

### Planning and Optimal Control 4. Markov Random Fields

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## Syllabus



#### A. Models for Sequential Data

- Tue. 22.10.(1)1. Markov ModelsTue. 29.10.(2)2. Hidden Markov Models
- Tue. 5.11. (3) 3. State Space Models
- Tue. 12.11. (4) 3b. (ctd.)

#### **B.** Models for Sequential Decisions

- Tue. 19.11. (5) 1. Markov Decision Processes
- Tue. 26.11. (6) 1b. (ctd.)
- Tue. 3.12. (7) 2. Introduction to Reinforcement Learning
- Tue. 10.12. (8) 3. Monte Carlo and Temporal Difference Methods
- Tue. 17.12. (9) 4. Q Learning
- Tue. 24.12. — Christmas Break —
- Tue. 7.1. (10) 5. Policy Gradient Methods
- Tue. 14.1. (11) tba
- Tue. 21.1. (12) tba
- Tue. 28.1. (13) 8. Reinforcement Learning for Games
- Tue. 4.2. (14) Q&A

## Outline



- 1. Markov Random Fields
- 2. Inference in MRFs
- 3. Learning MRFs
- 4. Partially Observed Markov Random Fields
- 5. Conditional Random Fields

### Outline



#### 1. Markov Random Fields

- 2. Inference in MRFs
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## Motivation



- models for sequential data often naturally can be written using conditional density / probability functions conditioning on the past
  - ► e.g., Markov models of type p(x<sub>t</sub> | x<sub>t-1</sub>) or the latent state transition model p(z<sub>t</sub> | z<sub>t-1</sub>)
- for other types of structured data there usually is no such marked direction
  - ▶ e.g., for images
- directed graphical models / Bayesian networks such as Markov Models and HMMs can be generalized to multidimensional data
  - multidimensional HMMs
  - ► require a direction to be marked, e.g., from top left to bottom right.
  - ► but it "feels" somewhat artificial
- $\rightsquigarrow$  use undirected graphical models / Markov random fields

## Stochastic Processes & Random Fields

Stochastic process / random process / random function:

• a collection of random variables  $X_i$  indexed by some index set I

 $\{X_i \mid i \in I\}$ 

- ► discrete-time:  $I = \{a, a + 1, a + 2, \dots, b\},$   $a \in \mathbb{Z} \cup \{-\infty\}, b \in \mathbb{Z} \cup \{\infty\}$
- ► continuous-time: I = [a, b],  $a \in \mathbb{R} \cup \{-\infty\}, b \in \mathbb{R} \cup \{\infty\}$
- **Random field**:  $I \subseteq \mathbb{R}^{K}$  or a grid (spatial) or a graph.
- = a density for structured data, on  $\mathcal{X}^{I}$



## Markov Random Fields



A random field p on an undirected graph I is called Markov if

▶ each variable is independent from all others given its neighbors

$$egin{aligned} X_i \perp \{X_i \mid i \in I\} \setminus N_i \setminus \{X_i\} \mid N_i \ N_i := \{X_j \mid j \in I, j ext{ is a neighbor of } i ext{ in } I\} \end{aligned}$$

## Hammersley-Clifford Theorem



A random field p on I is Markov iff

► p factorizes into non-negative functions over maximal cliques in I:

$$\exists (q_c)_{c \in C} : p(x) = \prod_{c \in C} q_c(x_c)$$
$$C := \{ c \subseteq I \mid c \text{ is a maximal clique} \}$$

► *q<sub>c</sub>* are called **potentials**.

Note: A set *c* of vertices is called a **clique** if all its nodes are linked in *I*. A clique *c* is called **maximal**, if there is no clique *d*:  $d \supseteq c$ .

## Pairwise MRF



- ► potentials can be defined on any subsets of maximal cliques
  - but not on supersets
- ▶ most simple non-trivial potentials: on every edge

$$p(x) = \prod_{i,j \in I \text{ linked}} q_{i,j}(x_i, x_j)$$

pairwise MRF



## Parametrizing Potentials I: Tables / Arrays

- potential functions q are parametrized
  - $\blacktriangleright$  so that parameters  $\theta$  can be learnt to fit the model to data
- ▶ if all variables in a potential q are discrete, the simplest parametrization is a table / a multidimensional array:

$$q(x_1,\ldots,x_{\mathcal{K}})=\theta_{x_1,\ldots,x_{\mathcal{K}}},\quad \theta\in(\mathbb{R}^+_0)^{\mathcal{X}_1\times\mathcal{X}_2\times\cdots\mathcal{X}_{\mathcal{K}}}$$

example:

	$x_2 \setminus x_1$	red	green	blue
$\theta = 0$	square	0.2	0.7	2.3
	circle	0.5	0.0	0.2

- ▶ potentials are not normalized (generally do not sum to 1).
  - ▶ for a general graph, there would be no guarantee that the product of however normalized potentials again is normalized.

## Example: Image Segmentation



- ▶ let  $I = \{1, ..., N\} \times \{1, ..., M\}$  be the coordinates of the pixels of an  $N \times M$  image
- ► let's define the graph on *I* to have an edge for neighboring pixels, i.e.,

$$(i,j) :\sim (i-1,j), (i+1,j), (i,j-1), (i,j+1)$$

- ► the state space X := {road, offroad, obstacle} are labels of the pixels denoting the type of object they belong to.
- ► here, the maximal cliques are just single edges
- ► an MRF could define its pairwise potentials via a table:

	$x_1 \setminus x_2$	road	offroad	obstacle
$(\mathbf{x}, \mathbf{y}) =$	road	0.9	0.1	0.2
$q_{1,2}(x_1, x_2) =$	offroad	0.1	0.9	0.01
	obstacle	0.2	0.01	0.9

## The Partition Function

- $\blacktriangleright$  potentials usually are not normalized / sum to 1.
  - even if they would, for general graphs it would not guarantee that their product is normalized.
- ▶ an MRF with parametrized potentials therefore is represented via

$$p(x \mid \theta) = \frac{1}{Z(\theta)} \prod_{c \in C} q_c(x_c \mid \theta_c)$$

•  $Z(\theta)$  is called **partition function** 

$$Z(\theta) := \sum_{x \in \mathcal{X}} \prod_{c \in C} q_c(x_c \mid \theta_c)$$

- Z makes the MRF p a proper probability function / sum to 1.
- ► Z in general depends on all parameters.
- ... but on none of the  $x_i$ .



## Parametrizing Potentials II: Features & Log-linear Models

- often array potentials do not work
  - e.g., because they have too many parameters if cliques are large or include nominal variables with many levels
  - cliques contain continuous variables
- ► alternative approach:
  - 1. define features  $\phi(x_1, \ldots, x_K)$  for the variables of a potential q
  - 2. define the potential as a log-linear model in the features:

$$q(x_1,\ldots,x_K \mid \theta) := e^{\theta^T \phi(x_1,\ldots,x_K)}$$
$$= e^{\sum_{\ell=1}^L \theta_\ell \phi_\ell(x_1,\ldots,x_K)}$$

aka maximum entropy model, maxent model

$$\log p(x \mid \theta) = \sum_{c} \theta_{c}^{T} \phi_{c}(x_{c}) - \log Z(\theta)$$



## Example: Image Segmentation (ctd.)

let's define the graph on I to have an edge for pixels up to L1-distance 2, i.e.,

$$\begin{array}{cccc} (i,j-2) \\ (i,j) :\sim & (i-2,j) \\ (i,j) :\sim & (i-2,j) \\ (i-1,j+1) \\ (i,j+2) \end{array} \begin{array}{cccc} (i,j-2) \\ (i,j-1) \\ (i+1,j-1) \\ (i,j+1) \\ (i,j+2) \end{array} \begin{array}{cccc} (i,j-2) \\ (i+1,j-1) \\ (i+1,j+1) \\ (i+1,j+1) \end{array} \end{array}$$

- now maximal cliques are a pixel (i, j) and its four distance 1 neighbors
- instead we could define features, e.g., the frequency of each label in the neighborhood:

 $\phi(x_c)_1 :=$  frequency of road in  $x_c$  $\phi(x_c)_2 :=$  frequency of offroad in  $x_c$  $\phi(x_c)_3 :=$  frequency of obstacle in  $x_c$ 

► and potentials as log-linear model in these features:

$$q_c(x_c \mid \theta) := e^{\theta_1 \phi(x_c)_1 + \theta_2 \phi(x_c)_2 + \theta_3 \phi(x_c)_3}$$



## Tables as Special Case of Log-Linear Models

► if we define a binary indicator feature for each joint variable value:

$$\phi(x_1,\ldots,x_{\mathcal{K}}) = (\mathbb{I}((x_1,\ldots,x_{\mathcal{K}}) = (x'_1,\ldots,x'_{\mathcal{K}})))_{(x'_1,\ldots,x'_{\mathcal{K}}) \in \mathcal{X}^{\mathcal{K}}}$$

then the log-linear model is just the array potential.

## Parametrizing Potentials III: Parameter Sharing

- often different potentials describe the same relation, just between different sets of variables
  - ► e.g., q<sub>1,2</sub> and q<sub>5,17</sub> describe the relation between a pixel and its neighbors, but for different image patches
    - one centered at (1,2), the other at (5,17)
- ▶ such potentials (and their parameters) often can be shared

$$q_c(x_c \mid \theta_c) = q(x_c \mid \theta)$$

- example: image segmentation
  - usually potentials will not depend on the reference pixel, but all be shared.
- ► parameter sharing allows to roll-out a MRF to graphs of different sizes
  - e.g., images of different width and height
  - ► MRF with shared parameters define MRF templates



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## **MRF** Inference



Inference in MRF (and generally graphical models) requires work:

- ► exact inference:
  - ► join tree algorithm
  - simpler (less efficient) algorithm:
    - variable elimination / bucket elimination
- ► approximate inference:
  - variational inference
  - ► inference via sampling / Monte Carlo inference

### Inference I: Margin Query

► MRF:

$$p(X) = \prod_{c \in C} q_c(X_c), \quad C \subseteq \mathcal{P}(I)$$

where I indices of variables,

$$\begin{array}{l} \mathcal{X}_i \text{ domain of variable } X_i \text{ for } i \in I, \\ \mathcal{X}_c = (X_i)_{i \in c}, \ x_c = (x_i)_{i \in c}, \ \mathcal{X}_c = \prod_{i \in c} \mathcal{X}_i \text{ for } i \in I, \\ \mathcal{C} \text{ set of cliques } c \subseteq I, \\ q_c : \mathcal{X}_c \to \mathbb{R}_0^+ \text{ clique potential of } c \in C \end{array}$$

- ► margin query:
  - target variables  $T \subseteq I$

$$p(X_T) = \sum_{x_R \in \mathcal{X}_R} p(X_T, X_R = x_R), \quad R := I \setminus T$$



## Variable eliminiation



- ► idea:
  - ▶ marginalize out one non-target variable X<sub>i</sub> at a time
  - collect all potentials containing this variable
  - $\blacktriangleright$  ... and replace them by their product
    - summing over all possible values for X<sub>i</sub>
    - materializing the product as array

#### Variable eliminiation / Algorithm 1 infer-mrf-varelim $(T, (q_c)_{c \in C})$ : while $\bigcup_{c \in C} c \setminus T \neq \emptyset$ : 2 choose $i \in \bigcup_{c \in C} c \setminus T$ arbitrarily 3 (C,q) := eliminate-variable(i, C, q)4 5 $p := \prod_{c \in C} q_c$ p := normalize(p)6 return p 7 \* eliminate-variable $(i, C, (q_c)_{c \in C})$ : 9 $D := \{c \in C \mid i \in c\}$ $c' := \bigcup_{c \in D} c \setminus \{i\}$ 10 $q_{c'} := \left( \sum_{x_i \in \mathcal{X}_i} \prod_{c \in D} q_c(x_i, (x_{c'})_{c \cap c'}) \right)_{x_i \in \mathcal{X}_i}$ 11 12 $C' := C \setminus D \cup \{c'\}$ return $C', (q_c)_{c \in C'}$ 13 where

- $T \subseteq I$  target variables to infer marginal of
- ▶  $(q_c)_{c \in C}$  MRF defined by a set of potentials on  $c \subseteq I$
- yields  $(p_{x_{\mathcal{T}}})_{x_{\mathcal{T}} \in \mathcal{X}_{\mathcal{T}}}$  marginal of variables T





## Inference / Variable eliminiation / Example

- $\blacktriangleright I := \{A, B, C, D, E, F\}$
- ▶  $C := \{\{A\}, \{A, B\}, \{A, C\}, \{B, D\}, \{B, C, E\}, \{C, F\}, \{F\}\}$
- $T := \{D\}$
- elimination sequence: F, E, C, A, B



## Inference / Variable eliminiation / Example

- $\blacktriangleright I := \{A, B, C, D, E, F\}$
- $C := \{\{A\}, \{A, B\}, \{A, C\}, \{B, D\}, \{B, C, E\}, \{C, F\}, \{F\}\}$
- $\blacktriangleright T := \{D\}$
- elimination sequence: F, E, C, A, B

► compute:  

$$q(C) := \sum_{F} q(C, F) q(F)$$

$$q(B, C) := \sum_{E} q(B, C, E) q(C)$$

$$q(A, B) := \sum_{C} q(B, C, E) q(C) q(A, B) q(A)$$

$$q(B, D) := \sum_{A} q(A, B) q(B, D)$$

$$q(D) := \sum_{B} q(B, D)$$

# Inference II: Conditional Probabilities p(A | B = b)



- ► in general, A and B could denote sets/vectors of variables:  $p(X_{i_1}, X_{i_2}, \dots, X_{i_N} | X_{j_1} = b_1, X_{j_2} = b_2, \dots, X_{j_M} = b_M\}$   $A = (X_{i_1}, X_{i_2}, \dots, X_{i_N})$   $B = (X_{j_1}, X_{j_2}, \dots, X_{j_M})$   $b = (b_1, \dots, b_M)$
- for each conditioning variable / value pair (B<sub>m</sub>, b<sub>m</sub>) = (X<sub>jm</sub>, b<sub>m</sub>) add an evidence potential epd<sub>jm,bm</sub>: epd<sub>i,b</sub> : X<sub>i</sub> → ℝ<sub>0</sub><sup>+</sup> x ↦ I(x = b)
- ▶ infer marginal of A for the potentials  $p' := p \cup \{ epd_{i,b} \mid (i,b) \in zip(B,b) \}$

Note:  $zip(A, B) := \{(A_i, B_i) \mid i = 1, ..., |A|\}$  for two sequences  $A \in \mathcal{X}^*, B \in \mathcal{Y}^*$  of equal length.

## Infering Conditional Probabilities / Example

- ► let us model the following rules:
  - ▶ if there is precipitation, roads are three times more likely to be slippery.
  - ► if there is frost, roads are two times more likely to be slippery.
- A: There is heavy precipitation.
  - B: There is frost.
  - C: Roads are slippery.

$$q(A, C) = \begin{pmatrix} 0.5 & 0.5 \\ 0.25 & 0.75 \end{pmatrix}, \quad q(B, C) = \begin{pmatrix} 0.5 & 0.5 \\ 0.3 & 0.7 \end{pmatrix}$$

What are the chances of the road to be slippery if there is precipitation, but no frost?

$$p(C \mid A = 1, B = 0)$$





## Infering Conditional Probabilities / Example

initial potentials:

$$q(A, C) = \begin{pmatrix} 0.5 & 0.5 \\ 0.25 & 0.75 \end{pmatrix}, \quad q(B, C) = \begin{pmatrix} 0.5 & 0.5 \\ 0.3 & 0.7 \end{pmatrix},$$
$$q(A) = epd_{A,1}(A) = \begin{pmatrix} 0 & 1 \end{pmatrix}, \quad q(B) = epd_{B,0}(B) = \begin{pmatrix} 1 & 0 \end{pmatrix}$$

► eliminate *A*:

$$q(C) = \sum_{A} q(A, C)q(A) = ( 0.25 \ 0.75 )$$

► eliminate *B*:

$$q'(C) = \sum_{B} q(B, C)q(B) = ( 0.5 \ 0.5 )$$

► collect:  $q''(C) = q(C) \odot q'(C) = (0.25 \ 0.75) \odot (0.5 \ 0.$ =  $(0.125 \ 0.375)$ normalization $(q'')(C) = (0.25 \ 0.75)$ 

Planning and Optimal Control 2. Inference in MRFs

## Inference III: Expectations $\mathbb{E}(f(X_T))$

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for a general function

$$f: \mathcal{X}_T \to \mathbb{R}, \quad T \subseteq I$$

- infer marginal  $p(X_T)$
- compute array  $(f(X_T))_{x_T \in \mathcal{X}_T}$  elementwise
- sum all cells of the elementwise tensor product  $p(X_T) f(X_T)$

$$\mathbb{E}(f(x_T)) = \sum_{x_T \in \mathcal{X}_T} p(x_T) f(x_T)$$

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 $\sim$ 



## Learning Maxent Models via Gradient Descent

▶ gradients for maxent models are straight-forward to derive:

$$\ell(\theta; x) := \log p(x \mid \theta) = \sum_{c} \theta_{c}^{T} \phi_{c}(x_{c}) - \log Z(\theta)$$

$$\nabla_{\theta_{c}} \ell(\theta; x) = \phi_{c}(x_{c}) - \nabla_{\theta_{c}} \log Z(\theta)$$

$$Z(\theta) := \sum_{x \in \mathcal{X}} \prod_{c \in C} e^{\theta_{c}^{T} \phi_{c}(x_{c})}$$

$$\nabla_{\theta_{c}} \log Z(\theta) = \frac{1}{Z(\theta)} \sum_{x \in \mathcal{X}} \prod_{c \in C} e^{\theta_{c}^{T} \phi_{c}(x_{c})} \phi_{c}(x_{c})$$

$$= \sum_{x \in \mathcal{X}} p(x \mid \theta) \phi_{c}(x_{c}) = \mathbb{E}(\phi_{c}(X_{c}))$$

$$\Rightarrow \quad \nabla_{\theta_{c}} \ell(\theta; x) = \phi_{c}(x_{c}) - \mathbb{E}(\phi_{c}(X_{c}))$$

▶ but it requires inference in the model to compute  $\mathbb{E}(\phi_c(X_c))$  !

## Learning Maxent Models via Gradient Descent

1 learn-mrf-gd(
$$x, (q_c)_{c \in C}, \eta, K, \epsilon$$
):  
2 for  $c \in C$ :  $\theta_c := 1_{\Theta_c}$   
3 for  $k := 1 : K$ :  
4 for  $c \in C$ :  $f_c := 0$   
5 for  $n = 1 : N$ :  
6 for  $c \in C$ :  
7  $f_c += \phi(x_{n,c})/N$   
8 for  $c \in C$ :  
9  $p_c := infer-mrf(c, (q_c(\theta_c))_{c \in C})$   
10  $g_c := 0$   
11 for  $v \in \mathcal{X}^C$ :  
12  $g_c += p_c(v) \cdot \phi(v)$   
13  $\Delta \theta_c := f_c - g_c$   
14 if  $\sum_c ||\Delta \theta_c||_2 < \epsilon$ :  
15 return  $(\theta_c)_{c \in C}$   
16 for  $c \in C$ :  
17  $\theta_c := \theta_c - \eta \Delta \theta_c$   
19  $return (f_c) = 0$ 

where

- ▶  $x \in (\mathcal{X}')^*$  data
- (q<sub>c</sub>)<sub>c∈C</sub> potentials of cliques, having parameters θ<sub>c</sub> ∈ Θ<sub>c</sub>
- C ⊆ 2<sup>I</sup> variables of the potentials / maximal cliques of graph I
- $\blacktriangleright~\eta$  steplength
- *K* maximal number of iterations
- $\epsilon$  minimum gradient norm

yields  $(\theta_c)_{c \in C}$  parameters of the potentials



Planning and Optimal Control 3. Learning MRFs



### **Optimality Criterion: Matching Moments**

$$\ell(\theta; x_{1:N}) := \frac{1}{N} \sum_{n=1}^{N} \log p(x_n \mid \theta)$$

$$\nabla_{\theta_c} \ell(\theta; x) = \frac{1}{N} \sum_{n=1}^{N} \phi_c(x_{n,c}) - \nabla_{\theta_c} \log Z(\theta)$$

$$= \mathbb{E}_{p_{emp}}(\phi_c(x_c)) - \mathbb{E}_p(\phi_c(x_c))$$

$$\ell(\theta; x) = 0;$$

thus at  $\nabla_{\theta_c} \ell(\theta; x) = 0$ :  $\mathbb{E}_{p_{emp}}(\phi_c(x_c)) = \mathbb{E}_p(\phi_c(x_c))$ 

#### moment matching



## Learning Maxent Models via Iterative Proportional Fitting

► for array potentials

$$\mathbb{E}_{p}(\phi_{c}(x_{c})) = \mathbb{E}_{p}(\mathbb{I}(x_{c} = x'))_{x' \in \mathcal{X}_{c}} = p(x_{c} \mid \theta) \propto \theta_{c,x_{c}}$$
$$\mathbb{E}_{p_{emp}}(\phi_{c}(x_{c})) = \mathbb{E}_{p_{emp}}(\mathbb{I}(x_{c} = x'))_{x' \in \mathcal{X}_{c}} = p_{emp}(x_{c}) = \frac{1}{N} \sum_{n=1}^{N} \mathbb{I}(x_{n,c} = x_{c})$$

► fixpoint iteration:

$$heta_{c,x_c}^{(t+1)} = heta_{c,x_c}^{(t)} rac{p(x_c \mid heta^{(t)})}{p_{\mathsf{emp}}(x_c)}, \quad x_c \in \mathcal{X}_c$$

approximate inference



## Learning Maxent Models via Iterative Proportional Fitting

1 learn-mrf-ipf(x, 
$$(q_c)_{c \in C}$$
):  
2 for  $c \in C$ :  
3  $\theta_c := 1_{\Theta_c}$   
4  $p_{emp,c} := (\frac{1}{N} \sum_{n=1}^{N} \mathbb{I}(x_{n,c} = x'_c))_{x'_c \in \mathcal{X}_c}$   
5 repeat  
6 for  $c \in C$ :  
7  $p := infer-mrf(c, (q_c(\theta_c))_{c \in C})$   
8 for  $x_c \in \mathcal{X}_c$ :  
9  $\theta_{c,x_c} := \theta_{c,x_c} \frac{p_{x_c}}{(p_{emp,c})_{x_c}}$   
10 until convergence  
11 return  $(\theta_c)_{c \in C}$ 

where

- ▶  $x \in (\mathcal{X}')^*$  data
- ► (q<sub>c</sub>)<sub>c∈C</sub> potentials of cliques, having parameters θ<sub>c</sub> ∈ Θ<sub>c</sub>
- C ⊆ 2<sup>I</sup> variables of the potentials / maximal cliques of graph I

yields  $(\theta_c)_{c\in C}$  parameters of the potentials

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## Learning via EM Algorithm



Learning from complete data we just discussed in the last section.

For incomplete data use EM:

- ► E-step: complete the data using inference
  - ► inference for every instance individually
  - ▶ joint marginals for variables cooccurring in the same clique/potential
  - every instance is split into possible completions
    - ► the probability of the completion figures as caseweight for the M-step
    - possibly different splittings for every clique
- M-step: update parameters  $\theta$  using a method for learning from complete data.
  - ► e.g., gradient descent



Case weight for joint completions  $\mathcal{X}_{c\cap Z}$  of instance *x*:

$$w_{c,x} := p(c \cap Z \mid X = x_c), \quad c \in C, x \in \mathcal{X}$$

where

$$egin{aligned} X &:= (X_1, \dots, X_M) & ext{observed variables} \ Z &:= (Z_1, \dots, Z_K) & ext{latent variables} \end{aligned}$$

$$\nabla_{\theta_c} \ell(\theta; x) = \phi_c(x_c) - \mathbb{E}(\phi_c(X_c))$$
  
$$\rightsquigarrow \quad \nabla_{\theta_c} \ell(\theta; x, z) = \sum_{z_c \in \mathcal{X}_{c \cap Z}} w_{c,x} \left(\phi_c(x_c, z_c) - \mathbb{E}(\phi_c(X_c, z_c))\right)$$
$$= \left(\sum_{z_c \in \mathcal{X}_{c \cap Z}} w_{c,x} \phi_c(x_c, z_c)\right) - \mathbb{E}(\phi_c(X_c))$$

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## The Sequence Labeling Problem



Given data  $\mathcal{D}^{\text{train}}$  of N pairs  $(x_n, y_n)$  of sequences  $x_n \in \mathcal{X}^*, y_n \in \mathcal{Y}^*$  of same length,

- ► *x<sub>n</sub>* called **predictor sequence**,
- y<sub>n</sub> called target sequence

and a loss function  $\ell:\mathcal{Y}^*\times\mathcal{Y}^*\to\mathbb{R},$  learn the parameters  $\theta$  of a model

$$p(y \mid x, \theta)$$

s.t. for yet unseen data  $\mathcal{D}^{\text{test}}$  the loss

$$\ell(\hat{y}; \mathcal{D}^{\mathsf{test}}) = rac{1}{|\mathcal{D}^{\mathsf{test}}|} \sum_{(x,y) \in \mathcal{D}^{\mathsf{test}}} \ell(y, \hat{y}(x))$$

is minimal.

## The Sequence Labeling Problem / Example

Part of speech tagging:

- ▶ predictor sequence *x*: words of a sentence.
  - e.g., At the banks Jim is catching a big fish.
- ► target sequence y: part of speech classes of each word.
  - e.g., pre art N N V V art adj N
  - ► a label for each element of the sequence:

At	the	banks	Jim	is	catching	а	big	fish.
pre	art	Ν	Ν	V	V	art	adj	Ν

► usually 9 different POS classes/tags/labels for English:

noun:	car	pronoun:	she	adjective:	yellow
verb:	to drive	adverb:	gracefully	preposition:	under
conjunction:	and	interjection:	hurray	article:	the



## Shiversiter Shideshain

## Label Sequencing Models 1: HMMs

► model targets y<sub>t</sub> by hidden states z<sub>t</sub>, predictors x<sub>t</sub> by observations x<sub>t</sub>.

$$p(x_{1:T}, y_{1:T} \mid \theta) = p(y_1 \mid \theta) \prod_{t=2}^{T} p(y_t \mid y_{t-1}, \theta) \prod_{t=1}^{T} p(x_t \mid y_t, \theta)$$

- ► learning:
  - simple, from fully observed data.
- prediction:
  - compute MAP  $p(z_{1:T} | x_{1:T})$  (decoding)
- but HMMs are generative models
  - spend data to learn generative models of the predictors  $x_t$
  - ► like Linear Discriminant Analysis vs. Logistic Regression



## Label Sequencing Models 2: MEMMs

Maximum entropy markov model (MEMM)

$$p(y_{1:T} | x_{1:T}, \theta) = p(y_1 | x_1, \theta) \prod_{t=2}^{T} p(y_t | y_{t-1}, x_t, \theta)$$

- Markov chain with state transition conditionend on concurrent predictor
- ▶ but  $y_t$  does not depend on future predictors  $x_{t+1:T}$ 
  - $y_t$  and  $x_{t+1}$  are d-separated by v-connection at  $y_{t+1}$ .
  - ▶ in the POS example, x<sub>9</sub> = fish would not allow to recognize x<sub>3</sub> = banks as noun (riverbank) instead of as verb (to bank in the financial sense).
  - called "label bias problem"

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## Label Sequencing Models 3: CRFs

► Conditional Random Fields (CRFs)

$$p(y_{1:T} \mid x_{1:T}, \theta) = \frac{1}{Z(x_{1:T}, \theta)} \prod_{t=1}^{T} q(y_t \mid x_t, \theta) \prod_{t=2}^{T} q(y_t, y_{t-1} \mid x_t, x_{t-1}, \theta)$$

often with log-linear potentials

$$q(y_t \mid x_r, \theta) = e^{\theta_{(t)}^T \phi(x_t, y_t, y_{t-1})}$$
$$q(y_t, y_{t-1} \mid x_t, x_{t-1}, \theta) = e^{\theta_{(t,t-1)}^T \phi(x_t, x_{t-1}, y_t, y_{t-1})}$$

- $\blacktriangleright$  = MRF with potentials depending on all predictors
- in CRFs,  $y_t$  does depend on  $x_{t+1:T}$  (through  $y_{t+1}$ )
  - ► because  $q(y_{t+1}, y_t)$  is not conditioned on  $y_t$  as  $p(y_{t+1} | y_t)$  is.

Planning and Optimal Control 5. Conditional Random Fields

## Example: Handwriting Recognition





[source: Murphy 2012, p.686]

recognize handwritten texts

 $\begin{aligned} q(y_t \mid x_{1:T}, \theta) &:= q(y_t \mid x_t, \theta_1) := \text{deep neural network for letters} \\ q(y_t, y_{t-1} \mid x_{1:T}, \theta) &:= q(y_t, y_{t-1} \mid \theta_2) := \text{language bigram model} \end{aligned}$ 

## Conditional Random Fields

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- many CRFs are chain-structured as the ones discussed
- CRFs can be defined more generally on arbitrary targets y structured by a graph 1:

$$p((y_i)_{i\in I} \mid x, \theta) = \frac{1}{Z(x, \theta)} \prod_{c\in C} q(y_c \mid x, \theta)$$

often with log-linear potentials

$$q(y_c \mid x, \theta) = e^{\theta_c^T \phi(x, y_c)}$$

 $\blacktriangleright$  = MRF with potentials depending on all predictors

 $\sim$ 

## Learning CRFs via Gradient Descent

• gradients for CRFs are straight-forward to derive:

$$\ell(\theta; y, \mathbf{x}) := \log p(y \mid \mathbf{x}, \theta) = \sum_{c} \theta_{c}^{T} \phi_{c}(y_{c}, \mathbf{x}) - \log Z(\mathbf{x}, \theta)$$

$$\nabla_{\theta_{c}} \ell(\theta; y, \mathbf{x}) = \phi_{c}(y_{c}, \mathbf{x}) - \nabla_{\theta_{c}} \log Z(\mathbf{x}, \theta)$$

$$Z(\mathbf{x}, \theta) := \sum_{y \in \mathcal{Y}} \prod_{c \in C} e^{\theta_{c}^{T} \phi_{c}(y_{c}, \mathbf{x})}$$

$$\nabla_{\theta_{c}} \log Z(\mathbf{x}, \theta) = \frac{1}{Z(\theta} \sum_{y \in \mathcal{Y}} \prod_{c \in C} e^{\theta_{c}^{T} \phi_{c}(y_{c}, \mathbf{x})} \phi_{c}(y_{c}, \mathbf{x})$$

$$= \sum_{y \in \mathcal{Y}} p(y \mid \mathbf{x}, \theta) \phi_{c}(y_{c}, \mathbf{x}) = \mathbb{E}(\phi_{c}(y_{c}, \mathbf{x}))$$

$$\Rightarrow \quad \nabla_{\theta_{c}} \ell(\theta; y, \mathbf{x}) = \phi_{c}(y_{c}, \mathbf{x}) - \mathbb{E}(\phi_{c}(y_{c}, \mathbf{x}))$$

▶ requires *N* inferences in the model to compute  $\mathbb{E}(\phi_c(y_{n,c}, \mathbf{x_n}))$  !



Summary (1/3)



- Random fields / stochastic processes are densities for structured data
  - represented by a set of random variables indexed by a (undirected) graph.
- Markov random fields
  - each variable is independent from all others given its neighbors or equivalently
  - decompose in a product over the **maximal cliques**.
    - clique factors are called **potentials**.
- Potentials usually are parametrized:
  - parametrized as arrays:
    - ▶ an array with a value for every combination of values of the variables.
  - parametrized by features and a log-linear model:

$$q(x_{1:K} \mid \theta) = e^{\theta^T \phi(x_{1:K})}$$

- parameter sharing for potentials describing the same relation between different instances / sets of variables
  - ► see also Markov Logic networks

# Summary (2/3)



- The **partition function** enforces the marginal of the product of potentials to be 1.
  - depending on all parameters
  - ► it usually is given only implicitly as sum over all possible instances and thus cannot be computed but for very simple models.
- ► A simple method for **inference** in MRFs is **variable eliminiation**.
  - marginalize out one non-target variable at a time
  - multiplying all potentials containing this variable
  - observed variables are represented by evidence potentials.
- ► MRFs can be **learned** by **gradient descent**.
  - ► due to the partition function requires inference of the expected features
  - ▶ one inference per gradient step (and clique/potential)

## Summary (3/3)



- ► Partially observed MRFs may contain latent variables.
  - can be learned by EM.
  - M-step: gradient descent as for fully observed MRFs.
  - E-step: infer distribution of latent variables
    - for each clique containing a latent variable
    - joint distribution per clique
    - ▶ requires *N* inferences per EM step (and affected clique/potential)
- Conditional random fields make potentials depend on the predictors.
  - ► to ensure that a target can depend on future observations (for the sequence labeling problem; "label bias problem").
  - ▶ also can be learned by gradient descent as well.
  - also require N inferences per gradient step (and clique/potential)

## Further Readings

- Markov random fields:
  - Murphy 2012, chapter 19.





#### References

Kevin P. Murphy. Machine Learning: A Probabilistic Perspective. The MIT Press, 2012.

