Sampling rare fluctuations of discrete-time Markov chains

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We describe a simple method that can be used to sample the rare fluctuations of discrete-time Markov chains. We focus on the case of Markov chains with well-defined steady-state measures, and derive expressions for the large-deviation rate functions (and upper bounds on such functions) for dynamical quantities extensive in the length of the Markov chain. We illustrate the method using a series of simple examples, and use it to study the fluctuations of a lattice-based model of active matter that can undergo motility-induced phase separation.

I. INTRODUCTION

Quantifying the rare behavior of stochastic models is necessary to understand important physical processes such as phase separation and chemical reactions [1]. Computer simulation methods for quantifying rare behavior include umbrella sampling [2]; forward-flux sampling [3]; transition-path sampling [4]; and diffusion Monte Carlo methods [5,6], which often involve the use of an auxiliary or reference model whose typical behavior is in some sense equivalent to the rare behavior of the model of interest [5,7,8,10–15]. Motivated by this latter approach, we show that the rare fluctuations of a discrete-time Markov chain generated by a stochastic model of interest can be quantified in a simple way using a stochastic reference model whose parameters are simply related to those of the original model. Partial information about the original model’s rare fluctuations – meaning a bound on the large-deviation rate function for time-extensive observables – can be obtained from typical trajectories of the reference model, while complete information about the original model’s rare fluctuations can be obtained by considering constrained fluctuations of the reference-model trajectory ensemble. In Ref. [10] we use this procedure to study the rare behavior of models of growth; here we focus on Markov chains with well-defined steady-state measures.

In what follows we describe the method, and use a series of simple examples to illustrate the method’s application to Markov chains with well-defined steady-state measures (Section II). We then apply the method to the lattice-based model of active matter introduced in Ref. [17], in order to quantify that model’s rare dynamic fluctuations (Section III). We conclude in Section IV.

II. SAMPLING THE RARE FLUCTUATIONS OF DISCRETE-TIME MARKOV CHAINS

Consider a discrete-time Markov chain of \(K\) steps. At a given instant the system is in microstate (or ‘state’) \(C\), and with probability \(p(C \rightarrow C')\) moves to state \(C'\) (which may be \(C\)). Probability conservation requires that \(\sum_{C'} p(C \rightarrow C') = 1\). A trajectory \(x\) is a sequence of microstates \(x = \{C_0, C_1, C_2, \ldots, C_K\}\) generated by this dynamics, and its probability is

\[
P[x] = P_0(C_0) \prod_{k=0}^{K-1} p(C_k \rightarrow C_{k+1}),
\]

where \(P_0(C_0)\) is the probability of starting in state \(C_0\). The master equation for this discrete Markov chain is

\[
P(k + 1) = WP(k),
\]

where \(P(k)\) is the column vector whose elements are \(P(i,k)\), the probability of residing in state \(i\) after \(k\) steps of the dynamics, and \(W\) is the matrix whose \((j,i)\)th element is \(p(i \rightarrow j)\). For an \(M\)-state system, \(P\) has \(M\) elements and \(W\) has \(M \times M\) elements.

Consider now an observable \(A[x]\), called the ‘activity’, extensive in the length of the trajectory [12]. Upon moving from \(C\) to \(C'\) we increment \(A\) by an amount \(\alpha(C \rightarrow C')\), so that \(A[x] = \sum_{k=0}^{K-1} \alpha(C_k \rightarrow C_{k+1})\). Our aim is to compute the probability

\[
\rho(a, K) = \sum_x P[x] \delta(A[x] - Ka)
\]

that a trajectory of length \(K\) possesses a particular value \(a = A/K\) of the intensive counterpart of the observable \(A\). For many models this probability adopts a large-deviation form \(\rho(a, K) \sim e^{-K I(a)}\) for large values of \(K\), where \(I(a)\) is the large-deviation rate function [11]. Eq. (3) is the instruction to simulate the original model and count the trajectories that possess activity \(A = Ka\). When \(a\) is close to a ‘typical’ value \(a_0\), for which \(I(a_0) = 0\), such sampling is efficient. But when \(a\) is far from \(a_0\), so that it is rarely generated by direct simulation, a different strategy is required.

To motivate this strategy, note that the desired quantity \(\rho(a, K)\) can be regarded as the normalization factor (or ‘partition function’) of the microcanonical path ensemble [18,21]

\[
P_a[x] = \frac{P[x] \delta(A[x] - Ka)}{\rho(a, K)},
\]

which defines an ensemble of trajectories conditioned upon the constraint \(A[x] = Ka\). (The term ‘microcanonical’ comes from analogy with the fixed-energy ensemble
of equilibrium statistical mechanics; here it is activity that is fixed). Alternatively one can define a canonical path ensemble (or tilted ensemble or s-ensemble) \[11\] in which \( A[x] \) can fluctuate,

\[
P_s[x] = \frac{e^{-sA[x]}P[x]}{\sum_x e^{-sA[x]}P[x]}.
\]

The master equation in the s-ensemble is

\[
P_s(k + 1) = W_sP_s(k),
\]

where \( W_s \) is the matrix whose \((j,i)\)th element is

\[
p_s(i \rightarrow j) = e^{-sa(i \rightarrow j)}p(i \rightarrow j).
\]

From the canonical path ensemble one can in certain cases recover the desired quantity \( p(a, K) \) \[20, 21\]. The logarithm of the normalization factor in Eq. (3), \( \lambda(s) \equiv K^{-1}\log \sum_x e^{-sA[x]}P[x] \), can be obtained by taking the logarithm of the principal eigenvalue of the generator \( W_s \) of s-ensemble dynamics \[11, 12\]. Legendre transform of \( \lambda(s) \) yields a rate function

\[
I_s(a) = \max_s (-sa - \lambda(s)),
\]

which is equal to the true rate function \( I(a) \) when the latter is convex \[11, 22, 23\]. The s-ensemble has been used with great success, providing insight into the behavior of a large number of models \[12\], but it possesses two drawbacks. One is that \( W_s \) is not a stochastic generator (its columns do not sum to unity, i.e. \( \sum_j e^{-sa(i \rightarrow j)}p(i \rightarrow j) \neq 1 \)), and so considerable effort and ingenuity is sometimes needed to identify and simulate the s-ensemble \[5, 7, 8, 11, 15\]. More seriously, the calculation of probabilities is indirect, and fails when the rate function \( I(a) \) of the original model is non-convex \[11, 22, 23\]. This is so in many physically interesting cases, where there exist phase transitions and coexistence of trajectories. Here \[8\] returns only the convex hull of the true rate function \[11\]: information about the trajectory ensemble is missing.

The method described here and in Ref. \[16\] draws inspiration from the s-ensemble method but calculates \( p(a, K) \) directly, and so can recover non-convex rate functions. It is related to the method used in Ref. \[24\], generalized to Markov chains and applied to a probability-conserving model. Consider a stochastic reference model whose transition probabilities \( p_{ref}(i \rightarrow j) \) are normalized versions of those of the s-ensemble, \( p_s(i \rightarrow j)/\sum_j p_s(i \rightarrow j) \) (the idea of using such a model is noted in Eq. (4.24) \[25\]). The master equation of the reference model is then

\[
P_{ref}(k + 1) = W_{ref}P_{ref}(k),
\]

where element \((j,i)\) of \( W_{ref} \) is

\[
p_{ref}(i \rightarrow j) = \frac{e^{-sa(i \rightarrow j)}p(i \rightarrow j)}{\sum_j e^{-sa(i \rightarrow j)}p(i \rightarrow j)}.
\]

Note that the columns of \( W_{ref} \) sum to unity - \( \sum_j p_{ref}(i \rightarrow j) = 1 \) - and so \( W_{ref} \) is stochastic (probability-conserving). The idea of the scheme is then as follows. The reference model is not the s-ensemble, and is not intended to be. Simulations of the reference model (parameterized by \( s \)) produce values of \( a \) that for large \( K \) concentrate on \( a_s \), where \( a_s \) minimizes the rate function \( I_{ref}(a) \) of the reference model. Carrying out such simulations, and using the equations shown below, we calculate a piece \( I(a_s) \) of the rate function of the original model. Repeating the process for a different value of \( s \) yields another piece of \( I(a) \), and so on. We do not generate the microcanonical or canonical path ensembles. Instead, a simple modification of the original model (which admits all the processes of that model and no new ones) allows us to control which value of \( a \) is sampled, and from comparison of original and reference models we calculate the likelihood, Eq. (3), of generating those values of \( a \) using the original model.

To carry out this calculation note that Eq. (3) can be written

\[
\rho(a, K) = \sum_x P_{ref}[x]w[x]\delta(A[x] - Ka),
\]

where

\[
P_{ref}[x] = P_0(C_0) \prod_{k=0}^{K-1} p_{ref}(C_k \rightarrow C_{k+1})
\]

is the trajectory weight of the reference model, and \( w[x] \equiv P[x]/P_{ref}[x] \). It is convenient to write the transition probabilities of the original and reference models as \( p(C \rightarrow C') = W(C \rightarrow C')/R(C) \) and \( p_{ref}(C \rightarrow C') = W_{ref}(C \rightarrow C')/R_{ref}(C) \), where \( W(C \rightarrow C') \) and \( W_{ref}(C \rightarrow C') = e^{-sa(C \rightarrow C')}W(C \rightarrow C') \)

are rates for the original and reference models, respectively, and \( R(C) \equiv \sum_{C'} W(C \rightarrow C') \) and \( R_{ref}(C) \equiv \sum_{C'} W_{ref}(C \rightarrow C') \) are the models’ escape rates from a given state.

We then have \( w[x] = \prod_{k=0}^{K-1} w_k \), where

\[
w_k = e^{sa(C_k \rightarrow C_{k+1})}R_{ref}(C_k) / R(C_k),
\]

It is also convenient to define the quantity

\[
q[x] = K^{-1} \sum_{k=0}^{K-1} \ln R_{ref}(C_k) / R(C_k),
\]

using the pieces of \( w[x] \) not fixed by the delta-function path constraint.

The sum in Eq. (11) is the instruction to simulate the reference model, and take the arithmetic mean of values of \( w \) for trajectories that display activity \( a \) (the constraint imposed by the delta function). It is natural to consider the case \( a = a_s \), where \( a_s \) is a value typical of the reference model (for which \( I_{ref}(a_s) = 0 \). If we generate \( N \)
trajectories of the reference model, and the trajectories labeled \( i = 1, 2, \ldots, \mathcal{M} \leq \mathcal{N} \) are typical in this way, then \(^{[11]}\) reads

\[
\rho(a_s, K) = \frac{\mathcal{M}}{\mathcal{N}} \cdot \frac{1}{\mathcal{M}} (w_1 + \cdots + w_\mathcal{M}).
\]

From \(^{[14]}\) and \(^{[15]}\) we have \( w_k = e^{\kappa_K a_k K q} \), where \( \kappa \) labels trajectories. Upon taking logarithms of \(^{[16]}\) and assuming \( K \) large, so that \( K^{-1} \ln(\mathcal{M}/\mathcal{N}) \) is negligible, we get

\[
I(a_s) = -s a_s - q_s - K^{-1} \ln \int dq P_s(q|a_s) e^{Kq}.
\]

Here \( I(a_s) = -K^{-1} \ln \rho(a_s, K) \) is the large-deviation rate function of the original model evaluated at \( a_s \); \( q_s = \int dq P_s(q|a_s) \) is the mean value of \( q(x) \) for the ensemble of typical reference-model trajectories (those with \( A(x) = K a_s \)); \( P_s(q|a_s) \) is the probability distribution of \( q \) for the ensemble of typical reference-model trajectories; and \( q = q - q_s \).

By Jensen’s inequality the first two terms in \(^{[17]}\) provide a bound on the rate function \( I_0(a_s) \geq I(a_s) \), where

\[
I_0(a_s) = -s a_s - q_s.
\]

The full rate function can be recovered by evaluation of the integral, which is demanding in general \(^{[21]}\) (e.g. requiring a high-order cumulant expansion) but straightforward when \( P_s(q|a_s) \) is Gaussian or close to it (this is not guaranteed in general, but is realized in several cases we have encountered \(^{[10]}\)).

The computational cost \( C^{(0)} K \) of generating a trajectory of length \( K \) of the reference model is in general greater than the cost \( C^{(0)} K \) of generating a trajectory of similar length using the original model, because to use the reference model we must calculate the bias parameters \( \alpha(C \rightarrow C') \). However, this cost is usually more than offset by our ability to direct the reference model to rarely-sampled values \( a_s \) of the dynamic parameter: the cost to sample the piece \( I(a_s) \) of the rate function using the original model is \( C = C^{(0)} K e^{K I(a_s)} \), while the cost using the reference model is \( C_{\text{ref}} = C^{(0)} K e^{K I_{\text{ref}}(a_s)} = C^{(0)} K \). Under most conditions we have encountered, the reference model is much more efficient than the original model. E.g. for the lattice model of Section \(^{[III]}\), \( \alpha(C \rightarrow C') \) can be calculated via spatially local updates (updates involving only the neighborhood of the chosen lattice site), and the ratio \( C^{(0)}/C^{(0)} \) does not depend on the size of the system (it is of order 10). In this case the ratio \( C/C_{\text{ref}} = (C^{(0)}/C^{(0)}) e^{K I(a_s)} \) becomes large for any appreciable \( K \) (and nonzero \( I(a_s) \)), and so the reference-model method is more efficient than direct simulation. If for instance the computation of the bias \( \alpha(C \rightarrow C') \) required updates of the whole lattice, and so \( C^{(0)} \) grew with system size \( \mathcal{N} \), then there would exist an interval of \( K \) for which direct simulation may be more efficient.

In what follows we specialize the discussion to ergodic (irreducible, aperiodic) Markov chains possessing well-defined steady-state measures (probability of occupancy) \( \pi(C) \) and \( \pi_{\text{ref}}(C) \). These are obtained by requiring \(^{[2]}\) and \(^{[9]}\) to be stationary, and satisfy

\[
\pi_{\text{ref}}(C) = \sum_{C'} \pi_{\text{ref}}(C') p(C' \rightarrow C).
\]

The existence of such a measure is guaranteed when each microscopic process has a non-vanishing probability of reversal, which is true of the model of Ref. \(^{[17]}\), but not true, for instance, of irreversible growth processes \(^{[10]}\).

Eq. \(^{[17]}\) can be written

\[
I(a_s) = -K^{-1} \ln \langle \exp(\ln w_1 + \cdots + \ln w_K) \rangle_{\text{ref}},
\]

where \( \langle \cdot \rangle_{\text{ref}} \) denotes the average over the reference-model dynamics (for trajectories such that \( a = a_s \)). By Jensen’s inequality we have Eq. \(^{[18]}\), or

\[
I_0(a_s) = -K^{-1} \ln \langle \ln w_1 + \cdots + \ln w_K \rangle_{\text{ref}}
\]

\[
= -\langle \ln w \rangle_{\text{ref}}.
\]

The average \( \langle \ln w \rangle_{\text{ref}} \) can be calculated by keeping track of the terms \( \ln w_i \) associated with each move in a reference-model simulation,

\[
\theta(K) = K^{-1} \sum_{k=0}^{K-1} \ln w_k.
\]

For large \( K \), the series \( \theta(K) \) will converge to \( \theta_s \), where

\[
\theta_s = s a_s + q_s
\]

\[
= a \sum_{C} \pi_{\text{ref}}(C) \sum_{C'} p_{\text{ref}}(C \rightarrow C') \alpha(C \rightarrow C')
\]

\[
+ \sum_{C} \pi_{\text{ref}}(C) \ln \frac{R_{\text{ref}}(C)}{R(C)}
\]

\[
= \sum_{C} \pi_{\text{ref}}(C) \left\{ -s \frac{\partial}{\partial s} \ln R_{\text{ref}}(C) + \ln \frac{R_{\text{ref}}(C)}{R(C)} \right\}.
\]

Note that for the choice \( \alpha(C \rightarrow C') = B(C') \), made in Ref. \(^{[17]}\), we have

\[
a_s = \sum_{C} \pi_{\text{ref}}(C) \sum_{C'} p_{\text{ref}}(C \rightarrow C') B(C')
\]

\[
= \sum_{C'} B(C') \sum_{C} \pi_{\text{ref}}(C) p_{\text{ref}}(C \rightarrow C')
\]

\[
= \sum_{C} \pi_{\text{ref}}(C) B(C),
\]

in which case

\[
\theta_s = \sum_{C} \pi_{\text{ref}}(C) \left\{ s B(C) + \ln \frac{R_{\text{ref}}(C)}{R(C)} \right\}.
\]

For simple models for which states \( C \) can be enumerated explicitly, the above expressions provide an analytic
A simple example in which the Ref. [16].

The rate function calculated using the reference-model method is not constrained to be convex: Eq. (17), which is a direct evaluation of a probability, follows from simple arguments that do not depend on the form of the rate function in question. Convexity of the s-ensemble rate function, $I_s(a)$, is guaranteed by the structure of Eq. (3), which imposes a particular relationship between the two terms in that expression. Upon maximization, the quantity $a$ in the first term of (3) is given by the derivative of the second term, $-\partial \lambda(s) / \partial s$ (where this derivative exists). The same relationship does not exist between the terms of Eq. (17), or between the terms of its bound, Eq. (18), $I_0(a_s) = -sa_s - q_s$. To see this, note from Eq. (25) that

$$ a_s = - \sum_{C'} \pi_{\text{ref}}(C) \partial_s \ln R_{\text{ref}}(C), \quad (30) $$

and

$$ q_s = \sum_{C} \pi_{\text{ref}}(C) \ln[R_{\text{ref}}(C)/R(C)]. \quad (31) $$

Thus

$$ - \partial_s q_s = a_s - \sum_{C} (\partial_s \pi_{\text{ref}}(C)) \ln[R_{\text{ref}}(C)/R(C)], \quad (32) $$

and so $a_s$ and $q_s$ are not related in the way they would be if they were the elements of a Legendre transform (except in the special case of constant exit-rate ratio $R_{\text{ref}}(C)/R(C)$).

Thus the rate-function bound [18], which is simple to calculate, can be non-convex and so can give information about coexistence of trajectories (see e.g. Fig. 4 of Ref. [16]).

A. Example 1

To illustrate the reference-model method we start with a simple example in which the s-ensemble and reference-model method are equivalent. Consider the unbiased coin-toss model, represented in Fig. 1 as a two-state Markov chain. The microstates 0 and 1 correspond to a tail or a head, respectively, and the probability of generating either is 1/2. The master equation in the constant-K ensemble, Eq. (2), has generator (matrix of transition probabilities)

$$ W = \begin{pmatrix} 1 & 1 \\ 1 & 1 \end{pmatrix}. \quad (33) $$

We define a dynamic observable $A$ that increases by $+1$ upon any move into microstate 1, and by $-1$ upon any move into microstate 0. The observable $a = A/K$ is then the total number of heads minus the total number of tails, divided by the number of coin tosses, and its rate function is [20]

$$ I_{\text{coin}}(a) = \frac{1 - a}{2} \ln (1 - a) + \frac{1 + a}{2} \ln (1 + a). \quad (34) $$

The s-ensemble can be used to derive this result [11]. The s-ensemble of the coin-toss model has generator

$$ W_s = \begin{pmatrix} e^s & e^s \\ e^{-s} & e^{-s} \end{pmatrix}, \quad (35) $$

whose principal eigenvalue is $\cosh s$. Then $\lambda(s) = \ln \cosh s$, and Eq. (8) yields the rate function

$$ I_s(a) = \max_s (-sa - \ln \cosh s) $$$$ = a \tanh^{-1} a - \ln \cosh \tanh^{-1} a $$$$ = \frac{1 - a}{2} \ln (1 - a) + \frac{1 + a}{2} \ln (1 + a), \quad (36) $$

as expected.

Consider now the reference-model dynamics, which is generated by

$$ W_{\text{ref}} = \begin{pmatrix} e^s & e^s \\ e^{-s} & e^{-s} \end{pmatrix} / (2 \cosh s). \quad (37) $$

This stochastic process can be simulated using standard methods [27], in order to calculate Eq. (17) numerically [16]. For this problem the evaluation can also be done analytically. The steady-state measure of the reference model is $\pi(0) = e^s / (2 \cosh s)$ and $\pi(1) = 1 - \pi(0)$, and the typical activity is $a_s = \sum_{C'} \pi(C) \sum_{C' = C' \text{ ref}} (C \to C') \alpha(C \to C') = -\ln \cosh s$ (thus $a_s$ and $s$ have the relationship that $a_s$ and $s$ possess in Eq. (36)). The exit-rate ratio is always $q_s = \ln (R_{\text{ref}}(C)/R(C)) = \cosh s$, and so Eq. (17) is

$$ I(a_s) = a_s \tanh^{-1} a_s - \ln \cosh \tanh^{-1} a_s $$$$ = \frac{1 - a_s}{2} \ln (1 - a_s) + \frac{1 + a_s}{2} \ln (1 + a_s), \quad (38) $$

giving us one point on the rate-function curve $I_{\text{coin}}(a)$. Repeating the method for a range of values of $s$ (which here amounts simply to replacing $a_s \to a$) allows us to reconstruct the full rate-function curve.
Fig. 2. For the 4-state model shown we compute the large-deviation rate function $I(a)$ associated with the order parameter $a = A/K$, where $A$ is incremented by 1 for every transition along the blue arrows in the course of a trajectory of $K$ steps. The green line is the result obtained using the $s$-ensemble. The blue and red lines on the right are upper bounds [18] on the rate function computed using the reference-model method. We keep track of explicit configuration changes during numerical simulations (red line, Eq. (22)) or evaluate analytically Eq. (24) = Eq. (13) (blue line). The black dashed line is obtained by evaluating Eq. (41), which accounts for fluctuations of the weight for typical reference-model trajectories.

B. Example 2

A second example illustrates a case in which the $s$-ensemble and reference-model ensemble are different (but both can be used to obtain $\rho(a, K)$). Consider the 4-state model shown in Fig. 2, with each rate set to unity (indicated are the transition probabilities, all 1/2). Define the activity $A$ as the number of configuration changes from state 1 to 2 or state 2 to 1 in the course of a simulation of $K$ steps. By symmetry of the network the mean number of such configuration changes will be $A = K/4$, but some trajectories will exhibit more changes, and some less. We can use the $s$-ensemble and the reference-model method to calculate the logarithmic probability distribution $I(a) = -K^{-1} \ln \rho(a, K)$ associated with the quantity $a = A/K$.

The generator of the original model is

$$W = \begin{pmatrix} 0 & 1/2 & 0 & 1/2 \\ 1/2 & 0 & 1/2 & 0 \\ 0 & 1/2 & 0 & 1/2 \\ 1/2 & 0 & 1/2 & 0 \end{pmatrix},$$

and that of the $s$-ensemble is

$$W_s = \begin{pmatrix} 0 & e^{-s} \pi ref(1) & 0 & 1/2 \\ e^{-s} \pi ref(2) & 0 & 1/2 & 0 \\ 0 & 1/2 & 0 & 1/2 \\ 1/2 & 0 & 1/2 & 0 \end{pmatrix}.$$  

The principal eigenvalue of $W_s$ is $e^{\lambda(s)} = \frac{1}{4} \pi ref(1) \pi ref(2) e^{-s} \pi ref(1) \pi ref(2) + 5e^{2s} + 1 + e^{-s} + 1$, from which, using Eq. (8), $I(a)$ can be calculated numerically. The result is the green line in Fig. 2.

The reference-model generator is

$$W_{ref} = \begin{pmatrix} 0 & e^{-s} \pi ref(1) & 0 & 1/2 \\ e^{-s} \pi ref(2) & 0 & 1/2 & 0 \\ 1/2 & 0 & 1/2 & 0 \\ 1/2 & 0 & 1/2 & 0 \end{pmatrix},$$

which can be simulated using standard stochastic methods [27]. For a range of values of $s$ (from $-3$ to $3$, in intervals of 0.25) we carried out a single simulation of the reference model of $K = 5 \times 10^8$ configuration changes. We computed the rate-function bound [18] by a) keeping track of explicit configuration changes, as in Eq. (22), and b) by evaluating Eq. (24). To evaluate the latter, note that the steady-state measure is

$$\pi ref(1) = \pi ref(2) = \frac{1 + e^{-s}}{2(3 + e^{-s})}. $$

The typical activity of the reference model is then $a_s = (\pi ref(1) + \pi ref(2)) \times e^{-s}/(1 + e^{-s})$, from which we obtain $s = \ln[(1/a_s - 1)/3]$. The relationship between $a_s$ and $s$ is different to the relationship $a = -\theta_{s}(s)$ obtained from the $s$-ensemble: here the two methods produce different typical activity for given $s$.

From Eq. (24) we have

$$I_0(a) = -\theta_{s}(a) = -a \ln \frac{1 - a}{3a} - \frac{1 + 2a}{3} \ln \frac{1 + 2a}{2 - 2a}. $$

As shown in Fig. 2, both ways of calculating $I_0(a)$, analytically using Eq. (43) and numerically using [22], the blue and red lines in the figure, give the same result.

The full rate function can be estimated by evaluating numerically the integral in Eq. (17). To do so we took
each trajectory of $5 \times 10^8$ steps and split it into $5 \times 10^3$ pieces of length $K' = 10^3$. Of that set we retained only those pieces with extensive order parameter $A' = K' \sigma_s$ (it is necessary to explicitly enforce the delta-function constraint of Eq. (11) for short trajectories). The resulting distribution of $q_i P(q_i \mid a_i)$, is Gaussian with variance $\sigma_s^2$. In this case Eq. (17) reads

$$I(a_i) \approx -s a_i - q_i - K \sigma_s^2/2.$$  \hspace{1cm} (44)

Evaluation of Eq. (44) gives the dotted black line in Fig. 2 consistent with the result calculated via the s-ensemble.

**C. Example 3**

A third example illustrates some numerical difficulties associated with obtaining a rate function via Legendre transform, even when the rate function is convex. Consider the two-state Markov chain of example IV.4 of I1 (see also 29), shown in Fig. 3(a). We shall construct a dynamic observable $A$ by associating a value of $+1$ with any move into state 1, and $-1$ with any move into state 0. In this case we can consider the model to be a minimal representation of a one-dimensional active particle (see e.g. 29, 30), with 0 being a left-pointing, left-moving state, and 1 being a right-pointing, right-moving state. At each step the particle flips direction with probability $\alpha$ (and takes a step in the new direction), or else retains its current orientation and takes a step in the corresponding direction. $a = A/K$ is then the position of the particle after $K$ steps.

For $\alpha = 1/2$ we have the random walker (or unbiased coin-toss) of Section IIA. For $\alpha = 0$ the two states cannot interconvert (the Markov chain is reducible), and there exist two rate functions (which vanish for $a = 1$ or $a = -1$, and are infinite otherwise). For $0 < \alpha < 1/2$ we have an ‘active walker’ that moves persistently right or left between flips. In physical terms, fluctuations of $A/K$ for this model are related to fluctuations of magnetization $M/N$ in the one-dimensional Ising model in zero magnetic field, where $N$ is system size. Flips between runs $\cdots$LLLRRRLLL$\cdots$ occur independently with probability $\alpha$, much as domain walls between Ising spins $\cdots$+−−++−−$\cdots$ occur independently with probability $(1 + e^{\beta J})^{-1}$, where $J$ is the Ising coupling. For nonzero $\alpha$ (finite $J$) we expect domain lengths to be exponentially distributed with mean $\alpha^{-1}$ or $1 + e^{\beta J}$; for there to be no bias between L and R or between $−$ and $+$; and for the mean values of $A/K$ and $M/N$ in the limits of large $K$ and large $N$ to be zero. Thus the rate function $I(a)$ should be convex, symmetric, and have a (unique) minimum at zero.

We first calculate the rate function using the s-ensemble, following Ref. I1. The generator

$$W = \begin{pmatrix} 1 - \alpha & \alpha \\ \alpha & 1 - \alpha \end{pmatrix},$$  \hspace{1cm} (45)

becomes, in the s-ensemble,

$$W_s = \begin{pmatrix} (1 - \alpha)e^s & \alpha e^s \\ \alpha e^{-s} & (1 - \alpha)e^{-s} \end{pmatrix}.$$  \hspace{1cm} (46)
These results agree with those obtained using the Legendre transform using intervals of spacing 5\times10^{-5} and this agrees with the reference-model result. The red line was calculated by Legendre transform using coarser intervals of spacing 5\times10^{-3}, and this result has spurious flat portions. It is clear that this result is unphysical: the mean value of a of the original model is 0, and so the true rate function must have a unique minimum at zero. The smaller the value of \alpha, the finer the mesh required by the Legendre transform.

In Fig. 3(b) we show I(a) obtained from the reference-model method for three values of \alpha (black dotted lines). These results agree with those obtained using the s-ensemble.

For smaller values of \alpha the Legendre transform encounters numerical problems. In Fig. 3(c) we consider the case \alpha = 10^{-2} (the blue line shows the case \alpha = 1/2 for comparison). The rate function, calculated via the reference-model method, is shown in black. The same result can be obtained using the s-ensemble via numerical Legendre transform, but only if many closely-spaced values of s are used to calculate Eq. (8): the associated scaled cumulant-generating function is sharply kinked, and its numerical differentiation requires fine intervals of s. In Fig. 3(c) the green line was calculated by numerical Legendre transform using intervals of s of 5\times10^{-4}, and this agrees with the reference-model result. The red line was calculated by Legendre transform using coarser intervals of spacing 5\times10^{-2}, and the result has spurious flat portions. It is clear that this result is unphysical: the smaller the value of \alpha, the finer the mesh required by the Legendre transform.

From this we extract \lambda(s) in the usual way \cite{11}, and calculate I(a) via Legendre transform. Some results for three values of \alpha are shown in Fig. 3(b) (green lines).

We next calculate I(a) using the reference-model method. Its generator is

\[
W_{\text{ref}} = \begin{pmatrix}
(1-\alpha)e^s & (1-\alpha)e^{-s} & (1-\alpha)e^s + \alpha e^s \\
(1-\alpha)e^{-s} & (1-\alpha)e^s & (1-\alpha)e^{-s} + \alpha e^{-s} \\
(1-\alpha)e^{-s} + \alpha e^{-s} & (1-\alpha)e^s + \alpha e^s & (1-\alpha)e^{-s} + \alpha e^{-s}
\end{pmatrix}.
\] (47)

Calculation of I(a) can be done analytically, because the Jensen bound \cite{18} is exact for this model. This is so because each value of activity A is associated with a unique path weight: every move into state 1 (and only state 1) generates activity +1, and every move out of the state generates a piece \ln(R_{\text{ref}}(1)/R(1)) of the path weight. Similarly for state 0. By conservation of probability, every move into a state must be balanced by a move out of the state, and so the path weight takes a unique value for each value of A. Solving as before for the stationary measure \pi_{\text{ref}}(C), we evaluate the terms in Eq. (24) and write the rate function in the parametric form \(a_s(s), I(s)\); we get

\[
a_s(s) = \frac{(\alpha - 1) \sinh(2s)}{\alpha - (\alpha - 1) \cosh(2s)}
\] (48)

and

\[
I(s) = \frac{(\alpha - 1) (e^{2s} - 1) s + \alpha (e^{2s} - 1) \ln(2\alpha - 1) \sinh s + \cosh s)}{\alpha (e^{2s} - 1)^2 - e^{4s} - 1}
+ \frac{e^{3s}(-2\alpha \sinh s + \sinh s + \cosh s) \ln(-2\alpha \sinh s + \sinh s + \cosh s)}{\alpha (e^{2s} - 1)^2 - e^{4s} - 1}.
\] (49)
By contrast, the key quantities $a_s$ and $q_s$ of the reference-model method, panels (c) and (d) of Fig. 4, are not constrained to change sharply where original model’s rate function $I(a)$ is slowly varying. The reference-model rate functions shown in Fig. 3 are evaluated directly, by taking the combination $-s a_s - q_s$. Each point on the function can be evaluated independently, and the grid of $s$ can be as coarse or as fine as desired. The evaluation works when the rate function has very small curvature or is non-convex (see e.g., Fig. 4 of [16]). In the latter case the rate function cannot be recovered by Legendre transform [11].

In physical terms the results of this section relate to the probability distribution $P(a)$ of the position $a$ of an active particle, with rotation rate $\alpha$, in discrete space. This distribution adopts a large-deviation form $P(a) \sim e^{-K(l_a)}$ in the limit of a large number of steps $K$, with $I(a)$ given by Equations 45 and 49, shown in Fig. 3. For small $\alpha$, fluctuations of $a$ become large in the sense of having a large variance, but are to leading order Gaussian: $I(a) \approx a \alpha^2/2$, for $a, \alpha$ small. This distribution can be calculated using the $s$-ensemble, whose key quantities $\lambda(s)$ and $a(s)$ (see Fig. 4(a,b)) show a sharp jump when the rate function has small gradient. This is so for the geometrical reasons shown in Fig. 4 of [11]. The distribution $P(a)$ can also be calculated using the reference-model method, whose key quantities (see Fig. 4(c,d)) are not required to change sharply. Indeed, here the original model is similar to the 1D Ising model at nonzero temperature, for the reasons discussed above, and the set of reference models is similar to the 1D Ising model in a field $h \propto s$. Neither shows a sharp transition.

III. SAMPLING NONEQUILIBRIUM TRAJECTORIES OF A LATICE MODEL OF ACTIVE MATTER.

For models with larger state space, matrix diagonalization becomes impractical, and other methods are required to determine the $s$-ensemble. [5, 7, 8, 11, 13, 15]. The reference-model method is implemented in the same way, however, as we now illustrate using the lattice model of active particles introduced in Ref. 17.

Active particles propel themselves by consuming energy, and move by diffusion and by drift in a direction that fluctuates [31, 46]. Active particles form clusters and undergo phase separation even in the absence of inter-particle attractions [31, 57, 39, 44, 17, 18]. This phenomenon, called motility-induced phase separation, can be reproduced on the computer using simple model particles [30, 39, 47, 48] (motility-induced phase separation also occurs in the presence of velocity alignment [36, 49–51]). Associated with this phenomenon are various kinds of large fluctuations [33, 37, 52].

With an eye to studying such fluctuations in detail we introduced in Ref. 17 the following lattice model of active matter. Consider particles that move in isolation with the rates shown in Fig. 3(a). Particles possess both positions and orientations, and move on a two-dimensional square lattice. An isolated particle moves forward with rate $v_+$, backward with rate $v_-$, to either side with rate $v_0$, and rotates $\pi/2$ with rate $D_+$ or $D_-$ (here we set $D_+ = D_- \equiv D_{\text{rot}}$). An on-lattice particle of this nature moves in a manner similar to an off-lattice Brownian particle. As expected from the behavior of off-lattice particles [30, 39, 47, 48], on-lattice active particles undergo motility-induced phase separation: see Fig. 4(b) and Fig. 5(a). We used $N$ hard particles on a periodically-replicated lattice of size $L^2$ (for a packing fraction $\phi = N/L^2$), starting from disordered configurations, and propagated dynamics using a continuous-time Monte Carlo algorithm [27]. At each step of the simulation the system moves from the current configuration $C$ to a new configuration $C'$ with probability

$$p(C \rightarrow C') = \frac{W(C \rightarrow C')}{\sum_{C''} W(C \rightarrow C'')}$$

(50)

where the rates $W$ for each possible process are taken from Fig. 5. Typical trajectories of this model lead to phase separation at a density-dependent value of the Péctel number [17, 59].

To quantify the rare behavior of this model we chose a dynamic order parameter $\alpha$ motivated by studies of glasses. There, authors sometimes choose to count the number of events that occur in a particular time, with the mean time taken to leave configuration $C$ being $1/R(C)$. This choice allows the identification of phase transitions out of equilibrium [12, 53, 54]. Here we choose to measure...
the values of the escape rates $R(C)$ of states explored by a fixed number of configuration changes. With this choice the bias $\alpha(C \to C') = B(C')$, where $B(C') = R(C')$ is the escape rate from the state to which the model is moving, and the transition probabilities of the reference model are

$$p_{\text{ref}}(C \to C') = \frac{e^{-s(B(C') - B(C))}W(C \to C')}{\sum_{C'} e^{-s(B(C') - B(C))}W(C \to C')}.$$  

(51)

The factors $e^{sB(C)}$ in numerator and denominator have been introduced so that numbers appearing in the arguments of exponentials are not too large.

The coding overhead for the reference model is only slightly greater than that of the original model. One needs to keep track of the escape rate $R(C')$ of each configuration $C'$ accessible from the current configuration (rather than just the escape rate of the current configuration), but this can be done efficiently, in the course of the simulation, by noting which particles can touch the moving particle, before and after its move, one move into the future.

The order parameter against which dynamics is conditioned is $a = A/K = K^{-1} \sum_{k=0}^{K-1} B(C_{k+1})$, the mean relaxation rate of configurations comprising the trajectory, which we report in the form of a scaled ‘activity’, $\nu \equiv a/(\Sigma N)$. Here $N$ is the number of particles and $\Sigma \equiv v_+ + 2v_0 + v_- + 2D_{\text{rot}}$ is the sum of rates of processes of isolated particles. We used Eq. (22) to calculate the rate-function bound, which in this case reads

$$I_0(\nu) = -sK^{-1} \sum_{k=0}^{K-1} [B(C_{k+1}) - B(C_k)]$$

$$-K^{-1} \sum_{k=0}^{K-1} \ln \sum_{C'} e^{-s[B(C') - B(C)]}W(C_k \to C').$$

(52)

The first term on the right-hand side of (52) becomes negligible for large $K$. We evaluated this expression for trajectories of about $K = 10^8$ steps (the bound looks no different trajectories of $K \sim 10^7$ steps are used).

Results derived from these simulations are shown in Fig. 7 for packing fraction $\phi = 1/5$. Here the rate-function bound (which is not convex) shows that the typical activity of the model (where $I_0(\nu) = 0$) shifts from large $\nu$ to small $\nu$ as the drift rate $v_+$ of the active particles increases, because typical trajectories exhibit clustering (see Fig. 6) and (hence) reduced dynamism. In addition, trajectories show an active-to-less active crossover as a function of the bias parameter $s$. For model parameters far from the ‘typical’ nonequilibrium phase transition (which occurs near $v_+ \approx 7$ for $\phi = 1/5$) this crossover is seen in the ensemble of rare trajectories (for which $s = 0$), while for model parameters near the ‘typical’ transition the crossover occurs in the ensemble of typical trajectories, for which $s \approx 0$. This behavior resembles transitions observed in models of glasses [53] or proteins [55] using the $s$-ensemble.
FIG. 7. Activity \( \nu \) as a function of \( s \) (left graphs) and rate-function bound \( I_0(\nu) \) as a function of \( \nu \) (right graphs) from the reference-model method applied to the active lattice model. Packing fraction is \( \phi = 1/5 \) and drift rate \( v_+ = 2 \) (blue lines, top) or \( v_+ = 7 \) (red lines, bottom). Lattice size is \( 100^2 \). Trajectories prepared at \( s = 0 \) are typical of the original model, while trajectories prepared at \( s \neq 0 \) are rare in the original model; how rare can be understood from the right-hand graphs. For \( v_+ = 2 \) we are far from the ‘typical’ phase transition (see Fig. 10), and the crossover from active to inactive trajectories occurs for \( s \neq 0 \). For \( v_+ = 7 \), close to the ‘typical’ transition, the crossover occurs near \( s = 0 \). Shown right are typical and rare configurations of the model.

IV. CONCLUSIONS

We have described a method to sample the rare fluctuations of discrete-time Markov chains, and have applied it to the lattice-based model of active matter introduced in [17]. The method is inspired by the s-ensemble method but is distinct: we use a probability-conserving reference model to calculate directly the likelihood that a model displays a particular value of a dynamic order parameter \( a \). As a result, the method can be used to calculate non-convex rate functions, so giving information about trajectory coexistence near nonequilibrium phase transitions. The method is straightforward to implement for models for which continuous-time Monte Carlo dynamics can be carried out. Take the rates of the original model \( W(C \rightarrow C') \), change them to \( e^{-\alpha s(C \rightarrow C')} W(C \rightarrow C') \), run simulations and evaluate the quantities needed for equation [17] (repeat for a range of values of \( s \)). In general, as noted in Ref. [24], high-precision evaluation of integrals of the form appearing in Eq. [17] is demanding. For several models we have studied, however, calculation of the variance of \( q \) appears to give a good approximation of the rate function \( I(a) \) (because fluctuations of \( q \) are often approximately Gaussian), while calculation of the mean of \( q \) – which is straightforward – gives via Eq. [18] a physically meaningful, and potentially non-convex, rate-function bound. Thus the method provides a simple way of obtaining non-trivial information about a model and its phase transitions away from equilibrium, and so provides a complement to existing trajectory-sampling methods [5, 6, 10, 13, 53, 56, 59].

We end with a note about time. The reference-model method operates within the constant-event-number ensemble (the discrete-time Markov chain), because placing no restriction upon jump times makes the sampling procedure easier (see Appendix B of [16]). However, the procedure used to move through configuration space is the same as that used in a continuous-time Monte Carlo
scheme (except that no random number is drawn to compute the time of escape from a given state), and all temporal information can be recovered after the fact. The escape time $t_k$ from configuration $C_k$ is an exponentially distributed random variable with mean $1/R(C_k)$, and so the total time of a trajectory, $T = \sum_{k=0}^{K-1} t_k$, is a stochastic variable with a known distribution $P(T)$ [60]. This distribution reduces to the Erlang distribution if all $R(C_k)$ are identical, and to the hypoexponential distribution if all $R(C_k)$ are distinct [61]. Thus the complete distribution of trajectory times, for fixed number of events and specified activity $a$, can be recovered from the constant-event-number ensemble without additional simulation.

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