Statistical Natural Language Processing Sequence learning

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Some (typical) machine learning applications

	x (input)	y (output)
Spam detection	document	spam or not
Sentiment analysis	product review	sentiment
Medical diagnosis	patient data	diagnosis
Credit scoring	financial history	loan decision

The cases (input–output) pairs are assumed to be *independent and identically distributed* (i.i.d.).

Structured prediction

In many applications, the i.i.d. assumption is wrong

	x (input)	y (output)
POS tagging	word sequence	POS sequence
Parsing	word sequence	parse tree
OCR	image (array of pixels)	sequences of letters
Gene prediction	genome	genes

Structured prediction

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Structured/sequence learning is prevalent in NLP.

In this lecture ...

- Hidden Markov models (HMMs)
- A short note on graphical probabilistic models
- Alternatives to HMMs (briefly): HMEM / CRF

... and soon

Recurrent neural networks

Recap: chain rule

We rewrite the relation between the joint and the conditional probability as

$$P(X, Y) = P(X | Y)P(Y)$$

We can also write the same quantity as,

$$P(X, Y) = P(Y \mid X)P(X)$$

In general, for any number of random variables, we can write

$$P(X_1, X_2, ..., X_n) = P(X_1 | X_2, ..., X_n) P(X_2, ..., X_n)$$

Recap: (conditional) independence

If two variables X and Y are independent,

$$P(X | Y) = P(X)$$
 and $P(X, Y) = P(X)P(Y)$

If two variables X and Y are independent given another variable Z,

$$P(X,Y \mid Z) = P(X \mid Z)P(Y \mid Z)$$

An example: probability of a sentence

$$P(It's a beautiful day) = ?$$

• We cannot just count all occurrences of the sentence, and divide it to the total number of sentences in English

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- But we can base its probability to the probabilities of the words. Using chain rule

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\begin{split} P(It's\ a\ beautiful\ day) &= P(day\ |\ It's\ a\ beautiful) P(It's\ a\ beautiful) \\ &= P(day\ |\ It's\ a\ beautiful) P(beautiful\ |\ It's\ a) P(It's\ a) \\ &= P(day\ |\ It's\ a\ beautiful) P(beautiful\ |\ It's\ a) P(a\ |\ It's) P(It's) \end{split}
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```

• Did we solve the problem?

Markov chains

calculating probabilities

Given a sequence of events (or states), $q_1, q_2, \dots q_t$,

• In a *first-order* Markov chain probability of an event q_t is

$$P(q_t|q_1,...,q_{t-1}) = P(q_t|q_{t-1})$$

- Sometimes this equality is just an assumption
- In higher order chains, the dependence of history is extended, e.g., second-order Markov chain:

$$P(q_t|q_t, \dots, q_{t-1}) = P(q_t|q_{t-2}, q_{t-1})$$

Markov chains

definition

A Markov model is defined by,

- A set of states $Q = \{q_1, \dots, q_n\}$
- A special start state q₀
- A transition probability matrix

$$\mathbf{A} = \begin{bmatrix} a_{01} & a_{02} & \dots & a_{0n} \\ a_{11} & a_{12} & \dots & a_{1n} \\ \vdots & \vdots & \ddots & \vdots \\ a_{n1} & a_{n2} & \dots & a_{nn} \end{bmatrix}$$

where $\alpha_{\mathfrak{i}\mathfrak{j}}$ is the probability of transition from state \mathfrak{i} to state \mathfrak{j}

Back to sentence probability example

• With a first-order Markov assumption,

$$\begin{split} P(It's \ a \ beautiful \ day) &= P(day \ | \ It's \ a \ beautiful) P(beautiful \ | \ It's \ a) P(a \ | \ It's) P(It's) \\ &= P(day \ | \ beautiful) P(beautiful \ | \ a) P(a \ | \ It's) P(It's \ | \ \langle S \rangle) \end{split}$$

- Now the probabilities are easier to calculate
- The above approach is an example of *n-gram language models* that we will return very soon

Hidden/latent variables

- In many machine learning problems we want to account for unobserved/unobservable *latent* or *hidden* variables
- Some examples
 - 'personality' in many psychological data
 - 'topic' of a text
 - 'socio-economic class' of a speaker
- In most structured learning problems, the 'structure' is a hidden variable
- Latent variables make learning difficult: since we cannot observe them, how do we set the parameters?

Learning with hidden variables

(Another) informal/quick introduction to the EM algorithm

- The EM algorithm (or its variants) is used in many machine learning models with latent/hidden variables
- 1. Randomly initialize the parameters
- 2. Iterate until convergence:
- E-step compute likelihood of the data, given the parameters
- M-step re-estimate the parameters using the predictions based on the E-step

Hidden Markov models (HMM)

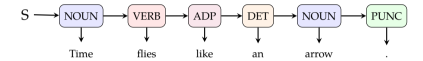
• HMMs are like Markov chains: probability of a state depends only a limited history of previous states

$$P(q_t|q_1,...,q_{t-1}) = P(q_t|q_{t-1})$$

- Unlike Markov chains, state sequence is hidden, they are not the observations
- At every state q_t , an HMM *emits* an output, o_t , whose probability depends only on the associated hidden state
- Given a state sequence $\mathbf{q} = q_1, \dots, q_T$, and the corresponding observation sequence $\mathbf{o} = o_1, \dots, o_T$,

$$P(\mathbf{o}, \mathbf{q}) = p(q_1) \left[\prod_{t=1}^{T} P(q_t | q_{t-1}) \right] \prod_{t=1}^{T} P(o_t | q_t)$$

Example: HMMs for POS tagging



- The tags are hidden
- Probability of a tag depends on the previous tag
- Probability of a word at a given state depends only on the current tag

HMMs: formal definition

An HMM is defined by

- A set of state $Q = \{q_1, \dots, q_n\}$
- The set of possible observations $V = \{v_1, \dots, v_m\}$
- A transition probability matrix

$$\mathbf{A} = \begin{bmatrix} a_{11} & a_{12} & \dots & a_{1n} \\ \vdots & \vdots & \ddots & \vdots \\ a_{n1} & a_{n2} & \dots & a_{nn} \end{bmatrix} \quad \begin{array}{l} a_{ij} \text{ is the probability of transition} \\ \text{from state } q_i \text{ to state } q_j \end{array}$$

- Initial probability distribution $\pi = \{P(q_1), \dots, P(q_n)\}$
- Probability distributions of

$$B = \begin{bmatrix} b_{11} & b_{12} & \dots & b_{1n} \\ \vdots & \vdots & \ddots & \vdots \\ b_{m1} & b_{m2} & \dots & b_{mn} \end{bmatrix} \quad \begin{array}{l} b_{ij} \text{ is the probability of emiting output o}_i \text{ at state q}_j \\ \end{array}$$

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A simple example

- Three states: N, V, D
- Four possible observations: a, b, c, d

$$\mathbf{A} = \begin{bmatrix} N & V & D \\ 0.2 & 0.7 & 0.1 \\ 0.5 & 0.1 & 0.4 \\ 0.8 & 0.1 & 0.1 \end{bmatrix} \begin{bmatrix} N \\ V \\ D \end{bmatrix}$$

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A simple example

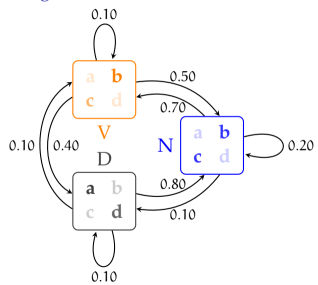
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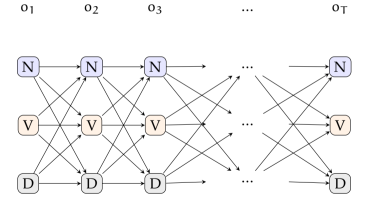
$$\pi = (0.3, 0.1, 0.6)$$

HMM transition diagram



Unfolding the states

HMM lattice (or trellis)



HMMs: three problems

Evaluation

Calculating likelihood of a given sequence

$$P(\mathbf{o} \mid M)$$

Recognition/decoding

Calculating probability of state sequence, given an observation sequence

$$P(\mathbf{q} \mid \mathbf{o}; M)$$

Learning

Given observation sequences, a set of states, and (sometimes) corresponding state sequences, estimate the parameters (π, A, B) of the HMM

Assigning probabilities to observation sequences

$$P(\mathbf{o} \mid M) = \sum_{\mathbf{q}} P(\mathbf{o}, \mathbf{q} \mid M)$$

- We need to sum over an exponential number of hidden state sequences
- The solution is using a dynamic programming algorithm
 - for each node of the trellis, store *forward probabilities*

$$\alpha_{t,i} = \sum_{j}^{N} \alpha_{t-1,j} P(q_i|q_j) P(o_i|q_i)$$

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Assigning probabilities to observation sequences

the forward algorithm

• Start with calculating all forward probabilities for t = 1

$$\alpha_{1,i} = \pi_i P(o_1|q_i)$$
 for $1 \le i \le N$

store the α values

• For t > 1,

$$\alpha_{t,i} = \sum_{j=1}^N \alpha_{t-1,j} P(q_i|q_j) P(o_i|q_i) \quad \text{for } 1 \leqslant i \leqslant N, 2 \leqslant t \leqslant T$$

• Likelihood of the observation is the sum of the forward probabilities of the last step

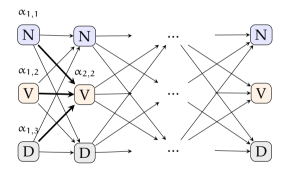
$$P(\mathbf{o}|M) = \sum_{i=1}^{N} \alpha_{i,T}$$

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Forward algorithm

HMM lattice (or trellis)





$$\alpha_{1,1} = \pi_N b_{xN}$$

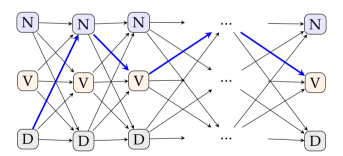
$$\alpha_{2,2} = \alpha_{1,1} a_{NV} b_{uV} + \alpha_{1,2} a_{VV} b_{uV} + \alpha_{1,3} a_{DV} b_{uV}$$

Determining best sequence of latent variables Decoding

- We often want to know the hidden state sequence given an observation sequence, $P(\mathbf{q} \mid \mathbf{o}; M)$
 - For example, given a sequence of tokens, find the most likely POS tag sequence
- The problem (also the solution, the *Viterbi algorithm*) is very similar to the forward algorithm
- Two major differences
 - we store maximum likelihood leading to each node on the lattice
 - we also store backlinks, the previous state that leads to the maximum likelihood

HMM decoding problem

a b c ...



Learning the parameters of an HMM

supervised case

- We want to estimate π , A, B
- If we have both the observation sequence o and the corresponding state sequence, MLE estimate is

$$\begin{split} \pi_i &= \frac{C(q_0 \rightarrow q_i)}{\sum_k C(q_0 \rightarrow q_k)} \\ \alpha_{ij} &= \frac{C(q_i \rightarrow q_j)}{\sum_k C(q_i \rightarrow q_k)} \\ b_{ij} &= \frac{C(q_i \rightarrow o_j)}{\sum_k C(q_i \rightarrow o_k)} \end{split}$$

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Learning the parameters of an HMM

• Given a training set with observation sequence(s) o and state sequence q, we want to find $\theta = (\pi, A, B)$

$$\underset{\boldsymbol{\theta}}{\operatorname{arg\,max}} P(\boldsymbol{o} \mid \boldsymbol{q}, \boldsymbol{\theta})$$

- Unlike i.i.d. case, we cannot factorize the likelihood over all observations
- Instead we use EM
 - Initialize θ
 - 2. Repeat until convergence

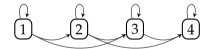
E-step given θ , estimate the hidden state sequence

M-step $\,$ given the estimated hidden states, use 'expected counts' to update θ

• An efficient implementation of EM algorithm is called *Baum-Welch algorithm*, or *forward-backward algorithm*

HMM variations

- The HMMs we discussed so far are called *ergodic* HMMs: all a_{ij} are non-zero
- For some applications, it is common to use HMMs with additional restrictions
- A well known variant (Bakis HMM) allows only forward transitions



• The emission probabilities can also be continuous, e.g., p(q|o) can be a normal distribution

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Directed graphical models: a brief divergence

Bayesian networks

 We saw earlier that joint distributions of multiple random variables can be factorized different ways

$$P(x, y, z) = P(x)P(y|x)P(z|x,y) = P(y)P(x|y)P(z|x,y) = P(z)P(x|z)P(y|x,z)$$

- Graphical models display this relations in graphs,
 - variables are denoted by nodes,
 - the dependence between the variables are indicated by edges
- Bayesian networks are directed acyclic graphs





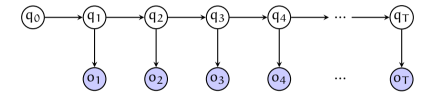


• A variable (node) depends only on its parents

Graphical models

- Graphical models define models involving multiple random variables
- It is generally more intuitive (compared to corresponding mathematical equations) to work with graphical models
- In a graphical model, by convention, the observed variables are shaded
- Graphs can also be undirected, which are called Markov random fields

HMM as a graphical model



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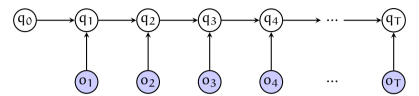
MaxEnt HMMs (MEMM)

- In HMMs, we model P(q, o) = P(q)P(o | q)
- In many applications, we are only interested in $P(q \mid o)$, which we can calculate using the Bayes theorem
- But we can also model $P(q \mid o)$ directly using a maximum entropy model

$$P(q_t \mid q_{t-1}, o_t) = \frac{1}{Z} e^{\sum w_i f_i(o_t, q_t)}$$

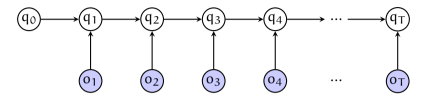
 f_i are features – can be any useful feature Z normalizes the probability distribution

MEMMs as graphical models

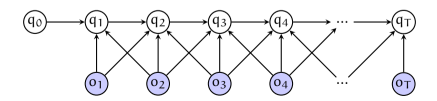


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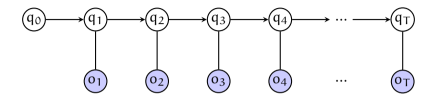
MEMMs as graphical models



We can also have other dependencies as features, for example



Conditional random fields



- A related model used in NLP is conditional random field (CRF)
- CRFs are undirected models
- CRFs also model $P(q \mid o)$ directly

$$P(\mathbf{q} \mid \mathbf{o}) = \frac{1}{Z} \prod_{t} f(q_{t-1}, q_t) g(q_t, o_t)$$

Generative vs. discriminative models

- HMMs are generative models, they model the joint distribution
 - you can generate the output using HMMs
- MEMMs and CRFs are discriminative models they model the conditional probability directly
- It is easier to add arbitrary features on discriminative models
- In general: HMMs work well when the state sequence, P(q), can be modeled well

Summary

- In many problems, e.g., POS tagging, i.i.d. assumption is wrong
- We need models that are aware of the effects of the sequence (or structure in general) in the data
- HMMs are generative sequence models:
 - Markov assumption between the hidden states (POS tags)
 - Observations (words) are conditioned on the state (tag)
- There are other sequence learning methods
 - Briefly mentioned: MEMM, CRF
 - Coming soon: recurrent neural networks

Next

Mon (after break) sequence learning with neural networks

Have nice break!