

# **Motif analysis**

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SciLifeLab (Long-term Bioinformatics Support)

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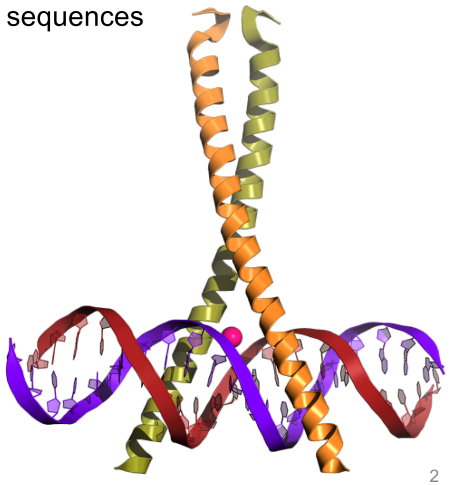
### The problem



From a transcription factor (TF) ChIP-seq experiment, find the DNA sequences recognized by the TF.

In this context: Motif = a set of nucleotide sequences

Typically 4-20 bp



### This talk



- What is a motif? How is it represented?
- De-novo motif discovery: What the problem is, principles behind the programs
- Examples of motif discovery programs
- Practical considerations: data size, how to handle repeats etc.

# How DNA sequence motifs be represented SciLifeLab

- 1. As a *sequence* of nucleotides, e.g. CTGGA
- 2. As a *regular expression*, taking into account ambiguity e.g. [C or G][C or T]GG[G or A]
- 3. As a *matrix*, based on nucleotide frequency in each position

Pos	1	2	3	4	5
A	0	~	0	0	5
С	5	4	0	0	0
G	4	0	10	10	4
Т	1	5	0	0	1

4. More complicated representations, taking dependencies between positions into account (HMMs, dinucleotide matrices, deep learning networks etc.)

## Position weight matrices



- A position weight matrix (PWM) is based on nucleotide frequencies in a set of aligned sequences.
- The frequencies are converted to probabilities, and then to loglikelihoods given a background model.

Pos	1	2	3	4	5
A	0	1	0	0	5
С	5	4	0	0	0
G	4	0	10	10	4
Т	1	5	0	0	1

Pos	1	2	3	4	5
A	0.0	0.1	0.0	0.0	0.5
С	0.5	0.4	0.0	0.0	0.0
G	0.4	0.0	1.0	1.0	0.4
Т	0.1	0.5	0.0	0.0	0.1

Pos	1	2	3	4	5
Α	-Inf	-1.32	-Inf	-Inf	1.0
С	1.0	0.68	-Inf	-Inf	-Inf
G	0.68	-Inf	2.0	2.0	0.68
Т	-1.32	1.0	-Inf	-Inf	-1.32

Position *frequency* matrix Position *probability* matrix

count nucleotides in each position

divide by total nr of sequences

divide by background freq. and log-transform  $-\log(m_{n,p}/b_n)$ 

Position weight matrix

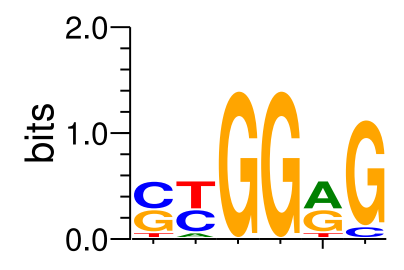
We might need to add a pseudo count to the frequency matrix, to avoid -Inf.

## Sequence logos



- Sequence logos are used to visualize PWMs.
- Nucleotide frequency and information content for each position can be represented.

Pos	1	2	3	4	5
Α	0	1	0	0	0
С	4	4	0	0	5
G	5	5	10	10	4
Т	1	0	0	0	1



Height: 2 – entropy = 
$$2 - \sum_{i=1}^n \mathrm{P}(x_i) \log_b \mathrm{P}(x_i)$$

### **Databases with TF binding site motifs**



- JASPAR (<a href="http://jaspar.genereg.net">http://jaspar.genereg.net</a>). Good, curated, free, data base with around 1500 motifs from all kinds of species.
- Transfac (<a href="http://genexplain.com/transfac/">http://gene-</a>
  regulation.com/pub/databases.html
  Good, curated, not free, data base with around 2800 motifs from all kinds of species.
  - Older version is free for academic use.
- Other databases
  - ChIPBase <a href="http://rna.sysu.edu.cn/chipbase/">http://rna.sysu.edu.cn/chipbase/</a>
  - HOCOMOCO (human only) <a href="http://hocomoco11.autosome.ru">http://hocomoco11.autosome.ru</a>
  - footprintDB (combining several databases)
    <a href="http://floresta.eead.csic.es/footprintdb/index.php">http://floresta.eead.csic.es/footprintdb/index.php</a>

### Scanning the genome with a PWM



 Every sequence can be scored on how well it matches the PWM, by adding up the scores for each position:

Pos	1	2	3	4	5
A	-Inf	-1.32	-Inf	-Inf	1.0
С	1.0	0.68	-Inf	-Inf	-Inf
G	0.68	-Inf	2.0	2.0	0.68
Т	-1.32	1.0	-Inf	-Inf	-1.32

GAGGG 
$$\rightarrow$$
 0.68 -1.32 + 2.0 +2.0 + 0.68 = 4.04  
CTGGG  $\rightarrow$  1.0 + 1.0 + 2.0 + 2.0 + 1.0 = 7  
CTGAG  $\rightarrow$  1.0 + 1.0 - Inf + 2.0 + 1.0 = - Inf

- The score represents the log likelihood of the sequence being a motif compared to bg
- High scores → likely strong TF binding → long time spent on DNA by TF
- Useful to have a cutoff on what we consider is a match. Setting cutoff can be tricky!

### Limitations of position weight matrices



- In 90% of tested cases, matrix based models perform as well as more complex models (Weirauch et al. Nature Biotech. 2013).
- But PWMs can be inaccurate if there is
  - Dependencies between nucleotides
  - Variable spacing between sequences

### De-novo motif finding



- Given a set of transcription factor binding sites (e.g. from ChIP-seq), are any motifs enriched?
- Some kind of background model is needed
  - A set of background sequences
    - Regions nearby the peaks (e.g. 2 Kbp away), with similar GC content
  - Nucleotide (or dinucleotide) frequencies
  - A bad background model will give strange and misleading results!



# **Motif finding methods**



- We need methods to search the space of possible motifs
- We also need a way to score motif candidates (e.g. enrichment, complexity)
- Optimal results are not guaranteed.

### **MEME**



#### Method:

- Starts with a guess, M, of what the motif might be. It then produces estimates, L, of where motif is located.
- Given L, the motif M is updated. Then L is updated with a new motif and so on, until the motif M doesn't change much.
- When the motif search has converged, the resulting motif is scored (based on enrichment and information content).
- To find more motifs, all occurrences of the motif are then removed from the input sequences, and the algortim is the re-run with a new start guess.

#### Output

- A set of PWMs, with scores and p-values
- Pros: Old, widely used method. Often works well.
- Cons: Slow, has trouble handling large inputs (>500 peaks)



### **DREME**



- Method:
  - Look at all 3-8mers to find the most enriched sequences (Fisher test)
  - Iteratively, try to make these more general with search
    - CTGGG
    - $\rightarrow$  CTGG[G or A]
    - → C[C or T]GG[G or A]
    - → [C or G][C or T]GG[G or A]
  - Convert this to PWM
- Output: PWMs, with p-values
- Pros: Very fast, good performance
- Cons: Restricted to short sequences (up to 8 bp). Does not take nucleotide frequency into account.



### Homer



#### Method

- Looks at all 8,10 and 12-mers to find the most enriched.
- The most enriched sequences are then converted to weight matrices are refined.

#### Output

- A set of PWMs, with info on e-values and which known motifit's similar to.
- If any known motifs are enriched in the given regions.

#### Pros

- Nice output, includes matching to known motifs
- Quite fast
- Usually works well

#### Cons

- The documentation is not good
- It's a bit hard to install, need to install genomes too.



### **Practical considerations**



- Less information content → harder problem
  - Short motifs are harder to find
  - Degenerate motifs are harder to find
- Which peaks to use?
  - Some methods will have problems handling tens of thousands of peaks.
  - Also, many weak peaks don't provide useful information
  - → often only the top 500 etc. peaks are used.
- Repeats (e.g. low complexity repeats) can throw the motif finding methods off. → Work on repeat masked sequences!

### How well do these methods work?

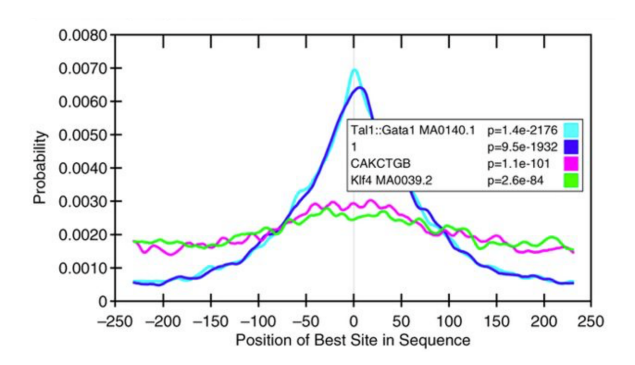


- There is no good benchmarking study on motif finding in ChIP-seq data, but usually finding the main motif is not that difficult
  - ChIP-seq gives short regions to look in
  - The top ChIP-seq peaks are typically very enriched for the motif of interest.
- There might also be co-factor motifs. These are harder to find.
- Compare this to analysis of promoters of co-regulated genes:
  - We have very long promoters to search for motifs
  - We have don't have as clear enrichment of the motifs.

# **Further analysis**



- PhyloGibbs incorporating sequence conservation in the motif finding.
- Ensemble methods combining the results from several motif finding programs
- TomTom Comparison of a new motif to a database of known motifs
- Centrimo Motif location.



### **Exercise**



- Takes sets of peaks from ENCODE
  - ChIP-seq against CTCF (human and mouse data sets)
  - ChIP-seq against REST, from previous lab
- Try a few different motif finders
  - DREME
  - MEME
  - HOMER
- Try a motif comparison tool, TomTom