Problem 5: LEARNING QUESTION

- Input: model structure M, Training Sequence S
- Output: Compute the parameters Θ
- Criteria: ML criterion
 - maximize $P(S | M, \Theta)$ HOW???

Problem 6: DESIGN QUESTION

- Input: Training Sequence S
- Output: Choose model structure M, and compute the parameters Θ
 - No reasonable solution
 - Standard models to pick from

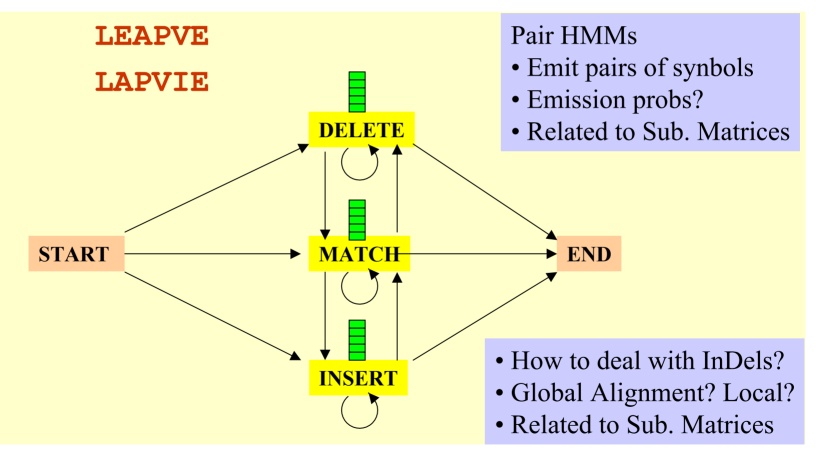
Iterative Solution to the LEARNING QUESTION (Problem 5)

- Pick initial values for parameters Θ_0
- <u>Repeat</u>

Run training set S on model M
Count # of times transition i ⇒ j is made
Count # of times letter x is emitted from state i
Update parameters Θ

• <u>Until</u>(some stopping condition)

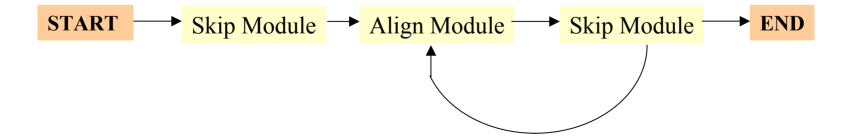
How to model Pairwise Sequence Alignment



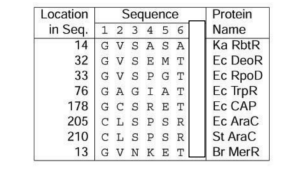
How to model Pairwise Local Alignments?

START → Skip Module — Align Module — Skip Module — END

How to model Pairwise Local Alignments with gaps?



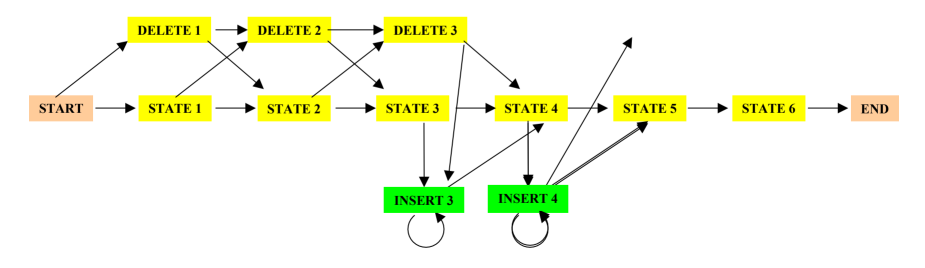
Profile HMMs



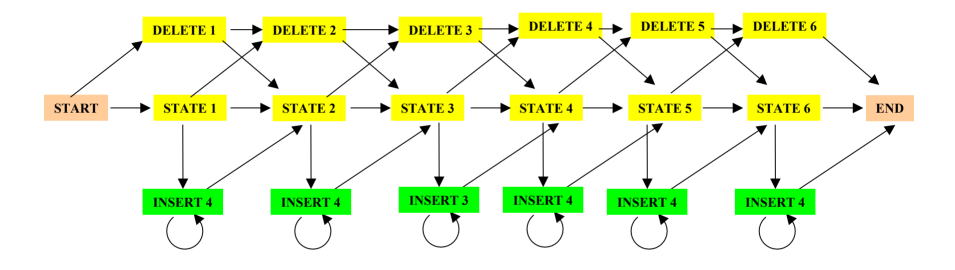


Profile HMMs with InDels

- Insertions
- Deletions
- Insertions & Deletions



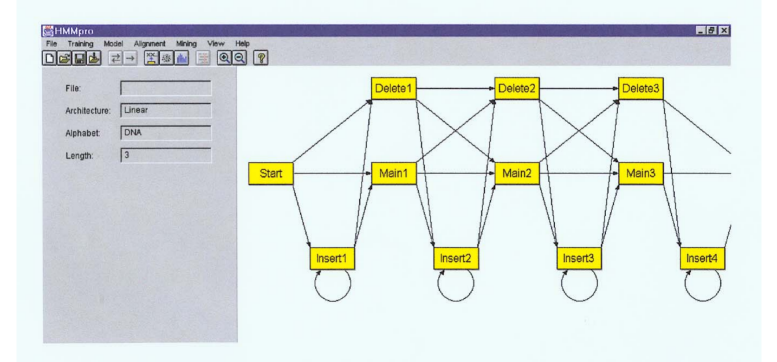
Profile HMMs with InDels



Missing transitions from DELETE j to INSERT j and from INSERT j to DELETE j+1.

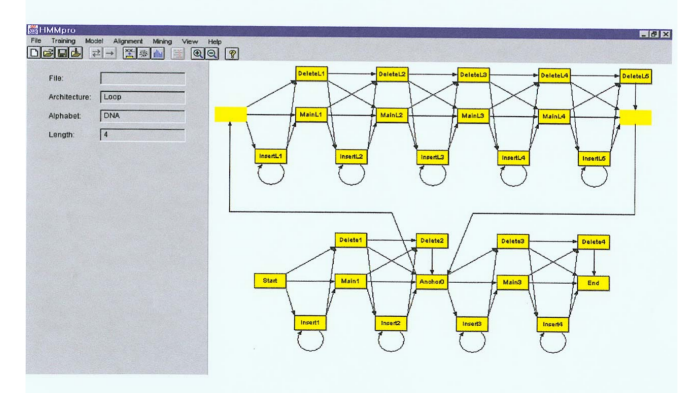
Standard HMM architectures

Linear Architecture



Standard HMM architectures

Loop Architecture



Standard HMM architectures

Wheel Architecture	
Contraction Image: Contract	Main10 Main10 Main2 Main3 Main3 Main3 Main4 Main4

Profile HMMs from Multiple Alignments

- HBA HUMAN VGA--HAGEY
- HBB HUMAN V----NVDEV
- MYG PHYCA VEA--DVAGH
- GLB3 CHITP VKG----D
- GLB5 PETMA VYS--TYETS
- LGB2 LUPLU FNA--NIPKH
- GLB1 GLYDI IAGADNGAGV

Construct Profile HMM from above multiple alignment.

Entropy

• Entropy measures the variability observed in given data.

$$E = -\sum_{c} p_{c} \log p_{c}$$

- Entropy is useful in multiple alignments & profiles.
- Entropy is max when uncertainty is max.

G-Protein Couple Receptors

- Transmembrane proteins with 7 α -helices and 6 loops; many subfamilies
- Highly variable: 200-1200 aa in length, some have only 20% identity.
- [Baldi & Chauvin, '94] HMM for GPCRs
- HMM constructed with 430 match states (avg length of sequences); Training: with 142 sequences, 12 iterations

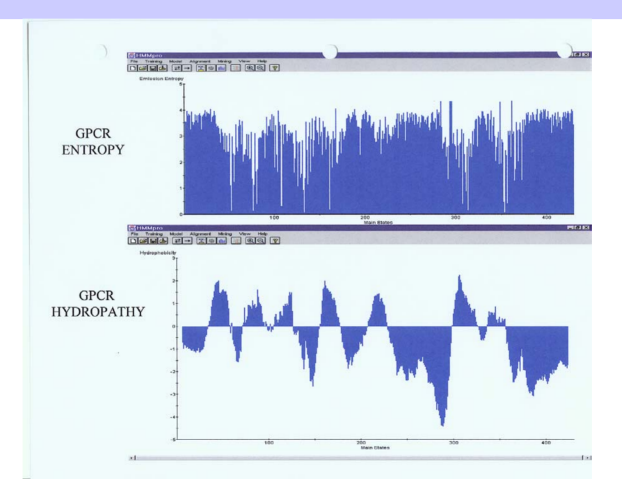
• Compute main state entropy values

$$H_i = -\sum_a e_{ia} \log e_{ia}$$

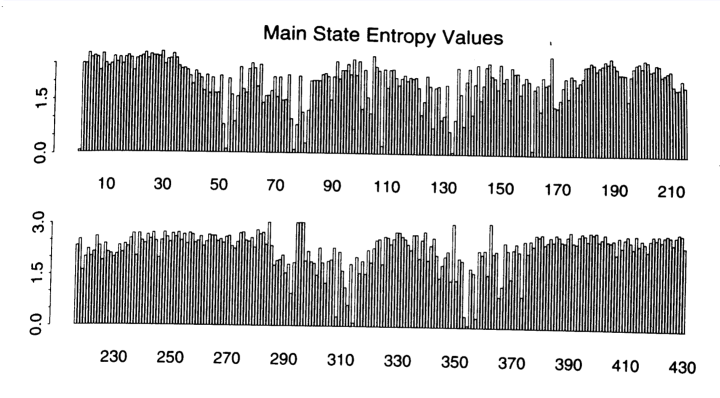
- For every sequence from test set (142) & random set (1600) & all SWISS-PROT proteins
 - Compute the negative log of probability of the most probable path π

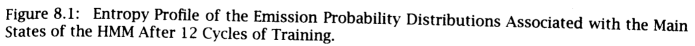
$$Score(S) = -\log(P(\pi \mid S, M))$$

GPCR Analysis



Entropy





GPCR Analysis (Cont'd)

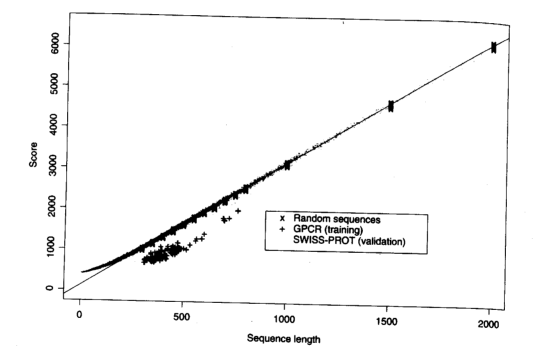


Figure 8.2: Scores (Negative Log-likelihoods of Optimal Viterbi Paths). Represented sequences consist of 142 GPCR training sequences, all sequences from the SWISS-PROT database of length less than or equal to 2000, and 220 randomly generated sequences with same average composition as the GPCRs of length 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800 (20 at each length). The regression line was obtained from the 220 random sequences. The horizonta³ distances in the histogram correspond to (malized scores (6).

Lecture 10

Applications of HMM for GPCR

- Bacteriorhodopsin
 - Transmembrane protein with 7 domains
 - But it is not a GPCR
 - Compute score and discover that it is close to the regression line. Hence not a GPCR.
- Thyrotropin receptor precursors
 - All have long initial loop on INSERT STATE 20.
 - Also clustering possible based on distance to regression line.

HMMs – Advantages

- Sound statistical foundations
- Efficient learning algorithms
- Consistent treatment for insert/delete penalties for alignments in the form of locally learnable probabilities
- Capable of handling inputs of variable length
- Can be built in a modular & hierarchical fashion; can be combined into libraries.
- Wide variety of applications: Multiple Alignment, Data mining & classification, Structural Analysis, Pattern discovery, Gene prediction.

HMMs – Disadvantages

- Large # of parameters.
- Cannot express dependencies & correlations between hidden states.