Hidden Markov Models Stochastic Regular Grammars

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Outline

- Single-sequence hidden Markov models
- Posterior probabilities for single-sequence HMMs
- Pair Hidden Markov models
- Evolutionary Hidden Markov models
- 5 Discriminative models and conditional random fields

Early motivation: isochores

- Long regions of uniform GC content (which is correlated with gene density, recombination frequency...)
 - e.g. Major Histocompatibility Complex (MHC) class II and class III sequences on human chromosome 6
 - Lengths 900.9 kb, 642.1 kb; GC-content 41%, 52%
- Gary Churchill: first Hidden Markov Model for isochore detection (1989)
- Earliest non-thermodynamic hit to "isochore" on PubMed is 1986, Alonso et al
- HMM analogy: occasionally dishonest casino (Durbin et al)



Notation

- Hidden Markov model: notation
 - Let x denote hidden state, y observed symbol. State space includes START and END
 - Let e(x, y) be probability of emitting character y in state x
 - Let t(i, j) be probability of transition to state j if currently in state i

- Idea of a particular partitioning as a "path" through the HMM
 - Let y_n be observed nucleotide at position n and let x_n be hidden state of position n
 - Let L be length of sequence
 - For convenience, set $x_0 = START$ and $x_{L+1} = END$
 - Let $Y = \{y_1 \dots y_L\}$ represent entire observed sequence, $X = \{x_0 \dots x_{L+1}\}$ hidden state sequence
 - Each $X = \{x_k\}$ represents a "path" (sketch); there are $\sim K^L$ possible paths for K states
 - Joint likelihood is

$$P(X, Y) = t(x_0, x_1) \prod_{k=1}^{L} e(x_k, y_k) t(x_k, x_{k+1})$$

- Note that $P(X, Y) \equiv P(X, Y|x_0)$. Conditioning on x_0 is assumed throughout
- Two questions, two algorithms. (i) Viterbi: what X maximises P(X, Y)? (ii) Forward: what is P(Y)?



Viterbi algorithm for finding ML hidden state path

$$\begin{array}{lll} \max_{X} P(X,Y) & = & \max_{X_{L}} \max_{X_{L-1}} \ldots \max_{X_{1}} t(x_{0},x_{1}) \prod_{k=1}^{L} e(x_{k},y_{k}) t(x_{k},x_{k+1}) \\ & = & \max_{X_{L}} t(x_{L},x_{L+1}) e(x_{L},y_{L}) \max_{X_{L-1}} t(x_{L-1},x_{L}) e(x_{L-1},y_{L-1}) \ldots \max_{X_{1}} t(x_{1},x_{2}) e(x_{1},y_{1}) t(x_{0},x_{1}) \\ & = & \max_{X_{L}} t(x_{L},\operatorname{END}) V_{L}(x_{L}) \end{array}$$

Summary

where

$$V_n(x_n) = \begin{cases} e(x_n, y_n) \max_{x_{n-1}} t(x_{n-1}, x_n) V_{n-1}(x_{n-1}) & \text{if } n > 0 \\ 1 & \text{if } n = 0 \end{cases}$$

Note that

$$V_n(x_n) = \max_{x_1 \dots x_{n-1}} P(x_1 \dots x_n, y_1 \dots y_n | x_0)$$

or, in words, $V_n(x)$ is the maximum likelihood of any path ending in state x and emitting symbols $y_1 \ldots y_n$. The ML state sequence, $\operatorname{argmax}_X P(X, Y)$, is recovered by **traceback** from V_L to V_1 .



Single-sequence hidden Markov models Posterior probabilities for single-sequence HMMs Pair Hidden Markov models Evolutionary Hidden Markov models

Discriminative models and conditional random fields Summary

Forward algorithm for summing over all possible hidden state paths

$$\begin{split} P(Y) &= \sum_{X} P(X,Y) \\ &= \sum_{x_{L}} \sum_{x_{L-1}} \dots \sum_{x_{1}} t(x_{0},x_{1}) \prod_{k=1}^{L} e(x_{k},y_{k}) t(x_{k},x_{k+1}) \\ &= \sum_{x_{L}} t(x_{L},x_{L+1}) e(x_{L},y_{L}) \sum_{x_{L}-1} t(x_{L-1},x_{L}) e(x_{L-1},y_{L-1}) \dots \sum_{x_{1}} t(x_{1},x_{2}) e(x_{1},y_{1}) t(x_{0},x_{1}) \\ &= \sum_{x_{l}} t(x_{L}, \text{END}) F_{L}(x_{L}) \end{split}$$

where

$$F_n(x_n) = \begin{cases} e(x_n, y_n) \sum_{x_{n-1}} t(x_{n-1}, x_n) F_{n-1}(x_{n-1}) & \text{if } n > 0 \\ 1 & \text{if } n = 0 \end{cases}$$

Note that

$$F_n(x_n) = P(x_n, y_1 \dots y_n | x_0) = \sum_{x_1 \dots x_{n-1}} P(x_1 \dots x_n, y_1 \dots y_n | x_0)$$

or, in words, $F_n(x)$ is the sum of likelihoods of all paths ending in state x and emitting symbols $y_1 \dots y_n$. By analogy to Viterbi traceback, a ML state sequence can be sampled by stochastic traceback from F_L to F_1 .

Implementation issues

- probability underflow for long sequences
- slow floating-point multiply
- possible solutions:
 - (possibly discretized) log-space scores
 - bignum types

Definition of the posterior probability that position *n* is in state *k*: sum over paths

$$P(x_n = k|Y) = \frac{\sum_X P(X, Y)\delta(x_n = k)}{P(Y)}$$

Splitting the path into three parts: < n, = n and > n

$$P(x_{n}|Y) = \sum_{x_{1}...x_{n-1}} \sum_{x_{n+1}...x_{L}} \frac{P(x_{1}...x_{n}, y_{1}...y_{n}|x_{0}) P(x_{n+1}...x_{L+1}, y_{n+1}...y_{L}|x_{n})}{P(Y)}$$

$$= \frac{F_{n}(x_{n}) B_{n}(x_{n})}{P(Y)}$$

$$B_{n}(x_{n}) = P(x_{L+1}, y_{n+1}...y_{L}|x_{n})$$

$$= \sum_{x_{n+1}...x_{L}} P(x_{n+1}...x_{L+1}, y_{n+1}...y_{L}|x_{n})$$

Likewise,

$$P(x_n, x_{n+1}|Y) = \frac{F_n(x_n)t(x_n, x_{n+1})e(x_{n+1}, y_{n+1})B_{n+1}(x_{n+1})}{P(Y)}$$

and (useful for compression)

$$P(y_{n+1}=k|y_1...y_n)=\frac{\sum_i\sum_jF_n(i)t(i,j)e(j,k)}{\sum_iF_n(i)}$$

The Backward algorithm

$$B_n(x_n) = \begin{cases} \sum_{x_{n+1}} t(x_n, x_{n+1}) e(x_{n+1}, y_{n+1}) B_{n+1}(x_{n+1}) & \text{if } n < L \\ t(x_L, \text{END}) & \text{if } n = L \end{cases}$$

The Baum-Welch training algorithm:

$$P(X, Y) = \left(\prod_{i,j} t(i,j)^{u(i,j)}\right) \left(\prod_{i,k} e(i,k)^{f(i,k)}\right)$$

where u(i, j) is the number of transitions $i \to j$ and f(i, k) is the number of emissions of character k from state i. Sufficient statistics for EM algorithm are therefore

$$\hat{t}(i,j) = \sum_{n=0}^{L} P(x_n = i, x_{n+1} = j | Y)$$

$$\hat{\mathbf{e}}(i,k) = \sum_{n=1}^{L} P(x_n = i|Y)\delta(y_n = k)$$

where \hat{t} and \hat{e} are the posterior expectations of u and f. Dirichlet (mixture) priors can be used for t and e.

Note that
$$\hat{t}(i,j) = \frac{\partial \log P(X)}{\partial \log t(i,j)}$$
 and $\hat{e}(i,k) = \frac{\partial \log P(X)}{\partial \log e(i,k)}$

Implementation issues: log-space probability addition

$$\log(e^a + e^b) = \max(a, b) + \oplus(|a - b|)$$

 $\oplus(|a - b|) = \log(1 + e^{-|a - b|})$

If scores are discretized, can implement \oplus as a lookup table for speed.

Null states

- Convenience: reduce number of transitions (e.g. delete states of profile HMM)
- Need to do a topological sort of null states so that they're filled in the correct order
- Awkwardness: null cycles. These break the toposort

Can always eliminate null states as follows

$$\mathbf{t} = \mathbf{a} + \mathbf{b} + \mathbf{c} + \mathbf{d}$$

$$\mathbf{a} = \begin{pmatrix} * & 0 \\ 0 & 0 \end{pmatrix}, \mathbf{b} = \begin{pmatrix} 0 & * \\ 0 & 0 \end{pmatrix}, \mathbf{c} = \begin{pmatrix} 0 & 0 \\ * & 0 \end{pmatrix}, \mathbf{d} = \begin{pmatrix} 0 & 0 \\ 0 & * \end{pmatrix}$$

$$\mathbf{t}' = \begin{pmatrix} * & 0 \\ 0 & 0 \end{pmatrix} = \mathbf{a} + \sum_{n=0}^{\infty} \mathbf{b} \mathbf{d}^n \mathbf{c} = \mathbf{a} + \mathbf{b} (\mathbf{I} - \mathbf{d})^{-1} \mathbf{c}$$

where **a** (and **t**') contain emit \rightarrow emit transitions, **b** emit \rightarrow null, **c** null \rightarrow emit and **d** null \rightarrow null.

- Factor graph representation of HMM; similarity to pruning/peeling and parsimony (Forward-Backward C Sum-Product)
- General-purpose HMM implementations: DART library (C++)

- Motivation: pairwise sequence alignment, pairwise genefinding, etc.
- Let x denote hidden state, y character in sequence Y, z character in sequence Z
- Let $\Delta y(x)$ be 1 if state x emits a character to Y, and 0 otherwise; likewise $\Delta z(x) = 1$ iff x emits to Z

- Emission probability e(x, y, z) is defined as follows:
 - If $\Delta y(x) = 1$ and $\Delta z(x) = 0$, then x is called a **delete** state and $e(x, y, z) \equiv e_d(x, y)$ is a function of x and y only
 - If $\Delta y(x) = 0$ and $\Delta z(x) = 1$, then x is called an **insert** state and $e(x, y, z) \equiv e_i(x, z)$ is a function of x and z only
 - If $\Delta y(x) = 1$ and $\Delta z(x) = 1$, then x is called a **match** state and $e(x, y, z) \equiv e_m(x, y, z)$ is a function of x, y and z
 - If $\Delta y(x) = 0$ and $\Delta z(x) = 0$, then x is called a **null** state and e(x, y, z) is a function of x only (typically just 1)
 - We will assume for now that there are no null states (apart from START and END).

- As before, t(i, j) is the probability of transition to state j if currently in state i
- Suppose sequence lengths are K, L so observed data are $Y = \{y_1 \dots y_K\}$ and $Z = \{z_1 \dots z_L\}$
- Again we have a state path x₁, x₂...x_N and for convenience we set x₀ =START and x_{N+1} =END.
 - Denote by Λ_{kl} the event that there exists a *break* at (k, l):

$$\Lambda_{kl} \Rightarrow \exists n : \sum_{i=1}^{n} \Delta y(x_i) = k, \sum_{i=1}^{n} \Delta z(x_i) = l$$

So Λ_{kl} means that, at some point n on the state path, the model has emitted k symbols to Y and l symbols to Z.



Viterbi

$$V_{kl}(x_n) = \max_{x_1...x_{n-1}} P(\Lambda_{kl}, x_1...x_n, y_1...y_k, z_1...z_l|x_0)$$

Recursion (assuming no null states)

$$V_{kl}(x_n) = \begin{cases} e(x_n, y_k, z_l) \max_{x_{n-1}} t(x_{n-1}, x_n) \\ \times V_{k-\Delta y(x_n), l-\Delta z(x_n)}(x_{n-1}) & \text{if } k > 0 \text{ or } l > 0 \\ 1 & \text{if } k = l = 0 \text{ and } x_n = \text{START} \\ 0 & \text{if } k = l = 0 \text{ and } x_n \neq \text{START} \\ 0 & \text{if } k < 0 \text{ or } l < 0 \end{cases}$$

Forward

$$F_{kl}(x_n) = P(\Lambda_{kl}, x_n, y_1 \dots y_k, z_1 \dots z_l | x_0)$$

=
$$\sum_{x_1 \dots x_{n-1}} P(\Lambda_{kl}, x_1 \dots x_n, x_{N+1}, y_1 \dots y_k, z_1 \dots z_l | x_0)$$

Recursion (assuming no null states)

$$F_{kl}(x_n) = \begin{cases} e(x_n, y_k, z_l) \sum_{x_{n-1}} t(x_{n-1}, x_n) \\ \times F_{k-\Delta y(x_n), l-\Delta z(x_n)}(x_{n-1}) & \text{if } k > 0 \text{ or } l > 0 \\ 1 & \text{if } k = l = 0 \text{ and } x_n = \text{START} \\ 0 & \text{if } k = l = 0 \text{ and } x_n \neq \text{START} \\ 0 & \text{if } k < 0 \text{ or } l < 0 \end{cases}$$

Backward

$$B_{kl}(x_n) = P(\Lambda_{kl}, x_{N+1}, y_{k+1} \dots y_K, z_{l+1} \dots z_L | x_n)$$

$$= \sum_{x_{n+1} \dots x_N} P(\Lambda_{kl}, x_{n+1} \dots x_{N+1}, y_{k+1} \dots y_K, z_{l+1} \dots z_L | x_n)$$

Recursion (assuming no null states)

$$B_{kl}(x_n) = \begin{cases} \sum_{x_{n+1}} t(x_n, x_{n+1}) e(x_{n+1}, y_{k+1}, z_{l+1}) \\ \times B_{k+\Delta y(x_n+1), l+\Delta z(x_n+1)}(x_{n+1}) & \text{if } k < K \text{ or } l < L \\ t(x_n, \text{END}) & \text{if } k = K \text{ and } l = L \\ 0 & \text{if } k > K \text{ or } l > L \end{cases}$$

Evidence, posterior probabilities & EM counts

$$\begin{split} P(Y,Z) &= \sum_{x} F_{KL}(x) t(x, \text{END}) \\ P(\Lambda_{kl}, x_{n} | Y, Z) &= \frac{F_{kl}(x_{n}) B_{kl}(x_{n})}{P(Y)} \\ P(\Lambda_{kl}, x_{n}, x_{n+1} | Y) &= \frac{F_{kl}(x_{n}) t(x_{n}, x_{n+1}) e(x_{n+1}, y_{k+1}, z_{l+1}) B_{k+\Delta y(x_{n+1}), l+\Delta z(x_{n+1})}(x_{n+1})}{P(Y)} \\ \hat{t}(i, j) &= \sum_{k=0}^{K} \sum_{l=0}^{L} P(\Lambda_{kl}, x_{n} = i, x_{n+1} = j | Y, Z) \\ \hat{e}_{m}(x, y, z) &= \sum_{k: y_{k} = y} \sum_{l=0}^{L} P(\Lambda_{kl}, x_{n} = x | Y, Z) \\ \hat{e}_{d}(x, y) &= \sum_{k: y_{k} = y} \sum_{l=0}^{L} P(\Lambda_{kl}, x_{n} = x | Y, Z) \\ \hat{e}_{i}(x, z) &= \sum_{k=0}^{K} \sum_{l: z_{l} = z} P(\Lambda_{kl}, x_{n} = x | Y, Z) \end{split}$$

Decision theory ("optimal accuracy").

- Decision theory: maximise expected "reward", making use of the posterior distribution
- Overlap score: an objective function (i.e. reward) that compares predicted alignment α with true alignment α'
 - Overlap score is $|\alpha \cap \alpha'|$, where an alignment is viewed as a set of match co-ords $\alpha = \{(k_1, l_1), (k_2, l_2) \dots\}$
 - Several other good objective functions (e.g. "Cline shift score"); overlap is simpler, albeit less realistic
 - NB also $\delta(\alpha=\alpha')$ which only rewards perfect alignments, yielding a multiplicative, Viterbi-like recursion
 - Example criteria: how good is alignment for structure prediction? homology detection? benchmark of choice?
 - e.g. PROBCONS (Batzoglou et al) uses the sum-of-pairs score, same as the BAliBASE benchmark



Posterior expectation of overlap score for an alignment (NB only match states have Δy(x)Δz(x) ≠ 0)

$$A[\alpha] = \sum_{(k,l)\in\alpha} P(\mathsf{match}, k, l)$$

$$P(\mathsf{match}, k, l) = \sum_{x} \Delta y(x) \Delta z(x) P(\Lambda_{kl}, x_n = x)$$

• Optimal accuracy recursion: let α_{kl} be any alignment up to (k, l), so $k' \le k$ and $l' \le l$ for all $(k', l') \in \alpha$. Then

$$\begin{array}{ll} O_{kl} &=& \max_{\alpha_{kl}} A\left[\alpha_{kl}\right] \\ &=& \left\{ \begin{array}{ll} \max\left(\begin{array}{l} O_{k-1,l-1} + P(\mathsf{match},k,l), \\ O_{k,l-1}, \\ O_{k-1,l} \end{array} \right) & \text{if } k,l \geq 0 \\ 0 & \text{otherwise} \end{array} \right. \end{array}$$

• Optimal alignment α recovered by traceback from O_{KL} .

- Dynamic programming algorithms whose finite state automata are almost or exactly Pair HMMs
 - Needleman-Wunsch; Smith-Waterman; Gotoh; Altschul, Proteins 1998
 - General implementations: DART library (C++), Exonerate (C), HMMoC (Java/C++), ...

- Can readily extend the Pair HMM to a multi-sequence HMM for multiple sequence alignment
 - Arbitrary number N of output sequences $Y^{(1)}, Y^{(2)}, Y^{(3)} \dots Y^{(N)}$ of lengths $L_1 \dots L_N$ (see e.g. Holmes 2003)
 - Dynamic programming time/memory complexity is $O(\prod_n L_n)$ —not cheap
 - Ultimately, would like to structure $\Delta Y^{(n)}(x)$, t(x, x') and $e(x, y^{(1)} \dots y^{(N)})$ according to some underlying phylogenetic tree
 - The DP algorithms can also be tree structured, c.f. "progressive alignment"
 - For now, we ignore phylogenetic structure of indels (Δ,t) and concentrate on substitution model (e)



- Initial, simplistic, restrictive concept of Evolutionary HMM, lacking a good gap model:
 - A single-sequence HMM that emits phylogenetically-correlated multiple alignment columns, instead of single characters.
 - Follows e.g. Goldman, Thorne & Jones, 1996 ("Using evolutionary trees in protein secondary structure prediction...")
 - For now, gaps will be glossed over heuristically (disallowed/treated as wildcards/excessively gappy columns discarded/etc.)
- For a tree with N leaves, the emitted symbol alphabet is Ω^N where Ω is the single-character alphabet



- Now the emission function e(x, y) for $y \in \Omega^N$ is expensive (and unnecessary) to tabulate
 - We can implement it as an instance of pruning instead: DP within DP
 - Assume an underlying continuous-time discrete-state Markov chain with rate matrix $\mathbf{R}^{(x)}$ and initial distribution $\pi^{(x)}$
 - Let e(x, y) be probability of observing characters y at (leaf) nodes of a phylogenetic tree, with this process
 - Tree, like alignment, will be specified as an input to our DP recursions
- The counts $\hat{t}(x, x')$ are still relevant, but $\hat{e}(x, \mathbf{y})$ are used to accumulate EM update counts for $\mathbf{R}^{(x)}$
 - Can update t's and R^(x)'s simultaneously or asynchronously; c.f. Neal and Hinton



- Many applications in genomic biology
 - Annotation of multiple alignments of DNA, RNA or protein sequences
 - Isochores; gene prediction; DNA-protein binding site modeling and analysis; protein transmembrane structure, signal peptide, domain profiles... etc.

- Implementation: xrate (distributed with the DART library)
 - S-expressions for the underlying alignment grammar
 - Here "alignment grammar" means the alphabet Ω , the HMM transition matrix t, the Δ 's, the \mathbf{R} 's and the π 's
 - More generally, can use stochastic context-free grammars as well as HMMs, & states can emit several co-evolving columns
 - Stockholm format for alignment, tree, Viterbi annotation, likelihoods and posterior probabilities
 - Allows internal nodes as well as leaves to be specified
 - By default, gaps in the multiple alignment are treated as wildcards (unobserved character is summed over)
 - A smarter handling of gaps soon leads us to indel rate models and so-called "Statistical Alignment"
 - Log messages (type xrate -help and xrate -loghelp)
 - Command-line options (type xrate -help)



- HMMs model $P(Y) = \sum_{X} P(X, Y)$ (generative modeling). ML training maximizes this probability.
- Intuitively, since we are interested in predicting X correctly, it may make more sense to model conditional probability P(X|Y) (discriminative modeling)
- Consider the conditional likelihood for an HMM, expressed in terms of the feature vector {u, f} implied by X:

$$\log P(X|Y) = \frac{1}{P(Y)} \exp \left(\sum_{i,j} u(i,j) \log t(i,j) + \sum_{i,k} f(i,k) \log e(i,k) \right)$$

where P(Y) is computed by the Forward algorithm.

• We can write down a likelihood P(X|Y) for a similarly trellis-structured graphical model as follows

$$P(X|Y) = \frac{1}{Z} \exp \left(\sum_{i,j} u(i,j) a(i,j) + \sum_{i,k} f(i,k) b(i,k) \right)$$

where Z is a partition function (computed by a Forward-like sum-product algorithm)

$$Z = \sum_{X'} \exp \left(\sum_{i,j} u_{X'}(i,j)a(i,j) + \sum_{i,k} f_{X'}(i,k)b(i,k) \right)$$

For equivalence with the HMM, set $a(i,j) = \log t(i,j)$ and $b(i,k) = \log e(i,k)$.

- A linear-chain conditional random field is essentially such a trellis-structured model, lacking the normalization constraints on the parameters $\theta = \{a, b\}$ that are implicit in the generative HMM
 - Sutton & McCallum, "An Introduction to Conditional Random Fields for Relational Learning"
 - Lafferty, McCallum & Pereira, "Conditional Random Fields: Probabilistic Models for Segmenting and Labeling Sequence Data"
- To avoid overfitting, and in place of the normalization constraints, it is useful to add a *regularization* term, e.g. a Gaussian prior on a() and b() with variance σ², penalizing large weights. Then the function to be optimized is

$$\ell(\theta) = \log P(X|Y) - \sum_{i,j} \frac{a(i,j)^2}{2\sigma^2} + \sum_{i,k} \frac{b(i,k)^2}{2\sigma^2}$$

- Parameter estimation proceeds by **numerical optimization** of $\ell(\theta)$, typically using quasi-Newton algorithms
 - e.g. the BFGS algorithm: Dimitri P. Bertsekas. Nonlinear Programming. Athena Scientific, 2nd edition, 1999
- The partial derivatives $\frac{\partial \ell}{\partial a(i,j)}$ and $\frac{\partial \ell}{\partial b(i,k)}$ are computed by direct analogy to $\hat{t}(i,j)$ and $\hat{u}(i,k)$ in Baum-Welch.

- In the absence of normalization constraints, we are free to add more "features" without having to directly account for them by subdividing the state space of the HMM. For example: a run of T's, a palindromic motif, etc. In the generative (HMM) view, all such features must be explicitly identified with a path through the model, so that nothing is counted more than once. In a discriminative framework, it doesn't matter if a residue is counted twice.
- In the trellis factor graph view, functions relating x_i and y_i are $P(x_i|Y)$ rather than $P(y_i|x_i)$
- HMMs and linear CRFs form what Ng and Jordan (2002) call a "generative-discriminative pair".
 - Other such pairs include naive Bayes *vs* logistic regression.



Summary

HMMs