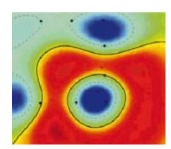
# mGene - a sophisticated Bioinformatics Application

(Transcriptionsstart, Splice Sites, PolyA Site Prediction) Structure Learning





Lecture by Klaus-Robert Müller, TUB 2015

### Contribution

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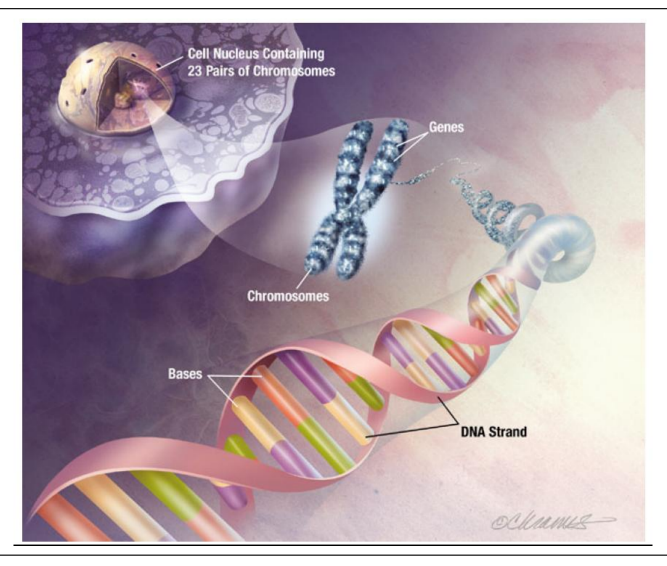
### **Bioinformatic- Benefits**

- Molecular medicine
  - More drug targets
  - Personalised medicine
  - Preventative medicine
  - Gene therapy
- Microbial genome applications
  - Waste cleanup
  - Climate change
  - Alternative energy sources
  - Biotechnology
  - Antibiotic resistance
  - Forensic analysis of microbes
  - Evolutionary studies
- Agriculture
  - Insect resistance
  - Improve nutritional quality
  - Grow crops in poorer soils and that are drought resistant

### **Bioinformatics - Applications**

- In Cell
  - which genes are on / off ?
  - in which tissue?
  - under which conditions ?
- Sequence Analysis on DNA/RNA ← in this Lecture
  - locate sequences (genes, start, stop, splice sites,...)
  - detect properties
  - how do individuals of same species differ (SNP's)
  - conservation
  - functional elements
- on Proteins
  - determine structure
  - determine function
  - find protein of similar functions
  - find binding sites (protein-protein, protein-dna)

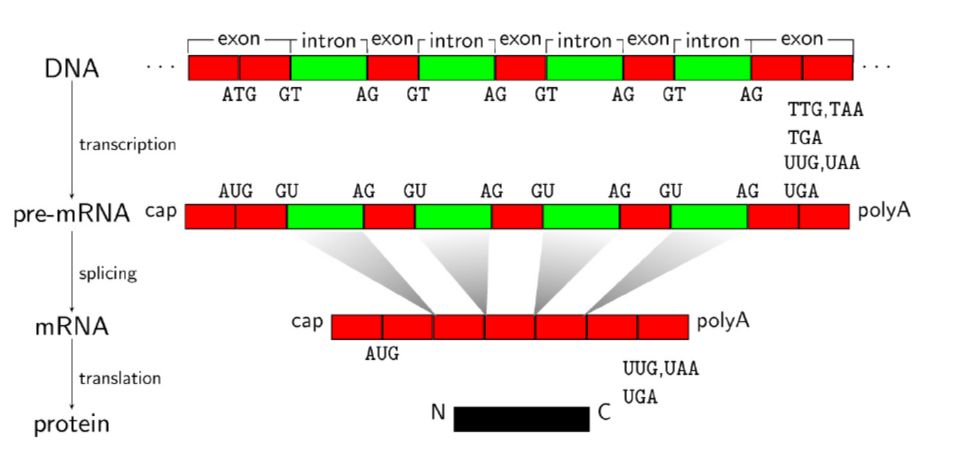
# **Bioinformatics – The Genome**







### Bioinformatics – from DNA to Protein

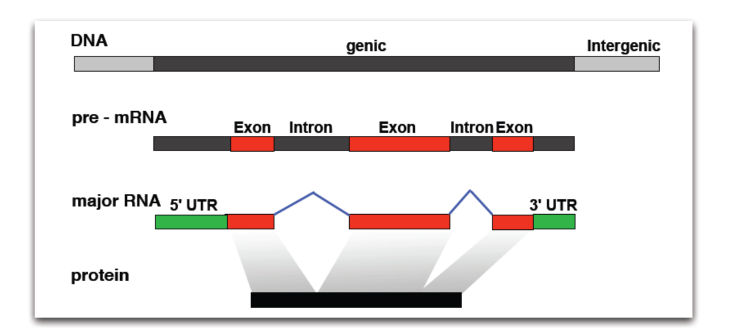






### **Finding Genes I**

- What is a Gene?
  - A segment on DNA that codes for a certain property (protein).
  - Proteins control everything, Enzymes (catalyze; involved in metabolism, DNA replication/repair, RNA synthesis)..., Cell signaling (Insulin), ligand binding (Haemoglobin),...

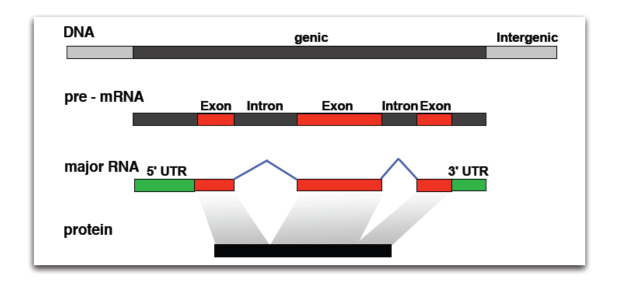






# **Finding Genes II**

- Sites to detect
  - Gene has a transcription start, transcription end only part from ATG...TAA,... is transcribed ⇒ pre-mRNA
  - Only **exons** code for protein, inserted **introns** are cut out in splicing  $\Rightarrow$  mRNA
  - Gene has a translation start and translation end that part is translated to ⇒ Protein







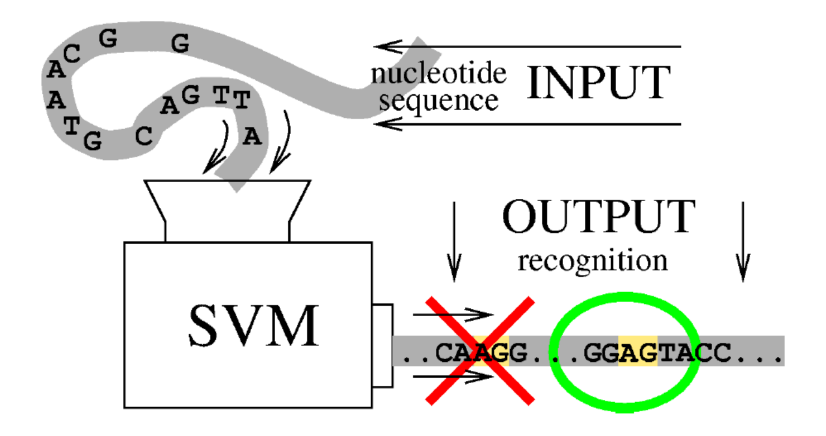
# **Finding Genes III**

- Requirements
  - human genome (all DNA) has 3 billion base pairs huge!!
  - method needs to be fast + fit in memory
- 2-step approach:
  - ① Detect Signals (focus on splice site and transcription start site prediction) ⇒ SVM on sliding windows
    - define kernels on strings
    - (spectrum kernel, weighted degree kernel)
  - 2 Learn Structure/Gene Segmentation (complex task)





# **Splice site detection**







# **Preparing data**

- Collecting data for training and evaluation is a complex, non-trivial task (half the work)
- two kinds, one for 1st pass (2-class classification positive/negative data); one for 2nd pass (correct segmentations)
- we assume data is given (others have done it for us :-)

# 2-class problem: solve with SVMs Classifier

$$f(\mathbf{x}) = \operatorname{sign}\left(\sum_{i=1}^{N} y_i \alpha_i \mathbf{k}(\mathbf{x}, \mathbf{x}_i) + b\right)$$

⇒ How to design the kernel?

### **Data classes**

- Position Independent (e.g. Which Tissue? Promoter Region)
   AAACAAAACGTAACTAATCTTTTAGAGAGAACGTTTCAACCATTTTGAGAGATTAACTCATCACAGATTTCAATACACAGATATAATTCAAAAATT
   CACTCCCCAAATCAACGATATTTAAAAATCACTAACACATCCGTCTGTGC
  - Task: separate DNA strings, '-' class random ACGT, '+' class contains 'AAAAA' motif
- - Task: separate DNA strings, '-' class random ACGT, '+' class 'AA' in the middle
- - ullet Task: separate DNA strings, '-' class random 'ACGT', '+' class 'AAA' in the middle shifted  $\pm 15$





# **Spectrum Kernel**

### To make use of position independent motifs:

- Idea: like bag of words kernel (text classification) but for Bioinformatics (words are now strings of length k (k-mers))
  - count k-mers in sequence A and sequence B.
  - Spectrum Kernel is sum of product of counts (for same k-mer)

### Example k = 3:

 $oldsymbol{x}$  AAACAAATAAGTAACTAATCTTTTAG $oldsymbol{\mathsf{G}}$ AAGAACGTTTCAACCATTTTGAG

x' TACCTAATTATGAAATTAAATTTCAGTGTGCTGATGGAAACGGAGAAGTC

3-mer	AAA	AAC	 CCA	CCC	 TTT
# in <b>x</b>	2	4	 1	0	 3
# in <b>x</b> '	3	1	 0	0	 1

$$k(x, x') = 2 \cdot 3 + 4 \cdot 1 + \dots + 1 \cdot 0 + 0 \cdot 0 \dots + 3 \cdot 1$$





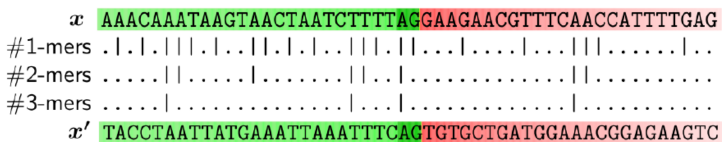
# Weighted degree kernel

### To make use of position dependent motifs:

$$\mathbf{k}(\mathbf{x},\mathbf{x}') = \sum_{k=1}^{d} \beta_k \sum_{l=1}^{L-k} \mathbf{I}(\mathbf{u}_{k,l}(\mathbf{x}) = \mathbf{u}_{k,l}(\mathbf{x}'))$$

- L length of the sequence x
- d maximal "match length" taken into account
- $\mathbf{u}_{k,l}(\mathbf{x})$  subsequence of length k at position l of sequence  $\mathbf{x}$

### Example degree d = 3:



$$k(\mathbf{x}, \mathbf{x}') = \beta_1 \cdot 21 + \beta_2 \cdot 8 + \beta_3 \cdot 4$$





# Weighted degree kernel

- for weighting we use  $\beta_k = 2 \frac{d-k+1}{d(d+1)}$ .
- effort is  $O(L \cdot d)$
- Speedup Idea: Reduce effort to O(L) by finding matching "blocks"

**Exercise:** Show that WD kernel and its "block" formulation are equivalent





### Weighted degree kernel with shift

### To make use of partially position-dependent motifs:

- If sequence is slightly mutated (Insertion, Deletion) WD kernel fails.
- Extension: Allow for some positional variance (shifts S(I))

$$\mathbf{k}(\mathbf{x}_{i}, \mathbf{x}_{j}) = \sum_{k=1}^{d} \beta_{k}^{L-k+1} \sum_{l=1}^{S(l)} \delta_{s} \ \mu_{k,l,s,\mathbf{x}_{i},\mathbf{x}_{j}},$$

$$\mu_{k,l,s,\mathbf{x}_{i},\mathbf{x}_{i}} = \mathbf{I}(\mathbf{u}_{k,l+s}(\mathbf{x}_{i}) = \mathbf{u}_{k,l}(\mathbf{x}_{j})) + \mathbf{I}(\mathbf{u}_{k,l}(\mathbf{x}_{i}) = \mathbf{u}_{k,l+s}(\mathbf{x}_{j})),$$

$$k(x_1,x_2) = w_{6,3} + w_{6,-3} + w_{3,4}$$
 $X_1 \longrightarrow CGAACG$ CTACGTATTATTTAGTCGGATTG  $\longrightarrow$ 





# Final signal and content sensors

- Exon vs. Intron Spectrum Kernel
- splice sites Weighted Degree Kernel
- transcription start, transcription stop Weighted Degree
   Kernel with shifts

### Perform Model Selection:

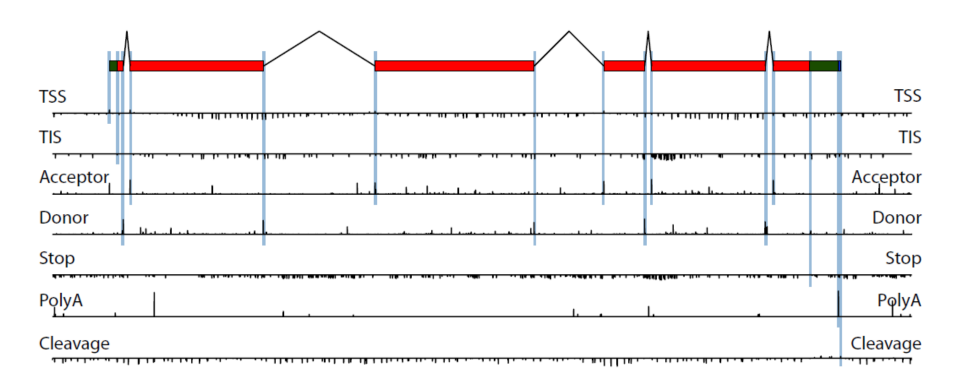
- window length
- k-mer length (spectrum kernel), degree, shift (WD-kernel)
- SVM regularization parameter C
- . . .
- takes a long time (cluster)

# We now have Signal and content sensors





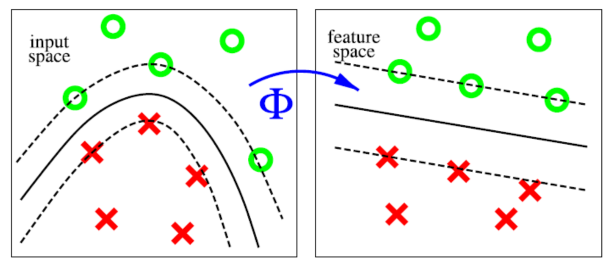
# **Example**







### What did we learn?



SVM decision function in kernel feature space:

$$f(\mathbf{x}) = \sum_{i=1}^{N} y_i \alpha_i \underbrace{\Phi(\mathbf{x}) \cdot \Phi(\mathbf{x}_i)}_{=k(\mathbf{x}, \mathbf{x}_i)} + b$$
(1)

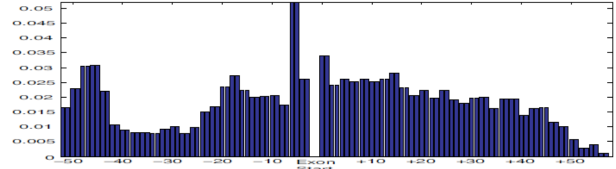
 learned parameters α by solving quadratic optimization problem

**Problem: Decision function** (2) is hard to interpret

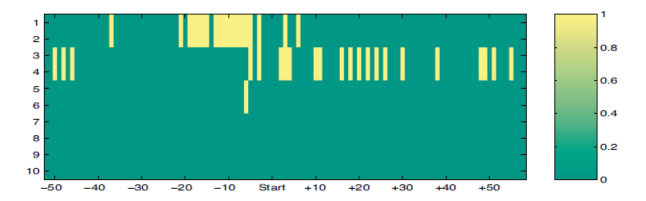
### **Understanding the SVM decision**

# Splice Sites

Which positions in the sequence are important for discrimination?



What characterizes those positions?



Which motifs at which position are important?

# **Optimize combinations of kernels**

Define Kernel as Convex Combination of Subkernels:

$$k(\mathbf{x}, \mathbf{y}) = \sum_{l=1}^{L} \beta_{l} k_{l}(\mathbf{x}, \mathbf{y})$$

e.g. Weighted Degree Kernel

$$k(\mathbf{x}, \mathbf{x}') = \sum_{l=1}^{L} \beta_l \sum_{k=1}^{d} \mathbf{I}(\mathbf{u}_{k,l}(\mathbf{x}) = \mathbf{u}_{k,l}(\mathbf{x}'))$$

- optimize weights \( \beta \) such that margin is maximized
  - $\Rightarrow$  determine  $(\beta, \alpha, b)$  simultaneously
  - ⇒ Multiple Kernel Learning (Bach, Lanckriet and Jordan 2004)





### Multiple kernel learning (MKL)

Possible solution We can add the two kernels, that is

$$k(\mathbf{x}, \mathbf{x}') := k_{sequence}(\mathbf{x}, \mathbf{x}') + k_{structure}(\mathbf{x}, \mathbf{x}').$$

Better solution We can mix the two kernels,

$$k(\mathbf{x}, \mathbf{x}') := (1 - t)k_{sequence}(\mathbf{x}, \mathbf{x}') + tk_{structure}(\mathbf{x}, \mathbf{x}'),$$

where t should be estimated from the training data.

In general: use the data to find best convex combination.

$$k(\mathbf{x}, \mathbf{x}') = \sum_{p=1}^{K} \beta_p k_p(\mathbf{x}, \mathbf{x}').$$

### **Applications**

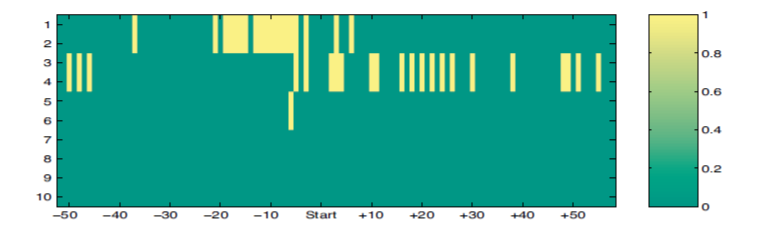
- Heterogeneous data
- Improving interpretability

# **Method of interpreting SVMs**

• Weighted Degree kernel: linear comb. of  $L \cdot D$  kernels

$$k(\mathbf{x}, \mathbf{x}') = \sum_{d=1}^{D} \sum_{l=1}^{L-d+1} \gamma_{l,d} \mathbf{I}(\mathbf{u}_{l,d}(\mathbf{x}) = \mathbf{u}_{l,d}(\mathbf{x}'))$$

Example: Classifying splice sites



See Rätsch & Sonnenburg 2006 for more details.

# Multiple Kernel Learning I

min 
$$\frac{1}{2} \left( \sum_{j=1}^{M} \beta_{j} \| \mathbf{w}_{j} \|_{2} \right)^{2} + C \sum_{i=1}^{N} \xi_{n}$$
w.r.t. 
$$\mathbf{w} = (\mathbf{w}_{1}, \dots, \mathbf{w}_{M}), \mathbf{w}_{j} \in \mathbf{R}^{k_{j}}, \boldsymbol{\xi} \in \mathbf{R}^{N}_{+}, \boldsymbol{\beta} \in \mathbf{R}^{M}_{+}, b \in \mathbf{R}$$
s.t. 
$$y_{i} \left( \sum_{j=1}^{M} \beta_{j} \mathbf{w}_{j}^{\top} \mathbf{x}_{i,j} + b \right) \geq 1 - \xi_{i}, \forall i = 1, \dots, N$$

$$\sum_{j=1}^{M} \beta_{j} = 1$$

**Properties:** equivalent to linear SVM for M = 1; solution sparse in "blocks"; each block j corresponds to one kernel





# **Multiple Kernel Learning II**

Dual Formulation (Bach, Lanckriet, Jordan 2004):

min 
$$\frac{1}{2}\gamma^{2} - \sum_{i=1}^{N} \alpha_{i}$$
w.r.t.  $\gamma \in \mathbf{R}, \alpha \in \mathbf{R}^{N}$ 
s.t.  $0 \le \alpha \le C, \sum_{i=1}^{N} \alpha_{i} y_{i} = 0$ 

$$\sum_{r=1}^{N} \sum_{s=1}^{N} \alpha_{r} \alpha_{s} y_{r} y_{s} K_{j}(\mathbf{x}_{r}, \mathbf{x}_{s}) - \gamma^{2} \le 0, \ \forall j = 1, \dots, M$$

$$=:S_{j}(\alpha)$$

"partial Lagrangian:"

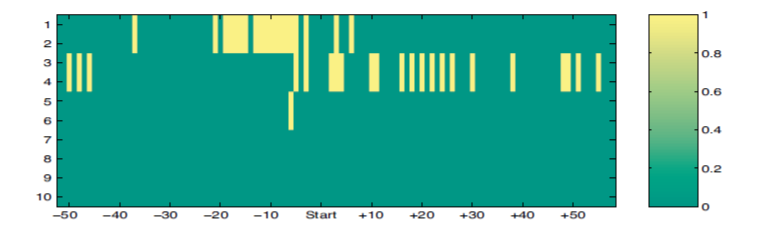
$$L := \frac{1}{2}\gamma^{2} - \sum_{i=1}^{N} \alpha_{i} + \sum_{j=1}^{M} \beta_{j}(S_{j}(\alpha) - \gamma^{2})$$

# **Method of interpreting SVMs**

• Weighted Degree kernel: linear comb. of  $L \cdot D$  kernels

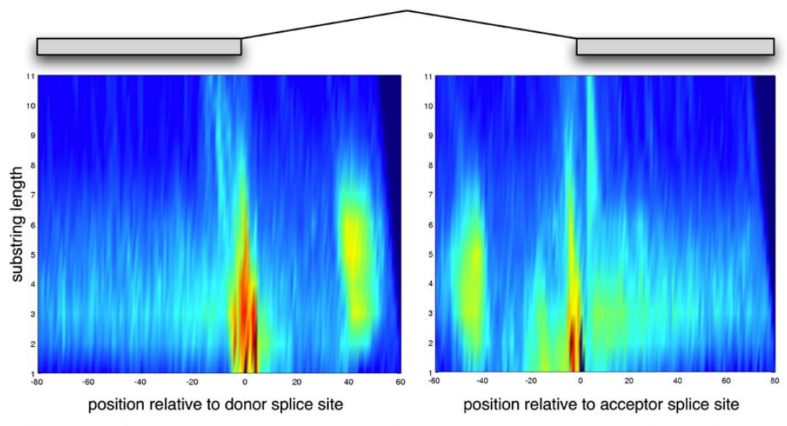
$$k(\mathbf{x}, \mathbf{x}') = \sum_{d=1}^{D} \sum_{l=1}^{L-d+1} \gamma_{l,d} \mathbf{I}(\mathbf{u}_{l,d}(\mathbf{x}) = \mathbf{u}_{l,d}(\mathbf{x}'))$$

Example: Classifying splice sites



See Rätsch & Sonnenburg 2006 for more details.

# **POIMs for splicing**



Color-coded importance scores of substrings near splice sites. Long substrings are important upstream of the donor and downstream of the acceptor site (Rätsch et.al 2007)





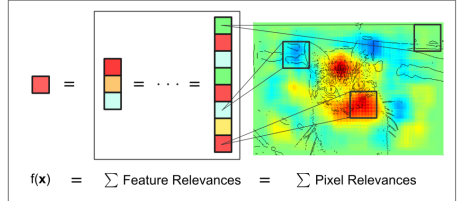
### Overview: Pixel-wise Explanation of the Classifier Decision



# 

Features

Pixel-wise Explanation



$$f(x) \approx \sum_{d=1}^{V} R_d$$

$$f(x) = \dots = \sum_{d \in l+1} R_d^{(l+1)} = \sum_{d \in l} R_d^{(l)} = \dots = \sum_{d} R_d^{(1)}$$

Classifier

output f(x)

# Layer-wise Relevance Propagation



Image x



# Structured output spaces

### Learning Task

For a set of labeled data, we predict the label.

### Difference from multiclass

The set of possible labels  $\mathcal Y$  may be very large or hierarchical.

### Joint kernel on ${\mathcal X}$ and ${\mathcal Y}$

We define a joint feature map on  $\mathcal{X} \times \mathcal{Y}$ , denoted by  $\Phi(\mathbf{x}, y)$ . Then the corresponding kernel function is

$$k((\mathbf{x}, y), (\mathbf{x}', y')) := \langle \Phi(\mathbf{x}, y), \Phi(\mathbf{x}', y') \rangle.$$

### For multiclass

For normal multiclass classification, the joint feature map decomposes and the kernels on  $\mathcal Y$  is the identity, that is

$$k((\mathbf{x}, y), (\mathbf{x}', y')) := [[y = y']]k(\mathbf{x}, \mathbf{x}').$$





# Joint feature map

### Interdependent Outputs

For example a hierarchy of classes like part of speech tagging.

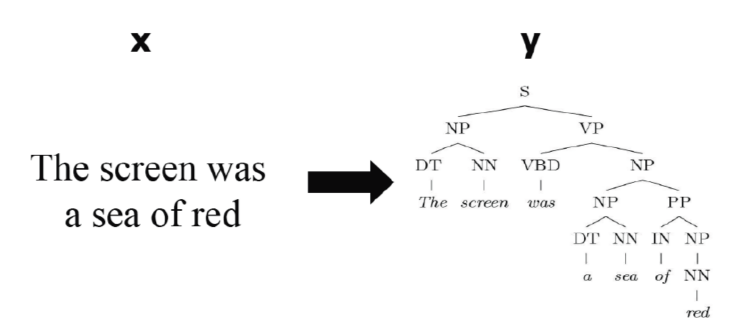
### Label Sequence Learning

Given an input sequence predict a label sequence annotating the input





# **Context free grammar parsing**



### **Recursive Structure**

From Klein & Taskar, ACL'05 Tutorial



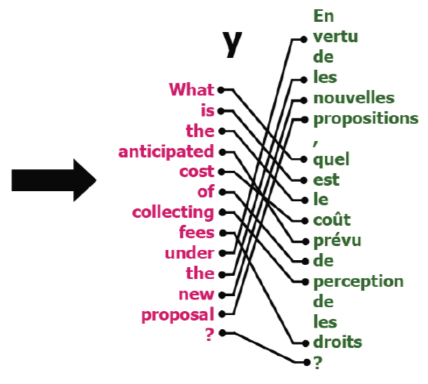


# Bilingual word allignment

X

What is the anticipated cost of collecting fees under the new proposal?

En vertu des nouvelles propositions, quel est le coût prévu de perception des droits?



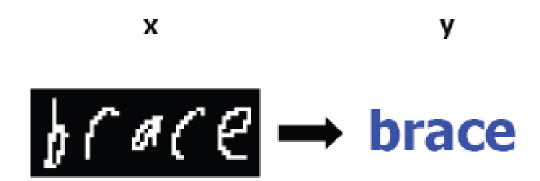
### **Combinatorial Structure**

From Klein & Taskar, ACL'05 Tutorial





# Handwritten letter sequences

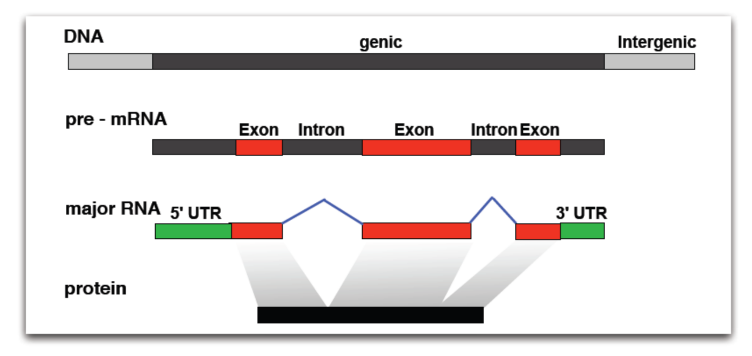






### Label sequence learning

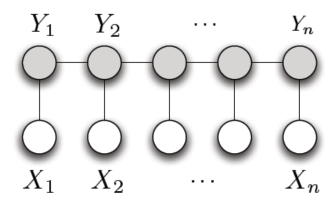
- Given: observation sequence
- Problem: predict corresponding state sequence
- Often: several subsequent positions have the same state
   ⇒ state sequence defines a "segmentation"
- Learn Segmentation for Gene Finding



### **Generative models**

- Hidden Markov Models (Rabiner, 1989)
  - State sequence treated as Markov chain
  - No direct dependencies between observations
  - Example: first-order HMM (simplified)

$$p(\mathbf{x},\mathbf{y}) = \prod_{i} p(x_i|y_i) p(y_i|y_{i-1})$$



Efficient dynamic programming (DP) algorithms





# Decoding via dynamic programming

$$\log p(\mathbf{x}, \mathbf{y}) = \sum_{i} (\log p(x_i|y_i) + \log p(y_i|y_{i-1}))$$
$$= \sum_{i} g(y_{i-1}, y_i, x_i)$$

with  $g(y_{i-1}, y_i, x_i) = \log p(x_i|y_i) + \log p(y_i|y_{i-1})$ .

**Problem:** Given sequence  $\mathbf{x}$ , find sequence  $\mathbf{y}$  such that  $\log p(\mathbf{x}, \mathbf{y})$ 

is maximized, i.e.  $\mathbf{y}^* = \operatorname{argmax}_{\mathbf{y} \in \mathcal{Y}^n} \log p(\mathbf{x}, \mathbf{y})$ 

Dynamic Programming Approach:

$$V(i,y) := \left\{ egin{array}{ll} \max\limits_{y' \in \mathcal{Y}} (V(i-1,y') + g(y',y,x_i)) & i > 1 \ 0 & ext{otherwise} \end{array} 
ight.$$

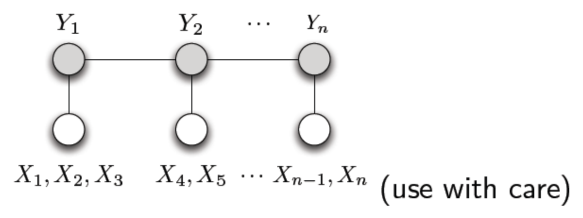




#### **Generative models**

- Generalized Hidden Markov Models
   Hidden Semi-Markov Models
  - Only one state variable per segment
  - Allow non-independence of positions within segment
  - Example: first-order Hidden Semi-Markov Model

$$p(x,y) = \prod_{j} p(\underbrace{(x_{i(j-1)+1}, \dots, x_{i(j)})}_{\mathbf{x}_{i}} | y_{j}) p(y_{j} | y_{j-1})$$



Use generalization of DP algorithms of HMMs

### Decoding via dynamic programming

$$\log p(\mathbf{x}, \mathbf{y}) = \prod_{j} p((x_{i(j)}, \dots, x_{i(j+1)-1})|y_j) p(y_j|y_{j-1})$$

$$= \sum_{j} g(y_{i-1}, y_i, \underbrace{(x_{i(j-1)+1}, \dots, x_{i(j)})}_{\mathbf{x}_j})$$

with  $g(y_{j-1}, y_j, \mathbf{x}_j) = \log p(\mathbf{x}_j | y_j) + \log p(y_j | y_{j-1})$ .

**Problem:** Given sequence  $\mathbf{x}$ , find sequence  $\mathbf{y}$  such that  $\log p(\mathbf{x}, \mathbf{y})$ 

is maximized, i.e.  $\mathbf{y}^* = \operatorname{argmax}_{\mathbf{v} \in \mathcal{V}^*} \log p(\mathbf{x}, \mathbf{y})$ 

Dynamic Programming Approach: V(i, y) :=

$$\begin{cases} \max_{y' \in \mathcal{Y}, d=1, \dots, i-1} (V(i-d, y') + g(y', y, \mathbf{x}_{i-d+1, \dots, i})) & i > 1 \\ 0 & \text{otherwise} \end{cases}$$

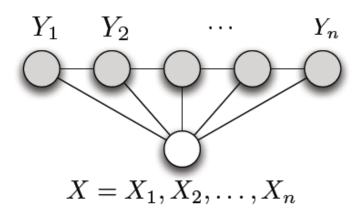




#### **Discriminative models**

- Conditional Random Fields (Lafferty et.al 2001)
  - conditional prob. p(y|x) instead of joint prob. p(x,y)

$$p(y|x, \mathbf{w}) = \frac{1}{Z(x, \mathbf{w})} \exp(\langle \mathbf{w}, \Phi(x, y) \rangle)$$



- can handle non-independent input features
- Semi-Markov Conditional Random Fields
  - introduce segment feature functions
  - dynamic programming algorithms exist

#### Max-margin structured output learning

- Learn function  $f(\mathbf{y}|\mathbf{x})$  scoring segmentations  $\mathbf{y}$  for  $\mathbf{x}$
- Maximize  $f(\mathbf{y}|\mathbf{x})$  w.r.t.  $\mathbf{y}$  for prediction:

$$\operatorname*{argmax}_{\mathbf{y}\in\mathcal{Y}^*}f(\mathbf{y}|\mathbf{x})$$

- Given N sequence pairs  $(\mathbf{x}_1, \mathbf{y}_1), \dots, (\mathbf{x}_N, \mathbf{y}_N)$  for training
- Determine f such that there is a large margin between true and wrong segmentations

$$\begin{aligned} & \min_{f} \quad C \sum_{n=1}^{N} \xi_n + \mathbf{P}[f] \\ & \text{w.r.t.} \quad f(\mathbf{y}_n | \mathbf{x}_n) - f(\mathbf{y} | \mathbf{x}_n) \geq 1 - \xi_n \\ & \text{for all } \mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*, \, n = 1, \dots, N \end{aligned}$$

Exponentially many constraints!

### Joint feature map

#### Recall the kernel trick

For each kernel, there exists a corresponding feature mapping  $\Phi(\mathbf{x})$  on the inputs such that  $k(\mathbf{x}, \mathbf{x}') = \langle \Phi(\mathbf{x}), \Phi(\mathbf{x}') \rangle$ .

#### Joint kernel on ${\mathcal X}$ and ${\mathcal Y}$

We define a joint feature map on  $\mathcal{X} \times \mathcal{Y}$ , denoted by  $\Phi(\mathbf{x}, y)$ . Then the corresponding kernel function is

$$k((\mathbf{x}, y), (\mathbf{x}', y')) := \langle \Phi(\mathbf{x}, y), \Phi(\mathbf{x}', y') \rangle.$$

#### For multiclass

For normal multiclass classification, the joint feature map decomposes and the kernels on  $\mathcal Y$  is the identity, that is

$$k((\mathbf{x}, y), (\mathbf{x}', y')) := [[y = y']]k(\mathbf{x}, \mathbf{x}').$$





#### Structured output learning with kernels

- Assume  $f(\mathbf{y}|\mathbf{x}) = \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \rangle$ , where  $\mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \in \mathcal{F}$
- Use  $\ell_2$  regularizer:  $\mathbf{P}[f] = ||w||^2$

$$\min_{\mathbf{w} \in \mathcal{F}, \boldsymbol{\xi} \in \mathbb{R}^N} \quad C \sum_{n=1}^N \xi_n + \|\mathbf{w}\|^2$$

$$\text{w.r.t.} \quad \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}_n) - \Phi(\mathbf{x}, \mathbf{y}) \rangle \ge 1 - \xi_n$$

$$\text{for all } \mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*, n = 1, \dots, N$$

- Linear classifier that separates true from wrong labelling
- Dual: Define  $\Phi_{n,\mathbf{y}} := \Phi(\mathbf{x}_n,\mathbf{y}_n) \Phi(\mathbf{x}_n,\mathbf{y})$

$$\max_{\boldsymbol{\alpha}} \qquad \sum_{n,\mathbf{y}} \alpha_{n,\mathbf{y}} - \sum_{n,\mathbf{y}} \sum_{n',\mathbf{y}'} \alpha_{n,\mathbf{y}} \alpha_{n',\mathbf{y}'} \langle \Phi_{n,\mathbf{y}}, \Phi_{n',\mathbf{y}'} \rangle$$
 w.r.t. 
$$\alpha_{n,\mathbf{y}} \geq 0, \sum_{\mathbf{y}} \alpha_{n,\mathbf{y}} \leq C \text{ for all } n \text{ and } \mathbf{y}$$

#### **Kernels**

- Recall:  $\Phi_{n,\mathbf{y}} := \Phi(\mathbf{x}_n,\mathbf{y}_n) \Phi(\mathbf{x}_n,\mathbf{y})$
- Then

$$\langle \Phi_{n,\mathbf{y}}, \Phi_{n',\mathbf{y}'} \rangle = \langle \Phi(\mathbf{x}_n, \mathbf{y}_n) - \Phi(\mathbf{x}_n, \mathbf{y}), \Phi(\mathbf{x}_{n'}, \mathbf{y}_{n'}) - \Phi(\mathbf{x}_{n'}, \mathbf{y}')$$

$$= k((\mathbf{x}_n, \mathbf{y}_n), (\mathbf{x}_{n'}, \mathbf{y}_{n'})) - k((\mathbf{x}_n, \mathbf{y}_n), (\mathbf{x}_{n'}, \mathbf{y}')) -$$

$$-k((\mathbf{x}_n, \mathbf{y}), (\mathbf{x}_{n'}, \mathbf{y}_{n'})) + k((\mathbf{x}_n, \mathbf{y}), (\mathbf{x}_{n'}, \mathbf{y})),$$

where

$$k((\mathbf{x}_n,\mathbf{y}),(\mathbf{x}_{n'},\mathbf{y}')) := \langle \Phi(\mathbf{x}_n,\mathbf{y}),\Phi(\mathbf{x}_{n'},\mathbf{y}') \rangle$$

Kernel learning (almost) as usual





## Special case: only two "structures"

• Assume  $f(\mathbf{y}|\mathbf{x}) = \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \rangle$ , where  $\mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \in \mathcal{F}$ 

$$\begin{aligned} \min_{\mathbf{w} \in \mathcal{F}, \boldsymbol{\xi} \in \mathbb{R}^N} \quad & C \sum_{n=1}^N \xi_n + \|w\|^2 \\ \text{w.r.t.} \quad & \langle \mathbf{w}, \Phi(\mathbf{x}, y_n) - \Phi(\mathbf{x}, 1 - y_n) \rangle \geq 1 - \xi_n \\ \text{for all } n = 1, \dots, N \end{aligned}$$

• Dual: Define  $\Phi_n := \Phi(\mathbf{x}_n, y_n) - \Phi(\mathbf{x}_n, 1 - y_n)$ 

$$\max_{\alpha} \qquad \sum_{n} \alpha_{n} - \sum_{n} \sum_{n'} \alpha_{n} \alpha_{n'} \langle \Phi_{n}, \Phi_{n'} \rangle$$
w.r.t. 
$$\alpha_{n} \geq 0, \alpha_{n} \leq C \text{ for all } n$$

Equivalent to standard 2-class SVM

#### **Optimization**

Optimization problem too big (dual as well)

$$\begin{aligned} \min_{\mathbf{w} \in \mathcal{F}, \boldsymbol{\xi}} \quad & C \sum_{n=1}^{N} \xi_n + \|\mathbf{w}\|^2 \\ \text{w.r.t.} \quad & \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}_n) - \Phi(\mathbf{x}, \mathbf{y}) \rangle \geq 1 - \xi_n \\ \text{for all } \mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*, n = 1, \dots, N \end{aligned}$$

- One constraint per example and wrong labeling
- Iterative solution
  - Begin with small set of wrong labellings
  - Solve reduced optimization problem
  - Find labellings that violate constraints
  - Add constraints, resolve
- Guaranteed Convergence

#### How to find violated constraints

Constraint

$$\langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}_n) - \Phi(\mathbf{x}, \mathbf{y}) \rangle \geq 1 - \xi_n$$

• Find labeling **y** that maximizes

$$\langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \rangle$$

Use Dynamic Programming Decoding

$$\mathbf{y} = \underset{\mathbf{y} \in \mathcal{Y}^*}{\operatorname{argmax}} \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \rangle$$

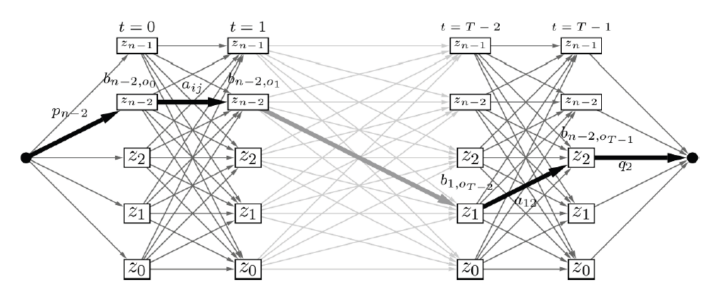
(DP only works if  $\Phi$  has certain decomposition structure)

- If  $\mathbf{y} = \mathbf{y}_n$ , then compute second best labeling as well
- If constraint is violated, then add to optimization problem

### **Dynamic programming**

- number of possible paths of length T for a (fully connected) model with n states is  $n^T$
- infeasible already for small T

### Solution: Use dynamic programming (Viterbi decoding)



• runtime complexity before:  $\mathcal{O}(n^T) \Rightarrow \text{NOW}$ :  $\mathcal{O}(n^2 \cdot T)$ 





## **Algorithm**

- $\mathcal{Y}_n^1 = \emptyset$ , for  $n = 1, \dots, N$
- Solve

$$(\mathbf{w}^{t}, \boldsymbol{\xi}^{t}) = \underset{\mathbf{w} \in \mathcal{F}, \boldsymbol{\xi}}{\operatorname{argmin}} \quad C \sum_{n=1}^{N} \xi_{n} + \|\mathbf{w}\|^{2}$$

$$\text{w.r.t.} \quad \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}_{n}) - \Phi(\mathbf{x}, \mathbf{y}) \rangle \geq 1 - \xi_{n}$$

$$\text{for all } \mathbf{y}_{n} \neq \mathbf{y} \in \mathcal{Y}_{n}^{t}, n = 1, \dots, N$$

**3** Find violated constraints (n = 1, ..., N)

$$\mathbf{y}_n^t = \underset{\mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*}{\operatorname{argmax}} \langle \mathbf{w}^t, \Phi(\mathbf{x}, \mathbf{y}) \rangle$$

If 
$$\langle \mathbf{w}^t, \Phi(\mathbf{x}, \mathbf{y}_n) - \Phi(\mathbf{x}, \mathbf{y}_n^t) \rangle < 1 - \xi_n^t$$
, set  $\mathcal{Y}_n^{t+1} = \mathcal{Y}_n^t \cup \{\mathbf{y}_n^t\}$ 

- If violated constraint exists then go to 2
- Otherwise terminate ⇒ Optimal solution





#### **Loss functions**

- So far 0-1-loss with slacks: If  $\mathbf{y} \neq \mathbf{y}$ , then prediction is wrong, but it does not matter how wrong
- Introduce loss function on labellings  $\ell(y, y')$ , e.g.
  - How many segments are wrong or missing
  - How different are the segments, etc





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- Extend optimization problem (Margin rescaling):

$$\begin{aligned} & \min_{\mathbf{w} \in \mathcal{F}, \boldsymbol{\xi}} & C \sum_{n=1}^{N} \xi_n + \|\mathbf{w}\|^2 \\ & \text{w.r.t.} & \langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}_n) - \Phi(\mathbf{x}, \mathbf{y}) \rangle \geq \ell(\mathbf{y}, \mathbf{y}') - \xi_n \\ & \text{for all } \mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*, n = 1, \dots, N \end{aligned}$$

• Finding violated constraints (n = 1, ..., N)

$$\mathbf{y}_n^t = \underset{\mathbf{y}_n \neq \mathbf{y} \in \mathcal{Y}^*}{\operatorname{argmax}} \langle \mathbf{w}^t, \Phi(\mathbf{x}, \mathbf{y}) \rangle + \ell(\mathbf{y}, \mathbf{y}_n)$$





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- Introduce loss function on labellings  $\ell(y, y')$ , e.g.
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Finding violated constraints more difficult





#### **Problems**

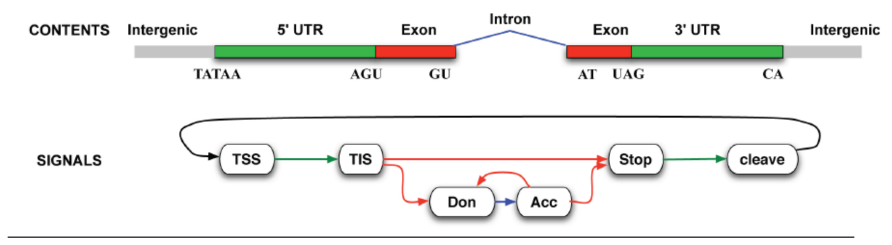
- Optimization may require many iterations
- Number of variables increases linearly
- When using kernels, solving optimization problems can become infeasible
- Evaluation of  $\langle \mathbf{w}, \Phi(\mathbf{x}, \mathbf{y}) \rangle$  in Dynamic programming can be very expensive
  - Optimization and decoding become too expensive
- Approximation algorithms useful
- Decompose problem
  - First part uses kernels, can be precomputed
  - Second part without kernels and only combines ingredients





#### Gene finding as a segmentation task

- Nodes correspond to sequence signals
  - Depend on recognition of signals on the DNA
- Transitions correspond to segments
  - Depend on length or sequence properties of segment
- Markovian on segment level, non-Markovian within segments
  - Allows efficient decoding and modeling of segment lengths







### Learning to predict segmentations

- Learn function f(y|x) scoring segmentations y for x
- f considers signal, content and length information
- Maximize  $f(\mathbf{y}|\mathbf{x})$  w.r.t.  $\mathbf{y}$  for prediction:  $\underset{\mathbf{y}}{\operatorname{argmax}} f(\mathbf{y}|\mathbf{x})$
- Determine f such that there is a large margin between true and wrong segmentations

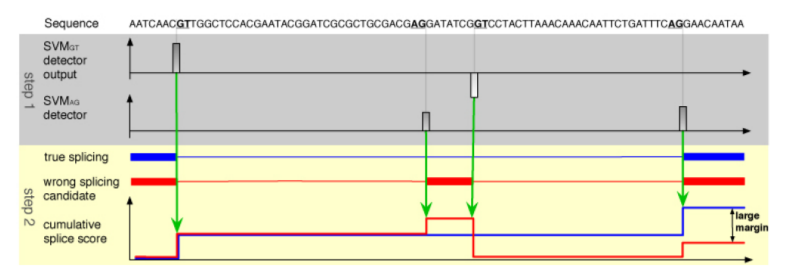
$$\begin{aligned} & \min_{f} & & \sum_{n=1}^{N} \xi_n + \mathbf{P}[f] \\ & \text{w.r.t.} & & f(\mathbf{y}_n | \mathbf{x}_n) - f(\mathbf{y} | \mathbf{x}_n) \geq 1 - \xi_n \\ & & \text{for all } \mathbf{y} \neq \mathbf{y}_n, n = 1, \dots, N \end{aligned}$$

- Use approximation (Rätsch & Sonnenburg, NIPS'06)
  - Train signal and content detectors separately
  - Combine in large margin fashion





## Large margin combination (simplified)



• Simplified Model: Score for splice form  $\mathbf{y} = \{(p_j, q_j)\}_{j=1}^J$ :

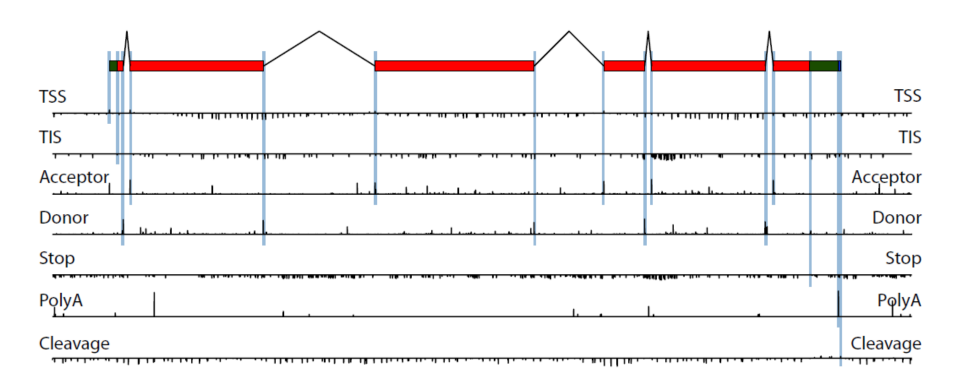
$$f(\mathbf{y}) := \underbrace{\sum_{j=1}^{J-1} S_{GT}(f_j^{GT}) + \sum_{j=2}^{J} S_{AG}(f_j^{AG})}_{\text{Splice signals}} + \underbrace{\sum_{j=1}^{J-1} S_{L_I}(p_{j+1} - q_j) + \sum_{j=1}^{J} S_{L_E}(q_j - p_j)}_{\text{Segment lengths}}$$

• Tune free parameters (in functions  $S_{GT}$ ,  $S_{AG}$ ,  $S_{L_E}$ ,  $S_{L_I}$ ) by solving linear program using training set with known splice forms





# **Example**







## **Results summary**

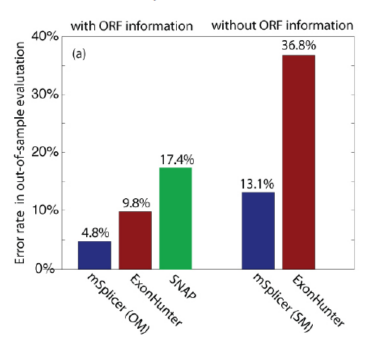
- Splicing only (Rätsch et al., PLoS Comp. Biol., 2007)
  - Comparison with other methods
  - Analysis of a few disagreeing cases
  - Results available on http://www.wormbase.org
- Full gene predictions
  - Relevant for the nGASP competition
  - Evaluation by organizers still pending





## Results I (Splice forms only)

- - 60% for training and validation
  - 40% for testing (exclude alt. spliced genes)
- Out-of-sample accuracy ( $\approx$ 1100 gene models):
  - Splice form error rate
    - 4.8% (coding)
    - 13.1% (mixed)
  - Much lower error rates than state-of-the-art
    - Exonhunter (Brejova et al., ISMB'05)
    - Snap (Korf, BMC Bioinformatics 2004)

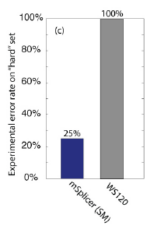


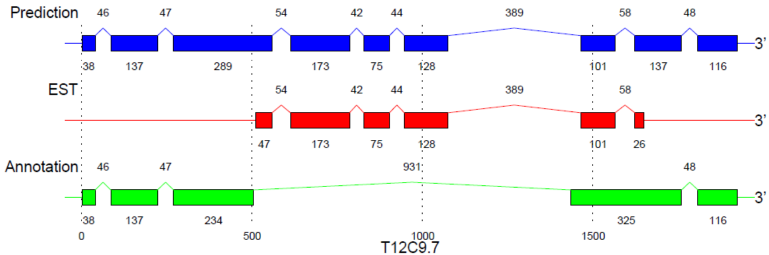




## Results II (Splice forms only)

- Validation by RT-PCR & direct sequencing
  - Consider 20 disagreeing cases
  - Annotation was never correct
  - 75% of our predictions were correct









#### mGene - Summary

# 2-step approach

- Content and Signal Sensors(transcription start,...)
  - Support Vector Machine with String Kernel (spectrum, weighted degree,...)
- Label Sequence (Segmentation) Learning
  - Joint feature maps for inputs and outputs
  - Related to (generalized) HMMs
  - Result in large optimization problems
    - Can be solved iteratively
    - But still too large for medium size problems
  - Decomposition of the Problem
    - Use efficient kernel-based two-class detectors
    - Integrate without kernels
- Beats HMM based approaches in Gene finding :-)