

Natural Language Processing (CSE 517): Conditional Random Fields

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Readings: Eisenstein (2019) 7 and 8

Motivation

Many tasks in NLP can be cast as **sequence labeling**, where each token (usually, word) gets its own label. Compare:

- ▶ Text classification: $\langle x_1, x_2, \dots, x_n \rangle \mapsto y \in \mathcal{L}$
- ▶ Sequence labeling: $\langle x_1 \mapsto y_1, x_2 \mapsto y_2, \dots, x_n \mapsto y_n \rangle$, each $y_i \in \mathcal{L}$
- ▶ Translation: $x \mapsto y \in \mathcal{V}_{target}^*$

Problems Typically Cast as Sequence Labeling

- ▶ supersense tagging (Ciaramita and Johnson, 2003)
- ▶ part-of-speech tagging (Church, 1988)
- ▶ morphosyntactic tagging (Habash and Rambow, 2005)
- ▶ segmentation into words (Sproat et al., 1996) or multiword expressions (Schneider et al., 2014)
- ▶ code switching (Solorio and Liu, 2008)
- ▶ dialogue acts (Stolcke et al., 2000)
- ▶ spelling correction (Kernighan et al., 1990)
- ▶ word alignment (Vogel et al., 1996)
- ▶ named entity recognition (Bikel et al., 1999)
- ▶ compression (Conroy and O'Leary, 2001)

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Ciaramita and Johnson (2003) and Ciaramita and Altun (2006) used a lexicon called WordNet to define 41 semantic classes for words.

- ▶ WordNet (Fellbaum, 1998) is a fascinating resource in its own right! See <http://wordnetweb.princeton.edu/perl/webwn> to get an idea.

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This represents a coarsening of the annotations in the Semcor corpus (Miller et al., 1993).

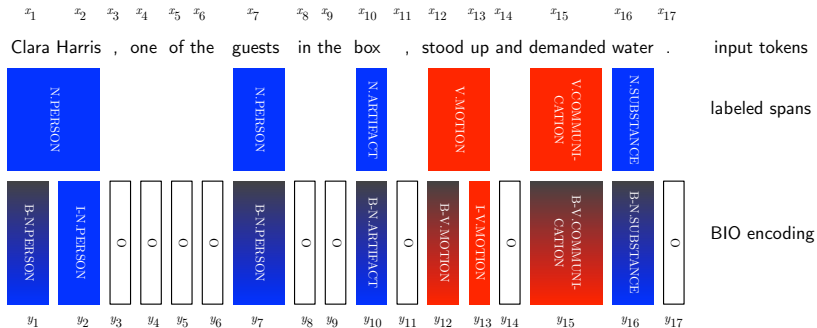
Example: *box's* Thirteen Synonym Sets, Eight Supersenses

1. box: a (usually rectangular) container; may have a lid. "he rummaged through a box of spare parts"
2. box/logge: private area in a theater or grandstand where a small group can watch the performance. "the royal box was empty"
3. box/boxful: the quantity contained in a box. "he gave her a box of chocolates"
4. corner/box: a predicament from which a skillful or graceful escape is impossible. "his lying got him into a tight corner"
5. box: a rectangular drawing. "the flowchart contained many boxes"
6. box/boxwood: evergreen shrubs or small trees
7. box: any one of several designated areas on a ball field where the batter or catcher or coaches are positioned. "the umpire warned the batter to stay in the batter's box"
8. box/box seat: the driver's seat on a coach. "an armed guard sat in the box with the driver"
9. box: separate partitioned area in a public place for a few people. "the sentry stayed in his box to avoid the cold"
10. box: a blow with the hand (usually on the ear). "I gave him a good box on the ear"
11. box/package: put into a box. "box the gift, please"
12. box: hit with the fist. "I'll box your ears!"
13. box: engage in a boxing match.

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Supsense Tagging Example



Observations

- ▶ Lots of subproblems: Which words have supersenses? Which words group together to form a multiword expression? For those that do, which supersense?
- ▶ Every word's label depends on the words around it, and their labels.
- ▶ Segmentation problems can be cast as sequence labeling (Ramshaw and Marcus, 1995):
 - ▶ Two labels, B and I, if every word must be in some segment
 - ▶ Three labels, B, I, and O, if some words are to be “discarded”
 - ▶ Variants for five labels (E for end, S for singleton), gaps/noncontiguous spans, and nesting, exist.

Concatenate B, I, etc., with labels to get labeled segmentation.

- ▶ Some sequences of labels might be invalid under your theory/label semantics.
- ▶ Evaluation: usually precision, recall, and F_1 on labeled segments.

Big Abstraction: Linguistic Analysis

Every linguistic analyzer is comprised of:

1. Theoretical motivation from linguistics and/or the text domain
2. An algorithm that maps \mathcal{V}^\dagger to some output space \mathcal{Y} .
 - ▶ Some \mathcal{Y} are very specialized, but others, like the one we discuss here, show up again and again.
3. An implementation of the algorithm
 - ▶ Once upon a time: rule systems and crafted rules
 - ▶ More robust: supervised learning from annotated data
 - ▶ Today: unsupervised pretraining followed by supervised finetuning

Sequence Labeling

Problem statement: given a sequence of n words \mathbf{x} , assign each a label from \mathcal{L} . Let $L = |\mathcal{L}|$.

Every approach we see today will cast the problem as:

$$\hat{\mathbf{y}} = \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} \operatorname{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta})$$

Naïvely, that's a classification problem where the number of possible 'labels' (output sequences) depends on the input and is $O(L^n)$ in size!

Sequence Labeling v. 0: Local Classifiers

Define score of a word x_i getting label $y \in \mathcal{L}$ in context: $\text{score}(\mathbf{x}, i, y; \boldsymbol{\theta})$, for example through a feature vector, $\mathbf{f}(\mathbf{x}, i, y)$. (Here, “ i ” indicates the position of the input word to be classified.)

Train a classifier to decode locally, i.e.,

$$\begin{aligned}\hat{y}_i &= \operatorname{argmax}_{y \in \mathcal{L}} \text{score}(\mathbf{x}, i, y; \boldsymbol{\theta}) \\ &\stackrel{\text{MLR}}{=} \operatorname{argmax}_{y \in \mathcal{L}} \boldsymbol{\theta}^\top \mathbf{f}(\mathbf{x}, i, y)\end{aligned}$$

The classifier is applied to each x_1, x_2, \dots in turn, but all the words can be made available at each position.

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We can do better when there are predictable relationships among labels.

Reflection

If we return to the original formulation,

$$\hat{\mathbf{y}} = \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} \text{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta}),$$

how can we write “Score” in terms of the notation on the last slide?

Local Classifiers (v. 0)



Lightweight; no need to learn anything new! But labels can't affect each other.

Sequence Labeling v. 1: Sequential Classifiers

Define score of a word x_i getting label y in context, *including previous labels*: $\text{score}(\mathbf{x}, i, \hat{\mathbf{y}}_{1:i-1}, y; \boldsymbol{\theta})$. (From here, we won't always write $\boldsymbol{\theta}$, but the dependence remains.)

Train a classifier, e.g.,

$$\hat{y}_i = \underset{y \in \mathcal{L}}{\operatorname{argmax}} \text{score}(\mathbf{x}, i, \hat{\mathbf{y}}_{1:i-1}, y)$$

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There is much literature on methods for training, and for decoding, with models like this. Important decoding method in NLP: beam search.

Beam Search for Sequential Classifiers

Input: \mathbf{x} (length n), a sequential classifier's scoring function score, and beam width k

Let H_0 score hypotheses at position 0, defining only $H_0(\langle \rangle) = 0$.

For $i \in \{1, \dots, n\}$:

- ▶ Empty C .
- ▶ For each hypothesis $\hat{\mathbf{y}}_{1:i-1}$ scored by H_{i-1} :
 - ▶ For each $y \in \mathcal{L}$, place new hypothesis $\hat{\mathbf{y}}_{1:i}y \rightarrow H_{i-1}(\hat{\mathbf{y}}_{1:i-1}) + \text{score}(\mathbf{x}, i, \hat{\mathbf{y}}_{1:i-1}, y)$ into C .
- ▶ Let H_i be the k -best scored elements of C .

Output: best scored element of H_n .

Notes on Beam Search for Sequential Classifiers

- ▶ Runtime is $O(n^2kL)$, space is $O(n^2k)$.
- ▶ You can improve runtime (e.g., to $O(nkL)$) if computation is shared across different i (often true with neural networks).
- ▶ Special cases:
 - ▶ $k = 1$ is greedy left-to-right decoding.
 - ▶ At $k = L^n$, you're doing brute force, exhaustive search.
- ▶ Generally: no guarantee.

Reflection

Suppose your label set is built out of BIO tags. For an output \hat{y} to be well-formed, it suffices to ensure that it contains no “OI” label bigrams.

How would you modify beam search to guarantee well-formedness?

Sequential Classifiers (v. 1)



Very powerful! Algorithms lack guarantees.

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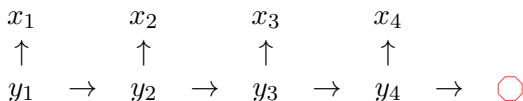
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$$x_4 \sim p_{\text{emission}}(X \mid y_4)$$

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The next approach should remind you of language models. It assumes that labeled sequences are generated according to the following story:



$$y_5 \sim p_{\text{transition}}(Y \mid y_4)$$

Sequence Labeling v. 2: Hidden Markov Models

By convention, $y_{n+1} = \text{○}$ is always the “stop label.”

$$\begin{aligned} p(\mathbf{X} = \mathbf{x}, \mathbf{Y} = \mathbf{y}) &= p_{start}(y_1) \cdot \\ &\quad \prod_{i=1}^n p_{emission}(x_i \mid y_i) \cdot p_{transition}(y_{i+1} \mid y_i) \\ \hat{\mathbf{y}} &= \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} p(\mathbf{Y} = \mathbf{y} \mid \mathbf{X} = \mathbf{x}) \\ &= \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} p(\mathbf{X} = \mathbf{x}, \mathbf{Y} = \mathbf{y}) \\ &= \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} \log p(\mathbf{X} = \mathbf{x}, \mathbf{Y} = \mathbf{y}) \end{aligned}$$

We can solve the global decoding problem *exactly* (i.e., find the model-optimal $\hat{\mathbf{y}}$) in $O(nL^2)$ time and $O(nL)$ space using the Viterbi algorithm (more later).

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$p_{emission}$ is a distribution over words, for each label. Many people find this counterintuitive! Estimation: counting occurrences of labels with words, and normalizing (per label, not per word).

$p_{transition}$ is exactly a bigram (first-order Markov) model over labels.

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With classical HMMs, the parameters ($p_{transition}$, $p_{emission}$, p_{start}) have a closed form if you have labeled data! The hardest part is implementing the algorithm for choosing the “argmax” label sequence. Downside:

- ▶ You don't get to design or learn features.

Reflection

The runtime of the model-optimal decoding algorithm for HMMs depends quadratically on the size of \mathcal{L} . For some problems (e.g., supersense tagging) the label set can be large. Can you think of a way to trade the guarantee of model-optimality for speed, while still using the HMM?

Hidden Markov Models (v. 2)



Algorithmically beautiful; lack of features is unsatisfying.

Sequence Labeling v. 3

To endow HMMs with features, we can replace the “lookup” probabilities ($p_{transition}, p_{emission}, p_{start}$) with scoring functions. This idea was explored by Berg-Kirkpatrick et al. (2010).

Classical HMM (v. 2):

$$\hat{y} = \operatorname{argmax}_{y \in \mathcal{L}^n} \log p_{start}(y_1) + \sum_{i=1}^n \left(\log p_{emission}(x_i | y_i) + \log p_{transition}(y_{i+1} | y_i) \right)$$

This approach (v. 3):

$$\hat{y} = \operatorname{argmax}_{y \in \mathcal{L}^n} s_{start}(y_1) + \sum_{i=1}^n s_{emission}(x_i, y_i) + s_{transition}(y_i, y_{i+1})$$

Each “ s ” could be a linear scoring function (like in MLR), perhaps using word or label vectors. For now, I’m hiding the parameters of each s .

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- ▶ No part of the the scoring function looks at neighboring words.



Brings features to HMMs, but learning is going to require more than just counting and normalizing.

Sequence Labeling v. 4

Let each scoring component (“ s ”) “see” the whole input. By convention, $y_0 = \bigcirc$ is always the “start label.”

$$\hat{\mathbf{y}} = \operatorname{argmax}_{\mathbf{y} \in \mathcal{L}^n} \overbrace{\sum_{i=0}^n s(\mathbf{x}, i, y_i, y_{i+1})}^{\text{Score}(\mathbf{x}, \mathbf{y})}$$

Note that \mathbf{x} can have arbitrary length, so we need “ s ” functions that are capable of adapting to variable-length input.

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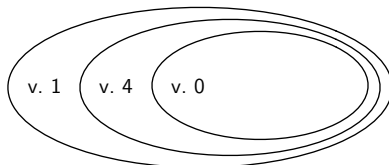
- ▶ Decoding is essentially the same as the HMM and v. 3: Viterbi algorithm.
- ▶ As with v. 3, learning is complicated and depends on the form of each “ s .”
- ▶ This model strictly generalizes local classifiers (v. 0), the HMM (v. 2), and v. 3.



Even better features for HMMs, with the promise of efficient decoding and learning.

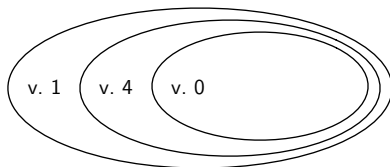
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Claim: As we move from v. 1 (sequential classifiers) to v. 4 to v. 0 (local classifiers), the scoring functions available become strictly less expressive.



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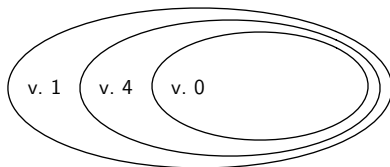
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Compare v. 1 and v. 4. What kinds of features can you use in v. 1 that you can't use in v. 4?

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




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Compare v. 1 and v. 4. What kinds of features can you use in v. 1 that you can't use in v. 4?

Now consider v. 4 and v. 0. What kinds of features can you use in v. 4 that you can't use in v. 0?

Where We Are

					
	0	1	2	3	4
Score decomp.	$s(\mathbf{x}, i, y_i)$	$s(\mathbf{x}, i, \mathbf{y}_{1:i})$	emission/ transition	$s(x_i, y_i) + s(y_i, y_{i+1})$	$s(\mathbf{x}, i, y_i, y_{i+1})$
learn	SGD	?	count & normalize	?	?
decode	local	beam search	Viterbi	Viterbi	Viterbi

The Main Dish

Two Problems to Solve

1. Decoding: the Viterbi algorithm for choosing \hat{y} .
 - ▶ Usually taught for classical HMMs (v. 2); I will teach it for v. 4, abstracting away “ s .”
2. Learning: estimating the parameters of each s function.
 - ▶ Depending on your choices here, you arrive at the structured perceptron, the classical conditional random field (CRF), neural CRFs, and more.

A Data Structure

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

The cell at row j , column i will hold information pertaining to choosing $\hat{y}_i = \ell_j$.

The End of the Sequence

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

$$\begin{aligned}\hat{y}_n &= \operatorname{argmax}_{y_n \in \mathcal{L}} \sum_{i=0}^n s(\mathbf{x}, i, y_i, y_{i+1}) \\ &= \operatorname{argmax}_{y_n \in \mathcal{L}} s(\mathbf{x}, i, y_{n-1}, y_n) + s(\mathbf{x}, i, y_n, \text{red circle})\end{aligned}$$

The decision about \hat{y}_n is a function of y_{n-1} , \mathbf{x} , and nothing else!

High-Level View of the Viterbi Algorithm

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Recurrence

First, think about the *score* of the best sequence.

Let $\heartsuit_i(y)$ be the score of the best label sequence for $\mathbf{x}_{1:i}$ that ends in y . It is defined recursively:

$$\heartsuit_{n+1}(\bigcirc) = \max_{y_n \in \mathcal{L}} s(\mathbf{x}, n, y_n, \bigcirc) + \boxed{\heartsuit_n(y_n)}$$

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\vdots

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
\vdots

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
\vdots

$$\heartsuit_1(y) = s(\mathbf{x}, 0, \text{green circle}, y)$$

Viterbi Procedure (Part I: Prefix Scores)


		input sequence				
		x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1					
	ℓ_2					
	\vdots					
	ℓ_L					
						

Viterbi Procedure (Part I: Prefix Scores)

input sequence					
	x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$			
	ℓ_2	$\heartsuit_1(\ell_2)$			
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$			
					


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Viterbi Procedure (Part I: Prefix Scores)

input sequence					
	x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$	$\heartsuit_2(\ell_1)$		
	ℓ_2	$\heartsuit_1(\ell_2)$	$\heartsuit_2(\ell_2)$		
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$	$\heartsuit_2(\ell_L)$		
					

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		input sequence			
		x_1	x_2	\dots	x_n
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	ℓ_2	$\heartsuit_1(\ell_2)$	$\heartsuit_2(\ell_2)$		$\heartsuit_n(\ell_2)$
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$	$\heartsuit_2(\ell_L)$		$\heartsuit_n(\ell_L)$
					

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
input sequence					
	x_1	x_2	\dots	x_n	
ℓ_1	$\heartsuit_1(\ell_1)$	$\heartsuit_2(\ell_1)$		$\heartsuit_n(\ell_1)$	
ℓ_2	$\heartsuit_1(\ell_2)$	$\heartsuit_2(\ell_2)$		$\heartsuit_n(\ell_2)$	
\vdots					
ℓ_L	$\heartsuit_1(\ell_L)$	$\heartsuit_2(\ell_L)$		$\heartsuit_n(\ell_L)$	
\bigcirc					$\heartsuit_{n+1}(\bigcirc)$

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
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Viterbi Procedure (Part I: Prefix Scores and Backpointers)

input sequence					
	x_1	x_2	\dots	x_n	
ℓ_1					
ℓ_2					
\vdots					
ℓ_L					
					


Viterbi Procedure (Part I: Prefix Scores and Backpointers)

		input sequence				
		x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$ $\text{bp}_1(\ell_1)$				
	ℓ_2	$\heartsuit_1(\ell_2)$ $\text{bp}_1(\ell_2)$				
	\vdots					
	ℓ_L	$\heartsuit_1(\ell_L)$ $\text{bp}_1(\ell_L)$				
						

$$\heartsuit_1(y) = s(\mathbf{x}, 0, \bigcirc, y)$$

$$\text{bp}_1(y) = \bigcirc$$


Viterbi Procedure (Part I: Prefix Scores and Backpointers)

		input sequence			
		x_1	x_2	\dots	x_n
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$ $\text{bp}_1(\ell_1)$	$\heartsuit_2(\ell_1)$ $\text{bp}_2(\ell_1)$		
	ℓ_2	$\heartsuit_1(\ell_2)$ $\text{bp}_1(\ell_2)$	$\heartsuit_2(\ell_2)$ $\text{bp}_2(\ell_2)$		
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$ $\text{bp}_1(\ell_L)$	$\heartsuit_2(\ell_L)$ $\text{bp}_2(\ell_L)$		
					

$$\heartsuit_i(y) = \max_{y_{i-1} \in \mathcal{L}} s(\mathbf{x}, i-1, y_{i-1}, y) + \boxed{\heartsuit_{i-1}(y_{i-1})}$$

$$\text{bp}_i(y) = \operatorname{argmax}_{y_{i-1} \in \mathcal{L}} s(\mathbf{x}, i-1, y_{i-1}, y) + \boxed{\heartsuit_{i-1}(y_{i-1})}$$


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input sequence					
	x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$ $\text{bp}_1(\ell_1)$	$\heartsuit_2(\ell_1)$ $\text{bp}_2(\ell_1)$	$\heartsuit_n(\ell_1)$ $\text{bp}_n(\ell_1)$	
	ℓ_2	$\heartsuit_1(\ell_2)$ $\text{bp}_1(\ell_2)$	$\heartsuit_2(\ell_2)$ $\text{bp}_2(\ell_2)$	$\heartsuit_n(\ell_2)$ $\text{bp}_n(\ell_2)$	
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$ $\text{bp}_1(\ell_L)$	$\heartsuit_2(\ell_L)$ $\text{bp}_2(\ell_L)$	$\heartsuit_n(\ell_L)$ $\text{bp}_n(\ell_L)$	
					

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Viterbi Procedure (Part I: Prefix Scores and Backpointers)

input sequence					
	x_1	x_2	\dots	x_n	
\mathcal{L}	ℓ_1	$\heartsuit_1(\ell_1)$ $\text{bp}_1(\ell_1)$	$\heartsuit_2(\ell_1)$ $\text{bp}_2(\ell_1)$	$\heartsuit_n(\ell_1)$ $\text{bp}_n(\ell_1)$	
	ℓ_2	$\heartsuit_1(\ell_2)$ $\text{bp}_1(\ell_2)$	$\heartsuit_2(\ell_2)$ $\text{bp}_2(\ell_2)$	$\heartsuit_n(\ell_2)$ $\text{bp}_n(\ell_2)$	
	\vdots				
	ℓ_L	$\heartsuit_1(\ell_L)$ $\text{bp}_1(\ell_L)$	$\heartsuit_2(\ell_L)$ $\text{bp}_2(\ell_L)$	$\heartsuit_n(\ell_L)$ $\text{bp}_n(\ell_L)$	
					$\heartsuit_{n+1}(\text{red octagon})$ $\text{bp}_{n+1}(\text{red octagon})$

$$\heartsuit_{n+1}(\text{red octagon}) = \max_{y_n \in \mathcal{L}} s(\mathbf{x}, n, y_n, \text{red octagon}) + \boxed{\heartsuit_n(y_n)}$$

$$\text{bp}_{n+1}(\text{red octagon}) = \operatorname{argmax}_{y_n \in \mathcal{L}} s(\mathbf{x}, n, y_n, \text{red octagon}) + \boxed{\heartsuit_n(y_n)}$$

Full Viterbi Procedure

Input: scores $s(\mathbf{x}, i, y, y')$, for all $i \in \{0, \dots, n\}$, $y, y' \in \mathcal{L}$

Output: $\hat{\mathbf{y}}$

1. Base case: $\heartsuit_1(y) = s(\mathbf{x}, 0, \bigcirc, y)$
2. For $i \in \{2, \dots, n+1\}$:
 - Solve for $\heartsuit_i(*)$ and $\text{bp}_i(*)$.

$$\heartsuit_i(y) = \max_{y_{i-1} \in \mathcal{L}} s(\mathbf{x}, i-1, y_{i-1}, y) + \heartsuit_{i-1}(y_{i-1}),$$

$$\text{bp}_i(y) = \operatorname{argmax}_{y_{i-1} \in \mathcal{L}} s(\mathbf{x}, i-1, y_{i-1}, y) + \heartsuit_{i-1}(y_{i-1})$$

(At $n+1$ we're only interested in $y = \bigcirc$.)

3. $\hat{y}_{n+1} \leftarrow \bigcirc$
4. For $i \in \{n, \dots, 1\}$:
 - $\hat{y}_i \leftarrow \text{bp}_{i+1}(\hat{y}_{i+1})$

Viterbi Asymptotics

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

Viterbi Asymptotics

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

Space: need to store s , and fill in the cells above.

Viterbi Asymptotics

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

Space: need to store s , and fill in the cells above. $O(nL^2)$ for s (in the most general case, often less), $O(nL)$ for cells

Viterbi Asymptotics

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

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Runtime: each cell requires an “argmax.”

Viterbi Asymptotics

		input sequence			
		x_1	x_2	\dots	x_n
labels in \mathcal{L}	ℓ_1				
	ℓ_2				
	\vdots				
	ℓ_L				

Space: need to store s , and fill in the cells above. $O(nL^2)$ for s (in the most general case, often less), $O(nL)$ for cells

Runtime: each cell requires an “argmax.” $O(nL^2)$

Why it Works

Viterbi exploits the distributivity property:

$$\begin{aligned}\max_{\mathbf{y}_{1:n}} \sum_{i=0}^n s(\mathbf{x}, i, y_i, y_{i+1}) &= \max_{y_n} s(\mathbf{x}, i, y_n, \text{red circle}) + \max_{\mathbf{y}_{1:n-1}} \sum_{i=0}^{n-1} s(\mathbf{x}, i, y_i, y_{i+1}) \\ &= \max_{y_n} s(\mathbf{x}, i, y_n, \text{red circle}) + \max_{y_{n-1}} s(\mathbf{x}, i, y_{n-1}, y_n) \\ &\quad + \max_{\mathbf{y}_{1:n-2}} \sum_{i=0}^{n-2} s(\mathbf{x}, i, y_i, y_{i+1})\end{aligned}$$

Max plus max plus max plus max plus . . .

Back to “ s ”

We haven't said much about the function that scores candidate label pairs at different positions, $s(\mathbf{x}, i, y, y')$.

This function is very important; two common choices are:

- ▶ Expert-designed, task-specific features $\mathbf{f}(\mathbf{x}, i, y, y')$ and weights θ
- ▶ A neural network that encodes x_i in context, y_i , and y_{i+1} and gives back a goodness score

Either way, let θ denote the parameters of s . From now on, we'll use $s(\mathbf{x}, i, y, y'; \theta)$ and $\text{Score}(\mathbf{x}, \mathbf{y}; \theta)$ to emphasize that “ s ” is a function of parameters θ we need to estimate.

Probabilistic View of Learning

As we've done before, we start with the principle of maximum likelihood to estimate θ :

$$\begin{aligned}\theta^* &= \arg \max_{\theta \in \mathbb{R}^d} \prod_{i=1}^T p(Y = \mathbf{y}_i \mid X = \mathbf{x}_i; \theta) \\ &= \arg \max_{\theta \in \mathbb{R}^d} \sum_{i=1}^T \log p(Y = \mathbf{y}_i \mid X = \mathbf{x}_i; \theta) \\ &= \arg \min_{\theta \in \mathbb{R}^d} \sum_{i=1}^T \underbrace{-\log p(Y = \mathbf{y}_i \mid X = \mathbf{x}_i; \theta)}_{\text{sometimes called "log loss" or "cross entropy"}}\end{aligned}$$

Next, we'll drill down into " $p(Y = \mathbf{y}_i \mid X = \mathbf{x}_i; \theta)$."

Conditional Random Fields

Lafferty et al. (2001)

CRFs are a tremendously influential model that generalizes multinomial logistic regression to structured outputs like sequences.

$$p_{\text{CRF}}(\mathbf{y} \mid \mathbf{x}; \boldsymbol{\theta}) = \frac{\exp \text{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta})}{Z(\mathbf{x}; \boldsymbol{\theta})}$$

$$Z(\mathbf{x}; \boldsymbol{\theta}) = \sum_{\mathbf{y}' \in \mathcal{Y}(\mathbf{x})} \exp \text{Score}(\mathbf{x}, \mathbf{y}'; \boldsymbol{\theta})$$

$$-\log p_{\text{CRF}}(\mathbf{y} \mid \mathbf{x}; \boldsymbol{\theta}) = - \underbrace{\text{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta})}_{\text{"hope"}} + \underbrace{\log Z(\mathbf{x}; \boldsymbol{\theta})}_{\text{"fear"}}$$

So, our “CRF”:

- ▶ Uses Viterbi for decoding (our v. 4 sequence labeler)
- ▶ Trains parameters to maximize likelihood (like MLR and NNs)

Conditional Random Field

Lafferty et al. (2001)



Sequence-Level Log Loss

Here's the maximum likelihood learning problem (equivalently, sequence-level log loss):

$$\boldsymbol{\theta}^* = \operatorname{argmin}_{\boldsymbol{\theta} \in \mathbb{R}^d} \sum_{i=1}^T -\text{Score}(\mathbf{x}_i, \mathbf{y}_i; \boldsymbol{\theta}) + \log Z(\mathbf{x}_i; \boldsymbol{\theta})$$

If we can calculate and differentiate (w.r.t. $\boldsymbol{\theta}$) the Score and Z functions, we can use SGD to learn.

Reflection

Given a training instance $\langle \mathbf{x}_i, \mathbf{y}_i \rangle$, what do you need to do to calculate $\text{Score}(\mathbf{x}_i, \mathbf{y}_i; \boldsymbol{\theta})$?

Calculating $Z(x; \theta)$

Good news! The algorithm that gives us Z is *almost exactly like* the Viterbi algorithm.

Forward algorithm: sums the expScore values for all label sequences, given x , in the same asymptotic time and space as Viterbi.

Let $\alpha_i(y)$ be the sum of all (exponentiated) scores of label prefixes of length i , ending in y .

Some Algebra

Given the decomposition

$$\text{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta}) = \sum_{i=0}^n s(\mathbf{x}, i, y_i, y_{i+1}; \boldsymbol{\theta}),$$

it holds that

$$\exp \text{Score}(\mathbf{x}, \mathbf{y}; \boldsymbol{\theta}) = \prod_{i=0}^n e^{s(\mathbf{x}, i, y_i, y_{i+1}; \boldsymbol{\theta})},$$

and therefore

$$Z(\mathbf{x}; \boldsymbol{\theta}) = \sum_{\mathbf{y}' \in \mathcal{Y}(\mathbf{x})} \prod_{i=0}^n e^{s(\mathbf{x}, i, y'_i, y'_{i+1}; \boldsymbol{\theta})}$$

Forward Algorithm

Input: scores $s(\mathbf{x}, i, y, y'; \boldsymbol{\theta})$, for all $i \in \{0, \dots, n\}$, $y, y' \in \mathcal{L}$

Output: $Z(\mathbf{x}; \boldsymbol{\theta})$

1. Base case: $\alpha_1(y) = e^{s(\mathbf{x}, 0, \text{green}, y; \boldsymbol{\theta})}$
2. For $i \in \{2, \dots, n+1\}$:
 - Solve for $\alpha_i(\ast)$.

$$\alpha_i(y) = \sum_{y_{i-1} \in \mathcal{L}} e^{s(\mathbf{x}, i-1, y_{i-1}, y; \boldsymbol{\theta})} \times \alpha_{i-1}(y_{i-1})$$

(At $n+1$ we're only interested in $y = \text{red}$.)

3. Return $\alpha_{n+1}(\text{red})$, which is equal to $Z(\mathbf{x}; \boldsymbol{\theta})$.

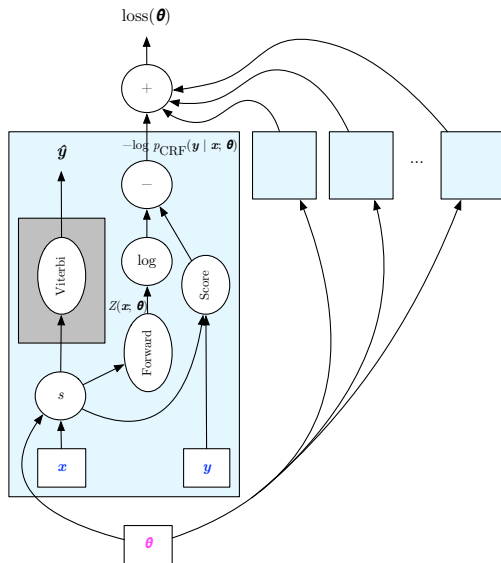
Intuitions about the Forward Algorithm

Just as Viterbi changes “scary max over big sum” to “max plus max plus max plus . . . ,”
the Forward algorithm changes “scary sum over big product” to
“plus times plus times plus times”

If you organize the operations in the other direction, you get the Backward algorithm.

You can differentiate Z with respect to s , because it's all just exp, addition, and multiplication. If you mechanically derive the partial derivatives, you will rediscover the Backward algorithm.

Computation Graph View of CRF



Reflection

Earlier in the lecture, I promised that learning would have some guarantees. Consider:

- ▶ The runtime and space requirements for calculating the loss and gradient, as a function of the data.
- ▶ The conditions under which we can confidently expect convergence to a global optimum of the likelihood if we use SGD.

An Alternative: Structured Perceptron

Recall that CRF = v. 4 + sequence-level log loss.

Perceptron loss (Collins, 2002):

$$\theta^* = \operatorname{argmin}_{\theta \in \mathbb{R}^d} \sum_{i=1}^T -\operatorname{Score}(\mathbf{x}_i, \mathbf{y}_i; \theta) + \max_{\mathbf{y}} \operatorname{Score}(\mathbf{x}_i, \mathbf{y}; \theta)$$

The structured perceptron = v. 4 + perceptron loss.

Regularization

Just as in classification with linear and non-linear models, you'll want to take steps to avoid overfitting.

The same tools (e.g., ℓ_2 and ℓ_1 penalties for linear model weights, and dropout for neural networks) can be used here.

Digestif: Connections and Generalizations

V. 2–4 are weighted finite-state machines (think of labels as states).

The models we saw today are all “first order” sequence models in the sense that each y_i only interacts with one immediate neighbor through s .

► Second-order: $\text{Score}(\mathbf{x}, \mathbf{y}) = \sum_{i=0}^n s(\mathbf{x}, i, y_i, y_{i+1}, y_{i+2})$

► m th-order: $\text{Score}(\mathbf{x}, \mathbf{y}) = \sum_{i=0}^n s(\mathbf{x}, i, \mathbf{y}_{i:i+m})$

Viterbi for m th order has $O(nL^{m+1})$ runtime.

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