A Constraint Optimization Approach to Causal Discovery from Subsampled Time Series Data

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Abstract

We consider causal structure estimation from time series data in which measurements are obtained at a coarser timescale than the causal timescale of the underlying system. Previous work has shown that such subsampling can lead to significant errors about the system's causal structure if not properly taken into account. In this paper, we first consider the search for system timescale causal structures that correspond to a given measurement timescale structure. We provide a constraint satisfaction procedure whose computational performance is several orders of magnitude better than previous approaches. We then consider finite-sample data as input, and propose the first constraint optimization approach for recovering system timescale causal structure. This algorithm optimally recovers from possible conflicts due to statistical errors. We then apply the method to real-world data, investigate the robustness and scalability of our method, consider further approaches to reduce underdetermination in the output, and perform an extensive comparison between different solvers on this inference problem. Overall, these advances build towards a full understanding

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of non-parametric estimation of system timescale causal structures from subsampled time series data.

Keywords: causality, causal discovery, graphical models, time series, constraint satisfaction, constraint optimization.

1 1. Introduction

Time-series data has long constituted the basis for causal modeling in many 2 fields of science (Granger, 1969; Hamilton, 1994; Lütkepohl, 2005). These data often provide very precise measurements at regular time points, but the underlying causal interactions that give rise to those measurements can occur at a much faster timescale than the measurement frequency. As just one example: fMRI 6 experiments measure neural activity (given various assumptions) roughly once per two seconds, but the underlying neural connections clearly operate much 8 more quickly. Time order information can simplify causal analysis since it can provide directionality, but time series data that undersamples the generating 10 process can be especially misleading about the true direct causal connections 11 (Dash and Druzdzel, 2001; Iwasaki and Simon, 1994). 12

For example, Figure 1a shows the causal structure of a process unrolled over 13 discrete time steps, and Figure 1b shows the corresponding structure of the 14 same process, obtained by marginalizing every second time step. If we do not 15 take into account the possibility of subsampling, then we would conclude that 16 Figure 1b gives the correct structure — and thus totally miss the presences of 17 all true edges. This drastic structure misspecification may lead us to perform a 18 possibly costly intervention on Z to control Y, when the influence of Z on Y is, 19 in fact, completely mediated by X and so, intervening on X would be a more 20 effective choice. Also, a (parametric) model with the structure in Figure 1b 21 gives inaccurate predictions when intervening on both X and Z: the value of Y 22 would be predicted to depend on Z and not on X, when in reality Y depends 23 on X and not on Z. 24

Standard methods for estimating causal structure from time series either fo-25 26 cus exclusively on estimating a transition model at the measurement timescale (e.g., Granger causality (Granger, 1969, 1980)) or combine a model of measure-27 ment timescale transitions with so-called "instantaneous" or "contemporane-28 ous" causal relations that aim to capture interactions that are faster than the 29 measurement process (e.g., SVAR (Lütkepohl, 2005; Hamilton, 1994; Hyvärinen 30 et al., 2010)), though only very specific types of interactions can be captured 31 with these latter models. In contrast, we follow Plis et al. (2015a,b) and Gong 32 et al. (2015), and explore the possibility of identifying (features of) the causal 33 process at the true timescale from data that subsample this process. 34

Plis et al. (2015a,b) developed algorithms that can learn the set of causal timescale structures that could yield a given measurement timescale graph, either at a known or unknown undersampling rate. While these algorithms show that the inference problem is solvable, they face a number of computational



Figure 1: (a) The structure of the causal system-scale time series. (b) The structure of the corresponding measurement scale time series if only every second sample is observed i.e. nodes at time slice t - 1 are marginalized. If subsampling is ignored and (b) is thought to depict the true causal structure, all direct causal relationships among $\{X, Y, Z\}$ are misspecified.

challenges that limit their use. They do, however, show the importance of constraints for this problem, and so suggest that a constraint satisfaction approach might be more effective and efficient. Gong et al. (2015) consider finding a linear SVAR from subsampled data. They show that if the error variables are non-Gaussian, the true causal effects matrix can be discovered even from subsampled data. However, their method is highly restricted in terms of numbers of variables and parametric form.

In this paper, we provide an exact discovery algorithm based on using a 46 general-purpose Boolean constraint solver (Biere et al., 2009; Gebser et al., 47 2011), and demonstrate that it is orders of magnitudes faster than the current 48 state-of-the-art method by Plis et al. (2015b). At the same time, our approach is 49 much simpler and, as we show, it allows inference in more general settings. We 50 then develop the approach to integrate possibly conflicting constraints obtained 51 from the data. In addition to an application of the method to the real-world 52 data, we investigate the robustness and scalability of our method, consider fur-53 ther approaches to reduce underdetermination in the output, and perform an 54 extensive comparison between different solvers on this inference problem. More-55 over, unlike the method by Gong et al. (2015), our approach does not depend 56 on a particular parameterization of the underlying model and scales to a more 57 reasonable number of variables. 58

The code implementing the approach presented in this article, including the answer set programming and Boolean satisfiability encodings, is available at

http://www.cs.helsinki.fi/group/coreo/subsampled/.

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This article considerably extends a preliminary version presented at the Inter-62 national Conference on Probabilistic Graphical Models 2016 (PGM 2016) (Hyt-63 tinen et al., 2016). Most noticeably, Sections 6–9 of this article provide entirely 64 new contents, including a real-world case study (Section 6), an evaluation of 65 the impact of the choice of constraint satisfaction and optimization solvers on 66 the efficiency of the approach (Section 7), and a discussion on learning from 67 mixed frequency data (Section 8). Furthermore, new simulations on accuracy 68 and robustness (Section 5, Figures 7-9) are now included. 69

70 2. Representation

We assume that the system of interest relates a set of variables \mathbf{V}^t = 71 $\{X^t, Y^t, Z^t, \ldots\}$ defined at discrete time points $t \in \mathbb{Z}$ with continuous $(\in \mathbb{R}^n)$ 72 or discrete $(\in \mathbb{Z}^n)$ values (Entrer and Hoyer, 2010). We distinguish the repre-73 sentation of the true causal process at the system or causal timescale from the 74 time series data that are obtained at the *measurement timescale*. Following Plis 75 et al. (2015b), we assume that the true between-variable causal interactions at 76 the system timescale constitute a first-order Markov process; that is, that the 77 independence $\mathbf{V}^t \perp \mathbf{V}^{t-k} | \mathbf{V}^{t-1}$ holds for all k > 1. The parametric models for 78 these causal structures are structural vector autoregressive (SVAR) processes or 79 dynamic (discrete/continuous variable) Bayes nets. Since the system timescale 80 can be arbitrarily fast (and causal influences take time), we assume that there 81 is no "contemporaneous" causation of the form $X^t \to Y^t$ (Granger, 1988). We 82 also assume that \mathbf{V}^{t-1} contains all common causes of variables in \mathbf{V}^t . These 83 assumptions jointly express the widely used causal sufficiency assumption (see 84 Spirtes et al. (1993)) in the time series setting. In this non-parametric setting, 85 we consider surgical interventions (on the observed variables in \mathbf{V}) that keep 86 variables fixed at the selected values through the (causal timescale) time steps. 87

The system timescale causal structure can thus be represented by a causal 88 graph G^1 (as in a dynamic Bayes net) with edges only of the form $X^{t-1} \to Y^t$, 89 where X = Y is permitted (see Figure 2a for an example). Since the causal 90 process is time-invariant, the edges repeat through t. In accordance with Plis 91 et al. (2015b), for any G^1 we use a simpler, rolled graph representation, denoted 92 by \mathcal{G}^1 , where for all $X, Y: X \to Y \in \mathcal{G}^1$ iff $X^{t-1} \to Y^t \in \mathcal{G}^1$. That is, the rolled 93 graph represents time only implicitly in the edges, rather than through variable 94 duplication. Both the unrolled and rolled representations contain exactly the 95 same structural information. Figure 2b shows the rolled graph representation 96 \mathcal{G}^1 of \mathcal{G}^1 in Figure 2a. 97

Time series data are obtained from the above process at the measurement 98 timescale, defined by some (possibly unknown) integral sampling rate u. The 99 measured time series sample \mathbf{V}^t is at times $t, t - u, t - 2u, \ldots$; we are interested 100 in the case of u > 1, i.e., the case of subsampled data. A different route 101 to subsampling would use continuous-time models as the underlying system 102 timescale structure. However, some series (e.g., transactions such as salary 103 payments) are inherently discrete-time processes (Gong et al., 2015), and many 104 continuous-time systems can be approximated arbitrarily closely as discrete-105 time processes. Thus, we focus here on discrete-time causal structures as a 106 justifiable, yet simple, basis for our non-parametric inference procedure. 107

The (causal) structure of this subsampled time series can be obtained (leaving aside sampling variation) from G^1 by marginalizing the intermediate time steps. Figure 2c shows the measurement timescale structure G^2 corresponding to subsampling rate u = 2 for the system timescale causal structure in Figure 2a. Each directed edge in G^2 corresponds to a directed path of length 2 in G^1 . For arbitrary u, X, Y, the formal relationship between G^u and G^1 edges is

$$X^{t-u} \to Y^t \in G^u \quad \Leftrightarrow \quad X^{t-u} \leadsto Y^t \in G^1,$$



Figure 2: Graph (a) shows the unrolled system timescale structure, where edges repeat through time steps. Graph (b) shows the rolled representation of the same structural information. Graph (c) shows the measurement timescale structure for subsampling rate u = 2, i.e. nodes at time slice t - 1 in graph (a) are marginalized. Graph (d) depicts the rolled representation of the same structural information as in graph (c).

¹¹⁴ where \rightsquigarrow denotes a directed path.

¹¹⁵ G^u must also represent "direct" connections between variables in the same ¹¹⁶ time step (Wei, 1994). The bi-directed arrow $X^t \leftrightarrow Y^t$ in Figure 2c is an ¹¹⁷ example: X^{t-1} is an unobserved (in the data) common cause of X^t and Y^t in ¹¹⁸ G^1 (Figure 2a). Formally, the system timescale structure G^1 induces bi-directed ¹¹⁹ edges in the measurement timescale G^u as follows:

$$X^t \leftrightarrow Y^t \in G^u \quad \Leftrightarrow \quad \exists Z, l < u : (X^t \leadsto Z^{t-l} \leadsto Y^t) \in G^1, \quad \text{where } X \neq Y.$$

¹²⁰ Just as \mathcal{G}^1 represents the rolled version of G^1 , \mathcal{G}^u represents the rolled version ¹²¹ of G^u : $X \to Y \in \mathcal{G}^u$ iff $X^{t-u} \to Y^t \in G^u$ and $X \leftrightarrow Y \in \mathcal{G}^u$ iff $X^t \leftrightarrow Y^t \in G^u$. ¹²² The relationship between \mathcal{G}^1 and \mathcal{G}^u —that is, the impact of subsampling— ¹²³ can be concisely represented using only the rolled graphs:

$$X \to Y \in \mathcal{G}^u \quad \Leftrightarrow \quad X \stackrel{u}{\rightsquigarrow} Y \in \mathcal{G}^1, \tag{1}$$

$$X \leftrightarrow Y \in \mathcal{G}^u \quad \Leftrightarrow \quad \exists Z, l < u : (X \stackrel{l}{\leftrightarrow} Z \stackrel{l}{\rightsquigarrow} Y) \in \mathcal{G}^1, \quad \text{where } X \neq Y.$$
(2)

Here $\stackrel{l}{\rightsquigarrow}$ denotes a path of length l. Using the rolled graph notation, the logical encodings in Section 3 are considerably simpler. ¹²⁶ Subsampling can also be interpreted as a transitive operation applied to ¹²⁷ graphs. For example, \mathcal{G}^6 is the graph that results from subsampling \mathcal{G}^2 by a ¹²⁸ further factor of 3. More generally, $\mathcal{G}^{u\cdot k}$ can be obtained by subsampling \mathcal{G}^k by ¹²⁹ (another) u steps according to:

$$\begin{split} X &\to Y \in \mathcal{G}^{u \cdot k} \quad \Leftrightarrow \quad X \stackrel{u}{\rightsquigarrow} Y \in \mathcal{G}^{k}, \\ X &\leftrightarrow Y \in \mathcal{G}^{u \cdot k} \quad \Leftrightarrow \quad \exists Z, l < u : (X \stackrel{l}{\leftarrow} Z \stackrel{l}{\rightsquigarrow} Y) \in \mathcal{G}^{k} \quad \lor \\ & \exists Z, W, l < u : (X \stackrel{l}{\leftarrow} Z \leftrightarrow W \stackrel{l}{\rightsquigarrow} Y) \in \mathcal{G}^{k}, \quad \text{where } X \neq Y. \end{split}$$

¹³⁰ Notice that in the latter equation, the bidirected edges in \mathcal{G}^k may induce addi-¹³¹ tional bidirected edges in $\mathcal{G}^{u \cdot k}$. These equations yield Equations 1 and 2 when ¹³² k = 1, since there are no bidirected edges in \mathcal{G}^1 .

In order to obtain a correspondence between the underlying causal structure and the distribution that gives rise to the observed data at measurement timescale, we assume for a given subsampling rate u that specific conditional independences correspond to the absence of specific causal connections:

$$X^{t-u} \perp Y^t \mid \mathbf{V}^{t-u} \setminus X^{t-u} \quad \Leftrightarrow \quad X \to Y \notin \mathcal{G}^u \tag{3}$$

$$X^{t} \perp Y^{t} \mid \mathbf{V}^{t-u} \quad \Leftrightarrow \quad X \leftrightarrow Y \notin \mathcal{G}^{u} \tag{4}$$

These assumptions are analogous to the combination of the Markov and faithfulness assumptions in the standard setting of causal discovery from crosssectional data. However, here the assumptions are restricted to the particular (in)dependence relations we require to determine the causal structure, i.e., we allow, for example, for canceling pathways, which would otherwise constitute a violation of faithfulness, at subsampling rates that we do not consider.

Danks and Plis (2013) demonstrated that, in the infinite sample limit, the 143 causal structure \mathcal{G}^1 at the system timescale is in general underdetermined, even 144 when the subsampling rate u is known and small. Consequently, even when 145 ignoring estimation errors, the most we can learn is an equivalence class of causal 146 structures at the system timescale. We define \mathcal{H} to be the estimated version of 147 \mathcal{G}^{u} , a graph over V obtained or estimated at the measurement timescale (with 148 possibly unknown u). Due to underdetermination, multiple $\langle \mathcal{G}^1, u \rangle$ pairs can 149 imply \mathcal{H} , and so search is particularly challenging when u is unknown. At the 150 same time, if \mathcal{H} is estimated from data, it is possible, due to statistical errors, 151 that no \mathcal{G}^u has the same structure as \mathcal{H} . With these observations, we are ready 152 to define the computational problems focused on in this work. 153

Task 1 Given a measurement timescale structure \mathcal{H} (with possibly unknown u), infer the (equivalence class of) causal structures \mathcal{G}^1 consistent with \mathcal{H} (i.e. $\mathcal{G}^u = \mathcal{H}$ by Eqs. 1 and 2) if such a \mathcal{G}^1 exists.

¹⁵⁷ We also consider the corresponding problem when the subsampled time series ¹⁵⁸ is directly provided as input, rather than \mathcal{G}^u .

Task 2 Given a dataset of measurements of V obtained at the measurement timescale (with possibly unknown u), infer the (equivalence class of) causal structures \mathcal{G}^1 (at the system timescale) that are (optimally) consistent with the data.

¹⁶³ Section 3 provides a solution to Task 1. Section 4 provides a solution to Task 2, ¹⁶⁴ including an explanation on how \mathcal{H} can be estimated from sample data in Sec-¹⁶⁵ tion 4.2. Later sections further consider generalizations of these two basic tasks.

¹⁶⁶ 3. Finding Consistent System Timescale Structures

We first focus on Task 1. We discuss the computational complexity of the underlying decision problem, and present a practical Boolean constraint satisfaction approach that empirically scales up to significantly larger graphs than previous state-of-the-art algorithms.

171 3.1. On Computational Complexity

¹⁷² Consider the task of finding even a single \mathcal{G}^1 consistent with a given \mathcal{H} . A ¹⁷³ variant of the associated decision problem is related to the NP-complete problem ¹⁷⁴ of finding a matrix root.

Theorem 1. Deciding whether there is a \mathcal{G}^1 that is consistent with the directed edges of a given \mathcal{H} is NP-complete for any fixed $u \geq 2$.

Proof. Membership in NP follows from a guess and check: guess a candidate 177 \mathcal{G}^1 , and deterministically check whether the length-u paths of \mathcal{G}^1 correspond to 178 the edges of \mathcal{H} (Plis et al., 2015b). For NP-hardness, for any fixed $u \geq 2$, there 179 is a straightforward reduction from the NP-complete problem of determining 180 whether a Boolean B matrix² has a *u*th root (Kutz, 2004): for a given $n \times n$ 181 Boolean matrix B, interpret B as the directed edge relation of \mathcal{H} , i.e., \mathcal{H} has 182 the edge (i, j) iff $A^u(i, j) = 1$. It is then easy to see that there is a \mathcal{G}^1 that is 183 consistent with the obtained \mathcal{H} iff $B = A^u$ for some binary matrix A (i.e., a uth 184 root of B). 185

If u is unknown, then membership in NP can be established in the same 186 way by guessing both a candidate \mathcal{G}^1 and a value for u. Theorem 1 ignores 187 the possible bi-directed edges in \mathcal{H} (whose presence/absence is also harder to 188 determine reliably from practical numbers of samples; see Section 5). Knowledge 189 of the presences and absences of such edges in \mathcal{H} can restrict the set of candidate 190 \mathcal{G}^1 s. For example, in the special case where \mathcal{H} is known to not contain any 191 bi-directed edges, the possible \mathcal{G}^1 s have a fairly simple structure: in any \mathcal{G}^1 192 that is consistent with \mathcal{H} , every node has at most one successor.³ Whether this 193 knowledge can be used to prove a more fine-grained complexity result for special 194 cases is an open question. 195

 $^{^2 \}rm Multiplication$ of two values in $\{0,1\}$ is defined as the logical-or, or equivalently, the maximum operator.

³To see this, assume X has two successors, Y and Z, s.t. $Y \neq Z$ in \mathcal{G}^1 . Then \mathcal{G}^u will contain a bi-directed edge $Y \leftrightarrow Z$ for all $u \geq 2$, which contradicts the assumption that \mathcal{H} has no bi-directed edges.

196 3.2. A SAT-Based Approach

Recently, the first exact search algorithm for finding the \mathcal{G}^1 s that are con-197 sistent with a given \mathcal{H} for a known u was presented by Plis et al. (2015b); it 198 represents the current state of the art. Their approach implements a specialized 199 depth-first search procedure for the problem, with domain-specific polynomial 200 time search-space pruning techniques. As an alternative, we present here a 201 Boolean satisfiability based approach. First, we represent the problem exactly 202 using a rule-based constraint satisfaction formalism. Then, for a given input \mathcal{H} , 203 we employ an off-the-shelf Boolean constraint satisfaction solver for finding a 204 \mathcal{G}^1 that is guaranteed to be consistent with \mathcal{H} (if such \mathcal{G}^1 exists). Our approach 205 is not only simpler than the approach of Plis et al. (2015b), but as we will show, 206 it also significantly improves the current state-of-the-art in runtime efficiency 207 and scalability. 208

We present our approach using answer set programming (ASP) as the con-209 straint satisfaction formalism⁴ (Niemelä, 1999; Simons et al., 2002; Gebser et al., 210 2011). It offers an expressive declarative modeling language, in terms of first-211 order logical rules, for various types of NP-hard search and optimization prob-212 lems. To solve a problem via ASP, one first needs to develop an ASP program (in 213 terms of ASP rules/constraints) that models the problem at hand; that is, the 214 declarative rules implicitly represent the set of solutions to the problem in a pre-215 cise fashion. Then one or multiple (optimal, in case of optimization problems) 216 solutions to the original problem can be obtained by invoking an off-the-shelf 217 ASP solver, such as the state-of-the-art Clingo system (Gebser et al., 2011) 218 used in this work. The search algorithms implemented in the Clingo system 219 are extensions of state-of-the-art Boolean satisfiability and optimization tech-220 niques which can today outperform even specialized domain-specific algorithms, 221 as we show here. 222

We proceed by describing a simple ASP encoding of the problem of finding 223 a \mathcal{G}^1 that is consistent with a given \mathcal{H} . The input—the measurement timescale 224 structure \mathcal{H} —is represented as follows. The input predicate node/1 represents 225 the nodes of \mathcal{H} (and all graphs), indexed by $1 \dots n$. The presence of a di-226 rected edge $X \to Y$ between nodes X and Y is represented using the predicate 227 edgeh/2 as edgeh(X,Y). Similarly, the fact that an edge $X \to Y$ is not present 228 is represented using the predicate no_edgeh/2 as no_edgeh(X,Y). The presence 229 of a bidirected edge $X \leftrightarrow Y$ between nodes X and Y is represented using the 230 predicate confh/2 as confh(X,Y) (X < Y), and the fact that an edge $X \leftrightarrow Y$ is 231 not present is represented using the predicate $no_confh/2$ as $no_confh(X,Y)$. 232

If u is known, then it can be passed as input using u(U); alternatively, it can be defined as a single value in a given range (here set to 1,..., 5 as an example):

urange(1..5). % Define a range of u:s 1 { u(U): urange(U) } 1. % u(U) is true for only one U in the range

⁴Note the comparison to other solvers using the propositional SAT formalism in Section 7.

Here the *cardinality constraint* $1 \{ u(U) : urange(U) \} 1$ states that the predicate u is true for exactly one value U chosen from those for which urange(U) is true.

Solution \mathcal{G}^1 s are represented via the predicate edge1/2, where edge1(X,Y) is *true* iff \mathcal{G}^1 contains the edge $X \to Y$. In ASP, the set of candidate solutions (i.e., the set of all directed graphs over *n* nodes) over which the search for solutions is performed, is declared via the so-called *choice construct* within the following rule, stating that candidate solutions may contain directed edges between any pair of nodes. If we have prior knowledge about edges that must (or must not) be present in \mathcal{G}^1 , then that content can straightforwardly be encoded here.

 $\{ edge1(X,Y) \} := node(X), node(Y).$

This is a so-called *choice rule* in the ASP syntax, which here states that edge1 245 can be true or false for any pair of nodes X, Y, as given by the predicate node. 246 The implied measurement timescale structure \mathcal{G}^u for a candidate solution \mathcal{G}^1 247 is represented using the predicates edgeu/2 and confu/2, which are derived in the 248 following way. First, we declare the mapping from a given \mathcal{G}^1 to the correspond-249 ing \mathcal{G}^u by declaring the exact length-L paths in a non-deterministically chosen 250 candidate solution \mathcal{G}^1 . For this, we declare rules that compute the length-L 251 paths inductively for all $L \leq U$, using the predicate path(X,Y,L) to represent 252 that there is a length-L path from X to Y. 253

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% Derive all directed paths up to length U
path(X,Y,1) :- edge1(X,Y).
path(X,Y,L) :- path(X,Z,L-1), edge(Z,Y), L <= U, u(U).</pre>
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The first rule states that an edge $X \to Y$ implies the existence of the (corresponding) path of length one. The second rule declares inductively, that the existence of a path of length L-1 from X to Z, and an edge $Z \to Y$, together imply the existence of a path of length L from X to Y.

Second, to obtain \mathcal{G}^u , we encode Equations 1 and 2 with the following rules that form predicates edgeu and confu describing the edges \mathcal{G}^1 induces on the measurement timescale structure \mathcal{G}^u . The first rule derives induced directed edges in \mathcal{G}^u from the length-U paths, and the second the bidirected edges based on the existence of pairs of confounding paths of length up to U - 1.

% Paths of length U, correspond to measurement timescale edges edgeu(X,Y) :- path(X,Y,L), u(L). % Paths of equal length (<U) from a single node result in bi-directed edges confu(X,Y) :- path(Z,X,L), path(Z,Y,L), node(X;Y;Z), X < Y, L < U, u(U).</pre>

Finally, we declare constraints that require that the \mathcal{G}^u represented by the edgeu and confu predicates is consistent with the input \mathcal{H} . This is achieved with the following *integrity* rules, which enforce that the edge relations of \mathcal{G}^u and \mathcal{H} are exactly the same for any solution \mathcal{G}^1 . In other words, the first two rules derive a contradiction in case the directed edge relations of \mathcal{G}^u and \mathcal{H} do not match; the third and fourth rules do the same for the bidirected edge relations of \mathcal{G}^u and \mathcal{H} . For example, if the **edgeh** is true in the input for some X and Y and the corresponding **edgeu** is not derived, the set of edges defined by **edge1** does not constitute a consistent graph for the input \mathcal{H} according to the first rule below.

:- edgeh(X,Y), not edgeu(X,Y).
:- no_edgeh(X,Y), edgeu(X,Y).
:- confh(X,Y), not confu(X,Y).
:- no_confh(X,Y), confu(X,Y).

Our ASP encoding of Task 1 consists of the rules just described. The set of solutions of the encoding correspond exactly to the \mathcal{G}^1 s consistent with the input \mathcal{H} . Note that before solving, these first-order rules are grounded for all possible instantiations of X, Y, Z and L relevant to the input.

277 3.3. Runtime Comparison

Both our proposed SAT-based approach and the recent specialized search algorithm MSL of Plis et al. (2015b) are correct and complete, so we focus on differences in efficiency, using the implementation of MSL by the original authors. Our approach allows for searching simultaneously over a range of values of u, but Plis et al. (2015b) focused on the case u = 2; hence, we restrict the comparison to u = 2.

The MSL algorithm starts by noting that every measurement timescale edge corresponds to a path of length u in \mathcal{G}^1 , where that path must be through another measured variable. MSL thus creates u - 1 "virtual" mediating nodes for



Figure 3: Running times for 10-node rolled graphs as a function of graph density for the state of the art (MSL) and our method (SAT). We used 100 graphs per density and a timeout of 100 seconds; both methods enumerate up to 1000 solutions.



Figure 4: Running times as function of the number of nodes for the state of the art (MSL) and our method (SAT). Left: 10%-dense graphs. Right: 15%-dense graphs. In both plots we use 100 graphs per size and a timeout of 1 hour; both methods enumerate up to 1000 solutions.

each measurement timescale edge, and then finds all ways of identifying virtual 287 nodes with actual nodes such that all-and-only the measurement timescale edges 288 are implied. Exhaustive search of all possible virtual to actual identifications is 289 computationally intractable, so MSL employs a branch-and-bound search pro-290 cedure, where a branch is bounded whenever it implies a "false positive" (i.e., 291 implies an edge that does not actually occur in the measurement timescale in-292 put). Because each edge requires u-1 virtual nodes, each of which must later 293 be identified with an actual node, MSL scales quite poorly as a function of u. 294

For the comparison, we simulated system timescale rolled graphs with vary-295 ing density and number of nodes (see Section 5 for exact details), and then 296 computed the implied measurement timescale structures for subsampling rate 297 u = 2. This structure was given as input to the inference procedures (including 298 the subsampling rate u = 2). Note that the input consisted here of graphs for 299 which there always is a \mathcal{G}^1 , so all instances were satisfiable. The task of the 300 algorithms was to output up to 1000 (system timescale) graphs in the equiva-301 lence class. The ASP encoding was solved by Clingo using the flag -n 1000 for 302 the solver to enumerate 1000 solution graphs (or all, in cases where there were 303 fewer than 1000 solutions). 304

The running times of the MSL algorithm and our approach (SAT) on 10-305 node (rolled) input graphs with different edge densities are shown in Figure 3. 306 Figure 4 shows the scalability of the two approaches in terms of increasing num-307 ber of nodes in the rolled input graphs and fixed 10% or 15% edge density. Our 308 declarative approach clearly outperforms MSL. 10-node rolled input graphs, 309 regardless of edge density, are essentially trivial for our approach, while the per-310 formance of MSL deteriorates noticeably as the density increases. For varying 311 numbers of nodes in 10% density input graphs, our approach scales up to 65 312



Figure 5: Left: Influence of input graph density on running times of our approach when the subsamping rate u = 2 is given as input and all solutions are enumerated. Right: Scalability of our approach when u is left to be determined by the method from interval $1, \ldots, 5$. All solutions over the range of us are enumerated.

nodes with a one hour time limit; even for 70 nodes, 25 graphs finished in one 313 hour. In contrast, MSL reaches only 35 nodes; our approach uses only a few sec-314 onds for those graphs. The scalability of our algorithm allows for investigating 315 the influence of edge density for larger graphs. Figure 5 (left) plots the running 316 times of our approach (when enumerating all solutions) for u = 2 (u = 2 was 317 given as input) on 20-node input graphs of varying densities. Note that here 318 the instances are sorted by the running time for each individual density (curve). 319 With a time limit of 1000 seconds we can solve 80% of the instances with 26%320 density, almost all of the instances with 25% density and all of the instances 321 with 24% density. Thus, the running time is increased for denser graphs: in 322 addition to more constraints, there are also more members in the equivalence 323 classes. Finally, Figure 5 (right) shows the scalability of our approach in the 324 more challenging task of enumerating all solutions over the range $u = 1, \ldots, 5$ 325 simultaneously. This also demonstrates the generality of our approach: it is not 326 restricted to solving for individual values of u separately. 327

328 4. Learning System Timescale Structures from Data

³²⁹ Due to statistical errors in estimating \mathcal{H} and the sparse distribution of im-³³⁰ plied \mathcal{G}^u in the space of possible undersampled graphs, the estimated \mathcal{H} will ³³¹ often have *no* \mathcal{G}^1 s with $\mathcal{G}^u = \mathcal{H}$. Given such an \mathcal{H} , neither the MSL algorithm ³³² nor our approach in the previous section can output a solution, and they simply

conclude that no solution \mathcal{G}^1 exists for the input $\mathcal{H}^{.5}$. In terms of our constraint 333 declarations, this is witnessed by conflicts among the constraints and the under-334 lying model space for any possible solution candidate. Given the inevitability 335 of statistical errors, we should not simply conclude that no consistent \mathcal{G}^1 ex-336 ists for such an \mathcal{H} . Rather, we should aim to learn \mathcal{G}^1 s that, in light of the 337 underlying conflicts, are "optimally close" (in some well-defined sense of "opti-338 mal") to being consistent with \mathcal{H} . We now turn to this more general problem 339 setting, and propose what (to the best of our knowledge) is the first approach 340 to learning, by employing constraint optimization, from undersampled data un-341 der conflicts. In fact, we can use the ASP formulation already discussed—with 342 minor modifications—to address this problem. 343

In this more general setting, the input consists of both the estimated graph \mathcal{H} , and also (i) weights $w(e \in \mathcal{H})$ indicating the reliability of edges present in \mathcal{H} ; and (ii) weights $w(e \notin \mathcal{H})$ indicating the reliability of edges absent in \mathcal{H} . Since \mathcal{G}^{u} is \mathcal{G}^{1} subsampled by u, the task is to find a \mathcal{G}^{1} that minimizes the objective function

$$f(\mathcal{G}^1, u) = \sum_{e \in \mathcal{H}} I[e \notin \mathcal{G}^u] \cdot w(e \in \mathcal{H}) + \sum_{e \notin \mathcal{H}} I[e \in \mathcal{G}^u] \cdot w(e \notin \mathcal{H}),$$

where the indicator function I(c) = 1 if the condition c holds, and I(c) = 0349 otherwise. Thus, edges that differ between the estimated input \mathcal{H} and the 350 \mathcal{G}^u corresponding to the solution \mathcal{G}^1 are penalized by the weights representing 351 the reliability of the measurement timescale estimates. In the following, we first 352 outline how to generalize the ASP encoding from the preceding section to enable 353 search for optimal \mathcal{G}^1 with respect to this objective function. We then describe 354 two alternatives for determining the weights w. In the following section, we 355 present simulation results on the relative performance of the different weighting 356 schemes. 357

358 4.1. Learning by Constraint Optimization

To model the objective function for handling conflicts, only simple modifi-359 cations are needed to our ASP encoding: instead of declaring hard constraints 360 that require that the paths induced by \mathcal{G}^1 exactly correspond to the edges in 361 \mathcal{H} , we soften these constraints by declaring that the violation of each individual 362 constraint incurs the associated weight as penalty. In the ASP language, this 363 can be expressed by augmenting the input predicates edgeh(X, Y) with weights: 364 edgeh(X,Y,W) (and similarly for no_edgeh, confh and no_confh), and by using 365 weighted soft rules syntactically represented via : ~ instead of :-. Here the ad-366 ditional argument W represents the weight $w((X \to Y) \in \mathcal{H})$ given as input. 367 The following expresses that each conflicting presence of an edge in \mathcal{H} and \mathcal{G}^{u} is 368 penalized with the associated weight W. The additional [W, X, Y, v] for v = 1, 2369

⁵For these cases, Plis et al. (2015b) ran MSL on graphs close to \mathcal{H} to try to find an input for which there is a \mathcal{G}^1 , but this strategy is not guaranteed to find an optimal solution, nor does it scale computationally.

syntactically enforce that a cost of W is incurred in case the corresponding rule is violated for a specific pair of nodes X, Y. The numbers $v \in \{1, 2\}$ at the end of the brackets enable the solver to distinguish the cost incurred due to bidirected and directed edges respectively.

```
:~ edgeh(X,Y,W), not edgeu(X,Y). [W,X,Y,1]
:~ no_edgeh(X,Y,W), edgeu(X,Y). [W,X,Y,1]
:~ confh(X,Y,W), not confu(X,Y). [W,X,Y,2]
:~ no_confh(X,Y,W), confu(X,Y). [W,X,Y,2]
```

This modification provides an ASP encoding for Task 2; that is, the optimal solutions to this ASP encoding correspond exactly to the \mathcal{G}^1 s that minimize the objective function $f(\mathcal{G}^1, u)$ for given u and input \mathcal{H} with weighted edges.

377 4.2. Weighting Schemes

We use two different schemes for weighting the presences and absences of edges in \mathcal{H} according to their reliability. To determine the presence or absence of a specific edge $X \to Y$ in \mathcal{H} , we simply test the corresponding independence $X^{t-1} \perp Y^t \mid \mathbf{V}^{t-1} \setminus X^{t-1}$. To determine the presence/absence of an edge $X \leftrightarrow Y$ in \mathcal{H} , we test the independence: $X^t \perp Y^t \mid \mathbf{V}^{t-1}$.

383 The simplest approach is to use uniform weights for the estimated \mathcal{H} :

$$\begin{aligned} w(e \in \mathcal{H}) &= 1 \quad \forall e \in \mathcal{H}, \\ w(e \notin \mathcal{H}) &= 1 \quad \forall e \notin \mathcal{H}. \end{aligned}$$

Uniform edge weights resemble the search on the Hamming cube of \mathcal{H} that 384 Plis et al. (2015b) used to address the problem of finding \mathcal{G}^1 s when \mathcal{H} did not 385 correspond to any \mathcal{G}^{u} , though our approach is much superior computationally. 386 A more intricate approach is to use pseudo-Bayesian weights following Mar-387 garitis and Bromberg (2009); Hyttinen et al. (2014); Sonntag et al. (2015). 388 They used Bayesian model selection to obtain reliability weights for indepen-389 dence tests. Instead of a *p*-value and a binary decision, these types of tests give 390 a measurement of reliability for an independence/dependence statement as a 391 Bayesian probability. We can directly incorporate their approach of using log-392 probabilities as the reliability weights for the edges. For details, see Section 4.3 393 of Hyttinen et al. (2014). Again, we only compute weights for the independence 394 tests mentioned above in the estimation of \mathcal{H} . 395

396 5. Simulations

We use simulations to explore the accuracy and runtime efficiency of our approach in various different settings. For the simulations, system timescale structures \mathcal{G}^1 and the associated data generating models were constructed in the following way. To guarantee connectedness of the graphs, we first formed a cycle of all nodes in a random order (following Plis et al. (2015b)). We then randomly sampled additional directed edges until the required density was



Figure 6: Accuracy of the optimal solutions when subsampling rate u = 2 is given as input (200 instances and 250 samples). The x-axis shows the different prior probabilities of independence in the utilized independence test. The two left columns give the accuracy of the estimation of the measurement timescale structure \mathcal{H} . The next two columns give the accuracy of our method with the two different weighting schemes. The rightmost column shows the accuracy of the baseline estimate that does not take subsampling into account (the directed edges of \mathcal{H} are directly interpreted as the system timescale edges).

⁴⁰³ obtained. Recall that there are no bidirected edges in \mathcal{G}^1 . We used Equations 1 ⁴⁰⁴ and 2 to generate the measurement timescale structure \mathcal{G}^u for a given u. When ⁴⁰⁵ sample data were required, we used linear Gaussian structural autoregressive ⁴⁰⁶ processes (order 1) with structure \mathcal{G}^1 to generate data at the system timescale, ⁴⁰⁷ where coefficients were sampled from the two intervals $\pm [0.2, 0.8]$. We then ⁴⁰⁸ discarded intermediate samples⁶ to get the particular subsampling rate.⁷

409 5.1. Accuracy

Figure 6 shows the accuracy of the different methods in one setting: subsamling rate u = 2 (given as input), network size n = 6, average degree 3 (density 25%), N = 250 samples, and 200 datasets in total. The positive predictions correspond to presences of edges; when the method returned several solutions with equal cost, we used the mean solution accuracy to measure the output accuracy. The x-axis numbers correspond to the adjustment parameter for the statistical independence tests (prior probability of independence). The two left

⁶All sample counts refer to the number of samples after subsampling.

 $^{^7 \}tt Clingo$ only accepts integer weights; we multiplied weights by 1000 and rounded to the nearest integer.



Figure 7: Accuracy of the optimal solutions when subsampling rate u = 2 is given as input (200 instances and 500 samples). The x-axis shows the different prior probabilities of independence in the utilized independence test. The two left columns give the accuracy of the estimation of the measurement timescale structure \mathcal{H} . The third column gives the accuracy of our method with the pseudo-Bayesian weighting scheme. The rightmost column shows the accuracy of the baseline estimate that does not take subsampling into account.

columns (black and red) show the true positive rate and false positive rate of 417 the \mathcal{H} estimation (compared to the true \mathcal{G}^2), for the different types of edges, 418 using different statistical tests. Given 250 samples, we see that the structure 419 of \mathcal{G}^2 can be estimated with a good tradeoff of TPR and FPR with the mid-420 dle parameter values, but not perfectly. The presence of directed edges can be 421 estimated more accurately. More importantly, the two rightmost columns in 422 Figure 6 (green and blue) show the accuracy of the \mathcal{G}^1 estimation. Both weight-423 ing schemes produce good accuracy for the middle parameter values, although 424 there are some outliers. The pseudo-Bayesian weighting scheme ("psbayesw", 425 shown in green) still outperforms the uniform weighting scheme ("uniformw", 426 shown in blue), as it produces high TPR with low FPR for a range of thresh-427 old parameter values (especially for 0.3). Both weighting schemes are superior 428 to the "baseline" shown in magenta on the right. This baseline \mathcal{G}^1 estimate 429 is formed by the directed edges of the estimated H, and thus corresponds to 430 estimating \mathcal{G}^1 without taking subsampling into account. 431

Figure 7 shows the accuracy when u = 3 (given as input), n = 6, average degree 3 (density 25%), N = 500, and 200 datasets. The accuracy for edge presences in the measurement timescale graph \mathcal{H} is lower than for u = 2, even though we have twice the number of samples (Figure 7, black, red). The problem is that measurement timescale edges here correspond to 3-edge paths, whose



Figure 8: Accuracy of the optimal solutions when subsampling rate u = 2 is given as input (200 instances and 250 samples), some samples are obtained at the adjacent timepoints. Due to previous simulations we used the prior probability of 0.3 for all methods. In more detail, the x-axis gives the probability that the sample was obtained at the correct time t, otherwise the sample was obtained either at the previous or the next time point, splitting the remaining probability. The two left columns give the accuracy of the estimation of the measurement timescale structure \mathcal{H} . The third column gives the accuracy of our method with the pseudo-Bayesian weighting scheme. The rightmost column shows the accuracy of the baseline estimate that does not take subsamping into account.

causal effects will be smaller (on average) than 2-edge paths for a fixed interval of system timescale edge coefficients (\pm [0.2, 0.8]), and so are harder to detect. Nevertheless, the constraint optimization procedure achieves a good tradeoff between TPR and FPR for system timescale edges (Figure 7, green). Larger subsampling rates (u) require more samples for accurate \mathcal{G}^1 structure discovery, but not several orders of magnitude more data.

443 5.2. Robustness of the subsampling rate

Figure 8 shows the accuracy of this method when some of the samples are not 444 obtained at the exact time assumed by the measurement timescale. Specifially, 445 the x-axis specifies the probability with which we obtain the correct sample 446 (for the given u = 2, which is given as input); otherwise, we take either the 447 sample before or the sample after (synchronously for all variables), splitting the 448 remaining probability. The results with probability 1 equal the result in Figure 6 449 with prior probability of independence 0.3 and N = 250 samples. These values 450 were used in all runs in this plot. Unsurprisingly, as the "jitter" in the sampling 451 process increases, the results deteriorate in terms of TPR and FPR. However, at 452 least for the models and subsampling rate of u = 2 tested here, the inference is 453



Figure 9: Accuracy when the true u is unknown. Two left boxplots show accuracy of the \mathcal{H} estimate as before. The next three boxplots show the accuracy of our approach (pseudo-Bayesian weights) when, regardless of the true u, u is fixed to 2, or to 3, or left for the procedure decision, respectively. In the second from right boxplot the true u was given as input, the rightmost boxplot shows the baseline that does not take subsampling into account.

not overly sensitive. When the probability of a correct sample is 0.9, the results 454 are still quite good, alleviating somewhat the dependence on the assumption 455 of an exact subsampling rate. Naturally, there are many further permutations 456 one could explore: jitter could affect variables independently of one another, 457 jitter could be represented by a more complex distribution, we could explore 458 the effect of jitter for different subsampling rates or when the subsampling rate 459 is unknown. Moreover, jitter could have a persistent, rather than a local effect, 460 in shifting subsequent measures as well. We have here only explored the simple 461 case mimicking the situation where the measurement device as a whole (i.e. 462 simultaneously for all variables) comes out of synch with the system at random 463 points without consequences for subsequent samples. 464

Figure 9 further examines the possibility to distinguish between different 465 subsampling rates. We generated 500 samples of data from 200 models (average 466 degree 3) with equal numbers of cases with u = 2 or u = 3. The two leftmost 467 boxplots show the accuracy of the estimated \mathcal{H} , which, given the mixture of u =468 2 and u = 3, is between the accuracy of \mathcal{H} obtained in previous simulations. The 469 next two boxplots show the accuracy of the \mathcal{G}^1 estimate, when the subsampling 470 rate u for the search procedure is fixed to 2 or 3, respectively, regardless of the 471 true u. As expected, the accuracy is mediocre in this case, since the method 472 assumes the incorrect subsampling rate u in half of the runs. But when the 473 method is left to determine the correct u by itself, the accuracy improves again, 474 as shown in the boxplots second form the right (the method was run with 475 u = 2...3). In fact, the accuracy comes close to that of the rightmost boxplots, 476 where the correct u was given as input to the procedure. Thus the procedure 477



Figure 10: Scalability of our constraint optimization approach (using Clingo) for different graph sizes, numbers of samples and weighting schemes. For each setting there are 100 instances that are sorted according to the solving time on each line.

is often able to recognize the correct u. The longer tails indicate that at times the determination of u is not perfect.

480 5.3. Scalability

Finally, the running times of our approach are shown in Figure 10 with 481 different weighting schemes, network sizes (n), and numbers of samples (N). 482 The subsampling rate was again fixed to u = 2 (and given as input), and 483 average node degree was 3. Figure 10 (left) shows that the pseudo-Bayesian 484 weighting scheme allows for much faster solving: for n = 7, it finishes all runs 485 in a few seconds (black line), while the uniform weighting scheme (red line) 486 takes several minutes in the longest runs. Thus, the pseudo-Bayesian weighting 487 scheme provides the best performance in terms of both computational efficiency 488 and accuracy. The number of samples has a significant effect on the running 489 times: larger number of samples take *less* time. Runs for n = 9, N = 200 (blue 490 line) take longer than for n = 9, N = 500 (Figure 10 left, magenta vs. cyan 491 lines). Intuitively, statistical tests should be more accurate with larger number 492 of samples, resulting in fewer conflicting constraints. For N = 1000, the global 493 optimum is found here for up to 12-node graphs (Figure 10 right), though in a 494 considerable amount of time. 495

496 6. Case Study: House data of Peters et al. (2013)

In order to demonstrate the applicability to real-world data, we analyzed the house temperature and humidity data of Peters et al. (2013). The data includes 7265 samples of hourly temperature and humidity measurements of six sensors placed in a house (SHED=in the shed, OUT=outside, KIT=kitchen



Figure 11: Results of the House data analysis for different subsampling rates (u) and measurement type. Edges with full lines are found to be present, absent edges are found to be absent, edges with dotted lines may be present or absent.

boiler, LIV=living room, WC=wc, BATH=bathroom) in the Black Forest. The house has heating, but the house is not in use for most of the year. This data was also partly analyzed by Gong et al. (2015). The measurements of this system were obtained at coarser intervals than the process of temperature and humidity changes are thought to take place. Since the data includes outside temperature and humidity measurements, the assumption of causal sufficiency at the system timescale seems a good approximation.

We analyzed the temperature and humidity components separately, and examined the differences of sequential measurements,⁸ as this removed trends from each univariate time series. The temperature measurement timescale graph (obtained at 0.9 prior probability of independence) includes a total of 20 (out of directed edges, and 8 (out of 15) bidirected edges, with varying pseudo-Bayesian weights. The humidity measurement timescale graph had the same total numbers of edges, although not the exact same edges.

As explained earlier, subsampling introduces underdetermination of the system timescale graph. Thus, we determined the presence of individual system timescale edges in the following way (Magliacane et al., 2016). For each edge in

⁸This may take out some of the influences of self-loops.

 \mathcal{G}^1 , we ran the inference procedure first enforcing its presence and then enforcing its absence.⁹ The difference in objective function values for the two outputs the optimal \mathcal{G}^1 s that do or do not contain the edge, respectively—indicates the support for the presence (absence) of the edge.

For the estimated \mathcal{H} , we computed \mathcal{G}^1 s edgewise for subsampling rates of 522 u = 2, 3. (Since the measurements were hourly, these correspond to time steps 523 of 30 and 20 minutes, respectively.) The two temperature graphs for u = 2524 and u = 3 (Figure 11a,b) differ substantially from one another, as do the two 525 humidity graphs (Figure 11d,e). These results provide empirical demonstrations 526 of the impact of subsampling, as different choices of u imply different structures. 527 At the same time, timesteps of 20 and 30 minutes arguably do not correspond 528 to realistic time steps for the temperature and humidity changes measured by 529 these data. 530

We thus considered larger subsampling rates u = 10..12, which correspond to 531 more realistic time steps of 5-6 minutes. As expected, there is more underdeter-532 mination for these u, but the results are also more plausible. Figure 11c suggests 533 that the temperature outside is not directly influenced by the temperature in 534 any of the rooms, but it directly influences the temperature in the shed. The 535 data do not, however, uniquely determine how the outside temperature directly 536 affects the temperatures in the rooms inside the house, nor the system timescale 537 causal dependencies between temperatures in the rooms. The algorithm output 538 is both intuitively sensible, and also points towards future targeted experiments 539 if the remaining underdetermination is to be resolved. 540

Similarly, the humidity structures for larger u are more plausible. Figure 11f 541 suggests that the humidity level in the WC is driven by both bathroom and 542 outside humidity, which is sensible since the WC is located next to the bathroom 543 and has a window, according to Peters et al. (2013). Similarly as Peters et al. 544 (2013), we find that the shed humidity affects bathroom humidity — for both 545 analyses this may be due to an inability to distinguish the shed humidity from 546 the outside humidity (they are particularly strongly correlated). The living 547 room and kitchen boiler humidities seem to depend on each other directly, so 548 the data suggest that the rooms may be adjacent, though that information was 549 not provided by Peters et al. (2013). The algorithm thus points to testable 550 predictions about the spatial house layout, and the mechanisms for humidity 551 transfer. 552

Overall, the processes controlling the temperature and humidity have differences and similarities. Determining the placement of sensors thus seems to require data from both measurement types. More importantly for our present paper, this case study shows that this algorithm can be applied to real-world data, provide intuitively sensible outputs, and provide novel experiments and measurements that would resolve remaining underdetermination.

⁹This can be done by adding a simple clause to the input code "edge(X,Y)." to enforce the presence and ":-edge(X,Y)." to enforce the absence of $X \to Y$.

559 7. Solver Performance Comparison

560 Thus far in this article we have considered Clingo as the only solver to find solutions to a declarative constraint encoding of the computational prob-561 lems considered here. This raises the question to what extent the choice of 562 the constraint solver affects the runtime performance of our approach. While 563 the high-level ASP syntax is relatively easy to understand and modify, our ap-564 proach can also be represented via propositional logic. The benefit of using 565 propositional logic is that various SAT solvers, as well as MaxSAT solvers (as 566 the Boolean optimization generalization of SAT), can be applied directly. In 567 this section we evaluate the impact of the choice of SAT and MaxSAT solvers 568 on the runtime efficiency of our approach. 569

570 7.1. Direct Propositional SAT Encoding

A direct propositional SAT encoding for finding a system timescale causal structure \mathcal{G}^1 consistent with a measurement timescale graph \mathcal{H} for a known uis presented in Eqs. 5–12.

$$\begin{split} \dot{h}_{X,Y} & \forall X,Y \in \mathbf{V} : X \to Y \in \mathcal{H} \\ \neg \dot{h}_{X,Y} & \forall X,Y \in \mathbf{V} : X \to Y \notin \mathcal{H} \end{split}$$
(5)

$$\begin{array}{l} h_{X,Y} \\ \neg \stackrel{\leftrightarrow}{h}_{X,Y} \end{array} \qquad \forall X,Y \in \mathbf{V} : X < Y, X \leftrightarrow Y \in \mathcal{H} \left(7 \right) \\ \forall X,Y \in \mathbf{V} : X < Y, X \leftrightarrow Y \notin \mathcal{H} \left(8 \right) \end{array}$$

$$\vec{h}_{X,Y} \iff \bigvee_{Z \in \mathbf{V}} (p_{X,Z}^{u-1} \wedge p_{Z,Y}^1) \qquad \forall X, Y \in \mathbf{V}$$
(9)

$$p_{X,Y}^{l+1} \Leftrightarrow \bigvee_{Z \in \mathbf{V}} (p_{X,Z}^l \wedge p_{Z,Y}^1) \qquad \forall X, Y \in \mathbf{V}, \ l \in \{1..u-2\}$$
(10)

$$\stackrel{\leftrightarrow}{h}_{X,Y} \Leftrightarrow \bigvee_{l=1}^{u-1} \stackrel{\leftrightarrow}{h^l}_{X,Y} \qquad \forall X, Y \in \mathbf{V} : X < Y$$
(11)

$$\stackrel{\leftrightarrow}{h^{l}}_{X,Y} \Leftrightarrow \bigvee_{Z \in \mathbf{V}} (p^{l}_{Z,X} \wedge p^{l}_{Z,Y}) \quad \forall X, Y \in \mathbf{V} : X < Y, \ l \in \{1..u-1\} \ (12)$$

Essentially, Eqs. 5–8 enforce the input constraints imposed by \mathcal{H} . Following the ASP encoding presented earlier, Eqs. 9–12 encode the mapping from the \mathcal{G}^{1} 's the edge relation of which is encoded as the length-1-path variables $p_{X,Y}^1$ —that are consistent with \mathcal{H} .

578 7.2. Solver Comparison: Finding Consistent System Timescale Structures

The results of a runtime performance comparison between Clingo and two state-of-the-art SAT solvers, Glucose (Audemard and Simon, 2009) and Lingeling (Biere, 2016), is presented in Figure 12 for u = 3 (given as input), edge density of 10% and the numbers of nodes ranging from 27 (on left) to 30 (on



Figure 12: Comparison of running times for different solvers finding a single graph in the equivalence class, when the subsampling rate u = 3 is given as input. Left: easier instances with 27 nodes. Right: harder instances with 30 nodes. Clingo uses the ASP encoding presented in Section 3.2, all others use the propositional SAT encoding in Section 7.1.

right). Note that the plots give the running times of each of the three solvers 583 sorted individually for each solver. In terms of runtime performance, the SAT 584 solvers Glucose and Lingeling, both working directly on the propositional SAT 585 encoding, exhibit noticeably improved performance over Clingo as the num-586 ber of nodes is increased (right plot). Thus, in terms of runtime efficiency 587 of our approach, it can be beneficial to apply current and future advances in 588 state-of-the-art SAT solvers directly on the propositional level for improved per-589 formance. In these simulations the ASP paradigm does not show any particular 590 computational advantage. 591

⁵⁹² 7.3. Solver Comparison: Learning System Timescale Structures from Data

As with the ASP encoding given earlier, the SAT encoding given as Eqs. 5– 593 12 is easily extended to solve the optimization problem underlying the task of 594 learning system timescale structure from undersampled data. In the language of 595 MaxSAT, the only change required is to make the constraints in Eqs. 5–8 soft, 596 and to declare that the cost incurred from not satisfying these individual con-597 straints equals that of $w(e \in \mathcal{H})$ (for Eqs. 5,7) or $w(e \notin \mathcal{H})$ (for Eqs. 6,8) for the 598 corresponding edge e. This enables a comparison of the runtime performance 599 of Clingo's default branch-and-bound based search for an optimal solution to 600 those of other MaxSAT solvers implementing alternative algorithmic approaches 601 on the direct propositional MaxSAT encoding. Results comparing the perfor-602 mance of Clingo to that of the modern MaxSAT solvers Eva500a (Narodyt-603 ska and Bacchus, 2014), LMHS (Saikko et al., 2016), MSCG (Morgado et al., 604 2015), Open-WBO (Martins et al., 2014), PrimalDual (Bjørner and Narodyt-605 ska, 2015), and QMaxSAT (Koshimura et al., 2012), as well as the commercial 606



Figure 13: Comparison of running times for different solvers finding the optimal graph, when the subsampling rate u = 3 is given as input. Left: easier instances with 6 nodes. Right: harder instances with 7 nodes. Clingo uses the ASP encoding presented in Sections 3.2 and 4.1, all others use the propositional SAT encoding in Section 7.1.

integer programming (IP) solver CPLEX run on a standard IP translation of 607 MaxSAT (Davies and Bacchus, 2013; Ansótegui and Gabàs, 2013), are shown 608 in Figure 13. Here we observe that Clingo's branch-and-bound approach is 609 among the best performing solvers (with the considered problem parameters). 610 However, the results also suggest that QMaxSAT, and so-called model-based 611 approaches using a SAT solver to search for an optimal solution over the ob-612 jective function range with a top-down strategy, can improve on the runtime 613 efficiency of our approach. These results clearly show that the choice of the 614 underlying Boolean optimization solver can indeed have a noticeable influence 615 on the practical efficiency of the approach. There is at least some potential for 616 further improving the runtime performance of our approach by making use of 617 advances in MaxSAT solver technology. 618

619 8. Learning from Mixed Frequency Data

In some contexts we may have obtained data from the same system at different subsampling frequencies. Two cases can be distinguished here: First, the subsampled time series may be anchored to the same underlying process such that one may know about the offset between the two.¹⁰ For approaches to this case see Tank et al. (2016), who treat this issue as a missing data problem in a parametric setting. The second case we consider here is one where the subsampled time series are taken at different times and cannot be coordinated to

¹⁰For example, in the special case with two simultaneously measured data sets with u = 2and 1 time step offset, we can combine the time series to give a dataset with no subsampling.



Figure 14: Example graphs for learning form mixed frequency data. Graph (a) shows the true system timescale causal structure. When this is subsampled by u = 2 or by u = 3, the result is also the structure (a) (this time in measurement timescale). System timescale structure (b) gives measurement timescale structure (a) when subsampling by u = 2. System timescale structure (c) gives measurement timescale structure (a) when subsampling by u = 3. However, if measurement timescale structures for u = 2 and u = 3 are given as (a) respectively, the true system timescale structure can in fact be identified as (a).

the same instance of an underlying time series. A natural question is how much 627 more can be learned by integrating information from multiple sampling rates. 628 If one sampling rate is an integer multiple of the other, then (provably) noth-629 ing additional can be learned. A more interesting situation arises when neither 630 sampling rate is an integer multiple of the other. For example, suppose the 631 causal system operates at a 1-second timescale. If the system is measured every 632 2 seconds in one dataset, and every 3 seconds in another dataset, then we have 633 $u_1 = 2/3 \cdot u_2$. More generally, if u_1/u_2 is non-integer, then when (if ever) is the 634 equivalence class of \mathcal{G}^1 that satisfies both $\mathcal{H}_1 \& \mathcal{H}_2$ smaller than the equivalence 635 class for either \mathcal{H} individually? We can start to answer this question using the 636 constraint satisfaction approach of this paper with only minor modifications. 637

For example, suppose the true system timescale structure is given in Fig-638 ure 14a. That is, the system includes four independent time series with self 639 loops. Undersampling does not change this graph, so the measurement timescale 640 structures for u = 2 and for u = 3 will also be the graph in Figure 14a. For this 641 measurement timescale graph, the system timescale structure is not uniquely 642 determined for either u = 2 or u = 3: for example, the system timescale struc-643 ture in Figure 14b produces Figure 14a with u = 2, and Figure 14c produces 644 Figure 14a with u = 3. In fact, any system timescale edge can be present or 645 absent given either of the measurement timescale graphs alone.¹¹ However, if 646 this measurement timescale graph is found at both u = 2 and u = 3, then the 647 system timescale structure can be uniquely determined: Figure 14b produces 648 a different measurement timescale graph for u = 3 and Figure 14c produces 649 a different measurement timescale graph for u = 2. And of course, the same 650 observations hold if the us are multiplied by a constant (e.g., if u = 4 and u = 6). 651 To examine the prevalence of this phenomenon, we exhaustively considered 652

all $65536(=2^{4\cdot4})$ different 4-variable \mathcal{G}^1 s, and compared the number of equiv-

¹¹The node labels in Figure 14b and c can be permuted.

alence classes given input at a single subsampling rate, versus given inputs at
two subsampling rates. A greater number of equivalence classes means a higher
chance that a random graph will be uniquely identifiable, and so the number of
equivalence classes is an approximate (inverse) measure of the extent of underdetermination.

For input at a single undersampling rate, for u = 2 we have 24265 equivalence classes; 7544 for u = 3; and 3964 equivalence classes for u = 4. These results with a single undersampled input graph thus replicate the known result that underdetermination is a significant problem, and it rapidly worsens as uincreases (Plis et al., 2015a,b).

If we instead have measurement timescale graphs for both u = 2, 3, then 664 we have 26720 equivalence classes, which is only slightly more than the number 665 for u = 2 by itself. That is, underdetermination is not substantially reduced 666 if we additionally measure at u = 3 when we already have measurements at 667 u = 2. Similarly, for u = 3, 4 we have 7814 equivalence classes; again, there is 668 a reduction in underdetermination compared to u = 3 by itself, but it is quite 669 small. This analysis assumes that all \mathcal{G}^1 are equally likely, and it is an open 670 question whether measurements at different undersampling rates would have 671 more impact for certain classes of \mathcal{G}^1 (e.g., connected graphs). 672

673 9. Discussion

We have assumed that all common causes of measured variables are them-674 selves measured, but this assumption is frequently violated in real-world data. 675 Constraint satisfaction methods have elsewhere been used with success to iden-676 tify causal relations in the presence of unobserved common causes or latent 677 variables (Hyttinen et al., 2014; Magliacane et al., 2016). For time series data, 678 dropping the assumption of causal sufficiency (in the system timescale) generates 679 complications. Even if the system timescale process including latent variables 680 is assumed to be first order Markov, the Markov order of the measurement 681 timescale (naturally without the latent variables) can be arbitrarily larger.¹² 682 That is, variables arbitrarily far in the past can (directly, in the measurement 683 timescale) cause variables at the current timestep. We would thus need to both 684 enrich the notation for \mathcal{G}^u to encode the time lags of direct causal effects, and 685 also modify the statistical tests used to estimate these connections. 686

Moreover, there can be more information contained in the pattern of time 687 lags (i.e., which past variables directly cause the present) than is given by the 688 Markov order of the system. As just one example, suppose $\{X^{t-2}, X^{t-4}, \ldots\} \rightarrow$ 689 Y^t . The simplest (in terms of number of latents) structure that explains these 690 influences (i) has a latent L through which X influences Y (i.e., $X^{t-2} \rightarrow L^{t-1} \rightarrow$ 691 Y^t); and (ii) L is part of a 2-loop with another latent M (i.e., $L^{t-1} \to M^t$ and 692 $L^t \leftarrow M^{t-1}$). In contrast, if we have $\{X^{t-2}, X^{t-3}, \ldots\} \to Y^t$, then the simplest 693 structure has only a single latent L through which X influences Y, but where L 694

¹²This complication is independent of undersampling, and arises even if u = 1.

has a self-loop (i.e., $L^{t-1} \rightarrow L^t$). The pattern of time lags for direct causes—in particular, the absence of certain time lags—thus contains information about the number and causal structure of the latent variables. Estimation of this pattern, however, can be quite complex statistically.

Subsampled time series data can be also particularly prone to violations 699 of faithfulness. For example, the underlying process unrolled over time may 700 include directed paths over many time steps that do not result in significant 701 statistical dependence in the observed data. In addition, variables observed 702 over subsequent time steps might be almost deterministically related. If $X^{t-1} \approx$ 703 X^{t-2} , then conditioning on X^{t-2} may render the statistical dependence through 704 $Y^t \leftarrow X^{t-1} \rightarrow Z^t$ undetectable from any realistic numbers of samples. In the 705 current framework, both of these situations are treated as estimation errors 706 in \mathcal{H} . Further modeling of these complications may help to achieve improved 707 accuracy. Another option could be to develop parametric approaches instead of 708 the non-parametric one presented in this paper. 709

710 10. Conclusion

In this paper, we introduced a constraint optimization based solution for the 711 problem of learning causal timescale structures from subsampled measurement 712 timescale graphs and data. Our approach considerably improves the state-of-713 art; in the simplest case (subsampling rate u = 2), we extended the scalability 714 by several orders of magnitude. Moreover, our method generalizes to handle 715 different or unknown subsampling rates in a computationally efficient manner. 716 Unlike previous methods, our method can operate directly on finite sample in-717 put, and we presented approaches that recover, in an optimal way, from conflicts 718 arising from statistical errors. We demonstrated the accuracy, robustness and 719 scalability of the approach through a series of simulations and applied it to 720 real-world time series data. We expect that this considerably simpler approach 721 will allow for the relaxation of additional model space assumptions in the fu-722 ture. In particular, we plan to use this framework to learn the system timescale 723 causal structure from subsampled data when latent time series confound our 724 observations. 725

726 Acknowledgments

We thank the anonymous reviews for comments that improved this paper. 727 AH was supported by Academy of Finland Centre of Excellence in Computa-728 tional Inference Research COIN (grant 251170) and Academy of Finland grant 729 295673. SP was supported by NSF IIS-1318759 & NIH R01EB005846. MJ was 730 supported by COIN (grant 251170) and Academy of Finland grants 276412, 731 284591; and Research Funds of the University of Helsinki. FE was supported 732 by NSF 1564330. DD was supported by NSF IIS-1318815 & NIH U54HG008540 733 (from the National Human Genome Research Institute through funds provided 734 by the trans-NIH Big Data to Knowledge (BD2K) initiative). The content is 735

solely the responsibility of the authors and does not necessarily represent the
 official views of the National Institutes of Health.

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