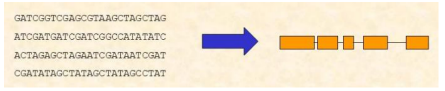
### SESSION 13. FINDING GENES

# In the world of snurps The Snurp



### Methods of gene prediction

- Computational gene prediction
  - De novo (or ab initio) approaches
    - Use statistical signals in DNA sequences that are characteristics of protein-coding genes
  - Homology-based approaches
    - Aligning to known mRNA or protein sequences or even HMMs to a genomic sequence
    - Require that known mRNAs and protein sequence information is available



cDNA sequencing(short or long)  $\rightarrow$  full-length gene

### Signals for de novo prediction

### □ Signals for de novo prediction

- Words (nucleotide k-mer) frequencies from coding sequences (Bishop 1994) + Start/Stop codons
- Exon-Intron boundary in Eukaryotes
- Length of exons and introns (not random)
- Polyadenylation signals (AATAAA)

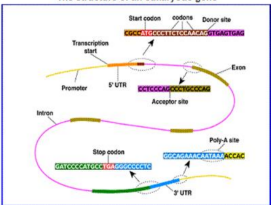


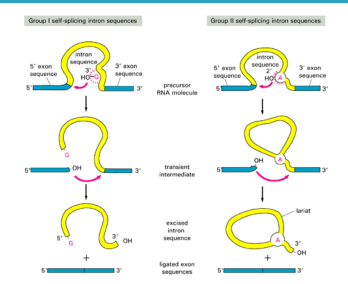


Figure 3. The structure of an eukaryotic gene (source: unknown).

### The splicing machinary

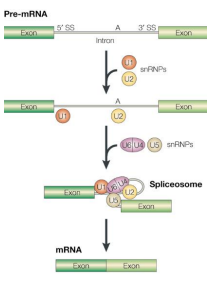
### □ Splicing

- Two sequential trans-esterification reactions by spliceosomes
- First, a branch site adenosine attacks the 5' splice site (donor site)
- Second, the free 3' end of the first exon attacks the 3' splice site (acceptor site)
- Then, two exons become covalently joined and the intron is released in the form of a lariat structure

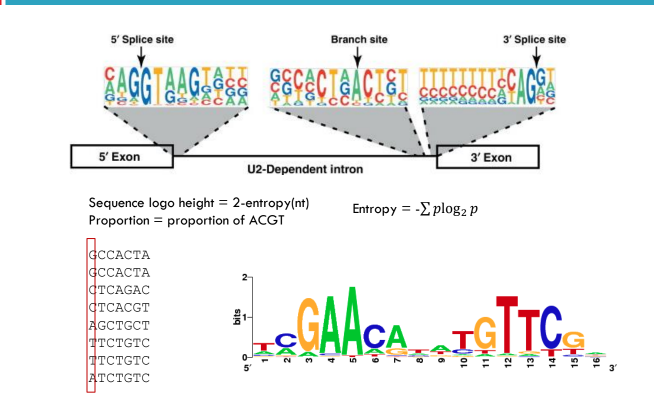


### The spliceosomes

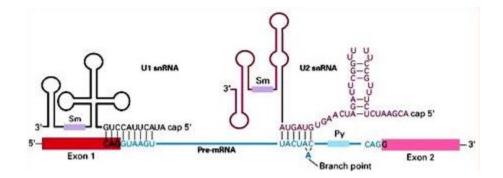
- □ Spliceosomes (RNA+protein complexes)
  - □ Includes a number of snRNPs (pronounced snurps)
  - **U**1, U2, U4, U5, and U6 snRNPs
  - U1 interacts with 5' splice site and U2 interacts with the branch site



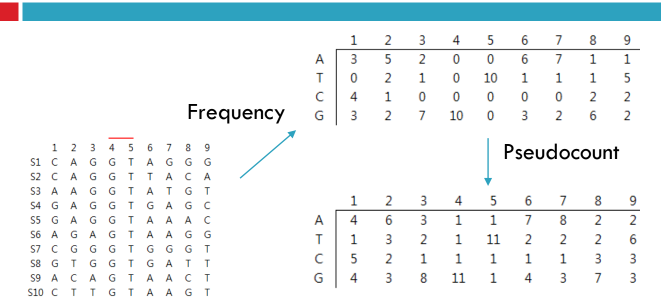
## Exon-junction motif analysis reveals splice signals (GU-AG + branch site)



## Exon-junction signals can be defined by U1 and U2 base pairing



### Splice site PSSM



### Splice site PSSM

	Probability									
	1	2	3	4	5	6	7	8	9	
А	0.29	0.43	0.21	0.07	0.07	0.50	0.57	0.14	0.14	
т	0.07	0.21	0.14	0.07	0.79	0.14	0.14	0.14	0.43	
С	0.36	0.14	0.07	0.07	0.07	0.07	0.07	0.21	0.21	
G	0.29	0.21	0.57	0.79	0.07	0.29	0.21	0.50	0.21	
Log-odd : log (Fij/Fexp)										
	1	2	3	4	5	6	7	8	9	TCCTGTCCCAGGTAGGGAA
А	0.13	0.54	-0.15	-1.25	-1.25	0.69	0.83	-0.56	-0.56	
Т	-1.25	-0.15	-0.56	-1.25	1.15	-0.56	-0.56	-0.56	0.54	
С	0.36	-0.56	-1.25	-1.25	-1.25	-1.25	-1.25	-0.15	-0.15	Fexp = .25
G	0.13	-0.15	0.83	1.15		0.13	-0.15	0.69	-0.15	Texp = .25
	1	2	3	4	5 5	)r 6	7	8	9	
А	0.06	0.46	-0.23	-1.33	-1.33	0.62	0.75	-0.64	-0.64	
Т	-1.20	-0.11	-0.51	-1.20	1.19	-0.51	-0.51	-0.51	0.59	
С	0.92	0.00	-0.69	-0.69	-0.69	-0.69	-0.69	0.41	0.41	
G	-0.20	-0.49	0.49	0.81	-1.59	-0.20	-0.49	0.36	-0.49	Fexp = Pi / Total P, i={A, C, G, T}

### make\_matrix.py

#### #!/usr/bin/python

```
import math, sys
splicefile = sys.argv[1]
number of sequences = 0 # we want to count the number of
                         # sequences in the file splice5.txt
for line in open(splicefile):
    line = line.rstrip()
    if number of sequences == 0:
        msa matrix = [[] * 9]
            # two dimensional array to
            # to store multiple alignment
            # create first empty row
        print msa matrix
    if number of sequences > 0:
        msa matrix.append([]) # add one row
    for j in range(0, 9): # fill the row with numbers
        msa matrix[number_of_sequences].append(line[j])
        if number of sequences==0: print msa matrix
    number_of_sequences += 1
```

```
GTGGTGAGT
                                                  ACAGTAAGT
                                                  CTTGTAAGT
                                                  AAGGTACTT
                                                  GAGGTAAGC
                                                  CAGGTTTGT
                                                  AAGGTAGGC
                                                  GAGGTGAGC
                                                  ACTGTACGT
                                                  AAGGTACCA
                                                  CAGGTCAGT
                                                   TCCGTGAGI
                                                  AAGGTAAAA
                                                  ACAGTAAGT
                                                  GAGGTATTT
                                                  AAGGTCAGT
                                                  CAGGTAATG
                                                  CAGGTGAGT
                                                  CAGGTACAG
                                                  TAAGTACGT
                                                  GAGGTAAGG
# produce count matrix
                                                  ATCGTAAGT
bases = ['A', 'T', 'C', 'G']
                                                  AAAGTAAGT
                                                  AAGGTGCGG
pssm = [[] * 9]
                                                  CAGGTAAGG
                                                  GAGGTAAGG
for i in range(0, 4):
                                                  AAGGTCAGT
                                                  CAGGTAATG
    if i > 0:
                                                  AAAGTAAGT
         pssm.append([])
                                                  CAGGTAAAA
    for j in range(0, 9):
                                                  TGGGTGAGT
         # add pseudocount = 1 to each of the AAGGTACGG
                                                  ACGGTGAGC
         pssm[i].append(1.0)
                                                  CAGGTGGGG
         # add counts to the pssm matrix
                                                  CIGGIGGGI
                                                  CAGGTGAGG
         for k in range(0, number_of_sequenc
GAGGIGGGG
              if msa matrix[k][j] == bases[i] CAGGTAAGG
                                                  TGGGTGAGT
                  pssm[i][i] += 1
                                                  GGAGTAAGT
                                                  AGGGTGAGG
# from count matrix produce PSSM by
                                                  ATGGTGAGT
                                                  "splice5.txt" 15169L, 151690C
# calculating the log odds values
for i in range(0, 4):
    for j in range(0, 9):
         pssm[i][j] = math.log(pssm[i][j] / (number of sequences + 4)
                                   * 4) / math.log(2)
         print pssm[i][j], # print PSSM
    orint "
```

GAGGTAAAC AAAGTAAGG CAGGTGGGT

python make matrix.py splice5.txt >matrix5.txt

1.417127811215887 1.27772888260454 -1.41348531150622 -11.8892187424726 -11.8892187424726 1.08160615889 1.49960579555099 -1.79910632280832 -0.674899621671848 -1.10703954864141 -0.810400792510635 -1.77547657642343 -11.8892187424726 1.99971472266178 -3.30050410689035 -1.39236495508457 -2.07063656499176 0.88289599579502 0.561992369359715 -0.932842585222944 -2.92632273713535 -11.8892187424726 -1.8892187424726 -3.18185961039173 -1.69815952794096 -2.15789971144755 -0.59085335738372 -0.465577547396757 -1.0547476924839 1.67826205867043 1.99971472266178 -11.8892187424726 0.740820810183483 -1.05159060930121 1.70043240404217 -0.209738643254714

#### score.py

```
863
                                                                                                            0.0421877149233
                                                                                                    864
                                                                                                            3.47543238438e-08
#!/usr/bin/python
                                                                                                    865
                                                                                                            8.90678251324e-10
import re, sys
                                                         print 'pos\tscore' # print header
                                                                                                    866
                                                                                                            7.87686655541e-11
# input file
                                                                                                    867
                                                                                                            1.69153038928
                                                          seq = seq.upper()
pssmfile = sys.argv[1]
                                                                                                    868
                                                                                                            0.000283434519102
                                                         bases = ['A', 'T', 'C', 'G']
# read matrix
                                                                                                    869
                                                                                                            0.000139474591032
i = 0
                                                                                                    870
                                                                                                            1.67929615407e-09
                                                         # score with the matrix
for line in open(pssmfile):
                                                                                                    871
                                                                                                            6.57510980056e-05
                                                         for k in range(0, len(seq) - 9):
    line = line.rstrip()
                                                                                                    872
                                                                                                            0.0018565833872
    if i == 0:
                                                              test = seq[k:k + 9]
                                                                                                    873
                                                                                                            1.16798994267e-09
        pssm = [[] * 9]
                                                              score = 0
                                                                                                    874
                                                                                                            3.63513668515e-07
    if i > 0:
                                                              for j in range(0, 9):
                                                                                                    875
                                                                                                            4.57861247817e-05
        pssm.append([]) # add one row
                                                                  base = test[j]
                                                                                                    876
                                                                                                            0.000274541015957
    col = re.split(' ', line)
                                                                  for b in range(0, 4):
                                                                                                    877
                                                                                                            1.2063875477e-09
    for j in range(0, 9): # fill the row with numbers
                                                                       if bases[b] == base:
                                                                                                    878
                                                                                                            0.000257659537677
        pssm[i].append(float(col[j]))
                                                                                                    879
                                                                                                            6.30368794006e-08
                                                                           score += pssm[b][j]
    i += 1
                                                                                                    880
                                                                                                            5.56689797646e-10
                                                                                    # convert log2<sup>881</sup>
                                                                                                            4.48265957774e-11
                                                              score = 2 ** score
# read sequence to be analyzed
                                                                                                    882
                                                                                                            0.0692197631
                                                                                         is the ex
seg = ''
                                                                                                    883
                                                                                                            1.46713237187e-08
                                                              pos = k + 3 # We want to print a
for line in open('amyloid.fa'):
                                                                                                    884
                                                                                                            1.03850108703e-10
                                                                            # next to the exon-in 885
                                                                                                            1.32711032479e-05
    if not re.search('>', line):
                                                                             # junction
                                                                                                    886
                                                                                                            3.39948840564e-07
        line = line.rstrip()
                                                              print pos. '\t', score
                                                                                                    887
                                                                                                            0.000426052487362
        seg += line
                                                                                                    888
                                                                                                            2.35157804612e-09
                                                                                                    889
                                                                                                            2.02971238834
```

862

9.10699121732e-05

### Visualization with R

#### plot the results of splice site prediction

#define some colours
rgb <- c("#009E73", "#D55E00", "#0072B2")</pre>

```
# make two graphs on top of each other
par(mfrow = c(2, 1))
```

# First consider the 5' splice site prediction and read the # output from # the Perl code data <- read.table("score5.txt", sep = "\t", header = TRUE)</pre>

# for the plot, we need to know about the sequence length
seqlen <- max(data\$pos)</pre>

# for the plot we need to know about the maximum score max\_score <- max(data\$score)</pre>

```
# make a plot for the 5' splice site data
plot(0, type = "n", lwd = 2, xlim = c(0, seqlen),
ylim = c(0, max_score * 1.1), main = "Splice site scoring",
xlab = "Position", ylab = "Score")
```

```
#print a legend
legend(seqlen * 0.7, max_score, "5prime", col = rgb[2],
    lwd = 1)
```

```
# plot the splice site scores
for (i in (1:seqlen)) {
    lines(c/data$pos[i], data$pos[i]), c(0, data$score[i]), col = rgb[2],
    lw = 2)
```

```
# plot the location of exons (we do not know these from the
# prediction)
lines(c(268, 331), c(max score/2, max score/2), col = rgb[1],
    1w = 4)
lines(c(447, 1054), c(max score/2, max score/2), col = rgb[1],
    1w = 4)
# Now consider the 3' splice site and read the output from
# the Perl code
data <- read.table("score3.txt", sep = "\t", header = TRUE)</pre>
max score <- max(data$score)</pre>
plot(0, type = "n", lwd = 2, xlim = c(0, seqlen),
    ylim = c(0, max score * 1.1), xlab = "Position", ylab = "Score")
legend(seqlen * 0.7, max_score, "3prime", col = rgb[3],
    1wd = 1)
for (i in (1:seglen)) {
    lines(c(data$pos[i], data$pos[i]), c(0, data$score[i]), col = rgb[3],
        1w = 2
}
# plot the location of exons as in the previous graph
lines(c(268, 331), c(max score/2, max score/2), col = rgb[1],
    1w = 4)
lines(c(447, 1054), c(max score/2, max score/2), col = rgb[1],
    1w = 4
```

Splice site scoring

