

CSC535: Probabilistic Graphical Models

Message Passing Inference

Prof. Jason Pacheco

Outline

- Sum-Product Belief Propagation
- Loopy Belief Propagation
- Variable Elimination
- Junction Tree Algorithm
- Max-Product Belief Propagation

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Sum-Product Belief Propagation

- Loopy Belief Propagation
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Why Graphical Models?

Structure simplifies both representation and computation



Representation Complex global phenomena arise by simpler-to-specify local interactions

<u>Computation</u> Inference / estimation depends only on subgraphs (e.g. dynamic programming, belief propagation, Gibbs sampling)

Suppose we have a chain graph...



...and want to calculate the marginal on B

$$P(D) = \sum_{a} \sum_{b} \sum_{c} P(a, b, c, D)$$

 \succ For K-valued variables this is $\mathcal{O}(K^3)$

For a Markov Chain on N variables calculating $P(X_N)$ takes $\mathcal{O}(K^{N-1})$

> We can do better by reordering operations...



Suppose we just care about marginal on D: $P(D) = \sum_{a} \sum_{b} \sum_{c} P(a)P(b \mid a)P(c \mid b)P(D \mid c)$ $= \sum_{c} P(D \mid c) \sum_{b} P(c \mid b) \sum_{a} P(a)P(b \mid a)$

(Distributive property)

$$= \sum_{c} P(D \mid c) \sum_{b} P(c \mid b) m_A(b)$$

$$=\sum_{c} P(D \mid c) m_B(c)$$

 $= m_C(D)$

Each message takes O(K²) time for total of O(3K²)

On a Markov Chain of N RVs takes O((N-1)K^2)



Convert Bayes net to MRF by ignoring local normalization:

 $P(A, B, C, D) \propto \psi(A)\psi(B, A)\psi(C, B)\psi(D, C)$



Convert Bayes net to MRF by ignoring local normalization:

 $P(A, B, C, D) \propto \psi(A)\psi(B, A)\psi(C, B)\psi(D, C)$

Repeat same procedure on MRF (we do not assume normalization):

$$P(D) \propto \sum_{c} \psi(c, D) \sum_{b} \psi(b, c) \sum_{a} \psi(a, b) \psi(a)$$

 $P(D) \propto \sum_{c} \psi(c, D) \sum_{b} \psi(b, c) m_A(b)$

 $P(D) \propto \sum_{c} \psi(c, D) m_B(c)$

 $P(D) \propto m_C(D)$

Markov Chain Revisited



Inference viewed as passing *messages* e.g. $B \rightarrow C$:

$$m_C(d) = \sum_c m_B(c) \psi(c,d)$$
 Incoming Message Compatibility Potential

- Only showed calculation of marginal at rightmost node
- Backward pass of messages calculates all marginals
- General inference on Markov chains called forward-backward alg.
- > Extension to other model structures called *sum-product algorithm*



Forward message:

$$\alpha_{n-1}(x_n) = \sum_{x_{n-1}} \alpha_{n-2}(x_{n-1})\psi(x_{n-1}, x_n)$$

Forward message:

$$\beta_{n+1}(x_n) = \sum_{x_{n+1}} \beta_{n+2}(x_{n+1})\psi(x_n, x_{n+1})$$

Marginal probability:

$$p(x_n) \propto \alpha_{n-1}(x_n)\beta_{n+1}(x_n)$$



Forward message:

$$\alpha_{n-1}(x_n) = \psi(x_n, y_n) \sum_{x_{n-1}} \alpha_{n-2}(x_{n-1}) \psi(x_{n-1}, x_n)$$

Forward message:

$$\beta_{n+1}(x_n) = \sum_{x_{n+1}} \beta_{n+2}(x_{n+1})\psi(x_n, x_{n+1})\psi(x_{n+1}, y_{n+1})$$

$$\begin{split} \alpha_{n-1}(x_n) &\propto p(y_1, \dots, y_n, x_n) \\ &= p(y_1, \dots, y_n \mid x_n) p(x_n) & (\text{Chain rule}) \\ &= p(y_n \mid x_n) p(y_1, \dots, y_{n-1} \mid x_n) p(x_n) & (\text{Conditional Independence}) \\ &= p(y_n \mid x_n) p(y_1, \dots, y_{n-1}, x_n) & (\text{Chain rule}) \\ &= p(y_n \mid x_n) \sum_{x_{n-1}} p(y_1, \dots, y_{n-1}, x_{n-1}, x_n) & (\text{Law of Total Probability}) \\ &= p(y_n \mid x_n) \sum_{x_{n-1}} p(y_1, \dots, y_{n-1}, x_{n-1}) p(x_n \mid x_{n-1}) & (\text{Chain rule + Conditional Independence}) \\ &\propto \psi(y_n, x_n) \sum_{x_{n-1}} \alpha_{n-2}(x_{n-1}) \psi(x_n, x_{n-1}) \end{split}$$

$$\begin{split} \beta_{n+1}(x_n) &\propto p(y_{n+1}, \dots, y_N \mid x_n) \\ &= \sum_{x_{n+1}} p(y_{n+1}, \dots, y_N, x_{n+1} \mid x_n) & \text{(Law of Total Probability)} \\ &= \sum_{x_{n+1}} p(y_{n+1}, \dots, y_N \mid x_n, x_{n+1}) p(x_{n+1} \mid x_n) & \text{(Chain rule)} \\ &= \sum_{x_{n+1}} p(y_{n+1}, \dots, y_N \mid x_{n+1}) p(x_{n+1} \mid x_n) & \text{(Conditional Independence)} \\ &= \sum_{x_{n+1}} p(y_{n+2}, \dots, y_N \mid x_{n+1}) p(y_{n+1} \mid x_{n+1}) p(x_{n+1} \mid x_n) & \text{(Chain rule)} \\ &\propto \sum_{x_{n+1}} \beta_{n+2}(x_{n+1}) \psi(x_{n+1}, y_{n+1}) \psi(x_n, x_{n+1}) \end{split}$$



Forward message gives the *filtered posterior*:

$$\alpha_{n-1}(x_n) \propto p(y_1, \dots, y_n, x_n) \propto p(x_n \mid y_1, \dots, y_n)$$

Smoothed posterior incorporates all observations:

$$p(x_n \mid y_1, \dots, y_N) \propto p(x_n \mid y_1, \dots, y_n) p(y_{n+1}, \dots, y_N \mid x_n)$$
$$\propto \alpha_{n-1}(x_n) \beta_{n+1}(x_n)$$



Forward-Backward extends to any tree-structured pairwise MRF

Marginal given by *incoming* messages (e.g. node C):

Pass messages from leavesto-root, then root-to-leaves

 $p(C) \propto \psi(C) m_A(C) m_F(C) m_G(C)$

Message updates depend only on Markov blanket...



Computational Complexity

$$m_{ts}(x_s) = \sum_{x_t} \psi_{st}(x_s, x_t) \psi_t(x_t) \prod_{k \in \Gamma(t) \setminus s} m_{kt}(x_t)$$
$$\phi(x_s, x_t)$$

For K-valued random variables X_s and X_t intermediate factor $\psi(x_s, x_t)$ is K-by-K matrix

Each message requires computation: $\mathcal{O}(K^2)$

There are |E| edges so total computation is: $\mathcal{O}(2|E|K^2)$



Non-Pairwise MRFs



Convert to tree-structured factor graph and redefine sumproduct messages

Notation Change

We will use slightly different notation for this section...



Sum-product extends to treestructured *factor graphs*

Key Observation

Any variable node X with N factors splits graph into N subgraphs with no shared variables

Approach

Recursively decompose into subtrees and marginalize them



Two kinds of computations marginalize different subtrees

Marginalize a sub-graph with a **variable node at its root** using the marginals of the sub-graphs attached to it.

Marginalize a sub-graph with a **factor node at its root** using the marginals of the sub-graphs attached to it.



Each root node (variable or factor) "waits" for all messages from its children before being marginalized out



Factor-to-variable message

Let X_s be the variables of the sub-graph attached to a factor, f_s (as root).

Denote the distribution of the sub-graph by $F_s(x, X_s)$

Define the factor-to-variable message from f_s to x by:



The message is the marginal of the subgraph with respect to all variables **except** *x*.

Variable-to-factor message

Let X_s be the variables in the sub-graph attached to a variable, x (as root).

Denote the distribution of the sub-graph by $G_s(x, X_s)$

Define the variable-to-factor message from x to f_s by:



The message is the marginal of the subgraph with respect to all variables **except** *x*.

What a variable node computes

The outgoing message to the factor, f_o , from x, is exactly the same marginal as the previous, except we exclude f_o .

$$\mu_{x \to f_o}(x) = \sum_{\mathbf{x}/x} \prod_{s \in ne(x)/f_o} F(x, X_s)$$



*This is **what** it computes, but not **how** it does it efficiently (i.e., as in the sum-product algorithm).

General variable node computation

The outgoing message to the factor, f_o , from x, is exactly the same marginal as the previous, except we exclude f_o .

$$\mu_{x \to f_o}(x) = \sum_{\mathbf{x}/x} \prod_{s \in ne(x)/f_o} F(x, X_s)$$



In the following, we will consider the first case, $\tilde{p}(x)$, but everything works the same for $\mu_{x \to f_o}(x)$.

What the **root** variable node computes



(*ne*(•) denotes neighbours)



Sum-product on a slide





Variable node x_m gathers messages, $\mu_{f_l \to x_m}$, and sends $\mu_{x_m \to f_s}(x_m) = \prod_{l \ni f_l \in n(x_m) \setminus f_s} \mu_{f_l \to x_m}(x_m)$

Factor f_s gathers messages $\mu_{x_m \to f_s}(x_m)$, and sends $\mu_{f_s \to x}(x) = \sum_{x_1} \sum_{x_2} \cdots \sum_{x_M} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in ne(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$

Marginal is product of incoming factor-to-variable messages:

$$p(x_m) \propto \prod_{f_l \in ne(x_m)} \mu_{f_l \to x_m}(x_m)$$

One point of confusion

The two products over messages look similar, but the first:

Variable node
$$x_m$$
 gathers messages, $\mu_{f_l \to x_m}$, and sends

$$\mu_{x_m \to f_s}(x_m) = \prod_{l \ni f_l \in n(x_m) \setminus f_s} \mu_{f_l \to x_m}(x_m)$$

is a product of vectors, each over the same variable, but the **second** has the variable as the index in the product:

Factor
$$f_s$$
 gathers messages $\mu_{x_m \to f_s}(x_m)$, and sends

$$\mu_{f_s \to x}(x) = \sum_{x_1} \sum_{x_2} \cdots \sum_{x_M} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in ne(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$$

One point of confusion (continued)

There are several ways to interpret the message product:

$$\mu_{f_s \to x}(x) = \sum_{x_1} \sum_{x_2} \cdots \sum_{x_M} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in ne(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$$

N-dimensional analogue of the outer product creates a tensor:



E.g. For two messages each element of the sum corresponding to (x, x_1, x_2) is $f(x, x_1, x_2) \cdot \mu_1(x_1) \cdot \mu_2(x_2)$

Computational Complexity

Factor f_s gathers messages $\mu_{x_m \to f_s}(x_m)$, and sends $\mu_{f_s \to x}(x) = \sum_{x_1} \sum_{x_2} \cdots \sum_{x_M} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in ne(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$ Intermediate factor $\phi(x, x_1, x_2, \dots, x_M)$

Assuming all variables are K-valued, intermediate factor with M+1 variables has $\mathcal{O}(K^{M+1})$ entries

Sum-product algorithm example

Let
$$\tilde{p}(\mathbf{x}) = f_a(x_1, x_2) f_b(x_2, x_3) f_c(x_2, x_4)$$



Sum-product algorithm example

Let
$$\tilde{p}(\mathbf{x}) = f_a(x_1, x_2) f_b(x_2, x_3) f_c(x_2, x_4)$$



The sum-product algorithm

First, pass messages from leaves to your chosen root node. If you want more than one marginal or plan to do other computation, store the results as you go.

Initialization: If leaf node is a variable node, then start with a unity message. If leaf node is factor, then start with the factor.





$$\mu_{x_{1} \to f_{a}}(x_{1}) = 1$$

$$\mu_{f_{a} \to x_{2}}(x_{2}) = \sum_{x_{1}} f_{a}(x_{1}, x_{2})$$

Recall the general case (don't confuse general variables with this example)

Factor f_s gathers messages $\mu_{x_m \to f_s}(x_m)$, and sends $\mu_{f_s \to x}(x) = \sum_{x_1} \sum_{x_2} \cdots \sum_{x_M} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in ne(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$


$$\mu_{x_4 \to f_c}(x_4) = 1$$

$$\mu_{f_c \to x_2}(x_2) = \sum_{x_4} f_c(x_2, x_4)$$

$$\mu_{x_{2} \to f_{b}}(x_{2}) = \mu_{f_{a} \to x_{2}}(x_{2})\mu_{f_{c} \to x_{2}}(x_{2})$$

$$\mu_{f_{b} \to x_{3}}(x_{3}) = \sum_{x_{2}} f_{b}(x_{2}, x_{3})\mu_{x_{2} \to f_{b}}(x_{2})$$

We now have the marginal at X_3 :

$$p(x_3) \propto \mu_{f_b \to x_3}(x_3)$$

Summary of messages from leaves to root

$$\mu_{x_{1} \to f_{a}}(x_{1}) = 1$$

$$\mu_{f_{a} \to x_{2}}(x_{2}) = \sum_{x_{1}} f_{a}(x_{1}, x_{2})$$

$$\mu_{x_{4} \to f_{c}}(x_{4}) = 1$$

$$\mu_{f_{c} \to x_{2}}(x_{2}) = \sum_{x_{4}} f_{c}(x_{2}, x_{4})$$

$$\mu_{x_{2} \to f_{b}}(x_{2}) = \mu_{f_{a} \to x_{2}}(x_{2})\mu_{f_{c} \to x_{2}}(x_{2})$$

$$\mu_{f_{b} \to x_{3}}(x_{3}) = \sum_{x_{2}} f_{b}(x_{2}, x_{3})\mu_{x_{2} \to f_{b}}(x_{2})$$





Next we want to set up for additional computations, we pass messages from root to leaves.

Candidate for the first and second ones?



Passing messages from root to leaves.

$$\mu_{x_3 \to f_b}(x_3) = 1$$

$$\mu_{f_b \to x_2}(x_2) = \sum_{x_3} f_b(x_2, x_3)$$



Candidate for third and fourth?



from going the other way.

$$\mu_{x_{2} \to f_{c}}(x_{2}) = \mu_{f_{a} \to x_{2}}(x_{2}) \mu_{f_{b} \to x_{2}}(x_{2})$$

$$\mu_{f_{c} \to x_{4}}(x_{4}) = \sum_{x_{2}} f_{c}(x_{2}, x_{4}) \mu_{x_{2} \to f_{c}}(x_{2})$$

(similar to previous one)

Summary of messages from root to leaves.

$$\mu_{x_{3} \to f_{b}}(x_{3}) = 1$$

$$\mu_{f_{b} \to x_{2}}(x_{2}) = \sum_{x_{3}} f_{b}(x_{2}, x_{3})$$

$$\mu_{x_{2} \to f_{a}}(x_{2}) = \mu_{f_{b} \to x_{2}}(x_{2})\mu_{f_{c} \to x_{2}}(x_{2})$$

$$\mu_{f_{a} \to x_{1}}(x_{1}) = \sum_{x_{2}} f_{a}(x_{1}, x_{2})\mu_{x_{2} \to f_{a}}(x_{2})$$

$$\mu_{x_{2} \to f_{c}}(x_{2}) = \mu_{f_{a} \to x_{2}}(x_{2})\mu_{f_{b} \to x_{2}}(x_{2})$$

$$\mu_{f_{c} \to x_{4}}(x_{4}) = \sum_{x_{2}} f_{c}(x_{2}, x_{4})\mu_{x_{2} \to f_{c}}(x_{2})$$





$$\tilde{p}(x_2) = \mu_{f_a \to x_2}(x_2) \mu_{f_b \to x_2}(x_2) \mu_{f_c \to x_2}(x_2)$$



$$\begin{split} \tilde{p}(x_{2}) &= \mu_{f_{a} \to x_{2}}(x_{2}) \mu_{f_{b} \to x_{2}}(x_{2}) \mu_{f_{c} \to x_{2}}(x_{2}) \\ &= \left(\sum_{x_{1}} f_{a}(x_{1}, x_{2}) \mu_{x_{1} \to f_{a}}(x_{1}) \right) \left(\sum_{x_{3}} f_{b}(x_{2}, x_{3}) \mu_{x_{3} \to f_{b}}(x_{1}) \right) \left(\sum_{x_{4}} f_{c}(x_{2}, x_{4}) \mu_{x_{4} \to f_{c}}(x_{1}) \right) \end{split}$$



$$\begin{split} \tilde{p}(x_{2}) &= \mu_{f_{a} \to x_{2}}(x_{2}) \mu_{f_{b} \to x_{2}}(x_{2}) \mu_{f_{c} \to x_{2}}(x_{2}) \\ &= \left(\sum_{x_{1}} f_{a}(x_{1}, x_{2}) \mu_{x_{1} \to f_{a}}(x_{1}) \right) \left(\sum_{x_{3}} f_{b}(x_{2}, x_{3}) \mu_{x_{3} \to f_{b}}(x_{1}) \right) \left(\sum_{x_{4}} f_{c}(x_{2}, x_{4}) \mu_{x_{4} \to f_{c}}(x_{1}) \right) \\ &= \left(\sum_{x_{1}} f_{a}(x_{1}, x_{2}) \right) \left(\sum_{x_{3}} f_{b}(x_{2}, x_{3}) \right) \left(\sum_{x_{4}} f_{c}(x_{2}, x_{4}) \right) \end{split}$$



$$\begin{split} \tilde{p}(x_{2}) &= \mu_{f_{a} \to x_{2}}(x_{2}) \mu_{f_{b} \to x_{2}}(x_{2}) \mu_{f_{c} \to x_{2}}(x_{2}) \\ &= \left(\sum_{x_{1}} f_{a}(x_{1}, x_{2}) \mu_{x_{1} \to f_{a}}(x_{1}) \right) \left(\sum_{x_{3}} f_{b}(x_{2}, x_{3}) \mu_{x_{3} \to f_{b}}(x_{1}) \right) \left(\sum_{x_{4}} f_{c}(x_{2}, x_{4}) \mu_{x_{4} \to f_{c}}(x_{1}) \right) \\ &= \left(\sum_{x_{1}} f_{a}(x_{1}, x_{2}) \right) \left(\sum_{x_{3}} f_{b}(x_{2}, x_{3}) \right) \left(\sum_{x_{4}} f_{c}(x_{2}, x_{4}) \right) \\ &= \sum_{x_{1}} \sum_{x_{3}} \sum_{x_{4}} f_{a}(x_{1}, x_{2}) f_{b}(x_{2}, x_{3}) f_{c}(x_{2}, x_{4}) \end{split}$$



 $\tilde{p}($

$$\begin{split} x_{2} &) = \mu_{f_{a} \to x_{2}} \left(x_{2} \right) \mu_{f_{b} \to x_{2}} \left(x_{2} \right) \mu_{f_{c} \to x_{2}} \left(x_{2} \right) \\ &= \left(\sum_{x_{1}} f_{a} \left(x_{1}, x_{2} \right) \mu_{x_{1} \to f_{a}} \left(x_{1} \right) \right) \left(\sum_{x_{3}} f_{b} \left(x_{2}, x_{3} \right) \mu_{x_{3} \to f_{b}} \left(x_{1} \right) \right) \left(\sum_{x_{4}} f_{c} \left(x_{2}, x_{4} \right) \mu_{x_{4} \to f_{c}} \left(x_{1} \right) \right) \right) \\ &= \left(\sum_{x_{1}} f_{a} \left(x_{1}, x_{2} \right) \right) \left(\sum_{x_{3}} f_{b} \left(x_{2}, x_{3} \right) \right) \left(\sum_{x_{4}} f_{c} \left(x_{2}, x_{4} \right) \right) \right) \\ &= \sum_{x_{1}} \sum_{x_{3}} \sum_{x_{4}} f_{a} \left(x_{1}, x_{2} \right) f_{b} \left(x_{2}, x_{3} \right) f_{c} \left(x_{2}, x_{4} \right) \\ &= \sum_{x_{1}} \sum_{x_{3}} \sum_{x_{4}} \tilde{p} \left(\mathbf{x} \right) \end{split}$$

 x_1

Outline

Sum-Product Belief Propagation

Loopy Belief Propagation

Variable Elimination

- > Junction Tree Algorithm
- Max-Product Belief Propagation

BP for Loopy Graphs

Suppose we have a graph with cycles...

Sum-product BP for tree-structured graphs relies on a leaf-to-root / rootto-leaf sequential update schedule

Graphs with cycles are "loopy" and have no obvious message ordering

Where do we even start? Every node requires initial messages...



BP for Loopy Graphs

Observe BP message update only depends on Markov Blanket:

$$m_{52}(x_2) = \sum_{x_5} \psi(x_2, x_5) \prod_{k \in \Gamma(5) \setminus 2} m_{k5}(x_5)$$

Where Γ is the set of neighbors:

 $\Gamma(s) = \{t : (s,t) \in \mathcal{E}\}$

Idea Initialize all messages (somehow) then iteratively update each message until "convergence".



What is convergence? Will this converge? If so, then to what?

Initialize Messages

Constant: $m_{st}^0(x_t) = \text{const.}$ Random: $m_{st}^0(x_t) \sim U([0,1])$

Parallel (Synchronous) Updates

At iteration *i* update *all messages in parallel* using current messages mⁱ⁻¹ from previous iteration:

$$m_{st}^{i}(x_t) = \sum_{x_s} \psi_{st}(x_s, x_t) \prod_{k \in \Gamma(s) \setminus t} m_{ks}^{i-1}(x_s)$$

- Store, both, the *previous* messages (from iteration *i*-1) and *current* messages (from iteration *i*)
- Many convergence results assume parallel updates



Initialize Messages

Constant: $m_{st}^0(x_t) = \text{const.}$ Random: $m_{st}^0(x_t) \sim U([0,1])$

Asynchronous (Sequential) Updates

$$m_{st}(x_t) = \sum_{x_s} \psi_{st}(x_s, x_t) \prod_{k \in \Gamma(s) \setminus t} m_{ks}(x_s)$$

- Simplifies updates since only need to keep track of one copy of messages
- Makes parallel processing trickier



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Initialize Messages

Constant: $m_{st}^0(x_t) = \text{const.}$ Random: $m_{st}^0(x_t) \sim U([0,1])$

Asynchronous (Sequential) Updates

Choose an ordering of nodes and update using the latest available messages:

$$m_{st}(x_t) = \sum_{x_s} \psi_{st}(x_s, x_t) \prod_{k \in \Gamma(s) \setminus t} m_{ks}(x_s)$$

- Simplifies updates since only need to keep track of one copy of messages
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Notice that each row can be computed in parallel

Initialize Messages

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Both directions are independent just like in forward-backward algorithm

Initialize Messages

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Upwards / downwards directions can also be done in parallel (holding rows fixed)

Pseudocode from Murphy's Textbook

Algorithm 22.1: Loopy belief propagation for a pairwise MRF

- 1 Input: node potentials $\psi_s(x_s)$, edge potentials $\psi_{st}(x_s, x_t)$;
- 2 Initialize messages $m_{s \to t}(x_t) = 1$ for all edges s t;
- 3 Initialize beliefs $bel_s(x_s) = 1$ for all nodes s;

4 repeat

5 Send message on each edge

$$m_{s \to t}(x_t) = \sum_{x_s} \left(\psi_s(x_s) \psi_{st}(x_s, x_t) \prod_{u \in \operatorname{nbr}_s \setminus t} m_{u \to s}(x_s) \right);$$

- 6 Update belief of each node $\operatorname{bel}_s(x_s) \propto \psi_s(x_s) \prod_{t \in \operatorname{nbr}_s} m_{t \to s}(x_s);$
- 7 **until** beliefs don't change significantly;
- 8 Return marginal beliefs $bel_s(x_s)$;

Loopy BP on Factor Graphs

Set of *neighbors* of node s: $\Gamma(s) = \{f \in \mathcal{F} \mid s \in f\}$

[Source: Erik Sudderth]

 $p_f(x_f) \propto \psi_f(x_f) \quad \overline{m}_{sf}(x_s)$

 $s \in f$



Marginal Distribution of Each Factor: *Clique of variables linked by factor.*

Low Density Parity Check (LDPC) Codes

Parity Check Factors



Evidence (observation) Factors



Each variable node is binary, so $x_s \in \{0, 1\}$

Parity check factors

equal 1 if the sum of the connected bits is even, 0 if the sum is odd (invalid codewords are excluded)

Unary evidence factors equal probability that each bit is a 0 or 1, given data. Assumes independent "noise" on each bit.

Loopy BP Convergence

Loopy BP works well empirically, but there are no guarantees:

- Not guaranteed to converge in general graphs
- BP marginal *beliefs* are **approximations**
- Empirically, when LBP converges it does so quickly and with good approximations

Convergence based on change in messages / marginal approximations:

$$\rho(m^{\text{old}}, m^{\text{current}}) < \epsilon \quad \text{or} \quad \rho(\text{bel}^{\text{old}}, \text{bel}^{\text{current}}) < \epsilon$$

Typical convergence measures are:

Max change: $\rho(m^{\text{old}}, m^{\text{current}}) = \max \{ |m^{\text{old}} - m^{\text{current}}| \}$ Total change: $\rho(m^{\text{old}}, m^{\text{current}}) = \sum |m^{\text{old}} - m^{\text{current}}|$

Loopy BP Convergence

Computation tree visualizes sequence of messages as BP proceeds...



Key Insight *T* iterations of BP equivalent to exact calculation in computation tree of height T+1. If edge strength sufficiently weak, then leaves will have minimal impact on root and BP converges.

Loopy BP Convergence

What can we do to improve convergence in a given model?

Message damping takes a *partial update* of messages each iteration,

$$m^{\text{new}} = (1 - \alpha)m^{\text{old}} + \alpha m^{\text{tmp}}$$

for damping factor $\alpha \in (0,1]$, e.g. $\alpha = 1$ is standard update

Message scheduling

- Asynchronous updates tend to converge faster than synchronous
- > Well-known Gauss-Seidel method does this in round-robin fashion (Bertsekas 97)
- Message update ordering also impacts convergence (e.g. disproportionate impact of nodes 2 & 3 in previous example)

Convergence depends largely on the existence of many small cycles

Example Ising model of ferromagnetism via atomic *spins:*

Binary *spin* variables: $x_i \in \{0, 1\}$

Interaction strength:

$$\psi_{ij} = \begin{pmatrix} \exp(J_{ij}) & \exp(-J_{ij}) \\ \exp(-J_{ij}) & \exp(J_{ij}) \end{pmatrix}$$

Field strength:

 $\psi_i = (\exp(h_i); \exp(-h_i))$



11x11 Ising model with random parameters



Source: D. Koller

Convergence of beliefs in 3 selected nodes



---- Synchronous — Asynchronous -- No Damping — True

Source: D. Koller

Oscillation in limit cycles is a typical failure mode of BP convergence



True

Source: D. Koller
Loopy BP Summary

- BP updates only depend on tree-structured Markov blanket
- **Approximate** BP inference in loopy graphs by iterating standard message updates until convergence (fixed point)
- No guarantees, but works well empirically in many instances
- Some techniques to improve convergence
 - Message damping
 - Asynchronous message update schedules

Outline

Sum-Product Belief Propagation

Loopy Belief Propagation

Variable Elimination

Junction Tree Algorithm

Max-Product Belief Propagation

Bayes Net → MRF





 $P(D, E, H, G, S, L, J) \propto \psi(D)\psi(E)\psi(G, D, E)$ $\psi(S, E)\psi(L, G)\psi(J, S, L)\psi(H, J, G)$

Нарру





$$\begin{split} P(D, E, H, G, S, L, J) &\propto \psi(D)\psi(E)\psi(G, D, E) \\ \psi(S, E)\psi(L, G)\psi(J, S, L)\psi(H, J, G) \end{split}$$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$



 $P(\mathbf{D}, E, H, G, S, L, J) \propto \psi(\mathbf{D})\psi(E)\psi(G, \mathbf{D}, \mathbf{E})$ $\psi(S, E)\psi(L, G)\psi(J, S, L)\psi(H, J, G)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$



$$\begin{split} P(E,H,G,S,L,J) \propto m_D(G,E)\psi(E) \\ \psi(S,E)\psi(L,G)\psi(J,S,L)\psi(H,J,G) \end{split}$$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate E (compute message $E \rightarrow (G,S)$): $m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$



 $P(\boldsymbol{E}, \boldsymbol{H}, \boldsymbol{G}, \boldsymbol{S}, \boldsymbol{L}, \boldsymbol{J}) \propto \boldsymbol{m_D}(\boldsymbol{G}, \boldsymbol{E}) \boldsymbol{\psi}(\boldsymbol{E})$ $\boldsymbol{\psi}(\boldsymbol{S}, \boldsymbol{E}) \boldsymbol{\psi}(\boldsymbol{L}, \boldsymbol{G}) \boldsymbol{\psi}(\boldsymbol{J}, \boldsymbol{S}, \boldsymbol{L}) \boldsymbol{\psi}(\boldsymbol{H}, \boldsymbol{J}, \boldsymbol{G})$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate E (compute message $E \rightarrow (G,S)$): $m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$ Eliminate H (compute message $H \rightarrow (G,J)$): $m_H(G,J) = \sum_h \psi(h,J,G)$



 $P(H, G, S, L, J) \propto m_E(G, S)\psi(L, G)$ $\psi(J, S, L)\psi(H, J, G)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$): $m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$ Eliminate **H** (compute message $H \rightarrow (G,J)$): $m_H(G,J) = \sum_h \psi(h,J,G)$



 $P(\boldsymbol{H}, \boldsymbol{G}, \boldsymbol{S}, \boldsymbol{L}, \boldsymbol{J}) \propto m_E(\boldsymbol{G}, \boldsymbol{S})\psi(\boldsymbol{L}, \boldsymbol{G})$ $\psi(\boldsymbol{J}, \boldsymbol{S}, \boldsymbol{L})\psi(\boldsymbol{H}, \boldsymbol{J}, \boldsymbol{G})$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate E (compute message $E \rightarrow (G,S)$): $m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$ Eliminate H (compute message $H \rightarrow (G,J)$): $m_H(G,J) = \sum_h \psi(h,J,G)$



 $P(G, S, L, J) \propto m_H(G, J) m_E(G, S) \psi(L, G)$ $\psi(J, S, L)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$):

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Eliminate **H** (compute message $H \rightarrow (G,J)$):

 $m_H(G,J) = \sum_h \psi(h,J,G)$



 $P(G, S, L, J) \propto m_H(G, J) m_E(G, S) \psi(L, G)$ $\psi(J, S, L)$

Eliminate **G**: $m_G(J, S, L) = \sum_g m_H(g, J) m_E(g, S) \psi(L, g)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$):

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Eliminate **H** (compute message $H \rightarrow (G,J)$):

	SAT
Letter	$m_G(J, S, L)$
	Job

 $P(S, L, J) \propto m_G(J, S, L)\psi(J, S, L)$

 $m_H(G,J) = \sum_h \psi(h,J,G)$ Eliminate **G**: $m_G(J,S,L) = \sum_g m_H(g,J)m_E(g,S)\psi(L,g)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$):

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Eliminate **H** (compute message $H \rightarrow (G,J)$):



```
P(S, L, J) \propto m_G(J, S, L)\psi(J, S, L)
```

 $m_H(G, J) = \sum_h \psi(h, J, G)$ Eliminate **G**: $m_G(J, S, L) = \sum_g m_H(g, J) m_E(g, S) \psi(L, g)$ Eliminate **S**: $m_S(J, L) = \sum_s m_G(J, s, L) \psi(J, s, L)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$):

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Eliminate **H** (compute message $H \rightarrow (G,J)$):



 $P(L,J) \propto m_S(J,L)$

 $m_H(G,J) = \sum_h \psi(h,J,G)$ Eliminate **G**: $m_G(J,S,L) = \sum_g m_H(g,J)m_E(g,S)\psi(L,g)$ Eliminate **S**: $m_S(J,L) = \sum_s m_G(J,s,L)\psi(J,s,L)$ Eliminate **L**: $m_L(J) = \sum_l m_S(J,l)$

Choose elimination ordering: D, E, H, G, S, L

Eliminate **D** (compute message $D \rightarrow (G,E)$):

 $m_D(G, E) = \sum_d \psi(d)\psi(d, G, E)$

Eliminate **E** (compute message $E \rightarrow (G,S)$):

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Eliminate **H** (compute message $H \rightarrow (G,J)$):



 $P(J) \propto m_l(J)$

 $m_H(G, J) = \sum_h \psi(h, J, G)$ Eliminate **G**: $m_G(J, S, L) = \sum_g m_H(g, J) m_E(g, S) \psi(L, g)$ Eliminate **S**: $m_S(J, L) = \sum_s m_G(J, s, L) \psi(J, s, L)$ Eliminate **L**: $m_L(J) = \sum_l m_S(J, l) \propto P(J)$

Accounting for Evidence

What if we observe a node (e.g. Letter=I)?

$$P(J \mid L = l) = \frac{P(J, L = l)}{P(L = l)}$$

Step 1: *Clamp* L = l in any factor with L:

$$\begin{split} P(D,E,H,G,S,L=l,J) \propto \psi(D)\psi(E)\psi(G,D,E) \\ \psi(S,E)\psi(L=l,G)\psi(J,S,L=l)\psi(H,J,G) \end{split}$$



Just treat these as new factors, since we don't care about normalizer:

 $\psi'(G) = \psi(L = l, G)$ and $\psi'(J, S) = \psi(J, S, L = l)$

Step 2: Remove L from elimination ordering

Main Points:

- Worst-case complexity of variable elimination is exponential in the number of latent variables
- Complexity is dependent on chosen elimination order

Consider eliminating **E** in the example...

$$m_E(G,S) = \sum_e m_D(G,e)\psi(e)\psi(S,e)$$

Multiplication creates intermediate factor:

$$\phi(S, G, E) = m_D(G, E)\psi(E)\psi(S, E)$$

Assuming all variables are K-valued, new factor $\phi(S, G, E)$ has K^3 entries requiring $\mathcal{O}(K^3)$



 $P(\boldsymbol{E}, \boldsymbol{H}, \boldsymbol{G}, \boldsymbol{S}, \boldsymbol{L}, \boldsymbol{J}) \propto \boldsymbol{m_D}(\boldsymbol{G}, \boldsymbol{E}) \boldsymbol{\psi}(\boldsymbol{E})$ $\boldsymbol{\psi}(\boldsymbol{S}, \boldsymbol{E}) \boldsymbol{\psi}(\boldsymbol{L}, \boldsymbol{G}) \boldsymbol{\psi}(\boldsymbol{J}, \boldsymbol{S}, \boldsymbol{L}) \boldsymbol{\psi}(\boldsymbol{H}, \boldsymbol{J}, \boldsymbol{G})$

Complexity determined by size of the largest intermediate factor

Elimination order D, E, H, G, S, L



 $\phi(D, E, G) = \mathcal{O}(K^3)$



 $\phi(E,G,S) = \mathcal{O}(K^3)$



 $\phi(E,G,S) = \mathcal{O}(K^3)$

Elimination order D, E, H, G, S, L



 $\phi(H,G,J) = \mathcal{O}(K^3)$

Elimination order D, E, H, G, S, L





$$\phi(G, S, L, J) = \mathcal{O}(K^4)$$

Elimination order D, E, H, G, S, L



$$\phi(S, L, J) = \mathcal{O}(K^3)$$

Elimination order D, E, H, G, S, L



 $\phi(L,J) = \mathcal{O}(K^2)$

Elimination order D, E, H, G, S, L



What if we choose a different elimination order?



 $\phi(L,J) = \mathcal{O}(K^2)$



 $\phi(G, D, E, L, H, J) = \mathcal{O}(K^6)$

The *induced graph* is the union of all graphs generated running variable elimination:

```
e.g. ordering D, E, H, G, S, L
```

Theorem (Informally) Given some elimination ordering:

- 1. Scope of every factor generated during variable elimination is a clique in the induced graph
- 2. Every **maximal clique** in the induced graph is a scope of some intermediate factor (of var. elim.)



Induced graph cliques Intermediate factors

Induced graph (and complexity) depend strongly on elimination order















Difficulty Effort Grade SAT Letter Job

- > Complexity of variable elimination is $\mathcal{O}(K^{w(\prec)+1})$
- > Lowest complexity given by the *treewidth:*

maximal cliques $\mathcal{C}(\prec)$ and *width*:

 $w(\prec) = \max_{c \in \mathcal{C}(\prec)} |c| - 1$

$$w^* = \min_{\prec} \max_{c \in \mathcal{C}(\prec)} |c| - 1$$

It is NP-hard to compute treewidth, and therefore an optimal elimination order (of course...)

Variable Elimination Summary

- > Variable elimination allows computation of marginals / conditionals
- > Algorithm is valid for **any graphical model**
- ➢ Suffices to show variable elimination for MRFs, since Bayes nets → MRFs by moralization
- Worst-case complexity is dependent on elimination order, and is exponential in number of variables
- Optimal ordering = treewidth, is NP-hard to compute

Outline

Variable Elimination

Sum-Product / Max-Product Algorithm

Junction Tree Algorithm

Loopy Belief Propagation

Some material adapted from Erik Sudderth lecture slides
Variable Elimination

Recall variable elimination sequentially marginalizes out variables...



Variable Elimination

Two major limitations of variable elimination:

- 1. Computation **exponential** in size of the largest intermediate factor (equivalently, largest clique in clique tree)
- 2. Computation is not reused for computing a series of marginals
- **E.g.** Suppose we use variable elimination to compute a marginal on an **HMM** with T nodes, each being K-valued
 - It takes $\mathcal{O}(TK^2)$ time to compute a single marginal
 - It takes $\mathcal{O}(T^2K^2)$ time to compute **all marginals**
 - We know forward-backward computes all marginals in $\mathcal{O}(TK^2)$

Marginal Inference Algorithms

One Marginal	All Marginals
Elimination applied to leaves of tree	Belief Propagation (BP) or sum-product algorithm
Variable Elimination	Junction Tree Algorithm BP on a junction tree (special clique tree)

Graph

Marginal Inference Algorithms

One Marginal	All Marginals
Elimination applied to leaves of tree	Belief Propagation (BP) or sum-product algorithm
Variable Elimination	Junction Tree Algorithm BP on a junction tree
	(special clique tree)

Graph

Elimination order: 6,5,4,3,2,1

Clique Tree









Elimination order: 6,5,4,3,2,1

Clique Tree







Junction Tree

Definition (Running intersection) For any pair of clique nodes V,W all cliques on the *unique path* between V and W contain shared variables



A junction tree is a clique tree with the running intersection property

Junction Tree

Clique tree edges are separator sets in original MRF...so clique tree encodes conditional independencies



Theorem A clique tree resulting from variable elimination satisfies the running intersection property and is thus a junction tree

Junction Trees and Triangulation



- A *chord* is an edge connecting two non-adjacent nodes in some *cycle*
- A cycle is *chordless* if it contains no chords
- A graph is *triangulated (chordal)* if it contains no chordless cycles of length 4 or more

Theorem: The maximal cliques of a graph have a corresponding junction tree *if and only if* that undirected graph is triangulated

Lemma: For a non-complete triangulated graph with at least 3 nodes, there is a decomposition of the nodes into disjoint sets A, B, S such that S separates A from B, and S is complete.

- > Key induction argument in constructing junction tree from triangulation
- Implies existence of *elimination ordering which introduces no new edges*

Induced Graph

Recall the **induced graph** is the union over intermediate graphs from running variable elimination

The induced graph is chordal thus:

- Maximal cliques of the induced graph form a junction tree
- It admits an elimination ordering that introduces *no new edges*
- Logic of junction tree algorithm:
 - 1. Triangulate the graph
 - a. Implies a junction tree
 - b. Induces an elimination order
 - 2. Run sum-product BP on junction tree to compute all clique marginals



Reminder: Pairwise Sum-Product BP





 $\begin{array}{rcl} K_t & \longrightarrow & number \ of \ discrete \ states \ for \ random \ variable \ x_t \\ p_t(x_t) & \longrightarrow & marginal \ distribution \ of \ the \ K_t \ discrete \ states \ of \ random \ variable \ x_t \\ m_{st}(x_t) & \longrightarrow & message \ from \ node \ s \ to \ node \ t, \ a \ vector \ of \ K_t \ non-negative \ numbers \\ m_{ts}(x_s) & \longrightarrow & message \ from \ node \ t \ to \ node \ s, \ a \ vector \ of \ K_s \ non-negative \ numbers \end{array}$

- Express algorithm via original variables x_s
- Messages depend on clique intersection (separators)
- Efficient schedules compute each message once



 X_2X_4

- Let x_{C_i} be variables in clique node C_j
- Let $x_{S_{ij}}$ be variables in separator such that:

$$x_{S_{ij}} = x_{C_i} \cap x_{C_j}$$

Let *residual* variables be:



 C_5

- Let x_{C_i} be variables in clique node C_j
- Let $x_{S_{ij}}$ be variables in separator such that:

 $x_{S_{ij}} = x_{C_i} \cap x_{C_j}$

Let residual variables be:

 $x_{R_{ij}} = x_{C_i} \backslash x_{S_{ij}}$

 Pass sum-product messages between clique nodes



Message:
$$m_{ji}(x_{S_{ji}}) \propto \sum_{x_{R_{ji}}} \psi_{C_j}(x_{C_j}) \prod_{k \in \Gamma(j) \setminus i} m_{kj}(x_{S_{kj}})$$

Marginal: $p_j(x_{C_j}) \propto \psi_{C_j}(x_{C_j}) \prod_{i \in \Gamma(j)} m_{ij}(x_{S_{ij}})$

S₁₂

S₂₃

 C_2

 C_3

 S_{34}

 C_5

S₄₅

 C_4

 C_6

S₄₆

- Express algorithm via original variables x_s
- Messages depend on clique intersection (separators)
- Efficient schedules compute each message once

Storage & Computational Cost

$$\mathcal{O}\left(\sum_{j}\prod_{s\in C_j}K_s\right)$$
, where $x_s\in\{1,\ldots,K_s\}$

Exponential in sizes of maximal cliques.

Message:
$$m_{ji}(x_{S_{ji}}) \propto \sum_{x_{R_{ji}}} \psi_{C_j}(x_{C_j}) \prod_{k \in \Gamma(j) \setminus i} m_{kj}(x_{S_{kj}})$$

Marginal: $p_j(x_{C_j}) \propto \psi_{C_j}(x_{C_j}) \prod_{i \in \Gamma(j)} m_{ij}(x_{S_{ij}})$

Summary: Junction Tree Algorithm



Junction Tree Algorithms for General-Purpose Inference

- 1. If necessary, convert graphical model to undirected form (*linear in graph size*)
- 2. Triangulate the target undirected graph
- > Any elimination ordering generates a valid triangulation (*linear in graph size*)
- > Finding an optimal triangulation, with minimal cliques, is *NP-hard*
- 3. Arrange triangulated cliques into a junction tree (at worst quadratic in graph size)
- 4. Execute sum-product algorithm on junction tree (*exponential in clique size*)

Message Passing Inference Summary

- Brute-force enumeration exponential regardless of graph
- Sum-Product BP
 - Exact inference in tree-structure graphs in O(TK²) time for T nodes, each taking K states
 - Reduces to Forward-Backward in HMMs
 - Same for Max-Product BP (reduces to Viterbi in HMMs)
- Variable elimination
 - Exact marginals in general graphs
 - Worst-case complexity exponential in size of largest clique
 - Need to rerun from scratch for each marginal
 - Complexity dependent on elimination order (NP-hard to optimize)

Message Passing Inference Summary

- Junction Tree Algorithm
 - Exact marginals in general graphs
 - Caches messages to compute all marginals
 - Worst-case complexity exponential in size of largest clique
 - Optimizing Jtree is NP-hard (corresponds to finding treewidth)
- Loopy BP: Just did this, did you forget already?

Maximum A Posteriori (MAP) Inference



Rather than marginalize sometimes we want to maximize, e.g.

$$(x_1^*, x_2^*, \dots, x_N^*) = \arg\max_{\mathbf{x}} p(\mathbf{x} \mid \mathbf{y})$$

Maximizing the log-joint is equivalent and numerically more stable:

$$(x_1^*, x_2^*, \dots, x_N^*) = \arg\max_{\mathbf{x}} \log p(\mathbf{x}, \mathbf{y}) + \text{const.}$$

Forward-Backward Algorithm

Recall the Forward-Backward algorithm messages...



Forward message:

$$\alpha_{n-1}(x_n) = \sum_{x_{n-1}} \alpha_{n-2}(x_{n-1})\psi(x_{n-1}, x_n)\psi(x_n, y_n)$$

Sum over state x_{n-1}

Maximize instead of marginalize...



Forward message:

$$\alpha_{n-1}(x_n) = \max_{x_{n-1}} \log \psi(x_n, y_n) + \alpha_{n-2}(x_{n-1}) + \log \psi(x_{n-1}, x_n)$$

Maximize over state x_{n-1} (in log-domain)

Maximize instead of marginalize...



Forward message:

$$\alpha_{n-1}(x_n) = \max_{x_{n-1}} \log \psi(x_n, y_n) + \alpha_{n-2}(x_{n-1}) + \log \psi(x_{n-1}, x_n)$$

We also store the argmax values:

$$x_{n-1}^*(x_n) = \arg\max_{x_{n-1}} \log\psi(x_n, y_n) + \alpha_{n-2}(x_{n-1}) + \log\psi(x_{n-1}, x_n)$$

Maximize instead of marginalize...



Final node gives maximum (up to const.) and maximizer of posterior:

$$\alpha_{N-1}(x_N) = \max_{x_1,\dots,x_{N-1}} \log p(x_1,\dots,x_N \mid \mathbf{y}) + \text{const.}$$

$$x_{N-1}^*(x_N) = \underset{x_1,\dots,x_{N-1}}{\arg\max} \log p(x_1,\dots,x_N \mid \mathbf{y}) + \text{const.}$$

Backwards pass reads off joint maximizer...



Backward Pass: $x_n^* = x_n^*(x_{n+1}^*)$

Joint maximizing sequence obtained at the end of backwards pass:

$$(x_1^*, x_2^*, \dots, x_N^*) = \arg\max_{\mathbf{x}} p(\mathbf{x} \mid \mathbf{y})$$

Max-Product (Max-Sum) Algorithm

Recall our decomposition of factor graph sub-trees...



Max-product on a slide





Variable x_m gathers incoming messages and sends:

$$\mu_{x_m \to f_s(x_m)} = \prod_{f_l \in \operatorname{ne}(x_m) \setminus f_s} \mu_{f_l \to x_m}(x_m)$$

Factor f_s gathers incoming messages and sends: $\mu_{f_s \to x}(x) = \max_{\mathbf{x} \setminus x} f_s(x, x_1, x_2, \dots, x_M) \prod_{m \in \operatorname{ne}(f_s) \setminus x} \mu_{x_m \to f_s}(x_m)$

Max-sum on a slide





Variable x_m gathers incoming messages and sends:

 $\log \mu_{x_m \to f_s(x_m)} = \sum_{f_l \in \operatorname{ne}(x_m) \setminus f_s} \log \mu_{f_l \to x_m}(x_m) \qquad \log \mu_{f_s \to x}(x) = \max \log f_s(x_m)$

Factor f_s gathers incoming messages and sends: $g \mu_{f_s \to x}(x) =$

$$\max_{\mathbf{x}\setminus x} \log f_s(x, x_1, x_2, \dots, x_M) + \sum_{m \in \operatorname{ne}(f_s)\setminus x} \log \mu_{x_m \to f_s}(x_m)$$

More numerically stable to work in log-domain (max-sum)...

Max-sum Example



Max-sum Example

- At the root we can record the argmax for its variable, but we do not know which variable choices produced it
 - Ties have the potential to make this particularly complicated
- We can "backtrack" to find this out provided that we stored what we need in the forward pass.
- If there are ties, they need to be handled consistently
 - In our example, we need to choose either $x_5 = 0$ or $x_5 = 1$ for both backtracking branches.



If we choose x₅=0, then we need a maximal configuration for x₅=0 for both pieces for a **consistent joint maximizing configuration**

The factor nodes must store enough information to evaluate any choice

WRONG



The configuration that we get backtracking pretending $x_5 = 0$, even though $x_5 = 1$ cannot compute to more than 0.1, and could be less, as the settings for the other variables are making the value as big as possible when $x_5 = 0$.








Message Passing Inference Summary

Forward-backward algorithm yields efficient marginal inference on HMM graph

G



 X_2

 $\beta_2(x_1)$

 $\alpha_1(x_2)$

 X_1

 $\beta_{N-1}(x_{N-2})$

 $\alpha_{N-2}(x_{N-1})$

 $\beta_N(x_{N-1})$

 $\alpha_{N-1}(x_N)$

 X_N

 Y_N

 $\beta_3(x_2)$

 $\alpha_2(x_3)$

Max-product / max-sum yields maximum a posteriori (MAP) inference in any treestructured model

F

В

D

Ε

Viterbi decoder is special case for HMM



And factor graphs