# Formal Analysis of Uncertain Continuous Markov Chains in Systems Biology

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- Keywords: Probabilistic Model Checking, Uncertain Continuous-Time Markov Chain, Computational Feasibility, Tractable, Phylogenetic Tree.
- Abstract: Data dependent abstraction for continuous-time Markov chain is often challenging given the incompleteness and imprecision of data. Uncertainty in the environment is modeled in the form of uncertain continuous-time Markov chain. In this work, a tractable model checking methodology, *stochastic partial model set checking* is formalized by approximation of the uncertain continuous-time Markov chain. The methodology was applied in querying to infer on a phylogenetic tree, constructed under uncertainty. Queries were posed on the formalism using continuous stochastic logic formula. Experimental results demonstrate the computational feasibility of the model.

## **1 INTRODUCTION**

Modeling of biological processes is an important research direction in systems biology. One of the problems in systems biology is the correct construction of phylogenetic trees from biological data. Phylogenetic studies are key in understanding the evolution of species. Often the construction of phylogenetic tree is not precise because of imprecision and uncertainty in the data. Stochastic models such as discrete-time Markov chains and continuous-time Markov chain have been used as modeling tools in problems in systems biology and formal verification methods such as model checking. Discrete-time Markov chain (DTMC) have been used as stochastic model in systems biology. The order of the processes in biology is abstracted in the order of the states in the dtmc. However biological applications, time runs continuously, and so discrete-time mathematical models are not always appropriate models. This is especially true in biology where organisms reproduce, infect each other, etc. in continuous time. Therefore, continuous-time Markov chains (CTMC) are often more suited to model abstraction. Continuoustime Markov chains (CTMC) are one of the foundational structure to model continuous probabilities in the modeling processes such as in- rates of execution

of chemical reaction and rates of epidemiology as in spread of diseases. However, rates are imprecise and are uncertain because it is challenging to get precise data given the experimental conditions are not always possible to replicate exactly and there is presence of noise. Parameter synthesis in formal modeling have been applied to address imprecise data. However, often times the modeler would introduce imprecision to quantify error and evaluate the constructed model abstraction in comparison with the real phenomena that is modeled. In this work, we create a tractable formalism for model checking on uncertain continuoustime Markov chain and evaluate the computational feasibility of the application of the model in inference of phylogenetic tree, by posing temporal logic queries.

- A novel inference formalism for temporal logic reasoning on uncertain continuous-time Markov chain is described. A stochastic partial model set is defined that provides the modeler the performance of the models. The methodology is an approximation to perform model checking on uncertain continuous-time Markov chain.
- 2. The formalism has been applied to model phylogenetic tree and computational feasibility of the abstraction on a published dataset of phylogenetic tree is evaluated.

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## 2 BACKGROUND

The foundation of this work is the intersection of probabilistic model checking for uncertain stochastic structures, formal modeling and inference of phylogenetic tree under uncertainty.

## 2.1 Formal Verification of Stochastic Models

Computation of reachability probabilities (Delahaye et al., 2016) for parametric Markov model has been studied by a closed form function (Hahn et al., 2011). Given the imprecision in data, models require parameters for effective inference. Computation of parameters and its estimation is an active area of research. The modeler executes temporal logic queries on the model and computes the parameter (Han et al., 2008). There are synthesis based approaches that have addressed uncertainty in data (Shavit et al., 2016). Approximations for analysis of biological systems using probabilistic model checking by removing the unlikely states, thereby reducing the state space has been studied (Neupane et al., 2019). Interval form of representation of probabilities on Markov chains to incorporate uncertainty has been formalized (Sproston, 2018). Here the intervals were open intervals. In our work, we focus on closed intervals of uncertain kinetic rates. Sampling techniques have been used to learn CTMC (Sen et al., 2004). In our formalism we use sampling on the subintervals after discretization of the interval representing the rates. Sampling having been applied to uncertain parametric CTMCs with parametric transition rates and scalability of the model is evaluated (Badings et al., 2022).

## 2.2 Formal Analysis in Systems Biology

There is a body of literature that described formal analysis of stochastic models of biochemical pathways using continuous-time Markov chains using probabilistic model checking (Calder et al., 2010; Calder et al., 2006; Heath et al., 2008) and different formalisms to address uncertainty in the environment. Quantitative analysis using continuous-time stochastic logic and have been compared with ordinary differential equation based model(Calder et al., 2006). However, the models do not take into account environmental variations in which the rate of reactions, i.e the speeds at which pathway (biochemical reaction) will execute is assumed to be exact. Recent work incorporating imprecision in CTMC is the formalization of uncertain CTMCs (Cardelli et al., 2023). The rates of the CTMCs have been assumed to vary nondeterministically with time represented by continuous intervals. The states of the CTMC are partitioned and the states are lifted in a standard CTMC setting but under uncertainty. The refinements were based on bisimulation of stochastic models (Baier et al., 2003; Neuhäußer and Katoen, 2007). Uncertain kinetic rates using probabilistic model checking has been described by creating a novel semantics (Barbuti et al., 2012). Model checking of uncertain continuous-time Markov chains have been reported(Bortolussi et al., 2016). Chemical reaction network has been modeled as parametric CTMC integrating with learning models with the goal of applying statistical model checking (Molyneux and Abate, 2020).

## 2.3 Construction of Phylogenetic Tree Under Uncertainty

Monte Carlo based method was studied as an inference method on phylogenetic tree by representing CTMC and approximate transition probabilities and compute the parameters (Hajiaghayi et al., 2014). Bayesian phylogenetic inference using annealed sequential Monte Carlo method have been proposed (Wang et al., 2020). This direction of research is ongoing and there is no substantial approximation that has been accepted as the standard in the community.

## **3 PRELIMINARIES**

In this section, we state the mathematical preliminaries that are foundations of our formalism. A continuous time Markov chain is a stochastic model with a discrete states with a continuous time as a parameter. In formal modeling, the continuous time Markov chain is described in the form of a state based system, called a labeled continuous-time Markov chain. For consistency throughout the paper, we will denote a labeled continuous-time Markov chain as CTMC.

**Definition 1.** A labeled CTMC (Baier et al., 2000) is a tuple,  $\mathcal{M} = \langle S, S_0, R, AP, L \rangle$  where:

- 1. S is a set of states.
- 2.  $S_0 \subset S$  is the set of initial states.
- *3.*  $R: S \times S \rightarrow \mathbb{R}_{\geq 0}$  as the rate matrix.
- 4.  $L: S \leftarrow 2^{AP}$  is the labeling function.

The labeled CTMC described in Definition 2 eliminates the requirement  $\mathbf{R}(s,s) = \sum_{s \neq s'} \mathbf{R}(s,s')$ , unlike non-state based definition of CTMCs. Self-loops are modeled by  $\mathbf{R}(s,s') > 0$ . **Definition 2.** (*Labeled Uncertain Continuous-time Markov Chain*) *A labeled uncertain CTMC (UCTMC) is a tuple,*  $\mathcal{K} = \langle S, S_0, R, \mathcal{A}, AP, L \rangle$  *where:* 

- 1. S is a set of states.
- 2.  $S_0 \subset S$  is the set of initial states.
- *3. A is the the set of actions.*
- 4.  $R: S \times \mathcal{A} \times S \rightarrow \mathbb{R}_{\geq 0}$  as the rate matrix and each,  $R_{i,j}$  is an interval,  $(m,n), m \leq n$  and  $m, n \in Q$ .
- 5.  $L: S \rightarrow 2^{AP}$  is a state labeling function.

A path in a UCTMC is a non-empty sequence,  $s_0a_0t_0s_1a_1t_1s_2...$  where  $R(s_i, a_i, s_{i+1}) > 0$  and  $t_i \in \mathbb{R}_{>0}$  for all  $i \leq 0$ . The value of  $t \in (\check{t}_i, \hat{t}_i)$ . The open interval is a way to add imprecision.

### 3.1 Probabilistic Model Checking

Model checking is performed on labeled finite state machine in the form of Kripke structure,  $\mathcal{M}$ . Formally,

**Definition 3.** (Model Checking (Clarke, 1997)) Given a model,  $\mathcal{M}$  and formula, $\phi$ , model checking is the process of deciding whether a formula  $\phi$  is true in the model, written  $\mathcal{M} \models \phi$ .

Here,  $\phi$  represents a temporal logic formula that represents the properties of a system. The model,  $\mathcal{M}$  represents the description of the system. The system description when represented by a stochastic structure,  $\mathcal{M}_s$ , then the model checking is performed on a stochastic structure and the validity of the formula is quantified by probability.

**Definition 4.** (*Probabilistic Model Checking*) Given a model,  $\mathcal{M}_s$  and formula, $\phi$ , model checking is the process of deciding whether a formula  $\phi$  is true in the model, written  $\mathcal{M} \models \phi$  and quantified by a probability.

Details on probabilistic computational tree logics can be found in the literature (Baier et al., 2008; Aziz et al., 1996) Probabilistic model checking on uncertain stochastic structures such as interval discrete Markov chains have been shown to be in PSPACE and NP-hard (Benedikt et al., 2013; Sen et al., 2006).

#### 3.1.1 Continuous Stochastic Logic

Stochastic Model Checking have been described (Kwiatkowska et al., 2007). Model checking on CTMC is performed by the logics known as continuous stochastic logic (CSL) (Aziz et al., 1996; Baier et al., 1999) The syntax of CSL (Aziz et al., 1996) Syntax of CSL is:

$$\begin{split} \phi &::= true \mid a \mid \phi \land \phi \mid \neg \phi \mid \mathcal{P}_{\oplus p}[\psi] \mid \mathcal{S}_{\oplus p}[\phi] \\ \psi &::= X \phi \mid \phi \mathcal{U}^{\leq k} \phi \mid \phi \mathcal{U} \phi \end{split}$$

where a is an atomic proposition,  $\oplus \in \{\leq, <, \geq, >$ 

 $\{p \in [0,1] \text{ and } k \in \mathbb{R}_{\geq 0}. \phi, \psi \text{ are state and path for-}\}$ mula respectively.  $\mathcal{P}_{\oplus p} \psi$  represents the probability of  $\phi$  satisfied from a given state satisfies the bound  $\oplus p$ . The bounded until operator  $\phi_1 \mathcal{U}^{\leq k} \phi_2$  is valid if  $\phi_2$  for a time instant in the interval [0, k] and  $\phi_1$  is valid at all preceding time instants. The path formula,  $X\phi$  is true if  $\phi$  is satisfied in the next state. The formula  $\phi_1 \mathcal{U}^{\leq k} \phi_2$ is true if  $\phi_2$  is satisfied within k time-steps and  $\phi_1$  is true at that point. Similar is the description of  $\phi_1 \mathcal{U} \phi_2$ where  $\phi_2$  is true some point in future till then  $\phi_1$  is true.  $\phi_2$  is satisfied at some time instant in the interval [0,k] and  $\phi_1$  holds at all preceding time steps. The S operator describes the steady-state behaviour of the CTMC.  $S_{\oplus}p[\phi]$  asserts that the steady-state probability of being in a state satisfying  $\phi$  meets the bound  $S_{\oplus p}$  The semantics of CSL are defined on CTMC and can be found (Aziz et al., 1996; Baier et al., 1999) for further reading.

## 4 STOCHASTIC PARTIAL MODEL SET CHECKING

Probabilistic model checking is performed on stochastic structures such as DTMC or CTMC. Model checking on UCTMC is challenging because the intervals for rates can generate infinite number of rates. The following are keys to approximate the UCTMC denoted by  $\mathcal{M}_s$  and construct a tractable computational model. A finite set of *stochastic partial model*,  $\mathcal{M}_{pp} = \{M_1, M_2, \dots, M_n\}$  where  $n \in \mathbb{N}$ . Each stochastic partial model  $M \in \mathcal{M}_{pp}$  has identical state space that of  $\mathcal{M}_s$  with the intervals for rates of M are the subintervals for rates in  $\mathcal{M}_s$ .

The model checking problem is phrased as given a model  $M_i$  and a formula,  $\phi$ , model checking is the process of deciding whether a formula  $\phi$  is true in the model and represented by  $M_i \models \phi$ . The generalization of the model checking on probabilistic partial models will be denoted by probabilistic partial model checking problem. Formally, given a set of probabilistic partial models, M and a formula,  $\phi$  model checking is the process of deciding whether a formula is true for each  $M \in \mathcal{M}_{pp}$ . and denoted by  $M \models \phi$ .

#### 4.1 Model Abstraction

The model abstraction is two step process: Initially, a UCTMC is constructed from the data. The next step is to construct set of stochastic partial models from the UCTMC is constructed.

*Construction of UCTMC Representing Phylogenetic Tree:* Given a set of species, *G* and set of rate of evolution (transitions),  $R = \{I_1, ..., I_n\}$  where  $n \in \mathbb{N}$  and

**Input:**  $\mathcal{K}_{s}(S_{0}, S, R, \mathcal{A}, AP, L)$  Number of subintervals, k

Output: Set of Stochastic Partial Models, R

- 1:  $S = \{s_o\}, \hat{S} = \emptyset, \hat{I}; \{S_O \text{ is the initial set and } \check{T} \text{ is}$ the set of transitions.}
- 2: IntervalSet  $\leftarrow 0$
- 3: OrderedSetRate  $\leftarrow \emptyset$
- 4: **for** each  $s_i \xrightarrow{r_{ij}} s_j$  where  $s_i, s_j \in S$  **do** 5: Construct *k* subintervals for each  $r_{ij}$ .
- IntervalSet  $\leftarrow$  IntervalSet  $\cup (r_1^{ij}, r_2^{ij}, \dots, r_k^{ij})$ 6: where  $r_{ij}$  is a subinterval.
- 7: **end for**

8: OrderdedTuples 
$$\leftarrow \prod^{|\text{IntervalSet}|}$$
 IntervalSet

9:  $\mathfrak{K} \leftarrow \emptyset$ 

- 10: for each  $T \in \text{OrderedTuples } \mathbf{do}$  $R \leftarrow T$ 11:
- $\mathcal{K}_{\mathcal{S}}\langle S_0, S, T, \mathcal{A}, AP, L \rangle$ 12:
- 13:  $\mathfrak{K} \leftarrow \mathfrak{K} \cup \mathcal{K}_{\mathfrak{s}}$
- 14: end for
- 15: K

Algorithm 1: StochasticPartialModel.

 $I_{am}$  is the interval representing *m*th. clade. UCTMC is constructed in the following way: Each state, s is labeled as specie and a transition to another state, s'implies species A evolves into species B. The label on the transition,  $I_m$  is the interval that represents the range of values the rate of evolution from species A to species B. The state, s' is labeled with species. Construction of Stochastic Partial Model Set: In the construction of a set of stochastic partial models, each intervals for rates on the transition in UCTMC is partitioned into subintervals. The partitions of the intervals are user defined value, k amd  $k \in \mathbb{N}$ . For example, an interval in  $\mathcal{K}_s$  between a transition  $s \to s'$  is given by I = (a, b) where  $a, b \in (N)$  and a < b, the intervals where  $s, s' \in \mathcal{K}_s$ . For k = 2, the subintervals are:  $(a, \frac{a+b}{2}], (\frac{a+b}{2}, b)$ . Similarly, for *n* intervals in  $\mathcal{K}_{s}$ , there will be *nk* subintervals and each stochastic partial model will represent a UCTMC with a series of subinterval from a series of intervals of transition rates in  $\mathcal{K}_s$ . There are  $k^n$  stochastic partial models. kcan be varied. Algorithm 1 shows the construction of set of stochastic partial model set. The fixed point algorithm constructs the set of stochastic partial models with subintervals.

#### 4.2 **Construction of Simulation Model**

The transition rates for probabilistic partial models are represented in the form of intervals. In order to make the models tractable, sampling on the intervals

are performed. Notation for upper bound and lower bound of an interval in a stochastic partial model is given by  $ub(I_p)$  and  $lb(I_p)$ , respectively and  $I_p$  represents an interval in a stochastic partial model and the set of intervals is  $\hat{I}_p$ . Algorithm 2 demonstrates the sampling of the rates from each intervals in a stochastic partial model. The models in the set, R form the basis of probabilistic model checking. If the values of the probabilities for consecutive subintervals are close to each other, then the intervals can be merged and hence, the number of models get reduced. The algorithm terminates because the number of intervals are finite and there are finite states in each stochastic partial model in the finite set,  $\Re$ . Each  $K_{sim} \in \Re_{sim}$ is a CTMC, an approximation of  $\mathcal{K}_{s} \in \mathfrak{K}$  and hence, CSL logic queries can be executed. The computational complexity of model checking is polynomial in the size of the model. Therefore, for a partial model set of size n, the model checking complexity is polynomial as the model checking is performed on each of the partial model.

#### Input: R

Output: Set of Sampled Stochastic Partial Models,

 $\Re_{sim}$ .  $\mathcal{K}_{sim} \leftarrow 0$ 1: 2: for each  $K \in \mathfrak{K}$  do for each  $r \in R$  do 3: 4:

- if r is partly closed interval then
- Generate random value, r such that 5:  $lb(I_p) < r \le up(I_p)$
- 6: else
- 7: Generate random value, r such that  $lb(I_p) \le r \le up(I_p)$
- SRate  $\leftarrow$  SRate  $\cup$  *r* where  $i = |\hat{I}_p|$ . 8:
- 9: end if
- 10:  $\mathcal{K}_{sim}(S_0, S, srate, \mathcal{A}, AP, L)$ .srate  $\in$  SRate.
- end for 11:
- $\mathfrak{K}_{sim} \leftarrow \mathfrak{K}_{sim} \cup \mathcal{K}_{sim}$ 12:
- 13: **end for**
- 14: £sim

Algorithm 2: Sampled Stochastic Partial Models.

#### **APPLICATION OF UCTMC IN** 5 PHYLOGENETIC TREE

The steps involved in reasoning for an abstraction of phylogenetic tree under uncertainty is given by :

- 1. Construct an UCTMC,  $\mathcal{K}_t$  representing a phylogenetic tree,  $\mathcal{T}$ .
- 2. Construct the set of stochastic partial models from the UTMC,  $\mathcal{K}$

3. Perform probabilistic model checking by posing queries using continuous stochastic logic on the set of sampled stochastic partial models.

The data used as the phylogenetic tree model was from published data by Baele et al. (Baele et al., 2021), as shown in Figure 1. The models were constructed and evaluated in PRISM (Kwiatkowska et al., 2011) The states in the model are labeled symbolically with species.



Figure 1: Fragment of Phylogenetic tree representing the speciation of eudicots, angiosperms, and seed plants (Baele et al., 2021).

Notation: The various species of plants are Arabidopsis Thaliana(at), Spinacia Oleracea(so), Oenothera Elata(oe), Lotus Corniculatus(lc), Nicotiana Tabacum(nt), and Atropa Belladonna(ab) in the model.

In the Figure 2, the boxplots show the variation amongst each model variation. The data is based on the randomly generated rates based on the ranges specified. The figure is provided for the sole purpose of conveying the variations among a select few models involved in the total calculations.

## 5.1 Evaluation of Temporal Logic Properties

The evaluation of specifications, represented by probabilistic temporal logics is performed using PRISM (https://www.prismmodelchecker.org/) model checker. The simulation was performed in a Mac with 2.3 GHz with quad core Intel i5 processor and 8GB RAM. In this model, we are focused on reachability properties and temporal logic properties. A sample of reachability biological queries represented as continuous stochastic logic formula are as follows:

**Property 1:** P=? (*ab* & *nt* & *oe*). The meaning of the statement is: Is there a state where all three species *ab*, *nt* and *oe* are present? Note: This

property's result will always evaluate to 0.0 because our model checks to see if a single species formed, but not multiple species.

- **Property 2:** P=? (*nt* = true) The meaning of the logic formula- is it possible to reach a state where species *nt* is present in the model.
- **Property 3:** P=? (*ab* = true ) The meaning of the logic formula- is it possible to reach a state where species *ab* is present in the model.
- **Property 4:** P=? (*at* = true). The meaning of the logic formula- is it possible to reach a state where species *at* is present in the model.
- **Property 5:** P=? (*so* = true ) The meaning of the logic formula- is it possible to reach a state where species *so* is present in the model.
- **Property 6:** P=? (*oe* = true ) This is a reachability property asking if it is possible to reach a state where species *oe* is present in the model.

Table 1: Execution Time of CSL Queries.

Model	Property	Time (in seconds)	
n=6	1	0.088	
	2	0.009	
	3	0.001	
	4	0.001	
	5	0.001	
	6	0	
n=10		0.018	
	2	0.202	
	3	0.202	
	4	0.245	
	5	0.23	
	6	0.227	
n=13	1	0.453	
	2	41.65	
	3	40.76	
	4	53.945	
	5	52.851	
	6	45.922	

Table 1 shows of the results obtained for model checking for different problem sizes input sizes and properties. The data consists of results obtained for seven different input sizes: n=6, n=10, and n=13, with varying properties. There are over 40 models, which refer to the varied rates specified earlier. Each variation has been checked for the same set of properties. For n=6, the results are further broken down into seven different variations of the input. For each input size and property combination, the "Time" for execution of the queries are presented. The properties checked are numbered from 1 to 6. The time taken



for model checking is recorded in seconds, and the result indicates whether the model satisfies the property checked. For example, for n=6 and property 2, the time taken for model checking was 0.009 seconds, and the result was 0.212459016, indicating that the model satisfied the property. The times of the execution of the logic queries demonstrate a correlation with the increase in the input sizes. For example, the time taken for model checking n=6 is significantly less than the time taken for n=13. However, there are some variations in time taken for the same input size, indicating that the properties of the input can also have an impact on the time taken for model checking.

#### 5.2 Analysis of Properties

Table 2 shows some properties are satisfied for all values of n, while others are not. Property 1 seems to be satisfied for all values of n, as indicated by the result of 0 for all entries. Here is the table with the average result for each individual property (averages are rounded to four decimal places):

The average result of each property reveals that some properties are more likely to be present in some models than others. Property 7 has the highest average probability of presence, with a 34.9% chance Table 2: Computed Probabilities from Properties.

Property	Average Result	
J PU	0.0	
2	0.227723931	
3	0.160131871	
4	0.077751529	
5	0.078001573	
6	0.160862186	

of eventually being present at some point in the model checking process. This could be due to various reasons, including but not limited to, the particular organism specified in the property and the values of the rates of transition associated with the property.

Analysis Results of Temporal Logic Queries: Table 3 shows the comparison of a property across all the models. Property 2 is defined as follows: P=? [F nt = true]. This property checks whether the state with nt is eventually reached during the execution of the system. The result of the property verification for different problem sizes. For the model with n=6, the model checker took 0.009 seconds to verify the property and returned a result of 0.212459. This means that there is a 21.25% chance that the system will eventually reach the state labeled with nt during execution. For the model with n=10, the model checker took 0.202

Model	Property	Time (seconds)	Result
n =6	2	0.009	0.212459
n =10	2	0.202	0.310325
n =13	2	40.76	0.220842
n =6 (variation 1)	2	0.001	0.187863

Table 3: Comparison of time of execution (in seconds) across different models.

seconds to verify the property and returned a result of 0.310325. This means that there is a 31.03% chance that the system will eventually reach the state nt during execution. For the model with n=13, the model checker took 40.76 seconds to verify the property and returned a result of 0.220842649. This means that there is a 22.08% chance that the system will eventually reach the state nt during execution. The model with n=6 (variation 1) represents a larger size (size of  $\pm 2$  ) of the interval for rates. The model checker took 0.001 seconds to verify the property and returned a result of 0.18786327, implying there is a 18.79% chance that the system will eventually reach the state nt during execution. The change in the size of the interval of the rates did not influence the time for verification of the specification. These results remain similar for each property, as the input size and model variables are altered. Based on these results, we can conclude that the larger the size of the model, the longer it takes for the model checker to verify the property. Additionally, we can see that the probability of reaching the *nt* state during system execution is higher in the model with n=10 compared to the other models. This information can be useful in determining the feasibility of the system and making design decisions to optimize the system performance.

## 6 CONCLUSION

In this paper, we describe an abstraction of feasible models, an approximation of a given uncertain continuous-time Markov chain. The approximated set of CTMC constructed is used for stochastic partial model set checking. The computational feasibility is evaluated on continuous stochastic logic queries, on a prototype of a phylogenetic tree from published data. The modeling technique can be improved by quantifying error for model checking of partial models and identifiability of the partial models (Browning et al., 2020). The formalism is promising to model other problems in systems biology where there data is imprecise and provide the modeler a tool to provide insights to the biological questions.

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