Outline

1. Markov properties and graphs
   - Conditional independence
   - Conditional independence graphs
   - Directed acyclic graphs
   - Roles for graphs

2. Structural learning

3. Decomposable graphs
   - Bayesian model determination in decomposable graphs
   - Priors on decomposable graphs
   - MCMC for structural/quantitative learning
   - Sampling junction trees

4. Non-decomposable graphs

5. Trees and forests

6. DAGs and BNs
Conditional independence

The key idea in understanding

- the structure of a multivariate distribution
- the structure of a sample of multivariate data

is conditional independence, a topic that has been extensively studied both in spatial statistics and in graphical modelling.

\(X \perp \perp Y \mid Z\)

means that if you already know the value of \(Z\), learning that of \(Y\) tells you nothing more about \(X\). Any dependence between \(X\) and \(Y\) is indirect, mediated through \(Z\).
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Conditional independence, probabilistically

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In terms of probability distributions, this means

$$p(x, y|z) = p(x|z)p(y|z)$$

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Markov random fields: the local Markov property
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Pairwise Markov property
Global Markov property
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Many years later, the theorem was superseded by a more complete understanding of Markov properties in undirected graphical models: we can distinguish Global, Local and Pairwise Markov properties, and relate all these to the Factorisation property of Gibbs distributions; in general

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F \implies G \implies L \implies P
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Under an additional condition implied by positivity of the joint density, \( G, L \) and \( P \) are all equivalent, and if the density is also continuous, \( F \) is equivalent too.
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Under an additional condition implied by **positivity** of the joint density, \( G, L \) and \( P \) are all equivalent, and if the density is also **continuous**, \( F \) is equivalent too.
The conditional independence graph $G$ of a multivariate distribution (for a random vector $X$, say) tells us much about the structure of the distribution. $G = (V, E)$ where the vertices $V$ index the components of $X$, and there is an (undirected) edge between vertices $i$ and $j$, written $i \sim j$ unless $X_i \perp \perp X_j | X_{V \setminus \{i,j\}}$

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Markov properties of DAGs

The directed local Markov property:

Variables are independent of their non-descendants, given their parents. There are also directed pairwise and global properties – the latter again involving separation. The directed factorisation property can be written

\[ p(X) = \prod_{v \in V} p(X_v | X_{pa(v)}) \]
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Roles for graphs in statistics

- Visualisation
- Modelling
- Inference - understanding, discovery of structure
- Algorithms
- ...
Graphs driving algorithms

- MCMC algorithms: Gibbs and Metropolis–Hastings
- INLA
- SMC
- Probability propagation
- Other message-passing algorithms
Association and causality

As we all know, association and causality are different, an important distinction blurred by some who call all DAGs ‘causal’.

Graphical model ideas can play a key part in investigating causality, in particular elucidating exactly when it is legitimate to infer causal understanding from an observational study.

The key difficulty is the problem of confounding:
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Structural learning of undirected graphs

Given i.i.d. observations on $X$, we are often interested in inferring $G$, the problem of structural learning (model selection, system identification). Why do we want to do this?

- $G$ may be of direct interest (e.g. constructing pedigrees, gene networks)
- looking for parsimony (e.g. covariance selection for stability of estimation)
- just for visual understanding

This is a model selection problem; it entails search in a huge discrete model space: there are $2^{\binom{v}{2}}$ graphs on $v$ vertices.
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Contingency tables

Prognostic factors for coronary heart disease

Analysis of a $2^6$ contingency table
(Edwards & Havranek, *Biometrika*, 1985)

- smoking?
- strenuous physical work?
- family history of CHD?
- blood pressure > 140?
- ratio of $\alpha$ and $\beta$ lipoproteins >3?
- strenuous mental work?
SNPs and gene expression

min BIC forest

from Lauritzen (2012).
S&P 500 equity data

Figure 10. Graphs build on S&P 500 stock data from Jan. 1, 2003 to Jan. 1, 2008. The graphs are estimated using (a) the glasso, (b) the nonparanormal and (c) forest density estimation. The nodes are colored according to their GICS sector categories. Nodes are not shown that have zero neighbors in both the glasso and nonparanormal graphs. Figure (d) shows the maximum weight spanning tree that results if the data are not Winsorized to trim outliers.

from Lafferty, Liu, Wasserman (2012).
Graphical model fitted to contingency table relating disease status (aff), SNPs – with Linkage disequilibrium, covariates, and 4 quantitative traits. Abel & Thomas, GAW17.
What is structural learning really supposed to deliver?

Does the absence of an edge really signifies conditional independence, or simply insignificant dependence?

c.f. sparsity in regression modelling/variable selection
Today, we will emphasise Bayesian methods, and specifically those that in principle deliver exact or approximate posterior probabilities (for some or all graphs), not simply optimise a possibly-Bayesian objective function.

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Decomposable graphs

Decomposable graphical models

The case where $G$ is decomposable has been much studied. Decomposability is a graph theory concept with statistical and computational implications.

Decomposable graphs are also known as triangulated or chordal: a graph is decomposable if and only if it has no chordless $k$-cycles for $k \geq 4$. 

Peter Green (UTS/Bristol)
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![not decomposable](image1.png) ![decomposable](image2.png)
A graph is decomposable if and only if it has a junction tree representation.

A junction tree is a graph whose vertices are cliques (maximal complete subgraphs), with the property that the cliques containing any prescribed set of vertices forms a connected sub-tree.

We label the links of a junction tree with the separators, intersections of the adjacent cliques. There may be many junction trees for a given decomposable graph.
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We label the links of a junction tree with the separators, intersections of the adjacent cliques. There may be many junction trees for a given decomposable graph.
If the distribution of a random vector $X$ has a decomposable conditional independence graph, then it has a remarkable representation in terms of (often low-dimensional) marginals:

$$p(X) = \frac{\prod_{C \in C} p(X_C)}{\prod_{S \in S} p(X_S)}$$

This is the ultimate generalisation of the fact that for an ordinary Markov chain

$$p(X) = p(X_0) \prod_{i=1}^{N} p(X_i|X_{i-1}) = \frac{\prod_{i=1}^{N} p(X_{\{i-1,i\}})}{\prod_{i=2}^{N-1} p(X_{i-1})}$$

For a general decomposable graph, the same kind of factorisation follows the branches of the junction tree.
Probabilistic significance of decomposability

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For a general decomposable graph, the same kind of factorisation follows the branches of the junction tree.
There are many consequences for computing with distributions on decomposable graphs, including junction tree algorithms (message passing/probability propagation) for Bayes nets (discrete graphical models).
Statistical significance of decomposability

Explicit Maximum likelihood estimates and exact tests for conditional independence for contingency tables and multivariate Gaussian distributions on decomposable graphs.

Clique–separator factorisation

\[ p(X) = \frac{\prod_{c \in C} p(X_c)}{\prod_{S \in S} p(X_S)} \]

yields dramatic speed-ups in structural learning.

Dawid & Lauritzen’s hyper-Markov laws - a framework for the construction of consistent prior distributions respecting the graphical structure.
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How restrictive is decomposability?

How many graphs are decomposable?

There are $2^{(v)}$ graphs altogether on $v$ vertices.

For $v \leq 3$ vertices, all are decomposable
for 4 vertices, $61/64$
for 6, $\approx 55\%$
for 8, $\approx 12\%$. 
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The 3 non-decomposable 4-vertex graphs:
Does that matter?

There is no reason why Nature should be kind enough to give us data from graphical models that are decomposable...

But given any (undirected) graphical model, we can add (‘fill in’) edges to make the graph decomposable.
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![Graphical Model Example](image)
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But given any (undirected) graphical model, we can add (‘fill in’) edges to make the graph decomposable.

So long as our model for the data, given the graph $G$, allows arbitrarily small interactions, we will lose little by assuming decomposability – we will merely tend to infer (hopefully, slightly) more complicated graphs than necessary.
Bayesian model determination with non-decomposable graphs

What happens if the true graph is not decomposable, but you conduct Bayesian structural learning assuming it is?

Fitch, Jones and Massam (2014, *Bayesian Analysis*) show that (for the 0-mean Gaussian case, HIW prior on the concentration matrix), asymptotically,

- The posterior will converge to graphical structures that are minimal triangulations of the true graph.
- The marginal log likelihood ratio comparing different minimal triangulations is stochastically bounded and appears to remain data dependent regardless of the sample size.
- The covariance matrices corresponding to the different minimal triangulations are essentially equivalent, so model averaging is of minimal benefit.
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And assuming decomposability has tremendous advantages....

- Computational advantages in fitting the model
- Evaluating the fit
- Prediction
- Sampling data from fitted model
Given \( n \) i.i.d. samples \( \mathbf{X} = (X_1, X_2, \ldots, X_n) \) from a multivariate distribution on \( \mathbb{R}^v \) parameterised by the graph \( \mathcal{G} \) and parameters \( \theta \), the usual formulation takes the form

\[
p(\mathcal{G}, \theta, \mathbf{X}) = \pi(\mathcal{G})p(\theta|\mathcal{G})p(\mathbf{X}|\mathcal{G}, \theta)
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and we perform joint structural/quantitative learning by computing the posterior \( p(\mathcal{G}, \theta|\mathbf{X}) \propto p(\mathcal{G}, \theta, \mathbf{X}) \).

What prior on \( \mathcal{G} \)? And on \( \theta|\mathcal{G} \)?
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What prior on $\mathcal{G}$? And on $\theta|\mathcal{G}$?
Many authors simply take a prior uniform over all valid graphs (makes sense even if we can’t count them!), or a binomial model conditioned on decomposability.

Armstrong et al, 2009, advocate specifying prior over the size of the graph (which is then uniform conditional on size).

Is there a canonical (e.g. conjugate) approach?
Priors on decomposable graphs

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Conjugate priors on decomposable graphs

Recall that in any decomposable graphical model the likelihood has the form

$$p(X|G) = \frac{\prod_{C \in C} p(X_C|G)}{\prod_{S \in S} p(X_S|G)}$$

So any prior on the graph $G$ that factorises similarly as a product over cliques divided by a product over separators will be conjugate.
A graph law $\pi(G)$ over the set $\mathcal{U}$ of undirected decomposable graphs on $V$ is *structurally Markov* (Byrne, 2011, Byrne & Dawid, 2015) if for any covering pair $(A, B)$, we have:

$$G_A \perp \perp G_B \mid \{G \in \mathcal{U}(A, B)\} \; [\pi],$$

where $\mathcal{U}(A, B)$ is the set of decomposable graphs for which $(A, B)$ is a decomposition.

- $(A, B)$ is a covering pair if $A \cup B = V$.
- $(A, B)$ is a decomposition if $A \cap B$ is complete, and separates $A \setminus B$ and $B \setminus A$. 
Byrne & Dawid’s structural Markov property

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\[ 	ext{Diagram of decomposable graphs} \]
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Byrne & Dawid show that a graph law is structurally Markov if and only if has the form

$$\pi(G) \propto \prod_{C \in C} \phi_C \prod_{S \in S} \phi_S$$

where $\{\phi_A : A \subseteq V\}$ are arbitrary positive set-indexed parameters.
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A new weak structural Markov property

A graph law $\pi(G)$ over the set $\mathcal{U}$ of undirected decomposable graphs on $V$ is weakly structurally Markov (WSM) if for any covering pair $(A, B)$, we have:

$$G_A \perp \perp G_B \mid \{G \in \mathcal{U}^A(A, B)\} \ [\pi],$$

where $\mathcal{U}^A(A, B)$ is the set of decomposable graphs for which $(A, B)$ is a decomposition, and $A \cap B$ is a clique in $G_A$.

This places fewer conditional independence conditions on $\pi$, so potentially corresponds to a richer class of graph priors – but we will see that we can still say something concrete about the form of these laws.
A new weak structural Markov property

A graph law \( \pi(G) \) over the set \( \mathcal{U} \) of undirected decomposable graphs on \( V \) is *weakly structurally Markov (WSM)* if for any covering pair \((A, B)\), we have:

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This places *fewer* conditional independence conditions on \( \pi \), so potentially corresponds to a richer class of graph priors – but we will see that we can still say something concrete about the form of these laws.
A weak structural Markov property

16 possibilities for $G_A$
(if $A \cap B$ remains a clique in $G_A$)

$G_A \perp \perp G_B | \{G \in \mathcal{U}_A(A, B)\}$ [$\pi$],

4 possibilities for $G_B$
We can show that a graph law is weakly structurally Markov if and only if has the form

$$\pi(G) \propto \prod_{C \in C} \phi_C \prod_{S \in S} \psi_S$$

where $$\{\phi_A : A \subseteq V\}$$, $$\{\psi_A : A \subseteq V\}$$ are arbitrary positive set-indexed parameters.

This more general form allows valuable extra flexibility in prior specification; this class of priors has also been studied by Bornn and Caron (2011).
Clique–separator factorisation graph laws

We can show that a graph law is weakly structurally Markov if and only if has the form

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Posterior using a prior with the weak structural Markov property

The posterior for $G$ is

$$p(G|X) \propto \frac{\prod_{C \in C} [\phi_C p(X_C|G)]}{\prod_{S \in S} [\psi_S p(X_S|G)]}$$

that is, a CSF law with parameters $\phi_A p(X_A|G)$ and $\psi_A p(X_A|G)$. 
Bayesian decomposable graphical model determination

For trees, there are explicit finite algorithms for computing MAP estimates; also perfect simulation is possible for random spanning trees, so a full Bayesian analysis can be conducted.

It would be interesting to find a way to extend these ideas to decomposable graphs, but that has not so far been successful.
Bayesian decomposable graphical model determination

For decomposable graphs, joint structural/quantitative learning currently requires MCMC sampling of the posterior $p(G, \theta | X) \propto p(G, \theta, X)$: this means running a Markov chain whose states have the form $(G, \theta)$ – a graph and a vector of parameters.

This chain is constructed to have equilibrium distribution $p(G, \theta | X)$ by ensuring that all moves have detailed balance with respect to this distribution, by using a Metropolis–Hastings sampler.
Pre-tests for maintaining decomposability

Conditions for maintaining decomposability in single-edge moves:

**Frydenberg & Lauritzen** Disconnecting \( x \) and \( y \) by removing an edge \((x, y)\) from \( G \) will result in a decomposable graph if and only if \( x \) and \( y \) are contained in exactly one clique.

**Giudici & Green** Connecting \( x \) and \( y \) by adding an edge \((x, y)\) to \( G \) will result in a decomposable graph if and only if \( x \) and \( y \) are contained in cliques that are adjacent in some junction tree of \( G \).

These extend to certain multiple-edge moves (completely connecting or disconnecting subsets of nodes).

These moves can be efficiently implemented using a junction-tree representation of the graph, since they make only local changes to the junction tree – but they can change its topology locally.
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These moves can be efficiently implemented using a junction-tree representation of the graph, since they make only local changes to the junction tree – but they can change its topology locally.
Pre-tests for maintaining decomposability

Conditions for maintaining decomposability in single-edge moves:

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Using the junction tree as the state

We recently found a simple way to speed up sampling dramatically, by ruling out the need to change the topology of the junction tree – we do this by using directly the junction tree $J$ as part of the model parameterisation, in place of the graph $G$.

This means augmenting the model so that, conditional on $G$, the junction tree $J$ is a priori drawn uniformly from among all equivalent junction trees, thus replacing the prior $\pi(G)$ on decomposable graphs by

$$\tilde{\pi}(J) = \frac{\pi(G(J))}{\mu(G(J))}$$

where $G(J)$ is the decomposable graph determined by $J$ and $\mu(G)$ is the number of equivalent junction trees representing $G$.

Fortunately, we have an efficient local method for evaluating $\mu(G)$, and a local method for occasionally randomising over equivalent junction trees.
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Whether two decomposable graphs are adjacent in the junction tree representation depends on the choice of junction tree.
Sampling decomposable graphs for posterior simulation

- Pre-testing for maintenance of decomposability
- Simplification of likelihood ratios and prior ratios from properties of decomposability
- Using the junction tree as the state
- Multiple-edge moves, randomising over junction trees

allow effective sampling of moderate sized graphs (?50–100 vertices).

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Giudici and Green (1999) and other early work conducts joint structural and quantitative learning by sampling from the joint posterior of graph and parameters – usually a variable-dimension problem, necessitating reversible jump or equivalent trans-dimensional sampling.

To try to scale up to larger problems, almost all authors have used conjugate hyper-Markov priors for parameters, so that these can be integrated out. Metropolis ratios for graph updates then simplify into a locally-computable form.

Thus, hyper-inverse-Wishart priors are used for variance matrices in gaussian models (Giudici and Green, 1999, Dobra, 2004, Jones et al, 2005), and hyper-Dirichlet for multinomial cell probabilities (Tarantola, 2004).

For the gaussian case, there has been a lot of detailed work on choice of hyperparameters in hyper-inverse-Wishart priors (e.g. Rajaratnam, Massam and Carvalho, 2008, Carvalho and Scott, 2009, Ben-David et al, 2011).
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- good at recovering low-dimensional features of the posterior, such as edge-inclusion probabilities, but
- in larger graphs, poor at global properties, as measured by log-posteriors.

This motivates the powerful idea of using online estimation of edge-inclusion probabilities to drive a ‘global’ stochastic search heuristic - the key idea in FINCS (feature inclusion stochastic search), Scott and Carvalho (2008), which combines local moves with resampling from the past history and jumps to new regions of graph space.

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... or rather, not necessarily decomposable graphs.

Even if $G$ is not decomposable, we still have the prime component factorisation

$$p(X) = \frac{\prod_{P \in \mathcal{P}} p(X_P)}{\prod_{S \in \mathcal{S}} p(X_S)}$$

where the prime components $P_i$ are the maximal subgraphs that cannot be decomposed: in a non-decomposable graph, at least one is not complete.
Non-decomposable graphs

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Bayesian model determination with non-decomposable graphs

The additional difficulties in sampling non-decomposable graphical models are (Jones et al, Stat. Sci., 2005):

- The normalising constants in the non-complete prime component marginals do not have closed form, so we need Monte Carlo methods to estimate them.
- These Monte Carlo calculated values have high variance.
- When you make single-edge perturbations to the graph, there is no guarantee of significant cancellations in likelihood ratios.

These difficulties hugely increase computing time – in their experiments, 420 times for a 12-node, 15-edge example; 5500 times for 15-node, 26-edge example (this is for Gaussian models, using conjugate priors on variances).
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Jones et al (Stat. Sci., 2005) conclude that sampling from the posterior is not practical for problems with much more than 15 nodes – and resort to (fast) heuristics like stochastic shotgun search to identify a graph with high posterior probability instead.

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Stochastic shotgun search

As a heuristic for computing part of the posterior, Jones et al (2005) propose iterating on

1. start with a graph $G$
2. select at random $n_1$ graphs that differ by one edge (neighbours), compute their unnormalized posterior probabilities and retain the top $n_2$
3. among the $n_2$ top neighbours, propose the $i$th graph $G_i$ as a new starting graph for an MCMC update with probability proportional to $p_i^\alpha$ where $p_i$ is the unnormalized posterior probability of graph $G_i$ and $\alpha$ is an annealing parameter.
4. return to step 2 and iterate. Maintain a list of the overall best $n_3$ graphs visited.

If the top $n_3$ graphs really capture most of the posterior probability, we might hope to get a good idea of the true posterior distribution.

This SSS method is more fully explored in Hans et al (2007); see also Ben-David et al (2011).
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Examples of Stochastic shotgun search vs MCMC

True, max posterior probability graphs: decomposable, unrestricted

<table>
<thead>
<tr>
<th>Method</th>
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<th>Max log posterior</th>
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</tr>
</thead>
<tbody>
<tr>
<td>MH-d</td>
<td>36</td>
<td>-2591.18</td>
<td>912</td>
<td>1</td>
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<tr>
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<td>792</td>
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<td>2773</td>
<td>-2590.94</td>
<td>13,266</td>
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</tbody>
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<tr>
<td>MH-d</td>
<td>93</td>
<td>15633.76</td>
<td>349,484</td>
<td>36</td>
</tr>
<tr>
<td>SSS-d</td>
<td>234</td>
<td>15633.76</td>
<td>33,495</td>
<td>9</td>
</tr>
<tr>
<td>MH-u</td>
<td>513,077</td>
<td>15633.83</td>
<td>666,425</td>
<td>309,222</td>
</tr>
<tr>
<td>SSS-u</td>
<td>5930</td>
<td>15636.38</td>
<td>82,845</td>
<td>112</td>
</tr>
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</table>
A sparse regression approach

...to *gaussian* model determination with non-decomposable graphs.

In a multivariate gaussian distribution, all conditional distributions are of course gaussian linear regressions. Dobra et al (2004) propose estimating these conditional regressions from data using (Bayesian) sparse regression techniques.

To ensure consistency of the estimated conditionals, one approach is take a pre-fixed order of the variables, and regress each one only on its predecessors. Sparse regressions then estimate a DAG, and we can deliver a corresponding undirected graph.

Non-Bayesian sparse regression methods such as the lasso have also been used in this way (Meinshausen and Buhlmann, 2006, Yuan and Lin, 2007)
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Subgraph containing two genes of interest in an example on expression data on 12588 genes in a breast cancer study, by Dobra et al (2004)
Other important contributions on non-decomposable graphs

- Lenkoski and Dobra (2011) – MOSS, a random uphill search method, constructs list of most probable graphs, subject to max clique size.
- Stingo and Marchetti (2015) adapt their perfect-ordering-representation to an approximate method for non-decomposable graphs, using a special mixture prior that models ’nearly decomposable’ graphs.
A tree is a **connected** undirected graph with no loops.
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Trees and forests are decomposable graphs, of course: for a tree, the junction tree is essentially isomorphic to the tree itself.
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If $G$ is a tree $T$, then

$$P(X|T) = \frac{\prod_{C \in C} p(X_C|T)}{\prod_{S \in S} p(X_S|T)} = \frac{\prod_{e \in E(T)} p(X_e|T)}{\prod_{v \in V_i} p(X_v|T)}$$

where $V_i$ are the non-leaf (interior) vertices, and $E(T)$ are the edges of $T$, so

$$p(T|X) \propto p(T) \prod_{(u,v) \in E(T)} \frac{p(X_{u,v})}{p(X_u)p(X_v)}$$

...a product of Bayes factors for dependence along edges.
Perfect simulation for posterior on trees and forests

For trees,

\[ p(T \mid X) \propto p(T) \prod_{(u,v) \in E(T)} \frac{p(X_{u,v})}{p(X_u)p(X_v)} \]

is amenable to perfect simulation (cf Propp and Wilson, 1998), using algorithms for random spanning trees (Guénoche, 1983; Broder, 1989; Aldous, 1990).

This extends to tree priors \( p(T) \) that are ‘decomposable’ - i.e. factorise as products of weights on edges (Meilă & Jaakkola, 2006).

Perfect sampling from random forest distributions of similar form is possible but much harder (Dai, 1998).
A directed acyclic graph (DAG) is a directed graph in which there are no directed loops. Equivalently, it is a directed graph in which edges from a node can only go to a higher-numbered node.
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DAGs

.. particularly important because of (real or more likely imagined) causal interpretation.

When the variables are all categorical, these models are often called Bayes(ian) net(work)s – a big focus of attention in the machine learning community, because of use in expert systems and availability of very fast algorithms.

(So in literature search, look for both DAGs and BNs).
Markov equivalence

Two DAGs are Markov equivalent if they imply exactly the same set of conditional independences.

A (graphical) test: they are Markov equivalent if they have the same skeleton (graph ignoring directions) and the same sets of unmarried parents (Verma and Pearl, 1990).
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Markov equivalence classes of DAGs can be represented graphically, in a form known as maximally oriented graphs (Meek, 1995), essential graphs (Andersson et al., 1997 or completed PDAGs (Chickering, 2002)).

A completed PDAG is a chain graph (graph with edges and arrows, but no directed cycles) that has the same skeleton as the original DAG, and edges that are directed if and only if they are directed in every equivalent DAG.

A DAG and its completed PDAG.
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MCMC on completed PDAGs

He, Jia & Yu (2013) construct a reversible Markov chain on Markov equivalence classes of DAGs, that is scalable to large sparse graphs. It uses operators (MCMC moves) chosen from these:

- **InsertU** $x - z$
- **InsertD** $x \rightarrow v$
- **MakeV** $z \rightarrow y \leftarrow u$
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Bayesian structural learning (in the sense of delivering posterior distributions) from i.i.d. data is a simply stated task.

... that proves amazingly hard to deliver computationally except on a modest scale.

... even with special choices of priors, etc.

... in spite of the best efforts of a lot of creative people!

We don’t really know what heuristics and approximations are delivering, quantitatively.

Perhaps time to focus effort (algorithmic and theoretical) more on edge-inclusion probabilities and other low-dimensional margins?

... and more effort on relaxation of the hard constraints of conditional independence?
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Wrap-up

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Bayesian structural learning (in the sense of delivering posterior distributions) from i.i.d. data is a simply stated task that proves amazingly hard to deliver computationally except on a modest scale, even with special choices of priors, etc. in spite of the best efforts of a lot of creative people! We don’t really know what heuristics and approximations are delivering, quantitatively. Perhaps time to focus effort (algorithmic and theoretical) more on edge-inclusion probabilities and other low-dimensional margins? And more effort on relaxation of the hard constraints of conditional independence?
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Bibliography


