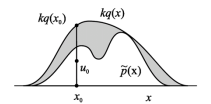


Probabilistic Graphical Models

Approximate Inference: Monte Carlo methods



Eric Xing Lecture 16, March 16, 2015



Reading: See class website

Approaches to inference



Exact inference algorithms

- The elimination algorithm
- Message-passing algorithm (sum-product, belief propagation)
- The junction tree algorithms

Approximate inference techniques

- Variational algorithms
 - Loopy belief propagation
 - Mean field approximation
- Stochastic simulation / sampling methods
- Markov chain Monte Carlo methods

How to represent a joint, or a marginal distribution?



- Closed-form representation
 - E.g., $(x_1, \dots, x_p)^T \sim \frac{1}{(2\pi)^{p/2} |\Sigma|^{1/2}} \exp\left(-\frac{1}{2}x \mu\right)^T \Sigma^{-1} (x \mu)\right)$

$$E_p(f(x)) = \int f(x)p(x)dx$$

• Sample-based representation:

Monte Carlo methods

- Draw random samples from the desired distribution
- Yield a stochastic representation of a complex distribution
 - marginals and other expections can be approximated using sample-based averages

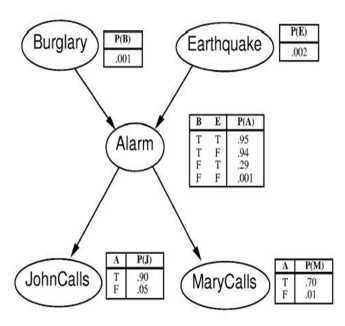
$$E[f(x)] = \frac{1}{N} \sum_{t=1}^{N} f(x^{(t)})$$

- Asymptotically exact and easy to apply to arbitrary models
- Challenges:
 - how to draw samples from a given dist. (not all distributions can be trivially sampled)?
 - how to make better use of the samples (not all sample are useful, or eqally useful, see an example later)?
 - how to know we've sampled enough?



Example: naive sampling

Construct samples according to probabilities given in a BN.



Alarm example: (Choose the right sampling sequence) 1) Sampling:P(B)=<0.001, 0.999> suppose it is false, B0. Same for E0. P(A|B0, E0)=<0.001, 0.999> suppose it is false...

2) Frequency counting: In the samples right, P(J|A0)=P(J,A0)/P(A0)=<1/9, 8/9>.

E0	В0	A0	MO	J0
E0	B0	A0	MO	J0
E0	B0	A0	MO	J1
E0	B0	A0	MO	J0
E0	B0	A0	MO	J0
E0	B0	A0	MO	J0
E1	В0	A1	M1	J1
E0	B0	A0	MO	J0
E0	В0	A0	MO	J0
E0	В0	A0	MO	J0

Example: naive sampling

Construct samples according to probabilities given in a BN.

Alarm example: (Choose the right sampling sequence)

- 3) what if we want to compute P(J|A1)? we have only one sample ... P(J|A1)=P(J,A1)/P(A1)=<0, 1>.
- 4) what if we want to compute P(J|B1)?

 No such sample available!

 P(J|A1)=P(J,B1)/P(B1) can not be defined.

For a model with hundreds or more variables, rare events will be very hard to garner evough samples even after a long time or sampling ...

E0	В0	A0	MO	J0
E0	В0	A0	MO	J0
E0	В0	A0	MO	J1
E0	В0	A0	MO	J0
E0	В0	A0	MO	J0
E0	В0	A0	MO	J0
E1	B0	A1	M1	J1
E0	В0	A0	MO	J0
E0	В0	A0	MO	J0
E0	В0	A0	MO	J0





Direct Sampling

- We have seen it.
- Very difficult to populate a high-dimensional state space

Rejection Sampling

- Create samples like direct sampling, only count samples which is consistent with given evidences.
- Likelihood weighting, ...
 - Sample variables and calculate evidence weight. Only create the samples which support the evidences.
- Markov chain Monte Carlo (MCMC)
 - Metropolis-Hasting
 - Gibbs

Rejection sampling

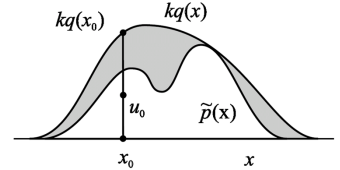
- Suppose we wish to sample from dist. $\Pi(X)=\Pi'(X)/Z$.
 - $\Pi(X)$ is difficult to sample, but $\Pi'(X)$ is easy to **evaluate**
 - Sample from a simpler dist Q(X)
 - Rejection sampling

$$x^* \sim Q(X)$$
, accept x^* w.p. $\Pi'(x^*)/kQ(x^*)$

Correctness:

$$p(x) = \frac{\left[\Pi'(x)/kQ(x)\right]Q(x)}{\int \left[\Pi'(x)/kQ(x)\right]Q(x)dx}$$
$$= \frac{\Pi'(x)}{\int \Pi'(x)dx} = \Pi(x)$$

Pitfall ...

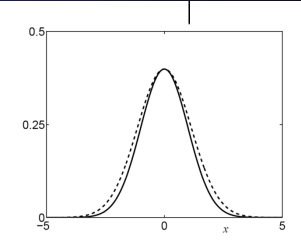


Rejection sampling



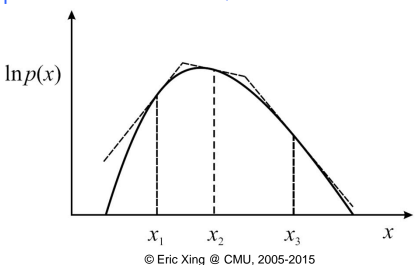
Pitfall:

- Using Q= $\mathcal{N}(\mu, \sigma_q^{2/d})$ to sample P= $\mathcal{N}(\mu, \sigma_p^{2/d})$
- If σ_q exceeds σ_p by 1%, and dimensional=1000,
- The optimal acceptance rate $k=(\sigma_q/\sigma_p)^d\approx 1/20,000$
- Big waste of samples!



Adaptive rejection sampling

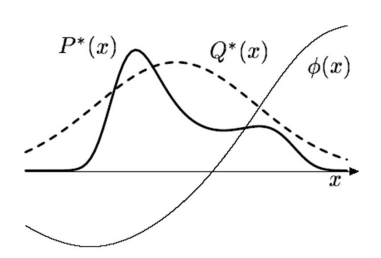
Using envelope functions to define Q



Unnormalized importance sampling



- Suppose sampling from $P(\cdot)$ is hard.
- Suppose we can sample from a "simpler" proposal distribution
 Q(-) instead.
- If Q dominates P (i.e., Q(x) > 0 whenever P(x) > 0), we can sample from Q and reweight:



$$\langle f(X) \rangle = \int f(x) P(x) dx$$

$$= \int f(x) \frac{P(x)}{Q(x)} Q(x) dx$$

$$\approx \frac{1}{M} \sum_{m} f(x^{m}) \frac{P(x^{m})}{Q(x^{m})} \quad \text{where } x^{m} \sim Q(X)$$

$$= \frac{1}{M} \sum_{m} f(x^{m}) w^{m}$$

What is the problem here?

Normalized importance sampling

- Suppose we can only evaluate $P'(x) = \alpha P(x)$ (e.g. for an MRF).
- We can get around the nasty normalization constant α as follows:

• Let
$$r(X) = \frac{P'(x)}{Q(x)}$$
 $\Rightarrow \langle r(X) \rangle_Q = \int \frac{P'(x)}{Q(x)} Q(x) dx = \int P'(x) dx = \alpha$

Now

$$\langle f(X) \rangle_{p} = \int f(x)P(x)dx = \frac{1}{\alpha} \int f(x) \frac{P'(x)}{Q(x)}Q(x)dx$$

$$= \frac{\int f(x)r(x)Q(x)dx}{\int r(x)Q(x)dx}$$

$$\approx \frac{\sum_{m} f(x^{m})r^{m}}{\sum_{m} r^{m}} \quad \text{where } x^{m} \sim Q(X)$$

$$= \sum_{m} f(x^{m})w^{m} \quad \text{where } w^{m} = \frac{r^{m}}{\sum_{m} r^{m}}$$

Normalized vs unnormalized importance sampling



Unormalized importance sampling is unbiased:

$$E_{\mathcal{Q}}[f(X)w(X)] =$$

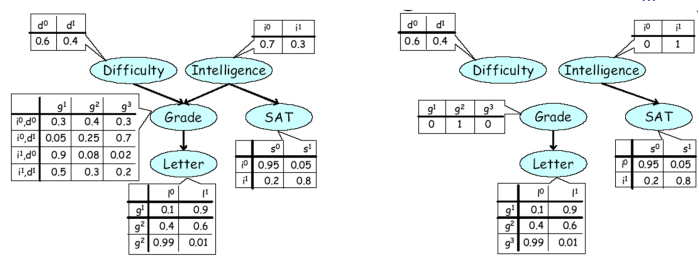
Normalized importance sampling is biased, e.g., for M = 1:

$$E_{\mathcal{Q}}\left[\frac{f(x^1)r(x^1)}{r(x^1)}\right] =$$

- However, the variance of the normalized importance sampler is usually lower in practice.
- Also, it is common that we can evaluate P'(x) but not P(x), e.g. P(x|e) = P'(x, e)/P(e) for Bayes net, or P(x) = P'(x)/Z for MRF.

Likelihood weighting

- We now apply normalized importance sampling to a Bayes net.
- The proposal Q is gotten from the mutilated BN where we clamp evidence nodes, and cut their incoming arcs. Call this P_M.



The unnormalized posterior is P'(x) = P(x, e). So for $f(X_i) = \delta(X_i = x_i)$, we get $\hat{P}(X_i = x_i | e) = \frac{\sum_m w_m \delta(x_i^m = x_i)}{\sum_{w_m} w_w}$ where $w_m = P'(x^m, e) / P_M(x^m)$.



Likelihood weighting algorithm

```
[x_{1:n}, w] = \text{function LW(CPDs, } G, E)
let X_1, \ldots, X_n be a topological ordering of G
w = 1
x = (0, \dots, 0)
for i = 1:n
   let u_i = x(Pa_i)
   if X_i \not\in E
   then sample x_i from P(X_i|u_i)
   else
        x_i = e(X_i)
        w = w * P(x_i|u_i)
```

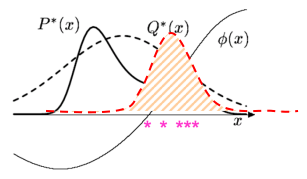
Efficiency of likelihood weighting



- The efficiency of importance sampling depends on how close the proposal Q is to the target P.
- Suppose all the evidence is at the roots. Then Q = P(X|e), and all samples have weight 1.
- Suppose all the evidence is at the leaves. Then Q is the prior, so many samples might get small weight if the evidence is unlikely.
- We can use arc reversal to make some of the evidence nodes be roots instead of leaves, but the resulting network can be much more densely connected.

Weighted resampling

- Problem of importance sampling: depends on how well Q matches P
 - If P(x)f(x) is strongly varying and has a significant proportion of its mass concentrated in a small region, r_m will be dominated by a few samples



• Note that if the high-prob mass region of Q falls into the low-prob mass region of P, the variance of $r^m = P(x^m)/Q(x^m)$ can be small even if the samples come from low-prob region of P and potentially erroneous .

Solution

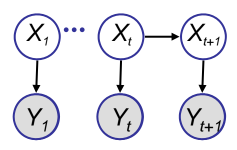
- Use heavy tail Q.
- Weighted resampling

$$w^{m} = \frac{P(x^{m})/Q(x^{m})}{\sum_{l} P(x^{l})/Q(x^{l})} = \frac{r^{m}}{\sum_{m} r^{m}}$$

Weighted resampling



- Sampling importance resampling (SIR):
 - Draw N samples from $Q: X_1 \dots X_N$
 - Constructing weights: $w_1 \dots w_N$, $w^m = \frac{P(x^m)/Q(x^m)}{\sum_{l} P(x^l)/Q(x^l)} = \frac{r^m}{\sum_{m} r^m}$ Sub-sample x from $\{X_1 \dots X_N\}$ w.p. $(w_1 \dots w_N)$
- Particular Filtering
 - A special weighted resampler
 - Yield samples from posterior $p(X_t|Y_{1:t})$
 - Also known as sequential Monte Carlo



Sketch of Particle Filters

The starting point

$$p(X_{t}|Y_{1t}) = p(X_{t}|Y_{t}, Y_{1t-1}) = \frac{p(X_{t}|Y_{1t-1})p(Y_{t}|X_{t})}{\int p(X_{t}|Y_{1t-1})p(Y_{t}|X_{t})dX_{t}}$$

• Thus $p(X_t|Y_{1:t})$ is represented by

$$\left\{ X_{t}^{m} \sim p(X_{t} \mid Y_{1t-1}), \quad w_{t}^{m} = \frac{p(Y_{t} \mid X_{t}^{m})}{\sum\limits_{m=1}^{M} p(Y_{t} \mid X_{t}^{m})} \right\}$$

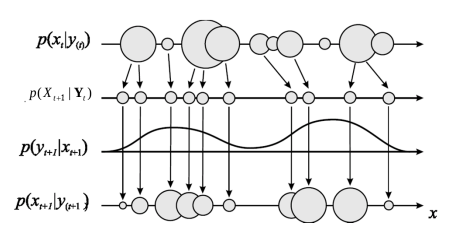
- A sequential weighted resampler
 - Time update

$$p(X_{t+1} | Y_{1t}) = \int p(X_{t+1} | X_t) p(X_t | Y_{1t}) dX_t$$

- $= \sum_{m} w_{t}^{m} p(X_{t+1} | X_{t}^{(m)})$ (sample from a mixture model)
 - Measurement update

$$p(X_{t+1}|Y_{1t+1}) = \frac{p(X_{t+1}|Y_{1t})p(Y_{t+1}|X_{t+1})}{\int p(X_{t+1}|Y_{1t})p(Y_{t+1}|X_{t+1})dX_{t+1}}$$

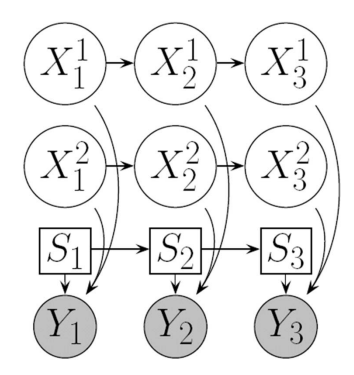
$$\Rightarrow \begin{cases} X_{t+1}^{m} \sim p(X_{t+1}|Y_{1t}), & W_{t+1}^{m} = \frac{p(Y_{t+1}|X_{t+1}^{m})}{\sum\limits_{m=1}^{M} p(Y_{t+1}|X_{t+1}^{m})} \end{cases}$$
 (reweight)







• Recall that the belief state has O(2t) Gaussian modes

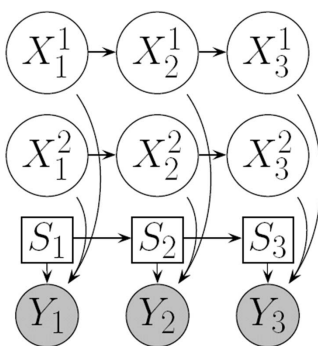


PF for switching SSM



 Key idea: if you knew the discrete states, you can apply the right Kalman filter at each time step.

- So for each old particle m, sample $S_t^m \sim P(S_t \mid S_{t-1}^m)$ from the prior, apply the KF (using parameters for S_t^m) to the old belief state $(\hat{x}_{t-1|t-1}^m, P_{t-1|t-1}^m)$ to get an approximation to $P(X_t \mid y_{1:t}, s_{1:t}^m)$
- Useful for online tracking, fault diagnosis, etc.



Rao-Blackwellised sampling

- Sampling in high dimensional spaces causes high variance in the estimate.
- RB idea: sample some variables X_p , and conditional on that, compute expected value of rest X_d analytically:

$$\begin{split} E_{p(X|e)}[f(X)] &= \int p(x_{p}, x_{d} \mid e) f(x_{p}, x_{d}) dx_{p} dx_{d} \\ &= \int_{x_{p}} p(x_{p} \mid e) \left(\int_{x_{d}} p(x_{d} \mid x_{p}, e) f(x_{p}, x_{d}) dx_{d} \right) dx_{p} \\ &= \int_{x_{p}} p(x_{p} \mid e) E_{p(X_{d} \mid x_{p}, e)} [f(x_{p}, X_{d})] dx_{p} \\ &= \frac{1}{M} \sum_{m} E_{p(X_{d} \mid x_{p}^{m}, e)} [f(x_{p}^{m}, X_{d})], \qquad x_{p}^{m} \sim p(x_{p} \mid e) \end{split}$$

This has lower variance, because of the identity:

$$\operatorname{var}\left[\tau(X_{p}, X_{d})\right] = \operatorname{var}\left[E\left[\tau(X_{p}, X_{d}) \mid X_{p}\right]\right] + E\left[\operatorname{var}\left[\tau(X_{p}, X_{d}) \mid X_{p}\right]\right]$$



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• Hence $\operatorname{var}\left[E\left[\tau(X_p,X_d)\mid X_p\right]\right] \leq \operatorname{var}\left[\tau(X_p,X_d)\right]$, so $\tau(X_p,X_d) = E\left[f(X_p,X_d)\mid X_p\right]$ is a lower variance estimator.

Summary: Monte Carlo Methods



- Direct Sampling
 - Very difficult to populate a high-dimensional state space
- Rejection Sampling
 - Create samples like direct sampling, only count samples which is consistent with given evidences.
- Likelihood weighting, ...
 - Sample variables and calculate evidence weight. Only create the samples which support the evidences.
- Markov chain Monte Carlo (MCMC)
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