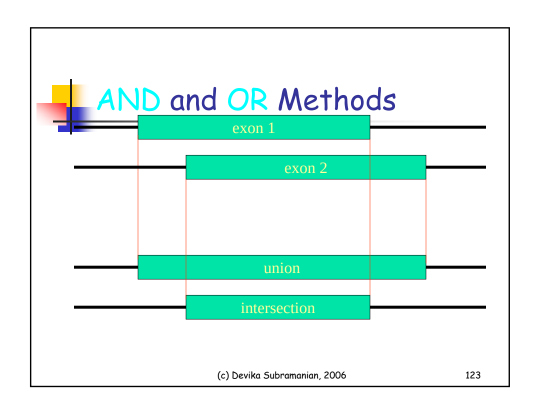


#### Integrated approaches for gene finding

- Programs that integrate results of similarity searches with ab initio techniques (GenomeScan, FGENESH+, Procrustes)
- Programs that use synteny between organisms (ROSETTA, SLAM)
- Integration of programs predicting different elements of a gene (EuGène)
- Combining predictions from several gene finding programs (combination of experts)

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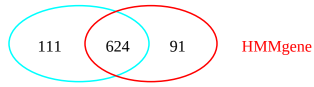




# Combining Genscan and HMMgene

 High prediction accuracy as well as reliability of their exon probability make them good candidates.

Genscan



 Genscan predicted 77% of exons correctly, HMMgene 75%, both 87%

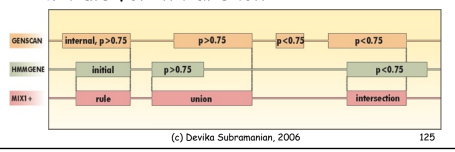
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#### **EUI** Method

(exon union - intersection)

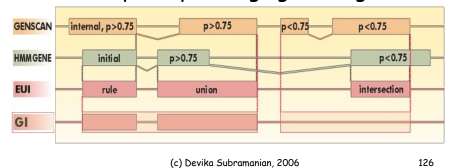
- Union of exons with p≥0.75
- 2. Intersection of exons with p<0.75
- 3 Rule for initial exon





#### Gene intersection (GI) method

- 1. Intersection of genes
- Apply EUI method to exons completely belonging to GI genes





## EUI with reading frame consistency

- Assign probabilities to GI genes.
   Determine position of acceptor and donor site in a reading frame.
- 2. GI gene with higher probability imposes the reading frame. Choose only EUI exons contained in GI genes that are in a chosen reading frame.

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## Results - Burset/Guigó dataset

METHODS RO	#no	Nucleotide accuracy			Exon accuracy				
	prediction	Su	Sp	AC	ESn	ESp	(ESn+Esp)/	ME	WE
Genscan	8	0.94	0.93	0.92	0.78	0.81	0.80	0.09 (203)	0.05 (188)
HMMgene	38	0.93	0.94	0.92	0.81	0.83	0.82	0.14 (308)	0.04 (139)
EUI	20	0.94	0.96	0.93	0.83	0.88	0.85	0.12 (250)	<b>0.03</b> (98)
GI	43	0.91	0.97	0.93	0.82	0.90	0.86	0.18 (386)	<b>0.02</b> (67)
EUI_frame	27	0.93	0.96	0.93	0.83	0.88	0.85	0.13 (286)	0.03 (87)

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### Summary: Eukaryotic gene finding

- Overall accuracy usually below 50%
  - Human gene finding is hardest
  - Very long introns, and lots of them
- Leading methods: HMMs and variants
- New ideas needed
- New opportunity: use sequence of related species

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#### Comparison of 1196 orthologous genes

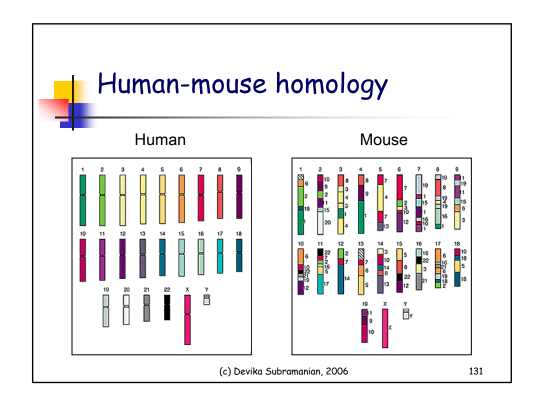
Sequence identity between genes in human/mouse

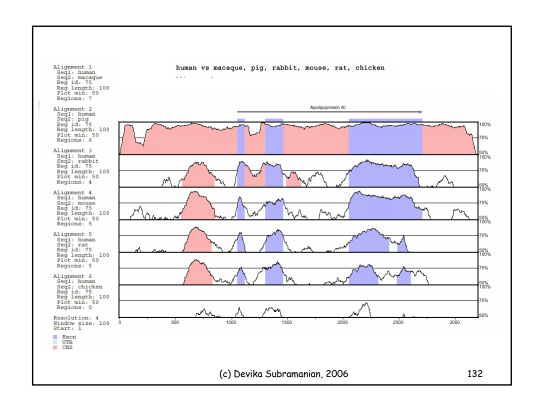
exons: 84.6%protein: 85.4%introns: 35%5' UTRs: 67%

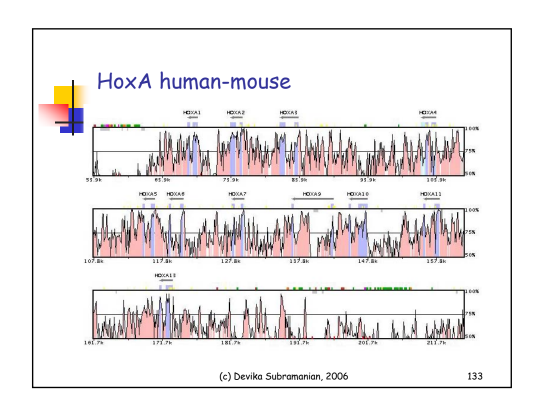
- 3' UTRs: 69%

• 27 proteins were 100% identical.

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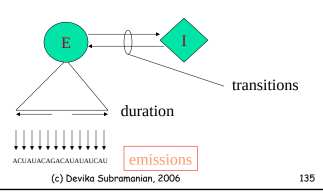
### Alignment

50	
247	GGTGAGGTCGAGGACCCTGCA CGGAGCTGTATGGAGGGCA AGAGC
	:       :    :        ::
368	GAGTCGGGGGAGGGGCTGCTGTTGGCTCTGGACAGCTTGCATTGAGAGG
100	
292	TTC CTACAGAAAAGTCCCAGCAAGGAGCCACACTTCACTG
	::  :   :  :  :
418	TTCTGGCTACGCTCTCCCTTAGGGACTGAGCAGAGGGCT CAGGTCGCGG
150	
332	ATGTCGAGGGGAAGACATCATTCGGGATGTCAGTG
467	
-0,	
200	
367	TTCAACCTCAGCAATGCCATCATGGGCAGCGGCATCCTGGGACTCGCCTA
517	TTCAATCTCAGCAACGCCATCATGGGCAGTGGAATTCTGGGGCTCGCCTA
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### Twinscan

 Twinscan is an augmented version of the Genscan HMM.





#### Twinscan Algorithm

- Align the two sequences (e.g. from human and mouse)
- 2. Mark each human base as gap ( ), mismatch ( : ), match ( | )

New "alphabet":  $4 \times 3 = 12$  letters

 $\Sigma = \{ A-, A:, A|, C-, C:, C|, G-, G:, G|, U-, U:, U| \}$ 

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#### Twinscan Algorithm

3. Run Viterbi using emissions  $e_j(k)$  where  $k \in \{A-, A:, A|, ..., T|\}$ 

Emission distributions  $e_j(k)$  estimated from real genes from human/mouse

 $e_{I}(x|) < e_{E}(x|)$ : matches favored in exons  $e_{I}(x-) > e_{E}(x-)$ : gaps (and mismatches) favored in introns

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### Example

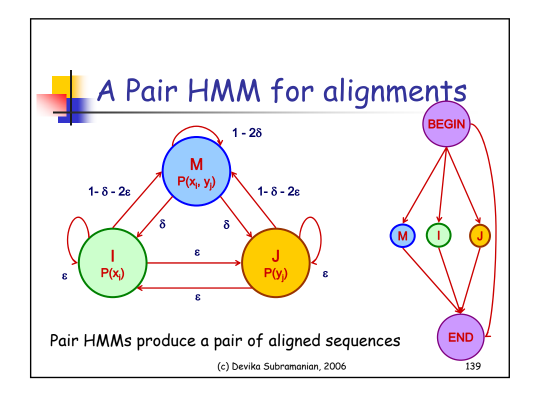
Human: ACGGCGACUGUGCACGU
Mouse: ACUGUGAC GUGCACUU
Alignment: ||:|:||-||||:|

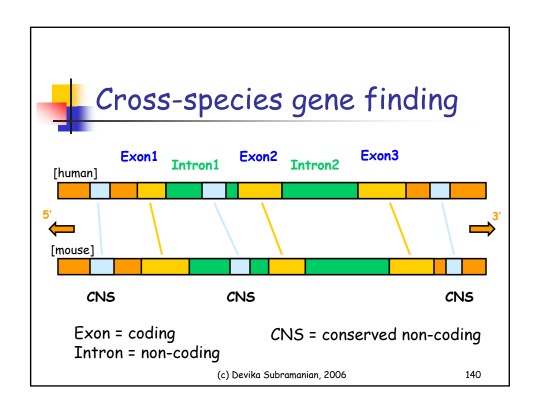
#### Input to Twinscan HMM:

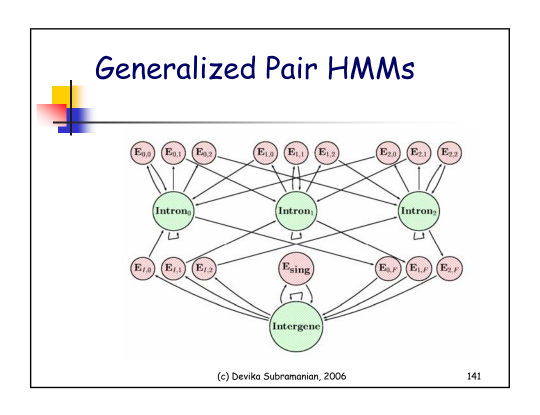
A C G: G C: G A C U- G U G C A C G: U

Recall, 
$$e_E(A|) > e_I(A|)$$
  
 $e_E(A-) < e_I(A-)$ 

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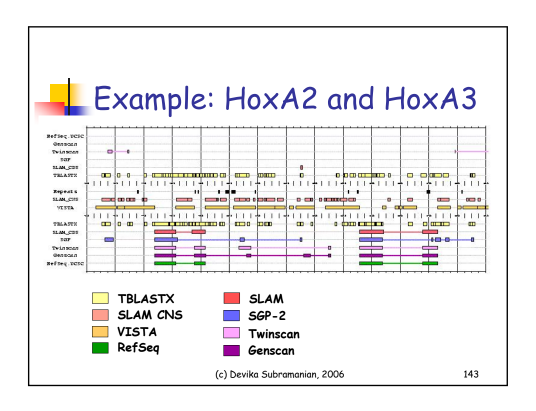




#### Ingredients in exon scores

- Splice site detection (VLMM)
- Length distribution (generalized)
- Coding potential (codon freq. tables)
- Isochore group

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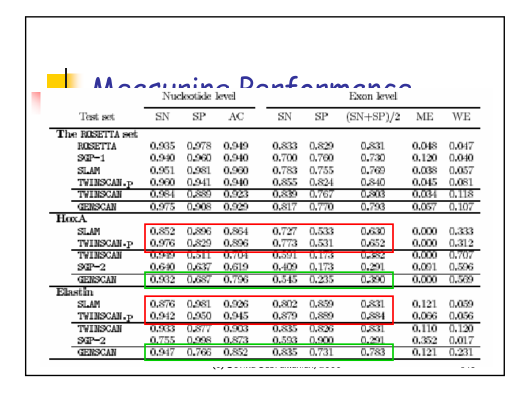




# What have we learned from comparative gene finding?

- conservation is a stronger splice site indicator than consensus
- intron lengths have diverged
- gene structure conservation is more powerful than sequence conservation for prediction
- · consensus for GC splice sites

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#### Priority organisms

#### Human-mouse gene finding not very high-impact

- lots of ancillary data gives better evidence
- most genes now known
- nonetheless, this problem is getting all the attention

#### Countless other species really need gene finders:

- Brugia malayi (causes lymphatic filariasis)
- Toxoplasma gondii
- Schistosoma mansoni (Schistosomiasis)
- Entamoeba histolytica (50 million cases/year)
- Tetrahymena thermophila (model organism)
- Plants: potato, maize, sorghum
- Mammals: chimp, dog, cow, pig

From the TIGR web site.
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### Genome scale gene finding

Strategy	Based on	Examples	
Ab initio prediction	Models of gene structure/comp	Genscan, GRAIL GenLang, hmmgene	
Microarray	Hybridization	Exon-scanning array	
Gene inference	Homology	GenomeScan	
Genomic:genomic alignment	Homology	ExoFish GLASS/Rosetta	
DNA:protein alignment	Homology	GeneWise	
cDNA sequencing	Sequencing	RIKEN	

C. Burge Nature Genet. 27, 5-7, 2001

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