Machine Learning and Bayesian Inference

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Part IV

Bayesian networks

Markov random fields

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Uncertainty: Probability as Degree of Belief

At the start of the course, I presented a *uniform approach* to *knowledge representation and reasoning* using *probability*.



The world is represented by RVs  $\mathbf{V} = \{V_1, V_2, \dots, V_n\}$ . These are partitioned:

- 1. Query variables  $\mathbf{Q} = \{Q_1, Q_2, \dots, Q_q\}$ . We want to *compute a distribution* over these.
- 2. Observed variables  $O = \{o_1, o_2, \dots, o_m\}$ . We know the values of these.
- 3. Latent variables  $\mathbf{L} = \{L_1, L_2, \dots, L_l\}$ . Everything else.

General knowledge representation and inference: the BIG PICTURE

The latent variables L are all the RVs not in the sets Q or O.



To compute a conditional distribution from a knowledge base Pr(V) we have to *sum over the latent variables* 

$$\Pr\left(\mathbf{Q}|o_{1}, o_{2}, \dots, o_{m}\right) = \sum_{\mathbf{L}} \Pr\left(\mathbf{Q}, \mathbf{L}|o_{1}, o_{2}, \dots, o_{m}\right)$$
$$= \boxed{\frac{1}{Z} \sum_{\mathbf{L}} \underbrace{\Pr\left(\mathbf{Q}, \mathbf{L}, o_{1}, o_{2}, \dots, o_{m}\right)}_{\text{Knowledge base}}}$$

### General knowledge representation and inference: the BIG PICTURE

*Bayes' theorem* tells us how to update an inference when *new information* is available.



For example, if we now receive a new observation O' = o' then

$$\underbrace{\Pr\left(\mathbf{Q}|o', o_1, o_2, \dots, o_m\right)}_{\text{After }O' \text{ observed}} = \frac{1}{Z} \Pr\left(o'|\mathbf{Q}, o_1, o_2, \dots, o_m\right) \underbrace{\Pr\left(\mathbf{Q}|o_1, o_2, \dots, o_m\right)}_{\text{Before }O' \text{ observed}}$$

Simple eh?

HAH!!! No chance...

Even if all your RVs are just Boolean:

- For *n* RVs knowing the knowledge base  $Pr(\mathbf{V})$  means storing  $2^n$  numbers.
- So it looks as though storage is  $O(2^n)$ .
- You need to establish  $2^n$  numbers to work with.
- Look at the summations. If there are n latent variables then it appears that time complexity is also  $O(2^n)$ .
- In reality we might well have n > 1000, and of course it's even worse if variables are non-Boolean.

And it *really is this hard*. The problem in general is *#P-complete*.

Even getting an *approximate solution* is provably intractable.

Having seen that in principle, if not in practice, the full joint distribution alone can be used to perform any inference of interest, we now examine a *practical technique*.

- We introduce the *Bayesian Network (BN)* as a compact representation of the full joint distribution.
- We examine the way in which a BN can be *constructed*.
- We examine the *semantics* of BNs.
- We look briefly at how *inference* can be performed.
- We briefly introduce the *Markov random field (MRF)* as an alternative means of representing a distribution.

A brief aside on the dangers of interpreting *implication* versus *conditional probability*:

- $\Pr(X = x | Y = y) = 0.1$  does *not* mean that if Y = y is then  $\Pr(X = x) = 0.1$ .
- $\Pr(X)$  is a *prior probability*. It applies when you *haven't seen* the value of Y.
- The notation  $\Pr(X|Y = y)$  is for use when y is the *entire evidence*.
- $\Pr(X|Y = y \land Z = z)$  might be very different.

Conditional probability is *not* analogous to *logical implication*.

# Implication and conditional probability

# In general, it is difficult to relate *implication* to *conditional probability*.

 $Pr(A \rightarrow B) = Pr(\neg A \lor B)$   $Pr(A|B) = \frac{Pr(A \land B)}{Pr(B)}$ 

Imagine that fish are very rare, and most fish can swim.

With implication,

$$\Pr\left(\texttt{fish} \to \neg\texttt{swim}\right) = \Pr\left(\neg\texttt{fish} \lor \neg\texttt{swim}\right) = LARGE$$
 With conditional probability,

$$\Pr\left(\neg\texttt{swim}|\texttt{fish}\right) = \frac{\Pr\left(\neg\texttt{swim} \land \texttt{fish}\right)}{\Pr\left(\texttt{fish}\right)} = \texttt{SMALL!}$$

One of the key reasons for the introduction of *Bayesian networks* is to let us *exploit independence*.

The initial pay-off is that this *makes it easier to represent* Pr(V).

A further pay-off is that it *introduces structure* that can lead to *more efficient inference*.

Here is a *very simple* example.

If I toss a coin and roll a die, the full joint distribution of outcomes requires  $2 \times 6 = 12$  numbers to be specified.

	•	•	•		••• ••	
H	0.014	0.028	0.042	0.057	0.071	0.086
T	0.033	0.067	0.1	0.133	0.167	0.2

Here  $\Pr(\text{Coin} = H) = 0.3$  and the die has probability i/21 for the *i*th outcome.

BUT: if we assume the outcomes are independent then Pr(Coin, Dice) = Pr(Coin) Pr(Dice)Where Pr(Coin) has two numbers and Pr(Dice) has six. So instead of 12 numbers we only need 8. A slightly more complex example:

	(	CP		¬CP
	SB	¬SB	SB	ΓSB
HD	0.024	0.006	0.016	0.004
¬HD	0.0019	0.0076	0.1881	0.7524

- HD = Heart disease
- CP = Chest pain
- SB = Shortness of breath

Similarly, say instead of just considering HD, SB and CP we also consider the outcome of the *Oxford versus Cambridge tiddlywinks competition* TC:

 $TC = \{Oxford, Cambridge, Draw\}.$ 

Now

 $Pr\left(\texttt{HD},\texttt{SB},\texttt{CP},\texttt{TC}\right) = Pr\left(\texttt{TC}|\texttt{HD},\texttt{SB},\texttt{CP}\right)Pr\left(\texttt{HD},\texttt{SB},\texttt{CP}\right).$ 

Assuming that the patient is not an *extraordinarily keen fan of tiddlywinks*, their cardiac health has nothing to do with the outcome, so

 $Pr\left(\texttt{TC}|\texttt{HD},\texttt{SB},\texttt{CP}\right) = Pr\left(\texttt{TC}\right)$ 

and  $2 \times 2 \times 2 \times 3 = 24$  numbers has been reduced to 3 + 8 = 11.

However although in this case we might not be able to exploit independence directly we *can* say that

$$\Pr\left(\texttt{CP},\texttt{SB}|\texttt{HD}\right) = \Pr\left(\texttt{CP}|\texttt{HD}\right)\Pr\left(\texttt{SB}|\texttt{HD}\right)$$

which simplifies matters.

*Conditional independence:*  $A \perp B | C$ 

• A is conditionally independent of B given C, written  $A \perp B | C$ , if

 $\Pr\left(A,B|C\right) = \Pr\left(A|C\right)\Pr\left(B|C\right).$ 

- If we know that C is the case then A and B are independent.
- Equivalently  $\Pr(A|B, C) = \Pr(A|C)$ . (Prove this!)

Although CP and SB are *not* independent, they do not directly influence one another *in a patient known to have heart disease*.

This is much nicer!

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\Pr\left(\text{HD}|\text{CP},\text{SB}\right) \propto \Pr\left(\text{CP}|\text{HD}\right)\Pr\left(\text{SB}|\text{HD}\right)\Pr\left(\text{HD}\right)
```

After a *regrettable incident* involving an *inflatable gorilla*, a famous College has decided to install an alarm for the detection of roof climbers.

- The alarm is *very* good at detecting climbers.
- Unfortunately, it is also sometimes triggered when one of the *extremely fat geese* that lives in the College lands on the roof.
- One porter's lodge is near the alarm, and inhabited by a chap with *excellent hearing* and a *pathological hatred* of roof climbers: he *always* reports an alarm. His hearing is so good that he sometimes thinks he hears an alarm, *even when there isn't one*.
- Another porter's lodge is a good distance away and inhabited by an *old chap* with *dodgy hearing* who likes to listen to his collection of *DEATH METAL* with the sound turned up.



Also called *probabilistic/belief/causal networks* or *knowledge maps*.

- Each node is a *random variable (RV)*.
- Each node  $N_i$  has a distribution

 $\Pr(N_i | \texttt{parents}(N_i))$ 

- A Bayesian network is a *directed acyclic graph*.
- Roughly speaking, an arrow from N to M means N directly affects M.

*Note that:* 

- In the present example all RVs are *discrete* (in fact Boolean) and so in all cases  $Pr(N_i | parents(N_i))$  can be represented as a *table of numbers*.
- Climber and Goose have only *prior* probabilities.
- All RVs here are Boolean, so a node with p parents requires  $2^p$  numbers.

A BN with n nodes represents the full joint probability distribution for those nodes as

$$\Pr(N_1 = n_1, N_2 = n_2, \dots, N_n = n_n) = \prod_{i=1}^n \Pr(N_i = n_i | \text{parents}(N_i)).$$

For example

 $\Pr(\neg \mathsf{C}, \neg \mathsf{G}, \mathsf{A}, \mathsf{L1}, \mathsf{L2}) = \Pr(\mathsf{L1}|\mathsf{A}) \Pr(\mathsf{L2}|\mathsf{A}) \Pr(\mathsf{A}|\neg \mathsf{C}, \neg \mathsf{G}) \Pr(\neg \mathsf{C})) \Pr(\neg \mathsf{G})$  $= 0.99 \times 0.6 \times 0.08 \times 0.95 \times 0.8.$ 

In general  $\Pr(A, B) = \Pr(A|B) \Pr(B)$  so

$$\Pr(N_1, ..., N_n) = \Pr(N_n | N_{n-1}, ..., N_1)$$
  $\Pr(N_{n-1}, ..., N_1)$ 

Repeating this gives

$$\Pr(N_1, \dots, N_n) = \Pr(N_n | N_{n-1}, \dots, N_1) \left| \Pr(N_{n-1} | N_{n-2}, \dots, N_1) \cdots \Pr(N_1) \right|$$
$$= \prod_{i=1}^n \Pr(N_i | N_{i-1}, \dots, N_1).$$

Now compare equations. We see that BNs make the assumption

$$\Pr(N_i|N_{i-1},\ldots,N_1) = \Pr(N_i|\operatorname{parents}(N_i))$$

for each node, assuming that parents $(N_i) \subseteq \{N_{i-1}, \ldots, N_1\}$ .

Each  $N_i$  is conditionally independent of its predecessors given its parents.

- When constructing a BN we want to make sure the preceding property holds.
- This means we need to take care over *ordering*.
- In general *causes should directly precede effects*.



Here, parents( $N_i$ ) contains all preceding nodes having a *direct influence* on  $N_i$ .

But its not quite that straightforward: what if we want to talk about nodes *other than predecessors and parents*?

For example, it is possible to show:



Any node A is conditionally independent of the  $N_i$ —its non-descendants—given the  $P_i$ —its parents.

It is also possible to show:



Any node A is conditionally independent of all other nodes given the *Markov* blanket  $M_i$ —that is, its parents, its children and its childrens' parents.

### Semantics: what's REALLY going on here?

There is a general method for inferring exactly what conditional independences are implied by a Bayesian network.

Let X, Y and Z be disjoint subsets of the RVs.

Consider a *path* p consisting of directed (in any orientation) edges from some  $x \in X$  to some  $y \in Y$ . For example



The path p is said to be *blocked* by Z if one of *three conditions* holds...

Semantics: what's REALLY going on here?

Path p is *blocked* with respect to Z if:

1. p contains a node  $z \in Z$  that is *tail-to-tail*:



2. p contains a node  $z \in Z$  that is *head-to-tail*:



(Similarly if the node is *tail-to-head*.)

3. *p* contains a node *N* that is *head-to-head*,  $N \notin Z$ , and none of *N*'s descendents is in *Z*:



### Semantics: what's REALLY going on here?

Finally:

1. X and Y are *d*-separated by Z if all paths p from some  $x \in X$  to some  $y \in Y$  are blocked.

2. If X and Y are *d*-separated by Z then  $X \perp Y | Z$ .

# More complex nodes

How do we represent

 $\Pr(N_i | \text{parents}(N_i))$ 

when nodes can denote general discrete and/or continuous RVs?

- BNs containing both kinds of RV are called hybrid BNs.
- Naive *discretisation* of continuous RVs tends to result in both a reduction in accuracy and large tables.
- $O(2^p)$  might still be large enough to be unwieldy.
- We can instead attempt to use *standard and well-understood* distributions, such as the *Gaussian*.
- This will typically require only a small number of parameters to be specified.

*Example:* a continuous RV with one continuous and one discrete parent.

 $\Pr(\text{Speed of car}|\text{Throttle position}, \text{Tuned engine})$ where SC and TP are continuous and TE is Boolean.

• For a specific setting of ET = true it might be the case that SC increases with TP, but that some uncertainty is involved

$$\Pr(\mathsf{SC}|\mathsf{TP},\mathsf{et}) = N(g_{\mathsf{et}}\mathsf{TP} + c_{\mathsf{et}},\sigma_{\mathsf{et}}^2).$$

• For an un-tuned engine we might have a similar relationship with a different behaviour

$$\Pr(\mathsf{SC}|\mathsf{TP},\neg\mathsf{et}) = N(g_{\neg\mathsf{et}}\mathsf{TP} + c_{\neg\mathsf{et}}, \sigma_{\neg\mathsf{et}}^2).$$

There is a set of parameters  $\{g, c, \sigma\}$  for each possible value of the discrete RV.

*Example:* a discrete RV with a continuous parent

 $\Pr\left(\text{Go roofclimbing}|\text{Size of fine}\right).$ 

We could for example use the *probit distribution* 

$$\Pr\left(\text{Go roofclimbing} = \text{true}|\text{size}\right) = \Phi\left(\frac{t - \text{size}}{s}\right)$$

where

$$\Phi(x) = \int_{-\infty}^x N(y) dy$$

and N is the Gaussian density with zero mean and variance 1.

# More complex nodes



#### Basic inference

We saw earlier that the full joint distribution can be used to perform *all inference tasks*:

$$\Pr\left(\mathbf{Q}|o_1, o_2, \dots, o_m\right) = \frac{1}{Z} \sum_{\mathbf{L}} \Pr\left(\mathbf{Q}, \mathbf{L}, o_1, o_2, \dots, o_m\right)$$

where

- Q is the query.
- $o_1, o_2, \ldots, o_m$  are the observations.
- L are the latent variables.
- 1/Z normalises the distribution.
- The query, observations and latent variables are a partition of the set  $\mathbf{V} = \{V_1, V_2, \dots, V_n\}$  of all variables.

#### **Basic inference**

As the BN fully describes the full joint distribution

$$\Pr\left(\mathbf{Q}, \mathbf{L}, o_1, o_2, \dots, o_m\right) = \prod_{i=1}^n \Pr(V_i | \text{parents}(V_i))$$

it can be used to perform inference in the obvious way

$$\Pr(\mathbf{Q}|o_1, o_2, \dots, o_m) \propto \sum_{\mathbf{L}} \prod_{i=1}^n \Pr(V_i| \text{parents}(V_i))$$

but this is in practice problematic for obvious reasons.

- More sophisticated algorithms aim to achieve this *more efficiently*.
- For complex BNs we resort to *approximation techniques*.

 $Pr(\mathbf{Q}, \mathbf{L}, o_1, \dots, o_m)$  has a particular form expressing conditional independences:



 $\Pr\left(C,G,A,L1,L2\right)=\Pr\left(C\right)\Pr\left(G\right)\Pr\left(A|C,G\right)\Pr\left(L1|A\right)\Pr\left(L2|A\right).$ 

Consider the computation of the query  $\Pr(C|l1, l2)$ 

We have

$$\Pr\left(C|l1,l2\right) \propto \sum_{A} \sum_{G} \Pr\left(C\right) \Pr\left(G\right) \Pr\left(A|C,G\right) \Pr\left(l1|A\right) \Pr\left(l2|A\right).$$

Here there are 5 multiplications for each set of values that appears for summation, and there are 4 such values.

In general this gives time complexity  $O(n2^n)$  for n Boolean RVs.

The naive implementation of this approach yields the *Enumerate-Joint-Ask* algorithm, which unfortunately requires  $O(2^n)$  time and space for n Boolean RVs.

The *enumeration-ask* algorithm improves matters to  $O(2^n)$  time and O(n) space by performing the computation *depth-first*.

However matters can be improved further by avoiding *duplication of computations*. Looking more closely we see that

$$\begin{split} \Pr\left(C|l1,l2\right) &\propto \sum_{A} \sum_{G} \Pr\left(C\right) \Pr\left(G\right) \Pr\left(A|C,G\right) \Pr\left(l1|A\right) \Pr\left(l2|A\right) \\ &= \frac{1}{Z} \Pr\left(C\right) \sum_{A} \Pr\left(l1|A\right) \Pr\left(l2|A\right) \sum_{G} \Pr\left(G\right) \Pr\left(A|C,G\right) \\ &= \frac{1}{Z} \Pr\left(C\right) \sum_{G} \Pr\left(G\right) \sum_{A} \Pr\left(A|C,G\right) \Pr\left(l1|A\right) \Pr\left(l2|A\right). \end{split}$$

There is some freedom in terms of how we *factorize* the expression.

This is a result of introducing assumptions about conditional independence.

Taking the second possibility:

$$\underbrace{\Pr\left(C\right)}_{C}\sum_{G}\underbrace{\Pr\left(G\right)}_{G}\sum_{A}\underbrace{\Pr\left(A|C,G\right)}_{A}\underbrace{\Pr\left(l1|A\right)}_{L1}\underbrace{\Pr\left(l2|A\right)}_{L2}$$

where C, G, A, L1, L2 denote the relevant factors.

The basic idea is to evaluate this from right to left (or in terms of the tree, bottom up) *storing results* as we progress and *re-using them* when necessary.

 $\Pr(l1|A)$  depends on the value of A. We store it as a table  $\mathbf{F}_{L1}(A)$ . Similarly for  $\Pr(l2|A)$ .

$$\mathbf{F}_{L1}(A) = \begin{pmatrix} 0.99\\ 0.08 \end{pmatrix} \mathbf{F}_{L2}(A) = \begin{pmatrix} 0.6\\ 0.001 \end{pmatrix}$$

as  $\Pr(l1|a) = 0.99$ ,  $\Pr(l1|\neg a) = 0.08$  and so on.

Similarly for Pr(A|C, G), which is dependent on A, C and G



Can we write  $\Pr(A|C, G) \Pr(l1|A) \Pr(l2|A)$  as  $\mathbf{F}_A(A, C, G) \mathbf{F}_{L1}(A) \mathbf{F}_{L2}(A)$ 

in a reasonable way?

Yes, provided multiplication of factors is defined correctly. Looking at

$$\Pr\left(C\right)\sum_{G}\Pr\left(G\right)\sum_{A}\Pr\left(A|C,G\right)\Pr\left(l1|A\right)\Pr\left(l2|A\right)$$

note that:

1. The values of the product

```
\Pr\left(A|C,G\right)\Pr\left(l1|A\right)\Pr\left(l2|A\right)
```

in the summation over A depend on the values of C and G external to it, and the values of A.

2. So

```
\mathbf{F}_A(A, C, G)\mathbf{F}_{L1}(A)\mathbf{F}_{L2}(A)
```

should be a table collecting values where correspondences between RVs are maintained.

This leads to a definition for *multiplication of factors* best given by example.

$$\mathbf{F}(A,B)\mathbf{F}(B,C) = \mathbf{F}(A,B,C)$$

where

A	B	$\mathbf{F}(A,B)$	B	C	$\mathbf{F}(B,C)$	A	B	C	$\mathbf{F}(A, B, C)$
$\top$	Τ	0.3		Τ	0.1		Τ	Τ	$0.3 \times 0.1$
	$\bot$	0.9		$\bot$	0.8		Т	$\bot$	$0.3 \times 0.8$
	$\top$	0.4		$\top$	0.8		$\bot$	$\top$	$0.9 \times 0.8$
	$\bot$	0.1		$\bot$	0.3		$\bot$	$\bot$	$0.9 \times 0.3$
							$\top$	$\top$	$0.4 \times 0.1$
							Т	$\bot$	$0.4 \times 0.8$
							$\bot$	$\top$	$0.1 \times 0.8$
							$\bot$	$\bot$	$0.1 \times 0.3$

This process gives us



How about

$$\mathbf{F}_{\overline{A},L1,L2}(C,G) = \sum_{A} \mathbf{F}_{A}(A,C,G) \mathbf{F}_{L1}(A) \mathbf{F}_{L2}(A)$$

To denote the fact that A has been summed out we place a bar over it in the notation.

$$\sum_{A} \mathbf{F}_{A}(A, C, G) \mathbf{F}_{L1}(A) \mathbf{F}_{L2}(A) = \mathbf{F}_{A}(a, C, G) \mathbf{F}_{L1}(a) \mathbf{F}_{L2}(a) + \mathbf{F}_{A}(\neg a, C, G) \mathbf{F}_{L1}(\neg a) \mathbf{F}_{L2}(\neg a)$$

where

$$\mathbf{F}_{A}(a,C,G) = \begin{bmatrix} C & G \\ \top & \top & 0.98 \\ \top & \bot & 0.96 \\ \bot & \top & 0.2 \\ \bot & \bot & 0.08 \end{bmatrix} \mathbf{F}_{L1}(a) = 0.99 \quad \mathbf{F}_{L2}(a) = 0.6$$

and similarly for  $\mathbf{F}_A(\neg a, C, G)$ ,  $\mathbf{F}_{L1}(\neg a)$  and  $\mathbf{F}_{L2}(\neg a)$ .



Now, say for example we have  $\neg c, g$ . Then doing the calculation explicitly would give

$$\begin{split} \sum_{A} \Pr(A|\neg c,g) \Pr(l1|A)) &\Pr(l2|A) \\ &= \Pr(a|\neg c,g) \Pr(l1|a) \Pr(l2|a) + \Pr(\neg a|\neg c,g) \Pr(l1|\neg a) \Pr(l2|\neg a) \\ &= (0.2 \times 0.99 \times 0.6) + (0.8 \times 0.08 \times 0.001) \end{split}$$

which matches!

Continuing in this manner form

$$\begin{split} \mathbf{F}_{G,\overline{A},L1,L2}(C,G) &= \mathbf{F}_G(G)\mathbf{F}_{\overline{A},L1,L2}(C,G) \\ \text{sum out } G \text{ to obtain } \mathbf{F}_{\overline{G},\overline{A},L1,L2}(C) &= \sum_G \mathbf{F}_G(G)\mathbf{F}_{\overline{A},L1,L2}(C,G), \text{ form} \\ \mathbf{F}_{C,\overline{G},\overline{A},L1,L2} &= \mathbf{F}_C(C)\mathbf{F}_{\overline{G},\overline{A},L1,L2}(C) \end{split}$$

and normalise.

What's the computational complexity now?

- For Bayesian networks with *suitable structure* we can perform inference in *linear time and space*.
- However in the worst case it is still *#P-hard*.

Consequently, we may need to resort to *approximate inference*.

*Markov chain Monte Carlo (MCMC)* methods also provide a method for performing *approximate inference* in *Bayesian networks*.

Say a system can be in a state S and moves from state to state in discrete time steps according to a probabilistic transition

$$\Pr\left(\mathbf{S} \to \mathbf{S}'\right)$$
.

Let  $\pi_t(\mathbf{S})$  be the probability distribution for the state after t steps, so

$$\pi_{t+1}(\mathbf{S}') = \sum_{\mathbf{s}} \Pr\left(\mathbf{s} \to \mathbf{S}'\right) \pi_t(\mathbf{s}).$$

If at some point we obtain  $\pi_{t+1}(\mathbf{s}) = \pi_t(\mathbf{s})$  for all  $\mathbf{s}$  then we have reached a *stationary distribution*  $\pi$ . In this case

$$\forall \mathbf{s}' \pi(\mathbf{s}') = \sum_{\mathbf{s}} \Pr\left(\mathbf{s} \to \mathbf{s}'\right) \pi(\mathbf{s}).$$

There is exactly one stationary distribution for a given  $Pr(S \rightarrow S')$  provided the latter obeys some simple conditions.

The condition of *detailed balance* 

$$\forall \mathbf{s}, \mathbf{s}' \pi(\mathbf{s}) \Pr\left(\mathbf{s} \to \mathbf{s}'\right) = \pi(\mathbf{s}') \Pr\left(\mathbf{s}' \to \mathbf{s}\right)$$

is sufficient to provide a  $\pi$  that is a stationary distribution. To see this simply sum:

$$\sum_{\mathbf{s}} \pi(\mathbf{s}) \Pr(\mathbf{s} \to \mathbf{s}') = \sum_{\mathbf{s}} \pi(\mathbf{s}') \Pr(\mathbf{s}' \to \mathbf{s})$$
$$= \pi(\mathbf{s}') \underbrace{\sum_{\mathbf{s}} \Pr(\mathbf{s}' \to \mathbf{s})}_{=1}$$
$$= \pi(\mathbf{s}')$$

If all this is looking a little familiar, it's because we now have another excellent application for the material in *Mathematical Methods for Computer Science*.

That course used the alternative term *local balance*.

Approximate inference for Bayesian networks

Recalling once again the basic equation for performing probabilistic inference

$$\Pr\left(\mathbf{Q}|o_1, o_2, \dots, o_m\right) \propto \sum_{\mathbf{L}} \Pr\left(\mathbf{Q}, \mathbf{L}, o_1, o_2, \dots, o_m\right)$$

where

- Q is the query.
- $o_1, o_2, \ldots, o_m$  are the observations.
- L are the latent variables.
- 1/Z normalises the distribution.
- The query, observations and latent variables are a partition of the set  $\mathbf{V} = \{V_1, V_2, \dots, V_n\}$  of all variables.

We are going to consider obtaining samples from the distribution  $Pr(\mathbf{Q}, \mathbf{L}|o_1, o_2, \dots, o_m).$  Approximate inference for Bayesian networks

The observations are fixed. Let the *state* of our system be a specific set of values for *a query variable and the latent variables* 

$$\mathbf{S} = (S_1, S_2, \dots, S_{l+1}) = (Q, L_1, L_2, \dots, L_l)$$

and define  $\overline{\mathbf{S}}_i$  to be the state vector with  $S_i$  removed

$$\overline{\mathbf{S}}_i = (S_1, \ldots, S_{i-1}, S_{i+1}, \ldots, S_{n+1}).$$

To move from s to s' we replace one of its elements, say  $s_i$ , with a new value  $s'_i$  sampled according to

$$s'_i \sim \Pr\left(S_i | \overline{\mathbf{s}}_i, o_1, \dots, o_m\right)$$

This has detailed balance, and has  $Pr(Q, L|o_1, ..., o_m)$  as its stationary distribution.

It is known as Gibbs sampling.

To see that  $Pr(Q, \mathbf{L} | \mathbf{o})$  is the stationary distribution we just demonstrate *detailed balance*:

$$\pi(\mathbf{s}) \Pr(\mathbf{s} \to \mathbf{s}') = \Pr(\mathbf{s}|\mathbf{o}) \Pr(s'_i|\overline{\mathbf{s}}_i, \mathbf{o})$$
  
=  $\Pr(s_i, \overline{\mathbf{s}}_i|\mathbf{o}) \Pr(s'_i|\overline{\mathbf{s}}_i, \mathbf{o})$   
=  $\Pr(s_i|\overline{\mathbf{s}}_i, \mathbf{o}) \Pr(\overline{\mathbf{s}}_i|\mathbf{o}) \Pr(s'_i|\overline{\mathbf{s}}_i, \mathbf{o})$   
=  $\Pr(s_i|\overline{\mathbf{s}}_i, \mathbf{o}) \Pr(s'_i, \overline{\mathbf{s}}_i|\mathbf{o})$   
=  $\Pr(\mathbf{s}' \to \mathbf{s}) \pi(\mathbf{s}').$ 

As a further simplification we can exploit *conditional independence*.

For example, sampling from  $\Pr(S_i | \overline{s}_i, \mathbf{o})$  may be equivalent to sampling  $S_i$  conditional on some smaller set.

### Approximate inference for Bayesian networks

So:

- We successively sample the query variable and the unobserved variables, conditional on the remaining variables.
- This gives us a sequence  $s_1, s_2, \ldots$  sampled according to Pr(Q, L|o).



Approximate inference for Bayesian networks

To see that the final step works, consider what happens when we estimate the expected value of some function of Q.

$$\begin{split} \mathbb{E}[f(Q)|\mathbf{o}] &= \sum_{q} f(q) \Pr\left(q|\mathbf{o}\right) \\ &= \sum_{q} f(q) \sum_{\mathbf{l}} \Pr\left(q, \mathbf{l}|\mathbf{o}\right) \\ &= \sum_{q} \sum_{\mathbf{l}} f(q) \Pr\left(q, \mathbf{l}|\mathbf{o}\right) \end{split}$$

so sampling using  $\Pr(q, \mathbf{l} | \mathbf{o})$  and ignoring the values for l obtained works exactly as required.

### Markov random fields

*Markov random fields (MRFs)* (sometimes called *undirected graphical models* or *Markov networks*) provide an *alternative approach* to representing a *probability distribution* while expressing *conditional independence assumptions*.

We now have:

- 1. An undirected graph G = (N, E).
- 2. *G* has a node  $N_i$  for each RV.
- 3. For each *maximal clique* c in G there is a *clique potential*  $\phi_c(N_c) > 0$  where  $N_c$  is the set of nodes in c.
- 4. The probability distribution expressed by G is

$$\Pr(N) \propto \prod_{c} \phi_c(N_c).$$

#### Markov random fields

*Example*: 3 maximal cliques of size 2, 2 of size 3 and 1 of size 4.



 $\Pr(N_1, \dots, N_9) \propto \phi_1(N_1, N_4) \times \phi_2(N_3, N_6) \times \phi_3(N_7, N_8) \times \phi_4(N_1, N_2, N_3) \\ \times \phi_5(N_3, N_8, N_9) \times \phi_6(N_4, N_5, N_6, N_7).$ 

### Markov random fields—conditional independence

The *test for conditional independence* is now simple: if X, Y and Z are disjoint subsets of the RVs then:

1. *Remove* the nodes in Z and any attached edges from the graph.

2. If there are *no paths from any variable in X* to *any variable in Y* then

 $X \perp Y | Z.$ 

Final things to note:

- 1. MRFs have their own algorithms for inference.
- 2. They are an *alternative* to *BNs* for representing a probability distribution.
- 3. There are *trade-offs* that might make a BN or MRF more or less favourable.
- 4. For example: *potentials offer flexibility* because they don't have to represent conditional distributions...
- 5....BUT you have to *normalize* the distribution you're representing.