Introduction to Causal Discovery

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Part 1: Classical Strategies and Concepts

What is a graphical model?

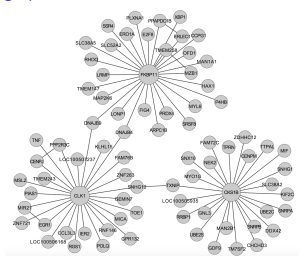


Figure: Gene associations related to asthma, an undirected graph from Wang et al. (2016).

What is a graphical model?

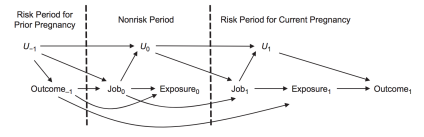


Figure: A directed acyclic graph from Johnson et al. (2019) that illustrates epidemiologic phenomena related to pregnancy risk.

What is a graphical model?

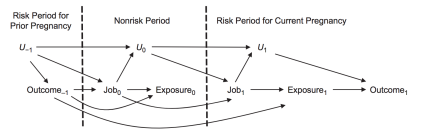


Figure: A directed acyclic graph from Johnson et al. (2019) that illustrates epidemiologic phenomena related to pregnancy risk.

Vertices (nodes) correspond to random variables and edges (arrows) correspond to statistical dependence of some kind. Graphical models are generally used to reason about multivariate joint distributions.

Graphs

A graph $\mathcal{G}=(V,E)$ where V is a set of vertices and E is a set of edges. Typically, the vertices index some random variables, i.e., let V=(1,...,p) with corresponding random variables $X_1,...,X_p$.

The set of edges depends on the type of graph.

In a directed graph, edges are formally ordered pairs of vertices.

$$ightharpoonup V_1
ightarrow V_2$$

In an undirected graph, edges are formally unordered pairs of vertices.

$$V_1 - V_2$$

NB: Sometimes we are a bit informal with notation and just let $V \equiv X = (X_1, ..., X_p)$ when this causes no confusion.

Example: Directed Acyclic Graphs (DAGs)



A directed acyclic graph (DAG) is a graph with only directed edges (\rightarrow) and no *cycles*, i.e., no paths from $X_i \rightarrow \cdots \rightarrow X_i$.

Directed Acyclic Graphs

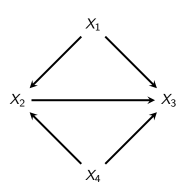
Definitions:

$$\begin{array}{l} \mathsf{Pa}(X_i,\mathcal{G}) \equiv \{X_j: X_j \to X_i \text{ in } \mathcal{G}\} & (\mathsf{parents of } X_i) \\ \mathsf{Ch}(X_i,\mathcal{G}) \equiv \{X_j: X_j \leftarrow X_i \text{ in } \mathcal{G}\} & (\mathsf{children of } X_i) \\ \mathsf{An}(X_i,\mathcal{G}) \equiv \{X_j: X_j \to \ldots \to X_i \text{ in } \mathcal{G}\} & (\mathsf{ancestors of } X_i) \\ \mathsf{De}(X_i,\mathcal{G}) \equiv \{X_i: X_i \leftarrow \ldots \leftarrow X_i \text{ in } \mathcal{G}\} & (\mathsf{descendants of } X_i) \\ \end{array}$$

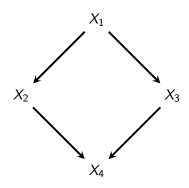
(By convention, X_i counts as an ancestor and descendent of itself.)

Statistical implications

Viewed purely as a statistical object, a DAG model represents a set of conditional independencies.



$$\{p(x_1,x_2,x_3,x_4): X_1 \perp X_4\}$$



$$\{p(x_1, x_2, x_3, x_4) : X_1 \perp \!\!\! \perp X_4 | (X_2, X_3) X_2 \perp \!\!\! \perp X_3 | X_1\}$$

Global Markov property

Let A, B, C be disjoint subsets of X. A distribution p(x) satisfies the global Markov property wrt DAG G if

$$A \perp_d B | C \implies A \perp \!\!\! \perp B | C.$$

where \perp_d denotes d-separation (defined on the next slide)

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where \perp_d denotes d-separation (defined on the next slide)

"A graphical criterion \implies conditional independence in the distribution"

d-separation

A vertex X_j is **active** on a path relative to C just in case either

- i) X_j is a collider $(\to X_j \leftarrow)$, and X_j or any of its descendents is in C, or
- ii) X_i is a noncollider and is not in C.

d-separation

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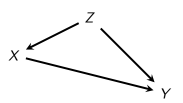
ii) X_j is a noncollider and is not in C.

A path π is **active** relative to C just in case every vertex on π is active relative to C.

 X_i and X_k are d-separated (\perp_d) given C just in case there is no path between X_i and X_k that is **active** relative to C.

 $A \perp_d B \mid C$ if $X_i \perp_d X_k \mid C$ for all $X_i \in A$ and $X_k \in B$.

Causal interpretations (two views)



Potential Outcomes Model

$\{Y(x,z) \mid \forall x,z\}, \{X(z) \mid \forall z\}, \text{ and } Z$ are mutually independent

$$\implies X \perp Y(x)|Z$$
 (conditional ignorability)

Nonparametric Structural Equation Model

$$Z = f_Z(\epsilon_z)$$

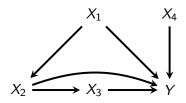
$$X(z) = f_X(z, \epsilon_x)$$

$$Y(x, z) = f_Y(x, z, \epsilon_y)$$

$$\perp (\epsilon_x, \epsilon_y, \epsilon_z)$$

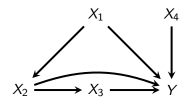
 \implies Z satisfies "back-door" criterion

What are causal graphical models good for?



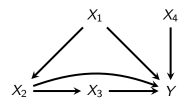
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- Qualitative causal conclusions ("what causes what? what are the pathways?")
- ▶ Valid (and efficient) covariate adjustment for estimating causal effects
- General identification theory for causal effects and mediation analysis

Experimental/problem design

▶ not always possible, feasible, or obvious

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Data

question is: how?

Approaches to graphical structure learning

Constraint-based: hypothesis tests of conditional independence (e.g., PC)

Greedy score-based: optimization of a model fit score (e.g., GES)

Semi-parametric methods: exploiting semi-parametric assumptions (e.g., ICA-LiNGaM)

Optimization-based: continuous optimization of a penalized likelihood or score (e.g., NOTEARS)

Hybrid methods: mix of above

Approaches to graphical structure learning (this lecture)

Constraint-based: hypothesis tests of conditional independence (e.g., PC)

Score-based: greedy optimization of a model fit score (e.g., GES)

Causal sufficiency & acylicity

In this lecture, we will assume *causal sufficiency*: there are no unmeasured (hidden, latent) variables that act as common causes of two or more measured variables.

We will also assume the data-generating process contains no feedback (i.e., is recursive). This means the graphical representation will be acyclic.

That is, the true data-generating process corresponds to some DAG with vertices corresponding *only* to $X = (X_1, ..., X_p)'$.

(In Part 2, we will dispense with causal sufficiency.)

Consider a DAG \mathcal{G} . By the global Markov property, we know that if

$$X_i \perp_d X_j | X_S \text{ in } \mathcal{G}$$

then
 $X_i \perp X_j | X_S \text{ in } p(x)$
here: $X_S \subseteq X \setminus \{X_i, X_i\}$

Thus we say that the graphical structure places an independence constraint on the data distribution. Constraint-based methods directly exploit those implied constraints on p(x) to try and infer backwards from data to graphical structure by a sequence of tests.

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Thus we say that the graphical structure places an independence constraint on the data distribution. Constraint-based methods directly exploit those implied constraints on p(x) to try and infer backwards from data to graphical structure by a sequence of tests.

Score-based methods instead proceed by assigning to every a graph \mathcal{G} some measure of how well the graph "fits" the data. Higher scores mean better fit, and a score-based approach can be viewed as an optimization task: search for the graph which has the highest score.

 \Rightarrow Neither of these approaches is universally "better" than the other, but they have their own virtues and drawbacks depending on the setting.

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 \Rightarrow There are also theoretical connections between these approaches, so not quite so distinct as may seem. (Will get to this later.)

Faithfulness

Markov property tells us a d-separation implies an independence $(X_i \perp_d X_j | X_S \implies X_i \perp X_j | X_S)$ but we can only test independence and hope to infer backwards to d-separation. What justifies this?

The faithfulness assumption imposes the additional constraint that "all observed conditional independence constraints follow from the graphical structure." This allows us to infer $X_i \perp \!\!\! \perp X_j | X_S \implies X_i \perp_d X_j | X_S$.

Faithfulness puts missing edges and conditional independence in a one-to-one correspondence. However, this is a substantive assumption on p(x) and may be violated!

Example: deterministic relationships among variables

Consider:

$$X_1 \longrightarrow X_2 \longrightarrow X_3$$

where $X_3 = 2 \times X_2$.

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where $X_3 = 2 \times X_2$.

 $X_3 \perp X_1 | X_2$ (by Markov property) but also $X_2 \perp X_1 | X_3$ (by determinism).

Example: exact "cancellation" or "balancing"

Consider:

$$X_1 \longrightarrow X_2 \longrightarrow X_3$$

where

$$X_1 = \epsilon_1$$

$$X_2 = \alpha X_1 + \epsilon_2$$

$$X_3 = \beta X_2 - \alpha \beta X_1 + \epsilon_3$$

$$\epsilon_1, \epsilon_2, \epsilon_3 \sim N(0, 1), \quad \alpha, \beta > 0$$

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$$\epsilon_1, \epsilon_2, \epsilon_3 \sim N(0, 1), \quad \alpha, \beta > 0$$

No independencies follow from Markov property, but $X_3 \perp X_1$ (by exact cancellation).

Faithfulness

It is common to rule out such "extra" or "non-graphical" conditional independencies, *by assumption*. This restricts the allowed set of distributions, and may not always be appropriate.

A distribution p(x) satisfies the faithfulness assumption wrt DAG ${\cal G}$ if

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A distribution p(x) satisfies the faithfulness assumption wrt DAG ${\cal G}$ if

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In conjunction with the global Markov property this means we're assuming $A \perp \!\!\! \perp B \mid C \iff A \perp_d B \mid C$.

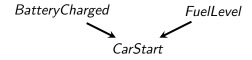
(Here A, B, and C are non-overlapping subsets of V.)

Colliders: a special graphical substructure



In constraint-based causal discovery, unshielded colliders play a central role.

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By d-separation: BatteryCharged \perp_d FuelLevel.

Colliders: a special graphical substructure

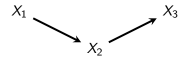


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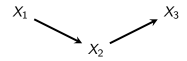
By d-separation: BatteryCharged \perp_d FuelLevel.

By faithfulness assumption: $BatteryCharged \not\perp FuelLevel \mid CarStart$.

Conditioning on *CarStart* makes *FuelLevel* informative about *BatteryCharged*.

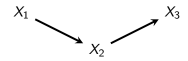


Very different from "chains."



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By d-separation: $X_1 \not\perp_d X_3$.

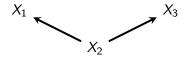


Very different from "chains."

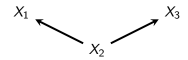
By d-separation: $X_1 \not\perp_d X_3$.

By d-separation: $X_1 \perp_d X_3 | X_2$.

Conditioning on X_2 makes X_1 non-informative about X_3 .

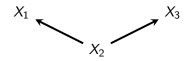


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Colliders: a special graphical substructure

This means that colliders can act as a "smoking gun" to determine some orientations, whereas forks and chains lead to some underdetermination.

How might you select a model if computational limitations weren't an issue?

Brute force enumerate all possible DAGs, derive independence implications for each, and test them.

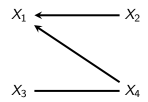
Markov equivalence

We typically identify causal structure only up to Markov equivalence.

$$X_1 o X_2 o X_3$$
 $X_1 \leftarrow X_2 \leftarrow X_3$ $X_2 \leftarrow X_2 o X_3$ $X_1 - X_2 - X_3$ $X_1 o X_2 \leftarrow X_3$ a) b) c)

Figure: a) Three Markov equivalent DAG models. b) The CPDAG representation of (a). c) A DAG that is not Markov equivalent to the graphs in (a).

CPDAG



Let $\mathcal G$ be a DAG. The CPDAG $\mathcal C$ (completed partial DAG) implied by $\mathcal G$ is a mixed graph (directed and undirected edges) that has the same adjacencies as $\mathcal G$ and:

- ▶ a directed edge $X_i \to X_j$ in $\mathcal C$ iff the edge $X_i \to X_j$ is common to all DAGs Markov equivalent to $\mathcal G$
- lacktriangle all other edges in ${\cal C}$ are undirected.

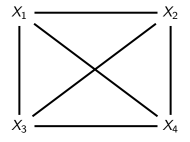
The *skeleton* of a (partially) directed graph is the undirected graph obtained by replacing all edges by undirected edges. Recall that Markov equivalent DAGs share the same skeleton and unshielded colliders.

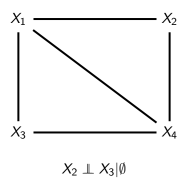
Which independence hypotheses to test?

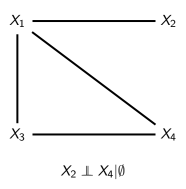
We want to search for separating sets which make each pair X_i, X_j conditionally independent. A naive approach would be to consider all possible subsets $X_S \subseteq X \setminus \{X_i, X_j\}$ to evaluate whether there exists a separating set.

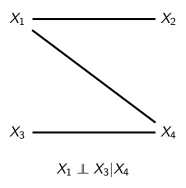
However, by the DAG Markov properties we know that $X_i \notin \operatorname{Adj}(X_j, \mathcal{G})$ if and only if $X_i \perp \!\!\! \perp X_j | \operatorname{Pa}(X_i, \mathcal{G})$ or $X_i \perp \!\!\! \perp X_j | \operatorname{Pa}(X_j, \mathcal{G})$.

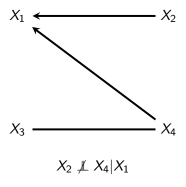
We don't know the parent set ahead of time (we don't know the graph!) so we look at $X_S \subseteq \operatorname{Adj}(X_i, \mathcal{G}')$ and $X_S \subseteq \operatorname{Adj}(X_j, \mathcal{G}')$ for some \mathcal{G}' which is a supergraph of the true unknown skeleton.

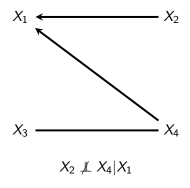












 $X_3 \rightarrow X_4$ and $X_3 \leftarrow X_4$ both possible

Algorithm 0.1: $PC(TEST, \alpha)$

Input: Samples of the vector $X = (X_1, ..., X_p)'$

Output: CPDAG \mathcal{G}

- **1**. Form the complete graph \mathcal{G} on vertex set X w/ undirected edges.
- **2**. Let s = 0
- 3. repeat
- **4. for all** pairs of adjacent vertices (X_i, X_j) s.t. $|\operatorname{Adj}(X_i, \mathcal{G}) \setminus \{X_j\}| \ge s$ and subsets $X_S \subseteq \operatorname{Adj}(X_i, \mathcal{G}) \setminus \{X_i\}$ s.t. |S| = s
- 5. **if** $X_i \perp X_j | X_S$ according to (TEST, α) **then** $\begin{cases} \text{Delete edge } X_i - X_j \text{ from } \mathcal{G}. \\ \text{Let sepset}(X_i, X_i) = \text{sepset}(X_i, X_i) = X_S. \end{cases}$
- 6. end
- **7**. Let s = s + 1
- **8**. **until** for each pair of adjacent vertices (X_i, X_j) , $|\operatorname{Adj}(X_i, \mathcal{G}) \setminus \{X_j\}| < s$.
- **9. for all** triples (i, k, j) s.t. $X_i \in \operatorname{Adj}(X_k, \mathcal{G})$ and $X_j \in \operatorname{Adj}(X_k, \mathcal{G})$ but $X_i \notin \operatorname{Adj}(X_j, \mathcal{G})$, orient $X_i \to X_k \leftarrow X_j$ in \mathcal{G} iff $X_k \notin \operatorname{sepset}(X_i, X_j)$.
- **10**. Exhaustively apply orientation rules (R1-R4) to orient remaining undirected edges.
- 11. return G.

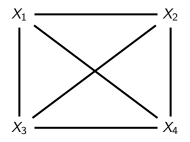
Orientation rules

R1: Orient $X_j - X_k$ into $X_j \to X_k$ whenever there is an arrow $X_i \to X_j$ such that X_i and X_k are nonadjacent.

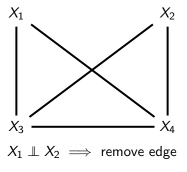
R2: Orient $X_i - X_j$ into $X_i o X_j$ whenever there is a path $X_i o X_k o X_j$.

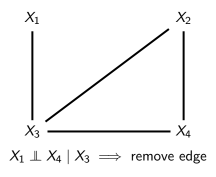
R3: Orient $X_i - X_j$ into $X_i \to X_j$ whenever there are two paths $X_i - X_k \to X_j$ and $X_i - X_l \to X_j$ such that X_k and X_l are nonadjacent.

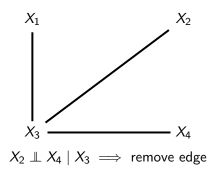
R4: Orient X_i-X_j into $X_i\to X_j$ whenever there are two paths $X_i-X_k\to X_j$ and $X_i-X_l\to X_k$ such that X_j and X_l are nonadjacent.

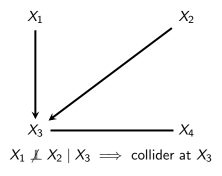


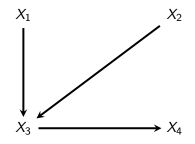
Begin with a "complete" graph











Additional orientation rule \implies directed away from X_3 (no colliders found with X_4 so $X_4 \rightarrow X_3$ ruled out)

The "logic" of PC:

- conditional independencies correspond to missing edges
- ▶ the "collider" rule enables orientation of triples $X_i o X_j \leftarrow X_k$
- other orientations follow from the acyclicity assumption
- remaining unoriented edges reflect the Markov equivalence of models consistent with the data

Theory

Computational complexity:

We note that for graphs with bounded degree, i.e., a bound on $d = \max_{i \in V} |\operatorname{Adj}(X_i, \mathcal{G})|$, the PC algorithm has a running time that is polynomial in the number of variables. The running time depends exponentially on the degree. Specifically the number of tests is bounded by $2\binom{p}{2} \sum_{i=0}^d \binom{p-1}{i}$ (in the worst case).

Theory

Correctness (soundness and completeness):

Theorem: Assume the distribution p(x) is Markov and faithful to some DAG \mathcal{G} . Let \mathcal{C} be the CPDAG implied by \mathcal{G} . The "oracle" PC algorithm returns \mathcal{C} .

Consistency:

Theorem (informal): Assume the distribution p(x) is Markov and faithful to some DAG $\mathcal G$. Let $\mathcal C$ be the CPDAG implied by $\mathcal G$. Let $\widehat{\mathcal C}$ be the output of PC with some consistent conditional independence test and level α_n . Then there is a sequence of $\alpha_n \to 0$ $(n \to \infty)$ such that $\lim_{n \to \infty} \mathbb P(\widehat{\mathcal C} = \mathcal C) = 1$.

Conditional independence tests

With finite data, the PC algorithm needs a procedure for deciding whether $X_i \perp X_j | X_s$.

In practice, test the null hypothesis

$$H_0: X_i \perp \!\!\! \perp X_j | X_S$$

and reject the null if some test statistic $T(x) < \alpha$, where α is a user-specified threshold (which may depend on the sample size). That is, if we reject the null hypothesis, we keep the edge, and if we fail to reject, we remove the edge.

Conditional independence tests

Note: α in PC has no straightfoward interpretation in terms of the Type I error, since in fact we are testing multiple hypotheses and an a priori unknown number of them.

So, α should be better thought of as a *tuning parameter* which controls the **sparsity** of the estimated graph. (Smaller $\alpha \implies$ more sparse)

Though it is not straightforward, one may try to be clever and control the FDR. In practice: make α small as the number of variables and sample size gets big. (More on choosing α later.)

Conditional independence tests

The choice of statistical test depends on the distribution p(x). If you're willing to assume that p(x) is in some nice parametric family (Gaussian, multinomial) then life is easier. However, nonparametric tests of conditional independence are also available (and in development).

First, consider the case where X is multivariate Gaussian.

Test for Gaussian data

Theorem. If X is multivariate Gaussian then $X_i \perp X_j | X_S$ iff $\rho_{ij.S} = 0$.

 $\rho_{ij.S}$ is called the partial correlation (of X_i and X_j given X_S) and can be defined as follows: for any $k \in S$,

$$\rho_{ij.S} = \frac{\rho_{ij.S\setminus k} - \rho_{ik.S\setminus k}\rho_{jk.S\setminus k}}{\sqrt{(1 - \rho_{ik.S\setminus k}^2)(1 - \rho_{kj.S\setminus k}^2)}}$$

This is a recursive definition that terminates at |S|=1. Leads to combos of regular pairwise correlations: $\rho_{ij}=\frac{\text{COV}(X_i,X_j)}{\sigma_i\sigma_i}$.

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This is a recursive definition that terminates at |S|=1. Leads to combos of regular pairwise correlations: $\rho_{ij}=\frac{\text{cov}(X_i,X_j)}{\sigma_i\sigma_j}$.

Easy to obtain estimate $\widehat{\rho}_{ij.S}$ based on the empirical covariance matrix.

Test for Gaussian data

Moreover define Fisher's Z-transform:

$$Z(\rho_{ij.S}, n) = \frac{1}{2}\sqrt{n - |S| - 3}\log\left(\frac{1 + \rho_{ij.S}}{1 - \rho_{ij.S}}\right)$$

under the null hypothesis $\rho_{ii.S} = 0$:

$$Z(\rho_{ij.S},n) - Z(\widehat{\rho}_{ij.S},n) \sim \mathring{N}(0,1) \text{ as } n \to \infty.$$

So for a level α test we reject if $Z(\widehat{\rho}_{ij.S}, n) > \Phi^{-1}(1 - \alpha/2)$

Other parametric tests

There are similar tests for multinomial discrete data based on the χ^2 and G^2 test statistics.

One may also use tests based on (logistic) regression or odds ratios.

Nonparametric tests

Tests which make minimal distributional assumptions are an active area of research. A well-known test based on Kernel matrices is the KCI (Kernel-based Conditional Independence) test.¹

KCI assumes the the variables are related by arbitrary square-integrable functions, plus some smoothness/simplicity conditions to test $X_i \perp X_j | X_S$ nonparametrically. The test statistic is:

$$\frac{1}{n} \operatorname{tr}(\widetilde{K}_{\ddot{X}_i|X_S} \widetilde{K}_{X_j|X_S})$$

where $\widetilde{K}_{A|B}$ is a conditional centralized kernel matrix constructed with some kernel function, and $\ddot{X}_i = (X_i, X_S)$.

¹Zhang et al. (2011) "Kernel-based conditional independence test and application in causal discovery" in UAI

Nonparametric tests

The KCI test is implemented in R (e.g., library(CondIndTests))

```
n <- 100
Z <- rnorm(n)
X <- 4 + 2 * Z + rnorm(n)
Y <- 3 * X^2 + Z + rnorm(n)
test1 <- CondIndTest(X,Y,Z, method = "KCI")
cat("These data come from a distribution, for which X and Y are NOT cond. ind. given Z.")
cat(paste("The p-value of the test is: ", test1$pvalue))</pre>
```

The problem is that the test is very computationally intensive with big n and big |S|.

There are other nonparametric tests in other packages, each have pros and cons

Other nonparametric independence tests

Shah and Peters $(2020)^2$ propose an independence test based on:

$$\mathbb{E}[\operatorname{cov}(X_i,X_j|X_S)]$$

which = 0 under the null, and

$$cov(X_i, X_j | X_S) = \mathbb{E}[X_i X_j | X_S] - \mathbb{E}[X_i | X_S] \mathbb{E}[X_j | X_S]$$

The test statistic is based on estimating residuals from semi/nonparametric regressions, i.e.,

$$\{x_i - \hat{f}(x_S)\}\{x_j - \hat{g}(x_S)\}$$

under the assumption that the true $f(x_S) = \mathbb{E}[X_i|X_S = x_S], g(x_S) = \mathbb{E}[X_j|X_S = x_S]$ satisfy some smoothness conditions.

²Shah & Peters (2020) "The hardness of conditional independence testing and the generalised covariance measure," Annals of Statistics, 48(3), 1514-1538.

Virtues and drawbacks of constraint-based search

Virtues:

- nonparametric (in principle)
- relatively scaleable
- lots of work on improvements, heuristics, generalizations

Drawbacks:

- statistical test errors propogate, can make a big difference to the output (can be mitigated by some stability techniques)
- does not handle conflicting statistical information in an intelligent way
- not that scaleable (cannot really parallelize)
- tackles problem only "locally," makes each decision only once, produces one estimated CPDAG rather than a range of good options (no "confidence intervals" or "posterior distribution")

Score-based model selection for DAGs

Score-based procedures view structure learning as an optimization problem: assign a score to every structure, and find the structure with the highest score.

A natural score to consider is the posterior probability of a particular structure given the observed data. More a posteriori probable structures "fit" the data "better" and we can imagine searching for the structure which has the highest posterior. How would we define this?

Posteriors over BN models

Statistically, a DAG model represents a set of distributions \mathcal{P} that satisfy the Markov condition wrt \mathcal{G} . Let $\theta_{\mathcal{G}}$ be the set of parameters which index the distributions in \mathcal{P} .

We define the postior probability of a structure $\mathcal G$ given data D as

$$p(G|D) = \frac{p(D|G)p(G)}{p(D)}$$

where $p(\mathcal{G})$ is a structure prior, $p(D|\mathcal{G}) = \int_{\Theta_{\mathcal{G}}} p(D|\theta_{\mathcal{G}},\mathcal{G}) p(\theta_{\mathcal{G}}|\mathcal{G}) d\theta_{\mathcal{G}}$, and $p(\theta_{\mathcal{G}}|\mathcal{G})$ is a parameter prior. $p(D|\mathcal{G})$ is called the *marginal likelihood*. Note that the denominator p(D) is the same for every structure so does not play a role in structure selection; we can thus ignore it.

Bayesian score

We could follow this reasoning to define a model score $S(\mathcal{G},D)$ which takes a candidate structure \mathcal{G} and a dataset D and returns a number proportional to the posterior (ignoring the denominator): the Bayesian score $S(\mathcal{G},D) \equiv \log p(D|\mathcal{G}) + \log p(\mathcal{G})$. We could even assume that every DAG in the space of DAGs has equal prior probability, to make that part easy.

That still leaves us with trying to evaluate $p(D|\mathcal{G}) = \int_{\Theta_{\mathcal{G}}} p(D|\theta_{\mathcal{G}},\mathcal{G}) p(\theta_{\mathcal{G}}|\mathcal{G}) d\theta_{\mathcal{G}}$ which depends on a prior for every $\theta_{\mathcal{G}}$.

 \Rightarrow this is very difficult to calculate outside of special cases.

Bayesian score

There was a lot of work in the early 90's on how to evaluate the marginal likelihood for special cases like binary (multinomial) or Gaussian random variables. Turns out if you don't pick the parameter prior right, you probably can't evaluate that integral.

For example, a lot of work was done (in the binary case) with the Dirchelet prior, $\theta_{\mathcal{G}} \sim Dir(\alpha)$ which has a density proportional to $\prod_{i=1}^n \theta_i^{\alpha_i-1}$. This isn't intuitive, but the Dirchelet prior is conjugate for the multinomial model, thus making $p(D|\mathcal{G})$ relatively easy to compute. There are few other examples of convenient priors like this.

BIC score

These days, it much more likely that one would use an approximation to the Bayesian score called the BIC criterion. More specifically, for distributions in the exponential family the BIC is an approximation to the marginal likelihood $p(D|\mathcal{G})$ under some weak assumptions about the prior).

$$\log p(D|\mathcal{G}) \approx \ell(D; \widehat{\theta}_{\mathcal{G}}) - \frac{d}{2} \log n + O(1)$$

where $\ell(\cdot)$ is the log-likelihood, $\widehat{\theta}_{\mathcal{G}}$ is the maximum likelihood estimate of the parameters, n is the sample size, and d is the model dimension. Assume a uniform prior over graphical structures we can just use this approximation as our model score:

$$S(G, D) \equiv \ell(D; \widehat{\theta}_G) - \frac{d}{2} \log n$$

BIC score interpretation

What's the interpretation here? If we assume a uniform prior over structures and a "smooth" prior over parameters, maximizing the BIC score is approximately equivalent to maximizing the posterior probability of the graph structure, given the data. That's the Bayesian interpretation.

Alternatively, we can view the BIC score as measure of model "fit" which is a penalized MLE – the second term penalizes complex models, preferring models of smaller dimension. Why? If we just used the MLE as measure of model fit, we would always choose the most complex graph, overfitting the data. The penalty prevents overfitting.

BIC score interpretation

You may wonder what would be the non-Bayesian justification of this particular sparsity penalty: $\frac{d}{2} \log n$. The answer is that this penalty makes the score consistent, i.e., maximizing the BIC score will select the true model in the limit. Some other penalties you may come up with will not be consistent (example: the Aikake Information Criterion or AIC score which has a penalty $\propto d$, independent of n, is not consistent!)

The BIC score is also easy to evaluate, because we know how to do MLE. For example: When $X \sim N(0, \Sigma)$

$$S(G, D) = \frac{n}{2} \log |\hat{\Sigma}| - \frac{d}{2} \log n$$

What might you do if computational limitations weren't an issue?

Brute force enumerate all possible DAGs, score each, and select the highest scoring one.

$$X_1 o X_2 o X_3$$
 $X_1 \leftarrow X_2 \leftarrow X_3$ $X_1 \leftarrow X_2 o X_3$ $X_1 \to X_2 \leftarrow X_3$ \downarrow \downarrow \downarrow \downarrow \downarrow \downarrow $S(\mathcal{G}_1, D)$ $S(\mathcal{G}_2, D)$ $S(\mathcal{G}_3, D)$ $S(\mathcal{G}_4, D)$

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$$S(\mathcal{G}_1, D) = S(\mathcal{G}_2, D) = S(\mathcal{G}_3, D)$$

since "same independence model \Rightarrow same likelihood" (for Gaussians) + same dimension

Greedy score-based model selection

Greedy score-based algorithms use **stepwise optimization** of the BIC score in such a way that we don't have to traverse the whole space of graphical structures, but which is still globally optimal.

Greedy score-based model selection

Greedy score-based algorithms use **stepwise optimization** of the BIC score in such a way that we don't have to traverse the whole space of graphical structures, but which is still globally optimal.

The Greedy Equivalence Search (GES) algorithm operates in the space of CPDAGs, to respect Markov equivalence.

Adds and removes directed edges to optimize a BIC score.

 X_1

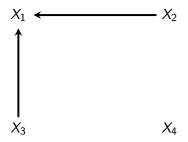
 X_2

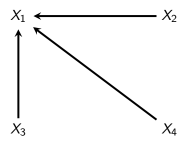
 X_3

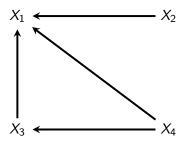
 X_4

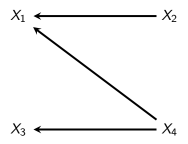
$$X_1 \longleftarrow X_2$$

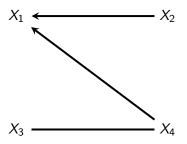
$$X_3$$











Algorithm 0.2: GES(Score)

Input: Samples of the vector $X = (X_1, ..., X_p)'$

Output: CPDAG \mathcal{G}

- 1. Form the empty graph \mathcal{G} on vertex set X.
- 2. Let $S(\mathcal{G}, D)$ be the Score for \mathcal{G} with data D.
- 3. $\langle \mathcal{G}, \mathcal{S} \rangle \leftarrow \text{ForwardEquivalenceSearch}(\mathcal{G}, \mathcal{S})$
- 4. $\mathcal{G} \leftarrow \text{BackwardEquivalenceSearch}(\mathcal{G}, \mathcal{S})$
- 5. return \mathcal{G} .

```
Input: Samples of the vector X = (X_1, ..., X_p)'
Output: CPDAG \mathcal{G}, Score S
1. while E_0 \neq \emptyset
        E_0 \leftarrow T_0 \leftarrow \emptyset, S_0 \leftarrow 0.
        for each candidate edge E = X_i \rightarrow X_i s.t. X_i \notin Adj(X_i, \mathcal{G})
3.
4.
             Let T' \leftarrow \text{vertices } X_k \text{ s.t. } X_k - X_i \text{ and } X_k \not\in \text{Adj}(X_i, \mathcal{G})
             for each subset T \in T'
5.
                   G' \leftarrow a DAG in G
                   S' \leftarrow S + \text{ScoreEdgeAddition}(G, E, T)
                   if S' > S and S' > S_0 and VALIDINSERT(G, E, T)
                   then \begin{cases} E_0 \leftarrow E \\ T_0 \leftarrow T \\ S_0 \leftarrow S' \end{cases}
9.
             end
10.
         end
11.
         if E_0 \neq \emptyset
          13. end
14. return \langle \mathcal{G}, \mathcal{S} \rangle.
```

This psuedocode follows Ramsey et al. (2010) "Six problems for causal inference in fMRI". NeuroImage 49: 1545-1558.

```
Input: Samples of the vector X = (X_1, ..., X_p)'
Output: CPDAG G
1. while E_0 \neq \emptyset
           E_0 \leftarrow H_0 \leftarrow \emptyset, S_0 \leftarrow 0.
          for each edge E = (X_i, X_i) in \mathcal{G}
3.
                   Let H' \leftarrow \text{vertices } X_k \text{ s.t. } X_k - X_i \text{ and } X_k \in \text{Adj}(X_i, \mathcal{G})
                   for each subset H \in H'
5.
                          G' \leftarrow a DAG in G
                          S' \leftarrow S + \text{ScoreEdgeDeletion}(G, E, H)
                          if S' > S and S' > S_0 and VALIDDELETE(G, E, H)
                           then \begin{cases} E_0 \leftarrow E \\ H_0 \leftarrow H \\ S_2 \leftarrow S' \end{cases}
9.
                   end
10.
             end
11.
             if E_0 \neq \emptyset
               \begin{aligned} & \text{then} \begin{cases} \text{Remove } E_0 \text{ from } \mathcal{G}. \\ & \text{for each } H \in H_0 \text{ if } X_i - H \text{ in } \mathcal{G}, \text{ orient } X_i - H \text{ as } X_i \to H. \\ & S \leftarrow S_0 \\ & \mathcal{G} \leftarrow \text{Rebuild}(\mathcal{G}) \end{aligned} 
13. end
14. return \mathcal{G}.
```

Algorithm 0.5: ValidInsert($\mathcal{G}, \mathcal{E}, \mathcal{T}$)

- 1. $NA_{ij} \leftarrow \text{vertices } X_k \text{ s.t. } X_k X_j \text{ and } X_k \in Adj(X_i, \mathcal{G})$
- 2. if $NA_{ij} \cup T$ is not a clique return FALSE
- 3. if some semidirected path from X_j to X_i does not contain any node in $NA_{ij} \cup T$ return FALSE
- 4. return TRUE

Algorithm 0.6: ValidDelete($\mathcal{G}, \mathcal{E}, \mathcal{H}$)

- 1. $NA_{ij} \leftarrow \text{vertices } X_k \text{ s.t. } X_k X_j \text{ and } X_k \in Adj(X_i, \mathcal{G})$
- 2. **if** $NA_{ij} \setminus H$ is not a clique **return** FALSE
- 3. else return TRUE

SCOREEDGEADDITION and SCOREEDGEDELETION are routines that calculate score differences based on new parent sets. Rebuild keeps track of Markov equivalence relations, basically transforms the current DAG in the procedure to corresponding CPDAG using (R1-R4) just as in PC.

For some omitted details see Chickering (2002) "Optimal structure identification with greedy search," Journal of Machine Learning Research 3: 507-554.

Part 1

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Constraint-based learning: the GES algorithm

The "logic" of GES:

- ▶ the highest scoring model will, in the limit $n \to \infty$, be a member of the true equivalence class
- by adding and removing edges to incrementally improve the score, can achieve a global optimum
- some rules that (un)orient edges to account for Markov equivalence and enforce acyclicity

Score

To make greedy search for CPDAGs globally optimal (converge on the highest scoring, and thus true, structure in the limit) the score we use should satisfy three abstract properties. It turns out that the BIC score satisfies these properties.

Consistency: the true structure \mathcal{G} will maximize the score, and all structures \mathcal{G}' that are not Markov equivalent to \mathcal{G} will have strictly lower score.

Score-equivalence: If $\mathcal G$ is Markov equivalent to $\mathcal G'$, then $S(\mathcal G,D)=S(\mathcal G',D)$.

Decomposability: $S(\mathcal{G}, D) = \sum_{i \in V} s(X_i, Pa(X_i, \mathcal{G}))$

Score properties

Assuming faithfulness, GES finds a global optimum for any consistent, score-equivalent, and decomposable score. It turns out the key property, which is a consequence of these 3, is *local consistency*:

Let \mathcal{G} be any DAG, and let \mathcal{G}' be the DAG that results from adding the edge $X_i \to X_j$. A scoring criterion $S(\mathcal{G}, D)$ is locally consistent if the following two properties hold:

- 1. If $X_i \not\perp X_i | \operatorname{Pa}(X_i, \mathcal{G})$ then $\lim_{n \to \infty} \mathbb{P}(S(\mathcal{G}', D) > S(\mathcal{G}, D)) = 1$
- 2. If $X_i \perp \!\!\! \perp X_j | \operatorname{Pa}(X_j, \mathcal{G})$ then $\lim_{n \to \infty} \mathbb{P}(S(\mathcal{G}', D) < S(\mathcal{G}, D)) = 1$

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Single-edge score differences ⇔ conditional independence tests

Scores and conditional independence

Single-edge score differences ⇔ conditional independence tests.

E.g., for Gaussian dist can show that

$$S(\mathcal{G}', D) - S(\mathcal{G}, D) = n \log(1 - \rho_{ij. \operatorname{Pa}(X_j, \mathcal{G})}^2) + \log(n)$$

Since GES only looks at score *differences* semi/nonparametric versions of this correspondence can be used to construct semi/nonparametric versions of GES (e.g., Huang et al. 2016).

Theory

The BIC score is consistent, score-equivalent (for simple parametric families), and decomposable. Thus it is also locally consistent.

Theorem: Assume the distribution p(x) is Markov and faithful to some DAG \mathcal{G} . Let \mathcal{C} be the CPDAG implied by \mathcal{G} . Let $\widehat{\mathcal{C}}$ be the output of GES using a locally consistent score. Then $\lim_{n\to\infty}\mathbb{P}(\widehat{\mathcal{C}}=\mathcal{C})=1$.

Why is greedy stepwise selection consistent here?

A key result that is that the global optimum of the score will be found (as $n \to \infty$) using only single edge additions/reversals.

A DAG $\mathcal H$ is an independence map (I-map) of a DAG $\mathcal G$ if every independence relationship in $\mathcal H$ holds in $\mathcal G$. We use $\mathcal G \leq \mathcal H$ to denote that $\mathcal H$ is an I-map of $\mathcal G$ and $\mathcal H$ contains more edges than $\mathcal G$.

Meek's conjecture³ states (informal here) that $\mathcal{G} \leq \mathcal{H}$ if and only if we can transform \mathcal{G} into \mathcal{H} by a sequence of (1) covered edge reversals and (2) single edge additions. \Longrightarrow If GES arrives at a local maximum of the score \mathcal{H} , can show that there exists a $\mathcal{G} \leq \mathcal{H}$ and there is a sequence of single-edge moves that transforms \mathcal{H} to \mathcal{G} while improving the score, contradicting that \mathcal{H} was a local maximum.

³Constructive proof in Chickering (2002)

Virtues and drawbacks of score-based search

Virtues:

- easy to parallelize, easy to scale to large numbers of variables
- less prone to statistical errors than constraint-based search, backwards stage can "recover" from some mistakes

Drawbacks:

- need to specify likelihoods, so not so easy to do nonparametrically (mostly restricted to exponential families, though there has been recent work on nonparametric scores)
- ightharpoonup no need⁴ to pick tuning parameter α
- somewhat less developed statistical properties

⁴but... may pick any positive constant factor to multiply the sparsity penalty (enforce more sparsity), which may be treated as a tuning parameter

Uniform consistency results in high-dim settings

There has been an interest in stronger guarantees of asymptotic convergence (rates), esp in settings where $p \gg n$. Strong guarantees invoke stronger assumptions:

- A1 The distribution $p_n(x)$ is multivariate Gaussian and faithful to the DAG \mathcal{G}_n for all n.
- A2 The dimension $p_n = O(n^a)$ for some $0 \le a < \infty$.
- A3 The maximal number of neighbors in the DAG \mathcal{G}_n is denoted by $q_n = \max_{1 \leq j \leq p_n} |\operatorname{Adj}(X_j, \mathcal{G}_n)|$ with $q_n = O(n^{1-b})$ for some $0 < b \leq 1$.
- A4 For all i, j, S, partial correlations are bounded from below and above:

$$\inf\{|p_{ij.S}|: i,j,S \text{ with } p_{ij.S} \neq 0\} \geq c_n,$$
 $c_n^{-1} = O(n^d) \text{ for some } 0 < d < b/2$
$$\sup_{i,j,S} |p_{ij.S}| \leq M < 1$$

Uniform consistency results in high-dim settings

Theorem: Asssume (A1-A4). Let $\widehat{\mathcal{C}}$ be the output of PC with Fisher's Z test of conditional independence test and level α_n . Then there is a sequence of $\alpha_n \to 0$ $(n \to \infty)$ such that $\mathbb{P}(\widehat{\mathcal{C}} = \mathcal{C}) = 1 - O(\exp(-Cn^{1-2d})) \to 1$ $(n \to \infty)$ for some $0 < C < \infty$.

Can choose $\alpha_n = 2(1 - \Phi(n^{1/2}c_n/2))$ and calculate (loose) bound the error.

Similar results exist for a wider range of distributions (semiparametric families, Gaussian copulas), so the Gaussianity is not necessary so long as you have statistical test with appropriate properties.

Very rough sketch of how the proof goes...

Let m_n denote the max size of the conditioning set considered by PC. Call this alg PC(m_n). Let $E_{ij.S}$ denote the event "an error occurs when testing $\rho_{ij.S} = 0$ ".

 $\mathbb{P}(\text{an error occurs in } \mathsf{PC}(m_n))$ $\leq \mathbb{P}\left(\bigcup_{i,j,S} E_{ij,S}\right) \leq O(p_n^{m_n+2}) \sup_{i,j,S} \mathbb{P}(E_{ij,S})$

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 $E_{ij.S} = E_{ij.S}^I \cup E_{ij.S}^{II}$ consists of two possible errors: type I and type II.

$$\sup_{i,j,S} \mathbb{P}(E_{i,j,S}^{I}) \leq O(n-m_n) \exp(-C_1(n-m_n)c_n^2)$$

$$\sup_{i,j,S} \mathbb{P}(E_{ij,S}^{il}) \leq O(n-m_n) \exp(-C_2(n-m_n)c_n^2)$$

Very rough sketch of how the proof goes...

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$$\mathbb{P}(\text{an error occurs in } \mathsf{PC}(m_n))$$

$$\leq \mathbb{P}\left(\bigcup_{i,j,S} E_{ij,S}\right) \leq O(p_n^{m_n+2}) \sup_{i,j,S} \mathbb{P}(E_{ij,S})$$

 $E_{ij.S} = E_{ij.S}^I \cup E_{ij.S}^{II}$ consists of two possible errors: type I and type II.

$$\sup_{i,j,S} \mathbb{P}(E_{ij,S}^{I}) \leq O(n - m_n) \exp(-C_1(n - m_n)c_n^2) \sup_{i,j,S} \mathbb{P}(E_{ij,S}^{II}) \leq O(n - m_n) \exp(-C_2(n - m_n)c_n^2)$$

$$\mathbb{P}(\text{an error occurs in } PC(m_n)) \leq O(p_n^{m_n+2}(n-m_n)\exp(-C_3(n-m_n)c_n^2)) \leq O(n^{a(m_n+2)+1}\exp(-C_3(n-m_n)n^{-2d}))$$

Uniform consistency results in high-dim settings

The previous result is from Kalisch and Buhlmann (2007). The combination of faithfulness and (A4) is called *strong faithfulness* and has been criticized as a strong assumption (Uhler et al. 2013).

Similar results (somewhat stronger assumptions) have been proved for GES (Nandy et al. 2016).

Uniform consistency results for a modified PC (SGS) have been proved under a weaker assumption called k-triangle-faithfulness (Spirtes and Zhang 2014).