Machine Learning Summer School

Lecture 2: Inference and Propagation Algorithms

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Inference in a graphical model

Consider the following graph:

\[ p(A, B, C, D, E) = p(A)p(B)p(C|A, B)p(D|B, C)p(E|C, D) \]

**Inference**: evaluate the probability distribution over some set of variables, given the values of another set of variables.

For example, how can we compute \( p(A|C = c) \)? Assume each variable is binary.

**Naive method:**

\[
\begin{align*}
p(A, C = c) &= \sum_{B, D, E} p(A, B, C = c, D, E) \quad \text{[16 terms]} \\
p(C = c) &= \sum_A p(A, C = c) \quad \text{[2 terms]} \\
p(A|C = c) &= \frac{p(A, C = c)}{p(C = c)} \quad \text{[2 terms]}
\end{align*}
\]

Total: \(16 + 2 + 2 = 20\) terms have to be computed and summed
Inference in a graphical model

Consider the following graph:

which represents:

\[ p(A, B, C, D, E) = p(A)p(B)p(C|A, B)p(D|B, C)p(E|C, D) \]

Computing \( p(A|C = c) \).

**More efficient method:**

\[
\begin{aligned}
p(A, C = c) &= \sum_{B,D,E} p(A)p(B)p(C = c|A, B)p(D|B, C = c)p(E|C = c, D) \\
&= \sum_B p(A)p(B)p(C = c|A, B) \sum_D p(D|B, C = c) \sum_E p(E|C = c, D) \\
&= \sum_B p(A)p(B)p(C = c|A, B) \quad [4 \text{ terms}] 
\end{aligned}
\]

Total: \( 4+2+2 = 8 \) terms

Belief propagation methods use the conditional independence relationships in a graph to do efficient inference (for singly connected graphs, exponential gains in efficiency!).
**Belief Propagation (in singly connected DAGs)**

**Definition:** A DAG is *singly connected* if its underlying undirected graph is a tree, *ie* there is only one undirected path between any two nodes.

**Goal:** For some node $X$ we want to compute conditional $p(X|e)$ given evidence (i.e. observed, visible variables) $e$.

Since we are considering singly connected graphs:

- every node $X$ divides the evidence into *upstream* $e^+_X$ and *downstream* $e^-_X$
- every edge $X \rightarrow Y$ divides the evidence into *upstream* $e^+_{XY}$ and *downstream* $e^-_{XY}$.
Three key ideas behind Belief Propagation

Idea 1: The probability of a variable $X$ can be found by combining upstream and downstream evidence:

$$p(X|e) = \frac{p(X,e)}{p(e)} = \frac{p(X,e^+_X,e^-_X)}{p(e^+_X,e^-_X)} \propto p(X|e^+_X) \times \frac{p(e^-_X|X,e^+_X)}{p(e^+_X|X,e^-_X)}$$

$X$ d-separates $e^-_X$ from $e^+_X$

$$= p(X|e^+_X)p(e^-_X|X) = \pi(X)\lambda(X)$$

Idea 2: The upstream and downstream evidence can be computed via a local message passing algorithm between the nodes in the graph.

Idea 3: “Don’t send back to a node (any part of) the message it sent to you!”
Factor graph propagation

Algorithmically and implementationally, it’s often easier to convert directed and undirected graphs into factor graphs, and run factor graph propagation.

\[ p(x) = p(x_1)p(x_2|x_1)p(x_3|x_2)p(x_4|x_2) \]
\[ \equiv f_1(x_1, x_2)f_2(x_2, x_3)f_3(x_2, x_4) \]

Singly connected vs Multiply connected factor graphs:
In a factor graph, the joint probability distribution is written as a product of factors. Consider a vector of variables \( x = (x_1, \ldots, x_n) \)

\[
P(x) = p(x_1, \ldots, x_n) = \frac{1}{Z} \prod_j f_j(x_{S_j})
\]

where \( Z \) is the normalisation constant, \( S_j \) denotes the subset of \( \{1, \ldots, n\} \) which participate in factor \( f_j \) and \( x_{S_j} = \{x_i : i \in S_j\} \).

**variables nodes**: we draw open circles for each variable \( x_i \) in the distribution.

**factor nodes**: we draw filled dots for each factor \( f_j \) in the distribution.
Propagation in Factor Graphs

Let $n(x)$ denote the set of factor nodes that are neighbors of $x$. Let $n(f)$ denote the set of variable nodes that are neighbors of $f$.

We can compute probabilities in a factor graph by propagating messages from variable nodes to factor nodes and vice versa.

**message from variable $x$ to factor $f$:**

$$\mu_{x\rightarrow f}(x) = \prod_{h \in n(x) \setminus \{f\}} \mu_{h\rightarrow x}(x)$$

**message from factor $f$ to variable $x$:**

$$\mu_{f\rightarrow x}(x) = \sum_{x \setminus x} \left( f(x) \prod_{y \in n(f) \setminus \{x\}} \mu_{y\rightarrow f}(y) \right)$$

where $x$ are the variables that factor $f$ depends on, and $\sum_{x \setminus x}$ is a sum over all variables neighboring factor $f$ except $x$. 
Propagation in Factor Graphs

\( n(x) \) denotes the set of factor nodes that are neighbors of \( x \).

\( n(f) \) denotes the set of variable nodes that are neighbors of \( f \).

**message from variable \( x \) to factor \( f \):**

\[
\mu_{x \rightarrow f}(x) = \prod_{h \in n(x) \setminus \{ f \}} \mu_{h \rightarrow x}(x)
\]

**message from factor \( f \) to variable \( x \):**

\[
\mu_{f \rightarrow x}(x) = \sum_{x \setminus x} \left( f(x) \prod_{y \in n(f) \setminus \{ x \}} \mu_{y \rightarrow f}(y) \right)
\]

If a variable has only one factor as a neighbor, it can initiate message propagation.

Once a variable has received all messages from its neighboring factor nodes, one can compute the probability of that variable by multiplying all the messages and renormalising:

\[
p(x) \propto \prod_{h \in n(x)} \mu_{h \rightarrow x}(x)
\]
initialise all messages to be constant functions

an example schedule of messages resulting in computing $p(x_4)$:

<table>
<thead>
<tr>
<th>message direction</th>
<th>message value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_1 \to f_1$</td>
<td>$1(x_1)$</td>
</tr>
<tr>
<td>$x_3 \to f_2$</td>
<td>$1(x_3)$</td>
</tr>
<tr>
<td>$f_1 \to x_2$</td>
<td>$\sum x_1 f_1(x_1, x_2) 1(x_1)$</td>
</tr>
<tr>
<td>$f_2 \to x_2$</td>
<td>$\sum x_3 f_2(x_3, x_2) 1(x_3)$</td>
</tr>
<tr>
<td>$x_2 \to f_3$</td>
<td>$\left(\sum x_1 f_1(x_1, x_2)\right) \left(\sum x_3 f_2(x_3, x_2)\right)$</td>
</tr>
<tr>
<td>$f_3 \to x_4$</td>
<td>$\sum x_2 f_3(x_2, x_4) \left(\sum x_1 f_1(x_1, x_2)\right) \left(\sum x_3 f_2(x_3, x_2)\right)$</td>
</tr>
</tbody>
</table>

where $1(x)$ is a constant uniform function of $x$
Propagation in Factor Graphs

an example schedule of messages resulting in computing \( p(x_4|x_1 = a) \):

<table>
<thead>
<tr>
<th>message direction</th>
<th>message value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( x_1 \to f_1 )</td>
<td>( \delta(x_1 = a) )</td>
</tr>
<tr>
<td>( x_3 \to f_2 )</td>
<td>( 1(x_3) )</td>
</tr>
<tr>
<td>( f_1 \to x_2 )</td>
<td>( \sum x_1 f_1(x_1, x_2) \delta(x_1 = a) = f_1(x_1 = a, x_2) )</td>
</tr>
<tr>
<td>( f_2 \to x_2 )</td>
<td>( \sum x_3 f_2(x_3, x_2) 1(x_3) )</td>
</tr>
<tr>
<td>( x_2 \to f_3 )</td>
<td>( f_1(x_1 = a, x_2) \left( \sum x_3 f_2(x_3, x_2) \right) )</td>
</tr>
<tr>
<td>( f_3 \to x_4 )</td>
<td>( \sum x_2 f_3(x_2, x_4) f_1(x_1 = a, x_2) \left( \sum x_3 f_2(x_3, x_2) \right) )</td>
</tr>
</tbody>
</table>

where \( \delta(x = a) \) is a delta function
Elimination Rules for Factor Graphs

- eliminating observed variables

If a variable $x_i$ is observed, i.e. its value is given, then it is a constant in all factor that include $x_i$.

We can eliminate $x_i$ from the graph by removing the corresponding node and modifying all neighboring factors to treat it as a constant.
Elimination Rules for Factor Graphs

- eliminating hidden variables

If a variable $x_i$ is hidden and we are not interested in it we can eliminate it from the graph by summing over all its values.

\[
\sum_{x_i} p(x) = \frac{1}{Z} \sum_{x_i} \prod_{j} f_j(x_{S_j})
\]

\[
= \frac{1}{Z} \prod_{j \notin n(x_i)} f_j(x_{S_j}) \left( \sum_{x_i} \prod_{k \in n(x_i)} f_k(x_{S_k}) \right)
\]

\[
= \frac{1}{Z} \prod_{j \notin n(x_i)} f_j(x_{S_j}) f_{\text{new}}(x_{S_{\text{new}}})
\]

where $f_{\text{new}}(x_{S_{\text{new}}}) = \sum_{x_i} \prod_{k \in n(x_i)} f_k(x_{S_k})$ and $S_{\text{new}} = \bigcup_{k \in n(x_i)} S_k \setminus \{i\}$.

This causes all its neighboring factor nodes to merge into one new factor node.
Inference in Hidden Markov Models and Linear Gaussian State-Space Models

- In HMMs, the states $X_t$ are discrete.
- In linear Gaussian SSMs, the states are real Gaussian vectors.
- Both HMMs and SSMs can be represented as singly connected DAGs.
- The forward-backward algorithm in hidden Markov models (HMMs), and the Kalman smoothing algorithm in SSMs are both instances of belief propagation / factor graph propagation.
Inference in multiply connected DAGs

The Junction Tree algorithm: Form an undirected graph from your directed graph such that no additional conditional independence relationships have been created (this step is called “moralization”). Lump variables in cliques together and form a tree of cliques—this may require a nasty step called “triangulation”. Do inference in this tree of cliques.

Cutset Conditioning: or “reasoning by assumptions”. Find a small set of variables which, if they were given (i.e. known) would render the remaining graph singly connected. For each value of these variables run belief propagation on the singly connected network. Average the resulting beliefs with the appropriate weights (given by normalizing constants).

Loopy Belief Propagation: just use BP although there are loops. In this case the terms “upstream” and “downstream” are not clearly defined. No guarantee of convergence, except for certain special graphs, but often works well in practice (c.f. “turbo-decoding” for error-correcting codes).
starting with a DAG...
moralize by marrying the parents of each node
remove edge directions
this results in an undirected graph with no additional conditional independence relations
triangulate so that there is no loop of length > 3 without a chord
this is necessary so that the final junction tree satisfies the running intersection property
find cliques of the moralized, triangulated graph
The Junction Tree Algorithm 5

- form **junction tree**: tree of (overlapping) sets of variables

- the **running intersection property** means that if a variable appears in more than one clique (e.g. $C$), it appears in all intermediate cliques in the tree.

- the junction tree propagation algorithm ensures that neighboring cliques have consistent probability distribution

- local consistency $\rightarrow$ global consistency
Summary

- inference consists of the problem of computing $p(\text{variables of interest}|\text{observed variables})$

- for singly connected DAGs, belief propagation solves this problem exactly.

- for factor graphs, the analogous algorithm is factor graph propagation.

- well-known algorithms such as Kalman smoothing and forward-backward are special cases these general propagation algorithms.

- for multiply connected graphs, the junction tree algorithm solves the exact inference problem, but can be very slow (exponential in the cardinality of the largest clique).

- one approximate inference algorithm is “loopy belief propagation”—we will see other approximate inference algorithms in a later lecture.
Appendix: Belief Propagation

top-down upstream evidence: (message $U_i$ sends to $X$)

$$\pi_X(U_i) = p(U_i|e_U^+_i X)$$

bottom-up downstream evidence: (message $Y_j$ sends to $X$)

$$\lambda_Y(X) = p(e_X^Y|X)$$

To update the probability of $X$ given the evidence:

$$\text{BEL}(X) = p(X|e) = \frac{1}{Z} \lambda(X) \pi(X)$$

$$\lambda(X) = \prod_j \lambda_{Y_j}(X)$$

$$\pi(X) = \sum_{U_1 \cdots U_n} p(X|U_1, \ldots, U_n) \prod_i \pi_X(U_i)$$
Belief Propagation (cont.)

top-down upstream evidence: (message $U_i$ sends to $X$)

$$\pi_X(U_i) = p(U_i|e_{U_iX}^+)$$

bottom-up downstream evidence: (message $Y_j$ sends to $X$)

$$\lambda_{Y_j}(X) = p(e_{XY_j}^-|X)$$

Bottom-up propagation, message $X$ sends to $U_i$:

$$\lambda_X(U_i) = \sum_X \lambda(X) \sum_{U_k:k \neq i} p(X|U_1, \ldots, U_n) \prod_{k \neq i} \pi_X(U_k)$$

Top-down propagation, message $X$ sends to $Y_j$:

$$\pi_{Y_j}(X) = \frac{1}{Z} \left[ \prod_{k \neq j} \lambda_{Y_k}(X) \right] \sum_{U_1 \ldots U_n} p(X|U_1, \ldots, U_n) \prod_i \pi_X(U_i) = \frac{1}{Z} \frac{\text{BEL}(X)}{\lambda_{Y_j}(X)}$$

$Z$ is the normaliser ensuring $\sum_X \pi_{Y_j}(X) = 1$
Appendix:

Fluffy and Moby: A Belief Propagation Demo
Fluffy = pet cat
Moby = pet fish

Fluffy ate Moby

Moby is dead

Fluffy is hungry

Fluffy's food bowl is full

Fluffy has flu

Fluffy has high temperature

1. Model Structure
2. Model Parameters

Fluffy = pet cat
Moby = pet fish

P(x₁=1)=0.01
Fluffy ate Moby

P(x₃=1|x₁=0)=0.1
P(x₃=1|x₁=1)=1.0

P(x₄=1|x₁,x₂)
x₁, x₂→x₄=1
0 0 0.9
0 1 0.1
1 0 0.1
1 1 0.01

P(x₂=1)=0.01
Fluffy has flu

P(x₅=1|x₂=0)=0.1
P(x₅=1|x₂=1)=0.9

P(x₆=1|x₄=0)=0.9
P(x₆=1|x₄=1)=0.1

Fluffy's food bowl is full
3. Propagating Evidence

1. Observe “Moby is dead”, i.e. $x_3 = 1$

2. Send $\lambda_{x_3}(x_1) \equiv p(e_{x_1 \rightarrow x_3} | x_1) = \begin{bmatrix} 0.1 \\ 1.0 \end{bmatrix}$ message $x_3 \rightarrow x_1$

3. $BEL(x_1 | x_3 = 1) = \frac{1}{Z} \begin{bmatrix} 0.99 \\ 0.01 \end{bmatrix} \odot \begin{bmatrix} 0.1 \\ 1.0 \end{bmatrix} = \begin{bmatrix} 0.91 \\ 0.09 \end{bmatrix}$
4. Propagating Evidence

4. Send $\pi_{x_4}(x_1) \equiv p(x_1|e^+_{x_1 \rightarrow x_4}) = \begin{bmatrix} 0.91 \\ 0.09 \end{bmatrix}$

5. Send $\pi_{x_4}(x_2) \equiv p(x_2|e^+_{x_2 \rightarrow x_4}) = p(x_2) = \begin{bmatrix} 0.99 \\ 0.01 \end{bmatrix}$ from $x_2 \rightarrow x_4$.

6. Compute $\pi(x_4) \equiv p(x_4|e^+_{x_4}) = \sum_{x_1, x_2} p(x_4|x_1, x_2)\pi_{x_4}(x_1)\pi_{x_4}(x_2) = \begin{bmatrix} 0.18 \\ 0.82 \end{bmatrix}$

7. $BEL(x_4|x_3 = 1) = \begin{bmatrix} 0.18 \\ 0.82 \end{bmatrix}$, whereas before observing $x_3 = 1$, $BEL(x_4) = \begin{bmatrix} 0.1 \\ 0.9 \end{bmatrix}$. 
5. Propagating Evidence

8. Observe “Fluffy’s Food Bowl is Full” \( x_6 = 1 \)!

9. Send \( \lambda_{x_6}(x_4) = \begin{bmatrix} 0.9 \\ 0.1 \end{bmatrix} \) message \( x_6 \to x_4 \)

10. \( BEL(x_4|x_3 = 1, x_6 = 1) = \frac{1}{Z} \begin{bmatrix} 0.18 \\ 0.82 \end{bmatrix} \odot \begin{bmatrix} 0.9 \\ 0.1 \end{bmatrix} = \begin{bmatrix} 0.66 \\ 0.34 \end{bmatrix} \)

11. Send \( \lambda_{x_4}(x_1) = \sum_{x_4} \lambda_{x_6}(x_4) \sum_{x_2} p(x_4|x_1, x_2) \pi_{x_4}(x_2) = \begin{bmatrix} 0.19 \\ 0.82 \end{bmatrix} \)

12. \( BEL(x_1|x_3 = 1, x_6 = 1) = \frac{1}{Z} \begin{bmatrix} 0.99 \\ 0.01 \end{bmatrix} \odot \begin{bmatrix} 0.1 \\ 1.0 \end{bmatrix} \odot \begin{bmatrix} 0.19 \\ 0.82 \end{bmatrix} = \begin{bmatrix} 0.70 \\ 0.30 \end{bmatrix} \) \( \Rightarrow \) Fluffy still innocent!
Appendix: Understanding BP equations

\[ p(X|e) = \text{BEL}(X) = \frac{1}{Z} \lambda(X) \pi(X) = p(e_X^-|X)p(X|e_X^+) \]  \hspace{1cm} (1)

\[ p(e_X^-|X) = \lambda(X) = \prod_j \lambda Y_j(X) = \prod_j p(e_{XY_j}^-|X) \]  \hspace{1cm} (2)

\[ p(X|e_X^+) = \pi(X) = \sum_{U_1\ldots U_n} p(X|U_1, \ldots, U_n) \prod_i \pi_X(U_i) \]  \hspace{1cm} (3)

\[ = \sum_{U_1\ldots U_n} p(X|U_1, \ldots, U_n) \prod_i p(U_i|e_{U_i}^+X) \]  \hspace{1cm} (4)

\( Z \) is a normalization constant.

All equations follow from the conditional independencies in the graph.