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### Rational causal induction from events in time

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### Abstract

A longstanding focus in the causal learning literature has been on inferring causal relations from contingencies, where these abstract away from time by collating independent instances or by aggregating over regularly demarcated trials. In contrast, individual causal learners encounter events in their daily lives that occur in a continuous temporal flow with no such demarcation. Consequently, the process of learning causal relationships in naturalistic environments is comparatively less understood. In this paper, we lay out a rational framework that foregrounds the role of time in causal learning. We work within the Bayesian rational analysis tradition, starting by considering how causal relations induce dependence between events in continuous time and how this can be modeled by stochastic processes from the Poisson–Gamma distribution family. We derive the qualitative signatures of causal influence, and the general computations needed to infer structure from temporal patterns. We show that this rational account can parsimoniously explain the human preference for causal models that invoke shorter, more reliable and more predictable causal influences. Furthermore, we show this provides a unifying explanation for human judgments across a wide variety of tasks in reanalysis of seven experimental datasets. We anticipate the framework will help researchers better understand the many manifestations of continuous-time causal learning across human cognition and the tasks that probe it, from explicit causal structure induction settings to implicit associative or reinforcement learning settings.

Keywords: causal induction; causal inference; continuous time; learning; Bayesian models

### Rational causal induction from events in time

Time is inherent to our understanding of the world, shaping how we link the things 1 that happen around us and the actions we take. We might judge that a backfiring car startled some birds, suspect that a new food gave us indigestion, infer from a boiling kettle 3 that someone was recently in the kitchen, predict that we will be sore the day after the gym, or anticipate that a storm will follow a pink sunrise. All these inferences leverage causal models linking events in virtue of their experienced and historical temporal proximity through the lens of our intuitive causal theories. The successes of everyday cognition, as well as the successes of our scientific theories and the technologies they support, suggest that people are capable of representing entities, 9 properties, relations, events, states, and data defined in terms of time. This has been 10 recognized since the earliest attempts by philosophers to define what it means to form 11 beliefs about the external world (Hume, 1740). Time's arrow continues to be a core feature of philosophical discussion around the metaphysics of causality (Cartwright, 1994; Ross, 13 2024; Woodward, 2021), the acquisition of knowledge (Gettier, 1963; Goodman, 1983), and 14 the functioning of intentional and volitional control (Dennett, 1971; Libet, 2009). It is not surprising then, that the study of human and animal learning has grown out of basic 16 notions of association and reinforcement whereby the closeness of actions and events in 17 time governs how we come to relate them in our minds (Gallistel et al., 2019; Gallistel & 18 Gibbon, 2000; Garcia et al., 1966; Gershman, 2015; Hamou et al., 2025; Mnih et al., 2015; 19 Rescorla & Wagner, 1972; Schultz et al., 1997; Tarpy & Sawabini, 1974). 20 In recent decades, cognitive psychologists used the approach of rational analysis 21 (Anderson, 1990) to study how people learn causal structure from different kinds of 22 environmental data (Griffiths & Tenenbaum, 2009). However, accounts of human causal 23 learning have predominantly focused on inferences from contingency data. In these settings, evidence is provided helpfully "prepackaged" in the form of multiple (typically independent) trials or observations in which causal variables take different states (Allan,

1980; Anderson & Sheu, 1995; Cheng, 1997; Griffiths & Tenenbaum, 2005; Rescorla & 27 Wagner, 1972). Consequently, the causal beliefs that emerge concern the probabilistic dependence between the states of causes and the states of effects, on average, without any 29 representation of time. One common paradigm involves presenting participants with a set 30 of independent samples in which putative causes and effects are either present or absent. 31 Cover stories have been used to contextualize this as data arising from experimental research in biology (Buehner et al., 2003; Lu et al., 2008), physics (Coenen et al., 2015; 33 Lagnado & Sloman, 2004), and psychology (Rottman & Keil, 2012), since multiple independent trials are often the data that scientists collect under laboratory conditions. A minimal example of this kind of task might involve pairs of patient outcomes (e.g., sick or 36 not) under different treatment assignments (e.g., vaccinated or unvaccinated). Having seen 37 some evidence, participants are asked to judge whether or to what extent the treatment 38 causally affected the outcome (Buehner et al., 2003; Stephan et al., 2021). A 2-by-2 contingency table can capture the prevalence of different combinations of putative cause 40 and effect states (Allan, 1980; Cheng, 1997), and where this indicates dependence there is 41 evidence for some form of causal relation. Researchers have proposed a variety of approaches for drawing causal inferences from this sort of data and integrating new 43 evidence with prior expectations (see Perales & Shanks, 2007, for a review) and 44 distinguishing sharply between naturally observed and experimentally manipulated states (Lagnado & Sloman, 2002).

While these settings put timing considerations to one side, they do not eliminate
them. Researchers in causal learning (Gong & Bramley, 2024; Greville & Buehner, 2007;
Lagnado & Speekenbrink, 2010; Pacer, 2016) and associative learning (Gallistel et al., 2014;
Gallistel & Gibbon, 2000; Hamou et al., 2025) have both recognized the problem of using
"trials" as the basic unit of measurement. Fundamental questions remain as to how to
determine an appropriate time window to measure outcomes, and how to ensure the
observations are sufficiently independent to be aggregated. Without supporting knowledge

about the relevant causal mechanisms, waiting too short a time before measuring an effect may not allow the influence to propagate or become apparent (e.g., the vaccine may not have taken effect yet), while waiting too long will tend to introduce confounding factors (e.g., the infection running its course, or the patient dying from natural causes). Equally, we need to determine the timing of interventions since some time-dependent factors (e.g., age) may also mediate the relationships between variables (Gong et al., 2023; Rottman, 2016). In order to curate scenarios and aggregate data into these simple contrasts, one must already draw on sophisticated prior causal beliefs about the relevant mechanisms and their temporal properties. Without this one could not be confident that an experimental protocol truly licenses abstraction to the level of contingency data. In short, time is integral to any general account of how we induce and represent causal models of our environment.

In this paper, we present a rational framework for causal induction from time. We 65 lay out a computational-level treatment of the problem (Marr, 1982), building this up from 66 basic principles of statistical dependence between events in time to formalize a grammar for continuous-time causal theories and a calculus for generating and comparing them with data. Our framework unifies the formalisms laid out in Griffiths and Tenenbaum (2009) 69 and Pacer and Griffiths (2012, 2015) with those used in Bramley, Gerstenberg, Mayrhofer 70 et al. (2018, 2019), Gong and Bramley (2023), Gong et al. (2023) (see also Bramley, Mayrhofer et al., 2017; Gong & Bramley, 2020, 2022; Stephan et al., 2020; Valentin et al., 2020, 2022). Many of the formal elements we use here appear in one or several of these 73 papers. However, none of these papers unpack this into a general theory, nor generalize their modeling across a wide class of time-based causal inference settings. We here 75 synthesize those works and for the first time formalize a general framework, demonstrating 76 its underlying rationale, the derivation of core principles, and showcasing its broad scope 77 and fit to behavioral data in a diverse array of tasks.

We situate our analysis within the Bayesian rational analysis tradition (Anderson, 1990), as this has proven very successful in developing a rational account of atemporal

causal induction settings (Griffiths & Tenenbaum, 2005, 2009; Pearl, 2000; Rottman & Hastie, 2014). The main difference from these is that we link causal influence with dependence between events in continuous time, rather than their coincidence in independent trials (i.e., contingency). We show that this formalism anticipates and grounds the foundational principles of causal induction laid out by Hume (Hume, 1740) and enshrined in theories of associative learning (Gallistel et al., 2019; Gallistel & Gibbon, 2000; Rescorla, 1968). We show there is a natural bridge between the time-dependence and contingency level focus of established tools for causal inference (Bramley et al., 2015; Pearl, 2000).

As a rational, computational-level model, the computations involved demand
assume accurate perception, infinite memory and computational resources. While
assumptions are not aligned with the limitations of human learners they serve to describe
the normative problem that heuristics and approximations should approximate (Anderson,
1990; Griffiths, 2020; Simon, 1982). We show how the normative analysis anticipates the
ceteris paribus human preference for causal explanations that connect events via shorter,
more reliable and more predictable causal influences. Furthermore, we present the first
computational model that can provide a unifying explanation for human judgments across
seven experimental datasets from the temporal causal learning literature.

Our analysis has deep connections with theories of time, rate, and conditioning, 99 foundational to the animal learning literature (cf. Gallistel & Gibbon, 2000; Hamou et al., 100 2025). We will highlight connections with these throughout the paper, but here highlight a 101 few ways in which we think our Bayesian treatment offers a novel and uniquely general 102 perspective on these basic learning phenomena. Associative and reinforcement learning 103 (whether model free or model based) are ultimately models of behavior, while Bayesian 104 models describe the interplay of inductive biases and evidence in the formation of beliefs. 105 Since causal models are by design, use-case-agnostic models of an agent's environment 106 (Craik, 1967), it feels natural to conceptually separate the analysis of how agents form

models, from analysis of how these models guide behavior, even if the conceptual 108 distinction is often blurred in cognitive processing. An often implicit assumption of 109 associative learning models is that we can rely on local link-by-link learning to build a 110 global understanding. However this is in general a heuristic that will lead to causal 111 misattributions (Btesh et al., in press; Fernbach & Sloman, 2009). By modeling causal 112 structure induction at the rational level as selecting the globally most probable causal 113 model, we can identify and explain these mistakes and biases as consequences of 114 approximations rather than risking treating them as the right answer to the wrong 115 question (Bramley, Dayan et al., 2017; Fernbach & Sloman, 2009; Griffiths & Tenenbaum, 116 2005, 2009; Pearl, 2000). Modeling structure induction as a model selection problem also 117 helps in thinking about the imputation of hidden causes (Gershman et al., 2010; Gershman 118 et al., 2015; Valentin et al., 2020). Bayesian learning models are also effective in describing 119 setting in which people make choices across a much larger hypothesis space (Bramley, 120 Dayan et al., 2017; Bramley & Xu, 2023; Griffiths & Tenenbaum, 2009); incorporating 121 structured priors, including domain-specific causal theories and mechanistic knowledge of 122 various kinds (Lu et al., 2008; Yeung & Griffiths, 2015); being sensitive to sample size 123 (Griffiths & Tenenbaum, 2005); and providing uncertainty or confidence estimates 124 (Kolvoort et al., 2025; O'Neill et al., 2022; O'Neill et al., 2024). We here focus on 125 incorporating temporal information into the Bayesian framework, but readers may refer to 126 the extensive body of previous research comparing Bayesian models with associative and 127 reinforcement learning models more broadly (e.g., Courville et al., 2006; Fernando, 2013; 128 Griffiths & Tenenbaum, 2005, 2009; Lake et al., 2017; Perales & Shanks, 2007; Tenenbaum 129 et al., 2006) and recent work about how the languages of Bayesian models and model-based 130 reinforcement learning models could relate to one another (e.g., Eckstein & Collins, 2020; 131 Gershman, 2015, 2017; Wang, 2021). Nevertheless, the Bayesian approach is not the only 132 method, in principle, that can provide predictions for temporal causal learning tasks. Many 133 of the high-level ideas in this paper are aligned with those championed by associative

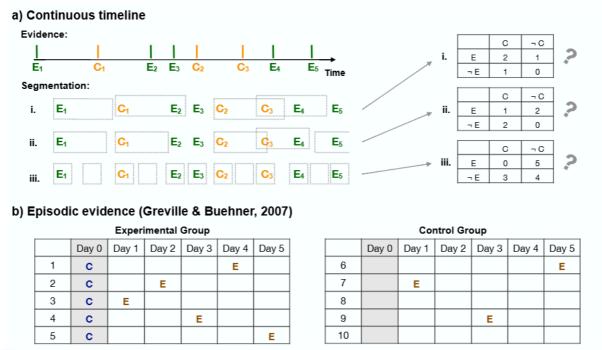


Figure 1

Continuous time evidence. a) Examples of the overly arbitrary decisions one could make when segmenting continuous timeline evidence into contingency evidence, along with the corresponding contingency tables. b) Examples of episodic evidence adapted from Greville and Buehner (2007). In the experiment, participants assessed the impact of a treatment (C) on the survival of bacterial cultures, considering culture death as the outcome (E).

learning literature, particularly Gallistel (Gallistel et al., 2014, 2019; Gallistel & Gibbon, 135 2000; Gallistel & Shahan, 2024; Gallistel & Wilkes, 2016) and Rescorla's earlier work 136 (Rescorla, 1968). Their models stem from animal associative learning paradigms, which are 137 not identical to human causal structure learning tasks. For example, all causal learning 138 studies we review in this paper ask participants to report their inner beliefs, rather than 139 analyzing beliefs indirectly through actions. As such, we mainly highlight the higher-level 140 similarities in the text while making a more detailed comparison between our Bayesian 141 account and temporal associative learning models (Gallistel et al., 2014; Gallistel & 142 Gibbon, 2000; Rescorla, 1968; Schultz et al., 1997) in the General Discussion.

### Desiderata for a Rational Theory of Causal Induction from Time

We demonstrate here four key desiderata for a rational model of causal induction from time: 1) providing predictions outside trial-based data settings; 2) capturing reasoning about dynamics and feedback; 3) grounding our core temporal-causal intuitions; 4) generalizing across a wide range of temporal causal learning tasks.

### Going beyond trials and contingencies

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We experience our environment in a single continuous timeline, making timing 150 considerations a ubiquitous aspect of inference. Data arriving in continuous time will 151 generally involve causes and effects that occur neither simultaneously, nor sufficiently 152 separated to allow for any principled segmentation into "independent, identically 153 distributed" (i.i.d.) scientific samples. This implies we must be able to litigate between 154 competing causal explanations linking multiple events, even as they occur and recur within 155 a single ongoing data-stream. For example, an everyday causal inference problem is trying 156 to identify the cause of a recurring stomach ache. As shown in Figure 1a, because any 157 decision about how to cluster and aggregate over potential trigger events and sickness 158 episodes is arbitrary, there is no unique or fully principled encoding of this continuous-time 159 data into a contingency table. An analogous problem in associative learning would be the difficulty of scoring the trials when more than one unconditioned stimulus (the effect) 161 occurs, or determining the boundary of trials when neither the conditioned stimulus nor 162 the unconditioned stimulus occurs (Gallistel, 2021; Gallistel et al., 2014). This brings home 163 that any analysis that focuses exclusively on a discretized trial structure cannot take full 164 advantage of the available metric information about the continuous time that has passed. 165 Worse, such a representation can result in different conclusions depending on one's choice 166 of measurement window. 167

The key to dealing with this problem is not to create pseudo-trials, but rather to shift the representational focus to explicitly model causal influences, in terms of how they shape the *delays* between particular causal events and, relatedly, how they shape the *rates*  at which events of different types occur (Gallistel & Gibbon, 2000).

### 172 Reasoning about dynamics and feedback

Many causal processes in the natural world are cyclic (Malthus, 1872), and people 173 frequently report causal beliefs that include feedback loops when allowed to do so in 174 experiments (Kim & Ahn, 2002; Nikolic & Lagnado, 2015; Rehder, 2017; Sloman et al., 175 1998). Cyclic systems can involve both excitatory or inhibitory feedback, which can result 176 in complex, periodic and chaotic behavior (Davis et al., 2020). For example, a cyclic 177 system might exhibit events occurring in a repeated alternating fashion: e.g., a 178 bidirectional relationship  $A \leftrightarrow B$  could generate a sequence of events A, B, A, B, A, ..., while 179 the same system plus an output component  $A \leftrightarrow B \rightarrow C$  could produce a variety of 180 temporal patterns depending on the relative delays and reliabilities of the individual 181 connections (A, B, C, A, B, C, ..., but also A, C, B, A, C, B, ..., or A, C, B, A, B, C, ...). 182 Recognizing, predicting, explaining or controlling the behavior of such cyclic causal systems 183 is only possible if one properly represents the temporal dimension. Contingency data, at 184 best, blurs this dimension and the Bayesian network formalism typically recruited for 185 causal analyses represents causal structure as inherently acyclic (DAGs; Griffiths & 186 Tenenbaum, 2009; Pearl, 2000; Rottman & Hastie, 2014).<sup>2</sup> In order to study how people 187 reason about real time causal systems, we need a framework that is able to represent these 188 dynamic and continuous features. 189

<sup>&</sup>lt;sup>1</sup> Formally, a causal mechanism is cyclic if it contains a feedback loop such that a causal variable in the system has itself as a descendant (Pearl, 2000).

<sup>&</sup>lt;sup>2</sup> Workarounds are sometimes used to model dynamic and cyclic structure with existing tools. For example, the dynamic Bayesian network "unrolls" a repeated temporal structure over equally spaced time steps (Dean & Kanazawa, 1989; Rottman & Keil, 2012; Valentin et al., 2022), where a chain graph can be used to model cyclic substructures with undirected edges in an otherwise directed causal network (Lauritzen & Richardson, 2002). However, these approaches impose significant constraints on representation. The former constraints the expression of temporal information to equally spaced discrete time points, allowing each type of event to occur only once at each time point, and implicitly modeling all causal influences as having the same latencies. Chain graphs do not represent dynamics of the causal feedback but only their equilibrium distribution. These limitations do not seem well matched to everyday causal reasoning where we may think that effects can occur at any moment, be separated by intervals of arbitrary and often variable length, and where the ability to anticipate when something will happen is likely to be important.

### Grounding our core causal intuitions

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There are many empirical findings regarding how people process temporal information to learn causal relationships. We here summarize common intuitions regarding delay information that people adopt when making causal judgments: short delays, reliable delays, and delay expectations. One of our goals in this paper is to explain these three intuitions from a rational perspective.

# 196 Intuition 1: "Contiguity" – (Relatively) shorter delays are more likely to be 197 causal

Perhaps the most foundational result in human and animal learning is that strength 198 of association between events depends on the delay between their presentations. This is the 199 contiguity effect in associative learning in human and animals, where the association 200 formed between two events decreases as the delay increases (see Gallistel et al., 2019; 201 Schultz, 2015; Tarpy & Sawabini, 1974, for reviews). Similarly, people tend to make 202 stronger causal attributions between events that occur close together than far apart, 203 especially when they don't have specific knowledge of the mechanisms involved (Buehner & 204 McGregor, 2006; Greville & Buehner, 2007, 2010, 2016; Lagnado & Sloman, 2006; Shanks 205 & Dickinson, 1991; Shanks et al., 1989).

This effect shows up when different cause candidates are studied under a shared 207 context. For example, it could be when there are competing causes in a system, people 208 tend to attribute an effect to the cause more closely preceding the effect. Lagnado and 209 Sloman (2006) found that when participants frequently observed events in the order 210 A-C-B, they were more likely to consider C to be the cause of B rather than A, even 211 though in some cases A and B co-occurred without C (see our later analysis of this 212 dataset; see also Bramley, Gerstenberg, Mayrhofer et al., 2018). It could also be that when 213 causes are learned in different trials, people give higher causal ratings in trials where they 214 observe shorter inter-event delays ("fast causes") compared to trials with longer delays 215 ("slow causes"; Buehner & May, 2003; Greville & Buehner, 2010, 2016; Shanks &

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Dickinson, 1991; Shanks et al., 1989). Although short- and long-delay causes are presented 217 in separate trials, typically some context was shared across trials making "slow causes" 218 slower in a relative as well as an absolute sense. For example, researchers used the same 219 observation duration for both short- and long-delay conditions and included the same 220 density of baserate effect events (Greville & Buehner, 2010, 2016; Shanks et al., 1989). 221 When the shared context is reduced, the contiguity effect may disappear. This has been 222 named as a time-scale invariance property in animal learning research by Gallistel and 223 colleagues (Gallistel & Gibbon, 2000; Gallistel & Shahan, 2024; Gallistel & Wilkes, 2016; 224 Kalmbach et al., 2019). For example, in Gallistel and Shahan (2024), rats learned 225 associations with delays up to 16 minutes, as long as the training was also scaled to be 226 longer. Lagnado and Speekenbrink (2010), whose findings we will model later, also found 227 that human participants drew similar conclusions about short- and long-delay causes when 228 the total observation time of a trial was scaled to match the causal delays (i.e., the observation period for long-delay trials was proportionally longer than that of short-delay 230 trials; see also Zhang & Rottman, 2024) and the baserate was matched accordingly (i.e., 231 the baserate was lower for long-delay trials; see later analysis).<sup>3</sup>

Researchers have used process-level factors to explain the short-delay intuition, such as the idea that the longer the delay, the harder it is for the cause to be sustained in working memory long enough to become associated (Ahn et al., 1995; Buehner & May, 2003; Einhorn & Hogarth, 1986). However, this explanation does not reconcile the contiguity results with the time-scale invariance results, which can instead be naturally

<sup>&</sup>lt;sup>3</sup> There are confounds in early studies to be considered when interpreting the contiguity results. For example, a free-operant procedure was often used where learners could decide when and how often to press a button to activate the cause. Participants were found to press less often when the causal delay was long (Buehner & May, 2003; Shanks & Dickinson, 1991; Shanks et al., 1989) (controlled in Greville and Buehner (2010) where a similar number of presses was found across conditions), which meant that participants tended to amass less evidence for long-delay causes. In some earlier studies, effects of later interventions were be masked if the effect of an early intervention had not yet been revealed (e.g., Shanks et al., 1989) (controlled in Buehner and May (2003) and Greville and Buehner (2010) where effects would never be masked), which would significantly impact the empirical causal strength of long-delay causes, as more ineffective interventions could have been made during a long intervention-effect interval.

reconciled within a rational Bayesian framework. In a later section, we will show that the
short-delay intuition is rational when short- and long-delay causes are learned with shared
contexts, i.e., when (1) causes compete within the same causal system; or (2) causes are
learned in different systems with the same observation duration and baserate. We will also
explain how the time-scale invariance property emerges when the shared context is
eliminated.

### 4 Intuition 2: Reliable delays are more likely to be causal

People tend to make stronger causal attributions when the delays between a 245 putative cause and effect are similar across repeated observations (Bramley, Gerstenberg, Mayrhofer et al., 2018; Gong et al., 2023; Greville & Buehner, 2010, 2016; Lagnado & 247 Speekenbrink, 2010). Greville and Buehner (2010) provides an anecdote that can serve as 248 an intuitive thought experiment: suppose you always encounter traffic lights that take a 249 very long time to change during your commute to work. You've heard a rumor that 250 flashing your car's headlamps might help because the traffic lights would respond to the 251 flashing lights of emergency vehicles. Now, suppose you try this, and the traffic lights do 252 indeed change after a consistent delay of around 10 seconds. Compare this to a situation 253 where sometimes the lights change very quickly after your headlamp flash, while at other 254 times they take much longer. In which situation would you be more likely to believe that 255 flashing the headlamps actually causes the lights to change? Greville and Buehner (2010) 256 indeed found that people give stronger causal ratings when the delays between a putative 257 cause and effect are drawn from a narrower distribution (e.g., 4.5-7.5 s), as opposed to a 258 wider distribution (e.g., 3-9 s) even when the average delay length is the same (i.e., 6 s). 259 Bramley, Gerstenberg, Mayrhofer et al. (2018) asked participants to select between two 260 causal structures based on episodic evidence with three types of event occurring in a 261 consistent orders (e.g., A - B - C) but variable temporal delays. They found that people 262 favored the "Chain" structure  $(A \to B \to C)$  when the delay between A and C was 263 variable but the delay between B and C was more reliable, and preferred the "Fork"

structure  $(B \leftarrow A \rightarrow C)$  when the delay between A and C was reliable but the delay between B and C was variable (see later analysis for this dataset).

Although a preference for reliable delays seems intuitive, it is challenging to explain 267 under associative or reinforcement learning theories. For example, Greville and Buehner (2010) demonstrated that, under the assumption of temporal-discounting reinforcement 269 learning (Chung, 1965; Myerson & Green, 1995), the expected sum of rewards for two 270 varied action-reward pairs should be greater than that for two unvaried action-reward pairs, which, counterintuitively, would lead to a preference for unreliable delays. The 272 reliable-delay intuition also cannot be explained by a simple difference in the learning rate 273 or the time required to reach the asymptote, as empirically, the preference for reliable 274 delays remains regardless of whether participants learned for 2 minutes or 4 minutes (Greville & Buehner, 2010). In contrast, we will demonstrate that our Bayesian model 276 naturally captures this intuition, as well as its stability to data exposure manipulations. 277

# 278 Intuition 3: Delays that match causal expectations are more likely to be 279 causal

People also tend to make stronger causal attributions when the delay between a 280 putative cause and effect is consistent with their causal-mechanistic understanding of the 281 situation at hand (Bramley, Gerstenberg, Mayrhofer et al., 2018; Buehner & McGregor, 282 2006; Gong & Bramley, 2023; Hagmayer & Waldmann, 2002; Stephan et al., 2020). For 283 example, Buehner and McGregor (2006) found that participants assigned higher causal 284 judgments to the insertion of a ball that turned on a light on a physical apparatus when 285 the light came on after a few seconds, rather than instantly, if they were aware that it 286 would take time for the ball to roll through the apparatus and reach the light switch (see 287 also Buehner & May, 2004). Similar results were found in 4-7-year old children (Mendelson 288 & Shultz, 1976; Schlottmann et al., 2013). Hagmayer and Waldmann (2002) found 289 participants judged whether an insecticide prevents mosquitoes by comparing prevalence of 290 mosquitoes in fields with and without the insecticide, but judged whether planting flowers 291

Table 1
Dataset features.

Name	Reference	Base Rate	Prevention	Cycle	Delay Prior		
Continuous timeline, effect specified:							
Earthquake	Lagnado and Speekenbrink (2010)	~	X	X	X		
Device: Prevention	Gong and Bramley (2023)	~	<b>V</b>	X	~		
Continuous timeline, effect unspecified:							
Device: Active Learning	Gong et al. (2023)	X	X	~	~		
Episodic evidence, effect specified:							
Bacteria	Greville and Buehner (2007)	~	<b>/</b>	X	X		
Future Bacteria	Gong and Bramley (2024)	~	<b>~</b>	X	X		
Episodic evidence, effect unspecified:							
Computer Virus	Lagnado and Sloman (2006)	X	X	~	X		
Device: Chain or Fork	Bramley, Gerstenberg, Mayrhofer et al. (2018)	X	X	X	Х		

Note: Human data are from Experiment 2 in Lagnado and Speekenbrink (2010), Experiment 1a in Gong and Bramley (2023), Experiment 1 in Gong et al. (2023), Experiment 1 in Greville and Buehner (2007), Experiment 1 in Gong and Bramley (2024), Experiment 1 in Lagnado and Sloman (2006), and Experiment 3 and 4 in Bramley, Gerstenberg, Mayrhofer et al. (2018).

prevents mosquitoes based on whether the prevalence of mosquitoes was affected the year
after the flowers were planted, presumably expecting that flowers would take longer to
influence the insect population than insecticide. We will show how this influence of
expectations fits neatly in the Bayesian rational analysis.

### Providing causal judgment predictions for various learning tasks

The final desideratum for a rational theory is that it should be able to offer 297 quantitative causal judgment predictions for a wide variety of temporal causal learning 298 tasks. Across the literature, different tasks have manipulated a wide variety of dimensions 299 from the number and nature of the causal events, how they are spaced, and what 300 participants have to infer. In these tasks, judgment patterns cannot simply be accounted 301 by one or two of the intuitions we highlight but require the complete set. One reason for 302 the variety of causal learning tasks is that temporal evidence can be accumulated in 303 different ways, depending on the context. Evidence might be collected within a single 304 extended encounter with a causal system, where all events occur within a single timeline, as depicted in Figure 1a. For instance, in Lagnado and Speekenbrink (2010), participants

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observed a geological system for several minutes, tracking the occurrence of "seismic wave" 307 events (potential causes) and earthquake events (the effect) unfolding over time. 308 Alternatively, evidence might also be gathered from multiple independent causal systems of 309 the same type, as shown in Figure 1b. For example, in the research conducted by Greville 310 and Buehner (2007), participants observed the timing of the death of multiple separate 311 bacteria culture samples (the effect) after receiving a particular treatment (the potential 312 cause). Instead of having multiple cause and effect events within a single timeline, there is 313 one cause event and one effect event each with its own timeline, with these independent 314 samples aligned by their cause or treatment time. In the rest of the paper, we refer to the 315 former situation as "continuous timeline", while the latter is termed "episodic evidence". 316

There is also a distinction based on whether the effect variables are specified. In some cases, the effect variables are specified, and participants are asked to diagnose which 318 of several candidate variables are causing them. This is similar to the traditional 319 associative learning task, in which different cues have different (positive or negative) 320 associations with the target reward, and animals learn to assigns credit accordingly (Gallistel et al., 2019; Rescorla & Wagner, 1972). In other cases, participants are tasked 322 with determining the existence of a connection between two or more variables and the 323 causal direction of these connections.

We will model seven human datasets that we categorized into four groups based on 325 the nature of the evidence (continuous or episodic) and whether the effect variables are 326 specified, as shown in Table 1. These datasets also vary in other task dimensions, 327 including: (1) base rate: whether the effect occurred without any endogenous causes; (2) 328 prevention: whether preventative causal relationships were involved in the response 329 options; (3) cycles: whether feedback loop relationships were involved in the response 330 options; and (4) delay expectations: whether participants were informed or pre-trained 331 about causal delays prior to the task. We will show that lay people's judgments 332 consistently align with the rational framework in these tasks in later modeling sections.

### A Rational Model of Causal Induction from Time

We lay out our theory in two steps. The first step demonstrates the qualitative 335 differences in temporal patterns when two variables are related versus when they are 336 unrelated to each other. The second step demonstrates how we can quantitatively infer 337 causal structures using temporal information. This logic is very similar to the theory of 338 Causal Bayesian Networks (Pearl, 2000, see below) for atemporal causal induction, which 339 specifies both how statistical independence serves as a qualitative way to determine which 340 variables are related, and how parameterized graphical models and interventional calculus 341 further assist in causal structure learning and downstream inference. To streamline the 342 descriptions, we provide equations for generative causal induction in the main text, while 343 equations incorporating preventative causation are largely relegated to Appendix A. 344

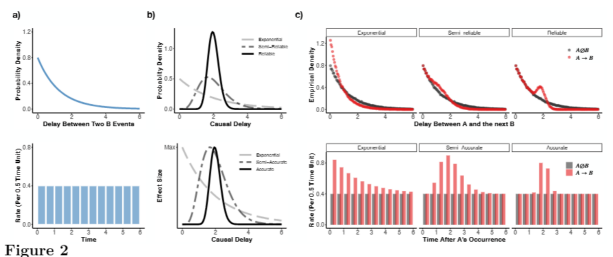
### Independence in time

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In grounding causal induction from contingencies, Pearl (2000) starts from a 346 principle of statistical independence. If A and B are perfectly independent, that is if P(B|A) = P(B), Pearl argues this is nearest we get to confidence that there is no causal 348 relation between A and B.<sup>4</sup> If A and B are dependent, that is if  $P(B|A) \neq P(B)$ , it follows 349 that there must be some causal explanation for this dependence. It could be because A 350 causes B, because B causes A, or because A and B share a (potentially distant) common 351 causal ancestor. Interventions allow one to rule between these possibilities, by statistically 352 disconnecting the intervened-on variable from its normal causes, such that if B depends on 353 the intervened-on occurrences of A, that is if  $P(B|do(A)) \neq P(B)$ , or  $P(B|do(A)) \neq P(B|do(\neg A))$  we can be confident that A is genuine causal parent or 355 ancestor of B. By combining data from a series of interventions one can thus identify 356 causal structure among a set of relata (Eberhardt et al., 2012; Steyvers et al., 2003). In 357 associative learning, Gallistel et al. (2014) has also argued that any effective cause should 358

<sup>&</sup>lt;sup>4</sup> This is known as the assumption of faithfulness (Scheines, 1997), meaning roughly the assumption that there are no additional statistically invisible causal relationships.



Causal dependence in time. a) The exponential distribution and constant rate are used to represent the base rate of B events. The same exponential distribution can be used to represent the delay between A and its next B when A and B have no relation. b) Different causal dynamics occur when the occurrence of A would generate either one extra B event (top) or a cluster of B events (bottom). c) How the data patterns differ when A causes B  $(A \rightarrow B)$  versus when A is not a cause of B  $(A \lozenge B)$ .

provide more information about the effect's occurrence compared to a random time point.

We will see later that our demonstration, especially under the special setting of an

uninformative base rate (represented by exponential distributions), echoes this point.

Similar to Pearl (2000), we ground the temporal causal induction problem by first articulating the principle of independence with respect to temporal position before identifying causality with departures from independence. However, rather than having a simple probability of occurrence P(B), we need to consider the pattern of B events occurring in time.<sup>5</sup> Without a model of its causes, a class of events might occur at subjectively unpredictable moments (e.g., receiving an email) or with some regularity or periodicity (e.g., receiving a repeat subscription delivery from Amazon). We here focus on the fully unpredictable cases for mathematical convenience, while our quantitative model is able to deal with periodic and otherwise more predictable event patterns (see our later

<sup>&</sup>lt;sup>5</sup> We restrict our focus to the problem of inferring models relating events discretized as occurring at a point in continuous time. That is, we assume the learner starts having already processed their experience into point events that they are able to locate precisely in an experienced timeline. We discuss the relationships with other representations in the General Discussion.

analysis of the dataset from Gong & Bramley, 2023). If a type of events B occur completely unpredictably, i.e., the timing of the most recent B event provides no information about the timing of the next B event, it means the *delay* between any two adjacent B events will follow an exponential distribution (see Figure 2a, top panel). Exponential distributions are "memoryless" — their expectation is constant, and so does not depend on how much time has already elapsed (Gallistel & Wilkes, 2016; Gong, 2023; Grabenhorst et al., 2019; Pishro-Nik, 2014). Under the exponential distribution, the expected *delay*  $P_d(.)$  from the present moment until the next event is always:

$$P_d(t|\lambda) = \lambda e^{-\lambda t} \tag{1}$$

The exponential distributions contain a rate parameter  $\lambda$  indicating how many B events one expects to observe on average; the delay between two adjacent B events is therefore  $\frac{1}{\lambda}$  on average.

When the occurrence of a type of events follows an exponential distribution, the

observed rate follows a Poisson process (see Figure 2a, bottom panel; Pishro-Nik, 2014). A Poisson process models the probability  $P_r(k|\lambda)$  of observing a particular quantity k of such independent events in a fixed time unit given a presumed r ate  $\lambda$ :

$$P_r(k|\lambda) = \frac{\lambda^k e^{-\lambda}}{k!} \tag{2}$$

Here,  $\lambda$  is the same parameter shaping the delays between individual events in Equation 1.
When delays are generated unpredictably at a constant rate  $\lambda$ , the Poisson process is
homogeneous. The Poisson process, along with its corresponding exponential waiting time,
has been widely used to model the arrival of events in previous cognitive research (Clarke,
1946; Gallistel & Wilkes, 2016; Grabenhorst et al., 2021; Grabenhorst et al., 2019; Griffiths

<sup>&</sup>lt;sup>6</sup> Mathematically, this means if we expect to wait an average of x minutes for an event to occur, but we have already waited for a couple of minutes and the event as not happened yet, the expected wait time is still x minutes (see p.77 in Gong, 2023, for a proof).

<sup>391</sup> & Tenenbaum, 2007). As we will see, maintaining these two representational perspectives <sup>392</sup> (delays vs. rates) is very useful since the causal influence of events can be conceptualized <sup>393</sup> as acting at either level, which can have subtle metaphysical and mechanistic consequences.

So far we have focused on the behavior of B while unperturbed by causal influences, i.e., the statistical patterns we expect to see between independent events. If A and B are 395 independent (and do not share even a distant common cause), the occurrence of A cannot 396 carry any information about the occurrence of B, so any inclusion of A in our model of B 397 is predictively impotent. Instead of having a simple contingent probability of P(B|A), we 398 here need to specify the pattern of B conditioned on the occurrence(s) of A: if A has no 399 influence on B, measuring the time from the occurrence of A until B occurs is equivalent to 400 measuring from any other arbitrary moment before the next B event. Due to the memoryless feature, the delay from As occurrence to the next B will follow the same 402 exponential distribution with the same parameter  $\lambda$  as that between one occurrence of B 403 and the next (Figure 2a, 2c, top panel). More intuitively, the rate of B occurrences after a 404 causally impotent A will remain the same as the base rate (Figure 2a, 2c, bottom panel). 405

### A qualitative understanding: Causal departures from independence

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With a definition of independence in hand, we can start to articulate departures from independence and how they reveal causal structure. More specifically, we consider how the statistical pattern would look different, if occurrences of one type of event A are able to generate occurrences of another type of event B.

We here illustrate two different generative processes: one-cause—one-effect and one-cause—many-effects (cf. Gallistel & Wilkes, 2016), using a simple example in Figure 3. Suppose a fictional substance called 5-HTP is used to treat insomnia. Consuming a 5-HTP capsule can cause a person to sleep, resulting in a one-cause—one-effect scenario. Here,

<sup>&</sup>lt;sup>7</sup> This may seem counter-intuitive: the fact that any A event inevitably occurs between two B events seems to suggest that the A-B delay would be shorter than B-B delay on average. However, to help build intuition, note that as long as A events are distributed independently from B, they are more likely to fall in a larger gap between successive B events, because these "take up more space" in the timeline.

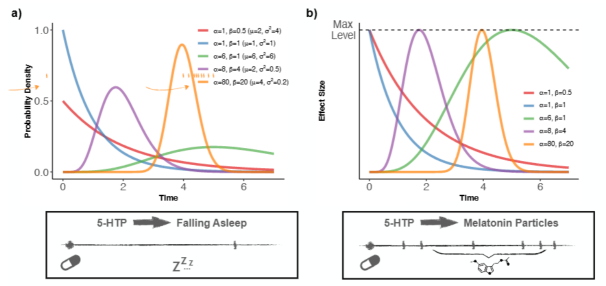


Figure 3

Examples of two types of function that could be used to model cause-effect delays and causal influences, respectively. Illustrative example relates a drug "5-HTP" and sleep. (a) A gamma probability density function capturing the delay between taking a drug and falling asleep and (b) scaled gamma density function capturing the rate of melatonin production after drug is administered. Different distributions demonstrate the functions' ability to capture various temporal dynamics. The orange distribution is the ground truth generative distribution. The orange effects in the timeline are those in fact generated by the drug while the gray effects are the base rate effects.

temporal information is embedded within the delay between the causal event of pill consumption and its effect event of falling asleep. The causal delay can vary across different mechanisms (see Figure 3a), analogous to our anticipation of certain medications (e.g., Adrenaline) taking effect rapidly and precisely, while others (e.g., painkillers) exhibit a delayed onset with some degree of variability. The gamma distribution can help us describe a variety of shapes and capture quick or slow temporal mechanisms. It is a generalization of the exponential distribution. It can be codified with a shape parameter  $\alpha$  along with the rate parameter  $\beta$ :

$$P_d(t|\alpha,\beta) = \frac{\beta^{\alpha}}{\Gamma(\alpha)} t^{\alpha-1} e^{-\beta t}$$
(3)

With  $\alpha = 1$  we recover the exponential, independent setting but with  $\alpha > 1$ , the gamma

distribution becomes increasingly peaked and increasingly normally distributed around its mean, or expected delay (Figure 3a).<sup>8</sup> The expectation  $\mu$  and the variance  $\sigma^2$  of a gamma distribution follow  $\mu = \frac{\alpha}{\beta}$  and  $\sigma^2 = \frac{\alpha}{\beta^2}$ .

It is also possible that one cause could generate many effects, for example, we might 427 think the same scenario more granularly in terms of a pill's production of Melatonin 428 particles over time (Figure 3b). One-cause-many-effect scenarios are prevalent in 429 epidemiology. For example, a single water pollution event might cause many individuals to fall ill at different points in time (Griffiths & Tenenbaum, 2007). In this case, effects are 431 generated at a rate level, which specify how many additional effect events we expect to be 432 generated by the cause per time unit, and how this rate change is, itself, spread over time. 433 This includes a functional form of the event's causal influence on the effect's rate over time, which may include an incubation period, peak, and a decay process (see Figure 3b). 435 Concretely we can also use a scaled gamma function to capture the fluctuation in the rate 436 of the effect, as a function of the time t since the cause happens and the peak rate  $\lambda_1$  of the particular causal influence:  $f(\lambda_1, t)$ . This is done by scaling the Gamma density function 438 via dividing by its mode, the density at  $(\alpha - 1)/\beta$ : 439

$$f(\lambda_1, t | \alpha, \beta) = \lambda_1 \cdot \frac{P_d(t | \alpha, \beta)}{P_d(\frac{\alpha - 1}{\beta} | \alpha, \beta)}$$
(4)

After scaling, the predicted value ranges from 0 to the peak rate  $\lambda_1$  (see Figure 3b).

<sup>&</sup>lt;sup>8</sup> To further build the intuition of what a gamma distribution models, it can be helpful to think of it as a sum of  $\alpha$  exponential delays each of rate  $\beta$ . Here the role of  $\beta$  is equivalent to the rate parameter  $\lambda$  in the exponential distribution. A classic example of an unpredictable event is a radioactive decay of the type measured by a Geiger counter. If one estimated how long it would take a Geiger counter (placed near a source of radiation) to reach a count of  $\alpha$ , the resulting delay distribution would be gamma with a shape of  $\alpha$  and a  $\beta$  reflecting the average gap between each individual event. The larger  $\alpha$ , the larger the mean but, relatively speaking, the narrower the spread of expected waiting times around that mean. Thus, a highly reliable delay is one that decomposes into a sum of many small independent unpredictable delays. This works similarly to how a normal distribution can be conceptualized as the sum of many independent errors with smaller errors producing a narrower distribution.

<sup>&</sup>lt;sup>9</sup> We assume this distribution for convenience, but in principle any function with  $(0, \infty)$  support can play this role. For instance, some causal influences might exhibit a step function, or remain at their peak level for an extended period before decaying, or be succeeded by a rebound effect. We will discuss these kinds

From a cognizer's point of view, the temporal dynamics are liable to be *variable* and uncertain. This is an inevitable feature of any model that abstracts away some of the detail, leaving unmodeled noise and complexity in the generative or measurement processes. The gamma family here helps to capture how abstract subjective probability distributions encode causal-model-based expectations about inter-event delays or event rates, and how these distributions can be shaped and sharpened with evidence, which provides a solid foundation for a rational model of time-based causal induction.

Figure 2b demonstrates three examples when the occurrence of A can result in 448 either one (top panel) or multiple (bottom panel) events of B. In the single-effect scenario, the delay between A and its effect event B could follow a memoryless exponential 450 distribution, a semi-reliable gamma distribution (with high variance), or a reliable gamma 451 distribution (with low variance). Regardless of the situation, the delay between A and the 452 next B would no longer resemble the scenario where A and B are independent, as depicted 453 in Figure 2c. Similarly, in the multiple-effect scenario, the causal influence could manifest 454 as an exponential shape (with no incubation time), a semi-accurate gamma shape (with a 455 wide spread), or an accurate gamma shape (with a narrow spread). As shown in Figure 2c, the rate of B's occurrence will deviate from the constant rate when A is a cause of B. As 457 such, the deviation in data patterns could help the learner obtain a qualitative 458 understanding of whether A causes B or not.

### A quantitative understanding: Structure inference from temporal information

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So far we have demonstrated, at a qualitative level, how we can expect the temporal pattern of events following a putative cause event A to differ depending on whether A is truly a cause of B. We introduced two causal generative processes (generating a single-effect or multiple-effects), which inspired two analysis approaches: one based on the

situation in analyzing one of the datasets (Gong & Bramley, 2023). However, we believe utilizing the gamma function as a generic basis for modeling temporal causal beliefs is a sensible default, capable of capturing many scenarios from previous experiments (Gong et al., 2023; Greville & Buehner, 2007; Lagnado & Sloman, 2006; Lagnado & Speekenbrink, 2010). In other contexts the functional form can be derived from mechanism knowledge.

delays between particular token cause and effect events (event-based scheme), and one
based on fluctuations in the rate of occurrence of the effect depending on occurrence of the
cause (rate-based scheme). We will further elaborate the similarities and differences
between these in this section. We will also show that event-based and rate-based
approaches are not necessarily tied to the generative processes that inspire them. Rather,
they are better understood as two ways of thinking.

A quantitative-level analysis is needed to formally infer causal structure. One can 471 try to directly derive some indices from the qualitative patterns. Gallistel et al. (2014) 472 pointed out the option of using the entropy between two effect distributions — measured 473 from the previous cause event vs. any random point before (see Figure 2c) — to measure 474 the associative strength. Although this seems sensible in pairwise association scenarios, it 475 is not clear it can work for structure induction in general since it will change depending on 476 the causal background, and because, unless it is an intervention, the information one 477 variable carries about another can often be due to their sharing a common cause. We here 478 follow the logic of inference to the most probable parameterized Causal Bayesian Network 479 (Griffiths & Tenenbaum, 2005, 2009; Pearl, 2000; Rottman & Hastie, 2014) to build our 480 quantitative approach, which will allow the model to both (1) infer the parameters 481 capturing how a cause generates or prevents the effect in time; (2) deal with situations 482 where multiple causes (or causes and base rates) combine with each other to influence the 483 occurrences of an effect. 484

Challenging our initial 5-HTP example, everyday continuous-time evidence is often more complicated: events can occur at any time point, and different potential causes can overlap in time leading to pervasive credit assignment questions. For instance, imagine that you are taking a pill for medical purposes but feel that you are frequently experiencing stomach discomfort afterward. You might wonder whether this discomfort is genuinely a side effect of the pill. We illustrate this evidence in Figure 4a: one might experience multiple stomach aches during the observation period both related and unrelated to the

medicine. Meanwhile, the pill may be consumed irregularly, and the stomach discomfort
events caused by a pill, if they exist, could occur even after ingestion of a subsequent pill.
Therefore, it is not possible to simply segment this kind of evidence into independent trials
to compare candidate causal models.

The causal question can be formalized as whether treating the pills as an additional 496 cause of stomach aches provides a better overall account of the evidence than treating the 497 stomach aches as happening spontaneously (i.e., due to unexamined causes). Two 498 hypothetical structures  $S_0$  and  $S_1$  are illustrated in Figure 4b. In  $S_0$ , only the base rate B499 causes the discomfort, while in  $S_1$ , both the base rate B and the pill taking C cause the 500 discomfort. However, various other factors can potentially complicate this picture. For 501 example, if the learner suspects that something else, such as their diet, may be 502 contributing to their stomach aches, there could be additional dietary events in the 503 timeline and we could incorporate their potential causal influence into the model 504 comparison. We can imagine other cases, such as if you recognizes that your pill-taking is 505 influenced by your stomach aches (e.g., if you avoid taking the pill when you already have a 506 stomach ache), if the two could have a potential common cause such as time of day, or if 507 the stomach aches could have their own feedback cycle that makes them occur with 508 regularity. All these will refine the causal structure induction problem in ways our model framework is equipped to handle. 510

How can we address the structure selection question given temporal evidence?

Three critical components have been highlighted for a rational account of causal induction

(Griffiths & Tenenbaum, 2009): (1) an ontology that outlines the entities under

investigation and their properties, (2) a set of plausible relations that suggest how entities

may be connected, and (3) the functional form that determines how causes influence their

effects under each type of relation. In the contingency setting, the ontology is a set of

variables, the set of plausible relations is a hypothesis space of causal Bayesian networks

and the functional form is often assumed to be noisy-OR combination of independent

generative or preventative influences. Despite differences in the data they operate over, temporal and atemporal causal induction share the same basic problem of articulating model selection within a hypothesis space of causal structures. The normative learner updates their prior belief over structures s in the hypothesis space S using the likelihood function  $P(\mathbf{d}|s; \mathbf{w})$  to arrive at a posterior distribution  $P(s|\mathbf{d}; \mathbf{w})$ , given data  $\mathbf{d}$  and a set of parameters  $\mathbf{w}$ :

$$P(s|\mathbf{d}) \propto \int_{\mathbf{w}} P(\mathbf{d}|s; \mathbf{w}) \cdot P(s; \mathbf{w}) d\mathbf{w}$$
 (5)

In the remainder of this section, we address the question of what constitutes an appropriate ontology, set of relations and functional form for the likelihood of causal event data in time  $P(\mathbf{d}|s; \mathbf{w})$ . Similar to the rule of qualitative patterns, we will demonstrate two approaches, with an event-based scheme that analyzes evidence at the individual delay level while a rate-based scheme analyzes the evidence at the rate level.

### An event-based scheme

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The event-based scheme we propose uses the concept of token-level "actual 531 causation" to map each event to its possible causes (Halpern, 2016), identifying which of 532 several candidate events actually caused the observed outcome (Gerstenberg et al., 2021; 533 Stephan et al., 2020). While we may have knowledge and expectations about the delay 534 between a cause and its effect (i.e., its mean and variance parameters), to derive these 535 empirically we have to also commit to a particular causal story about which cause event 536 actually produced which effect event in order to apply those expectations. Under this 537 scheme one can consider various possible causal pathways that could produce the observed 538 events, depending on the underlying causal mechanisms (Bramley, Gerstenberg, Mayrhofer 539 et al., 2018; Gong & Bramley, 2023; Hamou et al., 2025; Stephan et al., 2020; Valentin 540

<sup>&</sup>lt;sup>10</sup> Here we foreground the problem of structure selection rather than parameter estimation conditioned on a structure (Griffiths & Tenenbaum, 2005). That is, we assume that for each structure and functional form, the relevant parameters are theoretically marginalized over their prior and support if they are unknown. However the same mathematical formalism can straightforwardly be used for parameter estimation within a causal model.

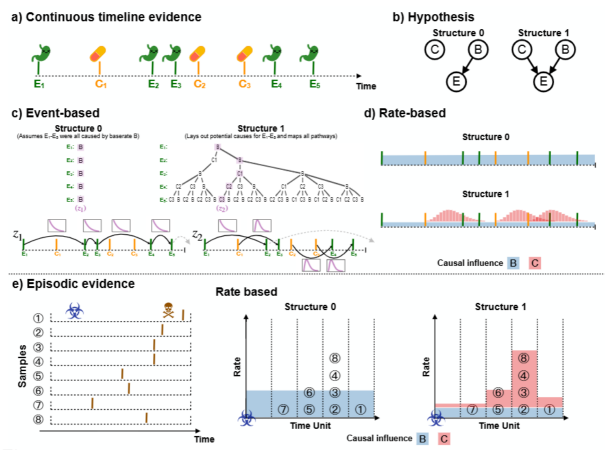


Figure 4

Causal inferences based on continuous-time causal evidence. (a) Evidence as events of stomach discomfort and pill taking unfolded in the timeline. (b) There are two causal structures in the hypothesis space. (c) The event-based scheme lays out all possible pathways (branches) that explain all effects under each hypothetical structure. (d) The rate-based scheme model in what way the rate of effects are expected to change under each hypothetical structure. (e) Episodic type of evidence where the cause and effect only happen once in each individual observation. Cases illustrate the situation in Greville and Buehner (2007) where the effect events across samples are assumed to follow exponential delays if the evaluated cause does not work. Under this situation, the evidence can be collapsed under the rate-based scheme.

et al., 2022). For example, in a causal structure s that includes an endogenous cause C, a hidden background cause B, and an effect E, each effect event could be caused by either Cor B, resulting in a pathway set  $\mathbb{Z}_s$  that contains a total of  $2^k$  possible pathways (where kis the number of effects). The event-based scheme allows for specific mechanistic constraints to be integrated into pathway construction. For instance, if we observe a sequence of events, such as  $\{C_1, E_1, E_2\}$ , and also believe that this is the kind of system within which one C event can only cause one E event (Ross, 2024), we can rule out the pathway that assumes both  $E_1$  and  $E_2$  were caused by  $C_1$ .

Given that conditional on a structural hypothesis, the potential actual causal pathways are mutually exclusive and exhaustive, it follows that the overall likelihood of each structure hypothesis is the sum of the individual likelihood of these pathways:

$$P(\mathbf{d}|s; \mathbf{w}) = \sum_{\mathbf{z} \in \mathbb{Z}_s} P(\mathbf{z}|s; \mathbf{w})$$
 (6)

To determine the likelihood of a specific actual pathway given a hypothesized type level causal structure  $P(\mathbf{z}|s; \mathbf{w})$ , we compute the likelihood of both observed effect events e and unobserved but predicted events h. For each observed effect e, we evaluate the probability that it was caused by its presumed generative cause event g (denoted  $g \to e$ ). Hidden (expected to be generated by a g but unobserved, denoted  $g \to h$ ) effects contribute to the likelihood wherever we do not observe the expected effect of a generative cause. This could be due to (1) the generative cause failing to produce the effect or (2) the effect not having occurred yet:

$$P(\mathbf{z}|s; \mathbf{w}) = \prod_{g \to e \in \mathbf{z}} \underbrace{w_g \cdot P_d(t_e - t_g | \alpha, \beta)}_{\text{Observed effects must have been generated}}$$

$$\times \prod_{g \to h \in \mathbf{z}} \underbrace{(1 - w_g) + w_g \cdot P_d(t_h > t_{end} | \alpha, \beta)}_{\text{Unobserved expected effects must have failed or be still-to-occur}}$$

$$(7)$$

In Figure 4c, the event-based scheme generates pathways for explaining stomach discomfort under different structure hypotheses. For  $S_0$ , all effect events are attributed to the base rate. For  $S_1$ , any effect event could be attributed to the base rate or to cause events that occurred previously.

### A rate-based scheme

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The rate-based scheme models causes that temporarily affect the rate of occurrence 565 of some effect. For a generative cause like 5-HTP, we expect the rate of its effect (melatonin) to temporarily increase from its base rate, and intuitively expect such rate 567 increases to be additive (unless there are also interactions between the base rate causes and 568 the focal cause). That is, an independent generative cause is something that adds extra 569 events to the timeline without affecting those that would have been there anyway (Gallistel 570 & Gibbon, 2000). For example, we might think of a large gathering causing rates of 571 infection with the Covid-19 virus to spike by contributing additional infection events. 572 The rate-based scheme employs a non-homogeneous Poisson process (Pacer & 573 Griffiths, 2012, 2015) to capture the likelihood of events in a setting where cause 574 temporarily affect the rate at which their effects occur. The likelihood depends on how the observed rates at each time bin are aligned with the expected rates: 576

$$P(\mathbf{d}|s;\mathbf{w}) = \prod_{t} P_r(d_t|f(\lambda,t))$$
(8)

We may be able to treat the base rate of an effect as a constant  $\lambda_0$  if we have no information to suggest it is periodic or structured across time. For any generative cause, 578 the causal influence can be modeled as modifying the effect's rate in a continuous fashion  $f(\lambda_1,t)$ . For example, an incubation-decay process is shown in Figure 3b, captured by the influence function in Equation 4.

Poisson processes have a desirable property known as "superposition", where the 582 union of two independent Poisson processes with rates  $\lambda$  and  $\lambda'$  is still a Poisson process with rate  $\lambda + \lambda'$ . The superposition property not only give us a simple answer to the 584 combination of a base rate and a (constant) causal influence, but also how a non-constant 585 causal influence implies a fluctuating rate. Combining a set of generative causes g with a base rate of  $\lambda_0$ , the total expected effect rate  $f(\lambda,t)$  at the time unit t can be represented

by accounting for superposition as follows:

$$f(\lambda, t) = \lambda_0 + \sum_{i \in \mathbf{g}} f(\lambda_i, t) \tag{9}$$

This could be seen as a continuous-time version of the noisy-OR logic gate used in modeling contingency data (Cheng, 1997). Prevention can be similarly captured as filtering  $\lambda_0$  resulting in a proportional temporary rate decrease that can be seen as a continuous-time version of the noisy-AND-NOT logic gate (see Appendix A). Figure 4d illustrates how the rate-based scheme models causally-induced rate changes to explain stomach discomfort. In  $S_0$ , the model assigns a constant base rate to account for the number of effect events per unit of time. In  $S_1$ , the model incorporates the

assumption that the effect rate dynamically changes following the occurrence of a cause

## $Summary\ and\ comparisons\ of\ two\ schemes$

event.

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We have introduced two schemes for thinking about causal induction in continuous 599 time. An event-based scheme involves reasoning at the level of individual events 600 adjudicating whether the delays between putative cause–effect pairs are causal ( $E_i$  was 601 actually caused by  $C_i$ ) or coincidental ( $E_i$  was actually caused by an unobserved exogenous 602 factor modeled by its base rate). The rate-based scheme involves modeling whether and 603 how the rate of occurrence of an effect changes after a cause occurs. These schemes 604 represent two different approaches to thinking about the temporal evidence, but there is a 605 continuity between the two built on a shared mathematical foundation. As we will see 606 below, there are cases where both perspectives coincide exactly or approximately in their predictions, regardless of the generative model of the evidence. Meanwhile, there are also 608 cases where they differ in their predictions or computational cost. The event-based scheme 609 can integrate detailed mechanistic knowledge, such as if cause produces exactly one effect, 610 while it often demands a costly marginalization over different causal pathways consistent

with evidence. The rate-based scheme implicitly marginalizes over these possibilities
allowing it to deal with larger event counts but as the cost of making it hard to
accommodate certain mechanistic principles.

### Human Generic Delay Intuitions: Short, Reliable, and Expected

We highlighted that people see short, predictable, and expected delays as cues to 616 causal connection. We now demonstrate that these intuitions all fall out from a Bayesian 617 analysis under our model framework. We first provide intuitive explanations for each 618 phenomenon and then back each up with simulation results. Each intuition found in 619 humans can manifest in two types of tasks: structure induction and causal diagnosis. For 620 example, the preference for the reliable delay can refer to learners' (1) higher causal ratings 621 for an evaluated causal relation in trials where the delays are more consistent, compared to 622 when they are less consistent, and (2) preference for Cause A over Cause B as an 623 explanation for the effect in a single learning trial, when the delay between A and the effect 624 is more consistent than the delay between B and the effect. This distinction matters more 625 for the short-delay intuition than the other two, as we will discuss in the intuitive 626 explanation section. In the simulation section, we primarily focus on structure induction 627 and demonstrate how the same intuition readily applies to diagnostic causal reasoning. 628

### 629 Intuitive explanations

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Delay length. In diagnostic causal reasoning (attributing the effect E to A over B 630 in the single learning trial), even in the absence of specific causal delay expectations, a 631 Bayesian learner has a preference for the diagnosis that implies shorter causal delays for at 632 least two reasons. The first reason arises from the fact that causal delays can range from 633 zero to infinity (with a limit on the lower side but not the upper side). As a result, any 634 proper delay density will be right-skewed, inherently favoring smaller values. The second 635 reason is that, when the learner don't have a specific prior for the delay variance, the 636 possible range of the delay variance becomes larger as the delay mean increases, making 637 long delays ceteris paribus harder to predict accurately. Figure 5 provides an example of

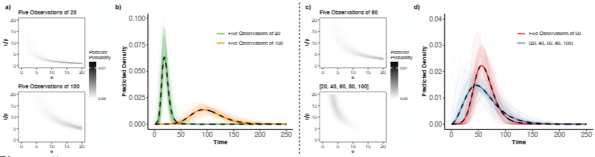


Figure 5

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Parameter posteriors and posterior predictions after observing a 20-time-unit inter-event delay five times vs. 100-time-unit inter-event delay five times (a,b) and after observing a 60-time-unit inter-event delay five times vs. 20-, 40-, 60-, 80-, 100-time-unit inter-event delays (c,d). (a and c) Parameter posteriors using the conjugate prior update process under the assumption of an initially weak prior ( $s=10^{-6}, r=10^{-6}, p=1, q=1$  for the distribution conjugate to the gamma likelihood with an intractable normalization constant; Fink, 1997). (b and d) The predicted density under sampled parameter combinations. Faint colored lines represent densities under posterior  $\alpha$  and  $\beta$  samples. Thick and dashed black lines represent the marginal or posterior predictive distribution for future inter-event delays.

the posterior on parameters  $\alpha$  and  $\beta$  (Figure 5a) and posterior predictive distribution after observing 20-time-unit delays five times vs. 100-time-unit delays five times (Figure 5b).<sup>11</sup> This illustrates that the posteriors are tighter after observing delays of 20 vs. 100. The delay prediction (Figure 5b) is also sharper (more confident) after observing delays of 20, in contrast to that following delays of 100. It highlights that a causal model that connects a set of events via shorter causal delays will assign higher likelihood to data than an alternative causal model that connects those events with longer causal delays, even when those delays are perfectly reliable (Greville & Buehner, 2010).

In structure induction (inferring the existence of short-delay or long-delay causes across different trials), the same preference appears because this comes down to a competition between the putative cause and the base rate to explain the effect's occurrences. The likelihood of the data under the baserate-only hypothesis ( $S_0$  in Figure 4b) remains constant, while the likelihood the causal variable explains the occurrence depends on the length of the delays it implies, following the same logic

<sup>&</sup>lt;sup>11</sup> We here used the analytical conjugate prior update process. In the rest of the paper, we will use simple Monte-Carlo sampling since analytic methods are not feasible.

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mentioned in diagnostic causal reasoning. The posterior distribution between causal and
baserate-only hypotheses, will differ accordingly. It is important to note though that if the
base rate and causal delays are scaled up equally, and the prior on causal delays is
uninformative, we will then have timescale invariance in the sense that a rational causal
learner will favor the causal hypothesis equally in the "slow" and the "fast" dataset. A
straightforward way to demonstrate this is to generate long-delay stimuli by scaling
short-delay stimuli by a constant (Gallistel & Shahan, 2024).

Delay reliability. The reliable-delay intuition can simply be explained by the

likelihood calculation. If observed delays exhibit great variability, the resulting gamma

distributions spread their expectations wider, resulting in a lower marginal likelihood

compared to less variable delays. As in the delay length section, we provide an example in

Figure 5 showing the parameter posteriors (Figure 5c) and delay predictions (Figure 5d)

after observing a consistent 60-time-unit delay five times, versus observing a set of varied

delays with a same average on 60 time units.

Delay variance would naturally be scaled with the delay length in a time-scale invariant environment. Otherwise, if the researchers force the long delays (e.g., 6 s) and short delays (e.g., 3 s) to have the same absolute standard deviation (e.g., 0.1 s), the relative variance will be smaller for the long delay, providing an advantage in the long-delay condition. We will further demonstrate this in the simulation section and when analyzing Lagnado and Speekenbrink (2010).

Delay expectation. The expected-delay principle can be understood as the influence of mechanistic knowledge or prior experience on people's *prior* distribution regarding a causal delay. For instance, if individuals strongly believe that a genuine switch should take approximately 4 seconds to turn on a device, a switch that takes 2 or 6 seconds would have a lower prior probability and consequently a less good explanation for an effect

<sup>&</sup>lt;sup>12</sup> However, a fully uniform prior over delays, e.g.,  $\exp(\frac{1}{\infty})$ , is improper because the range of possible delays is infinite meaning the prior has zero density everywhere, necessitating use of Markov Chain methods or weakly informative priors in practice.

Table 2
Symbols used and their meanings under three contexts.

	Synthetic data	Event-based scheme	Rate-based scheme
$m_u$	Mean of causal delays.	_	-
$i_u$	Half interval of causal delays.	_	_
$k_b$	Number of base rate effect events.	_	_
$k_c$	Number of cause events.	_	_
$w_c$	Cause's success probability.	Cause's success probability.	_
$\mu$	_	Mean of causal delays.	Mean of causal influence function.
$\sigma^2$	_	Variance of causal delays.	Variance of causal influence function.
$\mu_b$	_	Mean of base rate delays.	_
$\sigma_b^2$	_	Variance of base rate delays.	_
$\lambda_0$	_	_	Effect's base rate.
$\lambda_1$	_	_	Max generative causal influence.
ξ	_	_	Max preventative causal influence.

Note:  $\sigma_b^2$  is only used when modeling Gong and Bramley (2023) which included periodic base rates. In other cases, the base rate delay was modeled using exponential distributions, which only included one parameter.

compared to a switch that was pressed four second earlier. Correspondingly, a device activation 2 or 6 seconds after a switch press would be more likely to be caused by its base rate than a device activation 4 seconds after a switch press (Buehner & McGregor, 2006).

### 681 Simulation

To demonstrate that our account exhibits these features, we now simulate and model synthetic data. Given that most of the human empirical evidence was based on the one-cause—one-effect context, we focus on that context here. Note that the rate-based scheme generically assumes that a cause can produce multiple effect events and hence is slightly inconsistent with our simulation setting. Nevertheless we show that it still demonstrates a sensitivity to the change of delay lengths and delay variance. The meanings of symbols used in simulation and later dataset modeling are summarized in Table 2.

To show that this framework can handle data that are not exclusively generated

To show that this framework can handle data that are not exclusively generated from gamma distributions (just like humans, Greville & Buehner, 2010), we generate synthetic delay stimuli using uniform distributions, denoted as U(l, u), with a lower bound l and an upper bound l. We used the following procedure:

- Define each synthetic stimulus as lasting for 300 time units.
- 2. Generate  $k_b$  base rate effect events and place them at random times sampled from U(0,300).
- 3. Generate  $k_c$  cause events, again placing these on the timeline by sampling from U(0,300).
- 4. Generate effect events given the assumption that each cause event has a probability  $w_c$  of producing one effect event E with a delay sampled from  $U(m_u i_u, m_u + i_u)$ . In
  other words, the ground truth for the simulated data is always the structure that
  includes the causal link as well as the base rate.
- We then estimate the probability that there is a generative causal influence from C to E, i.e., judging the posterior probability of  $S_1$  over  $S_0$  in Figure 4b. The evidence that data d provide in favor of  $S_1$  over  $S_0$  can measured by the log-likelihood ratio:

$$\log \frac{P(\mathbf{d}|S_1; \mathbf{w})}{P(\mathbf{d}|S_0; \mathbf{w})} \tag{10}$$

We assume that all parameters used for simulating data are unknown to the learner 705 and hence the models marginalize over the parameters w. We achieve this via simple 706 Monte Carlo integration drawing m = 10,000 prior samples for all parameters. For the 707 event-based scheme, we assume a uniform prior on causal strength, meaning that cause succeeds in producing its effect with a probability  $w_c \sim U(0,1)$ . We also assume weak 709 priors on the parameters of the gamma causal delays between Cs and their effects E, such 710 that they have mean  $\mu \sim U(0,300)$  and variance  $\sigma^2 \sim U(0,\mu^2)$ . This means the prior on the mean causal delay is equally likely to be anything between zero and the full length of 712 the episode while the delay shape parameter  $(\alpha = \frac{\mu}{\sigma^2})$  can be anything between 1 and  $\infty$ .<sup>13</sup> 713

<sup>&</sup>lt;sup>13</sup> An alternative approach is to sample the mean  $(\mu)$  and shape  $(\alpha)$  parameters from very flat exponential distribution (Bramley, Gerstenberg, Mayrhofer et al., 2018). This produces the same qualitative results.

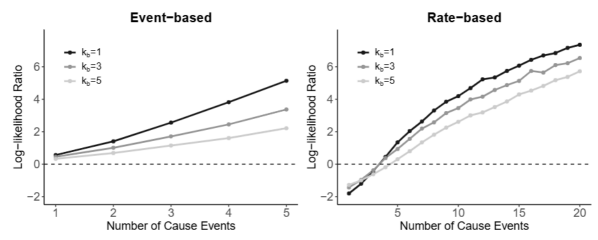


Figure 6 How log-likelihood ratio changes with the amount of cause events.

Similarly, the model assumes delays between base rate events follow an exponential 714 distribution with mean  $\mu_b \sim U(0, 300)$ .

For the rate-based approach, we specify priors on the base rate  $\lambda_0 \sim 1/U(0,300)$ , 716 the max causal influence  $\lambda_1 \sim U(0,1)$ . We assume causal influences dynamically inflate the rate according to a scaled gamma distribution with priors on the mean  $\mu \sim U(0,300)$  and the variance  $\sigma^2 \sim U(0, \mu^2)$ . 719

#### Synthetic data 720

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We first implemented a sanity check task to make sure that the model's preference 721 for  $S_1$  over  $S_0$  (1) increases as the number of cause events in the synthetic data increases 722 and (2) decreases as the base rate of effect events increases. Both of these properties are 723 based on the principle of distinguishing the (causal) signal from the noise and have been 724 demonstrated in the atemporal causal learning setting (Cheng, 1997; Griffiths & 725 Tenenbaum, 2005; Wu & Cheng, 1999). We here use  $w_c = 1$ ,  $m_u = 15$ ,  $i_u = 5$ , and consider 726 different numbers for baserate events  $k_b$  ( $k_b = \{1, 3, 5\}$ ). For the rate-based model, we 727 consider numbers for cause events  $k_c$  ranging from 1 to 20 (with a step of 1), while for the 728 event-based model, we limit the range to 1 to 5 due to computational cost. Results are shown in Figure 6. In both cases, the log-likelihood ratio in favor of  $S_1$  increases as the 730

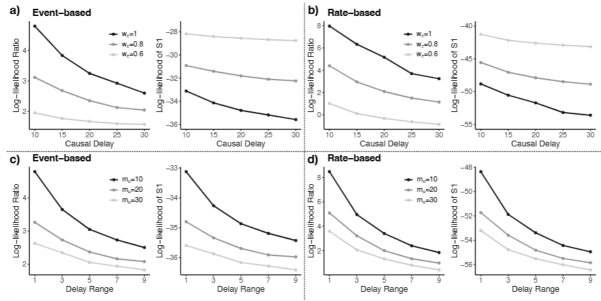


Figure 7
How log-likelihood ratio changes with the causal delays and delay ranges (and the corresponding log-likelihood of  $S_1$ ). A ratio above zero indicates that the model favors  $S_1$  (the causal structure) over  $S_0$  (the base rate structure).

number of cause events increases. Both models perform better when the base rate is low.

This reflects that the model's basic ability to learn causal structure from temporal data.

Compared to the event-based model, the rate-based model requires more cause events to favor  $S_1$  over  $S_0$ , indicating a higher requirement for data points under the relaxed constraints assumed by the model (it does not require specifying how many effect events would be generated by one cause event). Accordingly, we will use  $k_c = 5$  for the event-based model and  $k_c = 10$  for the rate-based model in the later simulations.

#### Delay length

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To illustrate the short-delay preference found in humans, we simulate stimuli arranged in a grid with  $w_c = \{0.6, 0.8, 1\}$ ,  $m_u = \{10, 15, 20, 25, 30\}$ ,  $i_u = 1$ , and  $k_b = 3$ .

Note that a fixed baserate events  $k_b$  over a fixed observation duration here is key to inducing this preference. Figure 7a demonstrates that the event-based model's preference for  $S_1$  over  $S_0$  diminishes as the duration of the true causal delays increase. This observation supports the notion that causal attribution is stronger when the delay is shorter

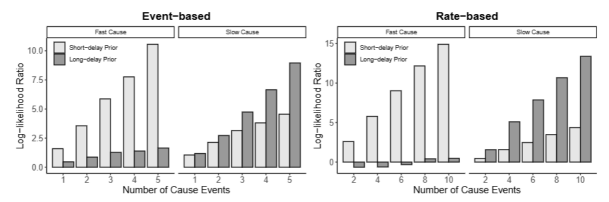


Figure 8
How log-likelihood ratio changes under different delay prior.

(i.e., structure induction). Additionally, the log-likelihood of  $S_1$  itself decreases as the delay length increases. This indicates that when faced with multiple potential cause candidates (i.e., diagnostic causal reasoning), the learner should tend to attribute the effect to the cause with the shortest delay. Similar patterns are replicated under the rate-based scheme (Figure 7b). We show in Appendix B how the time-scale invariance property would appear instead if we adapt the baserate and delay variance proportionally to the delay length.

## 51 Delay variance

To investigate the predictable-delay preference, we simulate stimuli arranged in a grid with  $m_u = \{10, 20, 30\}$ ,  $i_u = \{1, 3, 5, 7, 9\}$ ,  $w_c = 1$ ,  $k_b = 3$ . As shown in Figure 7c, the event-based model's preference for  $S_1$  over  $S_0$  diminishes as the range of delays increases. It reflects that causal attribution is stronger when the delays are reliable. Additionally, the log-likelihood of  $S_1$  decreases as the delay range expands, which suggests that when faced with multiple potential cause candidates, the learner should tend to attribute the effect to the cause with the most consistent or reliable delays. Similar results are observed in the rate-based scheme (Figure 7d). In Appendix C, we show how our model replicates the finding by Greville and Buehner (2010) that the effect of variance persists as the learning duration increases.

### Delay expectation

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natural advantage of shorter delays.

To investigate the influence of prior beliefs on causal judgments, we introduced two 763 different delay prior conditions instead of using the above uniform delay priors. For the 764 "short-delay prior", we set  $\mu$  to be sampled from a Gamma distribution with a mean of 10 765 and a standard deviation of 1, resulting in an assumed delay expectation of  $10\pm1$ . 766 Conversely, for the "long-delay prior", we assume  $\mu$  is sampled from a Gamma distribution 767 with a mean of 20 and a standard deviation of 1, representing an assumed delay 768 expectation of around  $20\pm1$ . Other model parameterizations remain the same as before. 769 For the synthetic data, we constructed scenarios in which a slow cause always 770 produced an effect with a delay sampled from  $m_u = 20$ ,  $i_u = 1$ , while a fast cause always 771 produced an effect with a delay sampled from  $m_u = 10$ ,  $i_u = 1$ . We set  $w_c = 1$  and  $k_b = 3$ 772 when simulating the data. Figure 8 demonstrates that in the fast-cause scenarios, the 773 preference for  $S_1$  in both event-based and rate-based models is stronger when one starts 774 with a short-delay prior, while in the slow-cause scenarios, the preference for  $S_1$  is stronger 775 when one starts with a long-delay prior. This confirms that the model will learn causal 776

# Human Performance in Learning from Continuous-time Evidence

strong as the tendency to favor the fast cause under the short-delay prior, highlighting the

relations more quickly when their time course aligns with expectations. It is worth noting

that the models' tendency to favor the slow cause under the long-delay prior is not as

We now reanalyze seven previous datasets that contain human performance in a
variety of temporal causal learning tasks, as shown in Table 1 (the corresponding
hypothesis spaces of causal structures are summarized in Figure 9). We will demonstrate
that our framework can accommodate all the variations across the scenarios probed across
these tasks. Furthermore, we will demonstrate a robust alignment with participants'
judgments and the predictions of the framework across these scenarios.

It is worth noting that all the datasets we model contain "interventions" — most of

them provided as pre-set interventions, while Gong et al. (2023) allowed participants to 789 intervene actively at moments of their choosing. The contingency-based learning literature 790 emphasizes the differences between intervention and covariation-only data (Bramley et al., 791 2015; Lagnado & Sloman, 2002; Lagnado et al., 2007; Pearl, 2000; Sloman & Lagnado, 792 2005; Waldmann & Hagmayer, 2005). In the contingency setting, interventions act as 793 "graph surgery" making observationally equivalent (aka. Markov equivalent structures) distinguishable. For example, casual structures such as  $X \to Y \to Z, X \leftarrow Y \leftarrow Z$ , and 795  $X \leftarrow Y \rightarrow Z$  have the same co-variation patterns (X and Y are always correlated, as are Y 796 and Z, and X and Z are unconditionally correlated but become uncorrelated once Y is controlled for), under different parameterizations all three models are equally good 798 accounts of contingency data with these independencies. People are able to select 799 interventions that allow them to distinguish these structures in causal learning experiments 800 (Bramley et al., 2015; Coenen et al., 2015). However, event timing can break the Markov 801 equivalence deadlock in some cases (Lagnado et al., 2007). For example, the temporal 802 pattern X - Y - Z is privileged under the first structure, Z - Y - X in the second, and 803 Y-X-Z or Y-Z-X in the third structure (even though high base rates or 804 preventative connections may complicate the pattern). Delay information will ultimately 805 tend to support whatever causal hypothesis most parsimoniously links the events, assigning 806 them the highest joint likelihood by implying the shorter, more reliable and expected causal 807 connections. However, whether the observationally most likely model is truly causally 808 correct, i.e., that the statistical pattern is not due to some unobserved or latent variable 809 (cf. Valentin et al., 2022), can be confirmed definitively through the use of interventions. 810

## Continuous, effect specified

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Lagnado and Speekenbrink (2010). Our first case study revisits the

"earthquake" experiment conducted by Lagnado and Speekenbrink (2010). Participants

were asked to investigate the effects of three types of seismic waves (red, yellow, and green)

on the occurrence of earthquakes. Unbeknownst to them, only one of the three types of

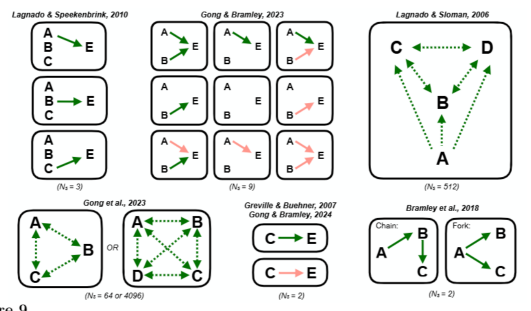


Figure 9
The hypothesis spaces used in different studies. Green arrows represent generative links and pink arrows represent preventative links. Dashed arrows  $A \to B$  represent two possibilities between two variables A and B: unconnected or  $A \to B$ , and dashed bidirectional arrows represent four possibilities between two variables A and B: unconnected,  $A \to B$ ,  $B \to A$ , or  $A \leftrightarrow B$ . Exogenous links (base rate) are ignored in all graphs.

waves (referred to as the cause) actually made earthquakes occur, while the other two 816 types (referred to as lures) had no effect on the earthquakes. In each trial, the cause wave 817 occurred 10 times and had a probability of 80% of resulting in an earthquake. Two other 818 lure causes also occurred 10 times each but had no effect on the earthquake. Two factors 819 were manipulated across trials within subjects: the delay length — the time between a 820 cause event and its effect event — could be either short  $(3\pm0.1 \text{ s})$  or long  $(6\pm0.1 \text{ s})$ ; the 821 probability of intervening events — how often the two other lure causes occurred between a 822 real cause and its effect — could be either high (65%) or low (35%). Four additional 823 earthquake events were sampled at random time points to serve as the base rate. The trials 824 lasted for an average duration of  $169\pm84$  s for the short-delay condition and  $318\pm157$  s for 825 the long-delay condition. Since the lures occur in the interval between the cause and effect, 826 a learner might mistake the lure for the true cause if they do not pay enough attention to 827 temporal information. Lagnado and Speekenbrink (2010) showed participants' ability to

figure out the genuine cause: they assigned higher ratings to the genuine cause compared to the two lures. Meanwhile, the judgments were influenced by the probability of intervening events rather than the mere length of delay (see Figure 10a).

There was a mixed set of factors predicting whether a short-delay intuition should 832 emerge. This study matched the base rate effect events (the long-delay condition had a 833 lower base rate due to a fixed number of background effects over a longer observation 834 period). However, the within-subject design provides a shared context liable to undermine 835 the invariance in the minds of participants. Moreover, the long-cause and short-cause 836 stimuli were created independently, rather than being directly scaled from one another. 837 Notably, the standard deviations for the two conditions were identical (0.1 s), meaning the 838 long-delay condition had a smaller relative variance than the short-delay condition, making 839 the causal relationships easier to identify from a normative perspective. As such, a formal 840 modeling procedure is necessary to determine the rationally predicted direction of the 841 delay effect in this task. 842

Participants were asked to provide both "absolute" and "comparative" ratings for 843 the causal properties of each wave. The "absolute" rating allowed participants to 844 independently rate each wave, while the "comparative" rating required participants to 845 allocate ratings for the three waves such that their ratings summed up to 100. Both types 846 of ratings revealed the same pattern with only a main effect of the probability of 847 intervening events but not the delay length (see Figure 10a). Here we model the 848 "comparative" rating, which can be interpreted as a comparison of the posteriors 849 associated with three causal structures as shown in Figure 9 (assuming that they each have 850 an equal prior). None of the delay or other parameters were explicitly disclosed to the 851 participants in the instructions, and the task included abstract visualizations and 852 gamification features, which might suggest to participants that it was more like a game 853 than a real-world situation. Therefore, we here do not speculate the prior knowledge 854 participants may use. We use the prior for parameters as follows:  $w_c \sim U(0,1)$  for the

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cause probability (and  $\lambda_1 \sim U(0,1)$  for the rate-based model),  $\mu_b \sim U(0,100)$  for the base 856 rate mean (and  $\lambda_0 \sim 1/U(0, 100)$  for the rate-based model),  $\mu \sim U(0, 100)$  for the cause 857 delay (or influence) mean, and  $\sigma^2 \sim U(0, \mu^2)$  for the cause delay (or influence) variance. 858 We generated a simple Monte Carlo sample of size m = 100,000 to approximate the 859 Bayesian inference process. We will use m = 10,000 for the remaining datasets (unless all 860 parameters were assumed to be known to the model) because these datasets had either fewer unknown parameters or narrower parameter ranges compared to this study.<sup>14</sup> 862

As shown in Figure 10a, both event-based and rate-based schemes successfully identified the genuine cause in each condition, similar to the participants. Also consistent with participants' responses, the models were more influenced by the probability of intervening events than by delay length. As mentioned above, a mixture of factors in the study could result in different predictions as to whether there should be a short-delay or long-delay preference. As a result, the rational model predicted similar strength judgments in both conditions, with a slight tendency to favor the long delay (Figure 10a). This is driven by the relatively smaller variance in the long-delay condition. Although this 870 tendency did not manifest clearly in participants' responses, we can see how helpful a rational model can be in isolating the influences of different factors, which is valuable for 872 designing experiments and further developing process-level models to account for things 873 like memory storage and perceptual noise. Further trial-level comparisons were not conducted with this dataset because of the lack of trial-level human judgments. 875

Gong and Bramley (2023). A somewhat similar dataset was collected in Gong 876 and Bramley (2023). Participants were presented with a causal device consisting of one 877 target component (Effect E) and two control components (Cause A and B). The

<sup>14</sup> The rate-based scheme requires a decision on the time bin configuration. The bins should be fine enough to satisfy that a cause should happen in a time bin before the time bins capturing most of its effects. Here we used 1 second for Gong and Bramley (2023), Gong et al. (2023), Lagnado and Sloman (2006) and Lagnado and Speekenbrink (2010) and 1 day for Gong and Bramley (2024) and Greville and Buehner (2007). Both choices can be regarded as natural. We used 300 milliseconds for Bramley, Gerstenberg, Mayrhofer et al. (2018) given that more coarse choices would compromise the accuracy.

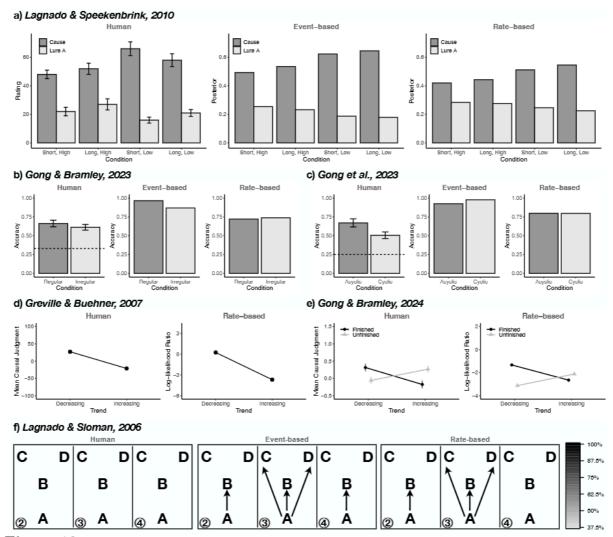


Figure 10

Aggregated results for six datasets. A temperature parameter of 15 was applied to Lagnado and Speekenbrink (2010) for visualization. Only one of the lures was reported in the original paper, presuming that the other lure could be calculated given the constraint in comparative ratings (LureB = 100 - Cause - LureA). Horizontal dashed lines in Gong and Bramley (2023) and Gong et al. (2023) indicate the chance-level performance. Ratings in Greville and Buehner (2007) are reversed so that they are aligned with Gong and Bramley (2024) where positive numbers indicated harmful influence and negative numbers indicated beneficial influence. The shading corresponds to the percentage of link selection in Lagnado and Sloman (2006); only links endorsed by more than 8 out of 24 participants or more than 50% chance by the models (after a softmax fitting) are shown; the occurrence orders were A-B-D-C, A-D-C-B, and A-B-CD (with the latter two occurring simultaneously) in Condition 2-4, respectively.

relationships between each control component and the target component could be 879 generative, non-causal, or preventative, resulting in 9 possible causal structures (see 880 Figure 9). A generative cause event would always produce an effect event after a gamma 881 distributed delay of 1.5±0.5 s. A preventative cause event would cancel any upcoming 882 effect events during a subsequent gamma distributed prevention window of  $3\pm0.5$  s. The 883 effect component could also activate spontaneously. Participants were randomly assigned to the regular base rate or the irregular base rate condition. Each base rate event occurred 885 semi-periodically, with gamma distributed  $5\pm0.5$  s delays after the previous one in the 886 regular condition, or unpredictably with exponentially distributed 5±5 s in the irregular 887 condition. Participants watched the device being intervened on by an artificial agent for a 888 total duration of 20 s during which there would always be three interventions on A and 889 three on B. 890

In the experiment we model here, participants were given video training to experience the delay parameters mentioned above. They were also explicitly told about the mean of the delays in the instruction. Therefore, we make the matching assumption that the model is also aware of these parameters. Specifically, for generative causes, we set  $w_c = 1$  (and  $\lambda_1 = 1$  for the rate-based model), the generative delay  $\mu = 1.5$  and  $\sigma^2 = 0.25$ . Regarding the base rate, we assume a mean of  $\mu_b = 5$  (and  $\lambda_0 = 1/5$  for the rate-based model).

In the case of preventative causes, the event-based scheme assumes the duration of preventative windows follow a gamma distribution with a mean of 3 and a variance of 0.25 (Figure 11a). All events occurring within a prevention window are assumed to be canceled. The rate-based scheme models the dynamics of preventative influence. It should be noted that the actual preventative mechanism employed here does not involve an incubation or decay process. Rather, the preventative window stays at its maximum level, effectively canceling all effects, for a certain duration. As such, under the rate-based scheme we accommodate this mechanism by modeling the preventative causal influence using the

gamma cumulative density function, as illustrated in Figure 11b, and assuming a maximum level denoted as  $\xi = 1$ .

The task instructions in Gong and Bramley (2023) imply three mechanistic features 908 of the scenario that can be implemented in the event-based scheme but less easily under the 909 rate-based. Firstly, a single generative cause event only produced one additional occurrence 910 of the effect component, which is consistent with the setup of the earthquake experiment 911 (Lagnado & Speekenbrink, 2010) described earlier. Secondly, in the regular condition, the 912 base rate events occurred semi-periodically. Therefore, instead of utilizing a memoryless 913 exponential distribution, the event-based model can employ a gamma distribution (with a 914 mean of 5 and a variance of 0.25), to model the semi-predictable delay between consecutive 915 base rate events. In contrast, since the rate-based model does not differentiate between effects generated by base rate events or generative causes, it is unable to leverage the 917 regularity of the base rate and thus treats the regular and irregular conditions in the same 918 manner. The third rule pertains to the preventative window. In the generative process, it 919 was the case that within a fixed preventative window, all expected effects would be 920 canceled, while any expected effects after the window would remain unaffected. 921 Consequently, the size of the true preventative window has to be smaller than the interval 922 between a preventative cause event and its nearest subsequent effect event E'. The absence 923 of an effect expected to occur after this E' can no longer be attributed to the preventative 924 causal influence. On the other hand, the rate-based model represents prevention as a 925 probabilistic influence, defining a soft window rather than a strict, deterministic one. 926

As shown in Figure 10b, aggregately the event-based scheme has higher accuracy compared to the rate-based scheme in identifying the causal models in Gong and Bramley (2023). It also shows a similar pattern of performing better in the regular condition compared to the irregular condition, which aligns with human performance. In contrast, the rate-based scheme demonstrates a slight tendency to perform better in the irregular condition, potentially attributable to the alignment of the base rate mechanism with the

933 model's assumptions.

For this and the subsequent datasets, we measured two types of correlation between 934 model judgments (the posterior probability of each answer) and human judgments (the 935 proportion of participants providing each answer). The first type is the Pearson correlation, which incorporates a softmax parameter to account for the stochastic nature of 937 judgments (Luce, 1959). We used a single parameter that we fit across all conditions for 938 each dataset. The second type is the Spearman correlation, which assesses the ranking 939 agreement between human and model judgments. This provides insight into how well the model captures the human dataset without introducing an additional free parameter. The 941 results are depicted in Figure 12a and 13a. Both the event-based and rate-based schemes 942 successfully captured human judgments, regardless of whether the conditions were regular or irregular. The event-based model demonstrated slightly superior correlations compared 944 to the rate-based schemes, suggesting that participants may have taken into account at 945 least some of the particular mechanistic constraints discussed above, during their reasoning 946 process. 947

We demonstrate here that people's judgments reflect rational considerations.

However, this does not mean the rational equations provide a good process-level account of
how people make these judgments. It is plausible that people relied on other algorithms to
approximate the rational solution. Gong and Bramley (2023) explored the types of
approximation algorithms people may use to choose among nine causal hypotheses after
observing a more-than-countable number of events short observation period. They found
that a summary-statistics approach, based on structurally local computation using
temporally local evidence, provided a better fit to participants' judgments than the

 $<sup>^{15}</sup>$  The softmax parameter  $\theta$  was used to maximize the log-likelihood between models' and participants' choices in Gong and Bramley (2023), Gong et al. (2023) and Lagnado and Sloman (2006), where participants' answers were binary about whether each causal connection existed or not. The parameter was used to maximize the linear correlation based on a non-linear transformation  $y=sign(x)|x|^{\theta}$  in Bramley, Gerstenberg, Mayrhofer et al. (2018), Gong and Bramley (2024) and Greville and Buehner (2007) where participants provided continuous ratings for how likely each connection existed or how strong each causal strength was.

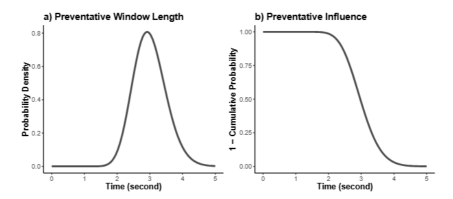


Figure 11

The preventative windows and preventative influences used to model Gong and Bramley (2023). a) The event-based scheme assumes the length of each preventative window is sampled from a gamma density function. b) To approximate this within the assumptions of the rate-based scheme, we assume the probability of prevention (what proportion of the effects will be prevented) is an inverse cumulative gamma function, reflecting a decreasing preventative influence over time.

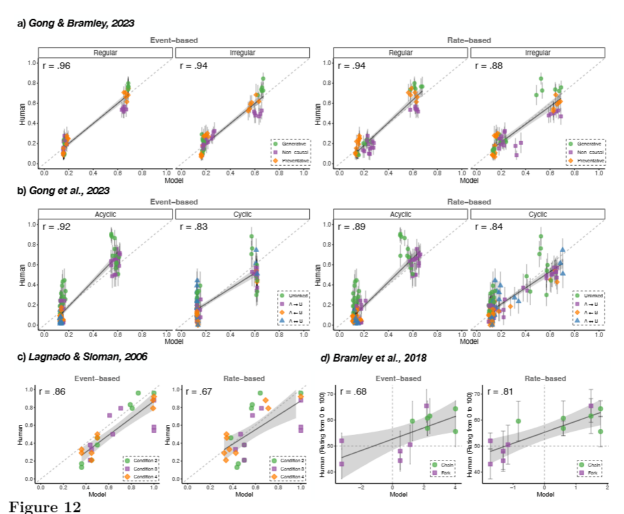
normative model. Readers interested in further details are referred to Gong and Bramley (2023). We further discuss process-level models as the future steps in the General Discussion.

## 959 Continuous, effect unspecified

Gong et al. (2023). When the effect variables are left unspecified, the number of potential causal structures increases quickly in the number of relata. Even when considering only generative relationships, there are four possible relationships between two variables: one-directional, reverse one-directional, bidirectional, and unconnected.

Consequently, for three variables, there are 64 potential structures, and for four variables, there are 4096 potential structures (refer to Figure 9). Gong et al. (2023) investigated how individuals learn about causal structures drawn from the large 3- and 4-variable hypothesis spaces by actively intervening in a causal system.

Although this is an active learning task, we use this model to analyze participants' judgment patterns rather than their intervention patterns. The rational framework we present here is an inference model, so it makes no distinction between learning from one's own interventions and learning from another person's interventions (these distinctions



Pearson correlations between model and human judgments. Y-axes indicate the proportion of human judgments in Gong and Bramley (2023), Gong et al. (2023), and Lagnado and Sloman (2006). Error bars indicate 95% confidence intervals of human judgments in the dataset whenever the raw data are available. Each individual point is the averaged response for a particular stimulus.

could be probed with a process-level model; Markant & Gureckis, 2014). To further
develop a rational account for how a learner should intervene to learn more effectively and
efficiently, we need to combine the current inference model with preposterior calculations
using information gain measures from information theory. Although this problem is beyond
the scope of this paper, readers are further referred to Gong et al. (2023) which makes and
tests proposals as to if and how people do this in continuous-time contexts.

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In the Gong et al. (2023) experiment, half of the ground truth structures contained

a cycle and half did not. Delay regularity was manipulated between participants. In each 979 causal system, for causally related components, an activated component would 980 probabilistically trigger the activation of each of its effect components once after a delay of 981  $1.5\pm0.1$  s in the regular condition, or after a delay of  $1.5\pm0.7$  s in the irregular condition. 982 All causal connections worked 90% of the time, and none of the components activated 983 spontaneously (i.e., there were no base rate activations). Participants were provided with six opportunities to activate a component in the system during a 45-second interval. 985 Considering the possible numerous connections and the cyclic structures, the number of 986 events recorded in this dataset was also significantly higher compared to the aforementioned Gong and Bramley (2023). 988

Participants in the study were pre-trained about the causal parameters with a similar procedure as in Gong and Bramley (2023), and we hence assume that models also know these parameters:  $w_c = 0.9$  (and  $\lambda_1 = 0.9$  for the rate-based model),  $\mu = 1.5$ , and  $\sigma^2 = 0.01$  or  $\sigma^2 = 0.49$  depending on the specific regular or irregular condition. No base rate was assumed by either model.

Human results showed a main effect of the structure cyclicity but no main effect of 994 the delay regularity, probably due to that the difference between regular and irregular 995 settings was not pronounced enough (Gong et al., 2023). Therefore, we here focus on the 996 results based on the cyclicity factor alone. In contrast to humans who performed better in 997 the acyclic condition than the cyclic condition, the event-based model demonstrates better 998 performance in the cyclic condition compared to the acyclic condition (Figure 10c). This 999 reflects the fact that the event-based model is able to leverage the larger amount of event 1000 information available in the cyclic structures via the one-effect per cause mechanistic 1001 constraint (Gong et al., 2023). Conversely, the rate-based model does not demonstrate the 1002 same tendency. Due to not enumerating actual causation pathways, this model fails to 1003 leverage the information available in abundance of cyclic events as effectively as the 1004 event-based model does.

In terms of both correlation measurements, the event-based model demonstrates 1006 better performance in capturing human judgments in acyclic structures, while the 1007 rate-based model performs better in capturing human judgments in cyclic structures (see 1008 Figure 12b and 13b). This may suggest that as the number of events increases, the exact 1009 enumeration computations become infeasible for people, necessitating the relaxation of the 1010 one-cause-one-effect constraint within the event scheme to enable more efficient 1011 approximations. The rate-based model's capability to capture human cyclic judgments 1012 highlights its ability to deal gracefully with larger number of events, providing a not perfect 1013 but more efficient approach in such scenarios. 1014

Once again, the analysis here demonstrated participants' rational thinking in solving the online causal structure learning problem, but this does not imply that participants can perform exactly as the rational models do. As shown in Figure 10c, participants' accuracy was lower than that of the models, and their accuracy decreased with the increasing number of events (Gong et al., 2023). This suggests that they may not be able to store and utilize the temporal information of all observed events as effectively as the rational framework.

#### Episodic, effect specified

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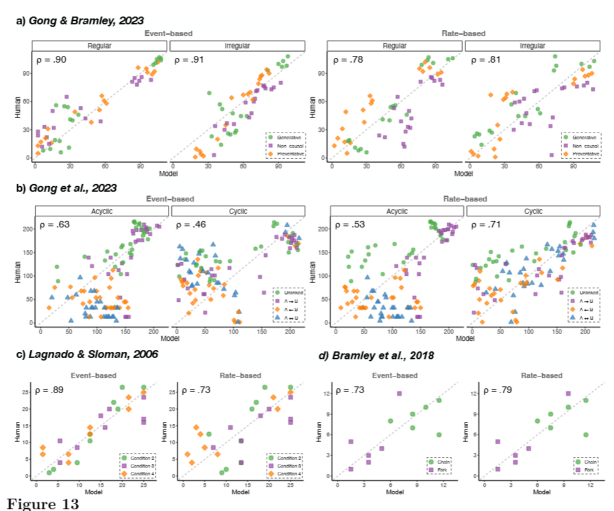
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Greville and Buehner (2007). We refer to "episodic evidence" as evidence 1023 gathered from multiple independent causal systems of the same type, where each sample 1024 has its own timeline. Episodic evidence can be seen as a combination of contingency and 1025 temporal information (Greville & Buehner, 2007). It involves the observation of multiple 1026 independent individuals, but with each observation lasting over a specific time period 1027 (Figure 1b). Research on episodic evidence often focuses on cases in which each type of 1028 event occurs at most once within the observed period (Bramley, Gerstenberg, Mayrhofer 1029 et al., 2018; Gong & Bramley, 2024; Greville & Buehner, 2007; Lagnado & Sloman, 2006). 1030 This means that the evidence within each individual's experience may not be very 1031 informative. However, by considering multiple cases, the reasoner can compensate for the 1032



Spearman correlations between model and human judgments. Axes are the ranks of judgments.

limited information within each instance and make more informed conclusions about their common causal structure.

In Greville and Buehner (2007), participants were asked to examine the influence of a ray treatment on the survival of bacterial cultures. Bacterial cultures were randomly assigned to the experimental group, which received a ray treatment at Day 0, or the control group, which did not receive any treatment. Each group consisted of 40 samples. Bacterial cultures were observed from Day 1 to Day 5. The number of new deaths occurring each day was recorded. Participants were asked to rate whether they perceived

 $<sup>^{16}</sup>$  The data were displayed as tabular records indicating whether each culture was still alive or not in

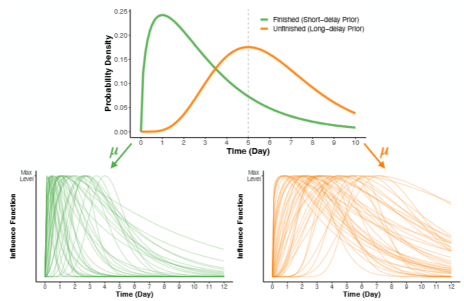


Figure 14
Short-delay and long-delay priors regarding the timing of when the cause will take effect on average (Gong & Bramley, 2024; Greville & Buehner, 2007). The parameter  $\mu$  is sampled from different prior distributions to form different causal influence functions.

the treatment as harmful or beneficial based on the observed outcomes in both the 1041 experimental and control conditions. The control condition always demonstrated a 1042 relatively constant death rate over time (e.g., 8, 8, 8, 8, 8), while the daily death rate in the 1043 experimental condition was manipulated to exhibit either a decreasing or an increasing 1044 trend. Results showed, after controlling for the total number of deaths over the 5-day 1045 period, participants judged the treatment as more harmful if there were more deaths right 1046 after the treatment (a decreasing trend; e.g., 16, 12, 8, 4, 0), and more beneficial if there 1047 were more deaths towards the end of the observation period (an increasing trend; e.g., 0, 4, 1048 8, 12, 16). As such, this study demonstrated that people not only care about the overall 1049 contingency data summarized from the entire observation period but also the detailed 1050 temporal dynamics (at a day-to-day level here). 1051

Greville and Buehner (2007), and as summarized counts of how many cultures died each day in the later Gong and Bramley (2024). Given that the results of Gong and Bramley (2024) replicated the same "finished" condition as in Greville and Buehner (2007), we do not discuss the potential influence of formats here.

Gong and Bramley (2024). While agreeing on the impact of temporal dynamics 1052 on judgments, Gong and Bramley (2024) proposed that some settings could produce a 1053 different pattern than the traditional notion of contiguity (Greville & Buehner, 2007). In 1054 this task, if in some conditions learners tended to assume that the causal process may be 1055 ongoing, an increasing trend might signal that the treatment will ultimately prove harmful. 1056 Gong and Bramley (2024) presented participants with more such ambiguous data, where a 1057 majority of the forty samples were still alive on Day 5 (e.g., 0, 1, 1, 3, 5 in the 1058 experimental condition and 1, 3, 2, 2, 2 in the control condition). Participants in the 1059 "Unfinished" condition were informed that the observation had not yet concluded, while 1060 participants in the "Finished" condition were told that the observation had finished (as in 1061 Greville & Buehner, 2007). Results in the Finished condition replicated Greville and 1062 Buehner (2007). However, in the Unfinished condition, participants interpreted an 1063 increasing trend in deaths as indicative of harm caused by the treatment, and a decreasing 1064 trend as indicative of benefit (see Figure 10e). 1065

These findings highlight the influence of instructional cues on participants' inductive 1066 biases and how they interpret the observed data. To model the human judgments, we here 1067 assume that instructions tend to influence the learner's prior expectation about causal 1068 delays as well the use of data. As shown in Figure 14, if participants are informed that the 1069 experiment ends at Day 5, they may tend to form a prior belief that the relevant causal 1070 influences are expected to occur within 5 days. When participants were led to believe that 1071 the observation had not finished, they anticipated the possibility of longer causal delays. 1072 Here we assume that, for the Finished instruction (Gong & Bramley, 2024; Greville & 1073 Buehner, 2007), the causal delay (or the expected time of the influential function in the 1074 rate-based context)  $\mu$  is sampled from a gamma distribution with mean of 3 (days) and a 1075 variance of 6. For the Unfinished instruction (Gong & Bramley, 2024),  $\mu$  is sampled from a 1076 gamma distribution with mean of 6 (days) and a variance of 6. While a range of values 1077 might be reasonable, we chose these because range of 0 to 5 would cover most of the 1078

sampled  $\mu$  under the Finished instruction (83%) while covering only a minority of the sampled  $\mu$  under the Unfinished instruction (38%, Figure 14).

The observed death of each bacteria culture in Greville and Buehner (2007) and 1081 Gong and Bramley (2024) could result from either the treatment or natural death (i.e., the 1082 base rate). Given that the cultures which died out in the same day were not 1083 distinguishable from each other, we here focus on the rate-based scheme and consolidate 1084 the data as shown in Figure 4e. Since the data were collapsed, the rate of how many events 1085 happened per day depends on the total sample size (i.e., forty in both studies). We assume 1086 that participants selected between the "harmful" and "beneficial" options by comparing the 1087 likelihood between a generative structure and a preventative structure (see Figure 9). To 1088 model the data, we set  $\lambda_0 \sim U(0,40)$ ,  $\lambda_1 \sim U(0,40)$  since there were at most forty cases in 1089 each group. We set  $\xi \sim U(0,1)$  for the max level of preventative influence (i.e., the 1090 beneficial influence). Similar to previous datasets, we set the variance  $\sigma^2 \sim U(0, \mu^2)$ . 1091

The aggregated results are shown in Figure 10d and 10e. Participants' inclinations 1092 in Greville and Buehner (2007) and Gong and Bramley (2024) were captured by the model. 1093 Under the Finished instruction (Gong & Bramley, 2024; Greville & Buehner, 2007), 1094 participants and the model both treated decreasing trends as more harmful than increasing 1095 trends, lining up with the contiguity-driven explanation. Under the Unfinished instruction 1096 (Gong & Bramley, 2024), participants and the model both treated increasing trends as 1097 more harmful than decreasing trends, lining up with the sensitivity to trends. Nevertheless, 1098 the model was less likely to demonstrate a cause as absolutely harmful (i.e., giving a rating 1099 above 0). This could be due to the fact that Gong and Bramley (2024) handcrafted the 1100 stimuli, where only a small number of bacterial cultures out of the sample (n=40) died 1101 during the observation, and the overall death rates were similar between the experimental 1102 and control groups (e.g., 3, 4, 2, 1, 0 in the experimental group and 2, 2, 1, 2, 3 in the 1103 control group). An ideal learner would expect the experimental condition to still exhibit a 1104 higher death rate towards the end of the observation if they were to claim a cause as an 1105

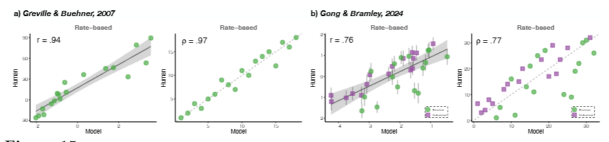


Figure 15
Pearson (r) and Spearman ( $\rho$ ) correlations between model and human jugdments in
Greville and Buehner (2007) and Gong and Bramley (2024). Error bars indicate 95%
confidence intervals of human judgments in Gong and Bramley (2024).

absolute harm. Participants' deviations from this prediction could have several
explanations. One speculation is that they did not simply represent harm as generative
(causative of death) and benefit as preventative, like the model. They may assess whether
a treatment is good or bad by also considering whether it advances or delays death, which
is a sensible interpretation, but falls outside the scope of our current model. We will return
to this point as a future direction in the General Discussion.

The trial-level results are shown in Figure 15. The rate-based model achieved a good fit with human judgments at the trial level in Greville and Buehner (2007) and Gong and Bramley (2024).

#### Episodic, effect unspecified

The final category we consider here is episodic evidence where the effect variable is unspecified (i.e., accepting a wider range of hypothetical structures rather than focusing on specifying one variable as the effect and finding its causes; Figure 9). The two datasets we consider here both involve scenarios where each kind of event can only happen once in each episode.

Lagnado and Sloman (2006). In Lagnado and Sloman (2006), participants were asked to imagine a situation in which a computer virus can spread through a network and told that the time at which a computer revealed its infection could occur after a variable delay, so later than the time at which the computer became infected. Participants were

told that each connection, if it existed, would spread the virus 80% of the time, and the virus could not reach a computer unless it had been sent from another (e.g., no spontaneous base rate infections would occur). Participants watched 100 clips, each showing an event sequence in which the virus appeared in different computers, and were asked to judge the existence of various potential causal links (i.e., directed network connections) in the system (see Figure 9).

The experiment included four conditions (in Condition 1 all events happened 1131 simultaneously so won't be modeled here), but the underlying ground truth structure was 1132 consistently: A was the cause of B, and B was the common cause of C and D. This meant 1133 that in each condition, computers C or D would never become infected without computer 1134 B being infected. Since the actual infection time was varied and unknown, the presumed 1135 rational solution is to rely on the conditional probability. However, the timing of virus 1136 appearance in each computer could be misleading. For example, in Condition 3 where 50% 1137 of trials followed the order of A - D - C - B, participants tended to judge the links 1138  $A \to D$ ,  $D \to C$ , and  $C \to B$  were more likely to exist than other links (see Figure 10f). 1139 Their answers cannot explain the other 50% of trials when only AB, ABC or ABD 1140 happened. This suggests that people's reliance on temporal information is so strong that it 1141 could not, in this case, be overshadowed by the contingency information. As such, in order to capture human judgments, what we will model here is not the objective rational 1143 solution, but the rational solution having disregarded the instruction that the temporal 1144 information is irrelevant.<sup>17</sup>

There was a one-second delay between events in subsequent time steps  $(t_1, t_2, t_3, t_4;$  see Table 2 in Lagnado & Sloman, 2006). As such, each trial lasted 4 s. We model the dataset using the parameters  $w_c = 0.8$  (and  $\lambda_1 = 0.8$  for the rate-based model) as in the instruction,  $\mu \sim U(0, 10)$ ,  $\sigma^2 \sim U(0, \mu^2)$ . The base rate is assumed to be zero. In this

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<sup>&</sup>lt;sup>17</sup> Note that after personal contact, we learned that the condition-by-condition human judgments published in Table 3 of Lagnado and Sloman (2006) were incorrect. Readers may refer to p.188 in Pacer (2016) or the GitHub repository of this paper for the corrected table.

dataset, the main difference between event-based and rate-based schemes is that the former 1150 address the rule that an event can only occur once for a specific equipment, which is 1151 consistent with the experimental design. Consequently, the event-based scheme 1152 outperforms the rate-based scheme in accurately capturing human judgments, as shown in 1153 Figure 12c and Figure 13c. Both models capture the aggregated human judgment patterns, 1154 as shown in Figure 10f. This includes the Condition 3 mentioned above, despite the fact 1155 that the models also tend to add  $A \to B$  and  $A \to C$  links which can help rationally 1156 account for the trials when B and C happened without D. Our framework demonstrates 1157 an ability to explain the phenomenon that temporal information can outweigh contingency information in human causal judgments. 1159

Bramley, Gerstenberg, Mayrhofer et al. (2018). Bramley, Gerstenberg, 1160 Mayrhofer et al. (2018) tested whether people can differentiate between two causal 1161 structures, chain and fork (see Figure 9), solely using delay information. Each trial 1162 consisted of 12 episodes, wherein events always occurred in the order A-B-C. However, 1163 there were variations in the delay variances between structures. In the chain structure 1164  $(A \to B \to C)$ , the delay variance between B and C was small, whereas the variance 1165 between A and C was large, as it encompassed the variability across both causal 1166 connections. Conversely, in the fork structure  $(B \leftarrow A \rightarrow C)$ , the delay variance between A 1167 and C was small, while the variance between B and C was large, as there was no direct 1168 causal link between the two variables. Participants were asked to judge by distributing 100 1169 percentage points across the two structures. 1170

In contrast to previous datasets, we used the "independent delay" parameterization, as described in the original paper (Bramley, Gerstenberg, Mayrhofer et al., 2018), which allowed for there to be distinct delay distributions between different links in the causal structure. This means that to choose between chain and fork, we only need to model the delays between B and C in the chain hypothesis and the delays between A and C in the fork hypothesis (since both hypotheses share the  $A \rightarrow B$  connection). Each episode lasted

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3 s. We assume  $w_c = 1$  (and  $\lambda_1 = 1$  for the rate-based model),  $\mu \sim U(0, 10)$ ,  $\sigma^2 \sim U(0, \mu^2)$ , and no base rate.

Results are shown in Figure 12d and 13d. Both models captured the general pattern 1179 of human judgments. The rate-based model demonstrated a better fit to human judgment 1180 compared to the event-based model. The event-based model showed a overall bias towards 1181 chains (judging all chain devices as chains and also judging some forks structures as 1182 chains). This is as expected and is due to the fact that A-C delays (calculated under the 1183 fork hypothesis) were always longer than the B-C delays (calculated under the chain 1184 hypothesis) in the stimuli. As a result, the ceteris paribus preference for the chain 1185 structure can be interpreted as an example of favoring the relatively shorter delay. In 1186 contrast, from the same evidence, the rate-based scheme, which assumes macro causal 1187 dynamic changes, has a greater tolerance for delays of causal influence than the 1188 event-based scheme. Participants may also have reasoned pragmatically that around half of 1189 the ground truths would be fork structures and so shifted their response threshold to favor 1190 the fork response more than the evidence supported. 1191

We note here that the computational cost of the rate-based model is not always lower than that of the event-based model. In the case of this dataset, the rate-based model may actually be more computationally demanding depending on the granularity of the time bins it uses. Conversely, the event-based model benefits from the fact that each type of event occurs only once in each episode, resulting in a small number of causal pathways to consider within each hypothesis.

#### General Discussion

In this paper, we developed a rational framework for the use of temporal information in causal inference. The framework leverages stochastic processes from the Poisson-Gamma family to model the (in)dependencies between events in time and drive selection and parametrization of causal structure hypotheses. To achieve this, we extended the causal graphical model formalism to incorporate likelihood functions for temporal

dynamics, before inverting these likelihoods to derive structural conclusions from evidence. 1204 We show how this general approach uncovers the underlying causal structure in all manner 1205 of complex continuous-time datasets. The framework is applicable to a wide range of 1206 temporal causal learning, associative and operant learning tasks, including scenarios where 1207 evidence comes in a long continuous timeline or from many shorter independent timelines, 1208 or when the causes can produce either one specific effect event or multiple effect events. As 1209 we demonstrate in modeling existing datasets, the framework accommodates variations 1210 such as the size of the hypothesis space, the involvement of background activity, 1211 preventative connections, cyclic dynamics, and whatever other information learners have 1212 about relevant causal delay distributions. To our knowledge, this is the first general 1213 computational framework for learning causal structure from events unfolding in continuous 1214 time. 1215

The framework anticipates three intuitions that have been frequently observed in 1216 human learning: causal attributions are, in general, stronger to the extent that the delays 1217 between a putative cause and effect tend to be shorter, more consistent, and more in line 1218 with preexisting mechanistic expectations. We demonstrate the rationale behind these 1219 intuitions falls directly out of the Bayesian framework, explaining why it makes sense for 1220 them to coexist and why a preference may fail to manifest itself under certain 1221 circumstances. Additionally, we demonstrate that the framework helps explain behavioral 1222 patterns across a range of learning tasks from the last 20 years. We find a high degree of 1223 consistency between judgments derived from the rational framework and the aggregate 1224 behavior of human participants. These analyses suggest that people are not only capable of 1225 utilizing temporal information in diverse causal learning situations, but also that they do 1226 so in systematic, predictable, and approximately rational ways. By providing a unified 1227 computational framework, we are finally able to consolidate empirical studies spanning 1228 many different tasks and better clarify these tasks' relationship with widely studied 1229 associative and reinforcement settings. This model offers a design space for locating

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different tasks within the temporal causal learning field and provides guidelines for further investigation of causal cognition (Almaatouq et al., 2022).

In the remainder of the paper, we compare our model with previous temporal associative learning models, discuss why we think the pluralistic "dual-aspect" view we present here makes sense for describing human temporal causal learning, consider the relationship between continuous and discrete value representations, and lay out several future research directions.

## A comparison to temporal associative learning models

We are not the first to point out the limitations of framing learning around 1239 trial-based contingencies, and consider how inferences operate on continuous-time data 1240 directly. This issue has been discussed in detail in the associative and animal learning 1241 literature. In particular, Gallistel and colleagues (Gallistel, 2021; Gallistel et al., 2014, 1242 2019; Gallistel & Gibbon, 2000; Gallistel & Wilkes, 2016) argue that the way in which time 1243 is segmented into trials for analysis, as well as the duration considered as a single trial, can 1244 dramatically alter the predictions from contingency-based associative learning models; if 1245 researchers' choices depart from whatever intuitive structure and discretizations their 1246 subjects make in understanding their tasks, their models are doomed to lack the 1247 representational expressivity needed to capture their subjects' learning processes. Like us, Gallistel et al. propose that learning depends on the rate at which the effect occurs after a 1249 cause, or operant behavior, takes place (Gallistel, 2021; Gallistel & Gibbon, 2000; Rescorla, 1250 1968). They further point out that temporal associative learning is not just determined by 1251 the frequency of temporal pairings (i.e., how often a presumed effect follows its presumed 1252 cause); it must also be sensitive to how often the effect occurs spontaneously without the 1253 cause's occurrence. 1254

In their detailed experimental analyses, Gallistel et al. focus at the process-level on the pairwise attribution problem. For instance, Gallistel and Gibbon (2000) show that a response behavior is triggered when the ratio of rates with or without a stimulus exceeds a 1268

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certain threshold. Gershman (2024) later suggest that a Rescorla-Wagner framework can 1258 be used to update the weights of different associative causes, by shifting from predictions 1259 about the presence or absence of effects to predictions about fluctuations in the rate of 1260 effects. Gallistel et al. (2014) also demonstrate that association strength depends on how 1261 much additional predictive information a presumed cause provides about its presumed 1262 effect, having accounted for the effect's base rate (Gallistel et al., 2014, 2019; Gallistel & 1263 Wilkes, 2016). In this more recent treatment, the casual inference no longer relies on 1264 detecting a rate change but instead on contrasting the random-timepoint  $\rightarrow$  next-reward 1265 delay distribution against a cue  $\rightarrow$  next-reward delay distribution, and using the entropy reduction between these distributions as a causal index. 1267

There are obvious connections between Gallistel et al.'s theoretical ideas and ours: both embrace rate representations and contrast causal against baseline effect patterns. Their models, like our Bayesian approach, predict the phenomenon of time-scale invariance, 1270 because the associative strength depends only on the relative, not absolute rates or delays in situations with or without the cause (Gallistel et al., 2019; Gershman, 2024; Rescorla, 1968). However, apart from time-scale invariance, it is unclear whether an associative model can explain why learners favor causal explanations that posit causal delay durations which are relatively shorter, more in line with prior expectations, and more regularly timed.

One feature that differed across the experimental datasets was whether the effect 1276 was pre-specified. When the effect is specified, the task is to identify the true (positive or 1277 negative) causes of this effect (Gong & Bramley, 2023, 2024; Greville & Buehner, 2007; 1278 Lagnado & Speekenbrink, 2010). This is similar to the credit assignment problem in 1279 associative (or reinforcement) learning, where learners credit conditional stimuli (or 1280 interventions) for a particular unconditional stimulus (or a kind of reward). Temporal 1281 associative learning models can potentially provide predictions in these tasks. The 1282 advantage of the Bayesian framework is that it applies equally to the open-ended 1283 "structure learning" tasks, prototypical in the causal cognition literature, where nothing is a priori specified as a cause or an effect. These scenarios require more global reasoning (as well as interventional data) to solve reliably and the Bayesian framework helps clarify the circumstances where locally focused heuristics are or are not sufficient to arrive at the right global model (Bramley, Dayan et al., 2017; Fernbach & Sloman, 2009).

Another characteristic of the Bayesian account is its flexibility to deal with varied 1289 temporal dynamics. What it actually compares here is how well different causal 1290 explanations fit. This can include explanations making different assumptions about 1291 functional form as well as about structure. For instance, current temporal associative learning models implicitly assume that if a cause produces multiple effects, it will produce 1293 them at a constant rate during an effective time window with hard onset and offset 1294 boundaries (Gallistel et al., 2019; Gallistel & Gibbon, 2000; Gallistel & Wilkes, 2016). In 1295 contrast, we illustrate in this paper how the Bayesian approach can handle whatever 1296 hypotheses are articulated. For example, we modeled a case where changes in the effect's 1297 rate followed a latent, peak, and decay process continuously, but could contrast this with a 1298 uniform generation window or any other mechanistic hypothesis. Through the event-based scheme, it also allows for the incorporation of other mechanistic constraints, such as the 1300 case where a cause can generate only one effect, or the possibility that the where baseline 1301 effects are not unpredictable, but periodic. In such a situation where the base rate itself is 1302 a moving target, it is unclear whether a simple entropy reduction index (Gallistel et al., 1303 2014, 2019) would provide a generalizable index of the power or strength of causal or 1304 relationship (cf. Cheng, 1997). 1305

Note that all advantages we mention pertain to the flexibility of the model space
that Bayesian inference is defined over. This is wholly compatible with the idea that at the
process level, we rely on the mechanisms of pairwise association or reinforcement among
other pragmatic, resource sensitive heuristics and approximations. Nevertheless, we hope
this rational analysis is useful for mapping out the space of continuous-time learning
problems including those classically used in associative learning tasks.

### 1312 A pluralistic view

We presented two schemes, event-based and rate-based, in parallel throughout this 1313 paper but introduced both as manifestations of a broader Poisson-Gamma framework for 1314 conceptualizing interevent dynamics. The existence of a pluralistic view is not a new concept in the field of causal cognition. For instance, in research on token-level causal 1316 attribution, where individuals are asked to make judgments regarding what was responsible 1317 for particular event rather than causative of a class of events in general (Halpern, 2016), 1318 researchers have debated the relative importance of covariation versus process (Gerstenberg 1319 et al., 2021; Lombrozo, 2010; Sloman, 2005; Wolff, 2007). The question arises whether 1320 people prioritize imagining how the outcome would have changed if the cause had been 1321 different (Icard et al., 2017; Sloman, 2005), or if they focus more on determining if there 1322 was a genuine physical exchange between the cause and effect (Talmy, 1988; Wolff, 2007). 1323 Instead of relying on a single level of abstraction, people are pluralist, considering both the 1324 occurrence of the outcome and the manner in which it occurred (Gerstenberg et al., 2021). 1325 This paper focuses on a type-level causal learning rather than token-level causal 1326 attribution, meaning we can benchmark the quality of a judgment against the true causal 1327 generative model that they are learning about. We next give two reasons why a pluralistic 1328 perspective is also important in the domain of type-level causation, especially when it 1329 comes to temporal evidence based learning. 1330

## $Mechanistic\ concerns$

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Causal structure learning can be driven by different types of evidence at different
levels of abstraction. As we orient away from highly abstracted atemporal contingencies
toward "raw" spatiotemporal dynamics, the richness of the data increases. Atemporal
evidence discards a lot of information by discretizing into finite sets of categories and time
points(Allan, 1980; Cheng, 1997; Griffiths & Tenenbaum, 2005; Perales & Shanks, 2007).
For instance, researchers examined causal inference from continuous spatiotemporal
evidence when asking individuals to make causal inferences about objects in 2D physical

scenes where it is unlikely that participants will ever see exactly the same thing happen twice (Bass et al., 2021; Bramley, Gerstenberg, Tenenbaum et al., 2018; Ludwin-Peery et al., 2021; Ullman et al., 2018). Due to the high dimensionality of the clips used in these studies, it is crucial to leverage one's pre-existing mechanistic theory (e.g., a familiarity with everyday intuitive physical dynamics) to discover latent causal features such as objects' masses or force relations within the space of a short observation.

We argue that temporal evidence shares characteristics with both atemporal and spatiotemporal evidence. Like atemporal data, temporal evidence permits some 1346 discretization and aggregation, as effect events may occur multiple times without the 1347 necessity of having individual identifications (e.g., the bacteria culture example in 1348 Figure 1b; Gong & Bramley, 2024; Greville & Buehner, 2007; Griffiths & Tenenbaum, 2005; Pacer & Griffiths, 2012). This allows for type-level reasoning, about how the rate of 1350 effect occurrence changes after a putative cause occurs (i.e., the rate-based scheme). At the 1351 same time, temporal information also invites token-level reasoning. When one cause 1352 produces a very limited number of effect events (e.g., only one per component), the precise 1353 delays between each cause and effect and the prior expectations about causal and 1354 non-causal delays becomes important. Type-level causal conclusions will arise from the 1355 detailed inference about which specific occurrence of a cause was responsible for this 1356 specific occurrence of the effect (i.e., the event-based scheme). As such, the general 1357 Bayesian inference framework allows us to express whatever mechanistic or ontological 1358 commitments we believe capture a particular causal inference domain.

#### $Computational\ cost\ concerns$

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Continuous time allows for precise temporal information, with each event having its unique time point and relationship with all other events. Events of different classes are often intermingled, and events of the same class may occur many times within the same observation. However, this precision and combinatorial credit assignment issue poses computational challenges and becomes infeasible when there are many events under

consideration. Strictly, observing a causal system in continuous time with uncertainty 1366 about the true causal delays, any event could theoretically be the result of any event that 1367 happened in the past. As a real-world example there are diseases, such as the bovine 1368 variant of Creutzfeldt-Jakob disease, that have very long incubation periods. The cause of 1369 a disease onset could be traced back to something eaten 15 years ago (Valleron et al., 1370 2001), but so many candidate events will occur within this period that it is impossible to 1371 consider them all. As such, a real cognizer should take seriously the trade-off between 1372 cost-of-computation and accuracy when reasoning about causal structure in their 1373 environment. The event-based and rate-based schemes we present here provide two levels 1374 at which one can process the same evidence, with the former generally more costly in its 1375 analysis of the micro-level delay details and the other a more abstracted and efficient way 1376 to capture the macro-level rate changes. By considering both approaches, a learner could 1377 flexibly choose or learn to represent a domain in a way that is sufficient and practical for 1378 their purposes. A rate-based scheme is especially useful when dealing with a large number 1379 of effects where it usually requires less computation. However, determining the appropriate 1380 granularity for rate calculation introduces another cost-benefit trade off that needs to be 1381 explored. 1382

#### 83 Abstraction and reduction: Moving between levels

We have focused on learning from events in continuous time, whereas other studies 1384 have examined causal learning from interactions with or observations of continuous valued 1385 variables varying in continuous time (Btesh et al., in press; Davis et al., 2020; Rehder 1386 et al., 2022; Soo & Rottman, 2018; Zhang & Rottman, 2023). Rather than viewing these as 1387 completely separate tasks, we think it is more fruitful to think of continuous and eventive 1388 representations as complementary ways of modeling and explaining causal phenomena. To 1389 illustrate this, consider the predator-prey relationship, such as that between lynx and 1390 hares. At a low level, we might model individual events such as a individual lynx catching 1391 and eating an individual hare. Abstracting this to a higher level, we might analyze how

populations of lynx and hares change over time, based on their populations, or similarly on 1393 their fluctuating birth and predation rates. At a higher level, we can investigate how each 1394 species experiences cyclic patterns of population-scale events and shocks such as "bloom" 1395 and "collapse", and analyze the progression of these an event representation again. By 1396 abstracting upwards or unpacking downwards, it seems that a flexible reasoner can cycle 1397 unboundedly between representations in terms of continuous values and those in terms of 1398 discretized events with the more appropriate choice determined by its utility in guiding 1399 action rather than adherence to metaphysical reality. 1400

Importantly, it is not necessary to limit causal reasoning to the level natively 1401 provided by the data. When modeling data from Gong et al. (2023), we showed that the 1402 rate-based model outperformed the event-based model in capturing human performance in 1403 identifying cyclic structures, although the evidence was actually generated following an 1404 event-based scheme. This kind of abstraction can be boundedly rational. People may spontaneously abstract to a relatively continuous representation (i.e., the rate) when it 1406 makes computational sense to do so even if this prohibits some subtler mechanistic 1407 considerations. They might also do the reverse. Rehder et al. (2022) model structure 1408 inference in a setting involving continuous variables varying in real time. Their modeling 1409 suggests participants are overwhelmed by the full dynamics and rather abstract these into 1410 a handful macro-scale "events" – essentially treating moments of dramatic increase or 1411 decrease as events and performing token-level causal inference about relationships between 1412 these. Taken together, this all suggests that people have the ability to adopt 1413 computationally sensible representations. The layout of the event-based scheme in this 1414 paper also suggests that, contrary to the common assumption in computer science and 1415 other time-related cognitive models, the representation and operation of temporally 1416 relevant data are not necessarily bound to the discretization of time into bins, which may 1417 lead to inefficiency — representing numerous empty bins where no events occur — or 1418 distortion due to inappropriate bin width, where multiple events per bin reduce temporal

accuracy and obscure order information. Reasoning from temporal evidence introduces a different and often more difficult computational challenge compared to the previously studied atemporal learning setting. As such, it can serve as a useful setting for studying the mechanisms that guide bounded rationality (Lieder & Griffiths, 2020; Simon, 1982).

## Limitations and future directions

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In the current paper, we treat events as instantaneous, occupying a single time 1425 point on the timeline. This means that a generative cause will always produce an 1426 observable effect even if that effect occurs close to another effect event. These point event representations could be seen as simplifications of everyday events that ignore their 1428 duration. Sometimes the duration of everyday events is too long to ignore: wet ground will 1429 stay wet for some time, tanned skin will fade slowly. In these cases, generative events can 1430 easily be overshadowed by already-occurring events (i.e., we might not notice that the 1431 sprinkler system came on because it had also been raining). This results in more complex 1432 causal inference scenarios such as preemption and over-determination (Gerstenberg et al., 1433 2021; Lombrozo, 2010). These sorts of situations can be handled by extensions to the Poisson-Gamma framework, especially using event-based scheme, by incorporating the 1435 relevant mechanistic knowledge to the model (cf. Bramley, Gerstenberg, Mayrhofer et al., 1436 2018). However, such situations are relatively unexplored in the temporal causal learning 1437 setting. When events have richer internal structure, such as a gradual onset and offset, the 1438 causal learning process becomes entangled with the question of people abstract continuous 1439 input into events. Future research could explore the possibility of integrating causal 1440 considerations with the theory of event segmentation (Altmann & Ekves, 2019) to build a comprehensive model of how discretized representations arise form continuous inputs. 1442 Causal cognition researchers have used linear regression (Rottman, 2016; Soo & 1443 Rottman, 2018) and the Ornstein-Uhlenbeck (OU) process (Btesh et al., in press; Davis 1444 et al., 2020; Gong & Bramley, 2022; Rehder et al., 2022; Uhlenbeck & Ornstein, 1930) to generate continuous-variable, continuous-time dynamics and treated inference within the

requisite model class as determining the normative solution to learn causal mechanisms. 1447 Although our event-level framework does not extend to continuous variables, linking it to 1448 this setting is a goal for future research. As an initial example, Gong and Bramley (2022) 1449 used the OU process to construct and test human inferences about continuous causal 1450 dynamics influences with a variety of properties. Similar to learning from event sequences, 1451 participants made stronger more confident judgments when the lag between changes to 1452 cause and effect variables' values was shorter and when the changes were more dramatic. 1453 This finding may suggest that combining the algorithms used in continuous-variable studies 1454 with the current Poisson-Gamma framework can help us better understand how people 1455 infer discrete event structure to explain continuous dynamics. 1456

Although we focus on the role of time in causal reasoning, the causal relationships 1457 we investigate in this paper still align with the primary focus of atemporal causal learning 1458 literature: generation and prevention. In fact, causal relationships can be much richer once the time dimension is taken into account. For example, a cause can be "zero sum", in the 1460 sense of merely altering the timing of subsequent effects without affecting their frequency 1461 (Bennett, 1987). In other words, something might have a large causal effect on something 1462 else, not because it generates or prevents events but instead modifies when those events 1463 occur, by hastening or delaying them, or otherwise influencing their occurrence in time 1464 (Greville et al., 2020). Hastening and delaying are two relationships that are less studied in 1465 causal literature but extremely common in daily life; for example, we regularly experience things like transport and logistics delays. This richer space of causal difference-making 1467 could be studied from a signal detection perspective, by examining the conditions under 1468 which people notice when an event has had a causal influence on the occurrence of another, 1469 and when they can recognize what functional form that influence has taken. These kinds of 1470 situations could also be studied from a causal language perspective (Beller & Gerstenberg, 1471 in press; Wolff, 2007). For example, there might be important differences between someone 1472 judging that a poison killed A than that it hastened A's death. With the temporal

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dimension in play, further research can investigate detection and representation of a broader range of causal influence patterns both empirically and computationally.

In this paper we showed that participants' judgments across seen experiments can 1476 be unified under a rational framework. This is just one part of the project of understanding 1477 how people make these judgments. Humans have limited cognitive resources for receiving, 1478 processing, and memorizing information. These constraints significantly impact temporal 1479 causal learning, especially when the time gap between a cause and its effect is long. This 1480 paper focuses on developing the computational-level account, while it is important not to 1481 stop there but to use this to help investigate the process level (Marr, 1982). Some of the 1482 shifting computational demands of causal inference show up as the normative account is 1483 applied to different settings that scale differently in terms of the number of relevant events, 1484 variables and constraints on their potential relationships and functional forms. We might 1485 break process-level investigation into least two aspects. One aspect is understanding how 1486 temporal evidence is processed and compressed under our memory and computational 1487 constraints (Gallistel & Gibbon, 2000; Gong et al., 2023). The other aspect concerns how the causal hypothesis space and priors are constructed and searched (Bramley & Xu, 2023; 1489 Buchanan et al., 2010; Gong et al., 2024). This issue of understanding progressive search 1490 over causal hypotheses is especially important in the temporal setting given the strong now-or-never pressure on online computation relative to self-paced evidence setting. We 1492 hope that future studies can use this rational analysis of temporal causal structure 1493 induction to further explore human learning mechanisms, and how the computations 1494 involved interact with temporal scale (second, hours, days; Willett & Rottman, 2021). 1495

A final point why understanding temporal causal reasoning is vital is that it not only drives the identification of causal mechanisms among known events but also determines when and where we direct our attention. For example, we can actively anticipate and look for events that are as yet unobserved but are predicted by the existence of our causal theories. This is a critical part of scientific practice: with a good mechanistic

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model researchers can decide when to measure the outcomes of their experiments, such as when a drug's influence on a person should be most apparent. Making these choices in a theory-guided way seems almost as important for causal discovery as the technique of random-assignment experiments, yet has received far less attention. Future studies should investigate how scientists and laypeople make observations or conduct experiments that consider the information measurable from the intermediate processes, as well as from the final outcome variables. By doing so, we can build a more comprehensive picture of scientific discovery as well as everyday cognition.

1509 Conclusion

We inhabit a complex environment filled with continuous spatiotemporal causal 1510 dynamics. In order to form a practical causal understanding of this dynamic world, it 1511 seems essential for our minds to process temporal information effectively and efficiently. 1512 Despite fruitful empirical findings regarding how individuals behave in various time-based 1513 causal learning tasks, there is a lack of a unified theoretical framework to integrate 1514 behavioral predictions across all these tasks. In this paper, we present such a rational 1515 framework for causal induction based on the Poisson-Gamma statistical distribution 1516 families. We show how this framework aligns with human causal judgments. The 1517 framework grounds the basic philosophical intuitions about causality, and captures core 1518 qualitative empirical patterns that have long been seen in human learning studies. 1519 Quantitatively, the model is a good fit with human judgments across seven very different 1520 datasets. By laying out this framework, we take a key step towards understanding the 1521 computational task faced by humans and other agents when inducing a model of their 1522 environment. We hope the framework will serve as a benchmark for further investigation of 1523 the cognitive processes involved in generating and adapting causal representations, as well 1524 as how and why these may differ across different domains and timescales. 1525

1526 References

- Ahn, W.-k., Kalish, C. W., Medin, D. L., & Gelman, S. A. (1995). The role of covariation versus mechanism information in causal attribution. *Cognition*, 54(3), 299–352.
- Allan, L. G. (1980). A note on measurement of contingency between two binary variables in judgment tasks. *Bulletin of the Psychonomic Society*, 15(3), 147–149.
- Almaatouq, A., Griffiths, T. L., Suchow, J. W., Whiting, M. E., Evans, J., & Watts, D. J.
- (2022). Beyond playing 20 questions with nature: Integrative experiment design in the social and behavioral sciences. *Behavioral and Brain Sciences*, 1–55.
- Altmann, G., & Ekves, Z. (2019). Events as intersecting object histories: A new theory of event representation. *Psychological Review*, 126(6), 817–840.
- Anderson, J. R., & Sheu, C.-F. (1995). Causal inferences as perceptual judgments. *Memory*6 Cognition, 23(4), 510–524.
- Anderson, J. R. (1990). The adaptive character of thought. Psychology Press.
- Bass, I., Smith, K. A., Bonawitz, E., & Ullman, T. D. (2021). Partial mental simulation
   explains fallacies in physical reasoning. Cognitive Neuropsychology, 38(7-8), 413–424.
- Beller, A., & Gerstenberg, T. (in press). A counterfactual simulation model of causal
   language. Psychological Review.
- Bennett, J. (1987). Event causation: The counterfactual analysis. *Philosophical Perspectives*, 1, 367–386.
- Bramley, N. R., Dayan, P., Griffiths, T. L., & Lagnado, D. A. (2017). Formalizing
   neurath's ship: Approximate algorithms for online causal learning. *Psychological Review*, 124(3), 301.
- Bramley, N. R., Gerstenberg, T., Mayrhofer, R., & Lagnado, D. A. (2018). Time in causal structure learning. *Journal of Experimental Psychology: Learning, Memory, and Cognition*, 44(12), 1880–1910.
- Bramley, N. R., Gerstenberg, T., Mayrhofer, R., & Lagnado, D. A. (2019). Intervening in time. *Time and causality across the sciences*, 86–115.

- Bramley, N. R., Gerstenberg, T., Tenenbaum, J. B., & Gureckis, T. M. (2018). Intuitive experimentation in the physical world. *Cognitive Psychology*, 195, 9–38.
- Bramley, N. R., Lagnado, D. A., & Speekenbrink, M. (2015). Conservative forgetful
- scholars: How people learn causal structure through sequences of interventions.
- Journal of Experimental Psychology: Learning, Memory, and Cognition, 41(3), 708–731.
- Bramley, N. R., Mayrhofer, R., Gerstenberg, T., & Lagnado, D. A. (2017). Causal learning
- from interventions and dynamics in continuous time. In G. Gunzelmann, A. Howes,
- T. Tenbrink & E. J. Davelaar (Eds.), Proceedings of the 39th annual conference of
- the cognitive science society (pp. 150–155).
- Bramley, N. R., & Xu, F. (2023). Active inductive inference in children and adults: A constructivist perspective. *Cognition*, 238, 105471.
- Btesh, V. J., Bramley, N., Speekenbrink, M., & Lagnado, D. (in press). Less is more: Local focus in continuous time causal learning. *Journal of Experimental Psychology:*
- Learning, Memory, and Cognition.
- Buchanan, D., Tenenbaum, J., & Sobel, D. (2010). Edge replacement and nonindependence in causation. In S. Ohlsson & R. Catrambone (Eds.), *Proceedings of the 32nd annual* conference of the cognitive science society (pp. 919–924).
- Buehner, M. J., Cheng, P. W., & Clifford, D. (2003). From covariation to causation: A test
   of the assumption of causal power. Journal of Experimental Psychology: Learning,
   Memory, and Cognition, 29(6), 1119.
- Buehner, M. J., & May, J. (2003). Rethinking temporal contiguity and the judgement of
  causality: Effects of prior knowledge, experience, and reinforcement procedure. The
  Quarterly Journal of Experimental Psychology Section A, 56(5), 865–890.
- Buehner, M. J., & May, J. (2004). Abolishing the effect of reinforcement delay on human causal learning. Quarterly Journal of Experimental Psychology Section B, 57(2), 179–191.

- <sup>1580</sup> Buehner, M. J., & McGregor, S. (2006). Temporal delays can facilitate causal attribution:
- Towards a general timeframe bias in causal induction. Thinking & Reasoning, 12(4),
- 1582 353-378.
- <sup>1583</sup> Carroll, C., & Cheng, P. (2009). Preventative scope in causation. In N. A. Taatgen &
- H. van Rijn (Eds.), Proceedings of the 31th annual conference of the cognitive
- science society (pp. 833–838).
- 1586 Cartwright, N. (1994). Nature's capacities and their measurement. Oxford University Press.
- <sup>1587</sup> Cheng, P. W. (1997). From covariation to causation: A causal power theory. *Psychological*
- 1588 Review, 104(2), 367.
- <sup>1589</sup> Chung, S.-H. (1965). Effects of delayed reinforcement in a concurrent situation 1. Journal
- of the Experimental Analysis of Behavior, 8(6), 439–444.
- <sup>1591</sup> Clarke, R. (1946). An application of the poisson distribution. Journal of the Institute of
- 1592 Actuaries, 72(3), 481–481.
- Coenen, A., Rehder, B., & Gureckis, T. M. (2015). Strategies to intervene on causal
- systems are adaptively selected. Cognitive Psychology, 79, 102–133.
- <sup>1595</sup> Courville, A. C., Daw, N. D., & Touretzky, D. S. (2006). Bayesian theories of conditioning
- in a changing world. Trends in Cognitive Sciences, 10(7), 294–300.
- 1597 Craik, K. J. W. (1967). The nature of explanation (Vol. 445). CUP Archive.
- Davis, Z., Bramley, N. R., & Rehder, B. (2020). Causal structure learning in continuous
- systems. Frontiers in Psychology, 11, 244.
- Dean, T., & Kanazawa, K. (1989). A model for reasoning about persistence and causation.
- 1601 Computational Intelligence, 5(2), 142–150.
- Dennett, D. C. (1971). Intentional systems. The Journal of Philosophy, 68(4), 87–106.
- Eberhardt, F., Glymour, C., & Scheines, R. (2012). On the number of experiments sufficient
- and in the worst case necessary to identify all causal relations among n variables.
- Proceedings of the Twenty-First Conference on Uncertainty in Artificial Intelligence.

- Eckstein, M. K., & Collins, A. G. (2020). Computational evidence for hierarchically structured reinforcement learning in humans. *Proceedings of the National Academy*of Sciences, 117(47), 29381–29389.
- Einhorn, H. J., & Hogarth, R. M. (1986). Judging probable cause. *Psychological Bulletin*, 99(1), 3–19.
- Fernando, C. (2013). From blickets to synapses: Inferring temporal causal networks by observation. *Cognitive Science*, 37(8), 1426–1470.
- Fernbach, P. M., & Sloman, S. A. (2009). Causal learning with local computations. *Journal*of Experimental Psychology: Learning, Memory, and Cognition, 35(3), 678–693.
- <sup>1615</sup> Fink, D. (1997). A compendium of conjugate priors (tech. rep.).
- https://www.johndcook.com/CompendiumOfConjugatePriors.pdf.
- Gallistel, C. R. (2021). Robert rescorla: Time, information and contingency. Revista de

  Historia de la Psicologia, 42(1), 7–21.
- Gallistel, C. R., Craig, A. R., & Shahan, T. A. (2014). Temporal contingency. *Behavioural Processes*, 101, 89–96.
- Gallistel, C. R., Craig, A. R., & Shahan, T. A. (2019). Contingency, contiguity, and causality in conditioning: Applying information theory and weber's law to the assignment of credit problem. *Psychological Review*, 126(5), 761–773.
- Gallistel, C. R., & Gibbon, J. (2000). Time, rate, and conditioning. *Psychological Review*, 1625 107(2), 289–344.
- Gallistel, C. R., & Shahan, T. A. (2024). Time-scale invariant contingency yields one-shot reinforcement learning despite extremely long delays to reinforcement. *Proceedings*of the National Academy of Sciences, 121(30), e2405451121.
- Gallistel, C. R., & Wilkes, J. T. (2016). Minimum description length model selection in associative learning. Current Opinion in Behavioral Sciences, 11, 8–13.
- Garcia, J., Ervin, F. R., & Kölling, R. A. (1966). Learning with prolonged delay of reinforcement. *Psychonomic Science*, 5(3), 121–122.

- Gershman, S. J. (2015). A unifying probabilistic view of associative learning. *PLoS* 1633 Computational Biology, 11(11), e1004567. 1634
- Gershman, S. J. (2017). Reinforcement learning and causal models. In M. Waldmann (Ed.), 1635 The oxford handbook of causal reasoning (pp. 295–306). Oxford University Press.
- Gershman, S. J. (2024). Rate estimation revisited. *PsyArXiv*. 1637
- Gershman, S. J., Blei, D. M., & Niv, Y. (2010). Context, learning, and extinction. 1638
- Psychological Review, 117(1), 197. 1639

1636

- Gershman, S. J., Norman, K. A., & Niv, Y. (2015). Discovering latent causes in 1640 reinforcement learning. Current Opinion in Behavioral Sciences, 5, 43–50.
- Gerstenberg, T., Goodman, N. D., Lagnado, D. A., & Tenenbaum, J. B. (2021). A 1642 counterfactual simulation model of causal judgments for physical events. 1643
- *Psychological Review*, 128(5), 936–975. 1644
- Gerstenberg, T., & Stephan, S. (2021). A counterfactual simulation model of causation by 1645 omission. Cognition, 216, 104842. 1646
- Gettier, E. L. (1963). Is justified true belief knowledge? Analysis, 23(6), 121–123. 1647
- Gong, T. (2023). Causal induction in time (Doctoral dissertation). University of Edinburgh. 1648
- Gong, T., & Bramley, N. R. (2020). What you didn't see: Prevention and generation in 1649 continuous time causal induction. In S. Denison, M. Mack, Y. Xu & B. Armstrong 1650 (Eds.), Proceedings of the 42ed annual conference of the cognitive science society 1651
- (pp. 2908–2914). 1652
- Gong, T., & Bramley, N. R. (2022). Intuitions and perceptual constraints on causal 1653
- learning from dynamics. In J. Culbertson, A. Perfors, H. Rabagliati & 1654
- V. Ramenzoni (Eds.), Proceedings of the 44th Annual Meeting of the Cognitive 1655 Science Society (pp. 1455–1461). 1656
- Gong, T., & Bramley, N. R. (2023). Continuous time causal structure induction with 1657 prevention and generation. Cognition, 240, 105530. 1658

- Gong, T., & Bramley, N. R. (2024). Evidence from the future. *Journal of Experimental*Psychology: General, 153(3), 864–872.
- Gong, T., Gerstenberg, T., Mayrhofer, R., & Bramley, N. R. (2023). Active causal structure learning in continuous time. *Cognitive Psychology*, 140, 101542.
- Gong, T., Valentin, S., Lucas, C. G., & Bramley, N. R. (2024). Paradoxical parsimony:
- How latent complexity favors theory simplicity. In L. Samuelson, S. Frank,
- M. Toneva, A. Mackey & E. Hazeltine (Eds.), Proceedings of the 46th annual conference of the cognitive science society (pp. 2428–3434).
- Goodman, N. (1983). Fact, fiction, and forecast. Harvard University Press.
- Grabenhorst, M., Maloney, L. T., Poeppel, D., & Michalareas, G. (2021). Two sources of uncertainty independently modulate temporal expectancy. *Proceedings of the National Academy of Sciences*, 118(16), e2019342118.
- Grabenhorst, M., Michalareas, G., Maloney, L. T., & Poeppel, D. (2019). The anticipation of events in time. *Nature Communications*, 10(1), 5802.
- Greville, W. J., & Buehner, M. J. (2007). The influence of temporal distributions on causal induction from tabular data. *Memory & Cognition*, 35(3), 444–453.
- Greville, W. J., & Buehner, M. J. (2010). Temporal predictability facilitates causal learning. *Journal of Experimental Psychology: General*, 139(4), 756–771.
- Greville, W. J., & Buehner, M. J. (2016). Temporal predictability enhances judgements of causality in elemental causal induction from both observation and intervention.
- Quarterly Journal of Experimental Psychology, 69(4), 678–697.
- Greville, W. J., Buehner, M. J., & Johansen, M. K. (2020). Causing time: Evaluating

  causal changes to the when rather than the whether of an outcome. *Memory & Cognition*, 48, 200–211.
- Griffiths, T. L. (2020). Understanding human intelligence through human limitations.

  Trends in Cognitive Sciences, 24(11), 873–883.

- Griffiths, T. L., & Tenenbaum, J. B. (2005). Structure and strength in causal induction.
- 1686 Cognitive Psychology, 51(4), 334–384.
- Griffiths, T. L., & Tenenbaum, J. B. (2007). From mere coincidences to meaningful discoveries. *Cognition*, 103(2), 180–226.
- Griffiths, T. L., & Tenenbaum, J. B. (2009). Theory-based causal induction. *Psychological Review*, 116(4), 661–716.
- Hagmayer, Y., & Waldmann, M. R. (2002). How temporal assumptions influence causal judgments. *Memory & Cognition*, 30(7), 1128–1137.
- Halpern, J. Y. (2016). Actual causation. MIT Press.
- Hamou, N., Gershman, S. J., & Reddy, G. (2025). Reconciling time and prediction error
   theories of associative learning. bioRxiv, 2025–01.
- Hume, D. (1740). A treatise of human nature. Oxford University Press (2000 reprint).
- Icard, T. F., Kominsky, J. F., & Knobe, J. (2017). Normality and actual causal strength.
   Cognition, 161, 80–93.
- Kalmbach, A., Chun, E., Taylor, K., Gallistel, C. R., & Balsam, P. D. (2019).
- Time-scale-invariant information-theoretic contingencies in discrimination learning.
- Journal of Experimental Psychology: Animal Learning and Cognition, 45(3),
- 1702 280–289.
- Kim, N. S., & Ahn, W.-k. (2002). Clinical psychologists' theory-based representations of
  mental disorders predict their diagnostic reasoning and memory. *Journal of*Experimental Psychology: General, 131(4), 451–476.
- Kolvoort, I., Davis, Z. J., Rehder, B., & van Maanen, L. (2025). Models of variability in probabilistic causal judgments. *Computational Brain & Behavior*, 8, 162–188.
- Lagnado, D. A., & Sloman, S. (2002). Learning causal structure. In W. Gray & C. Schunn (Eds.), Proceedings of the 24th annual meeting of the cognitive science society (pp. 560–565).

- Lagnado, D. A., & Sloman, S. A. (2004). The advantage of timely intervention. *Journal of*
- Experimental Psychology: Learning, Memory, and Cognition, 30(4), 856–876.
- Lagnado, D. A., & Sloman, S. A. (2006). Time as a guide to cause. Journal of
- Experimental Psychology: Learning, Memory, and Cognition, 32(3), 451–460.
- Lagnado, D. A., & Speekenbrink, M. (2010). The influence of delays in real-time causal
- learning. The Open Psychology Journal, 3(1), 184–195.
- Lagnado, D. A., Waldmann, M. R., Hagmayer, Y., & Sloman, S. A. (2007). Beyond
- covariation. In A. Gopnik & L. Schulz (Eds.), Causal learning: Psychology,
- philosophy, and computation (pp. 154–172). Oxford University Press.
- Lake, B. M., Ullman, T. D., Tenenbaum, J. B., & Gershman, S. J. (2017). Building
- machines that learn and think like people. Behavioral and Brain Sciences, 40, e253.
- Lauritzen, S. L., & Richardson, T. S. (2002). Chain graph models and their causal
- interpretations. Journal of the Royal Statistical Society Series B: Statistical
- Methodology, 64(3), 321–348.
- Libet, B. (2009). Mind time: The temporal factor in consciousness. Harvard University
- Press.
- Lieder, F., & Griffiths, T. L. (2020). Resource-rational analysis: Understanding human
- cognition as the optimal use of limited computational resources. Behavioral and
- 1729 Brain Sciences, 43, 1–60.
- Lombrozo, T. (2010). Causal–explanatory pluralism: How intentions, functions, and
- mechanisms influence causal ascriptions. Cognitive Psychology, 61(4), 303–332.
- Lu, H., Yuille, A. L., Liljeholm, M., Cheng, P. W., & Holyoak, K. J. (2008). Bayesian
- generic priors for causal learning. Psychological Review, 115(4), 955–984.
- Luce, R. D. (1959). *Individual choice behavior*. Wiley.
- Ludwin-Peery, E., Bramley, N. R., Davis, E., & Gureckis, T. M. (2021). Limits on
- simulation approaches in intuitive physics. Cognitive Psychology, 127, 101396.
- Malthus, T. R. (1872). An essay on the principle of population. Reeves & Turner.

- Markant, D. B., & Gureckis, T. M. (2014). Is it better to select or to receive? learning via active and passive hypothesis testing. *Journal of Experimental Psychology: General*, 143(1), 94–122.
- Marr, D. (1982). Vision: A computational investigation into the human representation and processing of visual information. MIT Press.
- Mendelson, R., & Shultz, T. R. (1976). Covariation and temporal contiguity as principles
   of causal inference in young children. *Journal of Experimental Child Psychology*,
   22(3), 408–412.
- Mnih, V., Kavukcuoglu, K., Silver, D., Rusu, A. A., Veness, J., Bellemare, M. G.,
- Graves, A., Riedmiller, M., Fidjeland, A. K., Ostrovski, G., et al. (2015).
- Human-level control through deep reinforcement learning. *Nature*, 518 (7540), 529–533.
- Myerson, J., & Green, L. (1995). Discounting of delayed rewards: Models of individual choice. *Journal of the Experimental Analysis of Behavior*, 64(3), 263–276.
- Nikolic, M., & Lagnado, D. A. (2015). There aren't plenty more fish in the sea: A causal network approach. *British Journal of Psychology*, 106(4), 564–582.
- O'Neill, K., Henne, P., Bello, P., Pearson, J., & De Brigard, F. (2022). Confidence and gradation in causal judgment. *Cognition*, 223, 105036.
- O'Neill, K., Henne, P., Pearson, J., & De Brigard, F. (2024). Modeling confidence in causal judgments. *Journal of Experimental Psychology: General*, 153(8), 2142–2159.
- Pacer, M. (2016). Mind as theory engine: Causation, explanation and time
  (Doctoral dissertation). UC Berkeley.
- Pacer, M., & Griffiths, T. L. (2012). Elements of a rational framework for continuous-time causal induction. In N. Miyake, D. Peebles & R. P. Cooper (Eds.), *Proceedings of the 34th annual conference of the cognitive science society* (pp. 833–838).
- Pacer, M., & Griffiths, T. L. (2015). Upsetting the contingency table: Causal induction over sequences of point events. In D. C. Noelle, R. Dale, A. S. Warlaumont,

1780

- J. Yoshimi, T. Matlock, C. D. Jennings & P. P. Maglio (Eds.), Proceedings of the 1765 37th annual conference of the cognitive science society (pp. 1805–1810). 1766
- Pearl, J. (2000). Causality. Cambridge University Press (2009 reprint). 1767
- Perales, J. C., & Shanks, D. R. (2007). Models of covariation-based causal judgment: A 1768 review and synthesis. Psychonomic Bulletin & Review, 14, 577–596. 1769
- Pishro-Nik, H. (2014). Introduction to probability, statistics, and random processes. Kappa 1770 Research, LLC Blue Bell, PA, USA. 1771
- Rehder, B. (2017). Reasoning with causal cycles. Cognitive Science, 41, 944–1002. 1772
- Rehder, B., Davis, Z. J., & Bramley, N. (2022). The paradox of time in dynamic causal 1773 systems. *Entropy*, 24(7), 863. 1774
- Rescorla, R. A. (1968). Probability of shock in the presence and absence of cs in fear 1775 conditioning. Journal of Comparative and Physiological Psychology, 66(1), 1–5. 1776
- Rescorla, R. A., & Wagner, A. R. (1972). A theory on pavlovian conditioning: Variations in 1777 the effectiveness of reinforcement and nonreinforcement. In A. Black & W. Prokasy 1778 (Eds.), Classical conditioning ii: Current theory and research (pp. 64–99). Appleton 1779 Century Crofts.
- Ross, L. (2024). Causal constraints in the life and social sciences. *Philosophy of Science*, 1781 *91*(5), 1068–1077. 1782
- Rottman, B. M. (2016). Searching for the best cause: Roles of mechanism beliefs, 1783
- autocorrelation, and exploitation. Journal of Experimental Psychology: Learning, 1784 Memory, and Cognition, 42(8), 1233. 1785
- Rottman, B. M., & Hastie, R. (2014). Reasoning about causal relationships: Inferences on 1786 causal networks. Psychological Bulletin, 140(1), 109-139. 1787
- Rottman, B. M., & Keil, F. C. (2012). Causal structure learning over time: Observations 1788 and interventions. Cognitive Psychology, 64 (1-2), 93–125. 1789
- Scheines, R. (1997). An introduction to causal inference. Carnegie Mellon University. 1790

- Schlottmann, A., Cole, K., Watts, R., & White, M. (2013). Domain-specific perceptual causality in children depends on the spatio-temporal configuration, not motion onset. Frontiers in Psychology, 4, 365.
- Schultz, W. (2015). Neuronal reward and decision signals: From theories to data.

  Physiological Reviews, 95(3), 853–951.
- Schultz, W., Dayan, P., & Montague, P. R. (1997). A neural substrate of prediction and reward. Science, 275(5306), 1593–1599.
- Shanks, D. R., & Dickinson, A. (1991). Instrumental judgment and performance under variations in action-outcome contingency and contiguity. *Memory & Cognition*, 1904, 353–360.
- Shanks, D. R., Pearson, S. M., & Dickinson, A. (1989). Temporal contiguity and the

  judgement of causality by human subjects. *The Quarterly Journal of Experimental*Psychology, 41(2), 139–159.
- Simon, H. A. (1982). Models of bounded rationality: Empirically grounded economic reason.

  MIT press.
- Sloman, S. A. (2005). Causal models: How people think about the world and its alternatives.

  Oxford University Press.
- 1808 Sloman, S. A., & Lagnado, D. A. (2005). Do we "do"? Cognitive Science, 29, 5–39.
- Sloman, S. A., Love, B. C., & Ahn, W.-K. (1998). Feature centrality and conceptual coherence. *Cognitive Science*, 22(2), 189–228.
- Soo, K. W., & Rottman, B. M. (2018). Causal strength induction from time series data.

  Journal of Experimental Psychology: General, 147(4), 485–513.
- Stephan, S., Mayrhofer, R., & Waldmann, M. R. (2020). Time and singular causation—a computational model. *Cognitive Science*, 44 (7), e12871.
- Stephan, S., Placì, S., & Waldmann, M. R. (2021). Evaluating general versus singular causal prevention. In T. Fitch, C. Lamm, H. Leder & K. Teßmar-Raible (Eds.),

- Proceedings of the 43th annual conference of the cognitive science society (pp. 1402–1408).
- Steyvers, M., Tenenbaum, J. B., Wagenmakers, E.-J., & Blum, B. (2003). Inferring causal networks from observations and interventions. *Cognitive Science*, 27(3), 453–489.
- Talmy, L. (1988). Force dynamics in language and cognition. *Cognitive Science*, 12(1), 49–100.
- Tarpy, R. M., & Sawabini, F. L. (1974). Reinforcement delay: A selective review of the last decade. *Psychological Bulletin*, 81(12), 984–997.
- Tenenbaum, J. B., Griffiths, T. L., & Kemp, C. (2006). Theory-based bayesian models of inductive learning and reasoning. *Trends in Cognitive Sciences*, 10(7), 309–318.
- Uhlenbeck, G. E., & Ornstein, L. S. (1930). On the theory of the brownian motion.

  Physical Review, 36(5), 823–841.
- Ullman, T. D., Stuhlmüller, A., Goodman, N. D., & Tenenbaum, J. B. (2018). Learning physical parameters from dynamic scenes. *Cognitive Psychology*, 104, 57–82.
- Valentin, S., Bramley, N. R., & Lucas, C. G. (2020). Learning hidden causal structure from temporal data. In S. Denison, M. Mack, Y. Xu & B. Armstrong (Eds.), *Proceedings* of the 42ed annual conference of the cognitive science society (pp. 1906–1912).
- Valentin, S., Bramley, N. R., & Lucas, C. G. (2022). Discovering common hidden causes in sequences of events. *Computational Brain & Behavior*, 1–23.
- Valleron, A.-J., Boelle, P.-Y., Will, R., & Cesbron, J.-Y. (2001). Estimation of epidemic
   size and incubation time based on age characteristics of vcjd in the united kingdom.
   Science, 294 (5547), 1726–1728.
- Waldmann, M. R., & Hagmayer, Y. (2005). Seeing versus doing: Two modes of accessing
   causal knowledge. Journal of Experimental Psychology: Learning, Memory, and
   Cognition, 31(2), 216–227.
- Wang, J. X. (2021). Meta-learning in natural and artificial intelligence. Current Opinion in
   Behavioral Sciences, 38, 90–95.

- Willett, C. L., & Rottman, B. M. (2021). The accuracy of causal learning over long timeframes: An ecological momentary experiment approach. *Cognitive Science*, 45(7), e12985.
- Wolff, P. (2007). Representing causation. Journal of Experimental Psychology: General,
   136(1), 82–111.
- Woodward, J. (2021). Causation with a human face: Normative theory and descriptive psychology. Oxford University Press.
- Wu, M., & Cheng, P. W. (1999). Why causation need not follow from statistical association: Boundary conditions for the evaluation of generative and preventive causal powers. *Psychological Science*, 10(2), 92–97.
- Yeung, S., & Griffiths, T. L. (2015). Identifying expectations about the strength of causal relationships. *Cognitive Psychology*, 76, 1–29.
- Zhang, Y., & Rottman, B. M. (2023). Causal learning with interrupted time series data.
   Judgment and Decision Making, 18, e30.
- Zhang, Y., & Rottman, B. M. (2024). Causal learning with delays up to 21 hours.

  Psychonomic Bulletin & Review, 31(1), 312–324.

### Appendix A

#### Prevention Causation

#### 1860 The event-based scheme

Considering preventative causation generally increases the space of possible 1861 explanations and would hence affect the likelihood calculation for specific actual pathways. 1862 Concretely, for each observed effect e, we must jointly evaluate the probability that (1) it 1863 was caused by its presumed generative cause event g as well as that (2) it was not 1864 prevented by any of the set of presumed preventative cause events p. Hidden (expected but 1865 unobserved) effects also contribute to the likelihood wherever we do not observe an 1866 expected effect of a generative cause. This could be due to (1) the generative cause failing 1867 to produce that effect, (2) that effect being prevented, or (3) that effect not having 1868 occurred yet: 1869

$$P(\mathbf{z}|s; \mathbf{w}) = \prod_{g \to e \in \mathbf{z}} \underbrace{w_g \cdot P_d(t_e - t_g | \alpha, \beta) \cdot (1 - P_p(e))}_{\text{Observed effects must have been generated and not prevented}}$$

$$\times \prod_{g \to h \in \mathbf{z}} \underbrace{(1 - w_g) + w_g \cdot P_d(t_h > t_{end} | \alpha, \beta) + w_g \cdot P_p(h)}_{\text{Unobserved expected effects must have failed or been prevented, or be still-to-occur}}$$
(A1)

The event-based scheme provides also allows for flexibility in dealing with preventative causation  $P_p(e)$  (the probability that e should have been prevented) based on different mechanisms of prevention. For instance, a preventative cause might block an effect from occurring at all for a specific time window. Alternatively, it might block the subsequent N effects from occurring before being "used up". A preventative cause might block all effects indiscriminately (e.g., operate on the effect variable), or selectively block effects from a particular cause (e.g., operate on the edge between two variables; Carroll & Cheng, 2009; Gerstenberg & Stephan, 2021).

# The rate-based scheme

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When a cause is generative, we expect the rate of its effect to temporarily increase, whereas we expect preventative causes to temporarily decrease the rate of their effects.

Intuitively, a preventative causal influence can be thought of as defeating some of

the effects that would otherwise have occurred, meaning that it will have a proportional

effect on the rate. As such, we assume preventative causes decrease the effect rate by a

proportion ranging from 0 to a maximum level of  $\xi$  (0 <  $\xi$  < 1). A preventative influence

 $^{1885}$  can also follow an incubation-decay process and be represented by a function of time

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$$f(\xi,t) = \xi \cdot \frac{P_d(t|\alpha,\beta)}{P_d(\frac{\alpha-1}{\beta}|\alpha,\beta)}$$
.

This means preventative causation can be viewed as "thinning" processes that selectively filter out some effect events with a probability of  $\xi'$ . This contrasts with the natural way to think of generative causation as "superposition" where more events are added to the timeline. Combining multiple causes with a base rate of  $\lambda_0$ , the expected effect rate  $f(\lambda, t)$  at the time unit t can be represented similar to the noisy-OR and noisy-AND-NOT principles by accounting for superposition and thinning as follows:

$$f(\lambda, t) = (\lambda_0 + \sum_{i \in \mathbf{g}} f(\lambda_i, t)) \prod_{j \in \mathbf{p}} (1 - f(\xi_j, t))$$
(A2)

### Appendix B

# Time-scale Invariance Simulation

To demonstrate when the time-scale invariance property applies, we synthesize three 1893 conditions, each with causal delays  $m_u = \{10, 12, 14, 16, 18\}$ . The "no-match" condition 189 follows the procedure in main text to use a fixed observation duration of 300 time units, 1895 with  $w_c = 1$ ,  $k_b = 3$ , and  $i_u = 1$  (similar to the main text we apply the number of cause 1896 events  $k_c = 5$  for the event-based model and  $k_c = 10$  for the rate-based model). In the 1897 "baserate-match" condition, the observation duration is scaled according to the causal 1898 delay. We retain 300 time units when  $m_u = 10$ , but adjust to 360, 420, 480, and 540 time 1899 units for the other delays, respectively. Causal events and base rate events are sampled 1900 within these new observation durations, ensuring that each causal delay has the same 1901 number of observed events  $(k_c \text{ and } k_b)$ , while the rate of background effect events scales 1902 with the causal delay. Specifically, for  $m_u = 10$  the baserate is 3 per 300 time units (an 1903 average delay between baserate effects of 75 time units), and for  $m_u = 12$  the baserate 1904 adjusts to 3 per 360 time units (with an average delay between base rate effects of 90 time 1905 units). The prior is also scaled accordingly. For example, if the observation duration is 360, 1906 we use  $\mu \sim U(0,360)$  and  $\mu_b \sim U(0,360)$  instead. In the "all-match" condition, we further 1907 scale the delay variance by setting  $i_u = \{1, 1.2, 1.4, 1.6, 1.8\}$  for each causal delay 1908 respectively. 1909

As shown in Figure B1, the tendency to favor a causal structure can remain at the same level even when the causal delay increases, as long as the environment is time-scale invariant (i.e., the observation duration, the base rate, and the delay variance are all scaled/matched according to the length of the causal delays). The tendency to favor the causal hypothesis decreases as the causal delay increases if the contextual factors do not scale, and it increases with the causal delay if the baserate-relevant factors are scaled but the delay variance is not.

The other intuitive way to think why the time-scale invariance would exist is to

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think, if we change the length scale of an event sequences from minutes to hours (and then the baserate will change from k per minute to k per hour). Accordingly, to create equally weak priors, we can for example, apply a uniform prior from 0 to 100 minutes (assuming the upper bound here is larger than the practically possible causal delay) in the short-delay condition and from 0 to 100 hours in the long-delay condition. In this case, the time unit becomes the only difference all calculations, but this does not matter because the time unit is pre-set arbitrarily before any calculation.

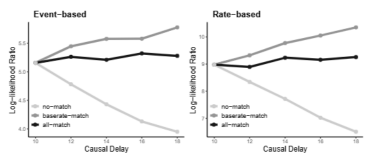


Figure B1

The log-likelihood ratio changes given the causal delays. A ratio above zero indicates that the model favors  $S_1$  (the causal structure) over  $S_0$  (the base rate structure).

### Appendix C

# The Persistent Effects of Delay Variance

Greville and Buehner (2010) found that the preference for unvaried delays persisted when the learning duration was doubled from 2 min to 4 min, which cannot be well explained by 1926 a difference in learning rates under the associative learning model (Chung, 1965; Rescorla, 1927 1968). We here show that this result is easily captured by the Bayesian framework. We 1928 simulate different learning durations [100, 150, 200] and set the cause events to occur every 1929 25 time units on average, with baserate effect events occurring every 50 time units on 1930 average. Each cause event generates an effect event with a delay sampled from 1931  $U(m_u - i_u, m_u + i_u)$  and a probability of  $w_c$ . As shown in Figure C1, when the average 1932 delay length  $m_u$  and the causal probability  $w_c$  are fixed, the effect of delay variance does 1933 not decrease as the learning duration increases. It instead increases as the evidence 1934 accumulates.

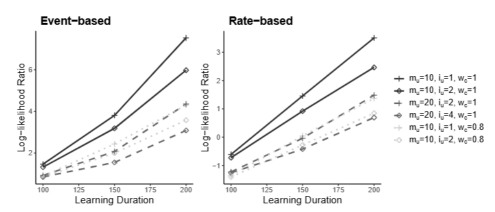


Figure C1

The log-likelihood ratio changes given the learning duration. A ratio above zero indicates that the model favors  $S_1$  (the causal structure) over  $S_0$  (the base rate structure).