Bayesian parameter synthesis of Markov population models

Master Project on Modeling of Complex, Self-organizing systems

Huy Phung University of Konstanz

July 19, 2020

Contents

1	Preliminaries 4						
	1.1	Discrete time Markov Chain	4				
		1.1.1 Definition	4				
		1.1.2 Parametric Discrete-time Markov Chain	5				
		1.1.3 Markov population model	6				
	1.2	Bayesian inference	6				
		1.2.1 Bayesian parameter estimation	6				
		1.2.2 Posterior conjugation	7				
		1.2.3 Metropolis-Hastings algorithm	9				
		1.2.4 Selection of prior distribution	10				
		1.2.5 Bayesian Parameter Estimation	10				
			11				
2	Rel	ated works	11				
3	Mo	deling of bees colony	12				
	3.1	· ·	12				
	3.2	Markov population models for bee colony	12				
		· · · · · · · · · · · · · · · · · ·	16				
			17				
4	Bayesian parameter synthesis of Markov population model 1						
	4.1	Inference of terminal state distribution	18				
	4.2	Evaluation of terminal states distribution	18				
	4.3	Inference of model parameter with Metropolis-Hastings algo-					
		rithm	19				
5	Ben	nchmark	21				
	5.1	Data synthesis	21				
	5.2	Evaluation measures	22				
	5.3	Inference of terminal state distribution	22				
		5.3.1 Experiment setup	22				
			23				
	5.4	Compare DTMC sampling and rational functions	24				
		5.4.1 Experiment setup:	24				
			24				

		5.4.3 Analysis of long run property	25					
	5.5	inference of model parameters						
		5.5.1 Experiment setup:	25					
		5.5.2 Results	25					
	5.6	5.6 Impacto of Metropolis-Hastings parameter selection 25						
	5.7	Impact of data quality to inferences	25					
	5.8	Linear model	25					
6	Implementation 25							
	6.1	Class design	25					
	6.2	Test environment	26					
	6.3	Directory structure	26					
7	Con	nclusion 2	.7					

Abstract

We study the collective behavior of a bee colony. Each bee in a colony possibly stings after observing a threat in the surrounding environment, and warn other bees by releasing a special substance, pheromone. By sensing the pheromone released in the environment, other bees in the colony may also sting. However, since stinging leads to the termination of an individual bee, it reduces the total defense capability as well. With parametric Discrete-time Markov chain as the model, we study how the actions of a single bee change with regarding to the colony size of and pheromone amount. This project propose and discuss Bayesian methods to estimate parameters of bees population models. It shows that the proposed methods is scalable and can deliver estimations of model parameters.

1 Preliminaries

1.1 Discrete time Markov Chain

Assume that each bee in a colony decides its next action (to sting or not to sting) based only on the current state of the environment, and the number of bees who sting or not sting can be modeled as a Markov process. To reduce the complexity of the model, we make another assumption that the states of the bees colony are observed after uniform time duration, hence the model is of discrete-time.

1.1.1 Definition

Definition 1.1 (Discrete Time Markov Chain). A Discrete Time Markov Chain (DTMC) is a tuple $(S, \mathbf{P}, S_{init}, AP, L)$ [2]

- S is a countable non-emty set of states
- $P: S \times S \rightarrow [0,1]$ is the transition probability function, s.t

$$\sum_{s' \in S} \mathbf{P}(s, s') = 1$$

• $S_{init}: S \to [0,1]$ is the initial distribution, s.t

$$\sum_{s' \in S} S_{init}(s') = 1$$

- AP is a set of atomic propositions
- $L: S \to 2^{AP}$ is the labelling function on states.

The transition probability function in DTMC is given as a stochastic matrix **P**, which satisfies the following properties

- $oldsymbol{\cdot}$ **P** is a square matrix.
- $\mathbf{P}_{i,j} \in [0,1].$
- $\sum_{i} \mathbf{P}_{i,j} = 1$

Definition 1.2 (Strongly Connected Component). Let $\mathcal{M} = (S, \mathbf{P}, S_{init}, AP, L)$ a DTMC. A subset $S' \subset S$ is strongly connected if and only if for every pair $s_1, s_2 \in S'$ there is a path between s_1 and s_2 which consists of only of state in S'. If there exist no $S'' \subseteq S$, such that $S \subset S''$ and S'' is strongly connected, then S' is a Strongly Connected Component, or SCC in short.

Definition 1.3 (Bottom Strongly Connected Component). Let $\mathcal{M} = (S, \mathbf{P}, S_{init}, AP, L)$ a DTMC and $S' \in S$ a Strongly Connected Component. S' is also a Bottom Strongly Connected Component, or BSCC for short, if and only if there exist no state $s \in S$ S' that is reachable from any state in S'.

1.1.2 Parametric Discrete-time Markov Chain

In order to generalize the stochastic matrix to encompass unknown information of the system, we introduce parametric DTMC. Let $\theta = (\theta_1, \dots, \theta_n) \in [0, 1]^n$, we represent each element in matrix **P** as a polynomial function of θ . Let \mathbf{Pol}_{θ} be the set of polynomials $P : [0, 1]^n \to [0, 1]$. We define parametric Discrete-Time Markov Chain, or pMC for short, as follow:

Definition 1.4 (Parametric DTMC). A parametric Discrete Time Markov Chain (pMC for short) is a tuple $(S, \theta, \mathbf{P}_{\theta}, S_{init}, AP, L)$

- S is a countable nonemty set of states
- θ is the set of model parameters
- $\mathbf{P}: S \times S \to \mathbf{Pol}_{\theta}$ is the transition probability function that map a transition relation between two states to a polynomial function of θ

• $S_{init}: S \to [0,1]$ is the initial distribution, s.t

$$\sum_{s' \in S} S_{init}(s') = 1$$

- AP is a set of atomic propositions
- $L: S \to 2^{AP}$ is the labelling function on states.

A concrete assignment of θ on pMC induces a DTMC. In this project, we concern about the problem of data-informed estimation of model parameter θ .

1.1.3 Markov population model

In the scope of this project we concern *population* of a bee colony with N individuals initially. We model the population of a bee colony using a pMC $\mathcal{M} = (S, \theta, \mathbf{P}_{\theta}, S_{init}, AP, L)$ with the following properties:

- Each state $s \in S$ represent number of individuals in the bee colony.
- There exists a set of terminal states $T \subset S$, such that t is a BSCC for all $t \in T$, and |T| = N + 1.

Assume we conduct a real biological experiment on a colony of N bees. At the end of the observation, we observes N' bees $(0 \le N' \le N)$ bees. Let $tSCC_i \in T$ denotes the terminal states with i bees left. As we conduct the experiment with the same colony size multiple times, we get a probability distribution over final states T.

1.2 Bayesian inference

1.2.1 Bayesian parameter estimation

Let D be observed data. In statistical inference, we assume that the observed data has a probability distribution of unknown parameter θ , i.e $D \sim P(D|\theta)$. In frequentist approach, the estimation of θ based on long-run property, that is, given a large enough sample size, expected value of parameter estimation $\hat{\theta}$ is equal to θ . Therefore, frequentist approach requires to gather a large

amount of data to deliver a close estimation $\hat{\theta}$. In Bayesian approach, we reuse the information beliefs gained from observed data to enhance the accuracy of the estimation of $\hat{\theta}$. The main advantage of Bayesian approach over frequentist approach is that it require less data to obtain an estimation $\hat{\theta}$. The beliefs obtained from prior knowledge of model parameter θ is repre-

sented by prior distribution $\pi(\theta)$. Also, we have probability distribution of observed data, given parameter θ , $P(D|\theta)$. This is also called *likelihood function*.

With Bayesian formula, we have

$$\pi(\theta|D) = \frac{P(D|\theta)\pi(\theta)}{\int_{\theta} P(D|\theta)\pi(\theta)d\theta}$$

 $\int_{\theta} P(D|\theta)\pi(\theta)d\theta$ is called marginal distribution. $\pi(\theta|D)$ is called posterior distribution. Computing posterior distribution is the essential part of Bayesian inference, since it gives us the estimation of parameter θ .

1.2.2 Posterior conjugation

Conjugated posteriors are special cases of Bayesian inference, in which the prior and posterior distribution belongs to the same family of distribution. Conjugated posteriors give us significant benefits

- 1. Tractability: we have analytical form of posterior distribution.
- 2. Computationally effective: updating model parameter is of linear time to the dimension of parameter.

We consider two conjugated posterior: Binomial-Beta and Dirichlet-Multinomial

Lemma 1 (Binomial-Beta Conjugation). Binomial distribution is conjugated to beta distribution.

Proof. The observed data $D = (x_1, \ldots, x_n)$ is sampled from $Binomial(k, \theta)$ function

$$P(D|\theta) = \prod_{i=1}^{n} {k \choose x_i} \theta^{x_i} (1-\theta)^{k-x_i}$$

The parameter θ is of $Beta(\alpha, \beta)$ distribution

$$\pi(\theta) = \theta^{\alpha - 1} (1 - \theta)^{\beta - 1}$$

We obtained:

$$\pi(\theta|D) \sim P(D|\theta)\pi(\theta)$$

$$\sim \theta^{\sum_{i=1}^{n} x_i} (1-\theta)^{nk-\sum_{i=1}^{n} x_i} \theta^{\alpha-1} (1-\theta)^{\beta-1}$$

$$= \theta^{\alpha-1+\sum_{i=1}^{n} x_i} (1-\theta)^{\beta-1+nk-\sum_{i=1}^{n} x_i}$$

Thus, the posterior is $Beta(\alpha + \sum_{i=1}^{n} x_i, \beta + nk - \sum_{i=1}^{n} x_i)$

Generalize this conjugation, we also have Multinomial-Dirichlet conjugation.

Lemma 2 (Multinomial-Dirichlet Conjugation). Multinomial distribution is conjugated to Dirichlet distribution.

Proof. The observed data $D = (x_1, \dots, x_n)$ is sampled from $Multinomial(n; \theta_1, \dots, \theta_n)$ function

$$P(x_1,\ldots,x_n|N,\theta_0,\ldots,\theta_n) = \frac{n!}{x_1!\ldots x_n!} \prod_{i=1}^n \theta_i^{x_i}$$

The parameter $(\theta_1, \ldots, \theta_n)$ is $Dirichlet(\alpha_1, \ldots, \alpha_n)$

$$\pi(\theta_1,\ldots,\theta_n) = \frac{1}{\mathbf{B}(\alpha_1,\ldots,\alpha_n)} \prod_{i=1}^n \theta_i^{\alpha_i-1}$$

We obtain

$$\pi(\theta_1, \dots, \theta_n | D) \sim P(D|\theta)\pi(\theta)$$

$$\sim \prod_{i=1}^n \theta_i^{x_i} \prod_{i=1}^n \theta_i^{\alpha_i - 1}$$

$$\sim \prod_{i=1}^n \theta_i^{\alpha_i - 1 + \sum_{i=1}^n x_i}$$

Thus, the posterior is $Dirichlet(\alpha_1 + x_1, \dots, \alpha_n + x_n)$

More detailed description in these cases can be found in [11] and [3]. We summarize the necessary results in the following table:

Likelihood	Prior	Posterior parameters
Binomial(n,k)	$Beta(\alpha, \beta)$	$\alpha' = \alpha + \sum_{i=1}^{n} x_i, \beta' = \beta + nk - \sum_{i=1}^{n} x_i$
$Multinomial(n; \theta_1, \dots, \theta_n)$	$Dirichlet(\alpha_1,\ldots,\alpha_n)$	$\alpha_i' = \alpha_i + x_i, 1 \le i \le n$

1.2.3 Metropolis-Hastings algorithm

In case the posterior distribution has no analytical form or its analytical form is difficult to sample from directly, we use Metropolis-Hastings algorithm (MH in short).

Metropolis-Hastings algorithm is a *Monte Carlo Markov Chain* algorithm. In its essential, Metropolis-Hastings algorithm draws Using the MH algorithm, we can estimate the parameter by posterior mean, without knowing the analytical form of posterior distribution itself.

```
Algorithm 1 Metropolis-Hastings Algorithm
```

```
Input: D is the observation data,
    Output: Trace is the set of accepted sampling point.
 1: procedure METROPOLIS-HASTINGS(D, maxIteration)
 2:
        Select a proposal distribution \pi(\theta)
        Draw a random initial point \theta
 3:
        Init empty trace Trace
 4:
        while maxIteration not reached do
 5:
            L \leftarrow P(D|\theta)
 6:
            Draw a point \theta' from the proposal distribution.
 7:
            L' \leftarrow P(D|\theta')
 8:
            if ln(L') - ln(L) > 0 then
 9:
                Add \theta' to Trace
10:
                \theta = \theta'
11:
            else
12:
                Draw a random number x from Uniform(0,1)
13:
                if x < \epsilon, (\epsilon very small, e.g 10^{-8}) then
14:
                    Add \theta' to Trace (avoiding local maxima)
15:
                    \theta = \theta'
16:
                end if
17:
            end if
18:
        end while
19:
20: end procedure
```

The likelihood function can be implemented as log-likelihood to avoid underflow error. Proposal distribution defines how do we proceed to the next parameter value on the parameter space; it can be of any distribution family. There are two advantages of using Markov Chain Monte Carlo in Bayesian

inference:

- 1. Parameter transition only needs the computation of likelihood function. Therefore, Monte Carlo Markov Chain can be used in general Bayesian inference, in which we are not guaranteed to have an analytical form of posterior.
- 2. Specifically in Metropolis-Hastings algorithm, marginal distribution is cancelled out, thus make Metropolis-Hastings a computationally efficient algorithm.

However, MH algorithm also has a drawback; its convergence becomes slower as the dimension of parameter θ increases.

1.2.4 Selection of prior distribution

Theoretically, prior can be of any distribution family. However, a selection of prior distribution that is too different than the actual distribution of parameter can leads to a false propagation of beliefs and degrade inference results. It is suggested by [9] that in case of no prior knowledge exists to help the selection of prior distribution, Uniform distribution is preferable since it is less likely to propagate false beliefs to the inference.

A systematic inference to select prior distribution family and prior distribution parameter (hyperparameters) is possible with *Hierarchical Bayes Models* [1].

1.2.5 Bayesian Parameter Estimation

With posterior distribution $\pi(\theta|D)$ we estimate the parameter $\hat{\theta}$ using Bayesian posterior mean:

$$\hat{\theta} = \mathbf{E}[\theta] = \int_{\theta} \theta \pi(\theta|D) d\theta$$

In case we have samples from posterior distribution, for example the Trace from Metropolis-Hastings algorithm, for example when we use MH algorithm, the discrete form of posterior mean is used:

$$\hat{\theta} = \mathbf{E}[\theta] = \sum_{\theta} \theta \pi(\theta|D)$$

1.2.6 Bayesian Parameter Estimation

Definition 1.5 (Bayesian Credible Set). Set C is a $(1\alpha)100\%$ credible set for the parameter θ if the posterior probability for θ to belong to C equals (1α) .

$$P(\theta \in C|D) = \int_C \pi(\theta|D)d\theta = 1 - \alpha$$

In this project, we use 0.95 credible set, i.e $\alpha = 0.05$

Definition 1.6 (Highest Posterior Density credible set). Highest Posterior Density $(1 - \alpha)100\%$ credible set (HPD for short) is the interval with minimum length over all Bayesian $(1 - \alpha)100\%$ Credible Set.

In this project, the HPD is calculated using algorithm from PyMC3 library [10]. For simplicity, we assume that in all cases which we concern, HPD is computed for unimodal distribution.

Algorithm 2 Compute Highest Posterior Density Interval

Input: S is samples from a distribution.

Input: $0 \le \alpha \le 1$

Output: HPD interval

1: **procedure** Compute HPD(S)

- 2: Compute interval width $w = |S| * \alpha$
- 3: Find modal (peak) of sample points.
- 4: Return minimal interval of size |S| w which contains the modal.
- 5: end procedure

2 Related works

The definition and model checking of DTMC and pMC is studied by [2], [5], and [7].

Bayesian inference of pMC parameters is studied in [9] and [6]. In [9], the authors developed methods to synthesize parameters to satisfy a given set of PCTL properties. In [6], the authors presented methods to perform model checking of biological system using Bayesian statistic. The authors in [6] uses a Bayesian hypothesis test, where H_0 is the null hypothesis that the model

satisfies a PCTL P, and alternative hypothesis H_1 is that the system does not satisfies P.

In this project, we use bee colony model semantics from [4]. The methods and implementation in this project is designed to extend the results of [4] and its tool DiPS.

3 Modeling of bees colony

3.1 Biological description

We consider the collective action of a bee colony. Each bee in a colony could possibly sting after observing a threat in the surrounding environment, and warn other bees by releasing pheromone. By sensing the pheromone released in the environment, other bees in the colony may also sting. Since stinging leads to the termination of an individual bee, it reduces the total defense capability as well. We studies how the actions of a bee changes with regarding to its surrounding the environment. There are 3 assumptions on the system:

- 1. Each bee release an unit amount of pheromone immediately after stinging.
- 2. A bee dies after stinging and releasing pheromone. In the other words, no bee can sting more than once.
- 3. Stinging behaviour only depends on the concentration of pheromone in the environment.

Under these assumption, a bee colony can be viewed as a set of agents (bees) interact with each other in a closed environment with the appearance of a factor *pheromone*. Each agent in the colony observe the following two factors of the environment: (1) amount of pheromone and (2) number of other agents. Afterward, the agent has probability to commit an action, namely *sting*. The agent is eliminated from environment after stinging.

3.2 Markov population models for bee colony

Assume that we have a colony of n bees initially. As aforementioned, an individual bee is terminated after it stings. Thus, at the end of experiment,

the number of bees is $n' \in \{0, 1, ..., n\}$. We model the bee colony with a DTMC $\mathcal{M} = (S, \mathbf{P}, S_{init}, AP, L)$, such that

- $|S_{init}| = 1$
- There exists n + 1 tSCCs which encode the population at the end of the experiment.

Semantics of Markov population models for bees colony are developed by [4]. In this report we summarize three models of bees population

- 1. Synchronous model
- 2. Asynchronous model
- 3. Semi-synchronous model

Asynchronous model

In asynchronous experiment we assume that there is almost improbable for two bees to sting simultaneously, and any bees release pheromone immediately after its death. By that observation we can assume that each bee sting at different level of pheromone.



Figure 1: Example asynchronous model of 3 bees, 2 parameters

Synchronous model

In *fully synchronous* experiment we assume that the number of stinging bees is only counted after a fixed amount of time, so that without loss of generality we can assume the pheromone diffuse almost immediately among the bee colony, and each bee decide to sting or not to sting immediately after sensing the pheromone concentration.

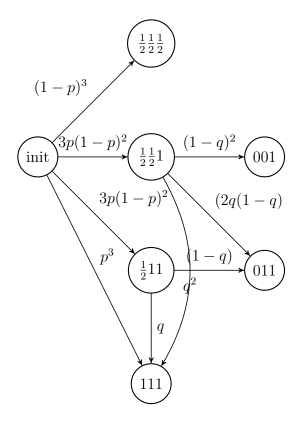


Figure 2: Synchronous model of 3 bees, 2 parameters.

Semi-synchronous model

In *Semisynchronous model*, we assume that the behaviour is initally synchronous. That means, at the initial states we do synchronous update. From all succeeding states from initial states, the updates are of asynchronous semantics.

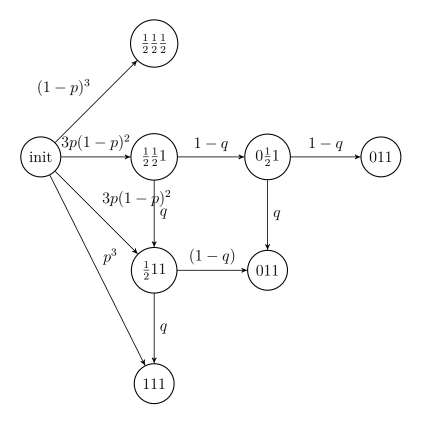


Figure 3: Semisynchronous model of 3 bees, 2 parameters

For a population of n bees, all three types of models share two properties:

- 1. Has exactly one initial state, $|S_{init}| = 1$. This assumption is obvious, since at the beginning of the experiment, all bees are alive.
- 2. Has n+1 tSCCs $(tSCC_0, \ldots, tSCC_n)$. This is because the number of dead bees cannot exceed the population size. It also follows that

$$\sum_{i=0}^{n} P(FG \quad tSCC_i) = 1$$

3.2.1 Multiparameters model

Multiparameters model of N bees has exactly N parameters r_0, \ldots, r_{N-1} . In the semantics of multiparameter models, we assume that r_0, \ldots, N is monotonically increasing, that is, $\forall i, j \in 0, \ldots, N-1 : i < j \implies r_i \leq r_j$.

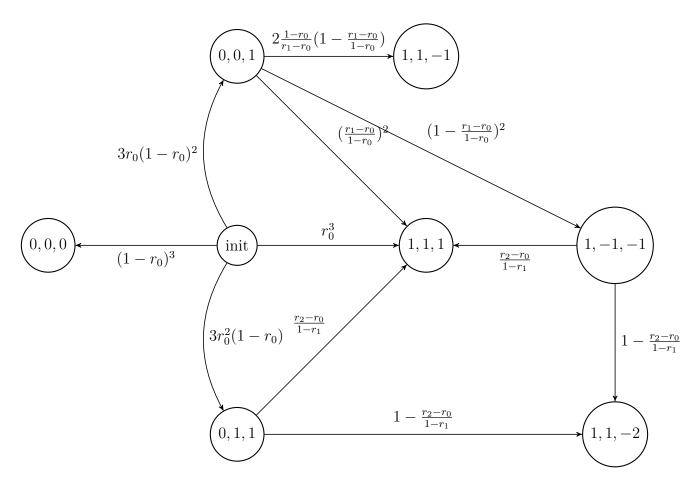


Figure 4: Synchronous model of 3 bees, multiparameters

3.2.2 Linear model

Linear model is similar to multiparameter models, except:

- There is no monotonically increasing constraint on r_0, \dots, r_{N-1}
- There are only 2 model parameters, denote as a and b
- Linear constraint:

$$r_i = a \times i + b$$

with $a, b \in [0, 1]$

i The purpose of imposing linear constraint to model parameters is to reduce the number of model parameters in total. In a linear model, the number of model parameters are always 2.

Bayesian parameter synthesis of Markov pop-4 ulation model

Inference of terminal state distribution 4.1

Using the Multinomial-Dirichlet conjugation, we can infer the terminal steady state distribution as we observe new experiment data $D = (a_0, \ldots, a_n)$, with a_i is the number of experiment in which the population at terminal state is i. Let $p_i = P(tSCC_i)$ be the terminal state distribution.

Algorithm 3 Estimation of terminal state distribution given a sample S

- 1: **procedure** Estimate $p_i = P(tSCC_i)(S)$
- Initialize $\alpha_0 = \alpha_1 = \ldots = \alpha_n = 1$
- Initialize $p_0 = p_1 = \ldots = p_n = 0$ 3:
- 4:
- Update $\alpha_i = \alpha_i + a_i, 1 \le i \le n$ Update $p_i = \alpha_i / \sum_{i=1}^n \alpha_i, 1 \le i \le n$
- 6: end procedure

As later shown in the implementation with synthetic data, the Bayesian estimation of terminal state distribution converges quickly to the true parameter used for data synthesis. This inference does not directly estimate the pMC model parameters. However, it can support the parameter estimation presented [4] to estimate the model parameters with narrower credible set. Furthermore, as the inference helps to estimate terminal state distribution with less data available.

4.2Evaluation of terminal states distribution

Given a concrete assignment of model parameter r_0, \ldots, r_{N-1} , there are 2 methods that are used in the project to evaluate terminal state distribution:

1. Rational functions.

2. DTMC sampling.

Rational functions are functions of model parameter that represent the probability of finally globally reach each terminal state. The function is delivered by PRISM model checker thanks to it capability of symbolic model checking [8].

However, it is not always possible to deduct rational functions from a given model, due to the technical limitation (time, memory) and the limitation of PRISM itself. In our conducted experiment, PRISM is capable of deliver rational functions up to a population of 15 bees. For a population of more bees, we use the second approach, *DTMC sampling*.

DTMC sampling has advantages over rational function. First, it is less computationally expensive to evaluate a parametric DTMC thanks to its simpler symbolic experession. Second, DTMC sampling is *parallelizable*; sampling can be done with as many processor cores as possible. The second advantage makes DTMC sampling *scalable*, compare to the rational function evaluation approach, which is not scallable due to its nature of deep recursion.

Algorithm 4 Evaluate terminal state distribution by DTMC sampling

```
Input: Population model \mathcal{M} of N individuals

Assignment of model parameter r

Number of DTMC samples nSamples

Output: D terminal state distribution

1: procedure Estimate p_i = P(tSCC_i)(S)

2: Initialize D = [0, \dots, 0] as an array of N + 1 entries

3: while not reached nSamples do

4: Get tSCC_i, 0 \le i \le N + 1 by simulating evaluated model \mathcal{M}

5: Increase D[i] by 1

6: end while

7: end procedure
```

4.3 Inference of model parameter with Metropolis-Hastings algorithm

We can also use Bayesian inference to estimate pMC model parameters directly. Given a pMC \mathcal{M} with $\theta = (\theta_1, \dots, \theta_K)$ as its model parameters, $\theta_i \in [0, 1]$. Note that $\theta_1, \dots, \theta_k$ are not simplex, thus it is not possible to use

Dirichlet prior in this case.

In this method, since there is no possible use of posterior conjugation, thus the hyperparameters estimation is not in the scope of this project. Let $f_i = P(FG \ tSCC_i)$ be estimation of terminal state distribution, we denote $\epsilon_i = f_i(\theta)$ as evaluations of the polynomial function with a concrete assignment of θ . The steady-state distribution has then $Multinomial(N, \epsilon = (\epsilon_0, \ldots, \epsilon_n))$ with N is the sample size of experiment data D. The posterior distribution has the following form:

```
\pi(\theta|D) \sim Multinomial(N, (\epsilon_0, \dots, \epsilon_n))\pi(\theta_1) \dots \pi(\theta_k)
```

As the posterior $\pi(\theta|D)$ has no analytical form, we use Metropolis-Hastings to sample from it.

Algorithm 5 Estimation of pMC parameters given a sample D

```
1: procedure METROPOLIS-HASTINGS(D, maxIteration)
        Select a proposal distribution \pi(\theta)
 2:
        Draw a random initial point \theta
 3:
 4:
        Init empty trace Trace
        while maxIteration not reached do
 5:
            L \leftarrow P(D|\theta)
 6:
            Draw a point \theta' from the proposal distribution.
 7:
            L' \leftarrow P(D|\theta')
 8:
            if \ln(L') - \ln(L) > 0 then
 9:
                Add \theta' to Trace
10:
                \theta = \theta'
11:
            else
12:
                Draw a random number x from Uniform(0,1)
13:
                if x \le \epsilon, (\epsilon very small, e.g 10^{-8}) then
14:
                    Add \theta' to Trace (avoiding local maxima)
15:
                    \theta = \theta'
16:
                end if
17:
18:
            end if
        end while
19:
20: end procedure
```

From the Trace returned by the algorithm, we estimate parameter

5 Benchmark

5.1 Data synthesis

In order to better evaluate the performance of Bayesian estimation of model parameters, we use synthetic data, generated from a model with known parameters. In case of rational functions are available, data synthesis for N bees population is conducted as follow:

- Select a pMC model \mathcal{M} to model the population of. Let $\theta = \theta_1, \ldots, theta_k$ is the model parameter.
- Assign a concrete value for θ which satisfies model's constraint for parameter, θ_{true}
- From \mathcal{M} , evaluate terminal state distribution of $f_i(\theta) = P(FG \ tSCC_i)$, $0 \le i \le N$.
- Draw sample S from multinomial distribution $Multinomial(N, (f_0(\theta), \dots, f_N(\theta)))$

Synthetic data can also be obtained directly from DTMC sampling. The using of synthetic data and known true parameter allow us to evaluate the inference result more precisely.

```
Model file: data/prism/bee_multiparam_synchronous_5.pm
Result file: data/prism/bee_multiparam_synchronous_5.txt
True parameters: [0.25, 0.3, 0.35, 0.45, 0.5]
Terminal state distribution:
    tSCC[0]=0.2373046875
    tSCC[1]=0.300125
    tSCC[2]=0.24029687500000008
    tSCC[3]=0.121000000000000022
    tSCC[4]=0.0769531249999996
    tSCC[5]=0.02432031249999996

Synthetic data
    [25, 27, 24, 12, 10, 2]
Synthetic data distribution
    [0.25, 0.27, 0.24, 0.12, 0.1, 0.02]
```

Figure 5: Example of data synthesis for 5 bees model.

5.2 Evaluation measures

In order to evaluate the inference result, we consider three aspects:

- 1. Accuracy: distance from estimated parameter to true parameter used to synthesize sample data.
- 2. *Precision:* width of the Highest Posterior Density corresponding to an estimation.
- 3. Cost: how does physical computation time increases as the size of population increases.

To measure *accuracy*, the distance from estimated parameter to true parameter, we use *Root Mean-Squared Error*.

Definition 5.1 (Root Mean-Square Error). Let $\theta = (\theta_1, \dots, \theta_n)$ and $\hat{\theta} = (\hat{\theta}_1, \dots, \hat{\theta}_n)$ be vectors of n real numbers, the Root Mean Square Error between p and \hat{p} is defined as follow:

$$RMSE(\theta, \hat{\theta}) = \sqrt{\frac{\sum_{i=1}^{n} (\theta_i - \hat{\theta}_i)}{n}}$$

In the following experiments, we use the three measures (accuracy, precision, and cost) to evaluate inference results.

5.3 Inference of terminal state distribution

5.3.1 Experiment setup

This experiment is conducted on multiparameters, synchronous model of 5 bees.

- Model file: bee multiparams synchronous5.pm
- PRISM result file: bee_multiparams_synchronous5.txt
- True parameter: [0.25, 0.3, 0.35, 0.45, 0.5]
- True tSCC distribution: [0.2373046875, 0.300125, 0.240296875, 0.1210, 0.076953125, 0.0243203125]

In order to compare with the work by [4], we also visualize the interval calculated by the following formula:

$$\theta_i \pm (z_{\alpha/2} \sqrt{\frac{\theta_i (1 - \theta_i)}{N}})$$

where N is the sample size.

5.3.2 Results

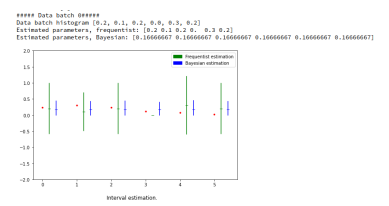


Figure 6: First batch of data. The Bayesian conjugation method suffers from initial fase beliefs.

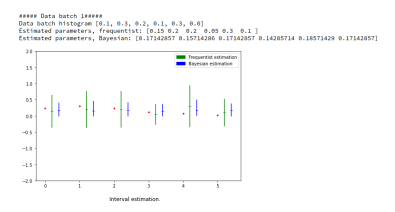


Figure 7: Second batch. Estimation changes significantly.

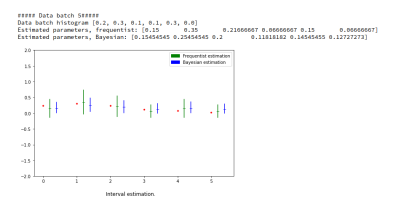


Figure 8: Sixth batch. Estimation improved; Bayesian conjugation gives narrower confidence intervals.

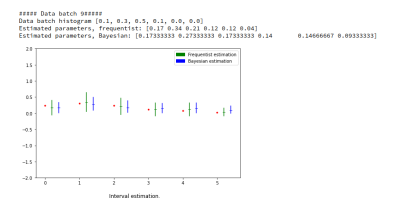


Figure 9: Tenth batch

5.4 Compare DTMC sampling and rational functions

5.4.1 Experiment setup:

5.4.2 Results

Sampling results

Runtime

Suggested number of chainrun: 1000 * number of bscc

- 5.4.3 Analysis of long run property
- 5.5 Inference of model parameters
- 5.5.1 Experiment setup:
- 5.5.2 Results

Sampling results Runtime

- 5.6 Impacto of Metropolis-Hastings parameter selection
- 5.7 Impact of data quality to inferences
- 5.8 Linear model
- 6 Implementation
- 6.1 Class design

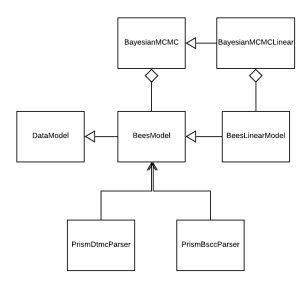


Figure 10: Class diagram of the project

Class explanation:

- 1. PrismModelParser and PrismBsccParser parses PRISM model file and symbolic model checking results. In details, the two classes do the following tasks:
 - Read PRISM model and result files.
 - Extract and preprocess expressions from model and result.
 - Factorize expressions if needed
 - Parse expressions in model and result file. store in Python byte-code.
 - Create DataModel object to store these parsed expressions.
- 2. DataModel, BeesModel, and BeesLinearModel store information about the population model.
- 3. BayesianMcmc and BayesianMcmcLinear implements Metropolis-Hastings algorithm.

6.2 Test environment

The Bayesian inference methods are implemented in Python 3 and tested in the followin system configuration:

- Intel Core i5-8265U @ 1.60GHz
- 16GB RAM
- OpenSUSE Tumbleweed 20200427
- Anaconda 3 2019.10 for linux x86 64

6.3 Directory structure

- data folder contains PRISM models and model checking results.
- docs folder contains final report
- scripts folder contains python implementation.
- examples folder contains experiment schemes.

7 Conclusion

The goal of this project is to experiment Bayesian approach on parameter inference of Markov population models. The results shows that Bayesian inference is efficient to deliver an estimation of model parameters. Given that in the biological system that is modelled, each parameters represent a real-world biological property, such as the probability of a bee sting, given a certain amount of pheromone, the parameter estimation presented in this project helps biologists to better understand the biological mechanism.

As it is shown on benchmark, the implementation in this project is capable of estimating parameters in both cases, when rational functions is available and is not available. As the size of rational function increases exponentially starting from populations of 15 bees, and PRISM fails to give rational function for model of 20 bees, the implementation in this project, which can work given only the model, shows an advantage. Also, with the parallelization of DTMC sampling, the implementation proves that it has significant scalability.

However compare to parameter synthesis method presented in [4], the implementation in this project does not give a detailed analysis of parameter space. Thus, [4] has several advantage for model of less than 15 bees, especially in the case of 2-parameters model, where it shows a very detailed analysis of accepted regions and rejected regions on parameter space.

There are still problems, which are not completely solved in the scope of this project:

- 1. Number or parameters: Accuracy and precision of an estimation decreases, while the computational cost increases, as the number of model parameter increases.
- 2. Selection of prior: a false selection of prior distribution and its parameters could propagate false beliefs over the whole inference. So far we have been using Uniform distribution by default to prevent However, this project has not coped with the problem of hyperparameters estimation.

References

- [1] Greg M Allenby, Peter E Rossi, and RE McCulloch. "Hierarchical Bayes Models: A Practitioners Guide. Grover R, Vriens M, eds". In: SSRN Electron J (2005).
- [2] Christel Baier and Joost-Pieter Katoen. *Principles of model checking*. MIT press, 2008.
- [3] Michael Baron. Probability and statistics for computer scientists. CRC Press, 2019.
- [4] Matej Hajnal et al. "Data-Informed Parameter Synthesis for Population Markov Chains". In: *International Workshop on Hybrid Systems Biology*. Springer. 2019, pp. 147–164.
- [5] Lisa Hutschenreiter, Christel Baier, and Joachim Klein. "Parametric Markov chains: PCTL complexity and fraction-free Gaussian elimination". In: arXiv preprint arXiv:1709.02093 (2017).
- [6] Sumit K Jha et al. "A bayesian approach to model checking biological systems". In: *International conference on computational methods in systems biology*. Springer. 2009, pp. 218–234.
- [7] Joost-Pieter Katoen. "The probabilistic model checking landscape". In: Proceedings of the 31st Annual ACM/IEEE Symposium on Logic in Computer Science. 2016, pp. 31–45.
- [8] M. Kwiatkowska, G. Norman, and D. Parker. "PRISM 4.0: Verification of Probabilistic Real-time Systems". In: Proc. 23rd International Conference on Computer Aided Verification (CAV'11). Ed. by G. Gopalakrishnan and S. Qadeer. Vol. 6806. LNCS. Springer, 2011, pp. 585–591.
- [9] Elizabeth Polgreen et al. "Data-efficient Bayesian verification of parametric Markov chains". In: *International Conference on Quantitative Evaluation of Systems*. Springer. 2016, pp. 35–51.
- [10] John Salvatier, Thomas V Wieckiâ, and Christopher Fonnesbeck. "PyMC3: Python probabilistic programming framework". In: *ascl* (2016), ascl—1610.
- [11] Stephen Tu. "The dirichlet-multinomial and dirichlet-categorical models for bayesian inference". In: Computer Science Division, UC Berkeley (2014).