Parameter Estimation in a Noisy 1D Environment via Two Absorbing Receivers

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Abstract—This paper investigates the estimation of different parameters, e.g., propagation distance and flow velocity, by utilizing two fully-absorbing receivers (RXs) in a one-dimensional (1D) environment. The time-varying number of absorbed molecules at each RX and the number of absorbed molecules in a time interval as time approaches infinity are derived. Noisy molecules in this environment, that are released by sources in addition to the transmitter, are also considered. A novel estimation method, namely difference estimation (DE), is proposed to eliminate the effect of noise by using the difference of received signals at the two RXs. For DE, the Cramer-Rao lower bound (CRLB) on the variance of estimation is derived. Independent maximum likelihood estimation is also considered at each RX as a benchmark to show the performance advantage of DE. Aided by particle-based simulation, the derived analytical results are verified. Furthermore, numerical results show that DE attains the CRLB and is less sensitive to the change of noise than independent estimation at each RX.

Index Terms—Molecular communication, parameter estimation, noise, absorbing receivers

I. INTRODUCTION

Nanoscale communication is critical for nanomachines to collaboratively execute complex tasks in biomedical applications, e.g., in vivo drug delivery and surgery. Molecular communication (MC) is envisaged as one of the most promising methods for nano-scale communication, where molecules act as information carriers. MC possesses important characteristics, such as low energy consumption and potential for biocompatibility, which makes it more suitable for in vivo applications than other nanoscale communication methods, e.g., electromagnetic methods [1].

Parameter monitoring in the in vivo environment is a potential application of MC, which is pivotal for healthcare, e.g., anomaly detection and targeted drug delivery [2]. For example, estimating the emission rate of biomarkers by a tumor could help to diagnose the stage of tumor development. Moreover, distance estimation could help to accurately locate a tumor and deliver drugs to the site. Furthermore, estimating the flow velocity and degradation rate of emitted biomarkers could help to evaluate the blood velocity and pH near the tumor, respectively, since the degradation rate varies with pH as discussed in [3]. Motivated by these applications, some previous studies have provided valuable insights into parameter monitoring, e.g., [4]–[7]. [4] and [5] considered estimation by a single transparent receiver (RX) and non-transparent RX, respectively, where a transparent RX is one that does not interact with molecules. [6] considered estimation in a one-dimensional (1D) pipe with two absorbing boundaries, where the estimation is based on particle concentration and flux in the pipe. [7] considered estimation by multiple non-transparent RXs based on data-fitting expressions of the channel impulse response (CIR), where the CIR is the expected number of absorbed molecules at each RX [8]. Although these studies stand on their own merits, they did not analytically derive CIRs at multiple non-transparent RXs and apply such CIRs to estimate parameters in the MC environment.

In this paper, we for the first time perform parameter estimation by using analytically derived expressions of CIR at two dependent non-transparent RXs. We consider a 1D environment, where a transmitter (TX) continuously releases molecules into the environment. The 1D environment is worthy of investigation since it can approximate many practical biological channels such as capillaries, blood vessels, active transportation channels, and communication on bio-chips [9], [10]. Moreover, we emphasize that some recent publications have also investigated a 1D model, e.g., [6], [11]. Furthermore, the intended TX might not be the only source of molecules. We also consider other sources that secrete the same type of molecules as the intended TX, which interfere with the parameter estimation at RXs. In this paper, we refer to the molecules not emitted from the intended TX as noisy molecules. To reduce the impact of noisy molecules on the estimation, we propose a novel estimation method, namely, difference estimation (DE). In this method, the difference between the number of absorbed molecules at two fully-absorbing RXs is used to estimate parameters, where fully-absorbing RXs are those that absorb molecules once they hit the RX surface. In addition, we consider independent maximum-likelihood (ML) estimation at each RX as a benchmark to show performance superiority of DE.

Our major contributions are summarized as follows. We first derive the time-varying expression for the expected number of absorbed molecules at each RX when the TX emits molecules continuously. We next derive the asymptotic expression for the number of absorbed molecules within a time interval at each RX after sufficient time has passed. We further derive the Cramer-Rao lower bound (CRLB) on the variance of different parameters. These parameters include distances, emission rate,
degradation rate of molecules, and flow velocity. Aided by particle-based simulation, we verify results and show that DE attains the CRLB. Importantly, our numerical results show that the impact of noise on the performance of DE is much less than that on independent estimation of each RX, which demonstrates the superiority of DE in reducing the impact of noise.

II. SYSTEM MODEL

In this paper, we consider a 1D unbounded environment, where a TX continuously releases molecules into the environment with a constant emission rate $\mu$ per second, as depicted in Fig. 1. We assume that the TX starts emission at $t = 0$ s. To estimate environmental parameters, two fully-absorbing RXs, i.e., RX$_1$ and RX$_2$, are placed at different sides of the TX with the distances $d_1$ and $d_2$ to the TX, respectively. Moreover, we consider a steady uniform flow $\bar{v}$ in the direction from RX$_1$ towards RX$_2$. Once molecules are released from the TX, they diffuse randomly with a constant diffusion coefficient $D$ and the constant flow. Additionally, molecules can degrade from the type $A$ into a new type of molecule $\phi$, i.e., $A \xrightarrow{k} \phi$, where $k$ is the degradation rate constant. The two RXs absorb type $A$ molecules as soon as they hit RX$_1$ and RX$_2$. In addition, we consider that the noisy molecules released by sources in addition to the intended TX may interfere with the estimation of environmental parameters. We emphasize that this additive noise is distinct from the randomness in the number of $A$ molecules absorbed by each RX due to diffusion, which may also interfere with the estimation. We note that one potential application of the system model in Fig. 1 is estimating parameters near a tumor in a blood-vessel environment.

In this system, estimation is based on the received signals of two RXs at time $t$, which is the number of absorbed molecules within the time interval $[t - \delta, t]$, where $\delta$ is the length of the time interval. We assume that each RX first performs $S$ observations separately, and then they perform estimation jointly based on the difference between observations at two RXs. Therefore, we name this estimation method as DE. As discussed in [12], the computation of difference can be conducted with the aid of another device. For example, the two RXs may transmit their observations to a fusion center that has easy access to computational resources. In addition, independent estimation at each RX is considered as a benchmark for DE.

In Fig. 2, we plot the expected received signals at two RXs in a noisy environment by the simulation. From this figure, we observe that the expected received signals become constant when $t > 4$ s. We refer to the stage where the expected received signal is constant as the asymptotic stage. We note that it is a short time for two RXs to reach the asymptotic stage after the TX starts releasing molecules. Therefore, we assume that RXs start to make observations at the asymptotic stage as in [13]. One advantage for using asymptotic observations is that they do not depend on time, such that time synchronization is not required between devices, which improves the practicality of the proposed estimation method.

III. DERIVATION OF CHANNEL IMPULSE RESPONSE

In this section, we first derive the expected number of absorbed molecules at each RX until time $t$ due to continuous emission by the TX. We then derive the closed-form expression for the number of absorbed molecules at each RX within the time interval $[t - \delta, t]$ as $t \to \infty$, which lays the foundation for parameter estimation in Section IV. We denote $N_j(t)$, $j = \{1, 2\}$, as the expected number of absorbed molecules at RX$_j$ by time $t$. Based on our previous work in [11], we derive $N_j(t)$ in the following theorem:

**Theorem 1**: The expected number of molecules absorbed at RX$_j$ by time $t$ due to the continuous emission of molecules at the TX is given by

$$N_j(t) = \mu \exp \left( \frac{d_j |\bar{v}|}{2D} \right) \sum_{i=0}^{\infty} \left[ R_i(2(i+1)d + d_j, t, 2) 
- R_i(2(i+2)d - d_j, t, 2) - R_i(2id + d_j, t, 0) 
+ R_i(2(i+1)d - d_j, t, 0) \right],$$

where $d = d_1 + d_2$ is the distance between the two RXs and $R_i(x, t, a)$ is given by

$$R_i(x, t, a) = \frac{\theta \kappa}{2} \left( (\alpha \omega(t) - \hat{\alpha} \nu(t)) - \frac{i + 1}{2} (\alpha \omega(t) + \hat{\alpha} \nu(t)) \right)$$

$$- \frac{\theta}{\sqrt{D \nu^2 + 4kD}} \left( \hat{\beta}(t) - \alpha \beta(t) \right) + (i + 1)t. \quad (2)$$

In (2), $\theta = d(i + 1)(i + a)$, $\kappa = \sqrt{\frac{\nu^2}{4D} + \frac{x^2}{4D}}$, $\alpha = \exp(x \kappa)$, $\hat{\alpha} = \exp(-x \kappa)$, $\beta(t) = \text{erfc} \left( \frac{x}{\sqrt{4Dt}} + \sqrt{k + \frac{\nu^2}{4D}} t \right)$, $\hat{\beta}(t) = \text{erfc} \left( \frac{x}{\sqrt{4Dt}} - \sqrt{k + \frac{\nu^2}{4D}} t \right)$, $\omega(t) = \int_0^t \beta(u) du$, and $\nu(t) = \int_0^t \hat{\beta}(u) du$. $\omega(t)$ and $\nu(t)$ are calculated numerically.
as a continuous random process, the time interval between releasing two successive molecules is a random variable (RV). Thus, the release time of each molecule is different, which means that the received signal follows a Poisson binomial distribution since each molecule has a different probability of being absorbed by the time the observation is made. As the Poisson binomial distribution is cumbersome to work with, we approximate it by a Poisson distribution. The approximation is accurate when the number of trials (emitted molecules) is large and the success probability $P_j(t)$ is small [15]. We assume that the number of absorbed noisy molecules within the time interval at each RX are identically and independently distributed Poisson RVs with constant mean $\xi$ as in [16]. This assumption is reasonable since sources in addition to the intended TX can also be regarded as TXs. Due to the additivity of two Poisson RVs, we model each observation of the received signal as a Poisson RV with mean $\tilde{N}_j + \xi$. As $g_{2,s}$ and $g_{1,s}$ follow a Poisson distribution, the difference between observations, i.e., $\tilde{g}_s$, follows a Skellam distribution with mean $N_2 - N_1$. According to the PMF of Skellam distribution in [17], the joint conditional PMF is given by

$$p(\tilde{g}|\varepsilon) = \prod_{s=1}^{S} \exp\left(-\left(\tilde{N}_1 + \tilde{N}_2\right)\right) \frac{\tilde{g}_s}{\tilde{N}_1^{\tilde{g}_s} \tilde{N}_2^{N_s-\tilde{g}_s}} I_{\tilde{g}_s} \left(2\sqrt{\tilde{N}_1\tilde{N}_2}\right),$$

(5)

where $\tilde{N}_j = \tilde{N}_j + \xi$ and $I_{\tilde{g}_s}(2\sqrt{\tilde{N}_1\tilde{N}_2})$ is the modified Bessel function of the first kind [18]. According to [14] Ch. 3], the CRLB of the variance of an unbiased estimator, denoted by $\operatorname{var}(\hat{\varepsilon})$, is $\operatorname{var}(\hat{\varepsilon}) \geq \frac{\mathbb{E}[\hat{\varepsilon}^2]}{\mathbb{E}[\hat{\varepsilon}]}$, where $\hat{\varepsilon}$ is the estimated value of $\varepsilon$, $\mathbb{E}[\hat{\varepsilon}^2]$ represents the expectation, and $L(\varepsilon)$ is the Fisher information that is given by [14] eq. (3.6)

$$L(\varepsilon) = -\mathbb{E}\left[\frac{\partial^2 \ln p(\tilde{g}|\varepsilon)}{\partial \varepsilon^2}\right],$$

(6)

where $\mathbb{E}[-]$ is taken with respect to $\tilde{g}_s$. We derive $L(\varepsilon)$ in the following theorem:

**Theorem 3:** The Fisher information of DE for unknown parameter $\varepsilon$ is derived as

$$L(\varepsilon) = \sum_{s=1}^{S} \frac{3\tilde{N}_s - \tilde{N}_1}{4N_2^2}\gamma_2(\varepsilon) + \frac{3\tilde{N}_s - \tilde{N}_2}{4N_1^2}\gamma_2(\varepsilon) - \frac{\tilde{N}_1 + \tilde{N}_2}{2N_1N_2}\gamma_1(\varepsilon)\gamma_2(\varepsilon),$$

(7)

where $\gamma_j(\varepsilon) = \frac{\partial \hat{N}_j}{\partial \varepsilon}$, $\zeta_1$ and $\zeta_2$ are found numerically as explained in the Appendix C. In Table. I, we list $\gamma_j(\varepsilon)$ for the parameters $d_2$, $v$, $\mu$ and $k$.

**Proof:** Please see Appendix C.
TABLE I
γ_j(ε) FOR EACH ESTIMATED PARAMETER

| γ_j(d_j) | μ ∂ε exp (−1) j d_j | \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) − \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) | \exp (−d_j \kappa) \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) − \exp (−d_j \kappa) \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) |
|--------|------------------------|---------------------------------|---------------------------------|
| γ_j(μ) | \frac{N_2 \kappa}{\mu} | \exp (−1) \kappa \kappa \exp (−1) \kappa \kappa | \exp (−1) \kappa \kappa \exp (−1) \kappa \kappa |
| γ_j(v) | \frac{\mu \partial ε \exp (−1) j d_j}{(1−\exp (−2d_j))} | \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) − \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) | \exp (−d_j \kappa) \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) − \exp (−d_j \kappa) \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) |
| γ_j(k) | \frac{\mu \partial ε \exp (−1) j d_j}{(1−\exp (−2d_j))} | \exp (−(2d_j + ε) \kappa) \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) − \exp (−(2d_j - ε) \kappa) \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) | \exp (−d_j \kappa) \left( \frac{d_j + 2 ε}{\sqrt{2} \text{v} \text{D}} \right) − \exp (−d_j \kappa) \left( \frac{d_j - ε}{\sqrt{2} \text{v} \text{D}} \right) |

B. Method of Moments
For DE, we apply the method of moments to estimate the unknown parameter, which is finding \( \hat{\xi} \) that makes 1) the analytical expression for the mean of \( \hat{g}_s \), and 2) the mean of \( \hat{g}_s \) calculated by observed data, equal [14 Ch. 9]. As each item in the vector \( \mathbf{g} \) is independent and follows the same distribution with the same mean, the mean of \( \hat{g}_s \) from observed data can be calculated by \( \frac{1}{S} \sum_{s=1}^{S} \hat{g}_s \). Since \( \hat{g}_s \) follows a Skellam distribution with mean \( \bar{N}_2 - \bar{N}_1 \), we have the following estimation criteria:

\[
\left( \bar{N}_2 - \bar{N}_1 \right) \bigg|_{\hat{\xi} = \xi} = \frac{1}{S} \sum_{s=1}^{S} \hat{g}_s, \tag{8}\]

where \( \xi \) is not included such that noise from sources in addition to the intended TX does not impact the estimation.

C. Benchmark
To highlight the superiority of DE in eliminating the impact of noise, we consider independent estimation at each RX as a benchmark, where each RX performs ML estimation based on their \( S \) observations in \( \mathbf{g}_j \). We recall that each observation is approximated by a Poisson distribution with mean \( \bar{N}_j \). Therefore, the joint conditional PMF of \( S \) observations at RX\(_j\), denoted by \( p(\mathbf{g}_j | \xi) \), is given by \( p(\mathbf{g}_j | \xi) = \prod_{s=1}^{S} \bar{N}_j^{g_{js}} \exp (-\bar{N}_j) / g_{js}! \). The principle of ML estimation is finding \( \hat{\xi} \) that maximizes \( p(\mathbf{g}_j | \xi) \). By solving \( \frac{\partial \ln p(\mathbf{g}_j | \xi)}{\partial \xi} = 0 \), we obtain the equivalent expression \( \bar{N}_j \bigg|_{\xi = \hat{\xi}} = \frac{1}{S} \sum_{s=1}^{S} g_{js} \). Since \( \xi \) is unknown in \( \bar{N}_j \), one approach for independent estimation is ignoring the noise. By doing so, we obtain the following estimation criteria:

\[
\bar{N}_j \bigg|_{\hat{\xi} = \hat{\xi}_{\text{ML}}} = \frac{1}{S} \sum_{s=1}^{S} g_{js}, \tag{9}\]

We highlight that (9) indicates that the ML estimation at RX\(_j\) is to find \( \hat{\xi}_{\text{ML}} \) that makes \( \bar{N}_j \bigg|_{\xi = \hat{\xi}_{\text{ML}}} \) and the mean of \( S \) observations in \( \mathbf{g}_j \) equal.

V. NUMERICAL RESULTS
In this section, we present numerical results to validate our theoretical analysis and provide insightful discussions. Particle-based simulation is used to simulate the random propagation of molecules. The simulation time step is \( t_{\text{sim}} = 0.001 \) s and all results are averaged over 2000 realizations. Throughout this section, we set \( d = 40 \) \( \mu \text{m} \), \( D = 7.94 \) \( \mu \text{m}^2/\text{s} \), \( \mu = 1000 \) \( \text{s}^{-1} \), \( k = 0.8 \) \( \text{s}^{-1} \), and \( \delta = 0.5 \) \( \text{s} \). In Fig. 3 we plot the received signals \( \Delta N_j(t) + \xi \) and \( \bar{N}_j \) versus time \( t \) at RX\(_j\) for three RXs versus time \( t \), where different flow velocities and distances between the TX and two RXs are considered to investigate the impact of the flow and distance on the received signals. We first observe that the simulation matches well with \( \Delta N_j(t) + \xi \) and \( \bar{N}_j \), which demonstrates the correctness of (11) and (4). Furthermore, comparing parameter sets 1) and 2), we observe that the received signals increase at RX\(_2\) and decrease at RX\(_1\) with an increase in the flow velocity. This is because we assume that the direction of the flow is from RX\(_1\) towards RX\(_2\). In addition, comparing parameter sets 2) and 3), we observe that the received signals increase with a decrease in the distance between the TX and RX.

In Fig. 4 we plot the normalized MSE for DE and two RXs and the CRLB for DE versus the number of observations, where \( \xi = 0 \) and \( \xi = 10 \) in this environment are considered. In this figure, we set \( d_1 = d_2 = 40 \) \( \mu \text{m} \) and \( v = 6 \) \( \mu \text{m/s} \). Both MSE and CRLB are normalized over \( \xi^2 \). First, we observe that the normalized MSE attains the CRLB for DE, which indicates that the method of moments applied in DE achieves the minimum variance. Second, we observe that the normalized MSEs at two RXs increase with the increase in noise. Third, in
Fig. 4. The normalized MSE versus the number of observations $S$, where $\xi = 0$ and $\xi = 10$ are considered. (a) Estimation of $d_2$. (b) Estimation of $v$. (c) Estimation of $\mu$. (d) Estimation of $k$.

Fig. 4(a) and Fig. 4(b), we observe that the normalized MSES at two RXs are close to each other when $\xi = 0$ and the MSE of DE is always lower than that at either of the RXs. This is because $d_2$ and $v$ have no impact on the number of molecules in this environment, which means that the performance of the estimation is not related to the number of molecules absorbed at each RX. As DE combines observations of both RXs, DE achieves a better performance. When noise increases, the impact on the estimation of DE is much less than that at either RX, which indicates the superiority of DE in eliminating the noise in estimation. Fourth, in Fig. 4(c) and Fig. 4(d), we observe a large gap between normalized MSES of either RX and the performance of RX2 is better than DE when $\xi = 0$. This is because $\mu$ and $k$ influence the number of molecules in the environment, which means that the number of absorbed molecules at either RX influences the performance of estimation. When $\xi = 0$, RX2 performs estimation based on a larger number of molecules than DE and RX1. Therefore, RX2 achieves the best performance. It is worth noting that the performance of RX2 becomes worse than DE for $S > 2$ when $\xi = 10$ and the increase of MSE in DE is much less than the increase at RX2, which still proves that DE is superior than independent estimation in eliminating the impact of noise. This superiority becomes obvious with the increase in $\xi$ and $S$.

In Fig. 5 we plot the minimum number of observations required for the MSE in DE to be less than the MSE at RX2 versus the mean of noise $\xi$. (a) Estimation of $\mu$. (b) Estimation of $k$.

Fig. 5. The minimum number of observations $S$ required for MSE in DE to be less than the MSE at RX2 versus the mean of noise $\xi$. (a) Estimation of $\mu$. (b) Estimation of $k$.

we first observe that the minimum number of observations required decreases with an increase in the noise mean, which indicates that the increase of MSE in DE is much slower than that at RX2, demonstrating that DE is less sensitive to the increase in noise than RX2. After $\xi > 13$ and $\xi > 17$ for the estimation of $\mu$ and $k$, respectively, the MSE in DE is always less than the MSE at RX2 since the minimum $S$ drops to one. Second, we observe that the minimum number of observations decreases with a decrease in $\mu$ or increase in $k$. This is because a decrease in $\mu$ or increase in $k$ results in a decrease in the received signal at RX2, which makes RX2 more susceptible to an increase in noise than DE.

**VI. Conclusion**

In this paper, we investigated parameter estimation by the cooperation of two RXs in a noisy 1D environment. We derived the analytical expressions for the CIR at both RXs. Moreover, we considered DE and derived the CRLB. To show the advantage of DE, independent estimation at each RX was also investigated. Our numerical results verified our analytical results. Numerical results also showed that DE is less sensitive to an increase in noise than independent estimation. For the estimation of $d_2$ and $v$, DE always achieves better performance than independent estimation. For the estimation of $\mu$ and $k$, DE becomes better than independent estimation with an increase in noise or in the number of observations. Future work includes considering mobile RXs and extending the estimation to a three-dimensional (3D) environment.

**Appendix A**

**Proof of Theorem 1**

We express $N_j(t)$ as $N_j(t) = \mu \int_0^t \hat{P}_j(u)du$, where $\hat{P}_j(t)$ is the fraction of absorbed molecules by time $t$ for an impulsive emission at the TX. To obtain $N_j(t)$, we first calculate $\hat{P}_j(t)$.

We denote $f_v(d,t,k,v)$ and $f_d(d,t,k)$ as the hitting rate with and without flow, respectively, when only one RX exists. According to [20], $f_v(d,t,k,v)$ is given by

$$f_v(d,t,k,v) = \exp\left(\frac{d|\vec{v}|}{2D}\right) \frac{d}{\sqrt{4\pi Dt}} \exp\left(-\frac{d^2}{4Dt} - \frac{v^2}{4D} + k\right)t\right),$$

where $d$ is the distance between the TX and RX. In (10), $|\vec{v}| = v$ if the direction of the flow is from the TX toward RX and $|\vec{v}| = -v$ for the opposite direction.
By setting \( v = 0 \) in (10), we obtain \( f(d, t, k) \) as
\[
   f(d, t, k) = \frac{d}{\sqrt{4\pi Dt^3}} \exp \left( -\frac{d^2}{4Dt} - kt \right). \tag{11}
\]

Based on (10) and (11), we obtain
\[
f_v(d, t, k, v) = \exp \left( \frac{dv}{2D} \right) f \left( \frac{d}{t}, t, k + \frac{v^2}{4D} \right). \tag{12}
\]

We denote \( F_v(d, t, k, v) \) and \( F(d, t, k) \) as the fraction of absorbed molecules by time \( t \) with and without flow, respectively, when only one RX exists, which are obtained by integrating \( f_v(d, t, k, v) \) and \( f(d, t, k) \) over time \( t \). Therefore, they have the same relationship as (12), which is
\[
   F_v(d, t, k, v) = \exp \left( \frac{dv}{2D} \right) F \left( \frac{d}{t}, t, k + \frac{v^2}{4D} \right). \tag{13}
\]

Similar to the method in [11 Sec. III], we can express \( \tilde{P}_1(t) \) and \( \tilde{P}_2(t) \) as (11) eqs. (12), (13)
\[
\hat{P}_1(t) = F_v(d_1, t, k, v) - \hat{P}_1(t) \ast f_v(d_1 + d_2, t, k, v), \tag{14}
\]
\[
\hat{P}_2(t) = F_v(d_1, t, k, v) - \hat{P}_1(t) \ast f_v(d_1 + d_2, t, k, v), \tag{15}
\]
where \( \ast \) stands for convolution. Substituting (12) and (13) into (15) and (14) and performing the Laplace transform for (15) and (14), we obtain
\[
   \tilde{P}_2(s) = \exp \left( \frac{v^2 s}{2D} \right) P_2 \left( s, \frac{v^2}{4D} + k \right), \tag{16}
\]
where \( \tilde{P}_2(s) \) is the Laplace transform of \( \hat{P}_2(t) \) and \( P_2(s, k) \) is the Laplace transform of \( P_2(t, k) \) that is given by (11) eq. (8). Performing the inverse Laplace transform of (16), we obtain \( \tilde{P}_2(t) = \exp \left( \frac{d_2 v}{2D} \right) P_2(t, \frac{v^2}{4D} + k) \). Substituting \( \tilde{P}_2(t) \) into \( N_2(t) = \mu \int_0^t P_2(u)du, \) we obtain \( N_2(t) \). \( N_1(t) \) can be obtained by exchanging \( d_1 \) and \( d_2 \) and replacing \( v \) with \(-v \) therein for \( N_2(t) \). Based on the expressions for \( N_1(t) \) and \( N_2(t) \), a unified formula can be written as (11).

**APPENDIX B**

**PROOF OF THEOREM 2**

According to the final value theorem, if \( \hat{P}_2(t) \) has a finite limit as \( t \to \infty \), we have
\[
   \lim_{t \to \infty} \hat{P}_2(t) = \lim_{s \to 0} s \tilde{P}(s). \tag{17}
\]
Substituting (16) into (17), we obtain \( \hat{P}_2(t) |_{t \to \infty} \) as
\[
   \hat{P}_2(t) |_{t \to \infty} = \exp \left( \frac{d_2 v}{2D} \right) \lim_{s \to 0} s \tilde{P}(s, \frac{v^2}{4D} + k) = \exp \left( \frac{d_2 v}{2D} \right) P_{2,asy} \left( k + \frac{v^2}{4D} \right), \tag{18}
\]
where \( P_{2,asy}(k) \) is given by (11) eq. (11)\(^2\) According to \( N_2(t) = \mu \int_0^t P_2(u)du, \) absorbed molecules at RX2 within the time interval \([t - \delta, t] \) at the asymptotic stage is
\[
   \tilde{N}_2 \approx \mu \Delta t \hat{P}_2(t) |_{t \to \infty}. \tag{19}
\]

\(^2\) Please note that \( k = 0 \) s\(^{-1} \) is not considered in this paper.

Substituting (18) into (19), we obtain \( \tilde{N}_2 \). \( N_1 \) can be obtained by exchanging \( d_1 \) and \( d_2 \) and replacing \( v \) with \(-v \) therein for \( N_2 \). Based on equations of \( N_1 \) and \( N_2 \), a unified formula can be written as (11).

**APPENDIX C**

**PROOF OF THEOREM 3**

For the CRLB to exist, the regularity condition [14] must be satisfied, which is \( \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] = 0 \). Substituting (5) into \( \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] \), we obtain
\[
   \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] = \sum_{s=1}^S -\gamma_1(\varepsilon) - \gamma_2(\varepsilon) + \frac{\mathbb{E}[\tilde{g}_4]}{2} \left( \frac{\gamma_2(\varepsilon)}{N_2} - \frac{\gamma_1(\varepsilon)}{N_1} \right) + \frac{\mathbb{E}[\tilde{g}_s]}{2} \left( \frac{\sqrt{N_2}}{N_1} \sqrt{\gamma_1(\varepsilon)} + \frac{\sqrt{N_1}}{N_2} \sqrt{\gamma_2(\varepsilon)} \right). \tag{20}
\]

Based on the definition of the expectation, \( \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] \) is calculated as
\[
   \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] = \sum_{s=1}^S \gamma_1(\varepsilon) \gamma_2(\varepsilon) + \frac{\mathbb{E}[\tilde{g}_4]}{2} \left( \frac{\gamma_2(\varepsilon)}{N_2} - \frac{\gamma_1(\varepsilon)}{N_1} \right) + \frac{\mathbb{E}[\tilde{g}_s]}{2} \left( \frac{\sqrt{N_2}}{N_1} \sqrt{\gamma_1(\varepsilon)} + \frac{\sqrt{N_1}}{N_2} \sqrt{\gamma_2(\varepsilon)} \right). \tag{21}
\]

where \( \eta \) is the summation of PMF of Skellam distribution. Therefore, \( \eta = 1 \). Substituting (21) and \( \mathbb{E}[\tilde{g}_s] = \tilde{N}_2 - \tilde{N}_1 \) into (20), we obtain \( \mathbb{E} \left[ \frac{\partial \ln p(\xi | \varepsilon)}{\partial \varepsilon} \right] = 0 \).

We then calculate \( L(\varepsilon) \). Substituting (5) into (6), we obtain
\[
   L(\varepsilon) = \sum_{s=1}^S -3 \tilde{N}_2 - \tilde{N}_1 \gamma_2(\varepsilon) + 3 \tilde{N}_1 - \tilde{N}_2 \gamma_1(\varepsilon) - \frac{\tilde{N}_1 + \tilde{N}_2}{2N_2 N_1} \gamma_1(\varepsilon) \gamma_2(\varepsilon) - \frac{\gamma_1(\varepsilon)}{N_1} \frac{\gamma_2(\varepsilon)}{N_2} + \frac{\gamma_1^2(\varepsilon)}{4N_1^2} + \frac{\gamma_2^2(\varepsilon)}{4N_2^2} + \frac{\gamma_1^2(\varepsilon)}{4N_1^2} \tag{22}
\]
Similar to the method in [21], we calculate $E \left[ \frac{\hat{g}_s I_{g_s-1}}{I_{g_s}} \right] = \frac{N_s}{N_1} \left( \frac{N_s}{N_1} - 1 \right)$, $E \left[ \hat{g}_s^2 \right] = N_2 - N_1$, and $E \left[ \frac{I_{g_s-2} \left( 2 \sqrt{N_1 N_2} \right)}{I_{g_s} \left( 2 \sqrt{N_1 N_2} \right)} \right] = \frac{N_s}{N_2}$.

In $L(\vartheta)$, $\vartheta$ represents the value of $E \left[ \frac{I_{g_s-2} \left( 2 \sqrt{N_1 N_2} \right)}{I_{g_s} \left( 2 \sqrt{N_1 N_2} \right)} \right]$ that is also calculated based on the definition of the expectation. For calculating $\vartheta$, we do not need the summation from $\hat{g}_s = -\infty$ to $\hat{g}_s = \infty$, and we focus on the range of $\hat{g}_s$ that makes the PMF of the Skellam distribution larger than a threshold that we choose to be $10^{-3}$ in our paper, i.e., $\exp \left( -\left( N_1 + N_2 \right) \right) \frac{N_s}{N_1} I_{g_s} \left( 2 \sqrt{N_1 N_2} \right) \geq 10^{-3}$. The validation of this threshold is performed in our numerical tests. After solving this expression, we have $\zeta_1 \leq \hat{g}_s \leq \zeta_2$. Substituting $E \left[ \frac{\hat{g}_s I_{g_s-1}}{I_{g_s}} \right]$, $E \left[ \hat{g}_s^2 \right]$, $E \left[ \frac{I_{g_s-2} \left( 2 \sqrt{N_1 N_2} \right)}{I_{g_s} \left( 2 \sqrt{N_1 N_2} \right)} \right]$, and $\vartheta$ into (22), we obtain (7).

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