

Inferring Models of cis-Regulatory Modules using Information Theory

BMI/CS 776

www.biostat.wisc.edu/bmi776/

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Overview

- Biological question
 - What is causing differential gene expression?
- Goal
 - Find regulatory motifs in the DNA sequence
- Solution
 - FIRE (Finding Informative Regulatory Elements)

Goals for Lecture

Key concepts:

- Entropy
- Mutual information (MI)
- Motif logos
- Using MI to identify cis-regulatory module elements

A Common Type of Question

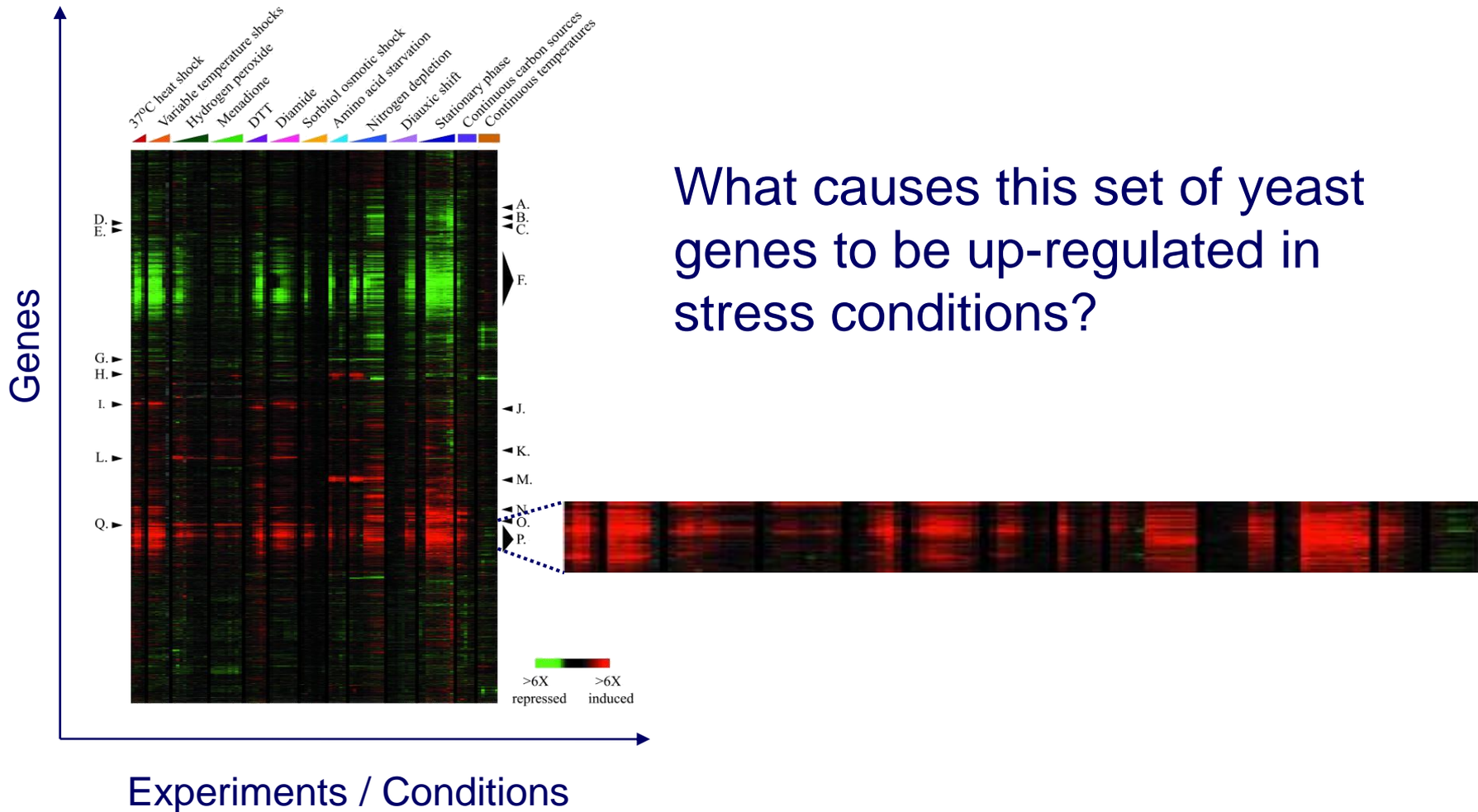
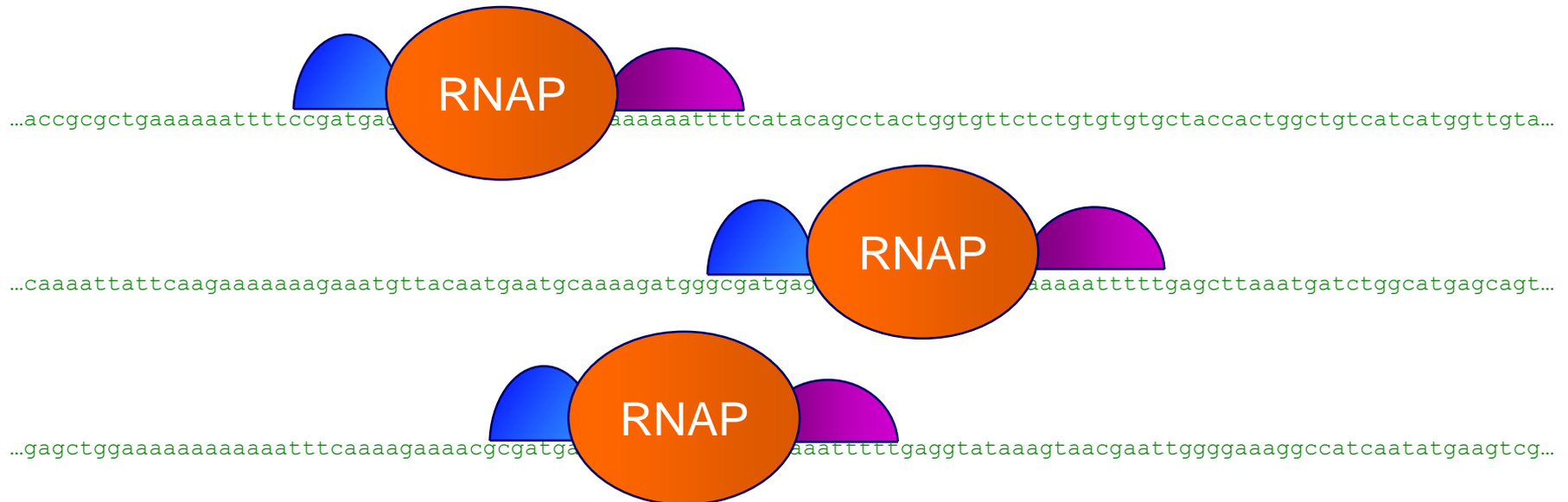


Figure from Gasch *et al.*, *Mol. Biol. Cell*, 2000

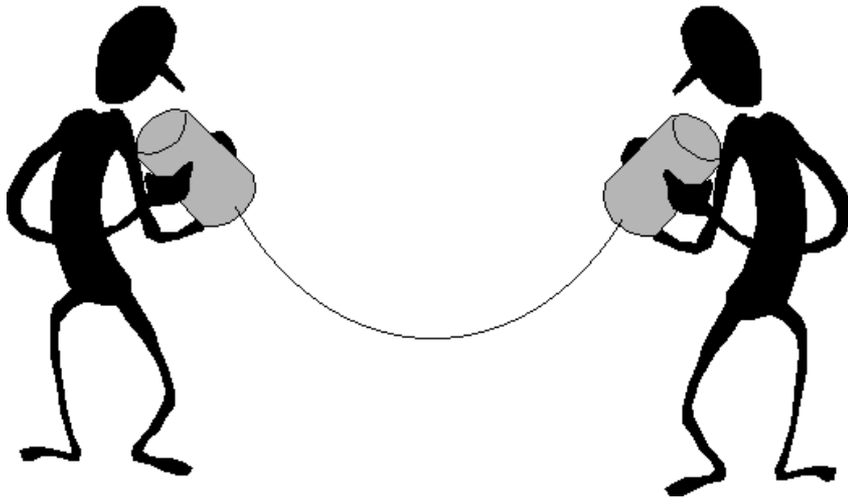
cis-Regulatory Modules (CRMs)

- Co-expressed genes are often controlled by specific configurations of binding sites



Information Theory Background

- Problem
 - Create a code to communicate information
- Example
 - Need to communicate the manufacturer of each bike



Information Theory Background

- Four types of bikes
- Possible code

Type	code
Trek	11
Specialized	10
Cervelo	01
Serotta	00

- Expected number of bits we have to communicate:
2 bits/bike

Information Theory Background

- Can we do better?
- Yes, if the bike types aren't equiprobable

Type, probability	# bits	code
$P(\text{Trek}) = 0.5$	1	1
$P(\text{Specialized}) = 0.25$	2	01
$P(\text{Cervelo}) = 0.125$	3	001
$P(\text{Serotta}) = 0.125$	3	000

- Optimal code uses $-\log_2 P(c)$ bits for event with probability $P(c)$

Information Theory Background

Type, probability	# bits	code
$P(\text{Trek}) = 0.5$	1	1
$P(\text{Specialized}) = 0.25$	2	01
$P(\text{Cervelo}) = 0.125$	3	001
$P(\text{Serotta}) = 0.125$	3	000

- Expected number of bits we have to communicate:
1.75 bits/bike

$$-\sum_{c=1}^{|C|} P(c) \log_2 P(c)$$

Entropy

- Entropy is a measure of uncertainty associated with a random variable
- Can be interpreted as the expected number of bits required to communicate the value of the variable

$$H(C) = -\sum_{c=1}^{|C|} P(c) \log_2 P(c)$$

entropy function for binary variable

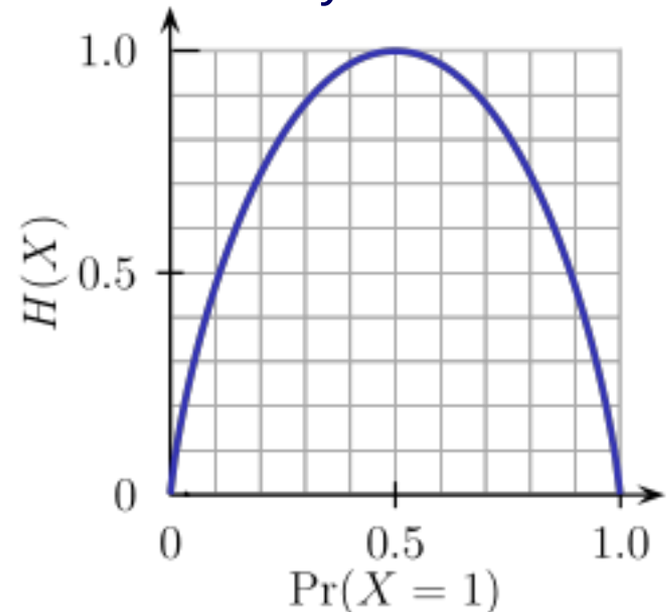
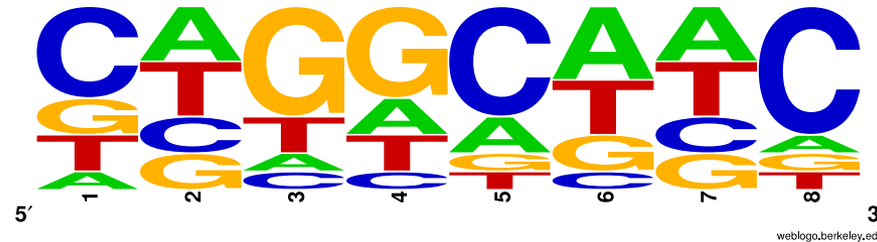


Image from Wikipedia

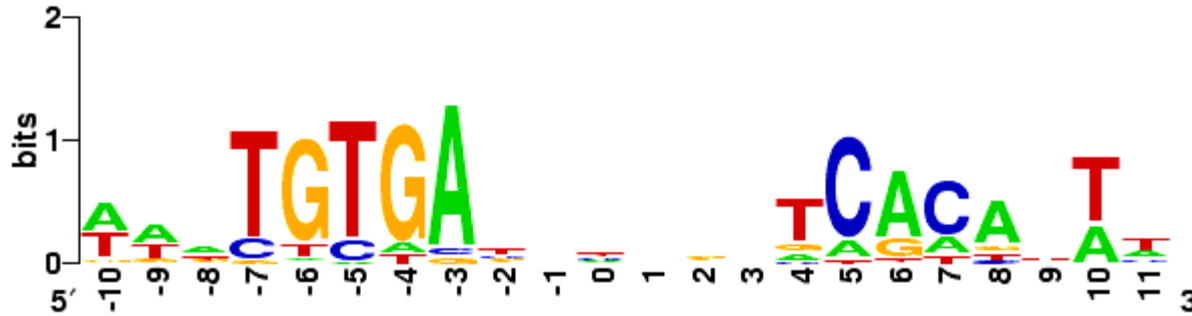
How is entropy related to
DNA sequences?

Sequence Logos



- Typically represent a binding site
- Height of each character c is proportional to $P(c)$

Sequence Logos

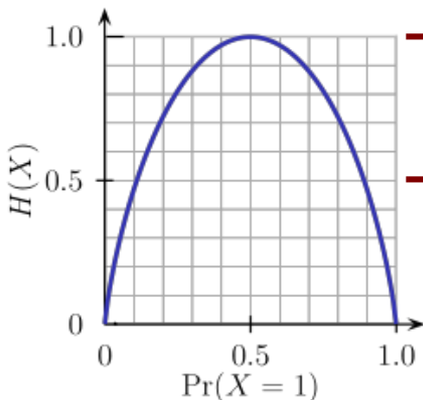


- Height of logo at a given position determined by decrease in entropy (from maximum possible)

$$H_{\max} - H(C) = \log_2 N - \left(- \sum_c P(c) \log_2 P(c) \right)$$

of characters in alphabet

decrease in entropy



FIRE

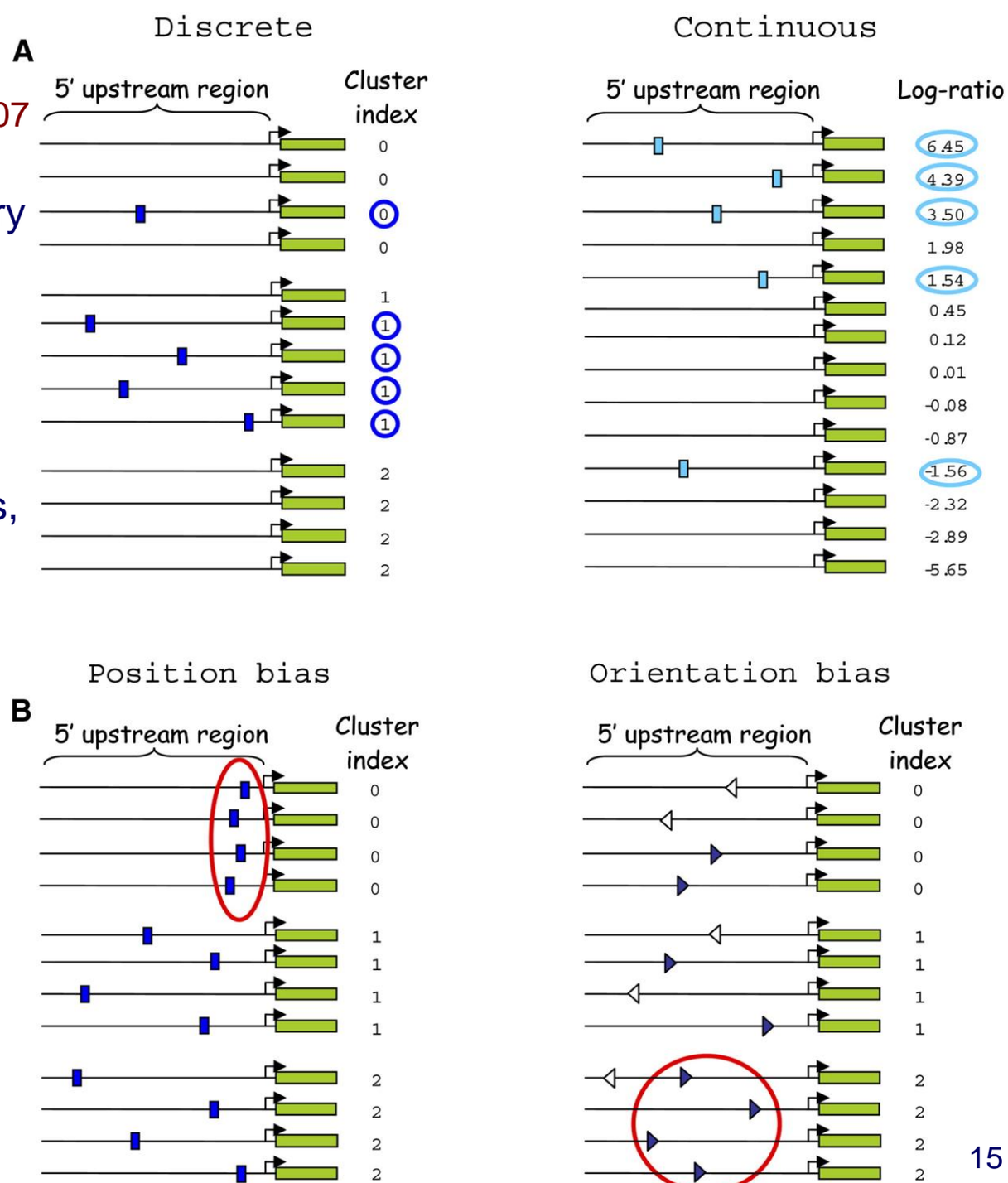
Elemento et al., *Molecular Cell* 2007

- Finding Informative Regulatory Elements (FIRE)

- Given a set of sequences grouped into clusters

- Find motifs, and relationships, that have high *mutual information* with the clusters

- Applicable when sequences have continuous values instead of cluster labels



Mutual Information in FIRE

- We can compute the mutual information between a motif and the clusters as follows

$$I(M; C) = \sum_{m=0}^1 \sum_{c=1}^{|C|} P(m, c) \log_2 \frac{P(m, c)}{P(m)P(c)}$$

$m=0, 1$ represent absence/presence of motif

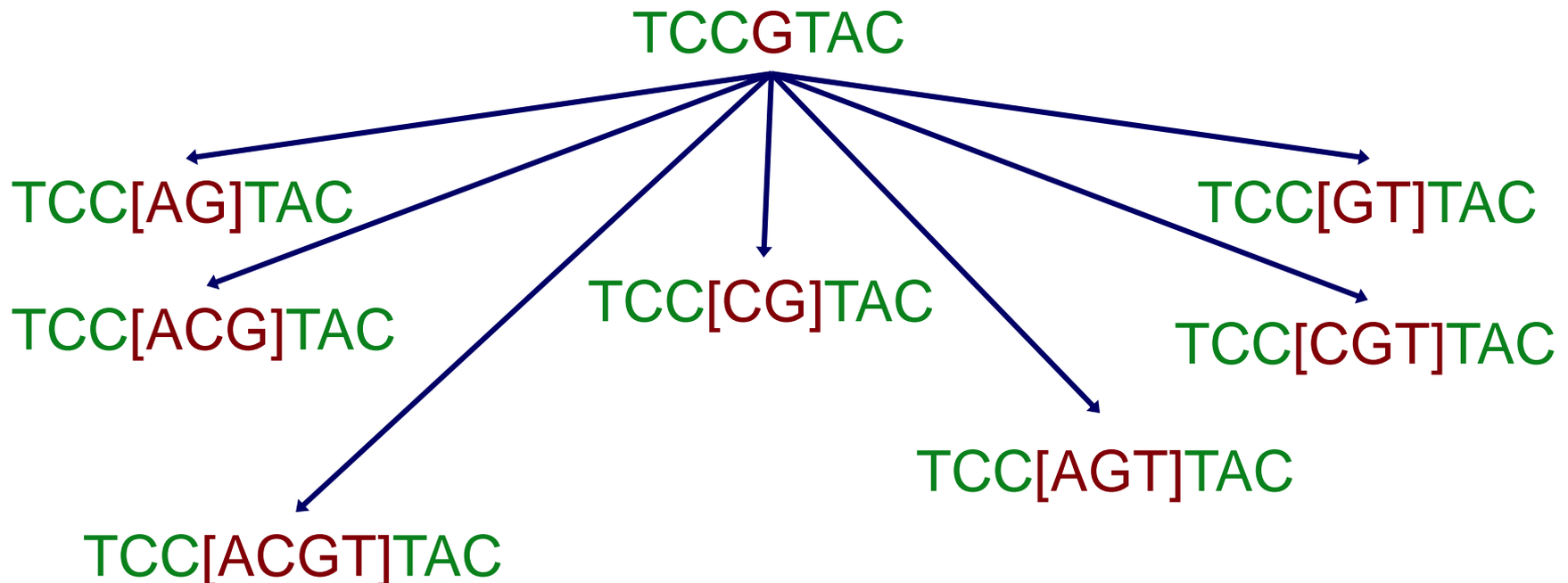
c ranges over the cluster labels

Finding Motifs in FIRE

- Motifs are represented by regular expressions; initially each motif is represented by a strict k -mer (e.g. **TCCGTAC**)
1. Test all k -mers ($k=7$ by default) to see which have significant mutual information with the cluster label
 2. Filter k -mers using a significance test to obtain motif seeds
 3. Generalize each motif seed
 4. Filter motifs using a significance test

Key Step in Generalizing a Motif in FIRE

- Randomly pick a position in the motif
- Generalize in all ways consistent with current value at position
- Score each by computing mutual information
- Retain the best generalization



Generalizing a Motif in FIRE

given: k -mer, n

$best \leftarrow \text{null}$

repeat n times

$motif \leftarrow k\text{-mer}$

 repeat

$motif \leftarrow \text{GeneralizePosition}(motif)$ // shown on previous slide

 until convergence (no improvement at any position)

 if $\text{score}(motif) > \text{score}(best)$

$best \leftarrow motif$

return: $best$

Generalizing a Motif in FIRE: Example

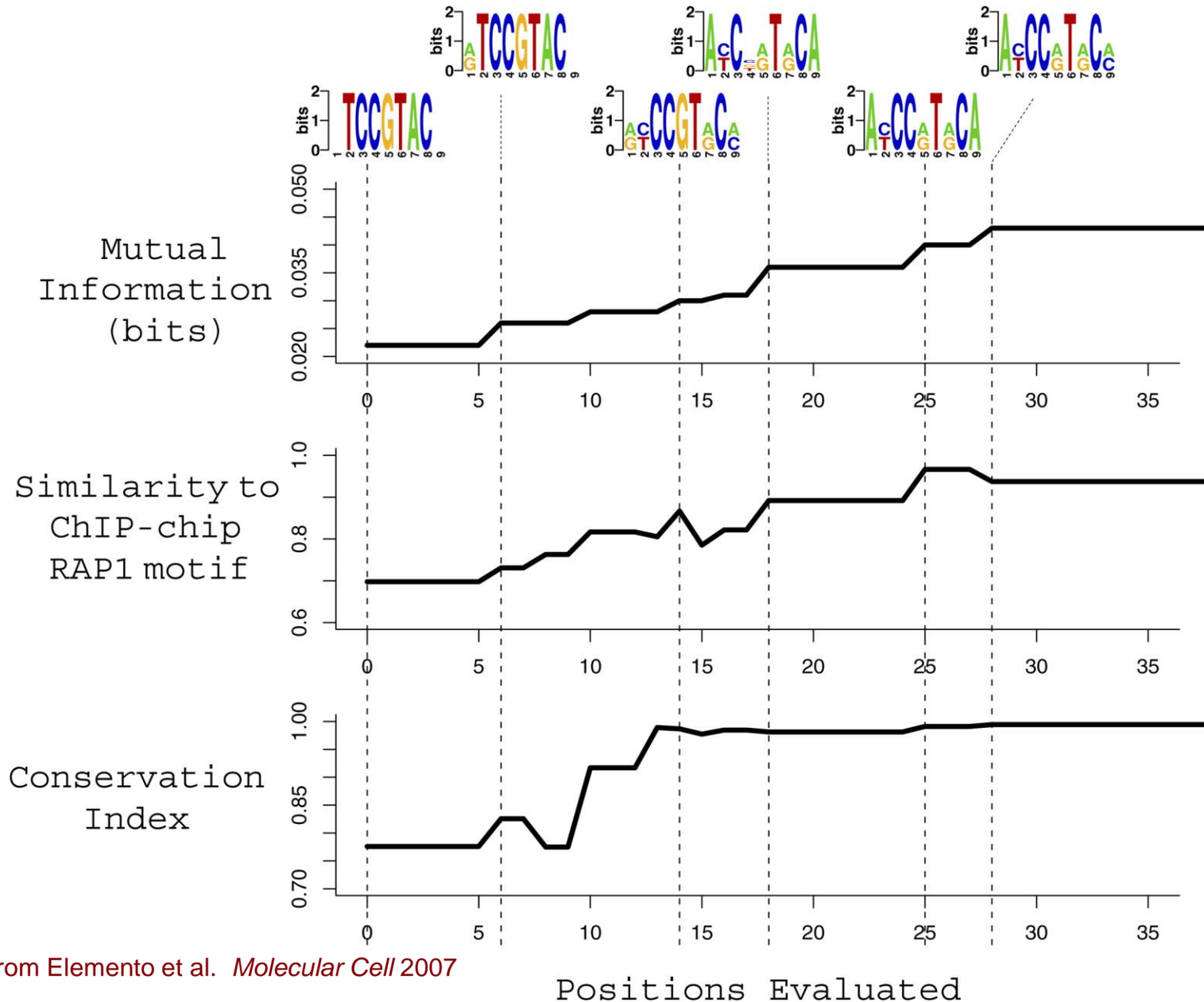


Figure from Elemento et al. *Molecular Cell* 2007

Avoiding Redundant Motifs

- Different seeds could converge to similar motifs

TCCGTAC
↓
TCC[CG]TAC

TCCCTAC
↓
TCC[CG]TAC

- Use mutual information to test whether new motif is unique and contributes new information

$$\frac{I(M; C | M')}{I(M; M')} > r$$

M' previous motif

M new candidate motif

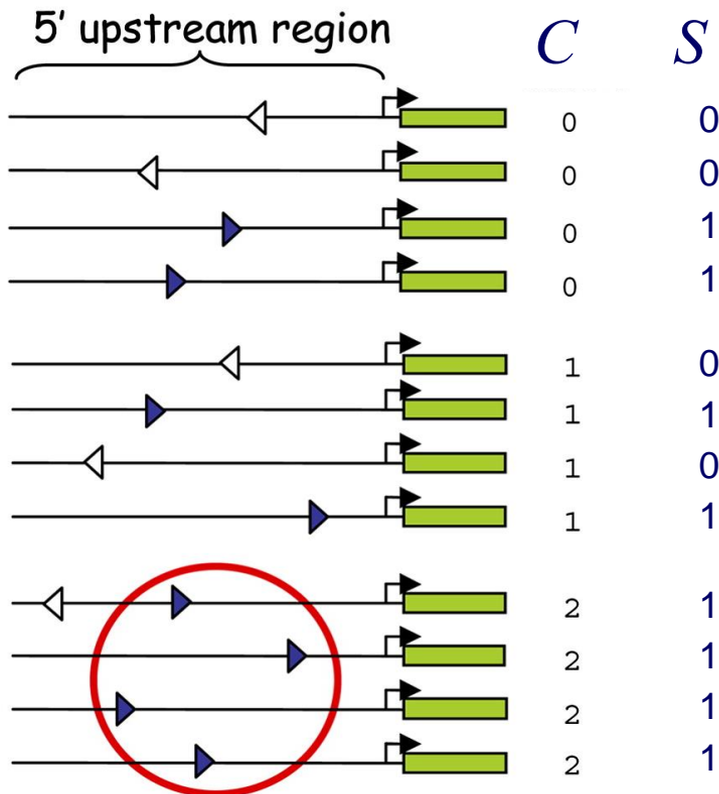
C expression clusters

Characterizing Predicted Motifs in FIRE

- Mutual information is also used to assess various properties of found motifs
 - orientation bias
 - position bias
 - interaction with another motif

Using MI to Determine Orientation Bias

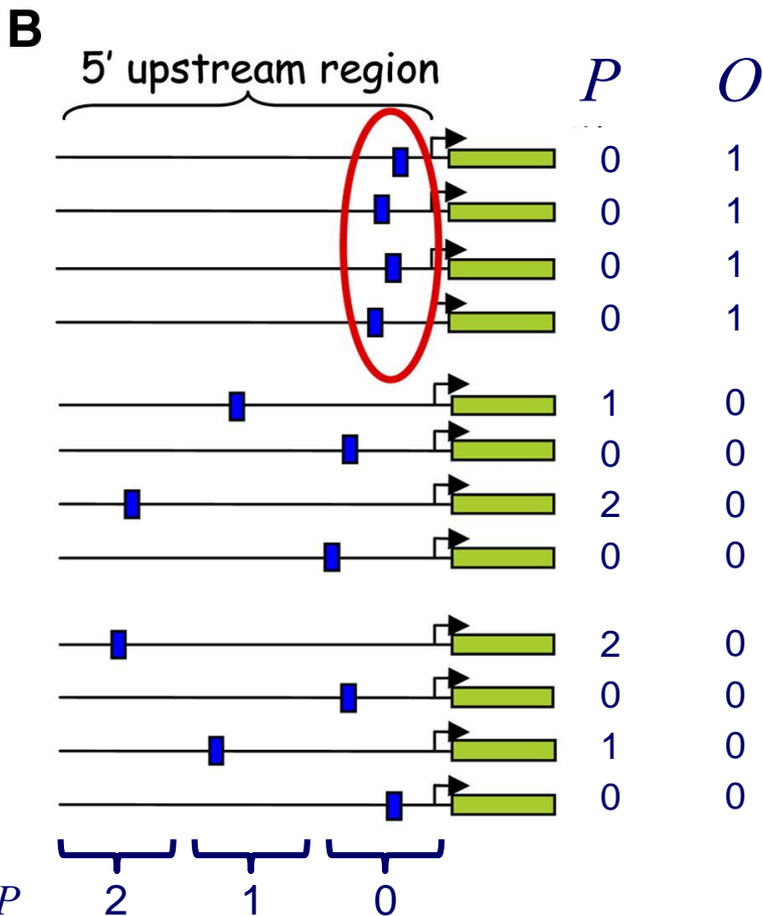
$I(S; C)$ C indicates cluster
 $S=1$ indicates motif present on transcribed strand
 $S=0$ otherwise (not present or not on transcribed strand)



Also compute MI where $S=1$ indicates motif present on complementary strand

Using MI to Determine Position Bias

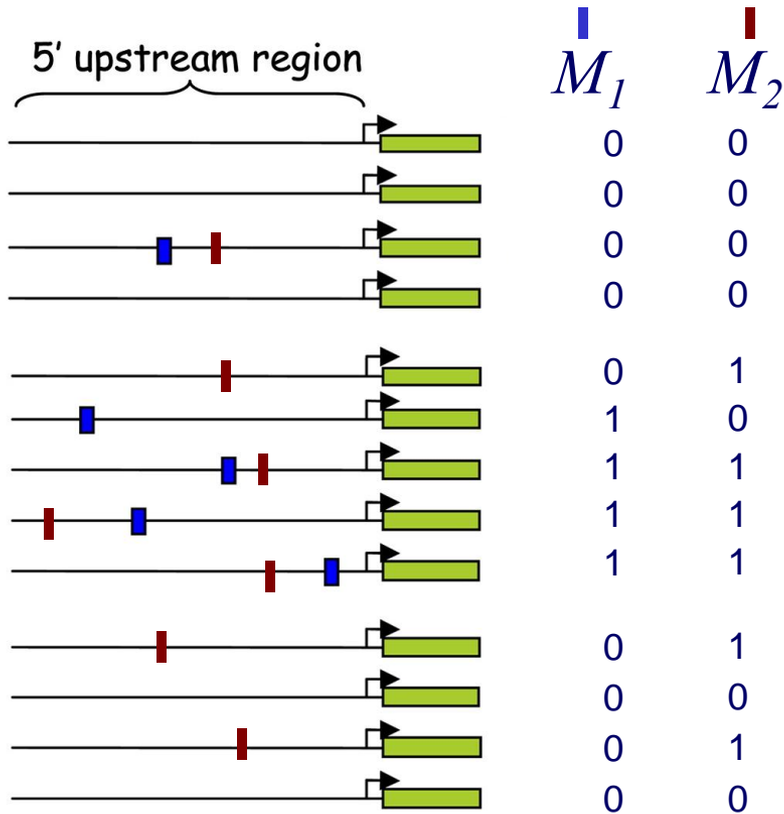
$I(P; O)$ P ranges over position bins
 $O=0, 1$ indicates whether or not the motif is over-represented in a sequence's cluster



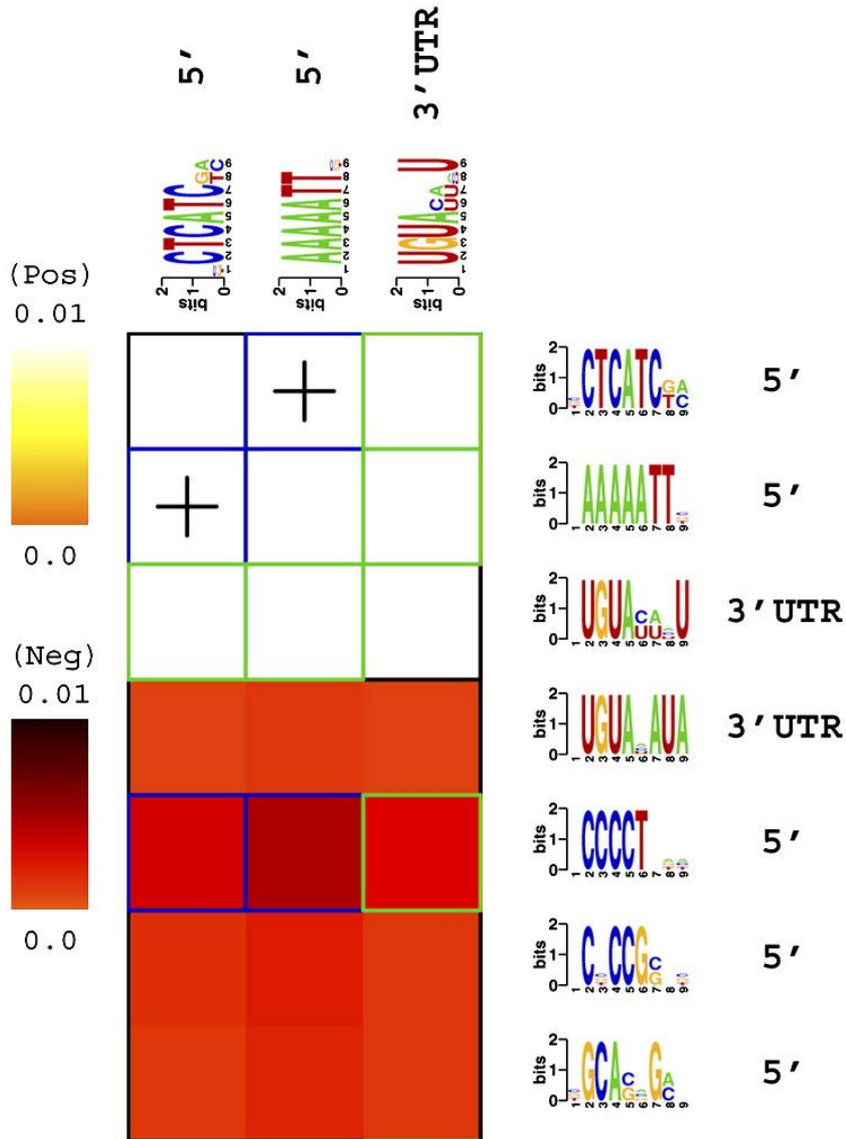
Only sequences containing the motif are considered for this calculation

Using MI to Determine Motif Interactions

$I(M_1; M_2)$ $M_i=0, 1$ indicates whether or not a sequence has the motif **and** is in a cluster for which the motif is over-represented; similarly for M_2



Using MI to Determine Motif Interactions



Yeast motif-motif interactions

White: positive association

Dark red: negative association

Blue box: DNA-DNA

Green box: DNA-RNA

Plus: spatial co-localization

Discussion of FIRE

- FIRE
 - mutual information used to identify motifs and relationships among them
 - motif search is based on generalizing informative k -mers
- Consider advantages and disadvantages of k -mers versus PWMs
- In contrast to many motif-finding approaches, FIRE takes advantage of *negative* sequences
- FIRE returns all informative motifs found
- Mutual information and conditional mutual information can also be useful for reconstructing biological networks
 - e.g., build gene-gene network where edges indicate high MI in genes' expression levels