For written notes on this lecture, please read Chapters 4 and 7 of *The Practical Bioinformatician*, and Koh & Wong, "Recognition of Polyadenylation Sites from Arabidopsis Genomic Sequences", *Proc GIW* 2007, pages 73--82

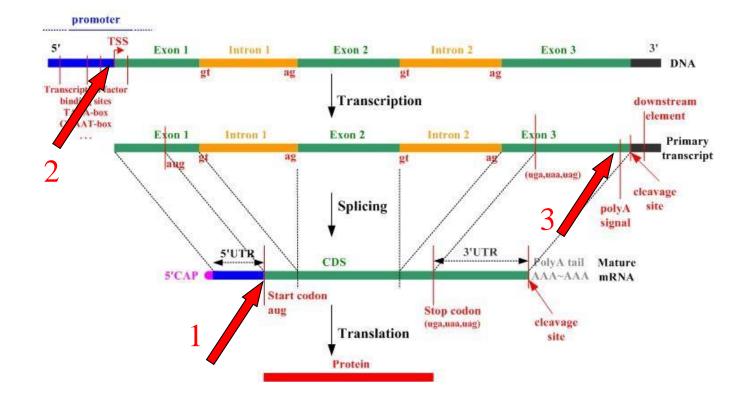
CS2220: Introduction to Computational Biology Unit 3: Gene Feature Recognition

Wong Limsoon 8 September 2016 15 September 2016



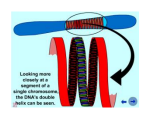
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Plan



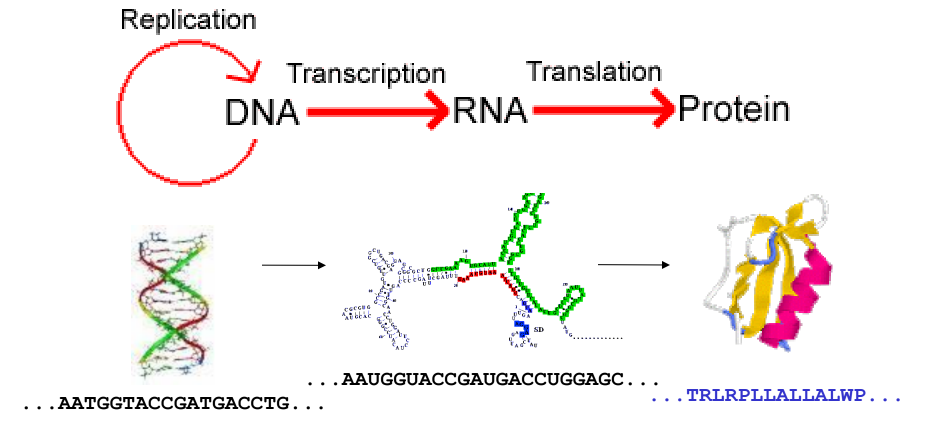
Some Relevant Biology

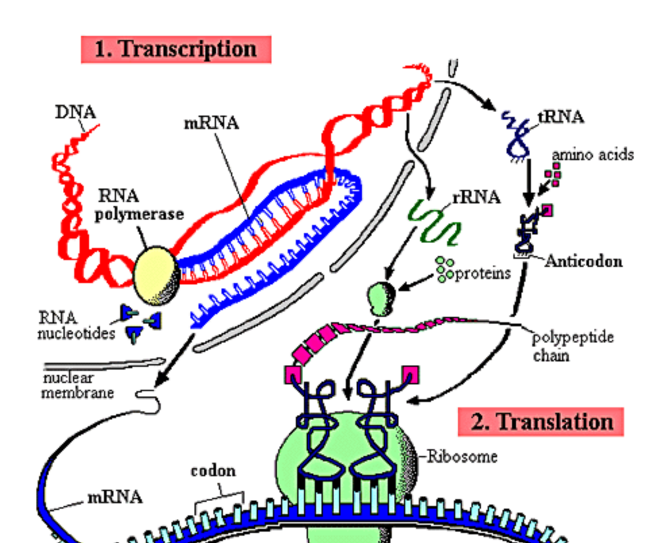




Central Dogma



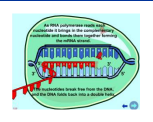




Protein synthesis



Players in protein synthesis

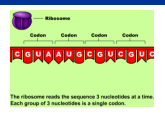


Transcription



- Synthesize mRNA from one strand of DNA
 - An enzyme RNA polymerase temporarily separates doublestranded DNA
 - It begins transcription at transcription start site
 - A → A, C→C, G→G, &
 T→U
 - Once RNA polymerase reaches transcription stop site, transcription stops

- Additional "steps" for Eukaryotes
 - Transcription produces pre-mRNA that contains both introns & exons
 - 5' cap & poly-A tail are added to pre-mRNA
 - RNA splicing removes introns & mRNA is made
 - mRNA are transported out of nucleus



Translation



- Synthesize protein from mRNA
- Each amino acid is encoded by consecutive seq of 3 nucleotides, called a codon
- The decoding table from codon to amino acid is called genetic code

- 4³=64 diff codons
- ⇒ Codons are not 1-to-1 corr to 20 amino acids
- All organisms use the same decoding table (except some mitochrondrial genes)
- Amino acids can be classified into 4 groups. A single-base change in a codon is usu insufficient to cause a codon to code for an amino acid in diff group

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Genetic code

- Start codon
 - ATG (code for M)
- Stop codon
 - TAA
 - TAG
 - TGA

		Second Position of Codon					
		T	C	A	G		
F	T	TTT Phe [F] TTC Phe [F] TTA Leu [L] TTG Leu [L]	TCT Ser [S] TCC Ser [S] TCA Ser [S] TCG Ser [S]	TAT Tyr [Y] TAC Tyr [Y] TAA Ter [end] TAG Ter [end]	TGT Cys [C] TGC Cys [C] TGA Ter [end] TGG Trp [W]	T C A G	T
r s t	С	CTT Leu [L] CTC Leu [L] CTA Leu [L] CTG Leu [L]	CCT Pro [P] CCC Pro [P] CCA Pro [P] CCG Pro [P]	CAT His [H] CAC His [H] CAA Gln [Q] CAG Gln [Q]	CGT Arg [R] CGC Arg [R] CGA Arg [R] CGG Arg [R]	T C A G	i r d
o s i t	A	ATT Ile [I] ATC Ile [I] ATA Ile [I] ATG Met [M]	ACT Thr [T] ACC Thr [T] ACA Thr [T] ACG Thr [T]	AAT Asn [N] AAC Asn [N] AAA Lys [K] AAG Lys [K]	AGT Ser [S] AGC Ser [S] AGA Arg [R] AGG Arg [R]	T C A G	o s i t
i o n	G	GTT Val [V] GTC Val [V] GTA Val [V] GTG Val [V]	GCT Ala [A] GCC Ala [A] GCA Ala [A] GCG Ala [A]	GAT Asp [D] GAC Asp [D] GAA Glu [E] GAG Glu [E]	GGT Gly [G] GGC Gly [G] GGA Gly [G] GGG Gly [G]	T C A G	n

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Example

Example of computational translation - notice the indication of (alternative) start-codons:

```
VIRTUAL RIBOSOME
Translation table: Standard SGCO
>Sea1
Reading frame: 1
  M V L S A A D K G N V K A A W G K V G G H A A E Y G A E A L
5' ATGGTGCTGTCTGCCGCCGACAAGGCCAATGTCAAGGCCGCCTGGGGCAAGGTTGGCGGCCACGCTGCAGAGTATGGCGCAGAGGCCCTG 90
                    KTYFPHFDLSHGSAQVKGHG
5' GAGAGGATGTTCCTGAGCTTCCCCACCACCACCACCACTTCCCCCACTTCGACCTGAGCCACGGCTCCGCGCAGGTCAAGGGCCACGGC 180
  .....>>>...)))......
  AKVAAALTKAVEHLDDLP
                                    A L
                                  G
.....))).....))).....)))
            V N F K L L S H S L L V T L A S H L
5' AAGCTGCGTGTGGACCCGGTCAACTTCAAGCTTCTGAGCCACTCCCTGCTGGTGACCCTGGCCTCCCCACTCCCCAGTGATTTCACCCCC 360
  ...)))......))
5' GCGGTCCACGCCTCCCTGGACAAGTTCTTGGCCAACGTGAGCACCGTGCTGACCTCCAAATACCGTTAA 429
  ......)))......***
Annotation key:
>>> : START codon (strict)
))) : START codon (alternative)
*** : STOP
```

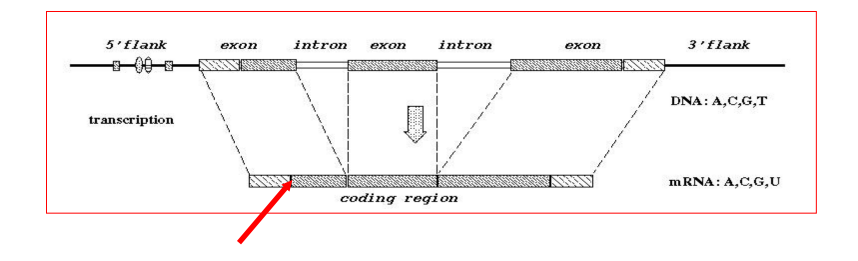
Recognition of Translation Initiation Sites

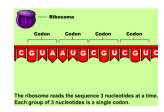
An introduction to the World's simplest TIS recognition system





Translation initiation site







A sample cDNA

299 HSU27655.1 CAT U27655 Homo sapiens	
CGTGTGTGCAGCAGCCTGCAGCTGCCCCAAGCCATGACACACTGACTCCCAGCTGTG	80
CCCAGGGCTTCAAAGACTTCTCAGCTTCGAGC <u>ATG</u> GCTTTTGGCTGTCAGGGCAGCTGTA	160
GGAGGCAG <u>ATG</u> AGAAGAGGGAG <u>ATG</u> GCCTTGGAGGAAGGGAAGGGGCCTGGTGCCGAGGA	240
CCTCTCCTGGCCAGGAGCTTCCTCCAGGACAAGACCTTCCACCCAACAAGGACTCCCCT	
	80
ieeeeeeeeeeeeeeee	160
EEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEEE	240
REFERENCE REFEREN	

What makes the second ATG the TIS?

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Approach

- Training data gathering
- Signal generation
 - k-grams, distance, domain know-how, ...
- Signal selection
 - Entropy, χ2, CFS, t-test, domain know-how...
- Signal integration
 - SVM, ANN, PCL, CART, C4.5, kNN, ...



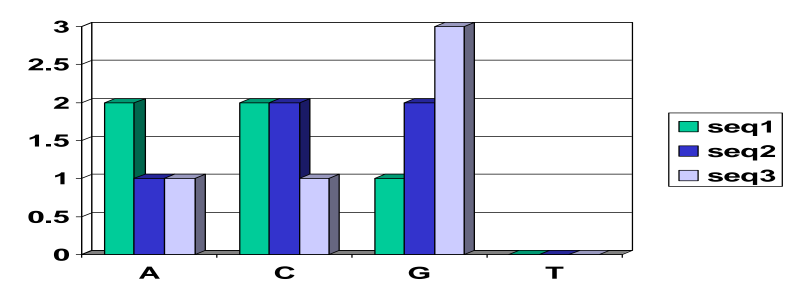
Training & testing data

- Vertebrate dataset of Pedersen & Nielsen [ISMB'97]
- 3312 sequences
- 13503 ATG sites
- 3312 (24.5%) are TIS
- 10191 (75.5%) are non-TIS
- Use for 3-fold x-validation expts



Signal generation

- K-grams (ie., k consecutive letters)
 - -K = 1, 2, 3, 4, 5, ...
 - Window size vs. fixed position
 - Up-stream, downstream vs. any where in window
 - In-frame vs. any frame





Signal generation: Example

299 HSU27655.1 CAT U27655 Homo sapiens

CGTGTGTGCAGCAGCCTGCAGCTGCCCCAAGCCATGGCTGAACACTGACTCCCAGCTGTG 80

CCCAGGGCTTCAAAGACTTCTCAGCTTCGAGCATGGCTTTTTGGCTCTCAGGGCAGCTGTA 160

GGAGGCAGATGAGAAGAGGGAGATGGCCTTGGAGGAAGGGGAAGGGGCCTGGTGCCGAGGA 240

CCTCTCCTGGCCAGGACCTTCCTCCAGGACAAGACCTTCCACCCAACAAGGACTCCCCT

- Window = ±100 bases
- In-frame, downstream

Any-frame, downstream

$$-GCT = 3$$
, $TTT = 2$, $ATG = 2$...

In-frame, upstream

$$-GCT = 2$$
, $TTT = 0$, $ATG = 0$, ...

Exercise: Find the in-frame downstream ATG

Exercise: What are the possible k-grams (k=3) in this sequence?

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Feature generation - Summary

Raw Data

206 BBCALCB.1 CAT X71666 Eukaryotae; mitochondrial eukaryotes; Metazoa; Chordata; CCGTCAGAGCGCCGACCGCCGACCGCCGAGCAGAATGGGAAATGAGGCAAGTTATCCT TTGGAAATGTGCTCACACTTTGATGCAGATGAAATTAAAAGGCTAGGAAAGAGATTTAAGAAGCTCGATTTGGACAATTC TGGTTCTTTGAGTGGAAAGAGTTCATGTCTCTACCTGAGTTACAA



An ATG segment – positive sample

> 206 +1_Index(56)



A feature vector --- upstream/downstream inframe 3 grams



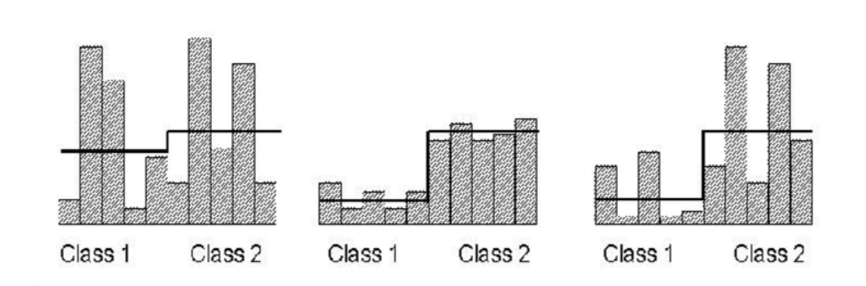
Too many features

- For each value of k, there are 4^k * 3 * 2 k-grams
- If we use k = 1, 2, 3, 4, 5, we have 24 + 96 + 384 + 1536 + 6144 = 8184 features!
- This is too many for most machine learning algorithms



Signal selection: Basic idea

- Choose a signal w/ low intra-class distance
- Choose a signal w/ high inter-class distance



Signal selection: E.g., t-statistics

The t-stats of a signal is defined as

$$t = \frac{|\mu_1 - \mu_2|}{\sqrt{(\sigma_1^2/n_1) + (\sigma_2^2/n_2)}}$$

where σ_i^2 is the variance of that signal in class i, μ_i is the mean of that signal in class i, and n_i is the size of class i.

Signal selection: E.g., MIT-correlation National University of Singapore

The MIT-correlation value of a signal is defined as

$$MIT = \frac{|\mu_1 - \mu_2|}{\sigma_1 + \sigma_2}$$

where σ_i is the standard deviation of that signal in class i and μ_i is the mean of that signal in class i.



Signal selection: E.g., χ2

The \mathcal{X}^2 value of a signal is defined as:

$$\mathcal{X}^{2} = \sum_{i=1}^{m} \sum_{j=1}^{k} \frac{(A_{ij} - E_{ij})^{2}}{E_{ij}},$$

where m is the number of intervals, k the number of classes, A_{ij} the number of samples in the ith interval, jth class, R_i the number of samples in the ith interval, C_j the number of samples in the jth class, N the total number of samples, and E_{ij} the expected frequency of A_{ij} ($E_{ij} = R_i * C_j/N$).



Example

 Suppose you have a sample of 50 men and 50 women and the following weight distribution is observed:

	obs	exp	(obs – exp)²/exp
НМ	40	60*50/100=30	3.3
HW	20	60*50/100=30	3.3
LM	10	40*50/100=20	5.0
LW	30	40*50/100=20	5.0

$$\chi 2=16.6$$

P = 0.00004,
df = 1
So weight and
sex are not indep

Is weight a good attribute for distinguishing men from women?



Signal selection: E.g., CFS

 Instead of scoring individual signals, how about scoring a group of signals as a whole?

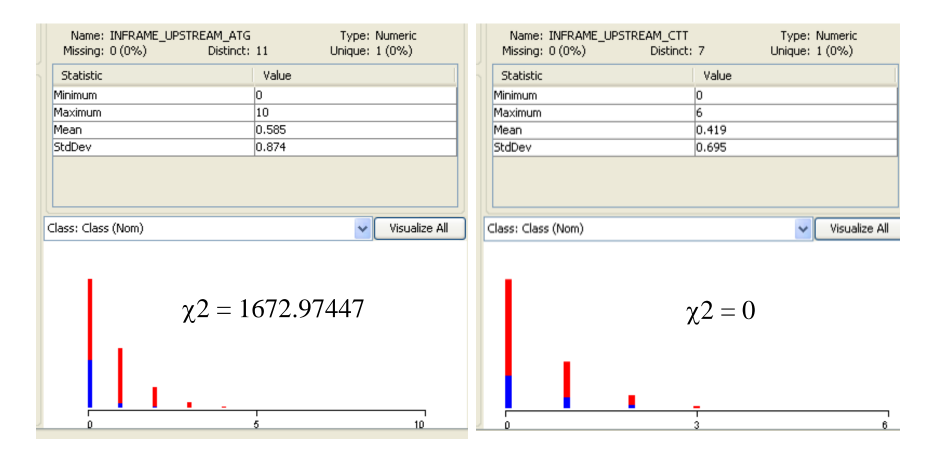
CFS

- Correlation-based Feature Selection
- A good group contains signals that are highly correlated with the class, and yet uncorrelated with each other

Exercise: What is the main challenge in implementing CFS?



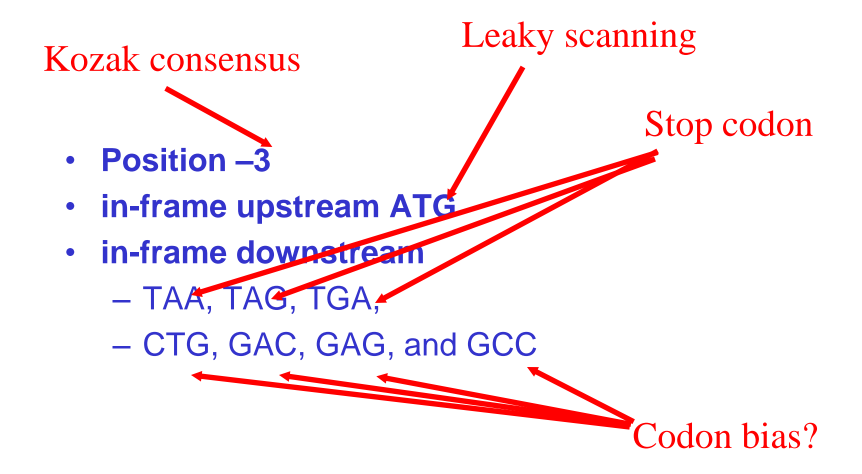
Distributions of two 3-grams



Which is the better one?

Sample k-grams selected by CFS NUS for recognizing TIS







Signal integration

kNN

 Given a test sample, find the k training samples that are most similar to it. Let the majority class win

SVM

- Given a group of training samples from two classes, determine a separating plane that maximises the margin of error
- Naïve Bayes, ANN, C4.5, ...



Results: 3-fold x-validation

	predicted	predicted
	as positive	as negative
positive	TP	FN
negative	FP	TN

Exercise: What is TP/(TP+FP)?

	TP/(TP + FN)	TN/(TN + FP)	TP/(TP + FP)	Accuracy
Naïve Bayes	84.3%	86.1%	66.3%	85.7%
SVM	73.9%	93.2%	77.9%	88.5%
Neural Network	77.6%	93.2%	78.8%	89.4%
Decision Tree	74.0%	94.4%	81.1%	89.4%



Improvement by voting

 Apply any 3 of Naïve Bayes, SVM, Neural Network, & Decision Tree. Decide by majority

	TP/(TP + FN)	TN/(TN + FP)	TP/(TP + FP)	Accuracy
NB+SVM+NN	79.2%	92.1%	76.5%	88.9%
NB+SVM+Tree	78.8%	92.0%	76.2%	88.8%
NB+NN+Tree	77.6%	94.5%	82.1%	90.4%
SVM+NN+Tree	75.9%	94.3%	81.2%	89.8%
Best of 4	84.3%	94.4%	81.1%	89.4%
Worst of 4	73.9%	86.1%	66.3%	85.7%



Improvement by scanning

- Apply Naïve Bayes or SVM left-to-right until first ATG predicted as positive. That's the TIS
- Naïve Bayes & SVM models were trained using TIS vs. Up-stream ATG

	TP/(TP + FN)	TN/(TN + FP)	TP/(TP + FP)	Accuracy
NB	84.3%	86.1%	66.3%	85.7%
SVM	73.9%	93.2%	77.9%	88.5%
NB+Scanning	87.3%	96.1%	87.9%	93.9%
SVM+Scanning	88.5%	96.3%	88.6%	94.4%



Performance comparison

	TP/(TP + FN)	TN/(TN + FP)	TP/(TP + FP)	Accuracy
NB	84.3%	86.1%	66.3%	85.7%
Decision Tree	74.0%	94.4%	81.1%	89.4%
NB+NN+Tree	77.6%	94.5%	82.1%	90.4%
SVM+Scanning	88.5%	96.3%	88.6%	94.4%*
Pedersen&Nielsen	78%	87%	-	85%
Zien	69.9%	94.1%	-	88.1%
Hatzigeorgiou	-	-	-	94%*

^{*} result not directly comparable



Technique comparison

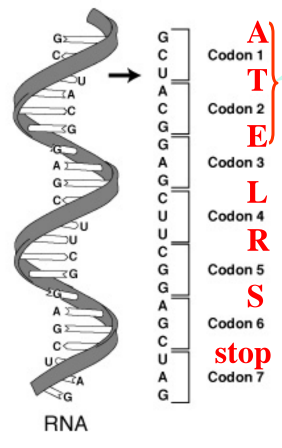
- Pedersen&Nielsen [ISMB'97]
 - Neural network
 - No explicit features
- Zien [Bioinformatics'00]
 - SVM+kernel engineering
 - No explicit features
- Hatzigeorgiou
 [Bioinformatics'02]
 - Multiple neural networks
 - Scanning rule
 - No explicit features

Our approach

- Explicit feature generation
- Explicit feature selection
- Use any machine learning method w/o any form of complicated tuning
- Scanning rule is optional



mRNA-protein



Ribonucleic acid

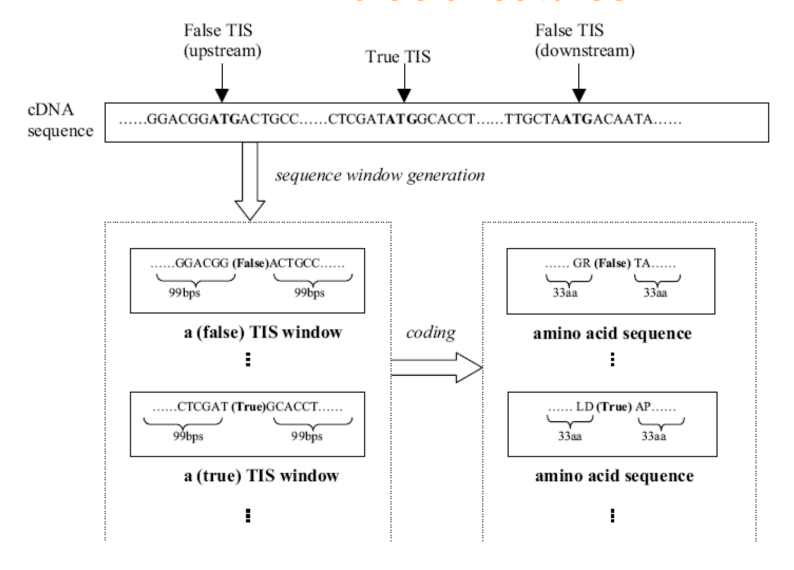
Exercise: List the first 10 amino acid in our example sequence

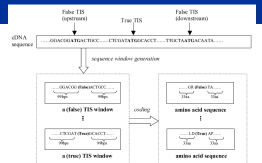
How about using k-grams from the translation?

First	U	С	A	G	Last
U	Phe F	Ser S	Tyr Y	Суѕ	U
	Phe	Ser	Tyr	Суѕ	С
	Leu T.	Ser	Stop (Ochre)	Stop (Umber)	A
	Leu	Ser	Stop (Amber)	Trp W	G
С	Leu	Pro p	His H	Arg R	U
	Leu	Pro	His	Arg	С
	Leu	Pro	Gln O	Arg	A
	Leu	Pro	Gln	Arg	G
A	lle 🕇	Thr T	Asn N	Ser	U
	Ile 📩	Thr	Asn	Ser	С
	Пе	Thr	Lys K	Arg	A
	Met M	Thr	Lys	Arg	G
G	Val V	Ala 🔼	Asp D	Gly G	U
	Val	Ala	Asp	Gly	С
	Val	Ala	Glu 🗜	Gly	A
	Val	Ala	Glu	Gly	G



Amino-acid features





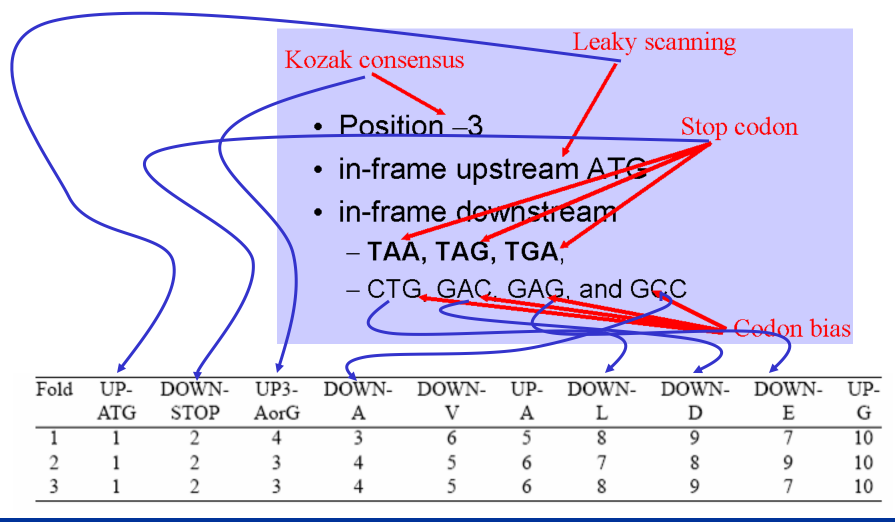


Amino-acid features

New feature space (total of 927 features + class label)						
42 1-gram amino acid patterns						
UP-A, UP-R, ,UP-N, DOWN- A, DOWN-R,, DOWN-N (numeric type)	,UP-N, DOWN-AA, DOWN-AA, DOWN-AR,, DOWN-NN		True, False			
	Frequency as values					
1, 3, 5, 0, 4,	6, 2, 7, 0, 5,	N, N, N,	False			
6, 5, 7, 9, 0, 2, 0, 3, 10, 0,		Y, Y, Y,	True			

Amino acid K-grams discovered by entropy







Independent validation sets

A. Hatzigeorgiou:

- 480 fully sequenced human cDNAs
- 188 left after eliminating sequences similar to training set (Pedersen & Nielsen's)
- 3.42% of ATGs are TIS

Our own:

 well characterized human gene sequences from chromosome X (565 TIS) and chromosome 21 (180 TIS)

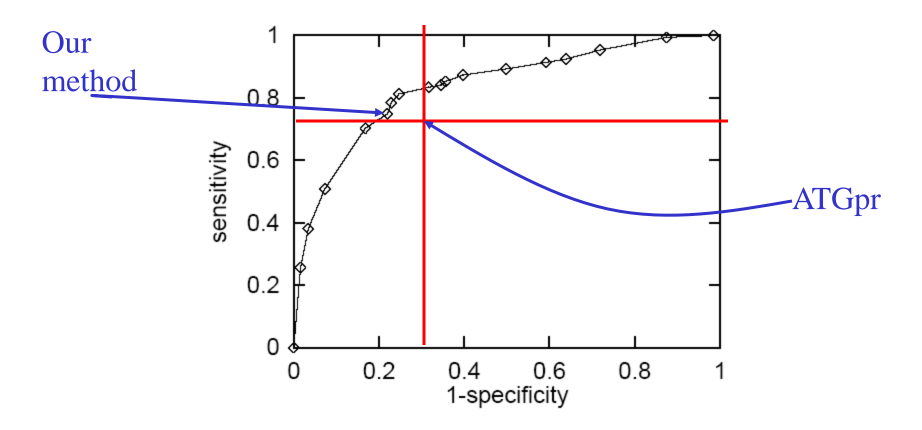
Validation results, on Hatzigeorgiou

Algorithm	Sensitivity	Specificity	Precision	Accuracy
SVMs(linear)	96.28%	89.15%	25.31%	89.42%
SVMs(quad)	94.14%	90.13%	26.70%	90.28%
Ensemble Trees	92.02%	92.71%	32.52%	92.68%
OT 73 6 /1'	05.010/	00 5407	21.000/	00.000/

 Using top 100 features selected by entropy and trained on Pedersen & Nielsen's dataset







 Using top 100 features selected by entropy and trained on Pedersen & Nielsen's

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About the inventor: Huiqing Liu

Huiqing Liu

- PhD, NUS, 2004
- Currently PI at Incyte
- Asian InnovationGold Award 2003
- New Jersey Cancer
 Research Award for
 Scientific Excellence
 2008
- Gallo Prize 2008



Recognition of Transcription Start Sites

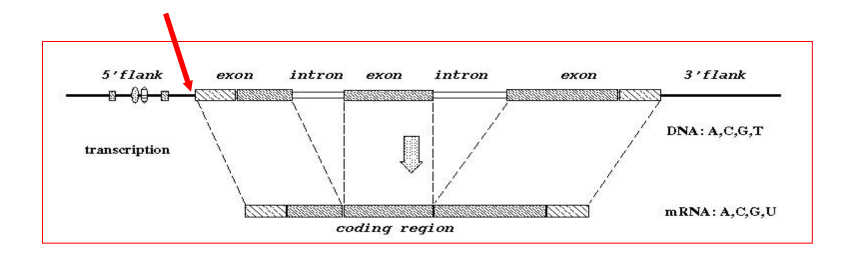
An introduction to the World's best TSS recognition system:

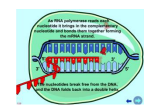
A heavy tuning approach





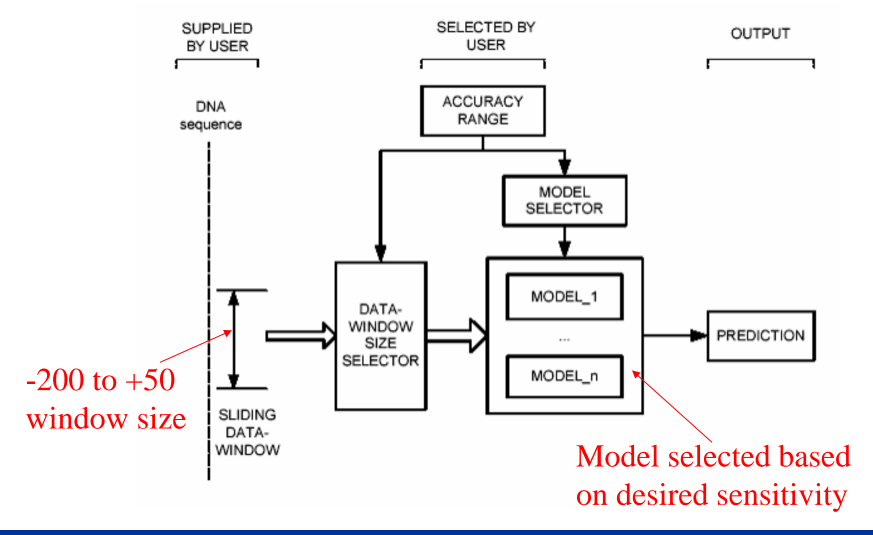
Transcription start site

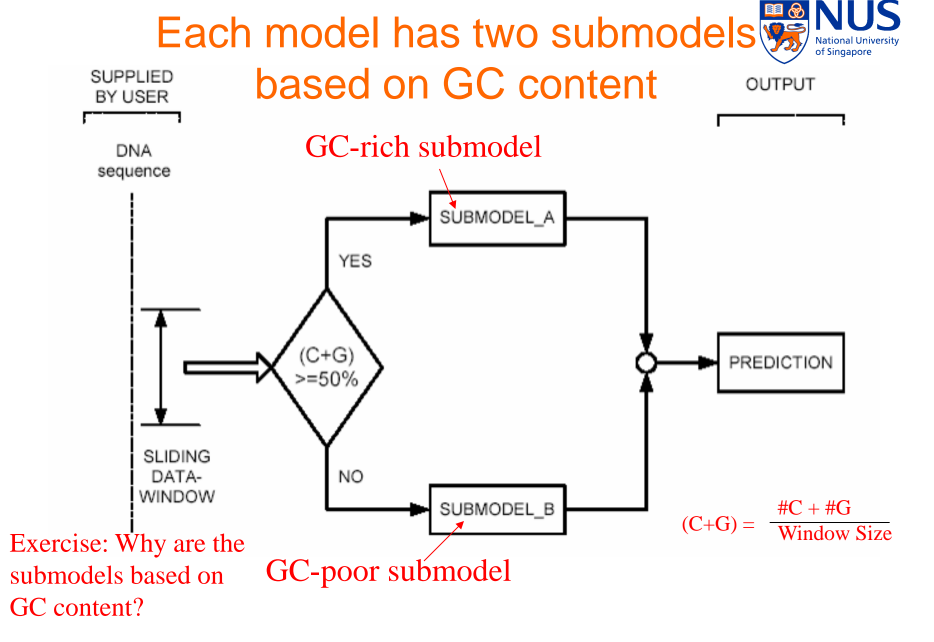




of Singapore

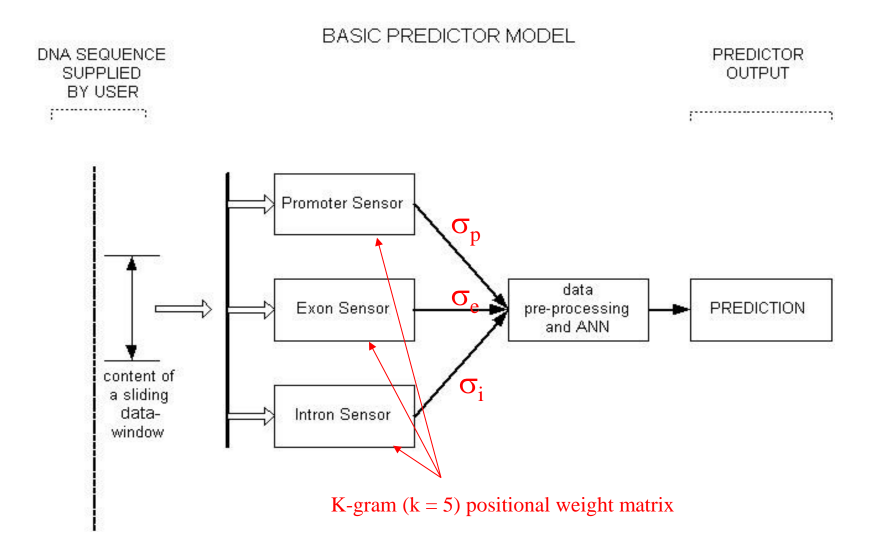
Structure of Dragon Promoter Finder







Data analysis within submodel





Promoter, exon, intron sensors

- These sensors are positional weight matrices of k-grams, k = 5 (aka pentamers)
- They are calculated as below using promoter, exon, intron data respectively

 Pentamer at ith

Window size
$$\boldsymbol{\sigma} = \underbrace{ \begin{bmatrix} \sum_{i=1}^{L-4} p_j^i \otimes f_{j,i} \\ \sum_{i=1}^{L-4} \max f_{j,i} \\ \end{bmatrix}}_{\text{Frequency of jth pentamer at ith position in training window}}, \quad p_j^i \otimes f_{j,i} = \underbrace{ \begin{cases} f_{j,i}, \text{ if } p_i = p_j^i \\ 0, \text{ if } p_i \neq p_j^i \\ \end{bmatrix}_{\text{jth pentamer at ith position in training window}}^{\text{position in input}},$$

Just to make sure you know what I mean ... Nation of Single

Give me 3 DNA seq of length 10:

- $Seq_1 = ACCGAGTTCT$
- Seq₂ = AGTGTACCTG
- Seq₃ = AGTTCGTATG

Then

1-mer	pos1	pos2	pos3	pos4	pos5	pos6	pos7	pos8	pos9	pos10
Α	3/3	0/3	0/3							
C	0/3	1/3	1/3		Exerc	ise: Fil	l in the	rest of t	he table	
G	0/3	2/3	0/3							
Т	0/3	0/3	2/3							

Just to make sure you know what I mean.

Give me 3 DNA seq of length 10:

- $Seq_1 = ACCGAGTTCT$
- Seq₂ = AGTGTACCTG
- $Seq_3 = AGTTCGTATG$
- Then

Exercise: How many rows should this 2-mer table have? How many rows should the pentamer table have?

2-mer	pos1	pos2	pos3	pos4	pos5	pos6	pos7	pos8	pos9
AA	0/3	0/3	0/3						
AC	1/3	0/3	0/3		Exerci	se: Fill	in the re	est of th	e table
			:						
TT	0/3	0/3	1/3				1/3		



Data preprocessing & ANN

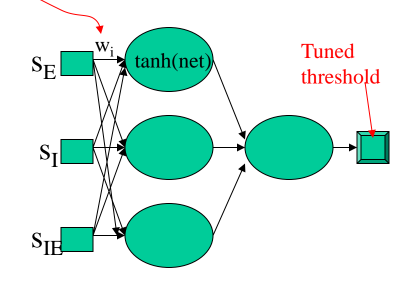
Tuning parameters

$$\begin{aligned} s_E &= sat(\sigma_p - \sigma_e, a_e, b_e), \\ s_I &= sat(\sigma_p - \sigma_i, a_i, b_i), \\ s_{EI} &= sat(\sigma_e - \sigma_i, a_{ei}, b_{ei}), \end{aligned}$$

where the function *sat* is defined by

$$sat(x,a,b) = \begin{cases} a, & \text{if } x > a \\ x, & \text{if } b \le x \le a. \\ b, & \text{if } b > x \end{cases}$$

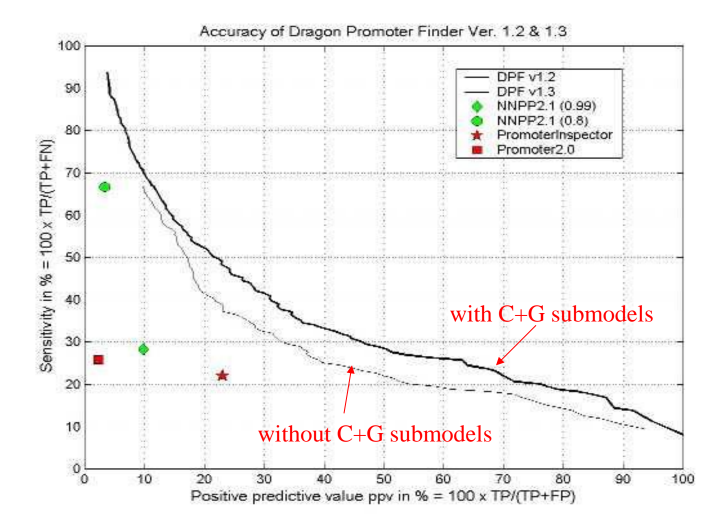
Simple feedforward ANN trained by the Bayesian regularisation method



$$tanh(x) = \frac{e^{x} - e^{-x}}{e^{x} + e^{-x}}$$
$$net = \sum s_{i} * w_{i}$$



Accuracy comparison



Training data criteria & preparation

- Contain both positive and negative sequences
- Sufficient diversity, resembling different transcription start mechanisms
- Sufficient diversity, resembling different nonpromoters
- Sanitized as much as possible

- TSS taken from
 - 793 vertebrate promoters from EPD
 - -200 to +50 bp of TSS
- non-TSS taken from
 - GenBank,
 - 800 exons
 - 4000 introns,
 - 250 bp,
 - non-overlapping,
 - <50% identities</p>



Tuning data preparation

 To tune adjustable system parameters in Dragon, we need a separate tuning data set

TSS taken from

- 20 full-length gene seqs
 with known TSS
- 200 to +50 bp of TSS
- no overlap with EPD

Non-TSS taken from

- 1600 human 3'UTR seqs
- 500 human exons
- 500 human introns
- 250 bp
- no overlap

Testing data criteria & preparation

- Seqs should be from the training or evaluation of other systems (no bias!)
- Seqs should be disjoint from training and tuning data sets
- Seqs should have TSS
- Seqs should be cleaned to remove redundancy, <50% identities

- 159 TSS from 147 human and human virus seqs
- cummulative length of more than 1.15Mbp
- Taken from GENESCAN, Geneld, Genie, etc.



About the inventor: Vlad Bajic

Vladimir B. Bajic

- Principal Scientist,
 I²R, 2001-2006
- Currently Director &
 Professor,
 Computational
 Bioscience Research
 Center, KAUST

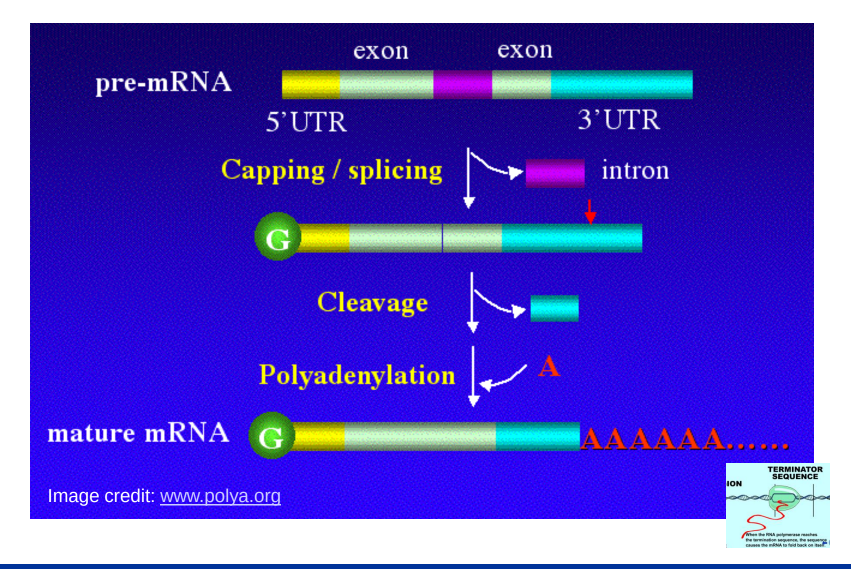


Recognition of Poly-A Signal Sites

A twist to the "feature generation, feature selection, feature integration" approach



Eukaryotic pre-mRNA processing



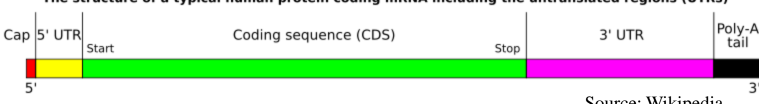


Polyadenylation in eukaryotes

- Addition of poly(A) tail to RNA
 - Begins as transcription finishes
 - 3'-most segment of newly-made RNA is cleaved off
 - Poly(A) tail is then synthesized at 3' end

- Poly(A) tail is impt for nuclear export, translation & stability of mRNA
- Tail is shortened over time. When short enough, the mRNA is degraded

The structure of a typical human protein coding mRNA including the untranslated regions (UTRs)



Source: Wikipedia

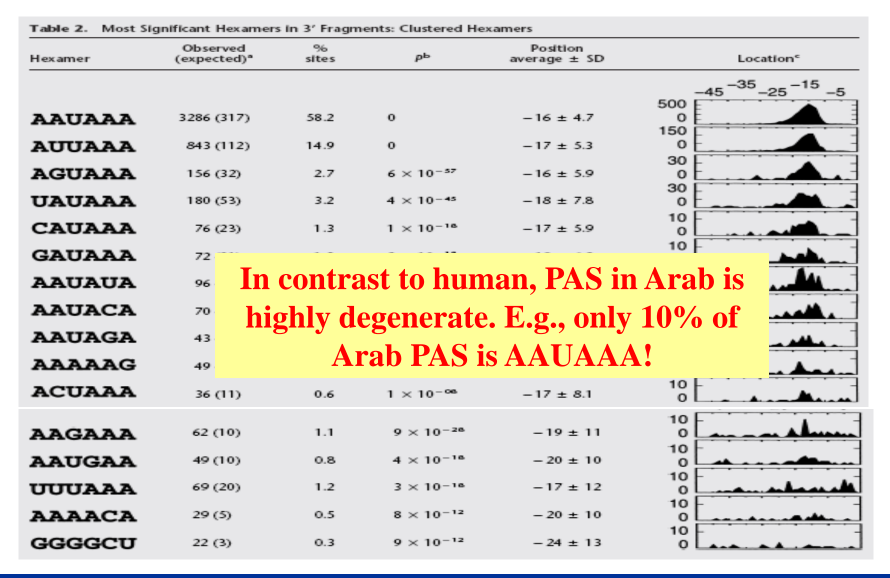
Poly-A signals in human (Gautheret et al., 200) National University of Singapore



Hexamer	Observed (expected) ^a	% sites	РР	Position average ± SD	Location ^c
					-45 ⁻³⁵ -25 ⁻¹⁵ -5
					500 F
AAUAA	3286 (317)	58.2	0	-16 ± 4.7	0 =
	042 (112)	14.0		17 . 53	150
AAAUUA	843 (112)	14.9	0	-17 ± 5.3	30
AGUAAA	156 (32)	2.7	6×10^{-57}	-16 ± 5.9	ő E
UAUAAA	180 (53)	3.2	4 × 10-45	-18 ± 7.8	30
UAUAAA	100 (33)	3.2	4 × 10 -	-10 ± 7.0	10
CAUAAA	76 (23)	1.3	1×10^{-18}	-17 ± 5.9	0
GAUAAA	72 (21)	1.3	2×10^{-16}	-18 ± 6.9	10
	, 2 (21)			-10 2 0.5	10 -
AAUAUA	96 (33)	1.7	2×10^{-19}	-18 ± 6.9	0
AAUACA	70 (16)	1.2	5 × 10 ⁻²³	-18 ± 8.7	10
					10 -
AAUAGA	43 (14)	0.7	1 × 10-9	-18 ± 6.3	0
AAAAAG	49 (11)	0.8	5×10^{-17}	-18 ± 8.9	10
ACUAAA	26.613	0.6	1 × 10-98	17 . 01	10
ACUAAA	36 (11)	0.6	1 × 10 ^{-∞}	-17 ± 8.1	0
	62 (I.O.)		0 10 - 28	10 . 11	10
AAGAAA	62 (10)	1.1	9 × 10 ⁻²⁶	-19 ± 11	10 -
AAUGAA	49 (10)	8.0	4×10^{-16}	-20 ± 10	0
AAAUUU	69 (20)	1.2	3 × 10 ⁻¹⁶	-17 ± 12	10
UUUAAA	05 (20)	1.2	3 ~ 10	-17 - 12	10 F
AAAACA	29 (5)	0.5	8×10^{-12}	-20 ± 10	0
GGGGCU	22 (3)	0.3	9×10^{-12}	- 24 ± 13	10

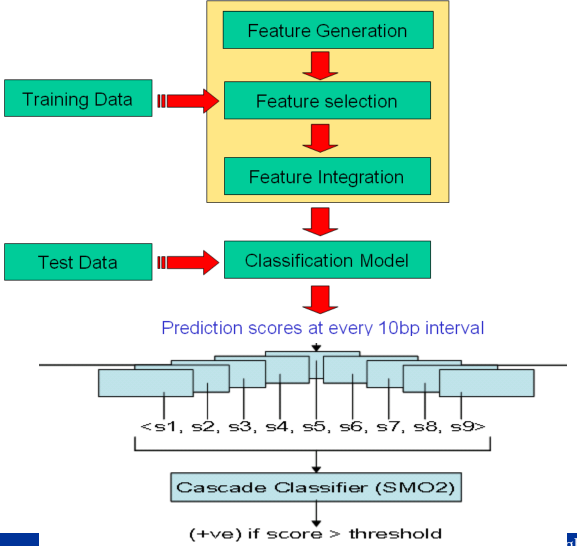
Poly-A signals in Arabidopsis





Approach on Arab PAS sites (I)







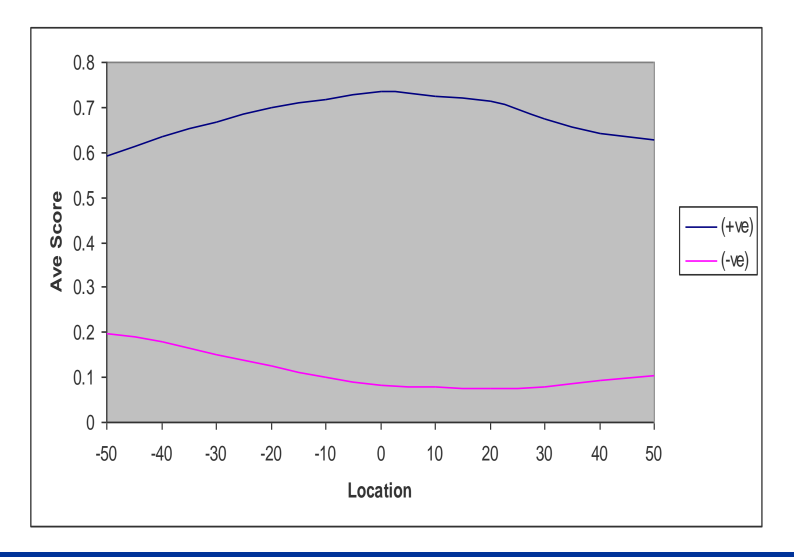


- Data collection
 - #1 from Hao Han, 811+ve seq (-200/+200)
 - #2 from Hao Han, 9742-ve seq (-200/+200)
 - #3 from Qingshun Li,
 - 6209 (+ve) seq (-300/+100)
 - 1581 (-ve) intron (-300/+100)
 - 1501 (-ve) coding (-300/+100)
 - 864 (-ve) 5'utr (-300/+100)

Feature generation

- 3-grams, compositional features (4U/1N. G/U*7, etc)
- Freq of features above in 3 diff windows: (-110/+5), (-35/+15), (-50/+30)
- Feature selection
 - $-\chi 2$
- Feature integration & Cascade
 - SVM

Score profile relative to candidate sites





Validation results

SN_0	SMO 1		SM	10 2	PASS 1.0	
Control	SN & SP	Threshold	SN & SP	Threshold	SN & SP	Threshold
Sequences						
CDS	90%	0.26	94%	0.24	95%	3.7
5'UTR	79%	0.42	85%	0.49	78%	5.5
Intron	64%	0.59	71%	0.67	63%	6.3

Table 2. Equal-error-rate points of SMO1, SMO2, and PASS 1.0 for SN_10.

SN_10	SMO 1		SM	10 2	PASS 1.0	
Control	SN & SP	Threshold	SN & SP	Threshold	SN & SP	Threshold
Sequences						
CDS	94%	0.36	96%	0.31	96%	4
5'UTR	86%	0.53	89%	0.6	81%	5.7
Intron	73%	0.68	77%	0.77	67%	6.6

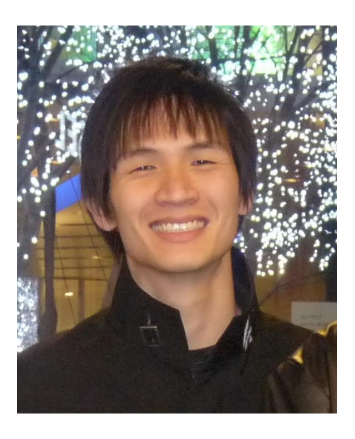
Table 3. Equal-error-rate points of SMO1, SMO2, and PASS 1.0 for SN_30.

SN_30	SMO 1		SMO 2		PASS 1.0	
Control	SN & SP	Threshold	SN & SP	Threshold	SN & SP	Threshold
Sequences						
CDS	97%	0.44	97%	0.37	97%	4.3
5'UTR	90%	0.62	92%	0.67	84%	6.2
Intron	79%	0.75	83%	0.81	72%	6.8

About the inventor: Koh Chuan Hod

Koh Chuan Hock

- BComp (CB), NUS,2008
- PhD, NUS, 2012
- Currently DataScientist at IndeedInc, Japan



Concluding Remarks...





What have we learned?

- Gene feature recognition applications
 - TIS, TSS, PAS
- General methodology
 - "Feature generation, feature selection, feature integration"
- Important tactics
 - Multiple models to optimize overall performance
 - Feature transformation (DNA → amino acid)
 - Classifier cascades

Any Question?





Acknowledgements

 The slides for PAS site prediction are adapted from slides given to me by Koh Chuan Hock



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