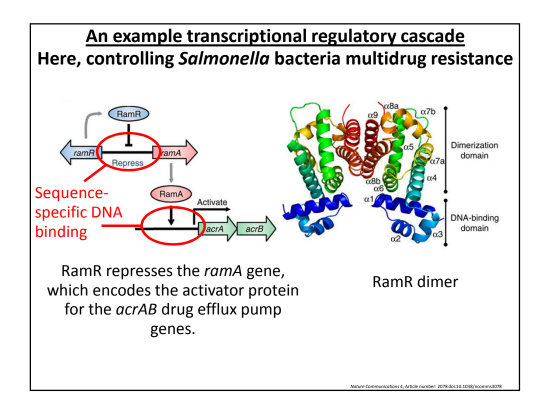
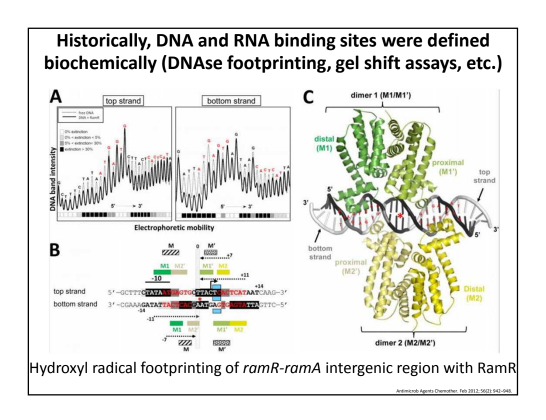
Motifs

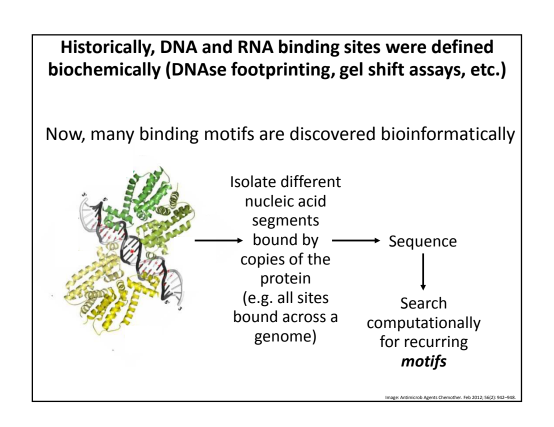
BCH364C/391L Systems Biology / Bioinformatics – Spring 2015

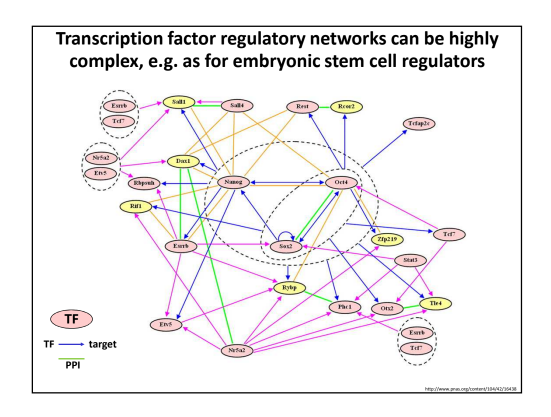
Edward Marcotte, Univ of Texas at Austin

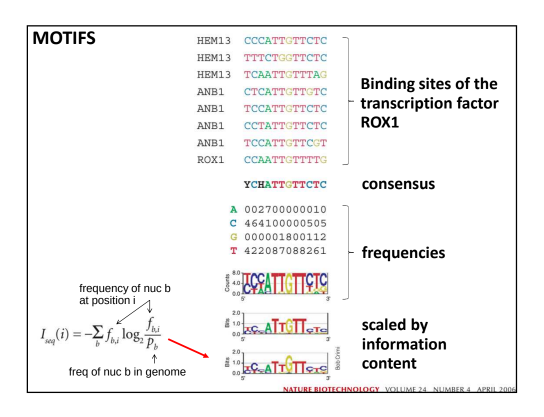
Edward Marcotte/Univ. of Texas/BCH364C-391L/Spring 201











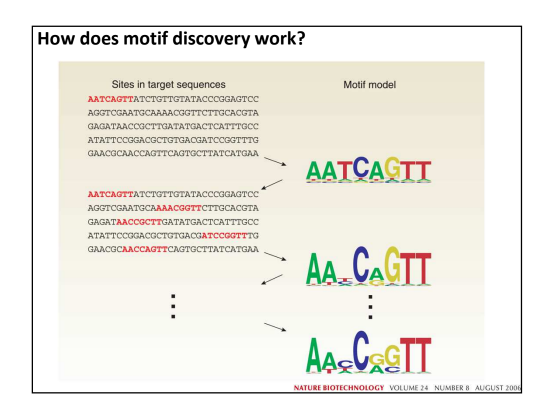
So, here's the challenge:

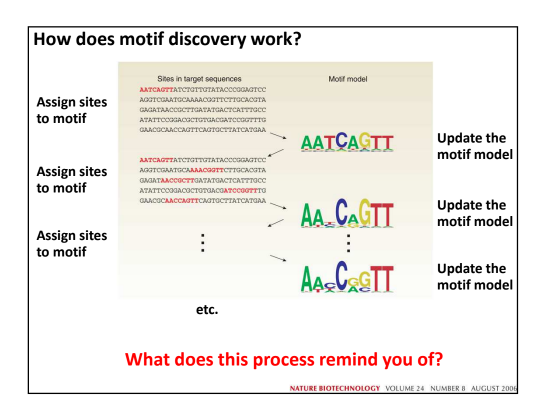
Given a set of DNA sequences that contain a motif (e.g., promoters of co-expressed genes), how do we discover it computationally?

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?

Motif finding often uses <u>expectation-maximization</u> (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.

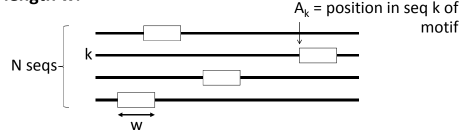
Searches the space of possible motifs for optimal solutions without testing everything.

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)

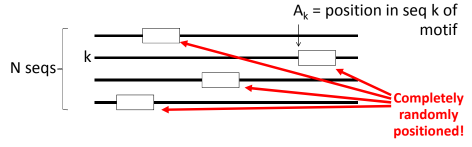
p_i = background probability of finding nucleotide (or aa) j

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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:



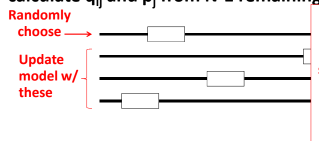
q_{ij} = probability of finding nucleotide (or aa) j at position i in motif i ranges from 1 to w j ranges across the nucleotides (or aa)

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<u>Predictive update step</u>: Randomly choose one sequence, calculate q_{ij} and p_i from N-1 remaining sequences



background frequency of count of symbol j at position i $q_{i,j} = \frac{c_{i,j} + b_j}{N-1+B}$

q_{ij} = probability of finding nucleotide (or as i ranges from 1 to w j ranges across the nucleotides (or a

p_j is calculated similarly from the counts <u>outside</u> the motifs

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

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