counts outside the motifs

SCIENCE • VOL. 262 • 8 OCTOBER 1993

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

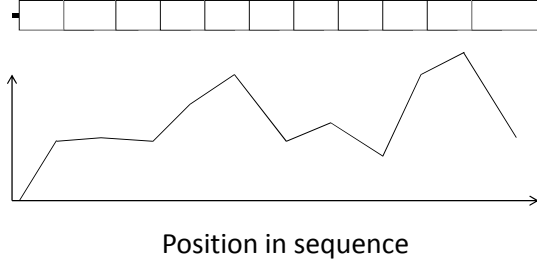
Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski,
Jun S. Liu, Andrew F. Neuwald, John C. Wootton

Stochastic sampling step: For withheld sequence, slide motif down sequence & calculate agreement with model

Withheld sequence →



Odds ratio of agreement with model vs. background



$$\frac{\prod (q_{ij})^{c_{xij}}}{\prod (p_j)^{c_{xij}}}$$

(see the paper for details)

SCIENCE • VOL. 262 • 8 OCTOBER 1993

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

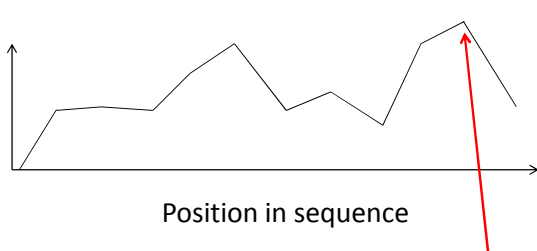
Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski,
Jun S. Liu, Andrew F. Neuwald, John C. Wootton

Stochastic sampling step: For withheld sequence, slide motif down sequence & calculate agreement with model

Withheld sequence →



Odds ratio of agreement with model vs. background



$$\frac{\prod (q_{ij})^{c_{xij}}}{\prod (p_j)^{c_{xij}}}$$

(see the paper for details)

Here's the cool part. DON'T just choose the maximum.
INSTEAD, select a new A_k position proportional to this odds ratio.

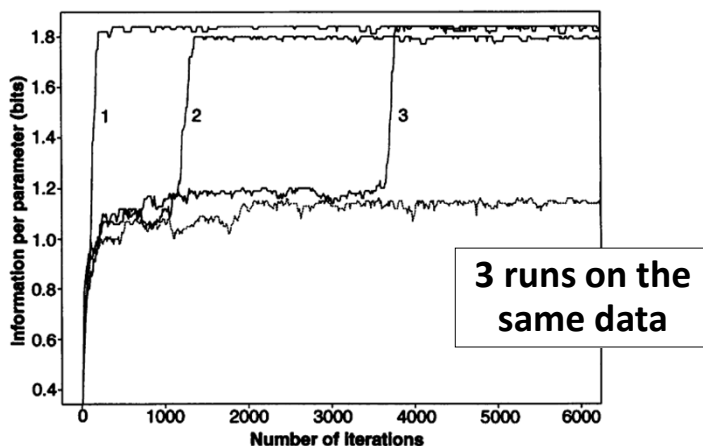
Then, choose a new sequence to withhold, and repeat everything.

SCIENCE • VOL. 262 • 8 OCTOBER 1993

Detecting Subtle Sequence Signals: A Gibbs Sampling Strategy for Multiple Alignment

Charles E. Lawrence, Stephen F. Altschul, Mark S. Boguski,
Jun S. Liu, Andrew F. Neuwald, John C. Wootton

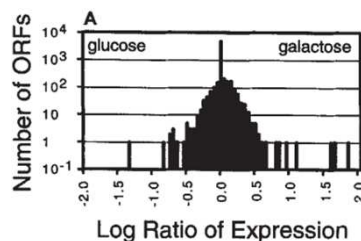
Over many iterations, this magically converges to the most enriched motifs. Note, it's stochastic:



SCIENCE • VOL. 262 • 8 OCTOBER 1993

Finding DNA regulatory motifs within unaligned noncoding sequences clustered by whole-genome mRNA quantitation

Frederick P. Roth^{1,*}, Jason D. Hughes^{1,2}, Preston W. Estep³, and George M. Church^{1,2}

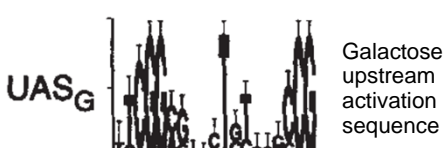


galactose vs. glucose → Measure mRNA abundances using DNA microarrays → Search for motifs in promoters of glucose vs galactose controlled genes

Discovered motifs



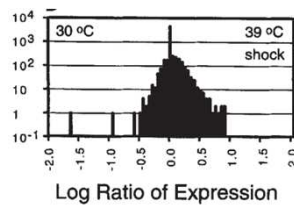
Known motif



"AlignAce" NATURE BIOTECHNOLOGY VOLUME 16 OCTOBER 1998

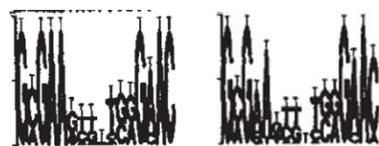
Finding DNA regulatory motifs within
unaligned noncoding sequences clustered
by whole-genome mRNA quantitation

Frederick P. Roth^{1,2}, Jason D. Hughes^{2,3}, Preston W. Estep², and George M. Church^{1,2*}

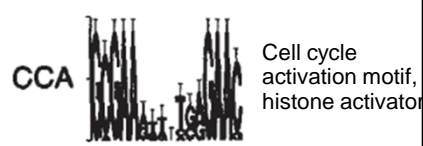


heat shock
vs.
30°C → Measure mRNA
abundances
using DNA
microarrays → Search for motifs in
promoters of heat-
induced and repressed
genes

Discovered motifs



Known motif



"AlignAce" NATURE BIOTECHNOLOGY VOLUME 16 OCTOBER 1998

If you need them, we now know the binding motifs for
100's of transcription factors at 1000's of distinct sites in
the human genome, including many new motifs.

e.g., <http://compbio.mit.edu/encode-motifs/>

Novel1	Novel2	Novel3	Novel6
BRCA1_disc1	EGR1_disc4	SP2_disc3	TATA_disc5
CHD2_disc1	ETS_disc1	TCF12_disc3	TATA_disc7
ETS_disc3	ETS_disc5	ZBTB7A_disc2	
ETS_disc6	ETS_disc7		
NR3C1_disc3	SETDB1_disc1		
ZBTB33_disc1	SIX5_disc1		
ZBTB33_disc2	SIX5_disc2		
ZBTB33_disc3	SIX5_disc3		
ZBTB33_disc4	SMARC_disc2		
	ZNF143_disc1		
	ZNF143_disc2		
	ZNF143_disc3		

2976-2987 Nucleic Acid Research, 2014, Vol. 42, No. 5
doi:10.1093/nar/gkt1249

Published online 11 December 2013

Systematic discovery and characterization of
regulatory motifs in ENCODE TF binding
experiments