Counterfactual Resimulation for Causal Analysis of Rule-Based Models

Jonathan Laurent\textsuperscript{1}, Jean Yang\textsuperscript{1}, Walter Fontana\textsuperscript{2}

\textsuperscript{1} Carnegie Mellon University \hspace{1em} \textsuperscript{2} Harvard Medical School

\{jonathan.laurent, jyang2+\}@cs.cmu.edu, walter_fontana@hms.harvard.edu

Abstract

Models based on rules that express local and heterogeneous mechanisms of stochastic interactions between structured agents are an important tool for investigating the dynamical behavior of complex systems, especially in molecular biology. Given a simulated trace of events, the challenge is to construct a causal diagram that explains how a phenomenon of interest occurred. Counterfactual analysis can provide distinctive insights, but its standard definition is not applicable in rule-based models because they are not readily expressible in terms of structural equations. We provide a semantics of counterfactual statements that addresses this challenge by sampling counterfactual trajectories that are probabilistically as close to the factual trace as a given intervention permits them to be. We then show how counterfactual dependencies give rise to explanations in terms of relations of enablement and prevention between events.

1 Introduction

Rule-based modeling languages for molecular biology, such as Kappa \cite{Danos2007} and BioNetGen \cite{Harris2016}, or organic chemistry, such as Mod \cite{Andersen2016}, can be used to write mechanistic models of complex reaction systems. These approaches consider entities that have a structure, and make a distinction between the transformation of a structure fragment (a pattern) specified by a rule and the reaction resulting from the application of the rule to a combination of entities contextualizing the fragment. The structure of bio-molecular entities is represented as a graph, and a rule is a graph-rewrite directive with a rate constant that determines its propensity to apply. The stochastic simulation of a rule collection generates a time series of rule applications—henceforth events—that might reach a state of interest in processes like the assembly of a molecular machine, the activation of a transcription factor, or the synthesis of a specific compound.

While rule-based models provide compactness, transparency, and the ability of handling combinatorial complexity, the perhaps most significant advantage lies in their suitability for causal analysis. This is because such analysis proceeds at the level of rules, not reactions, thereby avoiding contamination with context that defines a reaction yet is irrelevant to the application of the underlying rule. Due to the concurrent nature of events, it is typically far from obvious how a sequence attained a particular outcome. Biologists often refer to a causal account or explanation as a “pathway”, but have no formal framing for it.

Prior work in causal analysis \cite{Danos2012, Danos2007} takes advantage of rule structure to (i) compress a simulation trace into a minimal subset of events that are necessary and jointly sufficient to replicate the outcome of interest and (ii) highlight causal influences between events, exposing the extent of concurrency. Such analysis is performed on a sample of traces to the outcome, thus recovering the salient pathways as those that are statistically favored by the dynamics. This approach, however, suffers from two drawbacks. First, the focus on necessity in step (i) neglects events that are kinetically critical (in that they dramatically increase the probability of observing the outcome), yet are not logically necessary for achieving it. Second, step (ii) is limited to a narrow notion of causal influence that we may call enablement. Put simply, an event \( a \) (directly) enables event \( b \), if \( a \) modifies the state of the world so as to satisfy the requirements for \( b \) to occur. This positively tinted version of influence is blind to the ubiquitous role of inhibitory interactions in molecular biology. Indeed, an event \( a \) may cause an event \( b \) without (transitively) enabling it, but instead by preventing another event \( c \) that would have prevented \( b \). Clearly, uncovering such an explanatory narrative is challenging because it involves an event, \( c \) in this case, that did not occur in a simulation trace.

We here propose an approach that complements the existing causal analysis of event series generated from rule-based models by using counterfactual reasoning to answer questions of the kind: \textit{Had event} \( e_2 \) \textit{not occurred, would event} \( e_2 \) \textit{have happened?} Our contributions may be summarized as follows.

1. We provide a semantics for counterfactual statements in the context of rule-based models, where the standard definition of counterfactuals based on structural equations \cite{Pearl2009, Halpern2016} does not apply.

2. We show how such statements can be evaluated by sampling \textit{counterfactual traces} that are meant to probabilistically “hug” a given (factual) trace as much as an external...
intervention permits them to. To this end, we introduce an algorithm to generate counterfactual traces and provide an efficient implementation for the Kappa language.

3. We show how counterfactual dependencies between events can be systematically explained in terms of enablement and prevention relations that are more in line with biological reasoning.

2 Motivating Example
We provide some background on Kappa and introduce a toy example motivating the need for counterfactual reasoning in analyzing the causal structure of simulation traces.

2.1 Some Background on Kappa
Proteins are complex molecular machines that reversibly tag one another with molecular flags and reversibly bind each other to form transient associations. In this way, proteins come to have “state” that can control the interactions they engage in. In Kappa, a protein is modeled as an abstract agent with a name that designates its type and a signature of distinguishable sites at which it can be tagged or bound by other agents. A site can bind at most one agent at a time and must be in a definite state.

In our illustrative example, we consider two types of agents for which we use biological nomenclature. One type, $S$, is a substrate that receives a tag known as a phosphate group in a phosphorylation interaction. The other, $K$, is a kinase that phosphorylates $S$. Agents of both types feature a binding site at which they can bind one another and a site that can be in one of two possible states: unphosphorylated or phosphorylated. Thus, agents of type $K$ also have a phosphorylation state, but for the sake of simplicity we will have them acquire it “spontaneously”.

A mixture is a multiset of agents whose state at each site is fully specified. A mixture represents the state of a system and can be thought of as a (potentially large) graph consisting of many connected components. In a mixture, agents of the same type are distinguished by a unique global identifier.

![Figure 1: A mixture graph. Sites are here identified by their position on agents instead of a name. A grey site indicates a phosphorylated state. Numbers are global agent identifiers in the mixture.](image)

Interactions between agents are modeled by local rewriting rules. A rule $r$ is defined by a triple $(L_r, R_r, \lambda_r)$, where $L_r$ is the left-hand side specifying a pattern (the pre-condition), $R_r$ the right-hand side (or post-condition), and $\lambda_r$ a firing rate. The basic idea is that the “location” at which the mixture matches $L_r$ is rewritten in place by $R_r$, changing the state of the system. (Technically, a rule also requires a function from agents in $L_r$ to agents in $R_r$ to make the rewrite unambiguous.)

In our toy model, agents are subject to the rules depicted in Figure 2. Rule $b$ states that kinases and substrates can bind, provided their requisite binding sites are free (unbound). Note that $L_b$ is a pattern: It omits mentioning the sites that carry phosphorylation state, which are therefore not considered when matching the mixture to $L_b$. Rules $u$ and $u^*$ define unbinding events that depend on the phosphorylation state of the kinase $K$. The distinction is motivated by kinetics: Rule $u$ fires at a much faster rate than $u^*$. Rule $p$ specifies that a substrate can be phosphorylated when it is bound to a kinase. For the sake of simplicity, we model the phosphorylation of a kinase as a spontaneous process (rule $pk$).

By virtue of the $\lambda_r$, the rules of a model, together with an initial mixture, constitute a dynamical system that we describe shortly. We do so in a slightly nonstandard way by introducing the auxiliary concept of an event template. This is to prepare for the insight in section 3.1 that the stochastic and deterministic aspects of a model’s dynamics can be cleanly separated, thus enabling counterfactual analysis.

An event template is a pair $(r, \xi)$ where $r$ is a rule and $\xi$ a function from agents in $L_r$ to global identifiers. We say that $(r, \xi)$ is realizable in mixture $m$ if the global identifiers assigned by $\xi$ exist in $m$ and the agents bearing them match $L_r$. In this case, we write $m \vdash (r, \xi)$ and call $\xi$ an embedding of $L_r$ into $m$: $\text{EMB}_r(m) \triangleq \{ \xi : m \vdash (r, \xi) \}$. Whenever $m \vdash (r, \xi)$, we write $m \cdot (r, \xi)$ the mixture obtained by realizing $(r, \xi)$, i.e. by rewriting the agents with identifiers in the codomain of $\xi$ into $R_r$. The realization of an event template at a particular time creates an (actual) event, formally defined as a pair $(e, t)$ with $e$ an event template and $t$ its time of realization.

A model induces a continuous-time Markov chain (CTMC) over the set of reachable mixtures, where state $m$ transitions to state $m \cdot (r, \xi)$ at rate $\lambda_r$ for every event template $(r, \xi)$ that is realizable in $m$. The rate of leaving state $m$ by an application of rule $r$ is called the activity $\alpha_r(m)$ of rule $r$ in mixture $m$ and is equal to the product of the rule’s firing rate by the number of embeddings of $L_r$ into $m$: $\alpha_r(m) = \lambda_r | \text{EMB}_r(m) |$. For example, in Figure 1, rule $b$ has activity $2\lambda_0$ and rule $u$ has activity 0. The total activity of a mixture is defined as $\alpha(m) = \sum_r \alpha_r(m) = \sum_r \lambda_r | \text{EMB}_r(m) |$. 

![Figure 2: A motivating toy model. Sites not mentioned in a rule are left unchanged by it. Firing rates are not specified, but dotted (solid) arrows indicate slow (fast) reactions ($\lambda_u \gg \lambda_u^* \approx \lambda_p$).](image)
The CTMC induced by a model can be simulated with the Doob-Gillespie algorithm [Gillespie, 1977], which loops over the following steps: (1) draw a time interval δ to the next event from an exponential distribution with parameter \( \alpha(m) \) and increment the simulated system time by δ, (2) draw a rule \( r \) with probability \( \alpha_r(m) / \alpha(m) \) and (3) pick uniformly an embedding \( \xi \in \text{EMB}(m) \) of \( L_r \) into \( m \) and realize the event template \( \langle r, \xi \rangle \). This algorithm is efficiently implemented for rule-based models in Kappa as described in [Danos et al., 2007b; Boullier et al., 2017]. It outputs a sequence of events, called a trace.

2.2 Where Existing Analysis Falls Short
Consider our toy model and assume, for the sake of illustration, an initial mixture \( I \) with only a single kinase and a single substrate whose sites are unbound and unphosphorylated. We then ask: Starting from \( I \), how is \( p \) typically achieved? We are not merely looking for an account of reachability but for a causal explanation, i.e. a collection of events connected by causal influences.

For example, a simulation might produce the following trace (events are labeled by the rules that induced them):

\[
\begin{align*}
&b, \ u, \ pk, \ b, \ p, \ u^*, \ \cdots
\end{align*}
\]  

(1)

Current techniques [Danos et al., 2012; Danos et al., 2007a] generate a causal account by first computing a sub-trace of (1) that is (i) valid in the sense that each of its events can be triggered in turn starting from the initial mixture and (ii) minimal in the sense that none of its valid sub-traces features \( p \). The relations of enabling among events in the minimal sub-trace yield a directed acyclic graph. Although trivial in our toy example, minimization is an NP-hard problem. Carrying out this approach on (1), one notes that the first occurrence of \( b \) sets the necessary conditions for \( p \), but these are subsequently undone by \( u \) only for the second occurrence of \( b \) to re-introduce them. This illustrates why minimization compresses a trace into events that are necessary for the outcome—which requires eliminating futile cycles. In our case, the causal account for \( p \) starts with the initial condition, symbolized by the init event, and skips to the last \( b \) before the \( u \). Figure 3 depicts the resulting explanation graph, whose arrows denote enablement as defined informally in section 1 and formally in section 4.

Figure 3: A causal explanation for \( p \) in trace (1). Events are labelled by the rules that induced them. The init node corresponds to a special event that sets the mixture to its initial state.

The problem is that the explanation depicted in Figure 3 fails to recognize the critical role of \( pk \) in the original trace. Given the rules of the model, one notes that \( p \) is slow and the average time that \( K \) remains bound to \( S \) depends on the phosphorylation state of \( K \). The kinase \( K \) is phosphorylated in event \( pk \), causing the complex between \( K \) and \( S \) to be sticky, giving the slow phosphorylation \( p \) a chance to occur. It seems reasonable to assert that \( p \) would probably not have happened had \( pk \) not happened, since the opportunity for \( p \) would have been curtailed by a fast unbinding event. Thus, \( pk \) should be part of the explanation, although it neither enables \( b \) nor \( p \) directly (both rules are independent of \( K \)’s phosphorylation state). Reasoning of this kind is counterfactual [Lewis, 1974; Pearl, 2009].

In section 3, we give a rigorous semantics to this line of reasoning and introduce an algorithm for simulating counterfactual scenarios. In section 4, we show how counterfactual dependencies between events can be systematically explained in terms of a combination of enablement and prevention arrows, leading to the explanation shown in Figure 4.

3 Evaluating Counterfactual Statements
In our example, the counterfactual statement to be assessed is: “Had \( pk \) not happened, \( p \) would not have happened.” Our account in the previous section suggests that \( pk \) played a role, but it is also clear that given the stochastic nature of rule firing \( p \) could well have happened even in the absence of \( pk \); it just is unlikely. In a stochastic setting, counterfactual statements are not either true or false, but have degrees of likelihood. To assess that likelihood is our task.

Given an original (factual) trace \( \tau \), a naive approach might be to sample counterfactual traces, each of which starts with the state of the system attained in \( \tau \) just before event \( pk \) happened, but in which we skip over \( pk \) and then run an unconstrained simulation from that point onward. In this approach traces would quickly diverge from the original, distorting the causal role that \( pk \) played specifically in it. The question here is not what causal role \( pk \) can play in principle, but what role it actually did play in \( \tau \). Counterfactual statements are undetectable from the context in which they are formulated.

Pearl’s standard account of counterfactuals [Pearl, 2009] is based on performing “surgical interventions” on a structural equation model (SEM). A SEM features a finite sequence \((x_1, \ldots, x_n)\) of variables, each associated to a functional equation of the form \( x_j = f_j(x_1, \ldots, x_{j-1}, u_k) \), where \( f_j \) is a deterministic function and \( u_k \) a random variable. Ideally, each \( f_j \) defines an independent and autonomous physical mechanism. This is partially enforced by the requirement that the \( u_k \) must be mutually independent. Given some observation \( e \), the probability of the counterfactual statement “had \( x_j \) been equal to \( a \), \( \psi \) would have been true” is evaluated following a three-step process: (abduction) compute the distribution \( p_e \) of values for \( \bar{u} \) given observation \( e \), then (action) intervene in the model by replacing the defining equation for \( x_j \) by “\( x_j = a \)” and finally (prediction) compute the probability that \( \psi \) is true in this new model when \( \bar{u} \) is distributed according to \( p_e \).

Because of their dynamic nature, rule-based models are not readily expressible in terms of structural equations. However, Pearl’s construction generalizes to our setting, assuming a structural refinement of Kappa’s probabilistic semantics where deterministic causal mechanisms are separated from stochastic aspects.
3.1 A Refined Semantics for Kappa

The factual trace to which a counterfactual statement is tied includes random contingencies. These contingencies reflect the stochasticity underlying the specific sequence of events and its timing. Executing a counterfactual experiment requires that we separate this randomness from the deterministic action of rules in order to properly condition on the randomness of the factual trace. We propose such a decomposition, which is motivated by a standard construction in physics that justifies the Doob-Gillespie algorithm used in simulating reaction systems [Gillespie, 1977].

Our refined semantics reconceptualizes the CTMC induced by a model as follows: (i) Consider all possible event templates \( (r, \xi) \), where \( \xi \) maps into a large enough set of global identifiers. For each event template, imagine an independent Poisson process in which a bell rings at time intervals drawn independently from an exponential distribution with parameter \( \lambda_r \). These Poisson processes are all gathered in a random variable \( \Sigma \). A realization of \( \Sigma \) is called a schedule and it features a sequence of ringing times for every bell. (ii) With every schedule \( \sigma \), we associate a unique trace \( \theta(\sigma) \) that is generated as follows: starting with the initial mixture and moving through time, whenever a bell rings, its associated event template \( e \) is realized (by transforming the current mixture \( m \) into \( m \cdot e \)) if and only if \( m \vdash e \). For example, if the current mixture \( m \) is given as in Figure 1 and the bell linked to “apply rule b on substrate 3 and kinase 4” rings, a bond is created between these two agents. In contrast, the bell linked to “apply rule b on substrate 1 and kinase 2” would have no effect. (iii) Finally, the dynamic behavior of our model is captured by the random trace \( T \triangleq \theta(\Sigma) \), which can also be sampled efficiently using the Doob-Gillespie algorithm introduced in section 2.1. Note that the assumption underlying this whole construction is entirely contained in the existence and definition of \( \theta \). Given \( T \), the statistical properties of \( \Sigma \) (including the independence of our metaphorical bells) are consequences of Kappa’s original probabilistic semantics.

Intuitively, \( \Sigma \) determines when the opportunity for a reaction happens and between which molecules. It plays the same role as the random vector \( \bar{u} \) in a SEM. In contrast, \( T \) is a deterministic function that controls whether a reaction can occur when given the opportunity and what it does when it occurs. It corresponds to the \( f_i \) in a SEM and is likewise the target of interventions.

3.2 A Semantics for Counterfactuals

We define an intervention \( \iota \) as a predicate \( \text{BLOCKED}_i[.] \) ranging over events. The purpose of the predicate is to act as a filter preventing the occurrence of selected events. Given a predicate \( \psi \) over traces, we write the statement “Had intervention \( \iota \) happened in trace \( \tau \), \( \psi \) would have been true” as \( \tau \models [\iota] \psi \), borrowing a notation from [Halpern, 2016].

For an intervention \( \iota \) and a schedule \( \sigma \), we define the altered trace \( \theta(\sigma) \) much in the same way as \( \theta(\sigma) \), but also requiring \( \text{BLOCKED}_i[(e, t)] \) to be false for \( e \) to be realized when its bell rings at time \( t \). Then, we define \( T_\iota \triangleq \theta(\Sigma) \).

Given an observed trace \( \tau \), an intervention \( \iota \) and a predicate \( \psi \), the probability of \( \tau \models [\iota] \psi \) can now be defined according to Pearl’s three-step strategy: (abduction) condition the distribution of \( \Sigma \) by the observation that \( \theta(\Sigma) = \tau \), then (action) alter the behavior of \( T \) with intervention \( \iota \) and (prediction) consider the probability that \( \psi \) holds on \( \theta(\Sigma) \). This results in the following definition.

Definition 1 (Semantics of counterfactual statements). For \( \tau \) an observed trace, \( \iota \) an intervention and \( \psi \) a predicate on traces, the probability of the counterfactual statement “had intervention \( \iota \) happened in trace \( \tau \), predicate \( \psi \) would have been true” is defined as:

\[
P(\tau \models [\iota] \psi) \triangleq P(\psi(T_{\iota}) \mid T = \tau).
\]

Following Definition 1, we estimate the probability of the counterfactual statement \( \tau \models [\iota] \psi \) by sampling instances of the random variable \( T_{\iota} \mid \{T = \tau\} \). These are called counterfactual traces. Intuitively, they give an account of what else trace \( \tau \) could have been, had intervention \( \iota \) happened.

3.3 An Example

Let us illustrate our definitions by manually sampling a counterfactual trace for the example trace \( \tau \) given in (1) and the intervention \( \iota \) that consists in blocking every application of rule \( pk \).\text{BLOCKED}_i[(r, \xi, t)] = (r = pk).

For this, we must draw an instance of \( T_{\iota} \), conditioned on the observation \( T = \tau \).

Let us assume that \( T = \tau \). Then, the first event of \( T_{\iota} \) has to coincide with the first event of \( \tau \) (namely \( b \)). Indeed, suppose that \( (e, t) \) belongs to \( T_{\iota} \), with \( t \) prior to the time of the first event of \( \tau \). Thus, \( e \) is scheduled in \( \Sigma \) at time \( t \) and realizable in the initial mixture, which is shared between \( T_{\iota} \) and \( T \). As a consequence, \( (e, t) \) also belongs to \( T \) and therefore to \( \tau \), which is a contradiction. Continuing this line of reasoning, \( T_{\iota} \) and \( \tau \) must coincide until an event of \( \tau \) is blocked by \( \iota \).

After \( pk \) is blocked in \( T_{\iota} \), the current mixtures in \( T \) and \( T_{\iota} \) start diverging (the kinase is phosphorylated in the former and unphosphorylated in the latter). We call these mixtures factual mixture and counterfactual mixture, respectively. The next event to happen in \( \tau \) is the second binding event \( b \).

We argue that it also has to be the next event to happen in \( T_{\iota} \). Indeed, the only way an event \( (e, t) \) can happen in \( T_{\iota} \) before \( b \) while not happening in \( T \) is if \( e \) is realizable in the counterfactual mixture and not in the factual one. This is only true if \( e \) is an instance of rule \( pk \) and applications of this rule – if scheduled – would be blocked by \( \iota \) anyway.

After \( b \) happens in both \( T_{\iota} \) and \( \tau \), the event template associated with rule \( u \) (fast unbinding) becomes realizable in the counterfactual mixture, but not in the factual one. Therefore, the observation \( T = \tau \) provides no useful information about whether or not \( u \) is scheduled in \( \Sigma \) before \( p \) happens in \( \tau \). In fact, the probability that this is not the case is exactly equal to \( \exp\{-\lambda_u \delta\} \), where \( \delta \) is the length of the time interval between \( b \) and \( p \) in \( \tau \). Given these rates in our model, \( \delta \) is typically of the order of \( (\lambda_{u\ast} + \lambda_p)^{-1} \). Therefore, \( \lambda_u \delta \gg 1 \) and it is very likely that a fast unbinding event happens in \( T_{\iota} \) before \( p \) happens in \( \tau \), preventing \( p \) to happen in \( T_{\iota} \). This gives us the following counterfactual trace:

\[
b, u, p, b, u, p, \cdots
\]  

where events that are struck out are events of \( \tau \) that do not appear in \( T_{\iota} \) and events in bold are proper to \( T_{\iota} \).
3.4 The Counterfactual Resimulation Algorithm

We introduce Algorithm 1, a variation of the Doob-Gillespie algorithm, to sample a counterfactual trace efficiently given a reference trace $\tau$ and an intervention $\iota$. We call it counterfactual resimulation, since it works by going through every event of $\tau$, simulating only those parts of $\tau$ that are affected by $\iota$. In particular, when $\iota$ is the trivial intervention (BLOCKED,$\iota = \text{false}$), it returns $\tau$.

This algorithm relies on a modified notion of activity we call divergent activity. We define the set of divergent embeddings of the left-hand side of a rule $r$ into mixture $m$ and relative to $m_0$ as $\text{EMB}_r(m, m_0) \triangleq \text{EMB}_r(m) \setminus \text{EMB}_r(m_0)$. Equivalently, a divergent embedding is an embedding whose codomain features a divergent site, that is, a site whose state differs across $m$ and $m_0$. The divergent activity of a rule $r$ in mixture $m$ relative to $m_0$ is then the product $\lambda_r|\text{EMB}_r(m, m_0)|$. The total divergent activity of the system, $\alpha'(m, m_0)$, is the sum of all divergent activities. Finally, we use the notation $\tau[t]$ to refer to the mixture at time $t$ in $\tau$.

**Algorithm 1** Counterfactual resimulation

Input: a model, a reference trace $\tau$ and an intervention $\iota$

Output: an instance of $T_r | \{T = \tau\}$ (counterfactual trace)

1. $t \leftarrow 0$
2. $m \leftarrow $ initial mixture
3. **while** $t < t_{\text{end}}$ **do**
   4. $m_0 \leftarrow \tau[t]$
   5. $(e_t, t_t) \leftarrow $ first event of $\tau$ in time interval $(t, \infty)$
   6. $\alpha' \leftarrow \sum_r \lambda_r|\text{EMB}_r(m, m_0)|$
   7. draw $\delta \sim \text{Exp}(\alpha')$
   8. $t_c \leftarrow t + \delta$
   9. **if** $t_c < t_f$ **then**
      10. draw a rule $r$ with prob. $\propto \lambda_r|\text{EMB}_r(m, m_0)|$
      11. draw a divergent embedding $\xi \in \text{EMB}_r(m, m_0)$
      12. $e \leftarrow (r, \xi)$
      13. $t \leftarrow t_c$
   14. **else**
      15. $e \leftarrow e_f$
      16. $t \leftarrow t_f$
   17. **if** $\neg \text{BLOCKED}_r((e, t))$ and $m \vdash e$ **then**
      18. update $m$ to $m \cdot e$ and log event $(e, t)$

The role and relevance of the concept of divergent activity in counterfactual resimulation is summarized by the following proposition, where $\tau \cap I = \emptyset$ is a shortcut for “no event of trace $\tau$ occurs in the time interval $I$”.

**Proposition 1** (Property of the divergent activity). For $\tau$ a trace and $\iota$ an intervention, let $I = (t, t + \delta)$ be a time interval such that $\tau \cap I = \emptyset$ and $m_0 = \tau[t]$. Then, we have

$$\mathbb{P} \{ T_r \cap I = \emptyset \mid T = \tau, T_r[t] = m \} = e^{-\alpha'(m, m_0) \delta}.$$  

At every iteration of Algorithm 1, the divergent activity $\alpha'$ determines the probability that an event happens in the counterfactual trace prior to the next event in the factual trace $\tau$ (test of line 9). A proof of Proposition 1 is given in Appendix A. It is the main step in establishing:

**Theorem 1.** The counterfactual resimulation algorithm correctly samples instances of $T_r | \{T = \tau\}$.

3.5 Implementation

There are two challenges in efficiently implementing counterfactual resimulation. The first is a suitable representation for the sets of divergent embeddings $\text{EMB}_r(m)$ to minimize the cost of their update at each iteration. Since the Kappa simulator solves exactly that problem for the sets of embeddings $\text{EMB}_r(m)$ [Danos et al., 2007b], we leverage most of that infrastructure. The second consists in avoiding excessively many iterations of Algorithm 1 in which time is advanced in tiny increments and the proposed event is rejected. Suppose, for example, that in our toy model $pk$ has a very high firing rate and we wish to block, from a specific time onward, all events in which the sole kinase becomes phosphorylated. Upon blocking one occurrence of the event, the same event would want to happen again, and we would keep rejecting it a huge number of times until a different rule fires. More generally, event templates whose realization is bound to be blocked should be removed efficiently before their realization is attempted and not be counted in the system’s divergent activity. We solve this problem for a class of interventions we call regular. Specifically, an intervention $\iota$ is regular if the predicate BLOCKED$_r([((r, \xi), t)]$ can be expressed as a finite disjunction of formulæ of the form $(r = r') \land F(\xi, e) \land (t \in I)$ or $G(r, \xi) \land (t = t')$, where $r'$ is a rule, $t'$ a time, $I$ a time interval, $\xi[e]$ the restriction of $\xi$ to a single connected component $c$ of $L_r$, and $F, G$ arbitrary predicates. For regular interventions, our implementation is guaranteed to either produce or consume an event at each iteration.

**Proposition 2.** Sampling a counterfactual trace for a regular intervention can be done in time $\mathcal{O}(n \cdot r \log |m|)$, where $n$ is the sum of the number of events in the reference trace and in the resulting counterfactual trace, $r$ is the number of rules in the model and $|m|$ the size of the reaction mixture.

For non-regular interventions, there is an additional time-complexity term of $\mathcal{O}(N_0 \cdot r \log |m|)$, where $N_0$ denotes the number of non-productive iterations of Algorithm 1. In Appendix C, we provide a benchmark of our implementation on a scaled-up version of our toy model. The average slowdown per event compared to the Kappa simulator does not exceed 50% for a variety of interventions. Also, we observe that $N_0$ is typically small for the type of non-regular intervention that we expect to be most useful in practice.

Returning to our running example, sampling counterfactual traces repeatedly for trace 1 would reveal that, with very high probability, “event $p$ would not have happened, had $pk$ not happened”. However, we can go further by using counterfactual traces to explain this observation using enablement and prevention arrows.

4 Counterfactuals And Prevention

The diagram shown Figure 4 extends the one in Figure 3 and explains the counterfactual dependency between $p$ and $p_1$.
Some definitions and theorems are provided in the text. For example, Definition 2 and Definition 3 are used to define enablement and prevention. The text also discusses the completeness of enablement and prevention.

4.2 Compression of Counterfactual Experiments

Counterfactual experiments produced with counterfactual resimulation are usually very large. They typically feature a lot of redundancy, including events that are irrelevant to the outcome of interest or futile cycles as discussed in section 2.2. A compression step is usually necessary before concise causal narratives can be extracted from such experiments. However, the two traces of a counterfactual experiment cannot be compressed separately following the procedure described in section 2.2, as there is no guarantee that the compressed traces can still be generated from a unique schedule to form a valid counterfactual experiment. Instead, compressing a counterfactual experiment consists in extracting a minimal valid sub-experiment. A counterfactual experiment \((\tau_1, \iota, \tau_1')\) is said to be a sub-experiment of \((\tau_2, \iota, \tau_2')\) if \(\tau_1\) is a sub-trace of \(\tau_2\) and \(\tau_1'\) is a sub-trace of \(\tau_2'\). Also, valid counterfactual experiments can be characterized as follows.

**Proposition 3.** A triple \((\tau, \iota, \tau')\) is a valid counterfactual experiment if and only if all of the following hold: (1) both \(\tau\) and \(\tau'\) are valid traces; (2) no event of \(\tau'\) is blocked by \(\iota\); (3) for every event \(e \in \tau\) such that \(e \notin \tau'\), then either \(e\) is not executable in \(\tau'\) or \(e\) is blocked by \(\iota\); (4) for every event \(e' \in \tau'\) such that \(e' \notin \tau\), then \(e'\) is not executable in \(\tau\).

Compressing a counterfactual experiment consists in finding a minimal valid sub-experiment such that (i) the outcome...
of interest appears in the factual trace but not in the counterfactual trace and (ii) events that are blocked by \( \iota \) or on which the outcome of interest was shown to be counterfactually dependent in previous analyses, are kept in the factual trace. Because these constraints along with the properties featured in Proposition 3 can be encoded as boolean satisfiability constraints, compressing a counterfactual experiment can be done using standard SAT-solving techniques.

5 Conclusions and Future Work

In this paper, we propose a method for giving meaning to counterfactual statements in mechanistic models of complex physical processes. We modify the continuous-time Monte Carlo algorithm that is used to generate traces in these models so as to sample counterfactual trajectories that stay probabilistically as close to the original (factual) trajectory as an intervention permits them to be. We then construct causal diagrams that explain counterfactual dependencies in terms of enablement and prevention relations between events. Enablement is a standard causal relation between events within factual or within counterfactual traces, whereas prevention shows up as a relation between pairs of events, one in the factual and the other in the counterfactual trace. In particular, prevention can involve events that were not observed in the factual trace. This results in explanatory diagrams that resonate more with the ubiquitous presence of inhibitory interactions in biology and that can capture subtle kinetic aspects of rule-based models. Our completeness result, according to which any counterfactual dependency can be "explained" in terms of enablement and prevention, increases our confidence in this approach to counterfactuals and connects two visions of causality that are typically treated disjointly.

Despite a sound theoretical foundation and an effective implementation, our technique is in need of rigorous practical assessment using large-scale models. Moreover, difficult practical questions remain. Most notably, which counterfactual experiments are worth trying? It is unclear a priori which interventions are informative for traces that include many millions of events. At present, we can only offer tentative directions for future study.

One way to identify interventions worth making without relying on expert knowledge is by developing heuristics, such as recognizing correlations between events in samples of factual traces (or in a single long trace). In our toy model, the occurrence of \( p \) is often preceded by the occurrence of \( pk \). This correlation, together with the absence of \( pk \) from some initial causal account—such as the one presently achievable in a fully automated fashion based on enablement alone, (Figure 3)—suggests to try a counterfactual experiment on \( pk \). More generally, if (i) a context \( C \) in which an event \( e \) occurs is frequently more specific than is required by the left-hand side of the underlying rule and (ii) this observation cannot be explained by the current causal narrative, then a counterfactual experiment in which we block the last event responsible for at least part of \( C \) seems worthwhile in order to assess whether the current causal narrative needs to be updated. Another important practical question is whether interventions should block single events or "knock out" related event templates for a defined timeframe. Our framework accommodates both approaches, as exemplified in Appendix C.

On a more conceptual side, we are investigating principled ways of "gluing" together all explanatory accounts (such as Figure 4) that correspond to different counterfactual experiments. This would summarize the causal structure of a system relative to an outcome of interest. One wonders whether such a summary diagram might constitute a basis for obtaining approximate structural equations for complex mechanistic models. Replacing a rule-based model with such equations could enable targeted statistical analysis to estimate model parameters, for which simulations would be too expensive.

Acknowledgments

This work was sponsored by the Defense Advanced Research Projects Agency (DARPA) and the U. S. Army Research Office under grant numbers W911NF-14-1-0367 and W911NF-17-1-0073. We are grateful to Pierre Boutilier for his help in implementing counterfactual resimulation. We thank Jérôme Feret, Matt Fredrikson, Mickaël Laurent and Jean Krivine for valuable discussions and insights. We thank Thomas Kolokotrones, Ariel Procaccia and the anonymous reviewers for valuable feedback.

A Proof of Proposition 1

Proposition (Property of the divergent activity). For \( \tau \) a trace and \( \iota \) an intervention, let \( I = (t, t+\delta) \) be a time interval such that \( \tau \cap I = \emptyset \) and \( m_0 = \tau [t] \). Then, we have

\[ P \{ T_i \cap I = \emptyset \mid T = \tau, T_i[t] = m \} = e^{-\alpha(m,m_0)\delta} . \]

Proof. Given that \( T_i[t] = m \), no counterfactual event happens in time interval \( I \) if and only if no event template that is realizable in mixture \( m \) is scheduled in \( I \). Therefore,

\[ P \{ T_i \cap I = \emptyset \mid T = \tau, T_i[t] = m \} = \prod_{m^i \in e} \left\{ \bigwedge_{m^i \in e} \{ e \notin \Sigma \cap I \mid T = \tau \} \right\} = \prod_{m^i \in e} P \{ e \notin \Sigma \cap I \mid T = \tau \} . \] (3)

Let \( e \) an event template such that \( m \vdash e \). The probability that \( e \) has not been scheduled for tentative realization in \( I \) given that \( T = \tau \) depends on whether or not \( e \) is realizable in mixture \( m_0 = \tau [t] \). Indeed, we assumed that \( \tau \) contains no event in time interval \( I \). Therefore, if \( m_0 \vdash e \), then \( e \) cannot be scheduled in \( I \), without which it would have been realized in \( \tau \). Thus,

\[ m_0 \vdash e \implies P \{ e \notin \Sigma \cap I \mid T = \tau \} = 1 . \] (4)

In addition, if \( m_0 \nvDash e \), then the observation \( \{ T = \tau \} \) gives no information on whether or not \( e \) has been scheduled in \( I \). Because event templates are scheduled for tentative realization according to Poisson processes, we have:

\[ m_0 \nvDash e \implies P \{ e \notin \Sigma \cap I \mid T = \tau \} = e^{-\lambda \delta} \] (5)
where $\lambda_e$ is the rate of the rule associated with event template $e$. Combining (4) and (5) with equation (3),
\[
\mathbb{P}\{T, \cap I = \emptyset \mid T = \tau, T_i[t] = m\} = \exp\left(- \sum_{m^e, m_0} \lambda_e \cdot \delta \right)
\]
Finally, we can recognize the divergent activity in the exponential above:
\[
\sum_{m^e, m_0} \lambda_e = \sum_{(r, \xi)} \lambda_r \sum_{m \vdash (r, \xi)} 1 \{m \vdash (r, \xi), m_o \not\vdash (r, \xi)\} = \sum_{(r)} \lambda_r |\text{EMB}_r'(m, m_0)| = \alpha'(m, m_0).
\]
This concludes the proof. \qed

B Proof of Theorem 2

It is convenient to prove the following result instead, which is slightly more general.

**Theorem.** Let $(\tau, \iota, \tau')$ be a counterfactual experiment. If $e$ is an event that belongs only to $\tau$ or $\tau'$, then there exists an event $\bar{e}$ in $\tau$ that is blocked by $\iota$ and there is a directed path from $\bar{e}$ to $e$.

**Proof.** We prove this theorem by induction on the number of events before $e$ in both $\tau$ and $\tau'$. Let’s consider $e \in \tau$ such that $\bar{e} \not\in \tau'$. If $\text{BLOCKED}_\iota[e]$ is true, then we are done. Otherwise, by item (3) of Proposition 3, $e$ is not executable in $\tau'$. Therefore, there exists a site $s$ such that event $e$ tests $s$ to a different value than it has in $\tau'[t]$. Let $t$ the time of occurrence of $e$. Moreover, let’s define $e_0$ the last event in $\tau$ modifying $s$ that occurs strictly before time $t$, and $e'_0$ the last event in $\tau'$ modifying $s$ that occurs strictly before time $t$. These events modify $s$ to different values so they cannot be the same. Therefore, writing $t_0$ and $t'_0$ their respective time of occurrence, we have $t_0 \neq t'_0$ with probability one.

- If $t_0 < t'_0$, then we have $e_0 \notin \tau'$ and so we can apply the induction hypothesis on $e_0$. As a consequence, there exists a path from an event $\bar{e}$ in $\tau$ which is blocked by $\iota$ to $e_0$. Moreover, $e_0$ enables $e$. Therefore, there is a path from $\bar{e}$ to $e$.

- If $t'_0 < t_0$, then we have $e'_0 \notin \tau$ and so we can apply the induction hypothesis on $e'_0$. As a consequence, there exists a path from an event $\bar{e}'$ in $\tau'$ which is blocked by $\iota$ to $e'_0$. Moreover, $e'_0$ prevents $e$. Therefore, there is a path from $\bar{e}$ to $e$.

The same proof holds for when $e \in \tau'$ and $e \notin \tau$, using item (4) of Proposition 3 instead of item (3). \qed

This result implies Theorem 2. Indeed, we only need to prove that the constructed path from $\bar{e}$ to $e$ contains an even number of prevention arrows, which is true for any such path as prevention arrows always go from $\tau$ to $\tau'$ or the other way around but cannot “stay” within a single trace.

C Benchmark

We assessed the performance of our implementation of counterfactual resimulation using the toy model of Figure 2, but with an initial mixture consisting of a large number of kinase and substrate instances. Although such a simple model is not of biological significance, it is adequate for an initial assessment of performance.

C.1 Experimental Protocol

We consider four kinds of intervention defined in terms of an event $e_0 = ((r_0, \xi_0), t_0)$:

1. **Singular block:** This blocks only the specific event $e_0$:
   \[
   \text{BLOCKED}_\iota[e] \triangleq (e = e_0).
   \]

2. **Template block:** This blocks every realization of the event template $(r_0, \xi_0)$ from time $t_0$ onward. In formal terms:
   \[
   \text{BLOCKED}_\iota[\{(r, \xi), t\}] \triangleq (r = r_0 \land \xi(\ell) \land M_0 \neq \emptyset \land t \geq t_0).
   \]

3. **Agent-dependent rule block:** This blocks, from $t_0$ onward, every event resulting from rule $r_0$ that tests an agent modified by $e_0$. For example, we might wish to prevent substrate $S_0^{22T}$ from being phosphorylated by any kinase. In formal terms:
   \[
   \text{BLOCKED}_\iota[\{(r, \xi), t\}] \triangleq (r = r_0 \land \xi(L_e) \land M_0 \neq \emptyset \land t \geq t_0)
   \]
   where $M_0$ is the set of agents modified by $e_0$.

4. **Rule block:** This outright disables rule $r_0$ from $t_0$ onward:
   \[
   \text{BLOCKED}_\iota[\{(r, \xi), t\}] \triangleq (r = r_0 \land t \geq t_0).
   \]

All four kinds of intervention might be useful for causal analysis in different settings.

Our experimental setup consists of the model in Figure 2 comprising $10^4$ substrates and $10^4$ kinases. Every agent starts out in an unbound and unphosphorylated state. The simulation is stopped as soon as the system contains more phosphorylated than unphosphorylated substrates. We then proceed as follows. (i) We first generate $n_r = 10$ reference traces and record the CPU time $T$ it took the Kappa simulator to generate each of them on a personal computer with a 2.7GHz Intel Core i5 processor and 16GB of random-access memory. For each trace, we also identify the first application of each $r \in \{b, u, u^*, p, pk\}$ and declare it to be the event $e_0$ underlying the intervention. (ii) We test the 20 possible interventions, based on $e_0$, that can be formed by combining each of the five rules $r$ with each of the four intervention types. (iii) For each such intervention $\iota$, we generate $n_{\tau_{\iota,r}} = 10$ counterfactual traces $\tau'$. (iv) For every counterfactual trace $\tau'$, we record the CPU time $T'$ used by our implementation to generate it. We also record the number $N_{\iota}$ of iterations that neither produced a counterfactual event nor consumed a factual event (non-productive cycles). Finally, we define the slowdown $S$ of counterfactual resimulation relative to simulation as the ratio of $T'$, normalized by the number of distinct events $|\tau \cup \tau'|$ in the counterfactual experiment $(\tau, \iota, \tau')$, to $T$, normalized by the same number $|\tau|$ of events in $\tau$: $S \triangleq 1 - \frac{|\tau|}{|\tau \cup \tau'|} \cdot \frac{T}{T'}$. The average value and standard deviation of these quantities is shown

1889
Moreover, interventions that often be handled efficiently anyway. Non-productive cycles for a similar reason, and therefore, can connected components of same substrate twice in a large mixture. More generally, an intervention can produce non-productive cycles in theory, it is highly unlikely for a kinase to bind the same substrate twice in a large mixture. Although this intervention can produce non-productive cycles, this is not cause major divergences at the population level. In fact, only the five rule-blocking interventions had a major impact.

References


| $\tau$ | $\tau'$ | $|\tau\setminus\tau'|$ | $|\tau'\setminus\tau|$ | $N_{\delta}$ |
|-------|--------|-----------------|-----------------|--------|
| $\mathrm{b}$ | 4.70 | 1.17 ± 0.03 | 2.0 | 0 |
| $\mathrm{u}$ | 4.70 | 1.17 ± 0.03 | 1.0 | 0 |
| $\mathrm{u}^*$ | 4.70 | 1.17 ± 0.04 | 8.3 | 5.1 |
| $\mathrm{p}$ | 4.70 | 1.17 ± 0.03 | 1.0 | 0.5 |
| $\mathrm{pk}$ | 4.68 | 1.17 ± 0.03 | 8.0 | 17.6 |
| $\mathrm{b}$ | 4.69 | 1.17 ± 0.04 | 2.0 | 0 |
| $\mathrm{u}$ | 5.52 | 1.38 ± 0.05 | 30.3 | 2.9 |
| $\mathrm{u}^*$ | 5.43 | 1.35 ± 0.04 | 30.3 | 6.9 |
| $\mathrm{p}$ | 5.50 | 1.37 ± 0.04 | 1.0 | 0.5 |
| $\mathrm{pk}$ | 5.41 | 1.35 ± 0.04 | 8.0 | 17.6 |
| $\mathrm{b}$ | 3.78 | 0.94 ± 0.03 | 1.4e5 | 0 |
| $\mathrm{u}$ | 4.43 | 1.02 ± 0.03 | 1.3e5 | 1.2e4 |
| $\mathrm{u}^*$ | 4.82 | 1.16 ± 0.03 | 2.5e4 | 5.3e3 |
| $\mathrm{p}$ | 4.82 | 1.20 ± 0.03 | 5.0e3 | 0 |
| $\mathrm{pk}$ | 5.19 | 1.08 ± 0.03 | 1.4e4 | 3.0e4 |

Table 1: A benchmark of counterfactual resimulation. On average, $T = 4.01 \pm .12$ s. In addition, $|\tau| = 1.6e5 \pm 1.5e3$. Results

The observed slowdown $S$ never exceeds 50% on average. No intervention produced a non-productive cycle. This is not too surprising, as all the interventions we considered are regular, with the only exception of the “template block” for rule $b$. Although this intervention can produce non-productive cycles in theory, it is highly unlikely for a kinase to bind the same substrate twice in a large mixture. More generally, an intervention $\iota$ that is irregular, because BLOCKED$_\iota([(\rho, \xi), t])$ features a conjunction of terms constraining $\xi$ on different connected components of $L_{\rho\iota}$, does not tend to induce many non-productive cycles for a similar reason and, therefore, can often be handled efficiently anyway.

As expected, we observe that the “stronger” the intervention, the bigger the divergence of counterfactual traces from their factual reference trace. Moreover, interventions that only affect a small number of agents in a large mixture do