Introduction to Causal Discovery

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Part 2: Unmeasured Confounding and More Recent Developments/Challenges

Latent (hidden) variables

In reality, the "true" causal process probably includes a bunch of variables not captured in our data. Unmeasured variables are called "latent" or "hidden" and these pose a real problem for causal inference and causal learning.

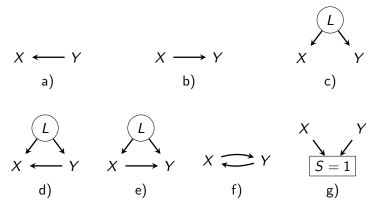
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For example, the underlying causal process may be described by a DAG $\mathcal{G}=(V,E)$ with vertices $V=X\cup L$, but we only observe X.

Latent variables from a structure learning point-of-view

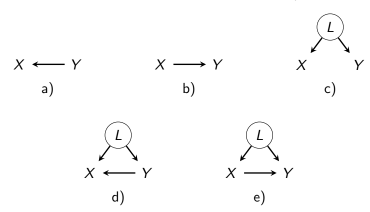
Consider two observed variables X and Y which are known to be dependent. What causal processes may *explain* this dependence?



+ combinations of f) & g) with the others.

Latent variables from a structure learning point-of-view

Consider two observed variables X and Y which are judged to be dependent. What causal processes may explain this dependence? (Let's exclude feedback and selection bias for the time being.)



How could we possibly distinguish between these possibilities from (observed) conditional (in)dependence facts alone?

Distinguishing "real" causality from latent confounding

In general, with just two variables (+ no background knowledge about the latents, no other assumptions) we cannot distinguish between those possibilities. They all imply the same restrictions on the distribution p(x,y): i.e., no restrictions at all.

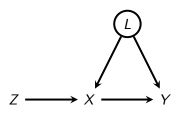
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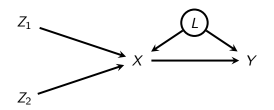
However, with > 2 variables, some *patterns of independence* may help narrow down the structure (assuming faithfulness).

For example:

$$Z \longrightarrow X \longrightarrow Y$$

 \Rightarrow Y and Z are independent given X. (Why?)

Patterns of independence constraints may rule out latent confounding



$$Z_1 \perp \!\!\! \perp Z_2$$

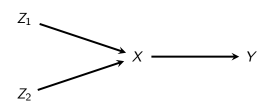
$$Z_1 \not\perp \!\!\! \perp Z_2 | X$$

$$Y \not\perp \!\!\! \perp \{Z_1, Z_2\}$$

$$Y \not\perp \!\!\! \perp Z_1 | X$$

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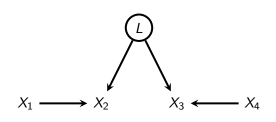
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Patterns of independence constraints may also *suggest* latent confounding



$$X_1 \not\perp X_2$$
 and $X_2 \not\perp X_3$ and $X_3 \not\perp X_4$
 $X_1 \perp X_4$ and $X_1 \perp X_3$ and $X_2 \perp X_4$
 $X_1 \not\perp X_3 \mid X_2$
 $X_2 \not\perp X_4 \mid X_3$

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 $X_1 \not\perp X_3 | X_2$
 $X_2 \not\perp X_4 | X_3$

 \Rightarrow may represent the independence model with a *mixed* graph

Constraint-based structure learning in the presence of latent variables

The assumption of causal sufficiency is rarely warranted in practice!

Fortunately, there exist procedures that allow for arbitrary latent variables. One constraint-based procedure, which follows similar logic to PC, is called the FCI (Fast Causal Inference) algorithm.

We don't want to perform search for the best DAG (or CPDAG) which "fits" the data, since in general no DAG over X will do. We have to consider searching over a different space of graphs.

Latent projections

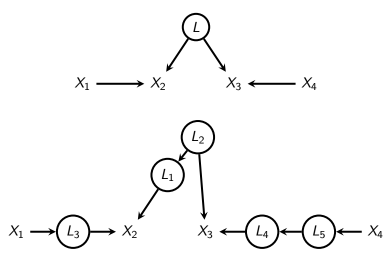
Beginning with a DAG \mathcal{G} , it is useful to think about the induced dependence structure when some variables have been marginalized out. The conditional independence relations in the *marginal* distribution are represented by an ADMG. One may construct an ADMG via the operation of *latent projection*.

Consider a DAG $\mathcal{G} = (V, E)$ with vertex set $V = X \cup L$. The latent projection $\mathcal{G}' = (V', E')$ is a mixed graph with vertex set V' = X such that:

- ▶ for any $X_i, X_j \in X$ there is an edge $X_i \to X_j$ if there exists a directed path from X_i to X_j in \mathcal{G} , with all intermediate nodes on the path in L
- ▶ there is an edge $X_i \leftrightarrow X_j$ if there exists a path from X_i to X_j of the form $X_i \leftarrow \cdots \rightarrow X_j$, where every intermediate node on the path is in L and no consecutive edges on the path are of the form $\rightarrow L_k \leftarrow$ for $L_k \in L$.

Latent projections

Note that an infinite number of distinct latent variable DAGs will share the same latent projection ADMG!



ADMGs

ADMGs preserve conditional independence relations among the *observed* variables. There is a separation criterion which generalizes d-separation to structures with bidirected edges: m-separation.

For A, B, C disjoint subsets of $X: A \perp_d^{\mathcal{G}} B | C \iff A \perp_m^{\mathcal{G}'} B | C$.

Direct structure learning of ADMGs is hard, not well-developed

Though ADMGs have a relatively clear causal interpretation, some nice properties, and lots of associated theory, they are not ideal targets for structure learning. Why?

- Markov equivalence for ADMGs is complicated
- Parameterization of ADMGs is complicated (except for binary variables)
- ADMGs are not maximal

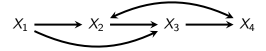
Maximality

Def. A graph $\mathcal G$ is said to be maximal if for every pair of vertices X_i, X_j

$$X_i \not\in \operatorname{Adj}(X_j,\mathcal{G}) \implies \exists X_S \subseteq X \setminus \{X_i,X_j\} \text{ such that } X_i \perp X_j | X_S$$

Thus a graph is maximal if every missing edge corresponds to at least one independence in the model. No additional edge may be added to a maximal graph without changing the independence model. (DAGs are maximal. ADMGs are not.)

Non-maximal ADMG



To make a PC-style search procedure possible, we focus on a class of graphs called Maximal Ancestral Graphs (MAGs). For the purposes of the present discussion, we can view MAGs as a special type of ADMG.

Def. If $X_i \leftrightarrow X_j$ in \mathcal{G} then $X_i \in \operatorname{Sp}(X_j, \mathcal{G})$.

Def. A (directed)¹ ancestral graph \mathcal{G} is a mixed graph (\rightarrow and \leftrightarrow edges) such that $\forall X_i \in X, X_i \notin An(Pa(X_i, \mathcal{G}) \cup Sp(X_i, \mathcal{G}), \mathcal{G})$. That is, an ancestral graph does not contain any directed or almost directed cycles.

Def. A MAG is an ancestral graph that is maximal.

 $^{^{1}}$ MAGs are actually more general than this: they can have undirected (-) edges to represent selection bias in addition to latent confounding, but I'm going to ignore that in this presentation.

MAGs have some pros and cons.

Pros:

- ► They are maximal, so if we find a conditional independence we can remove an edge in a PC-style search.
- We can characterize Markov equivalent MAGs.
- They have other "nice" properties of ADMGs, m-separation works out, etc.

Cons:

- They have a somewhat confusing interpretation!
- Less "informative" than ADMGs

 $X_i \to X_j$ in a MAG means that X_i is an ancestor of X_j in the underlying DAG \mathcal{G} .

 $X_i \leftrightarrow X_j$ means that X_i is not an ancestor of X_j and X_j is not an ancestor of X_i , which implies that there is a latent common cause of X_i and X_j in \mathcal{G} .

NB: An ancestral relationship + latent confounding can coexist! So just because $X_i \to X_j$ in a MAG does not mean there is no latent common cause between X_i and X_j .

$$X_i \longrightarrow X_j$$
 ... in a MAG



. may hide in the underlying DAG

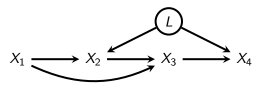
We can construct a MAG from a DAG by a procedure similar to latent projection.

Def. An *inducing path relative to L* is a path on which every vertex not in L (except for the endpoints) is a collider on the path and every collider is an ancestor of an endpoint of the path.

Start with a DAG $\mathcal G$ over $V=X\cup L$ and construct a MAG $\mathcal G'$ over V'=X:

- ▶ for each pair of variables $X_i, X_j \in X$, X_i and X_j are adjacent in \mathcal{G}' iff there is an inducing path between them relative to L in \mathcal{G} .
- ▶ for each pair of adjacent variables X_i, X_j in \mathcal{G}' , orient the edge as $X_i \to X_j$ in \mathcal{G}' if $X_i \in An(X_j, \mathcal{G})$; orient it as $X_i \leftarrow X_j$ in \mathcal{G}' if $X_j \in An(X_i, \mathcal{G})$; orient it as $X_i \leftrightarrow X_j$ in \mathcal{G}' otherwise.

A MAG will have "extra" adjacencies that do not correspond to adjacencies in the underlying DAG. These are adjacencies induced by the latent confounders, and which must be there to preserve maximality.





Partial Ancestral Graphs

A Markov equivalence class of MAGs is represented by a PAG. A PAG is a mixed graph that has $0 \rightarrow$ and $0 \rightarrow$ edges to represent uncertainty about edge endpoints. $0 \rightarrow$ can correspond to a "tail" or "arrowhead."

$$X_1 \circ \longrightarrow X_2 \hookrightarrow X_3 \leftarrow \circ X_4 \circ \longrightarrow \circ X_5$$

Just like a CPDAG represents a set of DAGs, a PAG represents a set of MAGs that each imply the same set of independence constraints.

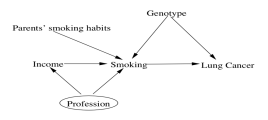


Figure 2: A causal DAG with a latent variable.

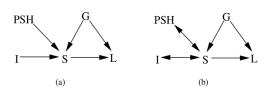


Figure 3: Two Markov Equivalent MAGs.

from Zhang (2008a)

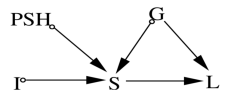


Figure 4: The PAG in our five-variable example.

from Zhang (2008a)

Algorithm 0.1: FCI(Test, α)

```
Input: Samples of the vector X = (X_1, ..., X_n)'
Output: PAG \mathcal{P}

    Form the complete graph P on vertex set X with o→ edges.

2. s \leftarrow 0
     repeat
             for all pairs of adjacent vertices (X_i, X_i) s.t. |\operatorname{Adj}(X_i, \mathcal{P}) \setminus \{X_i\}| > s
4.
               and subsets X_S \subset Adj(X_i, \mathcal{P}) \setminus \{X_i\} s.t. |S| = s
5.
                      if X_i \perp X_i \mid X_S according to (Test, \alpha)
                       then \begin{cases} \text{Delete edge } X_i \circ \circ X_j \text{ from } \mathcal{P}. \\ \text{Let sepset}(X_i, X_i) = \text{sepset}(X_i, X_i) = X_S. \end{cases}
6.
             end
             s \leftarrow s + 1
     until for each pair of adjacent vertices (X_i, X_i), |\operatorname{Adj}(X_i, \mathcal{P}) \setminus \{X_i\}| < s.
9. for all triples (X_i, X_k, X_i) s.t. X_i \in Adj(X_k, \mathcal{P}) and X_i \in Adj(X_k, \mathcal{P})
        but X_i \not\in Adj(X_i, \mathcal{P}), orient X_i *\to X_k \leftarrow *X_i iff X_k \not\in sepset(X_i, X_i).
10. for all pairs (X_i, X_i) adjacent in \mathcal{P} if \exists X_S s.t.
            X_S \in pds(X_i, X_i, \mathcal{P}) or X_S \in pds(X_i, X_i, \mathcal{P}) and X_i \perp X_i \mid X_S according to (Test, \alpha)
                       then \begin{cases} \text{Delete edge } X_i *** X_j \text{ from } \mathcal{P}. \\ \text{Let sepset}(X_i, X_j) = \text{sepset}(X_j, X_i) = X_S. \end{cases}

 Reorient all edges as o-o and repeat step 9.

12. Exhaustively apply orientation rules (R1-R10) in Zhang (2008b) to orient
            remaining endpoints.
```

Let $X \in \operatorname{pds}(X_i, X_j, \mathcal{G})$ if and only if $X \neq X_i$, $X \neq X_j$, and there is a path π between X_i and X in \mathcal{G} such that for every subpath $\langle X_m, X_l, X_h \rangle$ of π either X_l is a collider on the subpath in \mathcal{G} or $\langle X_m, X_l, X_h \rangle$ is a triangle in \mathcal{G} . A *triangle* is a triple $\langle X_m, X_l, X_h \rangle$ where each pair of vertices is adjacent.

Zhang (2008b) refers to "On the completeness of orientation rules for causal discovery in the presence of latent confounders and selection bias." Artificial Intelligence 172: 1873-1896.

13. return \mathcal{P} .

Testing subsets of $pds(X_i, X_i, \mathcal{G})$

FCI performs additional tests compared to PC. In particular, it tests subsets of the set $pds(X_i, X_i, \mathcal{G})$. This is because:

Theorem. If there exists an $X_S \subseteq X \setminus \{X_i, X_j\}$ s.t. $X_i \perp_m X_j | X_S$ in MAG \mathcal{G} , $X_S \subseteq \operatorname{pds}(X_i, X_j, \mathcal{G})$.

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This is the key to removing edges in FCI, and the first part of the algorithm really exists to in order to be able to compute $pds(X_i, X_j, \mathcal{G})$.

Example in R using pcalg package...

Visible edges

Def. Given a MAG \mathcal{M} / PAG \mathcal{P} , a directed edge $X \to Y$ in \mathcal{M} / \mathcal{P} is *visible* if there is a vertex Z not adjacent to Y, such that there is an edge between Z and X that is into X (has an arrowhead at X), or there is a collider path between Z and X that is into X and every non-endpoint vertex on the path is a parent of Y. Otherwise $X \to Y$ is said to be invisible. (All directed edges in a DAG/CPDAG are visible.)

Visible edges

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Directed edges that are visible do not "hide" confounders, they correspond to unconfounded causal effects.

Backdoor criterion for MAGs/PAGs

Def. Let X be a vertex in $\mathcal G$, where $\mathcal G$ represents a causal DAG, CPDAG, MAG, or PAG. Let $\mathcal R$ be a DAG or MAG represented by $\mathcal G$, in the following sense. If $\mathcal G$ is a DAG or MAG, we simply let $\mathcal R=\mathcal G$. If $\mathcal G$ is a CPDAG/PAG, we let $\mathcal R$ be a DAG/MAG in the Markov equivalence class described by $\mathcal G$ with the same number of edges into X as $\mathcal G$. Let $\mathcal R_X$ be the graph obtained from $\mathcal R$ by removing all directed edges out of X that are visible in $\mathcal P$.

Def. Let X and Y be two distinct vertices in mixed graph \mathcal{G} . We say that $V \in \mathsf{Dsep}(X,Y,\mathcal{G})$ if $V \neq X$ and there is a collider path between X and V in \mathcal{G} , such that every vertex on this path is an ancestor of X or Y in \mathcal{G} .

Def. If there is a possibly directed path from X to Y (or if X = Y) then Y is a possible descendent of X. Let $possDe(X, \mathcal{G})$ denote the set of possible descendents of X.

Backdoor criterion for MAGs/PAGs

Theorem.² Let X and Y be two distinct vertices in a causal DAG, CPDAG, MAG, or PAG \mathcal{G} . Let \mathcal{R} and $\mathcal{R}_{\underline{X}}$ be defined as above. If $Y \in \operatorname{Adj}(X,\mathcal{R}_{\underline{X}})$ or $\operatorname{Dsep}(X,Y,\mathcal{R}_{\underline{X}}) \cap \operatorname{possDe}(X,\mathcal{G}) \neq \emptyset$, then $p(y|\operatorname{do}(x))$ is not identifiable via the generalized backdoor criterion. Otherwise $\operatorname{Dsep}(X,Y,\mathcal{R}_{\underline{X}})$ satisfies the generalized backdoor criterion relative to (X,Y) and \mathcal{G} .

That is, when $\mathsf{Dsep}(X,Y)$ satisfies this criterion, it is sufficient to adjust for the variables in $\mathsf{Dsep}(X,Y,\mathcal{R}_{\underline{X}})$ to estimate the causal effect of X on Y.

 $^{^2}$ Maathuis and Colombo (2015) "A generalized back-door criterion," Annals of Statistics, 43(3), 1060-1088.

References on MAGs, PAGs, and FCI

Ali, Richardson, and Spirtes (2009) "Markov equivalence or ancestral graphs," Annals of Statistics 37(5B): 2808-2837.

Colombo, Maathuis, Kalisch, and Richardson (2012) "Learning high-dimensional directed acyclic graphs with latent and selection variables," Annals of Statistics 40(1): 294-321.

Maathuis and Colombo (2015) "A generalized back-door criterion," Annals of Statistics, 43(3), 1060-1088.

Perkovic, Textor, Kalisch, and Maathuis (2018) "Complete graphical characterization and construction of adjustment sets in Markov equivalence classes of ancestral graphs," JMLR 18: 1-62.

Richardson and Spirtes (2002) "Ancestral graph Markov models," Annals of Statistics 30(4): 962-1030.

Spirtes, Scheines, and Glymour (2000) Causation, Prediction, and Search, MIT Press.

Zhang (2008a) "Causal reasoning with ancestral graphs," JMLR 9: 1437-1474.

Zhang (2008b) "On the completeness of orientation rules for causal discovery in the presence of latent confounders and selection bias," Artificial Intelligence 172: 1873-1896.

Assumptions on the structural equations

In the methods discussed so far, we've allowed that the structural equations are *arbitrary* unknown functions (at least, in principle!):

$$X_i = f_i(\mathsf{Pa}(X_i, \mathcal{G}), \epsilon_i) \quad \forall i \in \{1, ..., p\}$$

However, an alternative approach to structure learning makes explicit assumptions on the structural equations. Such assumptions can imply asymmetries in the observed data, which can be used to tease apart different structures. For example, consider a linear model:

$$X_i = \sum_{X_j \in \mathsf{Pa}(X_i,\mathcal{G})} eta_j X_j + \epsilon_i \quad orall i \in \{1,...,p\}$$

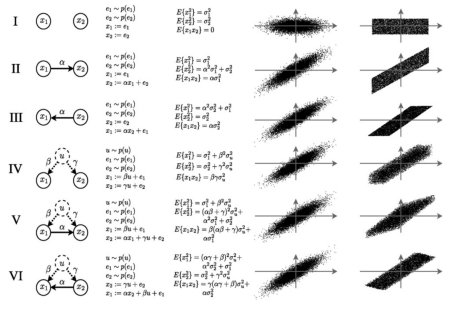
Linear models

Linear models are very common in some areas of applied analysis, particularly because they are convenient to analyze or estimate. However, it is easy to encounter examples for which linearity is obviously false, and **not** an appropriate assumption! If one *does* have reason to expect relationships to be linear, this can be used to significant advantage. Consider the case:

$$X_i = \sum_{X_j \in \mathsf{Pa}(X_i,\mathcal{G})} \beta_j X_j + \epsilon_i$$

 $\forall i$ with $\epsilon_1,...,\epsilon_p$ assumed to be mutually independent and **non-Gaussian**. The combination of linearity and non-Gaussianity³ makes it possible to identify the direction between variables.

 $^{^3}$ Note: Gaussian error terms + linear functions \implies Gaussian joint distribution. A Gaussian joint distribution \implies linear functions. However, linear functions by themselves do **not** imply Gaussianity: you can have models which are linear, with non-Gaussian errors, which \implies non-Gaussian joint distribution.



From Hoyer et al. (2008) "Estimation of causal effects using linear non-Gaussian causal models with hidden variables," Int. Jour. of Approx. Reasoning 49: 362-378. Last 2 columns show induced distributions over x_1, x_2 with Gaussian and Uniform noise, respectively.

LiNGAM

There are a number of algorithms based on the linear non-Gaussian acyclic model ("LiNGAM"), with or without allowing for latent variables.⁴ These typically use results from Independent Component Analysis (ICA) to identify a causal structure consistent with observed data.

When there are no latent variables, these algorithms may identify a unique DAG, rather than an equivalence class. That's because the algorithms exploit information besides conditional independence constraints: the implications of linearity and non-Gaussianity assumptions.

 \Rightarrow you may draw stronger conclusions if you make stronger assumptions; but, those stronger assumptions may be wrong!

⁴See Shimizu (2014) "LiNGAM: Non-Gaussian methods for estimating causal structures," Behaviormetrika 41(1): 65-98.

LiNGAM

Consider the case with no latent variables, as a matrix equation:

$$X = BX + \epsilon$$
$$X = A\epsilon$$

where $A = (I - B)^{-1}$ and ϵ 's are mutually independent. If $B_{ij} \neq 0$ then $X_j \to X_i$. LiNGAM methods use ICA to obtain an estimate of the mixing matrix A.

Actually, ICA typically focuses on estimating the inverse $W=A^{-1}$. Specifically the algorithm will find a matrix \widehat{W}_* such that:

$$\hat{\epsilon} = \widehat{W}_* X$$

with $\hat{\epsilon}$ mutually independent by minimizing $I(\hat{\epsilon}) = \sum_{i=1}^{p} H(\hat{\epsilon}_i) - H(\hat{\epsilon})$ where $H(\hat{\epsilon}) = \mathbb{E}[-\log p(\hat{\epsilon})]$. It can be shown that this mutual information metric is minimized when the elements of ϵ are mutually independent (which is what the model assumes).

LiNGAM

Since ICA only determines \widehat{W}_* up to a permutation of the columns and a scaling factor, the algorithm will permute and normalize the result appropriately to compute \widehat{B} , pruning coefficients close to zero if they are "small."

ICA solves the "cocktail party problem": recovering the "source" signals from "microphones" which linearly mix them. Fast algorithms for doing this have been explored in the engineering literature.

LV-LiNGAM

To allow for latent variables one may use *overcomplete* ICA: more "sources" than than "microphones." ⁵

Need to estimate A in X = Ae where A is non-square and we have only observed the variables in X. For example:

$$\begin{pmatrix} X_1 \\ X_2 \end{pmatrix} = \begin{bmatrix} \alpha & \beta & \gamma \\ \delta & \eta & \xi \end{bmatrix} \begin{pmatrix} e_1 \\ e_2 \\ u \end{pmatrix}$$

⁵See Hoyer et al. (2008) "Estimation of causal effects using linear non-Gaussian causal models with hidden variables," Int. Jour. of Approx. Reasoning 49: 362-378.

LV-LiNGAM

Very roughly...

Let $x^{(n)}$ denote the data matrix with sample size n and θ denote the full set of LV-LiNGAM parameters.

If the distribution of each error term is represented by a weighted mixture of Gaussians, then $p(x^{(n)}|\theta)$ can be expressed in closed form.

Under some assumptions, we can find the $\hat{\theta}$ that maximizes $p(x^{(n)}|\theta)$ using an Expectation-Maximization (EM) algorithm.

Practical issues

Since (overcomplete) ICA leaves indeterminacy wrt permutations of columns and does not produce exact zeros, various heuristics (search over possible permutations, shrinking "small" coefficients to zero, etc.) are involved in applying LV-LiNGaM.

Also in this case multiple models may be observationally equivalent, and so the procedure does not return a unique DAG.

LiNGAM etc.

Pro:

▶ By assuming linear non-Gaussian SEMs, algs can sometimes identify a unique DAG (assuming no unmeasured confounding) or a small equivalence class (allowing for unmeasured confounding)

Cons:

- Even with non-Gaussian errors, linearity assumption is very strong and unlikely to hold
- Statistical (asymptotic) properties of ICA-based algorithms are unknown/complicated, may depend on "degree of non-Gaussianity"
- ▶ If errors are too close to Gaussian or too non-Gaussian, may not perform well
- In practice, requires large sample sizes and small dimension p

Additive noise nodels

Another class of models assumes only that the noise terms enter into the function additively:

$$X_i = f_i(Pa(X_i, \mathcal{G})) + \epsilon_i$$

the functions f_i may be nonlinear (though are usually assumed to be differentiable). Somewhat surprisingly, assuming additive Gaussian noise + nonlinear functions is sufficient to identify the causal structure.

Note that the ANM is **not closed under marginalization**. If you start with an ANM over $X \cup L$, then the marginal model over X may no longer be in the ANM class. \implies in settings with latent variables, ANMs are difficult to justify.

Post-nonlinear causal models

Another class of models adds a nonlinear transformation on the additive noise model. Consider $X_i \to X_i$. The PNL model asserts

$$X_j = f_2(f_1(X_i) + \epsilon_j) \quad \epsilon_j \perp X_i$$

If the opposite direction $X_i \to X_i$ holds true, then

$$X_i = g_2(g_1(X_j) + \epsilon_i) \quad \epsilon_i \perp X_j$$

One may prove that under some technical conditions, $X_j \to X_i$ and $X_j \leftarrow X_i$ can be distinguished from the data.

Exploiting asymmetries

All of these semi-parametric methods impose some assumptions/restrictions on the structural equations, and derive some asymmetry in the observed data distribution from these assumptions. Then, check if the data exhibits the supposed asymmetry to try and infer backwards to the generating model.

In general, it difficult to establish properties of such methods and also computationally quite difficult to scale them up to large multivariate systems.

However, they can sometimes be combined with nonparametric methods like PC, etc to get more informative output. Also, new methods are being developed all the time.

Differentiable causal discovery

A recent approach to discovery combines score-based selection w/ ideas from continuous optimization (+ in practice, semi-parametric assumptions).

- Assume that the true DGP is an SEM with parameter matrix θ and $\mathcal{G}(\theta)$ the corresponding induced graph.
- Let \mathbb{G} denote the space of possible graphs (e.g., all DAGs over X).
- ▶ Finally, let $S(X; \theta)$ denote a consistent score that is minimized at the true θ .

Recast the discrete optimization problem as a continuous program:

$$\min_{\theta \in \Theta} S(X; \theta) \qquad \iff \min_{\theta \in \Theta} S(X; \theta)$$
s.t. $G(\theta) \in \mathbb{G}$ s.t. $h(\theta) = 0$.

 $h(\theta)$ is a differentiable function that = 0 iff $\mathcal{G}(\theta) \in \mathbb{G}$

Differentiable causal discovery for DAGs

Consider linear SEMs: $X_i = \sum_{j \in V} \theta_{ji} X_j + \epsilon_i \text{ w/ independent errors}$

and let $\ensuremath{\mathbb{G}}$ denote the space of DAGs. Can show that

$$h(\theta)=\operatorname{tr}(e^{\theta\circ\theta})-p=0$$

iff $\mathcal{G}(\theta)$ is a DAG, where \circ is the Hadamard product and e^A is the matrix exponential of A.

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s.t. $\mathcal{G}(\theta) \in \mathbb{G}$ s.t. $\operatorname{tr}(e^{\theta \circ \theta}) - p = 0$.

The gradient $\nabla h(\theta) = (e^{\theta \circ \theta})^T \circ 2\theta$ has closed form and so this can be solved by SOA constrainted optimization techniques (e.g., augmented Langrangian w/ dual ascent).

Differentiable causal discovery for ADMGs

Consider linear SEMs w/ correlated errors:

$$X_i = \sum_{j \in V} \delta_{ji} X_j + \epsilon_i \text{ and } \beta = \mathbb{E}[\epsilon \epsilon^T]$$

and let $\mathbb G$ denote the space of ancestral graphs. Can show that

$$h(\delta,\beta) = \operatorname{tr}(e^{\delta \circ \delta}) - p + \operatorname{sum}(e^{\delta \circ \delta} \circ (\beta' \circ \beta')) = 0$$

iff $\mathcal{G}(\delta, \beta)$ is ancestral, where $\beta'_{ij} = \beta_{ij}$ for $i \neq j$ and 0 otherwise.

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Autograd can be used to obtain analytic gradients for an augmented Lagrangian optimization scheme.

Solving the continuous program

$$\min_{\theta \in \Theta} S(X; \theta) + \frac{\rho}{2} |h(\theta)|^2 + \alpha h(\theta),$$

where ρ is the penalty weight and α is the Lagrange multiplier.

Then solve the dual equation: $\alpha^{k+1} \leftarrow \alpha^k + \rho^k h(\theta^k)$

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Lots to be said about the stability and convergence properties of diff optimization schemes here, but I am not knowledgable about this.

See, e.g., Ng et al. (2022) "On the convergence of continuous constrained optimization for structure learning" in AISTATS.

Scores for differentiable causal discovery

Zheng et al. (2018) us an ℓ_1 -penalized likelihood for (DAGs) whereas Bhattacharya et al. (2021) use (for ADMGs) an approximation to the BIC score.

NB: fitting likelihoods for (linear) ancestral ADMGs is a bit tricky – Bhattacharya et al. use the residual iterative proportional fitting (RICF) procedure (Drton et al. 2009).

NB: for both DAGs and ancestral ADMGs, the procedure is blind to equivalence class considerations. If the score is consistent and a global optima is achieved, then output should be Markov equivalent to the true \mathcal{G} . One option is to transform the output into the corresponding CPDAG or PAG. Local optima may be an issue.

Differentiable causal discovery for ADMGs

Bhattacharya et al. (2021) extend this idea to additional classes of ADMGs, i.e., Arid and Bow-free ADMGs, that can be more informative than ancestral graphs because they encode general equality constraints (Verma constraints) in addition to conditional independence constraints.

Learning Arid or Bow-free ADMGs is an interesting direction for causal discovery, but there are many challenges and unknowns, esp. because there is no convenient characterization of Markov equivalence for these graphs.

Differentiable causal discovery

- ► There is something very appealing about this approach b/c we can make use of impressive SOA continuous optimization algs
- ▶ In practice, people have pointed out some "undesriable behavior" or "sensitivity" of these methods:
 - ▶ Reisach et al. (2021) "Beware of the simulated DAG..." in NeurIPS.
 - ▶ Ng et al. (2022) "On the convergence of continuous constrained optimization for structure learning" in AISTATS.
 - ▶ Ng et al. (2024) "Structure learning with continuous optimization: a sober look and beyond" in CLeaR.
- ► There are open theoretical and practical questions about the properties of these algorithms, when we expect them to converge, etc.
- ▶ It is also not so obvious how to make things work when we weaken assumptions of linear SEM models (e.g., fully non-parametric SEM), non-uniqueness of solutions, etc.

Permutation-based causal discovery

Permutation-based algorithms formulate the discovery problem as a search over possible (partial) causal orderings of the vertices in a graph:

$$\underset{\pi \in \Pi}{\operatorname{arg max}} S(\mathcal{G}_{\pi})$$

where

$$S(\mathcal{G}_\pi) = egin{cases} -|\mathcal{G}_\pi| & ext{if } \mathcal{G}_\pi ext{ is Markov wrt } \mathcal{I}(\mathcal{G}_\pi) \ -\infty & ext{otherwise}. \end{cases}$$

each (partial) order in Π induces a graph \mathcal{G}_{π} and a set of conditional independence constraints $\mathcal{I}(\mathcal{G}_{\pi})$. The score compares graphs by their sparsity, and the major challenge is to find a way to efficiently traverse the space of orderings Π .

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In practice, we do not know $\mathcal{I}(\mathcal{G}_{\pi})$ but rather execute a sequence of conditional independence tests depending on the ordering π .

Permutation-based causal discovery

$$rg \max_{\pi \in \Pi} S(\mathcal{G}_{\pi})$$

Solus et al. (2018) present a greedy search procedure over the space of total causal orderings (DAGs) and show that it is consistent.

Bernstein et al. (2020) present a greedy search over partial causal (ancestral) orderings (MAGs).

Structure learning algorithms... incomplete list

DAGs/CPDAGs:

PC (+ variants), GES, ARGES, GSP, MMHC, ICA-LiNGAM, directLiNGAM, CAM, SAT-methods, NOTEARS (+ variants), GRaSP, BOSS

MAGs/PAGs/ADMGs:

► FCI, RFCI, FCI+, GFCI, GSPo, LV-LiNGAM, M3HC, SAT-methods, Differentiable CD

Cyclic graphs:

► CCD, LiNG, bcause, Two-Step, FASK, SAT-methods

constraint-based, score-based, hybrid, semiparametric, other

Software packages for structure learning

```
R:
pcalg
bnlearn
```

```
Python:
pcalg.py
causal discovery toolbox (cdt)
causal-learn
ananke
```

Java:

TETRAD

Matlab:

Bayes net toolbox

+ various implementations available from authors of papers

Open problems

- Better general-purpose nonparametric conditional independence tests
- Consistent, scaleable, nonparametric, and more accurate algs for learning MAGs/PAGs/ADMGs
- ► Characterizations of Markov equivalence for general ADMGs
- Better understanding and methods for learning graphs w/ confounding + cycles (DMGs)
- Consistency results that weaken "strong faithfulness" assumption
- Post-selection inference
- **.**..