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A Novel A Priori Simulation Algorithm for Absorbing Receivers in Diffusion-Based Molecular Communication Systems

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Abstract—A novel a priori Monte Carlo (APMC) algorithm is proposed to accurately simulate the molecules absorbed at spherical receiver(s) with low computational complexity in diffusion-based molecular communication (MC) systems. It is demonstrated that the APMC algorithm achieves high simulation efficiency since by using this algorithm, the fraction of molecules absorbed for a relatively large time step length precisely matches the analytical result. Therefore, the APMC algorithm overcomes the shortcoming of the existing refined Monte Carlo (RMC) algorithm which enables accurate simulation for a relatively small time step length only. Moreover, for the RMC algorithm, an expression is proposed to quickly predict the simulation accuracy as a function of the time step length and system parameters, which facilitates the choice of simulation time step for a given system. Furthermore, a likelihood threshold is proposed for both the RMC and APMC algorithms to significantly save computational complexity while causing an extremely small loss in accuracy.

Index Terms—Diffusion-based molecular communication, absorbing receivers, molecular communication simulation, Monte Carlo method.

I. INTRODUCTION

Molecular communication (MC) has emerged as an underpinning paradigm of exchanging and conveying information among nano-devices in very small dimensions or specific environments, such as water, tunnels, and human bodies [2]. Unlike electromagnetic wave-enabled communication, MC delivers information based on chemical changes within the environment. In MC, transmitters first send out information-carrying molecules to propagate within the environment. Later, such molecules are captured by receivers to allow them to obtain the carried information. Biological examples of MC include chemotactic signaling, calcium signaling, and bacterial migration [2]. Thus, MC offers the advantages of low energy consumption and potential for biocompatibility in liquid and gaseous media. Notably, the development of MC is envisioned to support transformational nano-applications, e.g., intra-body health monitoring, target drug delivery, and food and water quality monitoring [2].

Diffusion-based MC has been acknowledged as a simple but commonly adopted MC system within the nano-communication research community [3]. In this system, information molecules propagate using kinetic energy only, which preserves a high energy efficiency. One of the major challenges in designing and analyzing a diffusion-based MC system is receiver modeling. The majority of existing MC studies have considered two types of receivers: passive receivers and active receivers [3, 4]. Passive receivers do not impose any impact on molecule propagation, while active receivers have some mechanism for molecules to react either within or at the surface of the receiver. The active receiver model is more general and is better representative of typical biological receivers. This motivates us to investigate the properties of absorbing receivers, which are a common ideal approximation for active receivers.

The notion of diffusion with absorption is a long-existing biological phenomenon that has been investigated in the literature, e.g., [5, 6]. For example, [5] investigated the diffusion into adsorbers, where the adsorbed molecules stop diffusion upon hitting the receiver, but later may desorb and resume diffusion. Specifically, [5] examined the diffusion into a spherical adsorber, an ellipsoidal adsorber, and disk-like adsorber(s) in an infinite medium and obtained the diffusion currents for such adsorbers by solving Fick’s equations. It is noted that some conclusions drawn for adsorbers can be applied to absorbers, due to the similarity between adsorption and absorption. Unlike [5], [6] discussed the absorption of molecules that can either diffuse through or react chemically with the receiver surface. Based on such studies, [7, 8] considered a diffusion-based MC system with a single perfectly absorbing receiver within an unbounded three-dimensional (3D) environment. To be specific, [7] provided the numerical results of the hitting rate of molecules at different times and the fraction of molecules absorbed for a given time, the analysis of which was performed in [9], while [8] presented a simulation framework. Recently, [10, 11] evaluated the impact of a receiver with reversible adsorption on the performance of MC systems, where a molecule can be released back to the environment at some time after being captured by the receiver.

Apart from the theoretical analysis of MC systems, e.g., [7, 10, 11], the simulation of MC systems also serves as an effective means for performance evaluation. Against this background, a number of simulation frameworks, e.g., N3Sim [12], NanoNS [13], BiNS2 [14], and AcCoRD [15], have been developed to examine the behavior of information particles in various MC environments. The development of such
frameworks is based on two common simulation approaches, the microscopic approach and the mesoscopic approach [16]. Considering a sub-region of the simulation environment, the microscopic approach treats the molecules within the region individually, while the mesoscopic approach treats them in aggregate as uniformly distributed throughout the region. A widely recognized example of the microscopic approach in MC is the particle-based simulation that uses Brownian motion to characterize particle propagation. A detailed comparison between the microscopic and mesoscopic approaches and further information on how they have been implemented in MC frameworks were shown in [15, 17]. In this paper, we focus on the particle-based microscopic approach due to its prevalence for the physical simulation of an MC environment.

It is worthwhile noting that the existing microscopic algorithms incur high computational complexity in the simulation of the fraction of molecules absorbed in MC systems with absorbing receivers. This high complexity leads to a long simulation run time since the existing algorithms require a very small time step length to accurately model the absorption. To tackle this issue, we develop a new particle-based microscopic algorithm in this work. This algorithm aims to reduce the computational complexity for the simulation of an irreversible perfectly absorbing receiver in a 3D MC system where every molecule that arrives at the receiver is absorbed.

In the simulation procedure of the particle-based microscopic algorithm for diffusion-based MC systems, molecules are moved by adding Gaussian random variables (RVs) to their \(x\), \(y\), and \(z\)-coordinates at the end of every simulation time step. This makes the coordinates discrete functions of time, although in reality the movements of molecules undergoing Brownian motion are continuous over both time and space. In practice, molecules may actually diffuse into an absorbing receiver between two time steps. To determine this absorption, some existing simulation algorithms simply compare the observed coordinates of molecules after diffusing with the coordinates of the receiver [8, 10, 12]. As a consequence, the molecules that hit a receiver between time steps cannot be considered as “absorbed” by using these simulation algorithms. We name the probability that a molecule crosses the receiver’s boundary between time steps as the intra-step absorption probability. Recently, [15] and [18] investigated the possibility of intra-step absorption. Specifically, [15] declared a molecule as “absorbed” if its straight-line trajectory within a time step crossed an absorbing surface. Alternatively, [18] approximated the intra-step absorption probability for spherical receiver boundaries using the equation for flat planar receiver boundaries given by [19, Eq. (10)]. This approximation was referred to as the refined Monte Carlo (RMC) algorithm.

Our paper focuses on improving the accuracy and reducing the computational cost of simulating absorbing receivers. We first run simulations using the RMC algorithm and measure the change in accuracy as the time step length increases. Then, we fit the results to an expression to estimate the accuracy given the diffusion coefficient \(D\), time step length \(\Delta t\), receiver’s radius \(r_r\), and transmitter-to-receiver distance \(r_0\). After recognizing the poor accuracy of the RMC algorithm when \(\Delta t\) increases, we propose a new method for simulations of absorbing receiver(s) with a large time step length, namely, the \textit{a priori} Monte Carlo (APMC) algorithm. The APMC algorithm uses the \textit{a priori} probability of a molecule being absorbed to decide whether it is absorbed in the current simulation time step. If the molecule is determined as “absorbed” using the \textit{a priori} probability, then we omit the diffusion step. We show that this algorithm achieves very high accuracy when the diffusion step length is large relative to the size of the receiver. Despite that the RMC algorithm performs accurately for small time step size, we observe from MATLAB that the simulation run time of the RMC algorithm is higher than that of the APMC algorithm for small time step size. Furthermore, we identify that a major contributor to the computational complexity of the RMC and APMC algorithms is the generation of uniform RVs when assessing intra-step absorptions. Thus, we propose a likelihood threshold for both algorithms to reduce the computational cost caused by the generation of excessive RVs.

Our contributions extend our preliminary work in [1] and are summarized as follows:

1) As in [1], we present a new algorithm, i.e., the APMC algorithm, for MC simulation with a relatively large time step length. The advantages of the APMC algorithm are:

   a) For the case of a single perfectly absorbing receiver, we show that by using the APMC algorithm, the fraction of molecules absorbed precisely matches the corresponding analytical result when \(\sqrt{D\Delta t}/r_r\) is relatively large.

   b) For the case of two perfectly absorbing receivers, we show that by using the APMC algorithm, the fraction of molecules absorbed approaches the asymptotic analytical value when time grows large.

   We note that the advantages of the APMC algorithm in both cases cannot be achieved by the existing algorithms.

2) We propose polynomial fitting expressions to predict the accuracy of the RMC algorithm introduced in [18] for the case of a single perfectly absorbing receiver. Aided by numerical results, we show that the third order polynomial fitting expression is the most accurate one for accuracy prediction. This allows us to use this expression to characterize the accuracy of the simulation without running it.

3) We investigate the computational complexity of both the RMC algorithm and the APMC algorithm. Specifically, we compare their MATLAB run times and explore the trade-off between simulation accuracy and computational complexity for both algorithms, based on which we propose a likelihood threshold for the absorption probability to save computation time. Molecule absorption for the RMC algorithm and the APMC algorithm is possible if and only if the calculated absorption likelihood in the simulation is higher than the likelihood threshold. Aided by numerical results, we show that applying a likelihood threshold can save as much as 20% of the total number of generated RVs with an extremely small loss in simulation accuracy.

Comparing to our preliminary work [1], which only presented the received signals of the simplistic Monte Carlo
Absorbing molecules diffuse in the environment according to Brownian motion. Molecules propagate in the environment according to the RMC algorithm, the APMC algorithm, and the SMC algorithm for the system with a single absorbing receiver, this paper additionally presents the received signals for multi-receiver configurations, proposes accuracy prediction expressions for the RMC algorithm, and investigates the computational complexity of the RMC and APMC algorithms. Furthermore, the majority of the numerical results and discussions shown in this paper are not included in [1].

The rest of the paper is organized as follows. The system model of interests is described in Section II. Existing simulation algorithms, our proposed APMC algorithm, and the likelihood threshold are presented in Section III. Numerical results and discussions are provided in Section IV. In Section V, we present our conclusions.

II. SYSTEM MODEL

We consider a diffusion-based MC system within a 3D space, as depicted in Fig. 1. In this system, a point transmitter (TX) is located at the origin of the space and a spherical perfectly absorbing receiver (RX) is centered at location \((r_0, 0, 0)\). We denote \(r_s\) as the RX’s radius and \(\Omega_r\) as the RX’s boundary. At the beginning of a transmission process, the TX instantaneously releases \(N\) molecules. We assume that the molecules are small enough to be considered as points. Once released, molecules diffuse in the environment according to Brownian motion until hitting the RX’s boundary. We denote \(N_{\text{hit}}(\Omega_r, t|r_0)\) as the number of molecules released from the TX at time \(t_0 = 0\) s and absorbed by the RX by time \(t\). As per [9, Eq. (3.116)], we express \(N_{\text{hit}}(\Omega_r, t|r_0)\) as

\[
N_{\text{hit}}(\Omega_r, t|r_0) = \frac{N r_t}{r_0} \operatorname{erfc} \left( \frac{r_0 - r_t}{\sqrt{4Dt}} \right),
\]

where \(D\) is the diffusion coefficient and \(\operatorname{erfc}(\cdot)\) is the complementary error function. \(D\) describes the proportionality constant between the flux due to molecular diffusion and the gradient in the concentration of molecules.

The majority of the existing simulation algorithms for absorbing RXs did not consider the possibility of intra-step molecule absorption [18]. This absorption is depicted in Fig. 2, which shows that the actual trajectory of a molecule may cross the RX’s boundary during one simulation time step, even if its initial position at the beginning of the time step and its final position at the end of the same time step are both outside the absorbing RX. If this crossing occurs, the molecule is absorbed by the RX in practice. When the possibility of this absorption is ignored, the number of absorbed molecules is underestimated, thus deteriorating the accuracy of simulation.

In this work, we refer to the probability that a molecule is absorbed during a simulation time step as the intra-step absorption probability. For MC systems, the discussion of this probability was limited. Two of the papers that considered the intra-step absorption probability are [15, 18]. In [15], the absorption of a molecule was determined by whether its straight-line trajectory has crossed the absorbing surface. In [18], the intra-step absorption probability of a perfectly absorbing RX with a spherical boundary was approximated as that of a perfectly absorbing RX with an infinite planar boundary, given by [19, Eq. (10)]

\[
\Pr_{\text{RMC}} = \exp \left( -\frac{l_i}{D\Delta t} \right),
\]

where \(l_i\) is the initial distance of a molecule from the absorbing boundary at the beginning of a time step, \(l_i\) is the final distance of a molecule from the absorbing boundary at the end of the same time step, and \(\Delta t\) is the time step length.

Apart from the aforementioned MC system which contains only one absorbing spherical RX, we also consider a system containing two identical absorbing spherical RXs. In this system, TX stays at the origin of the space and two absorbing RXs are located on opposite sides of and equidistant from the TX. Focusing on this two-RX system, [20] evaluated the fraction of molecules absorbed, which is defined as the ratio between the number of molecules absorbed at the RX and the total number of molecules released. As the time elapsed goes to infinity, the asymptotic fraction of molecules absorbed by each RX is given by [20]

\[
\Pr_{\text{RMC}} \to \infty = \sqrt{2} \left( \cosh \mu - \cos \eta \right) \times \sum_{n=0}^{\infty} e^{-\left(n+\frac{1}{2}\right)\mu_1} \frac{\sinh \left(n+\frac{1}{2}\right) (\mu - \mu_2)}{\sinh \left(n+\frac{1}{2}\right) (\mu_1 - \mu_2)} P_n (\cos (\eta)),
\]

where \((\mu, \eta, \phi)\) are the bispherical coordinates corresponding to \((0, 0, 0)\) in the natural coordinate system, \(\mu_1 = \cosh^{-1} (r_0/r_s)\), \(\mu_2 = -\cosh^{-1} (r_0/r_s)\), \(r_s\) is the RX’s radius, \(r_0\) is the distance from the center of one RX to the TX, and \(P_n\)
is the \( n \)th-degree Legendre polynomial. The two-RX system allows us to show that our proposed simulation algorithm can be applied to not only a single-RX system but a multi-RX system, which will be demonstrated in Section IV.

### III. Simulation Algorithms

In this section, we first describe the common structure of the existing simulation algorithms for absorbing RXs. Then we propose polynomial functions to predict the accuracy of the RMC algorithm. After that, we present our proposed APMC algorithm and determine the likelihood threshold to be applied to the APMC and RMC algorithms. The algorithms discussed in this paper are summarized in Table I.

### A. Existing Simulation Algorithms

By observing the existing simulation algorithms for microscopic molecule absorption (such as those in [8, 15, 18]), we identify that they follow a common structure. This structure is presented in Algorithm 1. For simulating molecules that follow Brownian motion, \( \Delta t \) needs to be carefully chosen such that the diffusion step length is relatively small compared to the distances that define reflective or absorbing boundaries [19]. We note that the ratio of the step length to the distance affects the performance of the simulation algorithms, as will be demonstrated in Section IV.

We clarify that each algorithm has its own criterion for determining whether or not a molecule is absorbed by the RX, as given in Line 9 of Algorithm 1. Such determination criteria for algorithms in [8, 15, 18] are summarized as follows:

- As per the determination criterion in [8], the molecules being observed inside the RX at the end of a time step are absorbed. This criterion is referred to by [18] as the SMC algorithm. We note that the performance of the SMC algorithm is inaccurate, unless the time step length is very small. This is because the SMC algorithm ignores the possibility of intra-step absorption and thus, underestimates the number of absorbed molecules.
- As per the determination criterion in [15], a molecule is absorbed if the line segment from its initial position to its final position crosses the RX’s boundary. However, we note that the line segment crossing the RX’s surface is neither sufficient nor necessary to correctly detect intra-step absorption. For example, in Fig. 2 the molecule absorption that actually occurs cannot be detected by the criterion in [15]. In another case, shown in Fig. 3, molecule absorption is determined by the criterion in [15] but does not actually occur. The accuracy of the algorithm in [15] is better than but still comparable to the SMC algorithm.
- As per the determination criterion in [18], referred to as the RMC algorithm and described in Algorithm 2, (2) is used to calculate the intra-step absorption probability of an RX with a perfectly absorbing spherical boundary. As

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**Algorithm 1** A common structure of simulation algorithms for a single absorbing RX

1: Determine the end time of simulation.
2: for all simulation time steps do
3: \[ \text{if } t = 0 \text{ then} \]
4: \[ \text{Add } N \text{ molecules to environment.} \]
5: \[ \text{end if} \]
6: \[ \text{Scan all not-yet-absorbed molecules, i.e., molecules which are not absorbed by the RX.} \]
7: \[ \text{for all not-yet-absorbed molecules do} \]
8: \[ \text{Propagate each molecule for one step according to Brownian motion.} \]
9: \[ \text{Determine if the molecule is absorbed.} \]
10: \[ \text{end for} \]
11: \[ \text{end for} \]
Algorithm 2 Molecule absorption determination in [18]

1: if The molecule’s distance to \((r_0, 0, 0)\) is smaller than \(r_r\), then
2: The molecule is absorbed.
3: else
4: Calculate the intra-step absorption probability, \(Pr_{\text{RMC}}\), using (2).
5: Generate a uniform RV \(u\).
6: if \(Pr_{\text{RMC}} \geq u\) then
7: The molecule is absorbed.
8: end if
9: end if

depicted in Fig. 4, in the RMC algorithm, \(l_i\) denotes the shortest distance between the molecule’s initial position and the RX’s boundary and \(l_f\) denotes the shortest distance between the molecule’s final position and the RX’s boundary. As shown in [18], the accuracy of the SMC algorithm is comparable to that of the RMC algorithm when \(\Delta t\) is relatively small. When \(\Delta t\) increases while other parameters remain the same, the accuracy of the RMC algorithm becomes measurably higher than that of the SMC algorithm. However, in addition to the results presented in [18], we run simulations of the SMC and RMC algorithms and observe that the simulated fraction of absorbed molecules of the RMC algorithm deviates from the analytical one given by (2) when the root mean square (RMS) of the diffusion step length, \(\sqrt{2D\Delta t}\), is relatively larger than \(r_r\). We will show the impact of \(\Delta t\) on the accuracy of the RMC algorithm in Section IV.

B. Accuracy Prediction of RMC Algorithm

In this subsection, we propose a rule-of-thumb expression to predict the accuracy of the RMC algorithm when the parameters of the MC system are within specified ranges. Specifically, our aim is to make a rough but fast prediction of the algorithm’s accuracy for given parameters, without resorting to simulations. To collect necessary data for accuracy prediction, we adopt the following procedure (the intermediate plots mentioned in the procedure are not presented in this paper due to page limit):

1) For the parameters listed in Table II, we run simulations using the RMC algorithm for one time step. We note that the fraction of molecules absorbed at every current time step is based on the results generated in the previous time step. In order to eliminate the impact of results from previous time steps, we choose to test for the first simulation time step only, which means that we calculate with only two samples, one at the very beginning of the transmission, and another at the end of the first time step. Simulation is repeated for \(N\) molecules and the result obtained from the simulation of each molecule is called a realization. For each set of \(D, \Delta t,\) and \(r_0\), we plot \(R^2\) versus \(r_r\), where the measure of accuracy, \(R^2\), is calculated according to

\[
R^2 = 1 - \frac{\sum_{i=1}^{2} (Pr_{\text{hit}}(i-1) - Pr_{\text{sim}}(i-1))^2}{\sum_{i=1}^{2} (Pr_{\text{sim}}(i-1) - Pr_{\text{sim}})^2},
\]

where

\[
Pr_{\text{hit}}(i-1) = \frac{N_{\text{hit}}(\Omega_{r_i}, (i-1)\Delta t(r_0))/N,}
\]

is the analytical fraction of absorbed molecules at time \((i-1)\Delta t\), as obtained from (1). We define \(Pr_{\text{sim}}(i-1)\) as the simulated fraction of absorbed molecules by the RMC algorithm at time \((i-1)\Delta t\), which is obtained by averaging the total number of absorbed molecules out of all realizations at \((i-1)\Delta t\) over \(N\), and refer to it as the \(i\)th simulated sample. Additionally, we define \(Pr_{\text{sim}}\) as the mean of all simulated samples.

2) We plot reference lines that indicate different values of \(R^2\) in the same figure.

3) We observe the values of \(r_0, r_r, D,\) and \(\Delta t\) that achieve the selected values of \(R^2\).

4) For each selected value of \(R^2\), we plot a 3D figure where the \(z\)-coordinate is the recorded value of \(r_r\), the \(x\)-coordinate is the recorded value of \(D\Delta t\), and the \(y\)-coordinate is the recorded value of \(r_0\).

By observing the obtained results, we find that the accuracy of the RMC algorithm decreases when \(D\Delta t\) increases, \(r_0\) increases or \(r_r\) decreases. To predict the accuracy, we define a dimensionless variable \(\kappa\) that is calculated from the parameters of the MC system. We assume linear dependency between the variables, i.e., \(\kappa\) and \(r_r\), for the sake of a quick and simple calculation. Using the curve fitting tool in MATLAB, we fit the 3D plots to equations of \(\kappa\). Here, we define \(\kappa\) as

\[
\kappa = r_r (r_0 D \Delta t)^{-\frac{1}{3}},
\]

where the exponent of \(-\frac{1}{3}\) is chosen such that the right hand side is dimensionless. We next fit \(R^2\) to a polynomial function of \(\kappa\). We test the first, second, and third order polynomial fits\(^1\), which are given by

\[
R^2 \approx \frac{101\kappa + 47}{100},
\]

\[
R^2 \approx \frac{-372\kappa^2 + 392\kappa - 3}{100},
\]

and

\[
R^2 \approx \frac{979\kappa^3 - 1523\kappa^2 + 813\kappa - 51}{100},
\]

respectively. Given that \(R^2\) is the accuracy of the RMC algorithm, which implies \(0 \leq R^2 \leq 1\), we impose \(0.0726 \leq \kappa \leq 0.612\) for (9). If \(\kappa > 0.612\), then the RMC algorithm is predicted to have asymptotically high accuracy. If \(\kappa < 0.0726\), then the RMC is predicted to have asymptotically low accuracy. We then arrive at

\[
R^2 \approx \begin{cases} 
0, & \text{if } \kappa < 0.0726, \\
\frac{979\kappa^3 - 1523\kappa^2 + 813\kappa - 51}{100}, & \text{if } 0.0726 \leq \kappa \leq 0.612, \\
1, & \text{otherwise}.
\end{cases}
\]

\(^1\)We do not test higher order fits in this paper because we seek an easy-to-compute expression.
We will demonstrate in Section IV that the third order polynomial fit is the best match among (7), (8), and (10). The accuracy of the three polynomial fits will be shown in Fig. 8 and the corresponding measured RMS error (RMSE) will be shown in Table III.

C. New A Priori Monte Carlo Algorithm

In this subsection, we propose a new simulation algorithm for approximating the fraction of molecules absorbed at a perfectly absorbing RX when $\sqrt{D}\Delta t/r$, is larger than that considered for the RMC algorithm in [18]. We refer to the newly proposed algorithm as the APMC algorithm.

The procedure of the APMC algorithm is to first calculate, before the $j$th molecule diffuses, the probability that this molecule will be absorbed in the current time step. This probability depends on the distance between this molecule and the RX, $d_j$, and the time step length, $\Delta t$. Specifically, the probability that this molecule will be absorbed in the current time step is calculated as

$$\Pr_{\text{APMC}} = \frac{r_j}{d_j} \text{erfc} \left( \frac{d_j - r_j}{\sqrt{4D\Delta t}} \right),$$

which is obtained by scaling (1) by $N$, replacing the total simulation time $t$ with $\Delta t$, and replacing $r_0$ with $d_j$. In (11), $\Pr_{\text{APMC}}$ denotes the fraction of absorbed molecules released from a location $d_j$ away from the RX at time $t = 0 \text{s}$ and absorbed by the RX by time $\Delta t$. We note that (11) is calculated repeatedly in every time step for each of the free molecules with the molecule’s current updated location when it diffuses. Then the molecule absorption is determined by generating a uniform RV $u$, where $0 \leq u \leq 1$, and comparing its value with the probability obtained by (11). A molecule is marked as “absorbed” if $u \leq \Pr_{\text{APMC}}$. After determining the molecules absorbed, each not-yet-absorbed molecule is propagated according to Brownian motion. If any propagated molecule is inside the RX’s boundary at the end of the current time step, we revert the diffusion of this molecule and let it propagate again, until this molecule diffuses to a location outside the RX. This is because if a molecule propagates to a location inside the RX, it contradicts the preconditioning that the molecule is not absorbed. The APMC algorithm is detailed in Algorithm 3.

D. Likelihood Threshold for Simulation Complexity Reduction

In this subsection, we propose a likelihood threshold to reduce the computational complexity for the RMC algorithm and the APMC algorithm. For both algorithms, the computational complexity consists of three parts and can be written as

$$c_{\text{sim}} = O \left( \frac{t_{\text{end}}}{\Delta t} (c_{\text{diffuse}} + c_{\text{absorb}} + c_{\text{locate}}) \right),$$

where $c_{\text{diffuse}}$ denotes the complexity of diffusing all molecules, $c_{\text{absorb}}$ denotes the complexity of determining the molecules to be absorbed and absorbing these molecules, and $c_{\text{locate}}$ denotes the complexity of determining whether a molecule is inside the spherical RX. We observe from MATLAB run time profiles that the time for generating uniform RVs is one of the major factors of the run time costs. We also observe that an increase in the number of generated RVs corresponds to an increase in the total run time cost for both algorithms. Therefore, we use the number of generated RVs to characterize the computational complexity of both algorithms.

There are two types of RVs for the APMC algorithm and the RMC algorithm, namely, the uniform RVs used for testing the molecule absorption and the Gaussian RVs used for propagating molecules according to Brownian motion. Thus, we denote $N_u$ and $N_g$ as the number of uniform RVs and the number of Gaussian RVs, respectively. For each determination of molecule absorption, as shown in Line 6 in Algorithm 2 and Line 10 in Algorithm 3, we add 1 to $N_u$. For each molecule propagation, as shown in Line 8 in Algorithm 1 and Lines 15 and 18 in Algorithm 3, we add 3 to $N_g$ since there are three Gaussian RVs added to the $x$-, $y$-, and $z$-coordinates of a molecule. Here, we adopt the commonly used polar method to convert the number of generated Gaussian RVs to an equivalent number of generated uniform RVs [21].

### Algorithm 3 The APMC algorithm for molecule absorption

1. Determine the end time of simulation.
2. for all simulation time steps do
3. \hspace{1em} if $t = 0$ then
4. \hspace{2em} Release $N$ molecules into environment.
5. \hspace{1em} end if
6. \hspace{1em} Scan all not-yet-absorbed molecules.
7. \hspace{1em} for all not-yet-absorbed molecules do
8. \hspace{2em} Calculate the distance between the $j$th molecule to $(r_0, 0, 0)$, denote by $d_j$.
9. \hspace{2em} Calculate the absorbed probability $\Pr_{\text{APMC}}$ for each not-yet-absorbed molecule using (11) with $r_j$, $d_j$, $D$, and $\Delta t$.
10. \hspace{2em} if $\Pr_{\text{APMC}} \geq u$ then
11. \hspace{3em} The molecule is absorbed.
12. \hspace{1em} end if
13. \hspace{1em} end for
14. \hspace{1em} for all not-yet-absorbed molecules do
15. \hspace{2em} Propagate the molecule for one step.
16. \hspace{2em} while the molecule’s distance to $(r_0, 0, 0)$ $\leq r$ do
17. \hspace{3em} Revert the movement of this molecule to the location before propagation.
18. \hspace{3em} Propagate this molecule again.
19. \hspace{2em} end while
20. \hspace{1em} end for
21. end for
According to [21], the generation of a pair of Gaussian RVs requires 4/π pairs of uniform RVs. Thus, we use 4/π as a conversion factor and then the equivalent number of totally generated uniform RVs, which characterizes the computational complexity of simulation, is given by \( N_{t} = (4/\pi) N_{u} \).

Based on the aforementioned computational complexity characterization, we now propose a likelihood threshold, \( \xi \), to reduce the computational complexity. With this likelihood threshold, molecule absorption for the RMC algorithm is possible if and only if \( Pr_{\text{RMC}} \geq \xi \), while molecule absorption for the APMC algorithm is possible if and only if \( Pr_{\text{APMC}} \geq \xi \). This can significantly reduce \( N_{u} \) in simulation. Although the accuracy of the RMC and APMC algorithms may slightly decrease when \( \xi \) applies, an appropriate value of \( \xi \) can provide a good trade-off between simulation accuracy and computational complexity. Such trade-off of the RMC and APMC algorithms will be illustrated in Section IV.

**IV. NUMERICAL RESULTS**

In this section, we compare the fraction of molecules absorbed determined by the SMC algorithm, the RMC algorithm, and the APMC algorithm, with the aid of particle-based simulations, to show the benefits of our APMC algorithm. We also investigate the accuracy of the prediction expression obtained from empirical simulations in Section III-B by examining the performance of the RMC algorithm. Furthermore, we compare simulation run times of the RMC and APMC algorithms and explore the trade-off between simulation accuracy and computational complexity of algorithms. The algorithms are implemented in MATLAB.

Throughout this section, we denote \( M \) as the number of time-varying samples, and hence \( M - 1 \) as the number of time steps. Unless otherwise stated, we adopt the diffusion coefficient of \( D = 10^{-9} \text{m}^2/\text{s} \), the TX-RX distance of \( r_0 = 50 \mu \text{m} \), and the number of molecules released of \( N = 10^6 \).

**A. A Single Absorbing Receiver**

In this subsection, we focus on the MC system with a single absorbing RX. We first examine the impact of \( r_r \) and \( \Delta t \) on the fraction of molecules absorbed produced by the three algorithms in Fig. 5 and Fig. 6, respectively.

Fig. 5 plots the simulated fraction of molecules absorbed versus time \( t \) for \( M = 100, \Delta t = 0.1 \text{s} \), and \( r_r = 20 \mu \text{m} \) or \( 0.5 \mu \text{m} \). In this figure, the analytical result obtained using (1) is also plotted for examining the accuracy of the algorithms. From this figure, we observe that our APMC algorithm achieves a higher accuracy when \( r_r \) decreases while \( \Delta t, D \), and \( r_0 \) remain unchanged. Specifically, Fig. 5(a) shows that the RMC algorithm matches the fraction of molecules absorbed as predicted by the analytical result when \( \sqrt{D\Delta t}/r_r \) is small, which meets our expectation. Indeed, the performance of the RMC algorithm depends on the value of \( \sqrt{D\Delta t}/r_r \). When \( \sqrt{D\Delta t}/r_r \) approaches 0 and \( r_0 \) is slightly larger than \( r_r \), the surface area of the RX can be approximated by an infinite plane. Therefore, the probability of a molecule entering the RX between time steps is comparable to the probability of a molecule crossing a flat planar boundary between time steps.

We also observe from Fig. 5(a) that when \( \sqrt{D\Delta t}/r_r \) is small, the SMC algorithm and the APMC algorithm underestimate and overestimates the fraction of molecules absorbed, respectively. The overestimation of the APMC algorithm is due to the fact that the \textit{a priori} probability, given in (11), assumes that the molecule is released by the TX from its current location and calculates the probability that the molecule will be absorbed in the next time step, while actually the molecule is released by the TX from the origin of the 3D system at the beginning of the simulation and arrives at its current location at the current time step. If (11) is directly used, the assumption that the molecule has to first arrive at its current location is ignored. Therefore, the APMC algorithm may overestimate molecule absorption. Unlike Fig. 5(a), Fig. 5(b) shows that the APMC algorithm matches the fraction of molecules absorbed predicted by the analytical result while the other two algorithms do not. Particularly, the fractions of molecules absorbed produced by the RMC and SMC algorithms are very far away from the analytical result when \( t > 0 \text{s} \). Indeed, when \( \sqrt{D\Delta t}/r_r \) is large, the spherical RX’s boundary cannot be approximated by a flat planar boundary. Therefore, the RMC algorithm overestimates the fraction of molecules absorbed.

![Fig. 5: Comparison of the fraction of molecules absorbed results produced by the SMC algorithm, the RMC algorithm, and the APMC algorithm versus time for different \( r_r \) with \( M = 100 \) and \( \Delta t = 0.1 \text{s} \).](image1)

![Fig. 6: Comparison of the fraction of molecules absorbed results produced by the SMC algorithm, the RMC algorithm, and the APMC algorithm versus time for different \( \Delta t \) with \( M = 10 \) and \( r_r = 10 \mu \text{m} \).](image2)
Fig. 7: Distribution of the number of newly-absorbed molecules during each time step for the RMC and APMC algorithms versus time for \( \Delta t = 0.5 \) s, \( M = 10 \), and \( r_t = 10 \) \( \mu \)m. Simulations are repeated \( 10^3 \) times and \( N = 10^3 \) molecules are released each time. The spectrum bars represent observation probabilities.

TABLE III: RMSE Measurements for Polynomial Fits in Fig. 8

<table>
<thead>
<tr>
<th></th>
<th>1st order fit</th>
<th>2nd order fit</th>
<th>3rd order fit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fig. 8(a)</td>
<td>0.0638</td>
<td>0.1487</td>
<td>0.0440</td>
</tr>
<tr>
<td>Fig. 8(b)</td>
<td>0.0255</td>
<td>0.4378</td>
<td>0.0218</td>
</tr>
</tbody>
</table>

Fig. 8 plots the predicted accuracy given by (7), (8), and (10) as well as the measured accuracy of the APMC and RMC algorithms versus \( D\Delta t \) for \( r_0 = 40 \) \( \mu \)m, \( M = 2 \), and \( r_t = 15 \) \( \mu \)m or 20 \( \mu \)m. We observe from Fig. 8 that when \( D\Delta t \) increases, for a fixed \( r_t \), the accuracy of the RMC algorithm is at first higher than that of the APMC algorithm, but soon becomes much lower than that of the APMC algorithm. We also observe that the accuracy of the APMC stays close to 1 for most of the range of \( D\Delta t \) considered. This demonstrates the accuracy and robustness of our APMC algorithm when \( \sqrt{D\Delta t/r_t} \) is large.

To quantitatively assess the accuracy of the first, second, and third order polynomial fits in Fig. 8, we calculate the RMSE measurements of these fits via

\[
\text{RMSE} = \sqrt{\frac{\sum_{i=1}^{N} (Pr_{\text{hit}}(i-1) - Pr_{\text{sim}}(i-1))^2}{2}},
\]

where \( Pr_{\text{hit}}(i-1) \), \( Pr_{\text{sim}}(i-1) \), and \( i \) are defined the same as in (4). The RMSE measurements for both subfigures are given in Table III. Based on both subfigures and Table III, we find that the third order polynomial fit for \( R^2 \) is more accurate than the first and second order polynomial fits for \( r_t = 15 \) \( \mu \)m and 20 \( \mu \)m, since the third order polynomial fit is, on average, closer to the measured accuracy than the first and second order polynomial fits, as shown in both subfigures, and achieves the lowest RMSE fitting measurement, as shown in Table III. Therefore, in Figs. 9 and 10 we only consider the third order polynomial fit for the RMC algorithm.

Fig. 9 plots the measured accuracy and the predicted accuracy of the APMC algorithm versus \( D\Delta t \) for \( r_t = 20 \) \( \mu \)m, 25 \( \mu \)m, and 30 \( \mu \)m. For the measured accuracy at a given \( D\Delta t \), we run simulations for different combinations of \( D \) and \( \Delta t \) which achieve the same \( D\Delta t \), e.g., the combination of \( D = 2 \times 10^{-9} \) m\(^3\)/s and \( \Delta t = 0.5 \) s or the combination of \( D = 1 \times 10^{-9} \) m\(^3\)/s and \( \Delta t = 1 \) s. The simulated result
for each combination is displayed by a point (i.e., circle) in a scatter plot. We observe that the points overlap with each other for a given $D\Delta t$, which demonstrates that the performance of the RMC algorithm depends on $D\Delta t$, but not $D$ or $\Delta t$ separately. We further find that the average absolute value of the difference between the measured accuracy and the predicted accuracy is 5.66% for $r_r = 20\, \mu m$, 3.27% for $r_r = 25\, \mu m$, and 1.45% for $r_r = 30\, \mu m$, all of which are lower than 6%.

Fig. 10 plots the measured and predicted accuracy of the RMC algorithm together with the measured accuracy of the APMC algorithm versus $r_r$ for large time step lengths $\Delta t = 1\, s$ and $3\, s$. As we observe from this figure, the accuracy of the APMC algorithm stays close to 1 for most of the range of $r_r$ considered. This demonstrates that for large time step lengths, the APMC algorithm preserves a very high accuracy, which agrees with the discussion on Fig. 6(b), and the variance in accuracy is small. In addition, we observe that the measured accuracy of the RMC algorithm agrees well with the third order polynomial fit given in (10). This is in accordance with the observation made from Fig. 9 that the third order polynomial fit well approximates the measured accuracy when $D\Delta t \leq 3000$.

Finally, we compare the computational complexity of the APMC algorithm with that of the RMC algorithm. Fig. 11 plots the average run time per realization of MATLAB simulation for the APMC and RMC algorithms for $r_r = 20\, \mu m$, $N = 10^4$, $D\Delta t = 0.01\, s$, $0.05\, s$, $0.1\, s$, $0.5\, s$, $1\, s$, $2\, s$, and $5\, s$. We clarify that each run time is averaged over at least 20 realizations on an Intel i7 desktop PC. We first observe from the figure that the averaged run time per realization decreases when $\Delta t$ becomes higher. This is due to the fact that when $\Delta t$ increases, the number of time steps in the simulation decreases. This leads to a decrease in the number of propagation and absorption operations, which affects the computational complexity. Second, we observe that the APMC algorithm requires a slightly shorter run time to simulate a system transmission process than the RMC algorithm. This is due to the fact that the APMC algorithm overestimates the number of absorbed molecules under the parameters in Fig. 11. An overestimation means that more molecules are absorbed and removed permanently from the environment than expected, which leads to fewer molecules to be propagated and tracked by the system and thus reducing the computational complexity of the simulation. When the time step length increases to $\Delta t = 5\, s$, $\sqrt{D\Delta t/r_r}$ becomes large and the RMC algorithm leads to a more severe overestimation than the APMC algorithm. Therefore, the RMC algorithm requires a shorter time to simulate than the APMC algorithm.
2.0.850.95Accuracy, \( R \)

0.8
0.9
0.92
0.94
0.96\( \xi \)

0.9
0.92
0.94
0.96\( r \)

Fig. 13: Number of RVs generated and measured accuracy of the RMC algorithm versus \( \xi \) for \( r_0 = 50 \mu m, M = 10, N = 10^6, D = 10^{-9} m^2/s, \Delta t = 0.1 s, \) and \( r_v = 10 \mu m \) or \( r_v = 20 \mu m \).

0.8
0.9
0.92
0.94
0.96\( \xi \)

0.9
0.92
0.94
0.96\( r \)

Fig. 14: Comparison of the fraction of molecules absorbed for a system with two perfectly absorbing RXs and a system with four perfectly absorbing RXs.

In this subsection, we focus on the MC system with multiple absorbing RXs. We run simulations for the system with two absorbing RXs, placed symmetrically as described in the last paragraph of Section II, and the system with four absorbing RX, where the TX is a point located at the origin, and the RXs are located at \( (r_0, 0, 0), (-r_0, 0, 0), (0, r_0, 0), \) and \( (0, -r_0, 0) \) in a Cartesian coordinate system with the radius as \( r_r \). For both systems, we examine the fraction of molecules absorbed as produced by the RMC and APMC algorithms. Due to the symmetry of the systems, the probabilities that a molecule is absorbed by each of the RXs are identical.

In addition, we observe from Fig. 12 and Fig. 13 that when \( r_v \) increases from 10 \( \mu m \) to 20 \( \mu m \), the change in measured accuracy becomes less severe when applying the same \( \xi \). This is because a larger RX leads to a higher a priori absorption probability in the APMC algorithm and a higher intra-step absorption probability in the RMC algorithm. Therefore, the impact of likelihood threshold on the number of absorptions becomes smaller for a larger RX. We clarify that Figs. 12 and 13 reveal the impact of the likelihood threshold on the number of required RVs and the measured accuracy. With the aid of these figures and similar results, a suitable likelihood threshold can be identified to achieve an acceptable loss of accuracy for simulating other system parameters.

### B. Multiple Absorbing Receivers

In this subsection, we focus on the MC system with multiple absorbing RXs. We run simulations for the system with two absorbing RXs, placed symmetrically as described in the last paragraph of Section II, and the system with four absorbing RXs. We run simulations for the system with two absorbing RXs.

We also observe that when \( r_v \) increases from 10 \( \mu m \) to 20 \( \mu m \), the change in measured accuracy becomes less severe when applying the same \( \xi \). This is because a larger RX leads to a higher a priori absorption probability in the APMC algorithm and a higher intra-step absorption probability in the RMC algorithm. Therefore, the impact of the likelihood threshold on the number of absorptions becomes smaller for a larger RX. We clarify that Figs. 12 and 13 reveal the impact of the likelihood threshold on the number of required RVs and the measured accuracy. With the aid of these figures and similar results, a suitable likelihood threshold can be identified to achieve an acceptable loss of accuracy for simulating other system parameters.
$r_c$ increases to 40 μm, the fractions of molecules absorbed as produced by both algorithms approach the same value, which is similar to the system with two absorbing RXs.

V. Conclusion

In this paper, we proposed a new algorithm named the \textit{a priori} Monte Carlo (APMC) algorithm to use the \textit{a priori} approach for simulating the fraction of molecules being absorbed by spherical receiver(s). Based on numerical results, we demonstrated that the APMC algorithm outperforms an existing algorithm, the refined Monte Carlo (RMC) algorithm for large $\sqrt{\Delta t/r_f}$. Thus, the APMC algorithm is suitable to be used in simulations for both the single absorbing receiver case and the two absorbing receivers case when $\sqrt{\Delta t/r_f}$ is large. Moreover, for the single absorbing receiver case, we proposed an expression to predict the simulation accuracy of the RMC algorithm. We further investigated the MATLAB run time of the APMC and RMC algorithms and applied likelihood thresholds to reduce the computational complexity of both algorithms. This investigation revealed that the APMC algorithm significantly reduces the computational complexity of simulating absorbing receivers in molecular communication systems without compromising accuracy. Thus, it is worthwhile to extend the APMC algorithm to other reactive surfaces where the \textit{a priori} probability of surface interaction is known.

References


