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# 21 When can association graphs admit a causal interpretation?

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ABSTRACT

We discuss essentially linear structures which are adequately represented by association graphs called covariance graphs and concentration graphs. These do not explicitly indicate a process by which data could be generated in a stepwise fashion. Therefore, on their own, they do not suggest a causal interpretation. By contrast, each directed acyclic graph describes such a process and may offer a causal interpretation whenever this process is in agreement with substantive knowledge about causation among the variables under study. We derive conditions and procedures to decide for any given covariance graph or concentration graph whether all their pairwise independencies can be implied by some directed acyclic graph.

## 21.1 Introduction

Many of the most widely used statistical techniques center around the interpretation of the covariance matrix. The exponential family of the multivariate normal distribution provides a justification to reduce data to sample means,  $\bar{y}_i$ , i = 1, ..., p, and a covariance matrix, S, and for some purposes to sample means and a concentration matrix,  $S^{-1}$ , that is the inverse covariance matrix.

Under multivariate normality, a zero covariance,  $\sigma_{ij} = 0$ , means marginal independence and a zero concentration,  $\sigma^{ij} = 0$ , means conditional independence of variable pair  $Y_i, Y_j$  given all p - 2 remaining variables. In general, these conditions just indicate linear independencies, since the standardized versions of covariance and concentration are the marginal correlation coefficient,  $\rho_{ij}$ , and the partial correlation coefficient,  $\rho_{ij,k}$ , given all the remaining variables  $k = \{1, ..., p\} \setminus \{i, j\}$ :

$$\rho_{ij} = \sigma_{ij} (\sigma_{ii} \sigma_{jj})^{-1/2}, \rho_{ij,k} = -\sigma^{ij} (\sigma^{ii} \sigma^{jj})^{-1/2}.$$

Thus, data reduction to covariance or concentration matrix implies that essentially linear

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relations among the variables under study are of interest. These may be relevant not only for continuous but also for ordinally scaled variables or for dichotomous variables. But, since this is a fairly strong assumption in some applications, checks for nonlinearity or special linearisations are needed, such as those discussed in [Cox-Wer 92a, Cox-Wer 92b] and in [Wermuth-Cox 92].

A covariance matrix with some zero off-diagonal entries can be represented by an undirected graph with the corresponding edges missing [Cox-Wer 93] called **covariance graph**, abbreviated by  $G_{cov}$  here. Similarly, a concentration matrix with some zero entries can be represented by an undirected graph with the corresponding edges missing [Kiiveri-Spe 82] called **concentration graph**, abbreviated by  $G_{con}$  here.

A complete covariance or concentration graph has no missing edges, that is, it has exactly one edge for each variable pair, 1/2 p (p-1) in all. There can be a number of reasons for considering special incomplete undirected graphs. For instance, it could be that

- the linear independencies expressed thereby are themselves of substantive interest,
- reduction of dimensionality is needed to keep a reasonable balance between the number of parameters to be estimated and the number of available independent observations, or
- it is desired to decide whether the structure given by a covariance or concentration graph could have been generated by some stepwise process of univariate recursive regressions.

We concentrate here on the last aspect. This implies in particular that we consider covariance and concentrations graphs less as representing models of their own standing but, rather, a possible reflection of some directed acyclic graph, abbreviated as **dag**, which represents a recursive process for stepwise data generation. Such a process would admit a causal interpretation if the order of the variables involved in the process, especially their classification into response and explanatory variables, is in agreement with subject-matter knowledge about the causal process [Cox 92].

Each dag can be thought of as specifying dependence relationships between an ordered set of variables  $\{Y_1, ..., Y_p\}$  where j > i designates  $Y_j$  as potentially explanatory for  $Y_i$ ; it becomes actively explanatory in case an arrow is drawn from  $Y_j$  to  $Y_i$ . A missing edge for variable pair  $Y_i, Y_j$  has then the interpretation: response  $Y_i$  is (linearly) independent of  $Y_j$  given the remaining potential explanatory variables of  $Y_i, \{Y_{i+1}..., Y_{j-1}, Y_{j+1}, ..., Y_p\}$ . The set of missing edges defines the independence structure of the dag and permits<sup>3</sup> the stronger statement that  $Y_i$  is conditionally independent of all variables in  $\{Y_{i+1}, ..., Y_p\}$  that are not linked to  $Y_i$ , given those that are linked to  $Y_i$ . Each dag **implies** a set of pairwise marginal (linear) independencies, i.e. a covariance graph, and a set of pairwise conditional (linear) independencies given all p - 2variables, i.e. a concentration graph. By "implied" we mean independencies that must hold in *every* distribution that fulfills all the missing-link conditions shown in the dag. This excludes independencies that are just introduced by accidental equalities among the numerical values of some parameters in the model. From a given dag, these implied independencies may be read off directly, using the graphical criterion of d-separation [Pearl 88, page 117] and [Geiger etal 90].

For example the following Markov chain

<sup>&</sup>lt;sup>3</sup>Provided S is non-singular.

is a dag which implies a complete covariance graph, i.e. no marginal independencies, and a chainlike concentration graph, obtained by removing the arrow heads. If r, s, t, u denote correlations between pairs  $(Y_1, Y_2), (Y_2, Y_3), (Y_3, Y_4), (Y_4, Y_5)$ , then the correlation matrix implied by the Markov chain has the form

1	r	rs	rst	rstu
•	1	s	st	stu
•	•	1	t	tu
•	•	٠	1	u
•	٠	•	•	1.

It can be used to explain why it may be difficult to decide on marginal linear independence merely from observed marginal correlations. Suppose r = s = t = u and r = 0.3 then  $r^2 = 0.09$ ,  $r^3 = 0.027$ ,  $r^4 = 0.0081$  would be close to zero; even when r = 0.6,  $r^4 = 0.13$  would still not be far from zero. Thus, if we know that the data are generated by the above Markov chain we can conclude that all marginal correlation are nonzero but a correlation is weaker the further the variables are apart in the chain. However, if the process by which the data actually are generated is not known, and we have to rely merely on data inspection or associated tests, we may judge some of the smaller correlations to be zero apart from sample fluctuations i.e. we come to erroneous conclusions about the covariance structure (represented by the covariance graph) or the generating process. Similar arguments apply to the concentration graph.

Formal tests for agreement of an observed covariance matrix or concentration matrix with a corresponding hypothesized association graph are available. An arbitrary pattern of zeros in a covariance matrix is a special case of hypotheses linear in covariances [Anderson 73] and an arbitrary pattern of zeros in a concentration matrix is a covariance selection model [Dempster 72]. Maximum likelihood estimates and associated likelihood ratio tests are available under the assumption of multivariate normality and may for instance be computed with the help of LISREL [Jöreskog-Sör 78] for the former and with the help of MIM [Edwards 92] for the latter. The same estimates are called quasi-likelihood estimates if just the assumption of linearity is retained.

The problem of finding a causal explanation for a general distribution was treated in [Pearl-Wer 91] and, in the case of normal distributions, might require the processing of an exponential number of submatrices (i.e., all the majors of  $G_{cov}$ ). When an explicit list of *all* conditional and marginal independencies is available, the problem can be solved in time polynomial in the length of the list [Verma-Pea 92a] [Dori-Tar 92]. Still, the length of this list can be very large indeed. The current paper attempts to find a causal interpretation on the basis of a more limited information, assuming that the only vanishing dependencies available to the analyst are those corresponding to the zero entries of the covariance and concentration matrices.

To this end, we pose the following set of problems:

• Given a covariance graph,  $G_{cov}$ , decide whether it could have been generated by a directed acyclic graph

(i) with the same nodes and edges,

(ii) with the same nodes but fewer edges,

(*iii*) with some additional nodes (representing latent or hidden variables).

• Given a concentration graph,  $G_{con}$ , decide whether it could have been generated by a directed acyclic graph

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(iv) with the same nodes and edges,

(v) with the same nodes but fewer edges,

(vi) with additional (latent) nodes.

• Given a covariance and a concentration graphs, decide whether

(vii) both could have been generated by the same directed acyclic graph.

Theorems 1 through 7 of the next section present declarative and procedural solutions to these seven problems, respectively. Proofs are omitted, and will be included in an expanded version of this paper.

### 21.2 Main results

**Definition 1** A directed acyclic graph D is said to imply an independence I if I holds in every probability P that fulfills all the missing-link conditions of D, that is, choosing any variable ordering consistent with the direction of arrows in D, P renders each variable  $Y_i$  independent on its inactive explanatory variables (nonadjacent predecessors), given its active explanatory variables (adjacent predecessors). If D fails to imply an independence for pair (i, j) we say that D implies a dependence for that pair which, in turn, will induce an edge ij in  $G_{cov}$  or  $G_{con}$ .

**Lemma 1** D implies an independence I iff I satisfies the d-separation criterion in D [Pearl 88]. In particular, D implies a marginal independency  $\sigma_{ij} = 0$  if every path between nodes i and j in D traverses at least one pair of converging arrows (i.e.,  $\rightarrow \star \leftarrow$ ). Likewise, D implies a conditional independence given all remaining variables,  $\sigma^{ij} = 0$ , if every path between nodes i and j in D traverses at least one pair of non-converging arrows (i.e., either  $\rightarrow \star \rightarrow$  or  $\leftarrow \star \rightarrow$ ).

**Definition 2** (generate and coincide) A dag D is said to (a) generate a covariance graph  $G_{cov}$  if the set of marginal independencies implied by D matches the set of missing edges of  $G_{cov}$  and (b) to coincide with  $G_{cov}$  if D not only generates  $G_{cov}$  but also has the same edges as  $G_{cov}$ .

Similar definitions of generate and coincide apply to  $G_{con}$ . For example, the Markov chain discussed in Section 21.1 represents a dag which coincides with its  $G_{con}$  but not with its  $G_{cov}$ , since some edges in  $G_{cov}$  do not appear in the chain.

**Definition 3** ( $\lor$ -configuration) A triplet of nodes (a, b, c) in an undirected graph G is said to be a  $\lor$ -configuration (pronounced as vee-configuration), if ab and bc are edges in G while ac is not.

**Definition 4** (sink orientation) Given an undirected graph G, the sink orientation of G results from assigning arrows  $a \longrightarrow b \longleftarrow c$  to each  $\lor$ -configuration (a, b, c) in G.

Note that a sink orientation may contain bi-directed edges, since two opposing arrows may be assigned to the same edge. For example, the sink orientation of the 4-chain a - b - c - d is given by  $a \longrightarrow b \longleftrightarrow c \longleftarrow d$ , since both (a, b, c) and (b, c, d) are  $\lor$ -configurations. We mark that the sink orientation of a graph with n nodes can be formed in  $O(n^3)$  steps.

**Theorem 1** The following statements are equivalent. A covariance graph  $G_{cov}$  coincides with a dag D if and only if (i) no edge in the sink orientation of  $G_{cov}$  is bi-directed; (ii)  $G_{cov}$  contains no chordless 4-chain. Moreover, if the sink orientation of  $G_{cov}$  contains no bi-directed edges,

then every acyclic orientation of the remaining undirected edges constitutes a dag that coincides with  $G_{cov}$ .

**Example 1** Consider the graph



The 4-chain a-b-c-d is chordless, hence, there is no dag that coincides with  $G_{cov}$ . Indeed, the  $\lor$ -orientation of  $G_{cov}$  is



which renders the edge  $b \leftrightarrow c$  bi-directed.

Turning to problem (ii), we first define the notions of sink completion and exterior cliques.

**Definition 5** (sink completion) A sink completion of an undirected graph  $G_{cov}$  is any dag obtained from  $G_{cov}$  as follows: a) the sink orientation of  $G_{cov}$  is formed, b) all bi-directed edges are removed, c) the remaining undirected edges are oriented to form a dag.

Lemma 2 Every undirected graph has at least one sink completion.

This follows from the facts that (1) every sink orientation is acyclic and (2) any acyclic partial orientation of a graph can be completed to form a dag.

**Definition 6** (exterior cliques) A clique in an undirected graph is said to be exterior if it contains at least one node that is adjacent only to nodes in that clique. We call such a node "extremal".

Note that finding an extremal node in a graph with n nodes and E edges takes at most  $nE^2$  steps; for each node i we test whether every two neighbors of i are adjacent to each other. The same procedure can be used to enumerate all exterior cliques of a graph, since an extremal node must reside in one and only one clique.

**Theorem 2** The following statements are equivalent: (i)  $G_{cov}$  is generated by a directed acyclic graph; (ii) a sink completion of  $G_{cov}$  implies all edges of  $G_{cov}$ ; (iii) every edge of  $G_{cov}$  resides in an exterior clique.

**Example 2** Consider again the graph  $G_{cov}$  of Example 1. The sink completion of  $G_{cov}$  is given by the dag  $D = a \longrightarrow b \longleftarrow e \longrightarrow c \longleftarrow d$  which implies all edges of  $G_{cov}$ . In particular, the edge  $b \longrightarrow c$ , missing from D, is implied by the path  $b \longleftarrow e \longrightarrow c$ , which does not contain converging arrows (see Lemma 1.) Hence D generates  $G_{cov}$ . Indeed, following condition (*iii*), the exterior cliques of  $G_{cov}$  are (a, b), (b, c, d) and (c, d), which contain every edge of  $G_{cov}$ . A simple example of a graph  $G_{cov}$  that cannot be generated by a dag is the 4-chain  $a\_b\_c\_d$ . The sink completion of  $G_{cov}$  is the disconnected dag  $a \longrightarrow b = c \longleftarrow d$  which does not imply the edge  $b\_c$  in  $G_{cov}$ . Likewise,  $G_{cov}$  has only two exterior cliques, (a, b) and (c, d), thus violating condition (*iii*) relative to edge  $b\_c$ .

Turning to Problem (iii), we have:

**Theorem 3** Every covariance graph  $G_{cov}$  can be generated by a directed acyclic graph with additional nodes by replacing each edge ab of  $G_{cov}$  with a  $\leftarrow (*) \rightarrow b$ , where (\*) represents an unobserved variable.

Remark: To reduce the number of latent nodes, it is not necessary to replace each edge of  $G_{cov}$ , but only those which are not implied by the sink completions of  $G_{cov}$ . For example, the 4-chain  $a\_b\_c\_d$  is generated by the dag  $a \longrightarrow b \leftarrow (*) \longrightarrow c \leftarrow d$ , where the replaced edge bc is the only one that is not implied by the sink completion of this chain (see Example 2).

We now turn to problems (iv) - (vi) where we seek a dag that agrees with the structure of a given concentration matrix.

**Theorem 4** A concentration graph  $G_{con}$  coincides with some dag D if and only if  $G_{con}$  is a chordal graph, i.e. it contains no chordless n-cycle, n > 3. (Effective tests for chordality are well known in the literature [Tarjan-Yan 84].)

**Theorem 5** A concentration graph  $G_{con}$  can be generated by some dag if and only if there exists an ordering O such that the following procedure removes all edges of  $G_{con}$ :

- 1. Find an exterior clique, remove all its extremal nodes and their associated edges, mark all remaining edges in that clique, and add them to a list L of marked edges, in some order O.
- 2. Repeat the process on the reduced graph until either all nodes have been eliminated or, in case the subgraph contains no exterior clique, choose the first marked edge on L, and remove it from the graph.
- 3. Repeat step 1 and 2.

**Example 3** To see how the marking procedure helps identify a generating dag, consider the graph  $G_{con}$  below.



The cliques are: ABC, ACD, BCE, CDE and DEF. Only DEF is exterior, with extremal vertex F. Removing F together with edges DF and FE, leaves a subgraph with no exterior clique. However, edge DE is marked, so it can be removed, which creates new exterior cliques, allowing the process to continue till all edges are removed. Indeed, directing arrows toward each node removed in step 2, yields the dag  $D_1$  above, which generates  $G_{con}$ .

**Example 4** The following example (due to Verma) shows that the order of edge removals may be crucial, that is, failure to eliminate all edges in one ordering does not imply that no elimination ordering exists. Consider the following concentration graph:



The only extremal vertices are E, G, M, and O. If we remove these (in any order) it will result in a graph with no exterior cliques and with the following marked links: CF, DF, KN, and LN. If we first remove the marked edges FC and FD, the process will halt (because the cycle A - B - C - D - A cannot be eliminated.) However, if we first remove the marked edges LNand KN, then we will find a good elimination ordering: ..., N, H, J, L, K, I, F, ...

The fact that it is impossible by local means to decide which of the marked edges should be removed first renders the decision problem a difficult one. While in the example above it is clear that one should postpone the removal of FC and FD, because it leads to an impasse (the cycle A - B - C - D - A), such local clues are not available in the general case. Indeed, Verma and Pearl have shown that the problem is NP-Complete [Verma-Pea 92b]. Nevertheless, effective necessary conditions are available, which makes the decision problem feasible when the number of nodes is not too large.

**Lemma 3** The following are necessary conditions for a concentration graph  $G_{con}$  to be generated by a dag. (1) every chordless n-cycle of  $G_{con}$ , n > 3, must have at least one edge that resides in some k-clique, k > 2. (2)  $G_{con}$  must have at least one exterior clique.

**Example 5** The graph  $G_{con}$  below shows that these two conditions are not sufficient; it satisfies both, yet it cannot be generated by any dag.



The edge removal procedures of Theorem 5 can easily be converted into a backtrack algorithm that guarantees finding a generating dag, if such a dag exists. Whenever the procedure reaches an impasse, the algorithm backtracks to the state of last decision point, and selects another edge for removal. The efficiency of this algorithm can be improved by insisting that at any decision point we consider for removal only edges such that the reduced graph will satisfy the necessary conditions of Lemma 3. In general, we should also try to remove marked edges that belong to chordless *n*-cycles, n > 3, as early as possible, because we know that at least one such edge must eventually be removed if the process is to come to a successful end.

**Theorem 6** A concentration graph  $G_{con}$  can be generated by a dag with additional latent nodes if and only if it can be generated by a dag without latent nodes.

In other words, latent variables do not improve the power of dags to generate richer patterns of vanishing concentrations. This is in sharp contrast to patterns of vanishing covariances (Theorem 3); every such pattern can be generated by a dag with latent nodes.

Finally, turning to problem (vii) we have:

**Theorem 7** A covariance graph  $G_{cov}$  and a concentration graph  $G_{con}$  can both be generated by the same dag only if the following three conditions hold:

- 1.  $G_{cov}$  and  $G_{con}$  must each be generated by a dag.
- 2. The set of nodes of every exterior clique of  $G_{cov}$  must induce a subgraph of  $G_{con}$  that can be generated by a dag.
- 3. Every edge of  $G_{con}$  that does not reside in a larger clique must also be an edge of  $G_{cov}$ .

We are not sure at this point whether the conditions above are sufficient; so far we were not able to find a pair of undirected graphs that fulfill these three conditions and yet cannot be generated by a dag.

## 21.3 Conclusions

This paper provides conditions and procedures for deciding if patterns of independencies found in covariance and concentration matrices can be generated by a stepwise recursive process represented by some directed acyclic graph. If such an agreement is found, we know that one or several causal processes could be responsible for the observed independencies, and our procedures could then be used to elucidate the graphical structure common to these processes, so as to evaluate their compatibility against substantive knowledge of the domain.

If we find that the observed pattern of independencies does not agree with any stepwise recursive process, then there are a number of different possibilities. For instance,

- some weak dependencies could have been mistaken for independencies and led to the wrong omission of edges from the covariance or concentration graphs.

- the sampling procedure used in data collection was not properly randomized. Selection bias tends to induce symmetrical dependencies that cannot be emulated by a recursive process [Pearl 88, page 130].

- the process responsible for the data is non-recursive, involving aggregated variables, simultaneous reciprocal interactions, or mixtures of several recursive processes.

- some of the observed linear dependencies reflect accidental cancellations or hide actual nonlinear relations.

In order to distinguish accidental from structural independencies it would be helpful to conduct several longitudinal studies under slightly varying conditions. These fluctuating conditions would tend to affect the numerical values of the covariances but not the basic structural properties of the process underlying the data. Under such conditions, if the data were generated by a recursive process represented by some directed acyclic graph, those independencies that are "implied" by the dag (see Definition 1) would persist despite the fluctuating conditions, while accidental independencies would be perturbed and hence discarded. We regard this as a possibility to pursue Cochran's suggestion [Cochran 65, page 252] that "when constructing a causal hypothesis one should envisage as many different consequences of its truth as possible and plan observational studies to discover whether each of these consequences is found to hold."

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