Discovering Cyclic Causal Models in Psychological Research

Kyuri Park^{1*}, Lourens J. Waldorp², and Oisín Ryan¹

¹Department of Methodology and Statistics, Utrecht University ²Department of Psychology, University of Amsterdam

Abstract

Statistical network models have become popular tools for analyzing multivariate psychological data. In empirical practice, network parameters are often interpreted as reflecting causal relationships – an approach that can be characterized as a form of causal discovery. Recent research has shown that undirected network models are likely to perform poorly as causal discovery tools in the context of discovering acyclic causal structures, a task for which many alternative methods are available. However, acyclic causal models are likely unsuitable for many psychological phenomena, such as psychopathologies, which are often characterized by cycles or feedback loop relationships between symptoms. A number of cyclic causal discovery methods have been developed, largely in the computer science literature, but they are not as well studied or widely applied in empirical practice. In this paper, we provide an accessible introduction to the basics of cyclic causal discovery for empirical researchers. We examine three different cyclic causal discovery methods and investigate their performance in typical psychological research contexts by means of a simulation study. We also demonstrate the practical applicability of these methods using an empirical example and conclude the paper with a discussion of how the insights we gain from cyclic causal discovery relate to statistical network analysis.

Keywords: Constraint-based, cyclic causal discovery, directed cyclic graph (DCG), partial ancestral graph (PAG), statistical network model

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^{*}Correspondence concerning this paper should be addressed to: Kyuri Park, Department of Methodology and Statistics, Utrecht University, Padualaan 14, 3584 CH Utrecht, The Netherlands. e-mail: kyurheep@gmail.com

1 Introduction

A fundamental task in various disciplines of science is to understand the mechanisms or causal relations underlying the phenomena of interest. In psychology, one of the core questions is how psychopathology comes about, with the network theory positing that mental disorder is produced by a system of direct causal interactions between symptoms (Borsboom & Cramer, 2013). Given this theoretical framework, statistical network models have become popular tools for analyzing multivariate observational data (Robinaugh et al., 2020; Epskamp et al., 2018a). In practice, empirical researchers often interpret the conditional statistical relationships estimated in a network model as reflecting the causal relationships between symptoms of mental disorder and/or other psychological variables — an approach that can be characterized as a form of causal discovery (Spirtes et al., 2000; Peters et al., 2017; Ryan et al., 2022). However, it has been shown that network models are likely to perform poorly as causal discovery tools; relations in the network may not reflect the direct causal effects that researchers aim to discover, as they may be produced by unwittingly conditioning on common effects or by failing to account for unobserved confounders (Dablander & Hinne, 2019; Ryan et al., 2022).

In the field of causal discovery, using patterns of statistical (in)dependence estimated from observational data to infer causal structures is known as *constraint-based* causal discovery (Spirtes & Glymour, 1991). Ryan et al. (2022) suggest that network models could be replaced by purpose-built constraint-based causal discovery methods. However, the most popular and well-studied constraint-based methods assume that causal relationships are *acyclic*; that is if X causes Y, then Y does not cause X (Glymour et al., 2019). This is problematic since *cyclic* relationships or *feedback loops* are critical to the theoretical understanding of psychopathology (Borsboom, 2017). For example, Wittenborn et al. (2016) suggest that several different causal feedback loops, such as *perceived stress* \rightarrow *negative affect* \rightarrow *rumination* \rightarrow *perceived stress* play a key role in sustaining depression. Such theoretical expectations necessitate the use of *cyclic causal discovery* methods.

Although some constraint-based cyclic causal discovery algorithms have been developed (Mooij & Claassen, 2020; Richardson, 1996a; Strobl, 2019), they have not been as well studied as their acyclic counterparts. In part, this is due to the practical difficulties in fitting and interpreting cyclic causal models. For example, in the context of structural equation modeling, it is well known that models are usually identified as long as they are acyclic, whereas this is not necessarily true for cyclic models (Bongers et al., 2021; Bollen, 1989). Furthermore, interpreting the output of cyclic causal discovery algorithms is significantly more challenging than for acyclic counterparts, since direct statistical dependencies may not reflect direct causal dependencies when cyclic relationships are present (Bongers et al., 2018; Hyttinen et al., 2013). Recent work has explored the application of *invariance-based* algorithms — another type of cyclic causal discovery methods — to psychological data (Kossakowski et al., 2021). However, these methods require multiple datasets that measure the same variables but in different settings, such as a mix of observational and experimental data, which

is a disadvantage compared to constraint-based methods that can be applied using only a single observational dataset (Peters et al., 2016; Glymour et al., 2019).

To our knowledge, little to no research has been done on the applicability of constraint-based cyclic causal discovery methods in psychology, and much remains unknown about their performance. Therefore, in this paper, we aim to address the following question: How well do constraint-based cyclic causal discovery methods perform in typical psychological research contexts? First, we will provide an accessible overview of the different cyclic causal discovery methods, including the assumptions under which they are expected to work and how the output of these methods should be interpreted. Second, we investigate, by means of a simulation study, how well these different methods perform under various circumstances. In the simulation study, we study more and less ideal situations, by varying the sample size, the size and density of the underlying network, and the presence or absence of unobserved confounding variables. Third, we demonstrate the practical applicability of these methods by applying them to empirical data (McNally et al., 2017) and discussing how the insights we gain relate to the statistical network analysis of the same data.

2 Background

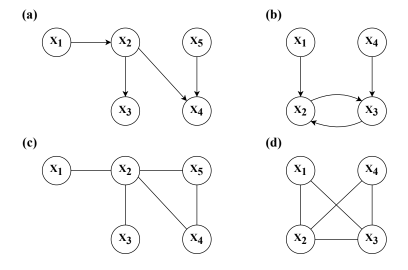
In this section, we will establish the basic concepts of graphical models and examine how constraint-based causal discovery methods operate. We will first introduce different types of graphical models, while demonstrating the differences between statistical and causal graphical models using example graphs. Then, we will explain the assumptions that underlie constraint-based causal discovery methods, as well as the technical difficulties that arise in the presence of cycles. Lastly, we will illustrate each step of the constraint-based causal discovery procedure using a simple example of a directed graph.

2.1 Graphical Models

A graph (\mathcal{G}) is a diagram made up of a set of vertices (\mathcal{V}) and edges (\mathcal{E}), describing the connections between the vertices, denoted as $\mathcal{G} = (\mathcal{V}, \mathcal{E})$. A probabilistic graphical model uses a graph to express the conditional (in)dependencies between random variables, where the vertices represent random variables, and the edges encode the conditional dependencies that hold among the set of variables (Lauritzen, 1996).

There exist various graphical models that differ in the types of edges they allow (e.g., directed vs. undirected) and how edges are interpreted in terms of statistical relationships. One commonly known graphical model in psychology is the *Pairwise Markov Random Field* (PMRF) — an undirected graph where edges indicate a statistical association between variables after controlling for all other variables — which forms the basis of statistical network models (Epskamp et al., 2018b; Epskamp & Fried, 2018). In PMRFs, edges are strictly undirected, and the presence of an edge

Figure 1. Example causal graphical models and corresponding PMRF models.



Note. (a) is the example directed acyclic graph (DAG). (b) is the example directed cyclic graph (DCG). (c) is the PMRF (Pairwise Markov Random Field) corresponding to the DAG in (a). (d) is the PMRF corresponding to the DCG in (b).

A - B indicates that A and B are statistically dependent, conditional on the set of all other variables in the network (Borsboom et al., 2021).

In a *causal* graphical model, on the other hand, the edges describe *causal* relationships between variables; the edges are typically directed, with $A \rightarrow B$ denoting that intervening on A results in a change in the probability distribution of B (Geiger & Pearl, 1990). The simplest and commonly used causal graph is a *directed acyclic graph* (DAG), also known as a *Bayesian network*, which consists of directed edges without any cycles (Pearl, 1988). When a causal graph contains cycles, it is referred to as a *directed cyclic graph* (DCG). Two examples of causal graphical models are shown in Figure 1. Figure 1(a) does not contain any cycles, whereas Figure 1(b) does; hence they are called a DAG and DCG, respectively.

Causal graphical models also describe patterns of statistical independencies, which can be read off from the graph using Pearl's *d-separation criterion* (Geiger et al., 1990). In principle, the d-separation criterion can be applied to both acyclic and cyclic causal graphs, as long as certain conditions, discussed in Section 2.2, are met. For instance, in Figure 1(a), we see a *chain* structure $X_1 \rightarrow X_2 \rightarrow X_3$, which implies that X_1 and X_3 are marginally dependent $(X_1 \not\perp X_3)$ but independent conditional on X_2 ($X_1 \perp X_3 \mid X_2$). More formally, we would say X_1 and X_3 are *d-connected* given the empty set but *d-separated* by X_2 . A *fork* structure $X_3 \leftarrow X_2 \rightarrow X_4$ implies the same pattern of independencies; X_3 and X_4 are marginally dependent ($X_3 \not\perp X_4$) but independent conditional on X_2 ($X_3 \perp X_4 \mid X_2$). However, a *collider* structure $X_2 \rightarrow X_4 \leftarrow X_5$ implies a contrasting pattern; here X_2 and X_5 are marginally independent ($X_2 \perp X_5$) but dependent conditional on X_4 ($X_2 \not\perp X_5 \mid X_4$).

This distinguishing characteristic of colliders is crucial when identifying the directions of causal relationships, as will be shown in section 2.3.

Having established the basics, we can now examine how PMRF-based statistical network models relate to different types of directed causal models. In Figure 1(c), we show the PMRF model that corresponds to the DAG in Figure 1(a), where an additional edge is introduced between $X_2 - X_5$ due to conditioning on the common effect (i.e., collider), X4. In Figure 1(d), the PMRF model corresponding to the DCG in Figure 1(b) is shown. Two spurious edges are induced in the PMRF network (e.g., $X_1 - X_3$ and $X_2 - X_4$) because of conditioning on the colliders X_2 and X_3 . These examples illustrate the limitations of statistical network models in inferring patterns of directed causal relationships. PMRFs can contain spurious edges resulting from conditioning on common effects, and the possibility of producing collider structures is likely to be higher with the presence of cycles, exacerbating this issue. Further elaborating on this, Ryan et al. (2022) suggest that causal discovery methods specifically designed for this task are likely to outperform statistical network models in learning the underlying causal structure, indicating that network models may not be desirable tools for discovering causal relationships. In the following sections, we will focus on how constraintbased causal discovery methods recover the causal structure while looking into the assumptions they require. Additionally, we will explore the practical and conceptual difficulties involved in performing causal discovery in the presence of causal cycles.

2.2 Acyclic vs. Cyclic Causal Graphs

The d-separation criterion described above applies to all acyclic graphs, but for graphs with cycles, it applies only under certain conditions. To understand these conditions, we first need to introduce some graph terminology. In the field of graphical models, we use kinship terminology to describe a graph structure, as follows:

if
$$\begin{cases} A \to B \\ A \leftarrow B \\ A \to \cdots \to B \text{ or } A = B \\ A \leftarrow \cdots \leftarrow B \text{ or } A = B \end{cases}$$
 in \mathcal{G} then A is a
$$\begin{cases} \text{parent} \\ \text{child} \\ \text{ancestor} \\ \text{descendant} \end{cases}$$
 of B and
$$\begin{cases} A \in pa_{\mathcal{G}}(B) \\ A \in ch_{\mathcal{G}}(B) \\ A \in an_{\mathcal{G}}(B) \\ A \in de_{\mathcal{G}}(B) \end{cases}$$
.

Also, when there exists an edge between two vertices A - B, A and B are said to be *adjacent*. For example, in Figure 1(b), $X_1 \in pa_{\mathcal{G}}(X_2)$, $X_2 \in ch_{\mathcal{G}}(X_1)$, $\{X_1, X_2, X_3, X_4\} \in an_{\mathcal{G}}(X_3)$, $\{X_1, X_2, X_3\} \in de_{\mathcal{G}}(X_1)$, and X_2 is adjacent to X_1 and X_3 . With this terminology in place, we can define the conditions that relate patterns of causal dependency in a causal graph to patterns of statistical dependency between random variables. First, we introduce the *global Markov* condition, which states that d-separation relations represented in causal graphs can be used to read off statistical independence relations such that:

if
$$X_A \perp \!\!\! \perp_G X_B \mid X_C \Longrightarrow X_A \perp \!\!\! \perp X_B \mid X_C$$
 for all disjoint subsets of X_A, X_B, X_C ,

where $\coprod_{\mathcal{G}}$ refers to d-separation in graphs, and \coprod refers to statistical independence between random variables. If causal graphs are *acyclic*, such as DAGs, then the *global Markov* condition holds regardless of the functional forms of causal relations and the distributions of variables involved (Lauritzen, 1996). In addition, in DAGs, the *global Markov* condition entails the *local Markov* condition, which states that a variable is independent of its non-descendants given its parents (Lauritzen, 2001). The fact that one Markov property implies the other comes in handy when reading off conditional independencies from a graph.

In contrast to the acyclic case, the situation is not as straightforward in *cyclic* graphs. In DCGs, the global Markov property does not always hold. Spirtes (1994) showed that this property does hold when causal relations are *linear* and the error terms are *independent*. However, even in this case, the global Markov property does not imply the local Markov property. For example, in Figure 1(b), the global Markov property is preserved $(X_1 \perp \!\!\!\perp_{\mathscr{G}} X_4 \mid \{X_2, X_3\} \Longrightarrow X_1 \perp \!\!\!\perp X_4 \mid \{X_2, X_3\})$, but the local Markov property is violated as $X_2 \perp \!\!\!\perp_{\mathscr{G}} X_4 \mid X_3$ (i.e., X_2 is *not* independent of its non-descendant X_4 given its parent X_3). This is because X_3 serves as both a parent of X_2 and a collider on the path $X_2 \longrightarrow X_3 \longleftarrow X_4$ at the same time.

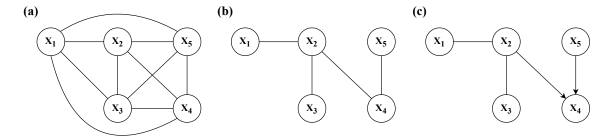
In the current paper, we limit the scope of our study to cyclic causal graphs that represent *linear* causal relations with jointly *independent* error terms, so for which the global Markov condition is satisfied. This type of assumption is common in many statistical modeling traditions in psychology and social sciences. For example, structural equation models and popular statistical network models, such as the Gaussian graphical model (GGM), also rely on similar assumptions about the linearity of statistical relations (Epskamp et al., 2018b; Bollen & Long, 1993). In addition, we make use of one more assumption, known as *faithfulness*, which is required for constraint-based causal discovery (Pearl, 2009; Spirtes et al., 2000). The *faithfulness* assumption is essentially the reverse of the global Markov condition, stating that statistical independencies map onto the structure of causal graphs such that:

$$X_A \perp \!\!\!\perp X_B \mid X_C \Longrightarrow X_A \perp \!\!\!\perp_G X_B \mid X_C.$$

In other words, it postulates that if two variables are (conditionally) statistically independent of each other, then this implies that they are causally independent of each other, ruling out, for instance, that different paths in a causal system exactly cancel one another out.¹ Together with the global Markov property, faithfulness enables us to make inferences about causal relationships represented in graphs by testing for the statistical independence among variables (Peters et al., 2017; Pearl, 1988).

¹For a discussion of this assumption in the context of psychological network analysis, see Ryan et al. (2022).

Figure 2. Steps of a constraint-based method.



Note. (a) shows the fully-connected graph for the example DAG from Figure 1(a), which is the initial starting point of the algorithm. (b) shows the estimated *skeleton* — an undirected graph of the underlying causal structure — after the first step. (c) shows the resulting graph after the second step, which represents the *Markov equivalence class* of DAGs (i.e., a set of DAGs that entail the same set of conditional independencies).

2.3 A Primer on Constraint-Based Causal Discovery

Under the aforementioned assumptions, constraint-based methods seek to recover the underlying causal structure by testing for conditional independence relations between variables from observational data (Scheines et al., 1998). Assuming linear relations with additive Gaussian errors, conditional independence can be tested using partial correlations (Lawrance, 1976), although notably a number of non-parametric conditional independence tests can also be used under less stringent assumptions (Li & Fan, 2020; Huang et al., 2016). Constraint-based methods typically employ a two-step procedure; first, establishing the *skeleton* — an undirected version of the underlying graph — and second, attempting to assign directions to the edges. In general, constraint-based techniques are unable to uniquely identify the underlying causal graph, but instead return a set of causal graphs that imply the same statistical independence relations (Spirtes et al., 2000).

To develop an intuition for how constraint-based methods work, we will examine how they operate on the relatively simple DAG from Figure 1(a). The method begins with a fully-connected graph, as shown in Figure 2(a). In the first step, the *skeleton* is estimated by testing for conditional independence; if two variables are independent when conditioning on *any* subset of the remaining variables (e.g., $X_1 \perp \!\!\!\perp X_3 \mid X_2, X_1 \perp \!\!\!\perp X_4 \mid X_2, X_1 \perp \!\!\!\perp X_3 \mid \{X_2, X_4\}, etc.$), then the edge between those two variables is removed (see Figure 2(b)). This is based on the principle that, in acyclic graphs, two variables are always statistically dependent (regardless of any conditioning set), if and only if a direct causal relation exists between them. In other words, we can identify the presence of direct causal paths between variables by testing whether they are statistically dependent given any subset of the remaining variables. However, while this principle allows us to detect the presence of an edge, it does not tell us the direction of that edge (e.g., $X_2 \rightarrow X_4$ or $X_2 \leftarrow X_4$). In the second step, we attempt to orient the edges in the skeleton by searching for *colliders* that induce distinctive

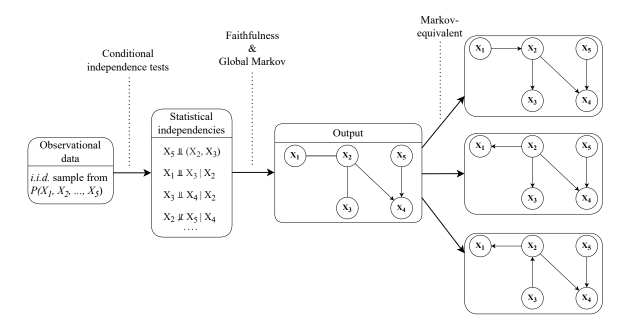


Figure 3. Summary of the constraint-based causal discovery procedure.

Note. A constraint-based algorithm starts with performing a series of conditional independence tests on observational (*i.i.d.*: independent and identically distributed) data. Under the faithfulness and global Markov assumption, the algorithm estimates a graph structure based on the observed statistical independence patterns. The output is a *partially directed* graph (as some edges remain undirected). It can represent multiple graphs that are *Markov equivalent*, meaning that they imply the same statistical independence relations. This set of equivalent graphs is referred to as a *Markov equivalence class*, and in this example, it consists of three different DAGs.

patterns of independencies (e.g., X_4 is identified as a collider since $X_2 \perp \!\!\! \perp X_5$ and $X_2 \not \perp X_5 \mid X_4$, thus $X_2 \rightarrow X_4 \leftarrow X_5$ is oriented; see Figure 2(c)). This procedure we have described is essentially the PC algorithm (Spirtes et al., 2000), and the output of the PC algorithm (Figure 2(c)) is called a complete partially directed acyclic graph (CPDAG). Note that the resulting CPDAG is not identical to the original true graph \mathcal{G} , as the two edges between $X_1 - X_2$ and $X_2 - X_3$ remain undirected. There are, in fact, three DAGs implied by the CPDAG, which are obtained by assigning directions to the undirected edges, excluding the combinations that introduce new colliders. These DAGs are called *Markov equivalent*, meaning that they encode the same conditional (in)dependencies (i.e., the same d-separation relations hold), and we call such a set of equivalent graphs a *Markov equivalence class*, denoted by $Equiv(\mathcal{G})$ (Spirtes et al., 2000). These Markov equivalent DAGs are shown in the right-hand side of Figure 3, which summarizes the constraint-based causal discovery procedure that we just described. This highlights a general difficulty in constraint-based causal discovery, which is that there are usually multiple graphs that are consistent with an observed set of statistical independencies.

Constraint-based methods for cyclic causal discovery operate under similar principles as those described earlier. However, cyclic causal discovery is in general more challenging, and the problem of having multiple graphs that are Markov equivalent is often exacerbated when cycles are allowed (Richardson & Spirtes, 1999). Consider how, in the DAG example above, we could identify the presence of direct causal relations between variables when they are statistically dependent given any subset of the remaining variables. Now suppose we apply the same rule to the DCG shown in Figure 1(b). In this cyclic graph, there is no direct causal path between X_1 and X_3 , but instead there is the path $X_1 \rightarrow X_2 \stackrel{\longleftarrow}{\longrightarrow} X_3$. When testing for conditional independencies, we find that X_1 and X_3 are not marginally independent because of the causal chain $X_1 \rightarrow X_2 \rightarrow X_3$. However, unlike in the acyclic case, we also find that X_1 and X_3 are not conditionally independent given X_2 , since X_2 also acts as a collider on the path $X_1 \rightarrow X_2 \leftarrow X_3$. In cyclic graphs, two variables can be statistically dependent conditional on every subset of the remaining variables, even when there is no direct causal relation between them. Therefore, when a cycle exists, a constraint-based method often cannot directly identify parental relations but recovers only up to ancestral relations, typically leading to a larger Markov equivalence class. This also means that the estimated skeleton of a cyclic graph represents ancestral relations, and is thus called the ancestral skeleton. Although the same principles of causal discovery that we described for DAGs can be adapted and used for DCGs, some additional constraints and orientation rules are required to address the complexities arising from the presence of cycles. In the next section, we will further elaborate on these constraints and rules while introducing several constraint-based cyclic causal discovery algorithms.

3 Causal Discovery Algorithms

In the previous section, we introduced the key concepts of graphical models and the fundamental principles of constraint-based causal discovery methods. We also outlined some challenges associated with estimating cyclic causal structures. In the following, we will provide a detailed description of three different constraint-based algorithms for cyclic graphs: *cyclic causal discovery* (CCD) (Richardson, 1996b), *fast causal inference* (FCI) (Mooij & Claassen, 2020), and *cyclic causal inference* (CCI) (Strobl, 2019). We will discuss their assumptions, steps involved, and output graphs along with their interpretation.

3.1 Assumptions of Algorithms

All three algorithms build upon the same principles and hence can be seen as extensions of the PC algorithm described in Section 2.3, but they entail slightly different assumptions. The CCD algorithm assumes *causal sufficiency*, which means that all common causes of variables have been measured, and there is no unobserved confounding. This is a critical assumption in causal discovery, as unobserved confounders can induce an edge between variables when there is no direct causal

	CCD	FCI	CCI
Global Markov condition	✓	\checkmark	✓
Faithfulness	\checkmark	\checkmark	\checkmark
Acyclicity	×	<u>_</u> a	X
Causal sufficiency	\checkmark	X	X
Absence of selection bias	\checkmark	<u>_</u> a	X
Linearity	\checkmark	<u>_</u> a	\checkmark

Table 1. Assumptions of cyclic causal discovery algorithms.

Note. ^a FCI was originally designed to infer causal structure in the presence of selection bias assuming acyclicity, but a recent study has proposed that it performs comparably well in the cyclic settings under certain conditions (Mooij & Claassen, 2020). Specifically, these conditions require that selection bias is *absent* and variables share *non-linear* relations.

relation between them (Lauritzen, 1996; Spirtes et al., 2013). Without this causal sufficiency assumption, causal discovery becomes more challenging, as an edge between two variables can result from a direct causal link but can also be induced by an unobserved common cause. The other two algorithms, FCI and CCI, relax this sufficiency assumption and account for the possibility of latent confounders. This makes them potentially more promising for psychological research, where the assumption of unobserved confounding is rarely warranted (Rohrer, 2018).

Another closely related concept to sufficiency is *selection bias*. Selection bias occurs when one conditions on an unobserved collider, for example, by selectively excluding a particular subgroup of samples, which leads to a similar problem of inducing spurious causal relations (Versteeg et al., 2022; Haslbeck et al., 2022). While the CCD algorithm assumes no selection bias, the FCI and CCI algorithms account for the possibility of selection bias. Although the FCI algorithm was initially designed for acyclic causal structures (Spirtes et al., 2000), it has been shown to perform well in cyclic settings under a more generalized Markov condition, while ruling out the presence of selection bias (Mooij & Claassen, 2020). Thus, we consider FCI as one of the cyclic causal discovery algorithms, but note that the suggested conditions by Mooij and Claassen (2020) hold under limited situations excluding linear and discrete cases. Table 1 summarizes the set of assumptions made by each of the algorithms, including the fundamental assumptions of global Markov condition, faithfulness, and acyclicity, as well as those related to the functional forms of causal relations and error terms.

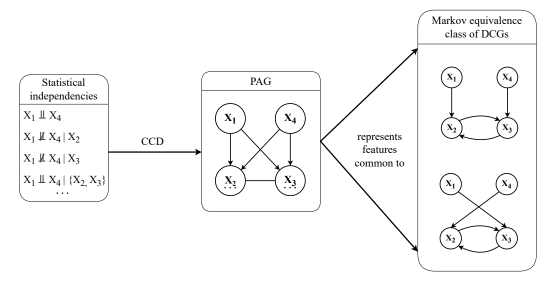
In Section 2.3, we learned that constraint-based causal discovery works by testing for patterns of conditional (in)dependence, resulting in an equivalence class of causal graphs that convey the same statistical (in)dependencies. The PC algorithm was used to illustrate this process, which involves two steps and yields a CPDAG as the output. For cyclic causal graphs, constraint-based algorithms follow a similar approach, but with some caveats. First, the specific steps taken are more complex

than those of the PC algorithm. Second, the output of these algorithms is not a CPDAG, but a different representation of equivalent cyclic graphs known as a *partial ancestral graph* (PAG). In the subsequent sections, we will elaborate on each cyclic causal discovery algorithm, beginning with an overview of its output, the PAG, and how it can be interpreted. We will then describe the specific steps taken to estimate the PAG.

3.2 CCD Algorithm

The CCD algorithm is considered relatively simple among the three algorithms, as it assumes that there is no unobserved latent confounding (i.e., *causal sufficiency*). The basic operation of CCD is summarized in Figure 4. The fundamental principles on which the CCD algorithm operates are similar to those of the PC algorithm, as illustrated in Figure 3. However, the output graph for CCD is a PAG, which represents the common features of equivalent directed cyclic graphs (DCGs). In what follows, we will explain how to interpret a PAG in detail using the example PAG shown in Figure 4.

Figure 4. Summary of CCD algorithm operation.



Note. Given the observed statistical independencies, CCD constructs a partial ancestral graph (PAG), which represents the *ancestral* features that are common to every directed cyclic graph (DCG) in a Markov equivalence class. In this particular example, the Markov equivalence class consists of two different DCGs.

3.2.1 CCD Output Representation: Partial Ancestral Graph (PAG)

As was the case for DAGs, there typically exist multiple DCGs that imply the same statistical independencies and so are statistically indistinguishable from one another. To represent a set of equivalent DCGs, the CCD algorithm uses a PAG that characterizes the common features shared by all equivalent DCGs, Equiv(G). As discussed in Section 2.3, the causal semantics of edges in PAGs become more complicated due to the presence of cyclic relations. In a CPDAG, an edge represents a direct causal relation between the corresponding vertices, while no edge implies its absence. In a PAG, the absence of an edge still indicates the absence of a direct causal relation, but the presence of an edge indicates causal *ancestry*, with $A \rightarrow B$ meaning that A is an *ancestor* of B. In the causal graphs we have looked at so far, the types of edges are limited to directed (\rightarrow) and undirected edges (\rightarrow). In PAGs, however, three different types of edge-endpoints { \circ , >, -} are utilized to represent the ancestral relations in Equiv(G). The interpretation of each edge-endpoint in a PAG is as follows:²

- 1. $A *\rightarrow B$ is interpreted as B is not an ancestor of A in every graph in Equiv(G).
- 2. A*-B is interpreted as B is an ancestor of A in every graph in Equiv(G).
- 3. $A*-\circ B$ is interpreted as the ancestral relation of B with regard to A is undetermined or varies across graphs in Equiv(G).

The PAG output of the CCD algorithm can also include a solid or dotted underlining to provide additional information about the causal relations in triplets. If there is a solid underlining $A*-*\underline{B}*-*C$, it indicates that B is an ancestor of (at least one of) A or C in every graph in Equiv(G). If there is a dotted underlining added to a collider structure such as $A \rightarrow \underline{B} \leftarrow C$, it indicates that B is *not* a descendant of a common child of A and C in every graph in Equiv(G). For example, from the PAG shown in Figure 4, we can read off the following:

- 1. X_2 and X_3 are not ancestors of X_1 and X_4 in every graph in Equiv(G).
- 2. X_1 and X_4 are both ancestors of X_2 and X_3 in every graph in Equiv(G).
- 3. X_2 is an ancestor of X_3 and X_3 is an ancestor of X_2 in every graph in Equiv(G), indicating the presence of a cyclic relationship between them.
- 4. X_2 and X_3 are not descendants of a common child of X_1 and X_4 in every graph in $Equiv(\mathcal{G})$. This means that it is not possible for both $X_1 \rightarrow X_2$ and $X_4 \rightarrow X_2$, or both $X_1 \rightarrow X_3$ and $X_4 \rightarrow X_3$ to coexist in any graph in $Equiv(\mathcal{G})$. For instance, if X_1 were to be a parent of X_2 , and considering that X_2 and X_3 are ancestors/descendants of each other, X_4 cannot also be a parent of X_2 ; otherwise this condition would be violated.

Given the causal ancestral relations represented by the example PAG described above, we can correspondingly derive the Markov-equivalent DCGs, which are shown in the right-hand side of Figure 4.

²In the description of the semantics for PAGs (Richardson, 1996b), * is used as a *meta-symbol*, indicating one of the three possible edge-endpoints. For instance, $A \longrightarrow B$ indicates any of the following edges: $A \longrightarrow B$, or $A \longrightarrow B$.

3.2.2 Steps of CCD Algorithm

In this section, we will provide a thorough demonstration of the CCD algorithm, which is the first theoretically well-founded constraint-based method that can be applied in a cyclic setting. While FCI and CCI share a similar procedure, they differ in their specific orientation rules. Therefore, we will not cover the steps of FCI and CCI as extensively as we do for CCD but rather highlight any changes or additions made by these algorithms. Note that some of the steps described below can be technical. While it is helpful to have a good understanding of the whole CCD procedure, it is not essential for comprehending the rest of the paper. Readers may choose to skim through some of the steps, particularly those that outline specific orientation rules of CCD (e.g., step 3, step 5, and step 6).³

The CCD algorithm consists of six steps. We illustrate each step using the example DCG from Figure 5(a), which is the same example DCG that we previously introduced in Figure 1(b). The algorithm starts with a fully-connected PAG with circle endpoints, as shown in Figure 5(b), which implies that the direction has not been determined yet. As it proceeds, (some) circles will be replaced by either an arrow head or a tail.

Step 1. The *ancestral* skeleton — an undirected graph of ancestral relations implied by the underlying structure — is estimated based on conditional independencies. This step is identical to the first step of the PC algorithm described above in Section 2.3; the algorithm tests whether two vertices, A and B, are statistically independent given any subset (S) of the remaining variables, and when such a subset is found (i.e., A and B are d-separated given S), then the algorithm removes $A \circ - \circ B$ and records $S = \mathbf{Sepset}\langle A, B \rangle = \mathbf{Sepset}\langle B, A \rangle$. A \mathbf{Sepset} , short for separation set, is a set of variables that d-separates two other sets of variables in a graph. Since $X_1 \perp \!\!\!\perp X_4 \mid \emptyset$ in our example DCG, $X_1 \circ - \circ X_4$ is removed and $\mathbf{Sepset}\langle X_1, X_4 \rangle = \mathbf{Sepset}\langle X_4, X_1 \rangle = \emptyset$ is recorded, which results in Figure 5(c).

Step 2. Again, the algorithm proceeds in a similar manner to the PC algorithm by searching for collider structures. In technical terms, the algorithm looks for triplets A*-*B*-*C where A and B are not directly connected $(A \perp\!\!\!\perp B \mid S)$ but $B \notin \mathbf{Sepset}(A, C)$, meaning that $A \not\!\!\!\perp C \mid B$. Once the algorithm identifies B as a collider, the triplet is oriented as $A \rightarrow B \leftarrow C$. Given that $X_2 \notin \mathbf{Sepset}(X_1, X_4)$ and $X_3 \notin \mathbf{Sepset}(X_1, X_4)$ in our example, $X_1 \circ -\circ X_2 \circ -\circ X_4$ and $X_1 \circ -\circ X_3 \circ -\circ X_4$ are oriented respectively as $X_1 \rightarrow X_2 \leftarrow X_4$ and $X_1 \rightarrow X_3 \leftarrow X_4$, resulting in Figure 5(d).

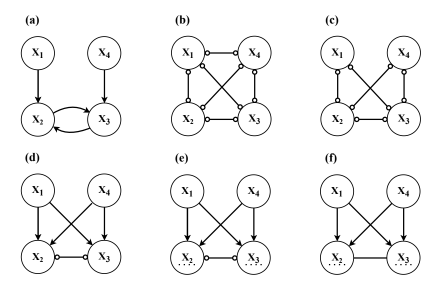
Step 3. The algorithm then checks for additional d-separating relations in triplets to perform additional orientations. If the algorithm finds a triplet $\langle A, B, C \rangle$ such that:

(i) A is not adjacent to B or C, (ii) B and C are adjacent, and (iii) $B \notin \mathbf{Sepset}(A, C)$,

then B*-*C is oriented as $B\leftarrow C$. This rule is based on the inference that when there is a d-connecting path from A to B given $\mathbf{Sepset}\langle A, C \rangle$, and there is no d-connecting path from A to C given $\mathbf{Sepset}\langle A, C \rangle$, then B cannot be an ancestor of C. If it were, the d-connecting path between A

³For a more concise and technical description of the CCD algorithm, please refer to Appendix A.

Figure 5. Trace of CCD algorithm.



Note. (a) shows the true directed cyclic graph. (b) shows the fully-connected PAG, which is the starting point of the algorithm. (c) shows the *ancestral* skeleton (i.e., an undirected version of the PAG) estimated in step 1. (d) shows the state of the PAG after step 2, where some of the edges are oriented given the identified colliders. (e) shows the state of the PAG after step 4, where the *Supsets* are identified and the corresponding colliders are dotted-underlined. (f) shows the final state of the PAG after step 5, where an additional edge between X_2 and X_3 is oriented.

and B could be extended to a d-connecting path between A and C given **Sepset** $\langle A, C \rangle$. As we find no such d-separating relations in our example, no further orientations are performed in step 3.

Step 4. In this step, the algorithm tries to refine the causal graph by adding underlinings to collider structures. Recall that underlinings in PAGs can convey additional information on causal relations in triplets, as mentioned in Section 3.2.1. Exploring the possibility of adding underlinings, the algorithm searches for **Supsets**, which stands for *super separation sets*. They are a set of variables that d-separates two endpoint vertices in a collider structure when conditioning on the collider. Hence, for each collider structure $A \rightarrow B \leftarrow C$, the algorithm checks if there is any **Supset**, that is, a set T including B that d-separates A and C. When such exists, $T = \text{Supset}\langle A, B, C \rangle$ is recorded and a dotted-underlining $A \rightarrow B \leftarrow C$ is added. Since $X_1 \perp \!\!\!\perp X_4 \mid \{X_2, X_3\}$ in our example, $\text{Supset}\langle X_1, X_2, X_4 \rangle = \text{Supset}\langle X_1, X_3, X_4 \rangle = \{X_2, X_3\}$ is recorded and each collider is dotted-underlined as $X_1 \rightarrow X_2 \leftarrow X_4$ and $X_1 \rightarrow X_3 \leftarrow X_4$, resulting in Figure 5(e). This indicates that X_2 and X_3 are not descendants of a common child of X_1 and X_4 .

Step 5. The last two steps of CCD concern additional orientation of the remaining undirected edges by examining **Supsets** that are found in the previous step 4. Now we expand and look for quadruplets — four ordered vertices $\langle A, B, C, D \rangle$ — where:

(i) $A \rightarrow B \leftarrow C$, (ii) $A \rightarrow D \leftarrow C$ or $A \rightarrow D \leftarrow C$, and (iii) B and D are adjacent. If $D \in \textbf{Supset}(A, B, C)$

in such quadruplets, orient B*-*D as B*-D, or else orient B*-*D as $B\to D$. This essentially makes use of the fact that $\mathbf{Supset}\langle A,B,C\rangle$ is not a descendant of a common child of A and C; if D belongs to the $\mathbf{Supset}\langle A,B,C\rangle$, then D is not a descendant of B, the common child of A and C, and hence is oriented as B*-D. In our example, there is such a quadruplet; (i) $X_1\to X_2\leftarrow X_4$, (ii) $X_1\to X_3\leftarrow X_4$, and (iii) X_2 and X_3 are adjacent. Since $X_2\in \mathbf{Supset}\langle X_1,X_3,X_4\rangle$ and $X_3\in \mathbf{Supset}\langle X_1,X_2,X_4\rangle$, $X_2\circ -\circ X_3$ is oriented as $X_2-\circ X_3$, and $X_2-\circ X_3$ is subsequently oriented as X_2-X_3 , resulting in Figure 5(f). This indicates that X_2 and X_3 are ancestors of each other, implying a cyclic causal relationship between them.

Step 6. In the final step, the algorithm continues to make additional inferences about ancestral relations in quadruplets by inspecting **Supsets**. This time, the algorithm searches for quadruplets $\langle A, B, C, D \rangle$, where $A \rightarrow \underline{B} \leftarrow C$ while D is not adjacent to either A or C. If A and C are d-connected given **Supset** $\langle A, B, C \rangle \cup D$, then orient $B* \leftarrow D$ as $B \rightarrow D$. This works based on the following inference; given A and C are d-separated by **Supset** $\langle A, B, C \rangle$ while $B \in \text{Supset}\langle A, B, C \rangle$, and if D is an ancestor of B, then it means that A and C are d-separated by **Supset** $\langle A, B, C \rangle \cup D$. In other words, if A and C are d-connected given **Supset** $\langle A, B, C \rangle \cup D$, then D is not an ancestor of B, and subsequently it can be oriented as $B \rightarrow D$. In our example, no such quadruplets exist.

The final PAG is shown in Figure 5(f), from which we can determine the Markov equivalence class of DCGs. To do so, we first need to identify all the ancestral relationships that are represented in the PAG, such as X_1 and X_4 being ancestors of X_2 and X_3 , and X_2 and X_3 being ancestors of each other. Furthermore, the dotted underlinings on X_2 and X_3 provide additional information that allows us to rule out cases where X_1 and X_4 are ancestors of both X_2 and X_3 simultaneously. By taking into account all the available information, we can derive two distinct DCGs that correspond to the given structural relationships, as depicted in the right-hand side of Figure 4.

3.3 FCI Algorithm

The FCI algorithm, originally proposed by Spirtes et al. (2000), is a constraint-based causal discovery method for directed acyclic graphs (DAGs), which takes into account the presence of latent confounding and possible selection bias. Recently, Mooij and Claassen (2020) demonstrated that the FCI algorithm can also be applied to cyclic causal discovery in the presence of latent confounding under more general faithfulness and Markov conditions, provided that the causal relationships are *non-linear*. For details on these conditions, we refer readers to Forré and Mooij (2017). In the following, we will outline the basic principles of the FCI algorithm and discuss how its output can be interpreted.

3.3.1 FCI Output Representation: Partial Ancestral Graph (PAG)

The FCI algorithm, like the CCD algorithm, aims to identify the underlying causal graph up to its Markov equivalence class and also employs a PAG to represent the common ancestral features

among the equivalent graphs. However, allowing latent confounders adds a complication; DCGs are not closed under marginalization over latent confounders, meaning that there exist infinitely many DCGs of observed variables (O) and latent confounders (L) that entail the same set of independencies (Richardson & Spirtes, 2002). This problem arises from the fact that we do not know how many latent confounders are involved, and the algorithm has to account for the possibilities of arbitrarily many latent confounders (Zhang & Spirtes, 2005).

To represent the presence of latent confounders in the infinite space of causal graphs, we need to introduce a new type of edge into the PAG representation. Specifically, we take bidirected edges (\leftrightarrow) to reflect the presence of latent confounders, with $A \leftrightarrow B$ denoting a confounding variable between A and B.⁴ The interpretation of edges in the PAGs estimated by the FCI algorithm is otherwise the same as in the CCD algorithm, except for the fact that in FCI PAGs, fully-connected vertices with circle endpoints $(\circ \multimap)$ may indicate a possible cyclic structure (Mooij & Claassen, 2020). In what follows, we will go over how the FCI algorithm estimates a PAG given the same example DCG used for the CCD algorithm (Figure 1(b)) and look at the interpretation of its output in more detail.

3.3.2 Steps of FCI Algorithm

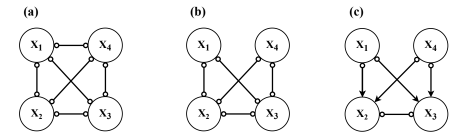
The algorithm consists of three main steps: skeleton discovery, collider structure orientation, and application of further orientation rules, where the first two steps are analogous to the CCD procedure. As with the CCD algorithm, the FCI algorithm begins with a fully-connected PAG with \circ — edges between every pair of variables (Figure 6(a)). Then, it estimates the ancestral skeleton (Figure 6(b)) by testing for statistical independence (see step 1 of CCD). Subsequently, the FCI algorithm searches for colliders in the same way as the CCD algorithm (see step 2 of CCD); when a collider (*B*) is identified, A*-*B*-*C is oriented as $A*-B\leftarrow*C$, resulting in Figure 6(c). Lastly, the FCI algorithm executes a set of orientation rules to further orient the edges. For a complete list of orientation rules (Zhang, 2008), see Appendix B. In this case, no additional endpoints are oriented in further steps, leaving Figure 6(c) as the final resulting PAG. Given this PAG, we can read off that:

- 1. X_2 and X_3 are not ancestors of X_1 and X_4 in every graph in Equiv(G).
- 2. X_2 and X_3 might be part of a cycle in \mathcal{G} as they are fully-connected with circle endpoints.

Notice that the PAG produced by FCI has more circle endpoints (o) than the PAG produced by CCD, which indicates a greater degree of uncertainty about causal ancestral relationships. This is because the FCI algorithm accounts for the possibility of latent confounding, leading to a larger space of possible graphs. Consequently, there are many more equivalent graphs that conform to the relational structure implied by the FCI PAG, resulting in a larger Markov equivalence class, as illustrated in the right-hand side of Figure 7.

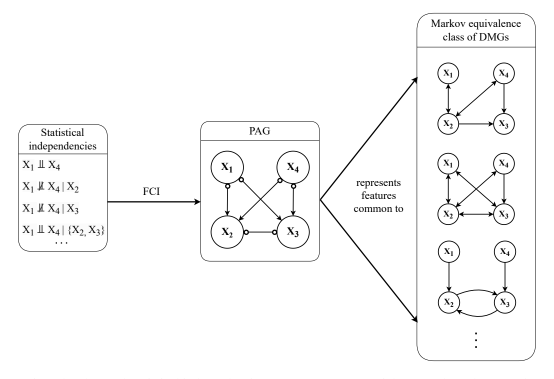
⁴The class of graphs that make use of bidirected edges (←) to represent latent confounding is called *directed mixed graphs* (DMGs), which can be seen as extensions of DCGs (Richardson, 2003).

Figure 6. Trace of FCI algorithm.



Note. (a) shows the fully-connected PAG, which is the starting point of the algorithm. (b) shows the *ancestral* skeleton estimated in the same manner as the CCD algorithm. (c) shows the state of the PAG after the orientation step using the collider structures identified in step 2.

Figure 7. Summary of FCI algorithm operation.



Note. Given the observed statistical independencies, FCI constructs a partial ancestral graph (PAG) that captures the common *ancestral* features of every directed mixed graph (DMG) in a Markov equivalence class. The PAG estimated by FCI has more circle endpoints than the one estimated by CCD in Figure 4, indicating a higher level of uncertainty about the causal relationships. This is because FCI accounts for the presence of latent confounders, which is represented by bidirected edges (\leftrightarrow) in the graph. As a result, the Markov equivalence class tends to be relatively large.

3.4 CCI Algorithm

The CCI algorithm combines features of both the CCD and FCI algorithms. It can identify cyclic causal structures, similar to CCD, and can handle latent confounding, similar to FCI (Strobl, 2019). Employing a combined approach, however, comes at a cost; the algorithm requires more complex edge-endpoint inferences and lengthy orientation rules. For a detailed explanation of each step of the CCI algorithm, see Appendix C.

3.4.1 CCI Output Representation: Partial Ancestral Graph (PAG)

As with the other two algorithms, CCI generates a PAG that captures the common ancestral relationships among equivalent graphs. To account for the presence of latent confounding in the infinite causal graph space, as described in Section 3.3.1, CCI also uses bidirected edges (↔). Apart from that, the interpretation of the other types of edges in PAGs estimated by CCI is the same as that described in Section 3.2.1 for the CCD output. In the following section, we will briefly outline the steps of CCI with the same example DCG (from Figure 1(b)) that has been used throughout the paper and examine the interpretation of the resulting PAG.

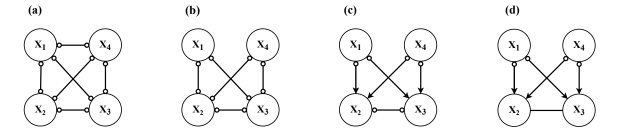
3.4.2 Steps of CCI Algorithm

The CCI algorithm consists of seven steps in total, the first two of which are identical to those of the other algorithms (i.e., skeleton discovery and collider structure orientation), and the remaining steps are similar to the further orientation rules implemented in CCD and FCI. Same as the others, CCI initiates with a PAG that is fully connected with \sim 0 edges between every pair of variables (Figure 8(a)). After running the skeleton discovery procedure (i.e., step 1), the ancestral skeleton is estimated, as shown in Figure 8(b). Upon orienting the edges based on identified colliders (i.e., step 2), CCI produces the output shown in Figure 8(c). In the following orientation step (i.e., step 5), the edge between X_2 and X_3 is oriented utilizing **Supsets** — similar to step 4 of the CCD algorithm — which results in Figure 8(d). Since no additional edges are oriented in the subsequent steps, Figure 8(d) remains the final PAG. From this PAG, we can read off the following:

- 1. X_2 and X_3 are not ancestors of X_1 and X_4 in every graph in Equiv(G).
- 2. X_2 is an ancestor of X_3 and X_3 is an ancestor of X_2 in every graph in $Equiv(\mathcal{G})$, implying a cyclic relationship between them.

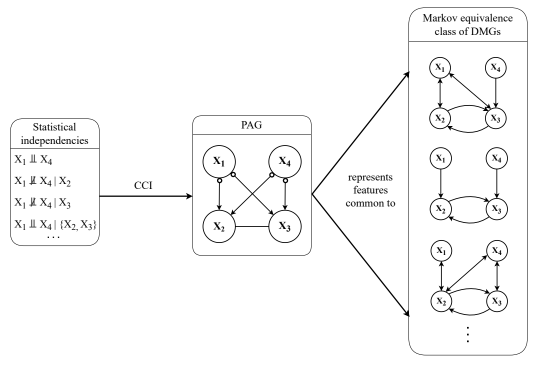
Similar to FCI, the PAG obtained from CCI has more circle endpoints compared to the one obtained from CCD, due to its consideration of latent confounding. Figure 9 summarizes the operation of CCI, which is similar to that of FCI shown in Figure 7, with a relatively large Markov equivalence class of graphs. However, unlike the PAG from FCI where cycles are implied by the fully-connected edge with circle endpoints (\circ — \circ), the PAG from CCI clearly indicates a cyclic relationship between X_2 and X_3 with an undirected edge (—), which implies the reciprocal ancestral relationship between the two variables.

Figure 8. Trace of CCI algorithm.



Note. (a) depicts the fully-connected PAG, which is the starting point of the algorithm. (b) depicts the *ancestral* skeleton estimated in the same way as the CCD and FCI algorithms. (c) depicts the state of the PAG after orienting the collider structures in step 2. (d) depicts the state of the PAG after applying the extra orientation rules in step 5. No further orientation takes place in the subsequent steps, leaving (d) as the final PAG.

Figure 9. Summary of CCI algorithm operation.



Note. CCI constructs a partial ancestral graph (PAG) based on the observed statistical independencies, which captures the *ancestral* features common to every directed mixed graph (DMG) in a Markov equivalence class. Similar to the FCI algorithm, CCI accounts for the potential presence of latent confounding, which is represented by bidirected edges (\leftrightarrow), leading to more circle endpoints in the PAG. This generally results in a larger Markov equivalence class, as illustrated in the right-side of the figure.

3.5 Overview of Algorithms

Thus far, we have introduced three constraint-based causal discovery algorithms for cyclic graphs and discussed their characteristics. Based on our example (DCG from Figure 1(b)), the CCD algorithm may appear to be the preferred option, as it has less uncertainty in its output and correspondingly returns a smaller set of equivalent graphs. However, recall that the main advantage of the other two algorithms — FCI and CCI — is their ability to handle the presence of latent confounding, which commonly occurs in psychological research in practice (Rohrer, 2018).

Figure 10 shows the resulting PAGs of each algorithm given two different example DCGs; one without a latent confounder and the other with a latent confounder. All three algorithms happen to generate the same PAG for these two different DCGs. Suppose the true underlying causal structure is the DCG without a latent confounder (top left in Figure 10). In that case, all three PAGs correctly depict the ancestral features of the true DCG. However, the PAG output by CCD is by far the most informative, as it contains no circle endpoints, thus representing the smallest equivalent set of graphs. Now, suppose the underlying causal structure is the DCG with a latent confounder (bottom left in Figure 10). Then, the PAG generated by CCD contains errors (i.e., X_1 is not an ancestor of X_2 and X_3), while the PAGs estimated by FCI and CCI correctly represent the ancestral features of the true DCG. In the following section, we will conduct a simulation study to compare the performance of each algorithm under various conditions and investigate which algorithm is most suitable for situations likely to arise in psychological research.

Figure 10. Comparison of the partial ancestral graphs (PAGs).

Note. Two different DCGs are displayed in the first column, with subsequent columns showing the corresponding PAGs estimated by each algorithm. In this particular case, all three algorithms happen to produce identical PAG estimates for both DCGs. LC = latent confounder; DCG = directed cyclic graph; PAG = partial ancestral graph.

4 Simulation

In the previous section, we outlined the characteristics of three causal discovery algorithms, and analyzed their respective assumptions and procedures. In this section, we present a simulation study to evaluate the performance of these algorithms under different conditions by varying the sample size, number of variables, density, and presence of latent confounders. We investigate how these factors affect each algorithm's performance and whether any algorithm outperforms the others in specific ways. We also perform sensitivity analyses to test the robustness of our findings to specific choices made regarding the simulation conditions. In the following, we will first discuss the data generation process, simulation design, and evaluation metrics. We will then present the results on each algorithm's performance using various metrics, followed by the findings of our sensitivity analyses.

4.1 Data Generation

We simulate data from the different cyclic models, as shown in Figure 11, all of which are characterized by *linear* relations and *independent Gaussian* error terms. These types of models are often used in psychological research, and for such cyclic models, the global Markov property — the necessary condition for constraint-based causal discovery — also holds, as discussed in Section 2.2.

To generate data, we first define a coefficient matrix **B**. A non-zero entry B_{ij} indicates an edge from X_j to X_i with a strength of B_{ij} . Thus, $X_1, ..., X_p$ can be generated according to the following equation:

$$X_i = \sum_{r=1}^p B_{ir} X_r + \varepsilon_i,$$

for i=1,...,p, where p is the number of vertices and ε are mutually independent $\mathcal{N}(0,1)$ random variables. The variables $X_1,...,X_p$ then have a multivariate Gaussian distribution with a mean vector of zero and a covariance matrix $\Sigma = (\mathbf{I} - \mathbf{B})^{-1}(\mathbf{I} - \mathbf{B})^{-T}$, where \mathbf{I} denotes the identity matrix. Note that this data generation scheme is possible provided that $(\mathbf{I} - \mathbf{B})$ is invertible, which is the case when the eigenvalues of \mathbf{B} are smaller than one in absolute value, $|\lambda| < 1$ (Eberhardt et al., 2010). While this is guaranteed if \mathbf{B} defines an acyclic model (Drton & Maathuis, 2017), for cyclic models, this does not always hold. To satisfy this condition, cyclic relations need to be not too strong so that the dynamical system converges to equilibrium (Rothenhäusler et al., 2015). We set fixed values for \mathbf{B} to make the simulation results easier to track and interpret. When specifying the \mathbf{B} matrices, we choose values within a range that is deemed reasonable (e.g., restricting the strength of cyclic relations to relatively small such that they do not diverge) and verify that the eigenvalues satisfy the equilibrium condition.⁵ When this condition is violated, we adjust the parameters until it is satisfied. In addition, to ensure that the simulation results are not dependent on the specific values of the coefficient matrix \mathbf{B} , we perform a sensitivity analysis. In this analysis, we randomly sample parameter values of \mathbf{B} from a uniform distribution on $[-0.9, -0.1] \cup [0.1, 0.9]$ in each iteration. We

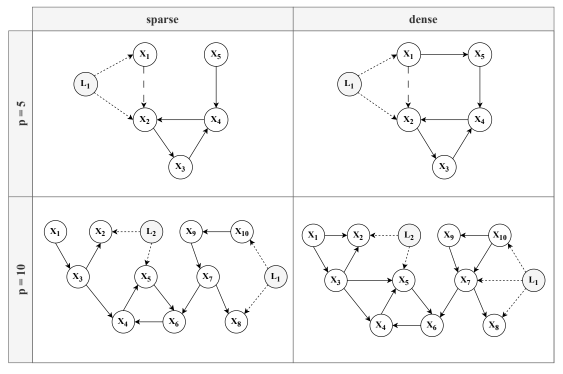
⁵For more details on the specific **B** matrices used in the simulations, please refer to Appendix D.

then check whether the equilibrium condition is met, and if not, we re-sample the parameters until it is satisfied. In the following, we provide a detailed description of the simulation setup, which is replicated across both simulation studies.

4.2 Simulation Setup

We test each algorithm under different conditions by varying the number of variables (rows in Figure 11) and the number of edges — the density (columns in Figure 11). We also explore the impact of unobserved confounding by introducing latent variables (L_1 and L_2 in Figure 11), as latent confounding is a common issue in psychological research, particularly when it comes to inferring causal relationships (Hallquist et al., 2021; Rohrer et al., 2022). Lastly, we vary the sample size ranging from small to moderately large, $N \in \{50, 150, 500, 1000, 2000, 3000, 4000, 5000, 7500, 10000\}$, for every simulated cyclic model. This leads to a $2 \times 2 \times 2 \times 10$ design; number of variables \times density \times latent confounder (presence/absence) \times sample size. We simulate each condition 500 times, estimating three PAGs per iteration using each algorithm. See Figure 11 for an overview of the different types of cyclic models used in the simulation.

Figure 11. Simulation settings.



Note. In our simulation study, we vary several factors such as the number of variables: $p \in \{5, 10\}$, the density: sparse / dense, the influence of a latent confounder: absence / presence, and the sample size: $N \in \{50, 150, 500, 1000, 2000, 3000, 4000, 5000, 7500, 10000\}$. This results in a $2 \times 2 \times 2 \times 10$ simulation design, with each combination of factors (except for N) yielding a unique model structure. Note that the edge between X_1 and X_2 (long-dashed line —) in the 5-variable models (top row) is only present in the conditions without a latent confounder.

To test for conditional independence, we employ partial correlations since the variables in our simulated data have linear relations with additive Gaussian errors. In such cases, conditional independence is equivalent to zero partial correlation (Lawrance, 1976). Throughout the study, we use a fixed alpha level of 0.01 ($\alpha = 0.01$) for conducting the conditional independence tests, which is a commonly used value in causal discovery studies with moderate sample sizes (Malinsky & Danks, 2018). However, we acknowledge that following the fixed α convention is not straightforwardly justified (Strobl et al., 2017). To address this, we perform another sensitivity analysis where we adjust the α level based on the sample size. Further elaboration on this analysis will be provided in Section 4.5. All simulations are performed using R software version 4.2.3 (R Core Team, 2023).

4.3 Evaluation Metrics

As discussed in Section 3, the algorithms attempt to recover the ancestral graphs implied by the underlying DCG. In order to evaluate the performance of each algorithm, we compare how well it recovers the true ancestral graph. To do this, we first need to construct a true ancestral graph for each simulated condition, which can then be compared to the estimated graphs. The true ancestral graph for each DCG considered in our simulation study is shown in Figure 12. For those interested, a step-by-step procedure for deriving a true ancestral graph from a DCG is provided in Appendix E.

For assessment, we use both *local* and *global* evaluation metrics. At the local level, we look at the individual edge-endpoints and compare whether they match the corresponding endpoint of the true graph. At the global level, we look at the graph structure as a whole and measure the distance between the true and estimated graph, assessing how closely the estimated graph's structure matches the true graph's structure. As the local metrics, we utilize *precision*, *recall*, and *uncertainty rate*.

$$Precision = \frac{True \ Positive}{True \ Positive \ + \ False \ Positive}$$

$$Recall = \frac{True \ Positive}{True \ Positive \ + \ False \ Negative}$$

$$Uncertainty \ rate = \frac{Number \ of \ circle \ endpoints \ (\circ)}{Total \ number \ of \ edge-endpoints}$$

Precision reflects the prediction accuracy (i.e., out of all predicted cases, how many are correct), and recall reflects the retrieval rate (i.e., out of all true cases, how many are retrieved). Each edge-endpoint in a resulting graph can fall into one of four categories: no edge-endpoint (null), arrow head (>), arrow tail (-), and circle (\circ). Given that circle endpoints indicate that an algorithm is unsure of the direction of causal relations, the uncertainty rate is defined as the proportion of the circle endpoints present in the output. For example, consider the example PAG shown in Figure 13(b). The uncertainty rate for this PAG can be calculated as $\frac{3}{20}$, where the number of circle endpoints is 3, and the total number of edge-endpoints is 20 ($\binom{5}{2} \times 2 = 20$). For the other *non-circle* endpoints,

sparse dense without LC X_2 b = 5 X_1 X_2 without LC p = 10with LC

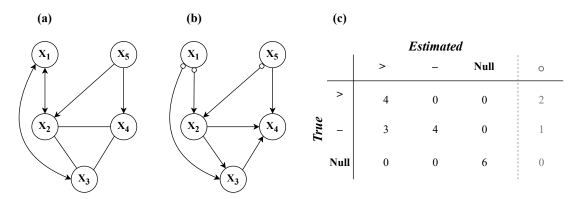
Figure 12. True ancestral graph for each condition.

Note. Each panel in the figure depicts the true ancestral graph for each simulated condition. Directed edges denote ancestral relationships $(A \rightarrow B \text{ means } A \text{ is an ancestor of } B)$, bidirected edges indicate the presence of latent confounders $(A \leftarrow B \text{ means there is a latent confounder between } A \text{ and } B)$, and undirected edges denote mutual ancestral relationships, thereby indicating cyclic relationships between the corresponding variables $(A - B \text{ means } A \text{ and } B \text{ are ancestors of each other, implying the presence of a cyclic relationship between them). For more information on how to derive the true ancestral graph from a DCG, see Appendix E. <math>p = \text{number of variables}$; LC = latent confounder.

we compute the precision and recall. As an example, suppose that Figure 13(a) is the true ancestral graph and Figure 13(b) is the estimated PAG output. We can then construct a confusion matrix of the estimated versus true edge-endpoints.⁶ Based on the confusion matrix shown in Figure 13(c), we can compute the precision and recall for each type of non-circle endpoints. For the arrow head (>), for instance, the precision and recall are computed as: $precision = \frac{4}{4+3+0}$ and $precision = \frac{4}{4+0+0}$. Note that the circle endpoints are not considered in the calculation of precision and recall and therefore they are greyed out in the confusion matrix below.

As the global metric, we use the *structural Hamming distance* (SHD) (de Jongh & Druzdzel, 2009). SHD quantifies the level of differences between two graphs by counting the number of edge insertions, deletions, and direction changes required to move from one graph (estimated graph $\hat{\mathcal{G}}$) to the other (true graph \mathcal{G}). It can be formulated as: SHD = A + D + C, where A, D, and C represent the number of added edges, deleted edges, and changes in edge-endpoints, respectively. A lower SHD value suggests that the estimated graph ($\hat{\mathcal{G}}$) is more closely aligned with the true graph (\mathcal{G}), indicating a better recovery of the true graph structure. For example, the SHD value for the PAG output in Figure 13(b) — provided that the true ancestral graph is Figure 13(a) — is 6, which is calculated by summing: 0(A) + 0(D) + 6(C).

Figure 13. Example performance evaluation.



Note. For the purpose of illustration, we consider the 5-variable sparse condition with a latent confounder shown in the top left panel of Figure 11. Panel (a) displays the true ancestral graph, while panel (b) presents an example of the estimated PAG output. Panel (c) shows the confusion matrix for the estimated versus true edge-endpoint, where the true endpoints are presented in rows and the estimated endpoints in columns. There are in total four possible edge-endpoints that can occur in an output: arrow head (>), arrow tail (-), null (no endpoint), and circle (o). The circle endpoints, however, are not counted toward the calculation of precision and recall, and thus are greyed out in the table.

⁶This example is based on the 5-variable sparse condition with a latent confounder shown in the top left panel of Figure 11.

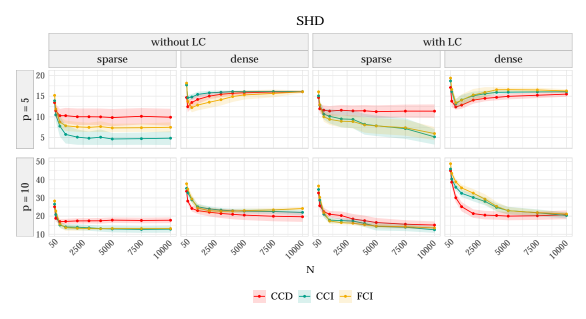
4.4 Simulation Results

In this section, we will report the results of the simulation study using the fixed values of **B** matrices. First, we will present the overall performance of the considered algorithms based on the structural Hamming distance (SHD), and then we will take a more detailed look into their performance using precision, recall, and uncertainty rate.

4.4.1 Performance on Global Metric

The SHD values for each algorithm per condition are shown in Figure 14, where the points and shaded area represent the average SHD values of 500 iterations and the corresponding interquartile range (IQR), respectively. As discussed in Section 4.3, a lower SHD value indicates a better performance, as it means that the estimated graph is closer to the true graph. Overall, the FCI and CCI algorithms perform better in sparse conditions, while the CCD algorithm outperforms the others in dense conditions. Additionally, the performance of all three algorithms is generally worse in dense conditions than in sparse conditions. Interestingly, in small models (p = 5) with high density (second and fourth columns from the top row of Figure 14), the SHD values momentarily decrease and then start increasing again as the sample size (N) becomes larger. This is partly against our expectation that the SHD values would decrease monotonically with increasing sample size. Furthermore,

Figure 14. Structural Hamming distance (SHD).



Note. The sample size (N) is shown on the x-axis, and the SHD values are shown on the y-axis. Each point represents the average SHD value across 500 simulations, while the shaded area represents the interquartile range (IQR). p = number of variables; LC = latent confounder.

we do not observe any significant contrasting patterns between conditions with and without latent confounders. This is contrary to our expectation that CCD would perform relatively better in conditions without latent confounders, while FCI and CCI would perform better in conditions involving latent confounders as they can handle latent confounding. In what follows, we will further examine their performance using local metrics while addressing some of the unexpected findings observed in the global metric analysis.

4.4.2 Performance on Local Metrics

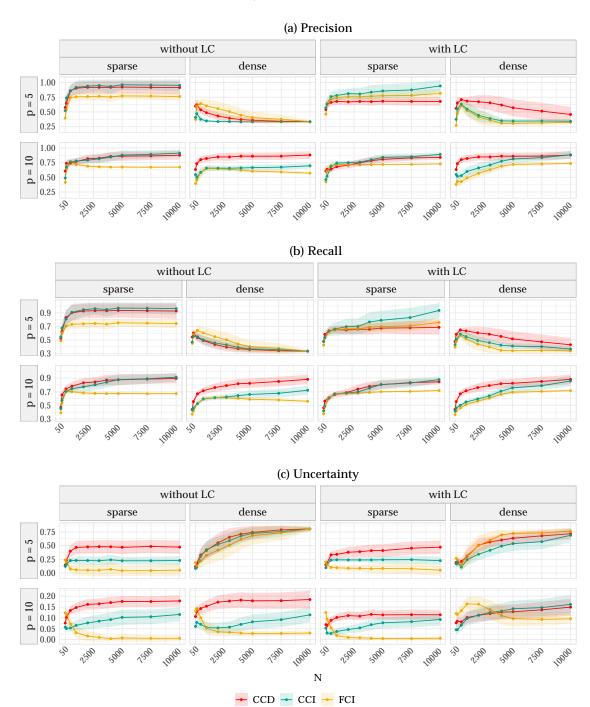
In this section, we will take a closer look at the performance of the algorithms based on precision, recall, and uncertainty rate. Additionally, we will analyze the unexpected results discussed previously in Section 4.4.1 by further breaking them down. These include (*i*) the observation that CCI and FCI outperform CCD in sparse conditions, even when latent confounders are not present and (*ii*) the SHD values increase instead of decreasing with increasing sample size in the *5-variable dense* conditions.

The results for precision, recall, and uncertainty rate for each algorithm are summarized in Figure 15, where each point represents the average value of the metrics over 500 iterations in the corresponding condition, and the shaded area represents the interquartile range (IQR) of the values obtained. As with the SHD results, we observe that the CCI algorithm generally exhibits high precision and recall in sparse conditions. On the other hand, the FCI algorithm records low precision and recall, even lower than CCD. This might seem contradictory at first glance, given that FCI has better SHD values than CCD in most conditions (except for a few dense cases). However, this can be explained by the overall low uncertainty rate of FCI as opposed to the high uncertainty rate of CCD, as shown in Figure 15(c). FCI tends to guess directions without necessarily outputting circles, while CCD tends to produce circles rather than guessing directions. This contrasting behavior consequently leads to FCI having better SHD values than CCD. To illustrate this point further, we examine the most frequently estimated PAGs from each algorithm in the 5-variable sparse condition without a latent confounder presented in Figure 16.7 In Figure 16(c), we can observe the typical orienting behavior of FCI, where it guesses the directions for every edge endpoint (i.e., not conservative), in contrast to CCD, which often produces circles (i.e., conservative), as shown in Figure 16(b). Based on the true graph shown in Figure 16(a), we can calculate the SHD: 9 for CCD (Figure 16(b)), 7 for FCI (Figure 16(c)), and 4 for CCI (Figure 16(d)). As such, FCI achieves a lower SHD value than CCD, despite having relatively lower precision and recall values due to some incorrectly predicted edge-endpoints.⁸ This highlights the distinct properties of the metrics used in our study. SHD is agnostic about whether an algorithm is conservative or less conservative, since

⁷These PAGs are constructed by assigning each edge-endpoint to the endpoint type that appears most frequently across all iterations

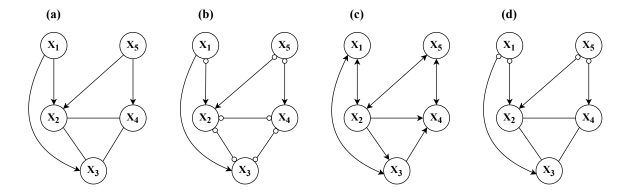
⁸Similar patterns are observed in the other sparse conditions, as shown in the first and third columns of Figure 15.

Figure 15. Precision, recall, and uncertainty rate.



Note. The sample size (N) is shown on the x-axis, and the corresponding metric values are shown on the y-axis. Each point on the graph represents the average of iteration-specific values of precision (top panel), recall (middle panel), and uncertainty rate (bottom panel) for each condition, with the shaded area indicating the interquartile range (IQR). p = number of variables; LC = latent confounder.

Figure 16. Frequently estimated PAGs in the 5-variable sparse condition without a latent confounder.

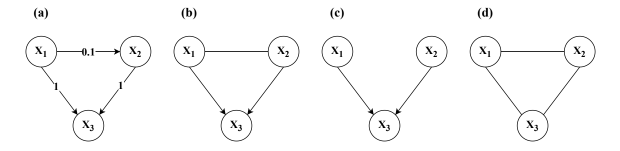


Note. Panel (a) shows the true ancestral graph of the 5-variable sparse condition without a latent confounder. Panels (b), (c), and (d) present the most frequently occurring PAGs in the 5-variable sparse condition without a latent confounder resulted from the CCD, FCI, and CCI algorithms, respectively. They were obtained by picking the most frequent type of edge-endpoints produced by each algorithm from 500 simulations with a sample size of 1000.

it measures all differences between the estimated and true graphs, including the circle marks. As a result, it penalizes an algorithm for being conservative as much as for making incorrect predictions. On the other hand, precision and recall only consider non-circle endpoints, while disregarding the circle marks. Therefore, they penalize an algorithm for making incorrect predictions but not for being conservative (i.e., producing circles).

In addition, the comparison between CCI and CCD based on precision and recall in Figure 15 reveals only a subtle difference in performance, even in the 5-variable sparse conditions where CCI outperformed CCD significantly in terms of SHD. This can be again attributed to CCD having a comparatively high uncertainty rate. From the PAG estimated by CCD in Figure 16(b), it can be observed that all of the directions predicted by CCD are correct, but the PAG contains a fairly large number of circle endpoints. These circle endpoints lead to a poorer SHD value for CCD, even though it exhibits comparably high precision and recall for correctly predicting directions. Moreover, further examination of the resulting PAGs shows that CCI performs better than CCD in detecting mutual ancestral relationships in cyclic structures. For instance, in Figure 16(b), CCD assigned all circle marks to the cycle involving X_2 , X_3 , and X_4 , while CCI correctly identified the mutual ancestral relationships in the cycle with undirected edges (—), as shown in Figure 16(d). This superiority of CCI in recovering mutual ancestry in cycles was consistently observed in other conditions as well, suggesting that this property is a significant factor contributing to CCI's overall better performance compared to other algorithms, even in the absence of latent confounding.

Figure 17. An example dense graph with a weak edge.



Note. Panel (a) shows the true graph. Panel (b) displays the desired output where the collider structure is correctly oriented while the weak edge between X_1 and X_2 is also identified. Panel (c) presents the output graph when an algorithm fails to detect the weak edge between X_1 and X_2 but correctly orients the collider structure. Panel (d) displays the output graph when an algorithm detects the weak edge between X_1 and X_2 . This results in a denser graph with less information on independencies, leading to the loss of orientation of the collider structure.

With respect to the unexpected increase in SHD values in the *5-variable dense* conditions, we also observe an unusual decrease in precision and recall as the sample size increases (second and fourth panels from the top rows of Figure 15(a) and Figure 15(b)), along with an increase in the uncertainty rate (second and fourth panels from the top row of Figure 15(c)), indicating that more circle endpoints appear in the resulting PAGs as the sample size grows. Learning a dense causal structure is generally more challenging, because there is less information available about the conditional independence relations when a large number of vertices are connected by edges (i.e., almost everything is dependent on everything else). This problem is further compounded in the presence of cycles, as all variables that are part of cycles are completely dependent on each other, making it even more challenging to obtain information about independence relations. Therefore, algorithms tend to fail to orient any edges and mostly output only circle endpoints in dense cyclic graphs. The decline in precision and recall, along with the rapid increase in uncertainty rate observed in Figure 15 for the *5-variable dense* conditions, are primarily due to this challenge in high-density situations — as the sample size becomes larger, relatively weak edges start getting picked up by the algorithms, leading to denser structures that make causal discovery exceedingly difficult.

However, interestingly, we also note a slight increase in both precision and recall for the 5-variable dense conditions when the sample size is relatively small, accompanied by a small drop in the SHD (second and fourth panels from the top row of Figure 14). This might seem counterintuitive, as we typically expect the algorithms to learn causal structures more accurately with larger sample sizes. But, in fact, it is possible to lose the correct edge orientation when the sample size is large, particularly in the case of a dense graph with weak edges (Eigenmann et al., 2017). Figure 17 illustrates a simple example case where the inclusion of a weak edge leads to the loss of

edge orientation. With a large sample size, algorithms are likely to pick the weak edge between X_1 and X_2 , resulting in an uninformative undirected graph, as shown in Figure 17(d). However, with a small sample size, algorithms are more likely to miss the weak edge and identify the collider X_3 , thereby obtaining the correct edge orientation, as shown in Figure 17(c). In our 5-variable dense scenarios, we run into a similar situation. There is a relatively weak edge between X_2 and X_5 , which is not detected by algorithms when the sample size is small, resulting in some correctly oriented edges. However, as the sample size becomes larger, the weak edge gets detected, and the algorithms fail to orient any edges, thus yielding completely undirected graphs (see Appendix F for a more detailed explanation of the results in our 5-variable dense cases). This explains the small dips at the beginning of the SHD line graphs (Figure 14) and brief spikes in the precision and recall graphs (Figure 15) for the 5-variable dense conditions with relatively small sample sizes.

To summarize, both CCD and CCI algorithms show good performance, with CCI slightly outperforming in sparse conditions and CCD in dense conditions. However, CCD tends to be more conservative in terms of edge orientation and produces more circle endpoints than CCI, which often results in higher SHD values. In contrast, FCI demonstrates poor performance across most conditions, which is not immediately obvious when considering only the global metric, SHD. Further analysis of local metrics reveals that FCI often makes quick directional inferences without producing circle marks, resulting in comparably good SHD values but low precision and recall values. To investigate the robustness of our findings against the specific parameter values chosen in this simulation, including the coefficients of $\bf B$ matrices and the value of α , we conducted additional simulations. In the next section, we will present the results of these additional analyses.

4.5 Sensitivity Analyses

In the previous section, we reported the results of a simulation study that used fixed parameter values for the weight matrix **B**. Those findings showed unusual trends in the *5-variable dense* conditions, suggesting that the performance of different causal discovery algorithms may be sensitive to the specification of **B**. To further explore this issue, we repeated the simulation study, but this time we randomly sampled the parameter values of **B** at each iteration.

The overall results are similar to the original simulation study, with some minor differences (see Appendix G for the complete results). First, CCD's performance improves slightly across conditions, showing comparable SHD values to those of CCI, even in sparse conditions. However, CCD's conservative nature and tendency to generate many circle endpoints persist, as observed in the original simulation. This leads us to the same conclusion as in the original simulation that CCI would be preferred when seeking the most information on causal directions with low uncertainty. Second, FCI exhibits greater stability and generates a reasonable number of circle endpoints, recording comparable uncertainty rates to those of CCD. However, FCI's relatively poor performance (low precision and recall) across conditions remains unchanged, consistent with what we observed in the

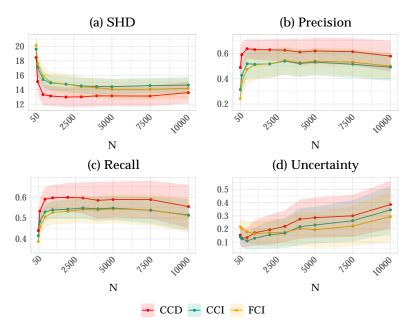


Figure 18. Performance in the 5-variable dense condition without a latent confounder.

Note. This figure illustrates the performance of each algorithm in the 5-variable dense condition without a latent confounder, where the coefficients of **B** were randomly sampled. The sample size (N) is shown on the x-axis, and the corresponding metric values are shown on the y-axis. Each point represents the mean value of each metric across 500 simulations, and the shaded area represents the interquartile range (IQR). Note that the performance under the *5-variable dense* condition *with* a latent confounder shows more or less the same patterns (see Appendix G for details).

original simulation. In the 5-variable dense conditions, however, we notice a significant change in the patterns, as we would expect based on our explanation of those results in Section 4.4.2. In contrast to the original simulation, we do not find any unusual kinks or dips in the performance curves with small sample sizes in the 5-variable dense conditions (see Figure 18). Instead, we find that the algorithms' performance steadily improves with increasing sample size, as demonstrated by the gradual decrease in SHD and increase in precision and recall until they level off, which aligns with our initial expectation. This result confirms that the peculiar patterns observed in the original simulation in the 5-variable dense conditions were indeed produced by the particular choices of parameter values.

In addition, in our main simulation study, we used a fixed alpha level of 0.01 ($\alpha=0.01$). However, it is commonly suggested to adjust α according to the sample size in order to ensure consistent results of conditional independence tests (Mooij et al., 2020; Colombo et al., 2012). If alpha is not adjusted appropriately based on the sample size, it can lead to either accepting the null hypothesis of independence ($H_0: X_i \perp \!\!\! \perp \!\!\! X_j \mid \mathbf{Y}$, where a set $\mathbf{Y} \subseteq \mathbf{X} \setminus \{X_i, X_j\}$) with small sample sizes, falsely declaring everything as independent, or rejecting it with large sample sizes, failing to find

any independence relations (Magliacane et al., 2017). To test the robustness of our results obtained using a fixed value of α , we conducted an additional simulation study as a sensitivity check. In this study, we varied the α level based on the sample size; decreasing α as the sample size (N) increased such that $\alpha_N \to 0$ as $N \to \infty$ at a suitable rate. The full results can be found in Appendix H. Overall, the results show no significant differences in the observed patterns across all conditions, indicating that the effect of adjusting α is negligible, and the algorithms are rather insensitive to the choice of α . Thus, we conclude that our findings are robust to the fixed α level chosen in the original simulation.

4.6 Intermediate Summary

Through the simulation study, we evaluated the performance of three different constraint-based cyclic causal discovery algorithms under various conditions. Based on the global metric, SHD, it appeared that CCI and FCI performed relatively well in sparse conditions, and CCD outperformed the others in dense conditions. However, upon examination of the local metrics — precision, recall, and uncertainty rate — it became apparent that FCI, in fact, performed rather poorly, and CCD outperformed FCI in most conditions, achieving higher precision and recall. This discrepancy was due to the difference in uncertainty rates between the two algorithms. While CCD tended to produce circle endpoints rather than erroneously orienting edges, which resulted in higher SHD values, FCI often guessed the directions of most edges without producing circles, leading to relatively low SHD values by occasionally hitting the directions correctly. This behavior was unexpected based on our discussion of FCI in Section 3, where we noted that FCI typically generates more circle endpoints as it accounts for unobserved confounding. One possible explanation for this unexpected behavior is the mismatch between the linear causal relations imposed in our simulation study and the non-linear causal relations suggested by Mooij and Claassen (2020). Overall, CCI exhibited good performance based on both global and local metrics. CCD performed relatively well in dense conditions, but it produced many circle endpoints, resulting in high uncertainty rates. FCI performed poorly, often incorrectly inferring directions for some of the edges. As part of our sensitivity analyses, we conducted two additional simulation studies. In one simulation, where we randomly sampled the parameters of the weight matrix **B**, we learned that the unusual patterns observed in the 5-variable dense conditions in the original simulation were due to the specific parameter values chosen. In another simulation, where we varied the α level according to the sample size, we found that adjusting α had minimal impact on the results, confirming the robustness of our findings that were based on the fixed α level. With these results in mind, in the following, we will present an empirical example where we apply the algorithms to real psychological data to assess their practical applicability.

⁹As our variables follow a multivariate normal distribution, we utilize partial correlations to identify the conditional independencies. Therefore, we test whether a partial correlation is equal to zero or not, where $H_0: \rho_{X_i,X_j|\mathbf{Y}} = 0$.

¹⁰When setting the α level, we adopted a heuristic value of $\alpha = \frac{1}{\sqrt{N}}$, given that the partial correlation can decay as $N^{-1/2+\epsilon}$ for any $0 < \epsilon < 1/2$ in multivariate Gaussian cases (Kalisch & Buehlmann, 2005).

5 Empirical Example

In this section, we will demonstrate the application and interpretation of the three causal discovery algorithms with an empirical example. We will begin by introducing the dataset and then present the output of each algorithm, along with the corresponding statistical network model. Our goal is to assess whether the output PAGs carry meaningful information about the causal structure and to compare the insights gained from these PAGs with those obtained from the statistical network model.

5.1 Data and Model Fitting

The dataset used in our empirical example is from McNally et al. (2017), where the authors focused on examining the causal relationships between symptoms of obsessive-compulsive disorder (OCD) and depression. Notably, the authors expressed a particular interest in investigating cyclic causal relationships. However, the causal discovery methods they were familiar with were only applicable to acyclic causal structures, leading them to use a statistical network model. Although they acknowledged that the network captures statistical rather than causal relationships, they argued that statistical network analysis addresses a key limitation of the acyclic causal discovery algorithm, which is excluding the possibility of feedback loops (cycles). Their explicit interest in cyclic causal discovery makes this dataset an ideal candidate for testing the cyclic causal discovery algorithms discussed in the current paper.

The dataset consists of 408 observations and 26 variables, comprising 16 depression symptoms and 10 OCD symptoms, all with no missing values. The severity of each symptom was measured using a four-point Likert scale, with 0 indicating no symptoms and 3 indicating extreme symptoms. The participants' age ranged from 18 to 69 years, with a mean of 31.1 and a standard deviation of 12.2. In this paper, we focus our analysis on a subset of the variables — depression symptoms — in order to keep the model size manageable and facilitate the interpretation of the estimated PAGs. For a summary of the depression symptom variables, please refer to Table 2.

To compare the results of the cyclic causal discovery algorithms with those of network analysis, we estimated a Gaussian graphical model (GGM). In a GGM, the edges signify partial correlations between pairs of nodes, controlling for the rest of nodes in the network (Epskamp et al., 2018b). To obtain a sparse network, we used the *graphical lasso* (glasso) method to regularize partial correlations, such that small partial correlations are driven to zero and therefore do not appear in the network (Friedman et al., 2008). For the causal models, we ran all three algorithms (CCD, FCI, and CCI) while setting the alpha level to 0.01 ($\alpha = 0.01$) as a rough way to correct for spurious edges resulting from a relatively large set of variables (Zhang et al., 2012). As in the simulation study, we used partial correlations to test for conditional independencies. In the following section, we will present the estimated models and interpret the findings from each model while comparing them to one another.

¹¹The data set is publicly available on the Psychological Medicine Journal webpage.

Table 2. Summary of depression symptoms.

Symptom (abbreviation)	Mean (SD)
1. Sleep-onset insomnia (<i>ons</i>)	1.20 (1.07)
2. Middle insomnia (<i>mdd</i>)	1.44 (1.07)
3. Early morning awakening (<i>lat</i>)	0.81 (1.07)
4. Hypersomnia (<i>hyp</i>)	1.01 (0.99)
5. Sadness (sad)	1.55 (0.94)
6. Decreased appetite (<i>dcp</i>)	0.49 (0.72)
7. Increased appetite (inc)	0.44 (0.87)
8. Weight loss (wghtl)	0.50 (0.94)
9. Weight gain (wghtg)	0.67 (1.04)
10. Concentration impairment (cnc)	1.48 (0.87)
11. Guilt and self-blame (glt)	1.56 (1.17)
12. Suicidal thoughts (scd)	0.63 (0.82)
13. Anhedonia (anh)	1.27 (1.05)
14. Fatigue (ftg)	1.33 (0.95)
15. Psychomotor retardation (<i>rtr</i>)	0.66 (0.81)
16. Agitation (agt)	1.10 (0.93)

Note. SD = standard deviation.

5.2 Empirical Analysis Results

The causal models estimated by each algorithm (i.e., PAGs) and the GGM are depicted in Figure 19. Upon examining the GGM presented in Figure 19(a), we can identify three symptom clusters that are highly interconnected within themselves but less connected with the rest. On the right-hand side of the network, we see symptoms related to physical weight and appetite: weight gain (wghtg), weight loss (wghtl), increased appetite (inc), and decreased appetite (dcp), with particularly strong partial correlations between inc - wghtg and dcp - wghtl. Towards the bottom of the network, we see three symptoms relating to insomnia: early morning awakening (late), sleep onset insomnia (ons), and middle insomnia (mdd). The remaining items towards the left side of the network form a single cluster. Here, we observe strong connections among several symptoms, including psychomotor retardation (rtr) – concentration impairment (cnc) – fatigue (ftg) – anhedonia (anh) – sadness (sad) – suicidal thoughts (scd). Especially, fatigue, anhedonia, and sadness emerge as the most central symptoms based on the number and strength of connections they have with the other nodes in the network. As McNally et al. (2017) note, this is in accordance with the general clinical observations where sadness and anhedonia, as the two gateway symptoms, are necessary for a diagnosis

of depression.

Looking at the PAG from the CCD algorithm shown in Figure 19(b), we can gain information on causal directions. Overall, the estimated structure is not too different from that of the GGM, but it is more sparse. The estimated PAG from CCD reveals that fatigue and anhedonia still remain central nodes. However, in the causal graph, we can now attribute their centrality to the fact that fatigue is the (indirect) effect of many variables, and anhedonia is the (indirect) cause of many variables — the directional information that could not be obtained from the statistical network model above. The three-cluster structure observed in the GGM is also present in this PAG, with only a few edges linking the separate clusters. For example, on the right side of the graph, we can see that increased appetite (inc) is an ancestor of guilt (glt), and weight loss (wghtl) is an ancestor of early morning awakening (lat). Anhedonia (anh) also plays an important role in linking these separate symptom clusters, acting as an ancestor of both early morning waking (lat) and decreased appetite (dcp). On the left side of the graph, we observe an intricate network where most of the variables are involved in a web of cyclic ancestral relationships, including chains such as fatigue $(ftg) \rightarrow$ concentration $(cnc) \rightarrow \text{psychomotor retardation } (rtr) \rightarrow \text{fatigue } (ftg), \text{ or longer chains like concentration } (cnc)$ \rightarrow agitation $(agt) \rightarrow$ guilt $(glt) \rightarrow$ sadness $(sad) \rightarrow$ suicidal thoughts $(scd) \rightarrow$ guilt $(glt) \rightarrow$ concentration (cnc). While it is tempting to interpret these ancestral patterns as those of direct cyclic causal relationships, readers should exercise caution. As previously demonstrated in Figure 4 from Section 3.2.1, there are a couple of points to keep in mind when mapping from ancestral to parental causal relationships. First, the cyclic causal relationships would be represented by undirected edges (—) in a PAG, not by patterns of directed ancestral relationships. Second, the dotted-underlining of the variables such as fatigue (ftg) and guilt (glt) likely indicates that not all the ancestral relationships shown in this PAG can be mapped onto direct causal relationships in the corresponding DCG. Hence, the structures that appear as cycles in this PAG may not necessarily imply the same direct causal cycles in the underlying DCG. Despite the rather complicated nature of interpreting the PAG, we believe that it provides more insightful information on the causal dynamics of depression symptoms when compared to the GGM, given that the PAG provides directional information, indicating the causal flow of variables, which cannot be obtained from the GGM.

However, it is important to note that CCD assumes causal sufficiency (i.e., the absence of unobserved confounders), which could lead to an overly optimistic PAG that does not accurately reflect the causal relationships. This limitation is addressed by the CCI and FCI algorithms, which allow for the presence of latent confounders. Figure 19(c) and Figure 19(d) show the estimated PAGs from CCI and FCI, respectively. The overall structure is reminiscent of the PAG obtained from CCD, with three distinct clusters. However, the PAGs generated by CCI and FCI contain fewer edges, resulting in more independent clusters. Also, these PAGs feature bidirected edges (\leftrightarrow), indicating the presence of latent confounders. The CCI algorithm, in particular, produced a significant number of bidirected edges, suggesting the presence of latent confounders between almost every variable (see Figure 19(c)). On the other hand, the PAG generated by FCI shown in Figure 19(d) has only a few

(b) (a) hyp ftg agt agt glt anh anh sad dcp dcp wghtl (c) (d) agt anh

Figure 19. A statistical network model and PAGs estimated from empirical data.

Note. Panel (a) shows the statistical network model (i.e., Gaussian graphical model) estimated from the empirical data on depression symptoms. Panels (b), (c), and (d) show the PAG estimated by the CCD, CCI, and FCI algorithm, respectively. ons = sleep onset insomnia; mdd = middle insomnia; lat = late (early morning awakening); hyp = hypersomnia; sad = sad; dcp = decreased appetite; inc = increased appetite; wghtl = weight loss; wghtg = weight gain; cnc = concentration impairment; glt = guilt; scd = suicidal thoughts; anh = anhedonia; ftg = fatigue; rtr = psychomotor retardation; agt = agitation.

bidirected edges but more circle endpoints. Both CCI and FCI identified some common bidirected edges, including $cnc \leftrightarrow ftg$, $agt \leftrightarrow cnc$, and $agt \leftrightarrow glt$, indicating that there are likely to be latent confounders present between these variables. This is consistent with previous research that has shown, for example, chronic physical illness to be a potential contributing factor to symptoms such as concentration impairment (cnc) and fatigue (ftg) (Menzies et al., 2021; Goërtz et al., 2021; de Ridder et al., 2008). In addition, both CCD and FCI predicted some common directional features, such as $rtr*\to ftg, hyp*\to ftg, scd*\to glt$, and $anh*\to dcp$. For example, $anh*\to dcp$ means that anhedonia may or may not cause decreased appetite, but decreased appetite does not cause anhedonia, which also partly aligns with a previous study, where appetite loss is identified as one of the features of anhedonia (Coccurello, 2019).

Overall, this example highlights that the PAGs produced by cyclic causal discovery algorithms provide unique insights into possible causal relationships that cannot be gained through network analysis alone. The directional information conveyed by the PAGs illustrates the causal flow of variables, and the sparsity of the PAGs indicates that not all variables are causally related to one another, emphasizing the importance of identifying the relevant variables when investigating the causal structure of depression symptoms. In addition, our findings show that the algorithms generally agree on many features, including the presence or absence of causal relations and some of the causal directions, which are in line with prior research, further enhancing their plausibility. However, the choice of which output PAG to pay most attention to depends on our degree of confidence in the assumptions underlying each algorithm. While the most informative PAG is generated by CCD, it makes strong assumptions about the absence of unobserved confounding and the linearity of causal relations. On the other hand, CCI suggests that most or all relationships between symptoms are likely influenced by latent confounding. FCI also flags the possibility of latent confounding but recovers different directed ancestral relations than CCI, as it makes use of different orientation rules. In general, FCI may be preferred over CCI when the linearity assumption does not hold, but CCI may be more appropriate in cases where the linearity assumption holds, as suggested by our simulation study. Finally, to ensure the reliability of our findings, we conducted an additional stability analysis by running the algorithms on multiple random subsets of the data and retaining only the causal relations that were consistently discovered. This analysis confirmed that the identified causal relations from our original analysis are reliable such that they are not undermined by small variations in the data. Further details regarding this analysis can be found in Appendix I.

6 Discussion

In this paper, we studied constraint-based cyclic causal discovery algorithms in typical psychological research settings, with a focus on identifying an effective algorithm for studying the underlying cyclic causal structure. We provided a comprehensive overview and didactic treatment of cyclic causal discovery by outlining the properties of three specific algorithms: CCD, FCI, and CCI. We assessed the performance of these algorithms under varying conditions through a simulation study, and we also demonstrated their practical applicability in psychological research by applying them to empirical data. Our results suggested that the CCI algorithm generally performed well, particularly in sparse conditions, and the CCD algorithm tended to outperform the others in dense conditions. The FCI algorithm performed poorly across all conditions, mostly guessing the directions of edges, though notably, our simulation was limited to studying systems with linear causal relations. Our empirical example showed that causal discovery methods provided more detailed and richer insights into the underlying causal dynamics of depression than the statistical network model, which in fact was found to contain numerous spurious edges, rendering it unsuitable for serving as a causal skeleton (Ryan et al., 2022).

With this paper, we aimed to provide empirical researchers with guidance on selecting an appropriate cyclic causal discovery algorithm for studying causal relationships. Our findings indicate that no single algorithm is suitable for all cases, and the choice of algorithm should be based on the characteristics of the causal system of interest. If the causal system is believed to be relatively sparse, then the CCI algorithm may be preferred, as it performs well under such conditions. Conversely, if the system is considered to be comparatively dense, researchers may opt for the CCD algorithm. Also, researchers need to consider their priorities when selecting an algorithm. If avoiding incorrect edge orientations is a priority, then the CCD algorithm, which is more conservative in edge orientation, would be the preferred choice. However, if acquiring more insights into causal relationships is a priority, even if it means accepting some incorrect edge orientations, then CCI would be the more suitable option. Although the impact of violating assumptions was not entirely clear from our simulation study, researchers should assess which assumptions are relevant and critical for the particular nature of the causal system they seek to study, as each algorithm makes different assumptions. Despite the complexity involved in using cyclic causal discovery methods, we emphasize that these techniques are more informative than statistical network analysis when exploring the underlying causal structure.

However, we acknowledge that causal discovery in cyclic settings entails theoretical and practical challenges, and that much future work remains to be done to gain a better understanding of the behavior of causal discovery algorithms. In our simulation study, we restricted ourselves to a set of fixed causal structures with fixed weights instead of randomly sampling graph structures, which is a more typical approach for simulating models (Mooij et al., 2020; Strobl, 2019; Colombo et al., 2012). We chose fixed structures to prioritize the explainability and interpretability of the results,

as using randomly sampled structures would not have allowed us to assess individual configurations and edges as thoroughly as we did with the fixed structures. However, our use of fixed structures does limit the generalizability of our findings, as we only studied a small number of structures. While our sensitivity analysis showed some level of robustness to variations in the weights of the causal structure, our limited exploration of structure remains a limitation of our study. One possible approach to improve the generalizability of our findings is to expand the simulation settings by incorporating a random graph structure at each iteration. However, this approach would be computationally intensive, as it requires verifying the cyclicity and equilibrium condition for each structure and iterating until satisfied. Future studies can explore the feasibility of this approach in more detail.

Furthermore, certain operational details of the algorithms were not considered in the assessment of their performance in our simulation study. For instance, the CCD algorithm generates additional underlinings that convey more information about the allowable patterns of direct causal relationships in the equivalence class of DCGs. However, this additional information is not provided by the CCI and FCI algorithms, and to make the algorithm outputs directly comparable, we did not account for this in our simulation study. While we examined the uncertainty rate to evaluate the informativeness of the resulting PAGs for each algorithm — more circle endpoints indicate greater uncertainty in the inferred causal relations, thereby implying a less informative PAG — it is possible that the extra underlinings from the CCD algorithm may have made its output PAG significantly more informative. To further investigate the usefulness of this extra information, future studies could explore alternative performance metrics, such as directly assessing the size of the equivalence class implied by each algorithm output. This would determine the extent to which the PAGs are informative in inferring the causal structure in general, with a smaller equivalence class indicating a more informative PAG. Though conceptually promising, it could be practically challenging as the search space can expand exponentially with a larger model, and it may not be straightforward to derive the size of the equivalence class when latent confounders are involved. Nonetheless, it can be a crucial validity check on our simulation results, where we found that CCD consistently generated less informative PAGs by producing more circle endpoints compared to the other algorithms.

Our simulation was also limited in scope as we only considered Gaussian linear cases, which had the advantage of satisfying the global Markov condition and allowing the use of partial correlations to test conditional independencies. Although Gaussian linear processes are commonly assumed in psychological research (Pek et al., 2018; Beller & Baier, 2013), this assumption may be oversimplified and too strict in practice. In fact, the data used in our empirical example, which is typical of those used in psychological network research measured on a Likert scale, deviated from the Gaussian distribution. Despite this, we proceeded with testing conditional independencies using partial correlations. Although the exact impact of this misspecification on our results remains unclear, it may have led to misleading findings (Baba et al., 2004), which requires further investiga-

¹²See Appendix J for the distributions of all depression symptom variables.

tion. As real-world applications often involve non-linear and non-Gaussian processes, future studies should explore more general scenarios beyond linear Gaussian cases to enhance the practical applicability and to gain a more comprehensive understanding of cyclic causal discovery algorithms. Various flexible conditional independence (CI) tests have already been developed to accommodate such cases (Li & Fan, 2020; Canonne et al., 2018), including non-parametric discretization-based CI tests (Huang, 2010) and kernel-based CI tests (Zhang et al., 2012), which can be applied without assuming a functional form between the variables or the data distribution. However, these testing methods are often more complex and require larger sample sizes, making their practical application challenging. Hence, it would be an interesting extension for future research to investigate the feasibility of implementing these methods and their effectiveness in real-world scenarios.

The network theory of psychopathology, which posits that mental disorders arise from direct causal interactions between symptoms and sustain themselves through feedback loops, has greatly contributed to the understanding of psychopathology (Borsboom, 2017). However, despite evidence demonstrating that network models are inadequate as causal discovery tools, many empirical researchers in psychology have attempted to generate causal hypotheses from estimated network models (Ryan et al., 2022). Alternatively, some researchers have tried to estimate a DAG even though the theory strongly suggests the presence of cycles/feedback loops (McNally et al., 2017; Briganti et al., 2022). In this paper, we propose a promising alternative, cyclic causal discovery methods, which are explicitly designed to recover causal structure with cycles. Although we acknowledge that these methods come with caveats, such as complicated algorithm steps, restrictive assumptions, and somewhat difficult output interpretations, we believe that these methods can offer a valid approach to understanding the causal mechanisms underlying dynamics of mental disorders. To advance the network approach to studying the dynamics of psychopathology, we argue that it is essential for researchers to gain familiarity with these methods and explore their potential applications in psychological research. We hope that the current paper will remove some barriers to entry and encourage a more widespread adoption of cyclic causal discovery methods in psychological science.

Materials

An entire research archive with R-code, results, and figures presented in this paper is available on GitHub (Park, 2023). The supplementary material for this paper can be found online at: https://kyurip.quarto.pub/discovering-cyclic-causal-models/.¹³

¹³Alternatively, you can access all the materials in the Open Science Framework (OSF) repository.

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Appendix A CCD Details

Algorithm 1 Cyclic Causal Discovery (CCD)

Input: A conditional independent oracle for a distribution \mathcal{P} , satisfying global directed Markov property and faithfulness conditions with respect to a directed graph \mathcal{G} with vertex set \mathcal{V} .

Output: A PAG Ψ for the Markov equivalence class of DCGs, Equiv(G).

- 1: Step 1. Form a complete graph (Ψ) with the edge \sim between every pair of vertices in \mathcal{V} .
- 2: n = 0
- 3: repeat
- 4: repeat
- Select an ordered pair of variables X and Y that are adjacent in Ψ such that the number of vertices in $\mathbf{Adjacent}(\Psi, X) \setminus \{Y\} \ge n$, and select a subset S of $\mathbf{Adjacent}(\Psi, X) \setminus \{Y\}$ with n vertices.
 - If $X \perp\!\!\!\perp Y \mid S$, then delete the edge $X \circ \neg \circ Y$ and record S in **Sepset** $\langle X, Y \rangle$ and **Sepset** $\langle X, Y \rangle$.
- 6: **until** all pairs of adjacent variables X and Y such that the number of vertices in $Adjacent(\Psi, X) \setminus \{Y\} \ge n$ and all sets S such that the number of vertices in S = n have been tested.

$$n = n + 1$$
;

- 7: **until** for all ordered pairs of adjacent vertices X and Y, **Adjacent**(Ψ , X)\{Y} < n.
- 8: *Step 2.* For each triple of vertices A, B, C such that each of the pair of A, B and the pair B, C are adjacent in Ψ but the pair A, C are not adjacent in Ψ , then:
- 9: (i) orient A*-*B*-*C as $A \rightarrow B \leftarrow C$ iff $B \notin \mathbf{Sepset}(A, B)$.
- 10: (ii) orient A*-*B*-*C as A*-*B*-*C iff $B \in \mathbf{Sepset}(A, B)$.
- 11: *Step 3.* For each triple of vertices A, X, Y in Ψ such that (i) A is not adjacent to X or Y, (ii) X and Y are adjacent, (iii) $X \notin \mathbf{Sepset}(A, Y)$, then orient X * *Y as $X \leftarrow Y$ if $A \perp \!\!\! \perp X \mid \mathbf{Sepset}(A, Y)$.
- 12: **Step 4.** For each vertex V in Ψ form the following set: $X \in \mathbf{Local}(\Psi, V)$ iff X is adjacent to V in Ψ , or there is a vertex Y such that $X \longrightarrow Y \longleftarrow V$ in Ψ .
- 13: m = 0
- 14: repeat
- 15: repeat
- Select an ordered triple $\langle A, B, C \rangle$ such that $A \rightarrow B \leftarrow C$, A and C are not adjacent, and $Local(\Psi, A) \setminus \{B, C\}$ has $\geq m$ vertices.
 - Select a set $T \subseteq \mathbf{Local}(\Psi, A) \setminus \{B, C\}$ with m vertices. If $A \perp \!\!\!\perp C \mid T \cup \{B\}$, then orient $A \longrightarrow B \longleftarrow C$ as $A \longrightarrow B \longleftarrow C$ and record $T \cup \{B\}$ in $\mathbf{Supset} \langle A, B, V \rangle$.

- 17: **until** for all triples such that $A \rightarrow B \leftarrow C$ (not $A \rightarrow B \leftarrow C$), A and C are not adjacent, **Local**(Ψ , A)\{B} has $\geq m$ vertices, every subset T with m vertices has been considered.
- 18: m = m + 1:
- 19: **until** all ordered triples $\langle A, B, C \rangle$ such that $A \rightarrow B \leftarrow C$, A and C are not adjacent, are such that **Local**(Ψ , A)\{B} have < m vertices.
- 20: **Step 5.** If there is a quadruple $\langle A, B, C, D \rangle$ of distinct vertices in Ψ such that:
- 21: (i) $A \rightarrow B \leftarrow C$,
- 22: (ii) $A \rightarrow D \leftarrow C$ or $A \rightarrow D \leftarrow C$,
- 23: (iii) B and D are adjacent,
- 24: then orient B*-*D ad $B\to D$ in Ψ if $D\notin \mathbf{Subset}(A,B,C)$. Else orient B*-*D as B*-D in Ψ .
- 25: Step 6. For each quadruple A, B, C, D in Ψ of distinct vertices such that:
- 26: (i) D is not adjacent to both A and C,
- 27: (ii) $A \rightarrow B \leftarrow C$,
- 28: if $A \not\perp D$ | **Supset** $\langle A, B, C \rangle \cup \{D\}$, then orient B * → D as B → D in Ψ .

Appendix B FCI Details

Algorithm 2 Fast Causal Inference (FCI)

Input: A conditional independent oracle for a distribution \mathcal{P} , satisfying global directed Markov property and faithfulness conditions with respect to a directed graph \mathcal{G} with vertex set \mathcal{V} .

Output: A PAG $\hat{\mathcal{G}}'$ for the Markov equivalence class of DMGs, $Equiv(\mathcal{G})$.

- 1: Step 1. Form the complete undirected graph Q on the vertex set V.
- 2: n = 0
- 3: repeat
- 4: repeat
- 5: Select an ordered pair of variables X and Y that are adjacent in Q such that the number of vertices in $\mathbf{Adjacent}(Q, X) \setminus \{Y\} \ge n$, and select a subset S of $\mathbf{Adjacent}(Q, X) \setminus \{Y\}$ with n vertices.

If $X \perp\!\!\!\perp Y \mid S$, then delete the edge $X \circ \neg \circ Y$ and record S in **Sepset** $\langle X, Y \rangle$ and **Sepset** $\langle X, Y \rangle$.

6: **until** all pairs of adjacent variables X and Y such that the number of vertices in

Adjacent(Q, X)\{Y} $\geq n$ and all sets S such that the number of vertices in S = n have been tested.

n = n + 1;

- 7: **until** for all ordered pairs of adjacent vertices X and Y, **Adjacent** $(Q, X) \setminus \{Y\} < n$.
- 8: **Step 2.** Let Q' be the undirected graph resulting from step 1.
- 9: For each triplet $\langle A, B, C \rangle$ such that each of the pair of A, B and the pair B, C are adjacent in Q' but the pair A, C are not adjacent in Q', then orient A*-*B*-*C as $A*-B\leftarrow *C$ iff $B \notin \mathbf{Sepset} \langle A, B \rangle$.
- 10: *Step 3.* For each pair of variables A and B adjacent in Q', if A and B are d-separated given any subset S of **Possible-d-sepset** $(A, B) \setminus \{A, B\}$ or any subset S of **Possible-d-sepset** $(B, A) \setminus \{A, B\}$ in Q', then remove the edge between A and B, and record S in **Sepset**(A, B) and **Sepset**(B, A).
- 11: Step 4. Execute the following orientation rules iteratively until none applies:
 - (i) If $A * \rightarrow B \circ \rightarrow * C$, and A and C are not adjacent, then orient the triple as $A * \rightarrow B \rightarrow C$.
 - (ii) If $A \rightarrow B * \rightarrow C$ or $A * \rightarrow B \rightarrow C$, and $A * \rightarrow C$, then orient $A * \rightarrow C$ as $A * \rightarrow C$.
 - (iii) If $A*\to B\leftarrow *C$, $A*\multimap D\circ -*C$, A and C are not adjacent, and $D*\multimap B$, then orient $D*\multimap B$ ad $D*\to B$.
 - (iv) If $u = \langle D, ..., A, B, C \rangle$ is a discriminating path between D and C for B, and $B \circ -*C$; then if $B \in \mathbf{Sepset}\langle D, C \rangle$, orient $B \circ -*C$ as $B \to C$; otherwise orient the triple $\langle A, B, C \rangle$ as $A \leftrightarrow B \leftrightarrow C$.
 - (v) For every (remaining) $A \circ \neg B$, if there is an uncovered circle path $p = \langle A, C, ..., D, B \rangle$ between A and B such that A, D are not adjacent and B, C are not adjacent, then orient $A \circ \neg B$ and every edge on p as undirected edges (—).
 - (vi) If $A B \rightarrow C$ (A and C may or may not be adjacent), then orient $B \rightarrow C$ as $B \rightarrow C$.
 - (vii) If $A \multimap B \multimap *C$ and A, C are not adjacent, then orient $B \multimap *C$ ad $B \multimap *C$.
 - (viii) If $A \rightarrow B \rightarrow C$ or $A \multimap B \rightarrow C$, and $A \multimap C$, orient $A \multimap C$ as $A \rightarrow C$.
 - (ix) If $A \circ \to C$, and $p = \langle A, B, D, ..., C \rangle$ is an uncovered potentially directed path from A to C such that C and B are not adjacent, then orient $A \circ \to C$ as $A \to C$.
 - (x) Suppose $A \circ \to C$, $B \to C \leftarrow D$, p_1 is an uncovered potentially directed path from A to B, and p_2 is an uncovered potentially directed path from A to D. Let m be the vertex adjacent to A on p_1 (m could be B), and w be the vertex adjacent to A on p_2 (w could be D). If m and w are distinct, and are not adjacent, then orient $A \circ \to D$ as $A \to D$.

Appendix C CCI Details

Algorithm 3 Cyclic Causal Inference (CCI)

Input: A conditional independent oracle for a distribution \mathcal{P} , satisfying global directed Markov property and faithfulness conditions w.r.t. a directed graph \mathcal{G} with vertex set \mathcal{V} . ($\mathcal{V} = \mathbf{O} \cup \mathbf{L} \cup \mathbf{S}$), where \mathbf{O} , \mathbf{L} , and \mathbf{S} refer to the sets of observable, latent, and selection variables, respectively.

Output: A PAG $\hat{\mathcal{G}}'$ for the Markov equivalence class of DMGs, $Equiv(\mathcal{G})$.

- 1: Step 1. Run FCI's skeleton discovery procedure.
- 2: Step 2. Run FCI's collider structure (v-structure) orientation procedure.
- 3: Step 3. For any triplet $\langle O_i, O_k, O_j \rangle$, such that we have $O_k \circ -*O_i$, if $O_i \perp\!\!\!\perp O_j \mid \mathbf{Sepset} \langle O_i, O_j \rangle \cup \mathbf{S}$, where $\mathbf{Sepset} \langle O_i, O_j \rangle$ is a separating set discovered in step 1, $O_k \notin \mathbf{Sepset} \langle O_i, O_j \rangle$, $O_i \perp\!\!\!\!\perp O_k \mid \mathbf{Sepset} \langle O_i, O_j \rangle \cup \mathbf{S}$ and $O_i \perp\!\!\!\!\perp O_k \mid \mathbf{Sepset} \langle O_i, O_j \rangle \cup \mathbf{S}$, then orient $O_k \circ -*O_i$ as $O_k \leftarrow *O_i$.
- 4: Step 4. Find additional non-minimal d-separating sets.
- 5: m = 0
- 6: repeat
- 7: repeat
- 8: Select an ordered triplet $\langle O_i, O_j, O_k \rangle$ with the collider structure $O_i * \rightarrow O_j \leftarrow *O_k$ such that $|\mathbf{PD\text{-}Sep}(O_i)| \ge m$
- 9: **repeat**
- 10: Select a subset

 $\mathbf{W} \subseteq \mathbf{PD\text{-}Sep}(O_i) \setminus \{\mathbf{Sepset}(O_i, O_k) \cup \{O_j, O_k\}\}\$ with m vertices

 $\mathbf{T} = \mathbf{W} \cup \mathbf{Sepset}\langle O_i, O_k \rangle \cup O_j$

if O_i and O_k are d-separated given $\mathbf{T} \cup \mathbf{S}$, then record the set \mathbf{T} in **Supset** $\langle O_i, O_j, O_k \rangle$

- 11: **until** all subsets
 - $\mathbf{W} \subseteq \mathbf{PD\text{-}Sep}(O_i) \setminus \{\mathbf{Sepset}\langle O_i, O_k \rangle \cup \{O_j, O_k\}\}\$ have been considered or a d-separating set of O_i and O_k has been recorded in $\mathbf{Supset}\langle O_i, O_i, O_k \rangle$;
- 12: **until** all triplets $\langle O_i, O_j, O_k \rangle$ with the collider structure $O_i * \rightarrow O_j \leftarrow *O_k$ and $|\mathbf{PD\text{-Sep}}(O_i)| \ge m$ have been selected;
- 13: **until** all ordered triplets $\langle O_i, O_j, O_k \rangle$ with the collider structure $O_i * \rightarrow O_j \leftarrow *O_k$ have $|\mathbf{PD\text{-Sep}}(O_i)| < m$;

- 15: **Step 6.** For any two vertices O_i and O_k , if we have $O_i \perp \!\!\! \perp O_k \mid \mathbf{W} \cup \mathbf{S}$ for some $\mathbf{W} \subseteq \mathbf{O} \setminus \{O_i, O_k\}$ discovered in step 1 or step 4 with $O_j \in \mathbf{W}$ but we have $O_i \perp \!\!\! \perp O_k \mid O_l \cup \mathbf{W} \cup \mathbf{S}$, then orient $O_l \circ \!\!\! \!\!\! * O_j$ as $O_l \leftarrow \!\!\! * O_j$.
- 16: *Step 7.* Execute orientation rules until no more endpoints can be oriented. The orientation rules are as follows:
 - (i) If we have $O_i * \to O_j \circ *O_k$ with O_i and O_k non-adjacent, then orient $O_j \circ *O_k$ as $O_j *O_k$. Furthermore, if $O_i * \to O_j$ is not potentially 2-triangulated w.r.t. O_k , then orient $O_j - \circ O_k$ as $O_j - \circ O_k$.
 - (ii) If we have $O_i *O_j \circ *O_k$ with O_i and O_k non-adjacent, and $O_j \circ O_k$ is not potentially 2-triangulated w.r.t. O_i , then orient $O_i \circ *O_k$ as $O_i *O_k$.
 - (iii) Suppose we have $O_i * \to O_j O_k$ with O_i and O_k non-adjacent, and $O_i * \to O_j$ is potentially 2-triangulated w.r.t. O_k . If $O_i * \to O_j$ can be potentially 2-triangulated w.r.t. O_k using only one vertex O_l in the triangle involve $\{O_i, O_j, O_l\}$, then orient $O_i * \to O_l$ as $O_i * \to O_l$, $O_j \circ *O_l$ as $O_j * \to O_l$ and/or $O_j * \to O_l$ as $O_j * \to O_l$. Next, if there exists only one potentially undirected path \prod_{O_l,O_k} between O_l and O_k , then substitute all circle endpoints on \prod_{O_l,O_k} with tail endpoints (-).
 - (iv) If $O_i * \to O_j *O_k$, there exists a path $\prod = \langle O_k, ..., O_i \rangle$ with at least $n \ge 3$ vertices such that we have $O_h *O_{h+1}$ for all $1 \le i \le n-1$, and we have $O_1 *O_n$, then orient $O_1 *O_n$ as $O_1 *O_n$.
 - (v) If we have the sequence of vertices $\langle O_1, ..., O_n \rangle$ such that $O_i O_{i+1}$ with $1 \le i \le n-1$, and we have $O_1 O_n$, then orient $O_1 O_n$ as $O_1 O_n$.
 - (vi) If we have $O_k * \multimap O_i$, there exists a non-potentially 2-triangulated path $\prod = \langle O_i, O_j, O_l, ..., O_k \rangle$ such that $O_k * \multimap O_i$ is not potentially 2-triangulated w.r.t. O_j , and $O_j * \multimap * O_k * \multimap O_k$ is an unshielded non-v-structure, then orient $O_k * \multimap O_i$ as $O_k * \multimap O_i$.
 - (vii) Suppose we have $O_i \circ -O_k$, $O_i *O_k * -O_l$, a non-potentially 2-triangulated path \prod_1 from O_i to O_j , and a non-potentially 2-triangulated path \prod_2 from O_i to O_l . Let O_m be a vertex adjacent to O_i on \prod_1 , and let O_n be the vertex adjacent to O_i on \prod_2 . If further $O_m * *O_i * *O_n$ is an unshielded non-v-structure and $O_i \circ *O_k$ is not potentially 2-triangulated w.r.t. both O_n and O_m , then orient $O_i \circ *O_k$ as $O_i *O_k$.

Appendix D Coefficient Matrices (B) Used in Simulation

In this appendix, we present the weight matrices **B** that were used in the simulation. There are eight matrices in total, with each matrix corresponding to a specific condition ($5p \ sparse = 5$ -variable low density, $5p \ dense = 5$ -variable high density, $5p \ sparse \ LC = 5$ -variable low density with a latent confounder, $5p \ dense \ LC = 5$ -variable high density with a latent confounder, $10p \ sparse = 10$ -variable low density, $10p \ dense = 10$ -variable high density, $10p \ sparse \ LC = 10$ -variable low density with latent confounders, $10p \ dense \ LC = 10$ -variable high density with latent confounders).

$$\mathbf{B}_{5p \ sparse} = \begin{bmatrix} 0 & 0 & 0 & 0 & 0 \\ 1 & 0 & 0.8 & 0 & 0 \\ 0 & 0 & 0 & 0.9 & 0 \\ 0 & 0.7 & 0 & 0 & 0.5 \\ 0 & 0 & 0 & 0 & 0 \end{bmatrix}$$

$$\mathbf{B}_{5p \ dense} = \begin{bmatrix} 0 & 0 & 0 & 0 & 0 \\ 1 & 0 & 0.8 & 0 & 0 \\ 0 & 0 & 0 & 0.9 & 0 \\ 0 & 0.7 & 0 & 0 & 1.5 \\ 1 & 0 & 0 & 0 & 0 \end{bmatrix}$$

 LC_1

	X_1	X_2	X_3	X_4	X_5	X_6	X_7	X_8	X_9	X_{10}
	0	0	0	0	0	0	0	0	0	0
	0	0	0.8	0	0	0	0	0	0	0
	0.4	0	0	0	0	0	0	0	0	0
	0	0	0.7	0	0	X ₆ 0 0 0 0 0.9 0 0 0 0 0 0 0	0	0	0	0
$\mathbf{B}_{10p \; sparse} = $	0	0	0	1	0	0	0	0	0	0
	0	0	0	0	0.8	0	0.5	0	0	0
	0	0	0	0	0	0	0	0	0.8	0
	0	0	0	0	0	0	1	0	0	0
	0	0	0	0	0	0	0	0	0	0.5
	0	0	0	0	0	0	0	0	0	0

	X_1	X_2	X_3	X_4	X_5	X_6	X_7	X_8	X_9	X_{10}	LC_1	LC_2	
	0	0	0	0	0	0	0	0	0	0	0	0	
	0	0	0.8	0	0	0	0	0	0	0	0	0.7	
	0.4	0	0	0	0	0	0	0	0	0	0	0	
	0	0	0.7	0	0	0.9	0	0	0	0	0	0	
	0	0	0	1	0	0	0	0	0	0	0	0.7	
$\mathbf{B}_{10p \; sparse \; LC} =$	0	0	0	0	0.8	0	0.5	0	0	0	0	0	
	0	0	0	0	0	0	0	0	0.8	0	0	0	
	0	0	0	0	0	0	1	0	0	0	0.8	0	
	0	0	0	0	0	0	0	0	0	0.5	0	0	
	0	0	0	0	0	0	0	0	0	0	0.6	0	
	0	0	0	0	0	0	0	0	0	0	0	0	
	0	0	0	0	0	0	0	0	0	0	0	0	

 X_2 X_3 X_5 X_6 X_7 X_8 X_9 X_{10} LC_1 LC_2 X_4 0.5 0.8 0.7 0.7 0.9 1.2 0.7 $\mathbf{B}_{10p\ dense\ LC} =$ 0.8 0.5 0.8 0.5 0.4 0.8 0.5

Appendix E True Ancestral Graph Derivation

This appendix provides a more detailed description of how to obtain the true ancestral graph from a DCG. The process consists of two steps. The first step step involves checking whether there exists an *inducing path* connecting a pair of observed vertices. If such a path exists, we make them adjacent by adding an edge between them. An inducing path is defined as follows.

Definition 1 (Inducing path) An undirected path U between observed vertices O_i and O_j is called an inducing path if and only if every observed vertex on U is a collider except for the endpoints O_i and O_j , every collider on U is an ancestor of either O_i and O_j , and every non-collider on U except for the endpoints is latent variable (Spirtes et al., 2000).

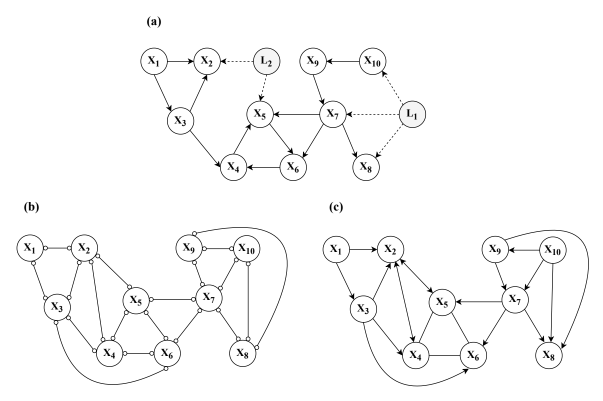
In the second step, we place either an arrow head (>) or tail (-) for every edge-endpoint based on the ancestral relationships between the corresponding vertices, as explained in Section 3.2.1.

To illustrate the procedure, we will use an example. Consider the DCG presented in Figure E1(a). By applying the previously defined concept of inducing path, we can identify the following inducing paths between the pairs of variables:

$$\begin{cases} X_2 \sim X_5: X_2 \leftarrow L_2 \rightarrow X_5 \\ X_2 \sim X_4: X_2 \leftarrow L_2 \rightarrow X_5 \leftarrow X_4 \\ X_3 \sim X_6: X_3 \rightarrow X_4 \leftarrow X_6 \\ X_5 \sim X_7: X_5 \rightarrow X_6 \leftarrow X_7 \\ X_7 \sim X_{10}: X_7 \leftarrow L_1 \rightarrow X_{10} \\ X_8 \sim X_{10}: X_8 \leftarrow L_1 \rightarrow X_{10} \\ X_9 \sim X_8: X_9 \rightarrow X_7 \leftarrow L_1 \rightarrow X_8. \end{cases}$$

Based on the list of identified inducing paths above, we can construct the ancestral skeleton, as shown in Figure E1(b). Then, the true ancestral graph can be derived by examining the ancestral relationships between pairs of vertices. For example, orient $X_1 \rightarrow X_2$ when X_1 is an ancestor of X_2 , orient $X_2 \leftarrow X_5$ when X_2 and X_5 are not ancestors of each other, and orient $X_4 \leftarrow X_5$ when X_4 and X_5 are ancestors of each other. Recall that bidirected edges (\leftarrow) denote mutual non-ancestral relations between vertices indicating the presence of latent confounders, and undirected edges (\leftarrow) indicate mutual ancestral relations implying cyclic relations between vertices. Once all the edge endpoints have been oriented, we can obtain the true ancestral graph, as shown in Figure E1(c).

Figure E1. Derivation of the true ancestral graph.



Note. Panel (a) shows an example DCG with two latent confounders L_1 and L_2 . Panel (b) depicts the ancestral skeleton of the DCG shown in panel (a). Panel (c) depicts the true ancestral graph of the DCG shown in panel (a).

Appendix F 5-variable Dense Case Examination

In Section 4.4, we reported an unexpected finding that the performance of the algorithms deteriorated (e.g., precision and recall decreased, while SHD increased) as the sample size *N* increased in the 5-variable dense conditions, which was contrary to our expectations. In this appendix, we look into this issue in more detail by examining the 5-variable dense condition without a latent confounder scenario using the CCD algorithm. Note that the following explanation applies to the other two algorithms as well.

Figure F1(b) and Figure F1(c) present the PAGs generated by the CCD algorithm using a relatively small sample size of 500 and a large sample size of 3000, respectively. We can see that with the small sample size, the algorithm failed to detect the edge between X_2 and X_5 , while with the large sample size, it did detect the edge. Upon the detection of an additional edge between X_2 and X_5 , however, the resulting graph became very dense, with almost every vertex connected to every other vertex. As a consequence, the algorithm failed to orient any edges, as shown in Figure F1(c).

(a) (b) (c) X_1 X_2 X_4 X_4 X_2 X_4 X_4 X_5 X_5

Figure F1. Graphs from the 5-variable dense condition without a latent confounder.

Note. Panel (a) shows the true ancestral graph for the 5-variable dense condition without a latent confounder. Panels (b) and (c) show the partial ancestral graphs (PAGs) estimated by CCD, using a relatively small sample size (N = 500) and a large sample size (N = 3000), respectively. The PAGs estimated by the FCI and CCI algorithms also exhibit a similar pattern.

Before the edge was detected, the algorithm was capable of performing orientation, and correctly oriented some edges by identifying the collider X_4 , despite some wrongly oriented edges like the incoming arrows towards X_1 . This explains the decrease in SHD and the slight increase in precision and recall with small sample sizes. Given the true graph shown in Figure F1(a), we can actually compute the SHD. The PAG with the small sample size in Figure F1(b) has an SHD value of 13, while the PAG with the large sample size in Figure F1(b) has an SHD value of 16.

To investigate why the algorithm failed to detect the edge with the small sample size, we examined all partial correlations between X_2 and X_5 across 500 simulations, using a large sample size of N=10000. The resulting distributions of partial correlations are shown in Figure F2. Notably, the results reveal a very small partial correlation between X_2 and X_5 when conditioned on X_3 ($\rho_{X_2,X_5|X_3}=0.04$). Also, we examined the results of conditional independence tests for all possible independence patterns between X_2 and X_5 , which are summarized in Table F1. The null hypothesis of independence was not rejected in more than half of the cases until the sample size reached 2000, which led to the omission of the edge between X_2 and X_5 . However, for sample sizes greater than 2000, the null hypothesis of independence was rejected, indicating that the algorithm was able to detect the edge between X_2 and X_5 . Consequently, the resulting graph became too dense to orient any edges, leading to a fully undirected ancestral graph, as shown in Figure F1(c).

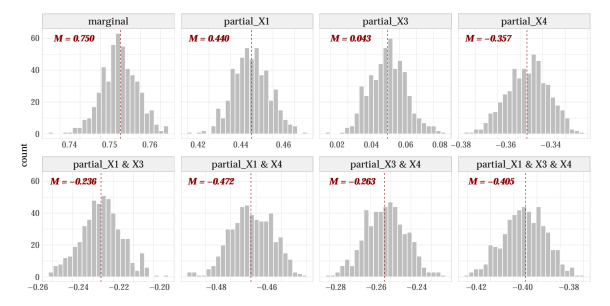


Figure F2. Distributions of marginal & partial correlations between X_2 and X_5 .

Note. The figure displays the distributions of the estimated marginal and partial correlations between X_2 and X_5 across 500 simulations based on a sample size of N=10000. The top left panel displays the distribution of marginal correlations between X_2 and X_5 , which reflects their correlation without controlling for other variables. The remaining panels represent the distributions of partial correlations between X_2 and X_5 while controlling for one or more variables, as indicated by the labels. M denotes the mean value of the marginal/partial correlations.

N	50	150	500	1000	2000	3000	4000	5000	7500	10000
Marginal	0	0	0	0	0	0	0	0	0	0
Cond. X_1	0.12	0	0	0	0	0	0	0	0	0
Cond. X_3	0.94	0.91	0.83	0.73	0.63	0.49	0.34	0.22	0.14	0.01
Cond. X_4	0.28	0	0	0	0	0	0	0	0	0
<i>Cond.</i> $\{X_1, X_4\}$	0.06	0	0	0	0	0	0	0	0	0
<i>Cond.</i> $\{X_1, X_3\}$	0.63	0.14	0	0	0	0	0	0	0	0
<i>Cond.</i> $\{X_3, X_4\}$	0.60	0.08	0	0	0	0	0	0	0	0
Cond. $\{X_1, X_3, X_4\}$	0.16	0	0	0	0	0	0	0	0	0

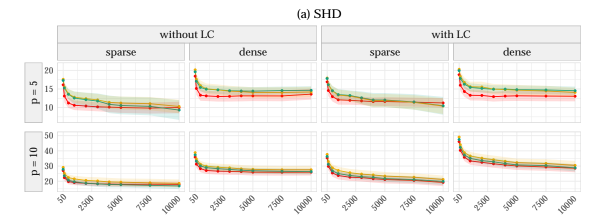
Table F1. Conditional independence test results (p > 0.01).

Note. The table shows all possible (conditional) independence patterns between X_2 and X_5 for different sample sizes (N) used in the simulation study. Each row corresponds to a specific independence relation, and each cell indicates the proportion of *p-values* that are greater than the α level of 0.01 (p > 0.01) out of 500 simulations, indicating that the null hypothesis of independence is not rejected. Marginal = not conditional on any variables; Cond. = conditional on.

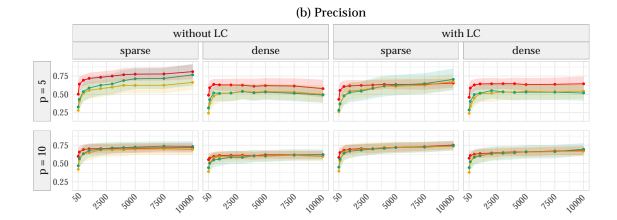
Appendix G Sensitivity Analysis 1: Randomly Sampling Weights

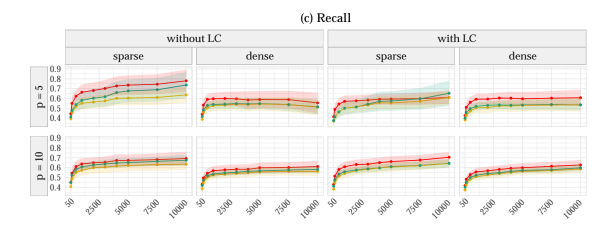
This appendix reports the complete results of the secondary simulation study where we randomly sampled parameters of B matrices. Figure G1 summarizes the results, which reveal several differences from the original study. First, CCD's performance appears to have improved across all conditions and shows comparable SHD values to CCI, even in sparse conditions (see Figure G1(a)). The outperformance of CCD is more evident when looking at precision (Figure G1(b)) and recall (Figure G1(c)). However, this is because CCD is more conservative and produces more circle endpoints than CCI, almost always recording higher uncertainty rates than CCI (see Figure G1(d)). Therefore, the overall conclusion remains the same as in the original simulation study; CCI is still the preferred choice when seeking to obtain the most information on causal directions with low uncertainty. Second, we observe that the FCI algorithm exhibits more stability and produces more circle endpoints, leading to similar levels of uncertainty as CCD across different conditions. However, its overall inferior performance compared to the other algorithms remains consistent with the findings of the original simulation study. Finally, we notice a significant change in the patterns in the 5-variable dense conditions. Instead of the odd dips and spikes observed previously in the original simulation study, we now see a steady decrease in SHD and an increase in precision and recall values until they plateau. As explained in Section 4.4.2, the odd drops and spikes were a result of the specific weights chosen in those cases, and therefore, such patterns are expected to disappear when weight parameters are randomly sampled. Based on these findings, we can conclude that the results of the original simulation study are not substantially influenced by the particular choices of **B** matrices, except for the atypical 5-variable dense cases. 14

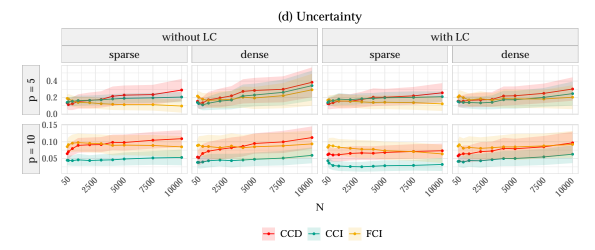
Figure G1. Performance analysis with randomly sampled weights for B matrices.



 $^{^{14}}$ Note that these results are based on sampling weights from both negative and positive values using a uniform distribution on $[-0.9, -0.1] \cup [0.1, 0.9]$. As an extra robustness check, we also performed the same analysis with only positive weights sampled from a Uniform([0.1, 0.9]). The outcomes were almost identical, with no discernible differences.





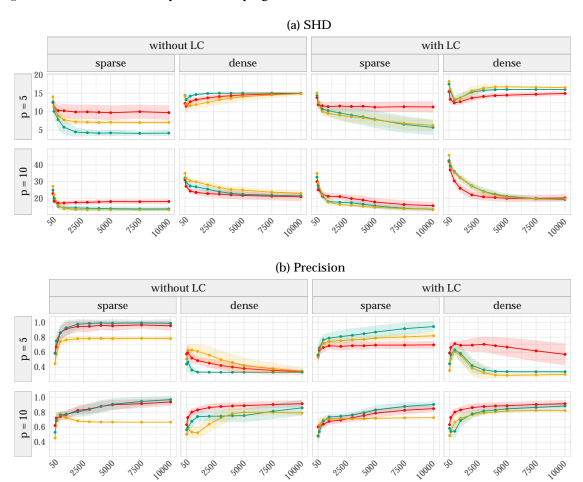


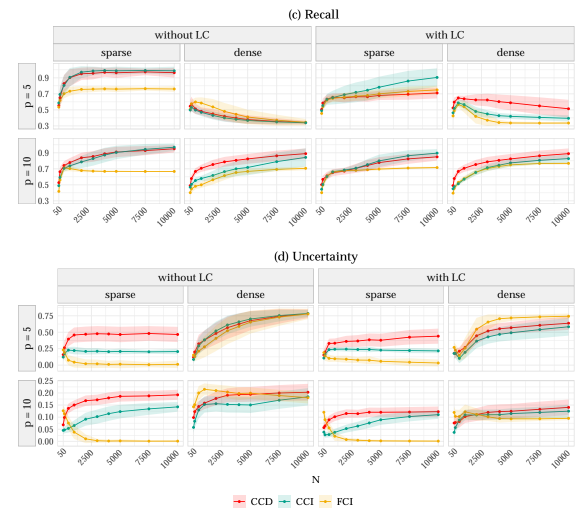
Note. Each point on the plot represents the average value of the corresponding metric across 500 iterations, while the shaded region represents the interquartile range (IQR) of the metric's distribution.

Appendix H Sensitivity Analysis 2: Varying α Levels

In this appendix, we present the findings of a secondary simulation study in which we varied the significance level (α) based on sample size. The α level plays a crucial role in constraint-based causal discovery (Spirtes et al., 2000), since it can be either too strict and indicate everything as independent or not strict enough and fail to find any independence relationships. To ensure consistent results from conditional independence tests, we lowered the α level as the sample size increased in this simulation study. The results can be found in Figure H1, which show that the observed patterns are almost identical to those from the original simulation study. Based on these findings, we can conclude that the results of our original simulation are robust to the fixed α level chosen in the study (i.e., $\alpha = 0.01$).

Figure H1. Performance analysis with varying α levels.





Note. Each point corresponds to the average value of a specified metric across 500 iterations, and the shaded area represents the interquartile range (IQR) of the metric's distribution.

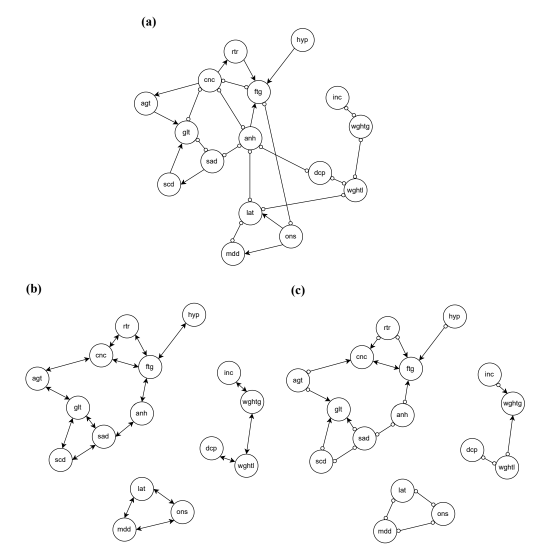
Appendix I Stability Analysis on Empirical Data

We performed a stability analysis on the empirical data in which we randomly sampled 90% of observations from the original dataset with N = 408, creating 1000 subsets with a sample size of 367 (408 × 0.9 = 367.2). Subsequently, we applied all three algorithms (CCD, FCI, and CCI) to each of the 1000 subsets and retained only the edge-endpoints that were identified more than 70% of the time. The PAGs resulting from each algorithm are presented in Figure I1. Overall, we observe that the causal structures found in the stability analysis are very similar to those obtained in the original analysis in Section 5.2. However, the resulting PAGs from the stability analysis are slightly more sparse, with a few missing edges. For example, the PAG estimated by CCD in Figure I1(a) no longer includes the edge between guilt (glt) and increased appetite (inc), which was present in the original analysis. Similarly, the PAGs estimated by CCI and FCI in Figure I1(b) and Figure I1(c),

¹⁵Note that the underlinings generated by the CCD algorithm are disregarded in this analysis.

respectively, do not show the edges between guilt (glt) and concentration (cnc) or anhedonia (anh) and decreased appetite (dcp). Additionally, there appears to be some loss of orientation in the PAGs, particularly in the one obtained from CCD. The CCD PAG in Figure I1(a) contains relatively more circle endpoints compared to the original result, which makes the cyclic structures observed in the original analysis less evident. Nevertheless, the results remained largely the same, except for only a couple of missing edges and a few arrow heads/tails replaced by circle marks. This suggests that our original results are stable and not significantly affected by minor variations in the data.

Figure 11. Estimated PAGs from stability analysis.



Note. Panels (a), (b), and (c) present the PAGs estimated by CCD, CCI, and FCI, respectively. Each PAG is obtained by selecting only the endpoints that were identified more than 70% of the times across 1000 subsets of the original data. ons = sleep onset insomnia; mdd = middle insomnia; lat = late (early morning awakening); hyp = hypersomnia; sad = sad; dcp = decreased appetite; inc = increased appetite; wghtl = weight loss; wghtg = weight gain; cnc = concentration impairment; glt = guilt; scd = suicidal thoughts; anh = anhedonia; ftg = fatigue; rtr = psychomotor retardation; agt = agitation.

Appendix J Distribution of Empirical Data

The distributions of depression symptoms are illustrated in Figure J1, which clearly indicates that most of them are skewed and do not follow a Gaussian distribution. Given that partial correlation is known to be a valid measure of conditional independence only in the case of multivariate Gaussian (Baba et al., 2004), it is possible that our analysis, which relied on partial correlations to test for conditional independence despite the non-Gaussian nature of the data, may have been misleading.

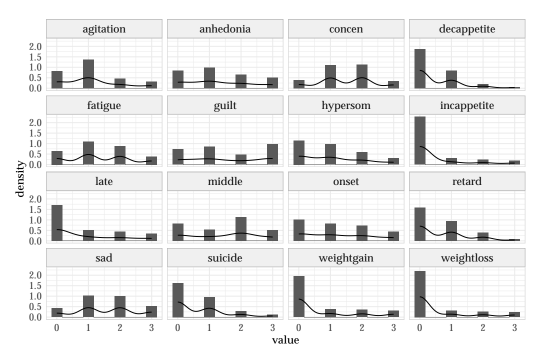


Figure J1. Distributions of depression symptoms.

Note. Each panel shows the distribution of individual depression symptom in the dataset from McNally et al. (2017). concen = concentration impairment; decappetite = decreased appetite; hypersom = hypersomnia; incappetite = increased appetite; late = late (early morning awakening); middle = middle insomnia; onset = sleep onset; retard = psychomotor retardation; suicide = suicidal thoughts.