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Effective inter-symbol interference mitigation with a limited amount of enzymes in molecular communications

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Abstract

In molecular communication via diffusion (MCvD), the inter-symbol interference (ISI) is a well-known severe problem that deteriorates both data-rate and link reliability. ISI mainly occurs because of the slow and highly random propagation of the messenger molecules, which causes the emitted molecules from the previous symbols to interfere with molecules from the current symbol. An effective way to mitigate the ISI is using enzymes to degrade undesired molecules. Prior work on ISI mitigation by enzymes has assumed an infinite amount of enzymes randomly distributed around the molecular channel. Taking a different approach, this paper assumes an MCvD channel with a limited amount of enzymes. The main question this paper addresses is how to deploy these enzymes in an effective structure so that ISI mitigation is maximized. To find an effective MCvD channel environment, this study considers optimization of the shape of the transmitter node, the deployment location and structure, the size of the enzyme deployed area, and the half-lives of the enzymes. It also analyzes the dependence of the optimum size of the enzyme area on the distance and half-life. Lastly, the paper yields interesting results regarding the actual shape of the enzyme region itself.

1 | INTRODUCTION

As the field of nanotechnology is increasingly gaining significance in areas such as bioscience or environmental engineering, researchers have produced diverse developments in nanoscale devices. For a possible means for nano-communication, molecular communication via diffusion (MCvD) emerged as a prime candidate technology.^{1–3} In the case of the dominantly used radio frequency (RF) communication, nano-range is difficult to implement because of the severe path-loss. ⁴ The drawback of the MCvD, however, is the high level of randomness in signal propagation that creates problematic nonlinear noise in macroscale applications. ⁵ Moreover, the heavy tail nature of the received signal in MCvD causes inter-symbol interference (ISI), which is detrimental to the capacity of an MCvD channel because ISI can increase the error-rate or decrease the data-rate. With a pre-decided symbol duration, ISI can be defined as 1 or more symbols from previous symbol periods interfering with the current symbol and causing noise at the receiver node.^{6–9}

Molecular communication via diffusion uses messenger molecules instead of electromagnetic waves as the transmitting signal between 2 nodes, Tx and Rx, for communication. This is the general model, and details of the system can be diversified by characteristics such as the shape of the Tx and Rx or distance between the nodes. For a molecular-concentration-based MCvD system,^{2,10,11} the Tx either emits molecules or does nothing at each pre-decided symbol period, depending on the intended message. Hence, the shape of the received signal at the Rx is very significant in analyzing the system capacity. ISI occurs when the signal intended for the previous symbol does not propagate fast enough directly to the receiver. Hence, 1 solution for ISI is to increase the symbol duration so that the system can wait until each of the messenger molecules reaches the Rx within its symbol period. This is done with the trade-off of data-rate. An improved method may be symbol interval optimization.¹² Another approach to combat ISI is to use decision feedback mechanisms in the amplitude modulation method.^{7,13} A more reasonable alternative, however, that does

Half-life of

the Enzyme

FIGURE 1 Steps of analyzing effective ways to deploy a limited amount of enzymes in MCvD for ISI mitigation. ISI, inter-symbol interference; MCvD, molecular communication via diffusion

Size of the

Enzyme Area

Half-life of

the Enzyme

not elongate the symbol period is using enzymes to destroy the ISI molecules. This has the trade-off against signal power because enzymes also decompose the molecules that make up the current signal, but the loss of power can easily be compensated by lowering the decoding threshold.

Several studies have proposed different ideas regarding the implementation of enzymes to mitigate.^{14–17} Authors Kuran. Yilmaz, and Tugcu¹⁴ proposed using "destroyer molecules" to decrease the mean and variance of the hitting time distribution. Here, the researchers deployed an unlimited amount of destroyer molecules inside a cylindrical tunnel structure-a direct and restricted path between the point Tx and the sphere Rx. Authors Noel, Cheung, and Schober¹⁵ also proposed using enzymes to mitigate ISI in a 3-dimensional MCvD channel with a non-absorbing receiver. An infinite quantity of enzymes were assumed to be spread throughout the channel, and favorable performance in ISI mitigation was shown by decreased bit-error-rate. Authors Heren et al¹⁶ presented an analytical function for the hitting probability of an MCvD channel with an infinite amount of enzymes deployed everywhere. All these studies have taken different approaches to using an infinite amount of enzymes for ISI mitigation. In resource perspective, enzymes could be used more efficiently in a limited amount. Indeed, assuming the deployment of an infinite amount of enzymes may not be practical in real implementation of MCvD with enzymes. In the study of Yilmaz et al,¹⁷ a fixed amount of enzyme scenario was considered to mitigate ISI for only a few cases of system environments, which presents a confined analysis of implementing limited enzymes.

This paper presents an analysis of effectively using a limited amount of enzymes in different system structures. If an unlimited number of enzymes are available, then there is no question of *where* to deploy them, as optimal ISI mitigation would occur when enzymes are deployed everywhere within an appropriate concentration. In a limited enzymes situation, however, a critical factor would be to deploy them in an effective location and structure. After verifying that using enzymes leads to a lower ISI than using no enzymes at all, this study compares different shapes of Tx (sphere and point). It then considers the deployment location-enzymes randomly deployed "everywhere"* versus specific locations. Afterwards, to find the optimum case for ISI mitigation, the study compares the results from specific areas (i.e., structures) "around Rx" and "around Tx." Lastly, the specific system parameters, the size and shape of the enzyme area and the half-lives of the enzymes, are taken into account to see which scenario mitigates ISI the most. Figure 1 summarizes the main aspects of the limited enzyme deployment issue analyzed in this research.

Size of the

Enzyme Area

The paper is organized as follows. Section 2 gives both quantitative and qualitative descriptions of the MCvD channel and enzyme dynamics. Section 3 expands the MCvD concept to the limited enzymatic MCvD channel specific to our paper including topology, geometry, and scenarios of how the limited enzymes are implemented. Section 4 elaborates on the simulation system used in this paper and Section 5 gives a specific analysis of the results. Section 6 concludes the paper.

2 | SYSTEM MODELING

2.1 Molecular communication via diffusion

In a general MCvD system, the transmitter node emits messenger molecules that freely diffuse by Brownian motion^{18–21} toward the receiver. Once the molecular signal is received by the receiver, the signal is accordingly decoded by the system's modulation scheme. Modulation can be done in different ways depending on properties such as the concentration, type, and time of release of the messenger molecules.^{10,11} Because path-loss for MCvD is proportional to d^{-3} being lower than that of RF that is d^{-2} , molecular communication has less path-loss distortion when used in nano-environments.^{4,22} The problem of MCvD is the long propagation time proportional to d^2 , which is square to that of RF.⁴ This means that

^{*}Note that this scheme of deploying a limited amount of enzymes "everywhere" is not exactly the same with the case of deploying an infinite amount of enzymes. In the case of the limited enzymes scenario, having enzymes everywhere yields to 0 concentration of enzymes asymptotically. Therefore, instead of exactly deploying the enzymes everywhere we consider a sphere with a big radius for the enzyme deployment area to make it comparable with the other limited enzyme cases.



FIGURE 2 A molecular communication via diffusion system with a limited amount of enzymes deployed around Rx for spheres Rx and Tx

molecules diffuse so slowly that they exceed their symbol period and interfere with the next symbol period's molecules, creating ISI. Figure 2 shows a diagram of the MCvD system with a limited amount of enzymes deployed around the Rx.

In analyzing an MCvD system, the important factor is how the molecular signal is perceived at the receiver.²³ The peak, tail, and duration of the received signal are directly related to the system's decoding scheme, error-rate, and data-rate. In a 3-dimensional MCvD system with a point Tx and an absorbing sphere Rx, the hitting probability of sent molecules to the receiver is

$$h(t) = \frac{r_r}{d + r_r} \frac{d}{\sqrt{4\pi Dt^3}} e^{-\frac{d^2}{4Dt}},$$
 (1)

where r_r , d, and D are the receiver radius, the shortest distance between the Rx and Tx, and the diffusion coefficient, respectively.²² The equation gives a general understanding of the messenger molecules' behavior inside the MCvD channel without enzymes.

2.2 | Enzyme dynamics

Enzymes catalyze and speed up reactions by decomposing certain substrates into different types of molecules or eliminating them entirely. The enzyme chemical reaction is defined by

$$E + S \xrightarrow{k_1} ES \xrightarrow{k_p} E + P, \qquad (2)$$

where *E*, *S*, *ES*, *P*, and k_n are the enzyme, substrate, enzyme-substrate compound, product, and rate of reactions, respectively. By applying the law of mass action ²⁴ to Equation 2, we get

$$\frac{d[S]}{dt} = -k_1[E][S] + k_{-1}[ES]$$

$$\frac{d[E]}{dt} = -k_1[E][S] + k_{-1}[ES] + k_p[ES]$$
(3)
$$\frac{d[ES]}{dt} = k_1[E][S] - k_{-1}[ES] - k_p[ES],$$

where $[\cdot]$ corresponds to the concentration operator. In this paper, a specific case of enzymatic reaction is considered under the following assumptions:

- $k_p \rightarrow \infty$ and $k_{-1} \rightarrow 0$, $\therefore S \rightarrow P$
- $[ES] \rightarrow 0$, $\therefore [ES] = d[ES]/dt = 0$.

These assumptions imply that we use the uni-molecular reaction for degradation, a very fast enzymatic reaction that can be realized by selecting the appropriate pairs of enzymes and messenger molecules. In the uni-molecular degradation we do not consider a 1-by-1 enzyme-to-molecule reaction but use the probability of degradation for each of the molecule accordingly.

Applying the assumptions to Equation 3, we get

$$\frac{d[S]}{dt} = -k_1[S][E]$$

$$\frac{d[ES]}{dt} = \frac{d[E]}{dt} = 0.$$
(4)

By solving Equation 4, the concentration of messenger molecules (substrate) at time *t*, namely, C(t), with the initial substrate concentration C_0 , is derived as an exponential decay function,

$$C(t) = C_0 e^{-\lambda t}.$$
(5)

 λ is the degradation factor of C(t) expressed as

$$\lambda = [S][E] = \frac{\ln 2}{\Lambda_{1/2}}.$$
(6)

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 $\Lambda_{1/2}$ corresponds to the half-life of the enzyme, which has a core role in controlling a constant amount of enzymes among different deployment scenarios. This is elaborated in detail in later sections of this paper.

If the hitting probability at time *t* to the receiver is $f_A(t)$, and the probability of degradation time *T* being greater than arrival time *t* is $P_B(T > t)$, the probability of messenger molecules hitting Rx before degradation is

$$f_A(t) \cdot P_B(T > t) \tag{7}$$

that is denoted by $h(t|\lambda)$ and equals to

$$h(t|\lambda) = \frac{r_r}{d+r_r} \frac{d}{\sqrt{4\pi Dt^3}} e^{-\frac{d^2}{4Dt} - \lambda t}.$$
(8)

Note that the mathematical formula in Equation 8 represents the concentration of received molecules for an enzymatic MCvD channel with a point Tx and an absorbing sphere Rx. The equations are used for scenarios with a point Tx, but do not directly correspond to cases with a sphere Tx. By indirectly using Equation 8, exponential decay for a channel with spheres Tx and Rx can easily be implemented and simulated. Although a separate analytical equation for the sphere Tx case is not induced, by changing the simulation so that molecules are emitted from the right-end of the spherical Tx (see Figure 2. "Tx Emit Point") and molecules that try to enter the Tx are reflected, we can implement a system for the sphere Tx. In Section 5.3, we verify that using different Tx shapes affects the received signal of molecules.

3 | CHANNEL ENVIRONMENTS

3.1 | Topology

This paper considers 2 different topologies: point Tx to sphere Rx and sphere Tx to sphere Rx. In the case of the Rx, a sphere shape is preferred to a point shape because better reception can be done with bigger shapes to a certain extent.^{22,25} For Tx, however, it is not yet clear which shape will be better for ISI mitigation. Therefore, the point Tx and sphere Tx with identical enzyme areas deployed around each of them are compared to evaluate which Tx is better for ISI mitigation.

3.2 Channel geometry and parameters

The specific geometry and important parameters of the MCvD channel analyzed in this paper are shown in Figure 3. Figure 3 shows the case where enzymes are deployed around the Rx with a sphere Tx for 3 different enzyme area cases. Other scenarios will have the same principle of geometry and system parameters with variation in topology or type and size of the enzyme area. In Figure 3, r_{enz} stands for the extended enzyme radius. The spheres Tx and Rx are both nonpassive. By nonpassive we mean that the Tx reflects the messenger





FIGURE 3 Detailed geometry and parameters of the molecular communication via diffusion channel with enzymes deployed around Rx for 3 different sizes of enzyme area

molecules that try to enter it by putting them back to their original positions and the Rx absorbs the messenger molecules that enter it by eliminating them from the channel after counting them. A point Tx will be passive in terms of interaction with the propagating molecules.

The enzyme area is an extending sphere shape being homocentric to the Rx or Tx (depending on the deployment structure). A limited amount of enzymes are only deployed within the enzyme area and the enzymes only affect the messenger molecules that are inside the designated enzyme area. Depending on the value of r_{enz} , the enzyme area's total volume will be decided. Note that the volume of the enzyme area is critical to implementing a constant number of limited enzymes into different systems. In this study, the value of r_r is fixed and identical for both the Rx and Tx, but the r_{enz} and dvary. The half-life of the enzymes is changed to see its affect on the system.

3.3 Limited enzyme implementation

Because of the fact that different channel scenarios with different enzyme area sizes are compared among each other, the amount of enzymes should always be constant for fair comparison. Note that we are keeping the number of enzymes constant, not the concentration. To be able to keep the amount of enzymes identical for all of the scenarios, the volume of the enzyme area, V_{totenz} , is used. Recall from Equation 6 that

$$[S][E] = \ln 2/\Lambda_{1/2}.$$
 (9)

The amount of enzymes is fixed to a unit 1, and we set $\ln 2/[S]$ as constant *c*. Then [*E*] is

$$[E] = 1/V_{\text{totenz}} = c/\Lambda_{1/2}.$$
 (10)

Therefore, by multiplying a certain V_{totenz} value to $\Lambda_{1/2}$, a constant unit number of enzymes will be maintained among different enzyme areas and deployment scenarios. This special type of $\Lambda_{1/2}$ is the effective half-life explained in Section 3.3.2.

3.3.1 | Total enzyme area

As inferred from the previous section, the total enzyme area is needed to calculate the effective half-life. The total enzyme area, V_{totenz} , should exclude any volumes of Tx or Rx that overlaps with the enzyme area. If V_{lp} is the volume of the overlapping area, then

$$V_{\rm totenz} = \frac{4}{3}\pi r_{\rm enz}^3 - V_{\rm lp} \;. \tag{11}$$

For finding the value of $V_{\rm lp}$, notice from Figure 3 that $V_{\rm lp}$ changes depending on the $r_{\rm enz}$. For a small $r_{\rm enz}$, that is, $r_{\rm enz} \le d$, $V_{\rm lp}$ only contains the volume of a single Rx or Tx as in *Enzyme Area 1* in Figure 3. When $r_{\rm enz}$ increases and is within the range of $d + 2r_r > r_{\rm enz} > d$, $V_{\rm lp}$ is the volume of a single Rx or Tx plus the lens-similar shape where the enzyme area and the Tx or Rx overlap partially. This lens-similar shape is a sphere-to-sphere intersection and can be calculated accordingly. ²⁶ This second case corresponds to the case of *Enzyme Area 2* in Figure 3. The last case of $V_{\rm lp}$ is when $r_{\rm enz} \ge d + 2r_r$. In this case $V_{\rm lp}$ is the volume of both Tx and Rx because the enzyme area overlaps with both (*Enzyme Area 3* in Figure 3). Hence, $V_{\rm lp}$ is

$$V_{\rm lp} = \begin{cases} \frac{8}{3}\pi r_r^3 , & \text{if } r_{\rm enz} \ge d + 2r_r \\ \frac{4}{3}\pi r_r^3 , & \text{if } r_{\rm enz} \le d \\ A + \frac{r_r^2 - r_{\rm enz}^2}{4d_c} + \frac{4}{3}\pi r_r^3 , & \text{otherwise} \end{cases}$$

$$A = \pi (r_{\rm enz} - d)^2 \frac{d_c^2 + 2d_c r_r - 3r_r^2 + 2d_c R_{\rm enz}}{12d_c} ,$$

$$d_c = 2r_r + d , \qquad R_{\rm enz} = r_r + r_{\rm enz} . \qquad (12)$$

Now, we can calculate V_{totenz} from Equation 11 that is used to evaluate the effective half-life for controlling a constant and limited number of enzymes.

3.3.2 | Effective half-life

To use Equation 10, where [*E*] is inversely proportional to V_{totenz} , a V_{totenz} must be multiplied to a reference $\Lambda_{1/2}$. Because our system's V_{totenz} changes depending on the scenario type and r_{enz} , we calculate a standard V_{totenz} , denoted as $V_{\text{totenz},1\mu\text{m}}$, and divide the current V_{totenz} with $V_{\text{totenz},1\mu\text{m}}$ and multiply the result to the original half-life. The result is the effective half-life as in Equation 14.

Assume a system where $r_{enz} = r_i$, half-life is $\Lambda_{1/2}^{r_i}$ with V_{totenz,r_i} . Then for 2 different cases of r_{enz} , the half-life can

be evaluated as

$$\Lambda_{1/2}^{r_2} = \Lambda_{1/2}^{r_1} \frac{V_{\text{totenz}, r_2}}{V_{\text{totenz}, r_1}}.$$
(13)

Now, we define references $\Lambda_{1/2}$ and V_{totenz} as $\Lambda_{1/2}^{\mu m}$ and $V_{\text{totenz},1\mu m}$, which are the standard half-life and standard total enzyme area when $r_{\text{enz}} = 1\mu m$. This way for any different V_{totenz} we can calculate the effective half-life as

$$\Lambda_{1/2}^{r_{\rm enz}} = \Lambda_{1/2}^{l\mu m} \frac{V_{\rm totenz, r_{\rm enz}}}{V_{\rm totenz, l\mu m}}.$$
 (14)

The effective half-life will accordingly change each time the scenario or r_{enz} changes. Note that enlarging the enzyme area has the drawback of reducing the degradation effect of enzymes due to the lowered enzyme concentration. On the other hand, a larger enzyme area increases the probability of the diffusing molecules entering the enzyme area. Hence, depending on the size of the enzyme area, there is a trade-off between degradation power and the probability of the molecules entering the enzyme area. Because only a limited amount of enzymes is available, how densely and in what size we make this enzyme region is a system design issue that accompanies trade-off. Therefore, such trade-off suggests that we should focus on finding the optimal deployment scenario and r_{enz} for system design.

Substituting $\Lambda_{1/2}^{r_{enz}}$ into Equation 5, we get the final probability of a messenger molecule inside the specified enzyme area not decaying for each Δt step as Equation 15. Now, we have formulated a degrading function for the limited number of enzymes case in a specified enzyme area.

$$\mathbf{P}(\text{no degradation } |\Lambda_{1/2}^{r_{\text{enz}}}) = e^{-\frac{ln(2)}{\Lambda_{1/2}^{r_{\text{enz}}}\Delta t}} = \frac{1}{2^{\Delta t/\Lambda_{1/2}^{r_{\text{enz}}}}}.$$
 (15)

3.4 | Enzyme deployment scenarios

There are mainly 4 different enzyme deployment scenarios analyzed in this paper: Point Tx Around Rx, Point Tx Around Tx, Sphere Tx Around Rx, and Sphere Tx Around Tx as depicted in Figure 4. For the rest of this paper, these are named as PT-ARx, PT-ATx, ST-ARx, and ST-ATx, respectively. These types of scenarios are compared among each other while having identical r_{enz} , $\Lambda_{1/2}^{l\mu m}$, t_s (symbol period), and *d* to make the channel environments identical except for the deployment type. Once the deployment with the best ISI mitigating performance is found, the optimum r_{enz} value for different $\Lambda_{1/2}^{l\mu m}$, t_s , and *d* will be analyzed.

4 | SIMULATION SYSTEM

In our simulation system, for each time frame Δt , every molecule emitted by the Tx moves by diffusion dynamics



FIGURE 4 Diagram and notations of the 4 different enzyme deployment scenarios

governed by the Gaussian distribution at each dimension, as follows:

$$\Delta \vec{r} = (\Delta x, \ \Delta y, \ \Delta z)$$

$$\Delta x \sim \mathcal{N}(0, \ 2D\Delta t)$$

$$\Delta y \sim \mathcal{N}(0, \ 2D\Delta t)s$$

$$\Delta z \sim \mathcal{N}(0, \ 2D\Delta t),$$

(16)

where $\Delta \vec{r}$, Δx , Δy , and Δz correspond to the displacement vectors and the displacements at *x*, *y*, and *z* dimensions at a time frame of Δt and $\mathcal{N}(\mu, \sigma^2)$ corresponds to the Gaussian distribution with mean μ and variance σ^2 .

At each Δt time step, each molecule is checked if it is inside the Tx node. If so, the molecule is put back to its original position that is outside Tx. Each molecule is checked again to see if it is inside the Rx node. The ones inside the Rx are counted and eliminated, ⁷ constituting the received signal. The last step for the simulation is to check for degradation of the remaining messenger molecules. For each molecule, the probability for not degrading Equation 15 is compared to a uniformly distributed random number for degradation check. This process is repeated until we reach t_{end} , the simulation end time. Note that because of the large number of molecules sent at 1 symbol period, which is approximately 5×10^4 , there is a risk of high complexity in simulation. It is, however, necessary to check through all the molecules for fine accuracy of the results. For less complexity, the operations may be performed in parallel by several simulators.

5 | RESULTS AND ANALYSIS

5.1 | Performance metrics and parameters

For each simulation type, 50 replications are done. In our simulations different r_{enz} values are considered with fixed r_r for both Tx and Rx. For every different r_{enz} , a different $\Lambda_{1/2}^{r_{enz}}$ is cal-

TABLE 1 Values and ranges of the parameters used in the simulations

Parameter	Value
Diffusion coefficient (D)	$100\mu m^2 s^{-1}$
Radius of the Rx/Tx (r_r)	5µm
Enzyme radius (r_{enz})	$2 \sim 26 \mu m$
Distance (d)	4,6,8,10µm
Molecules emitted for 1 t_s	5×10^4 molecules
Symbol period (t_s)	$0.1 \sim 1.0s$
Simulation end time (t_{end})	0.4,2.0 <i>s</i>
Unit half life $(\Lambda_{1/2}^{1\mu m})$	$0.002 \sim 0.008s$
Simulation step (Δt)	$10^{-5}s$
Replications for simulation	50



FIGURE 5 Received signals for ST-ARx system for 4 symbol periods when $t_s = 0.1s$. $(d = 4\mu m, r_r = 5\mu m)$

culated for maintaining a constant amount of limited enzymes and a different **P**(no degradation $|\Lambda_{1/2}^{r_{enz}}$) will be applied to the system.

For the evaluation of ISI, this study uses the interference-to-total-received (ITR) molecules metric. For a certain symbol period t_s , and simulation end time t_{end} , ITR is



FIGURE 6 Molecule absorption locations at the receiver for "Without Enzymes" and "With Enzymes" ($d = 6\mu m$, $r_r = 5\mu m$

defined as

$$ITR(t_s, t_{end}) = \frac{F(t_{end}) - F(t_s)}{F(t_{end})},$$
(17)

where $F(\cdot)$ indicates the total number of molecules received until time *t*. The parameter indicates the portion of ISI molecules to the total number of received molecules. In our case, a smaller ITR indicates a better ISI mitigation. In Table 1, we present the system parameters and their values or ranges that are used for the simulations and performance analysis.

5.2 Using enzymes

The received signals for 4 symbol periods when enzymes are used and not used are shown in Figure 5. In the received signal for using enzymes, the ISI molecules do not accumulate, so the height of the peak and tail of the signal is almost constant and low for all 4 symbol periods. On the contrary, when enzymes are not used, ISI molecules accumulate for each symbol period, causing the heights of the peak and tail of the signal to radically increase for each symbol period. This will more likely cause the receiver to erroneously decode the signal for a concentration-based modulation scheme. Hence, using enzymes prove to be more effective in ISI mitigation than not using them.

To get more understanding of how the enzymes affect the hitting probability, the point of hits for both of the cases, namely, with and without enzymes, is analyzed. Figure 6 shows the hitting locations from different view points. Upper and lower rows correspond to the cases without and with enzymes, respectively. More molecules are hitting from the receiver's back hemisphere for the without enzymes case compared to the enzyme added scenario. Molecules that are hitting from the back lobe travel longer distance than the other molecules that result in longer duration for reaching the receiver. Therefore, we can claim that the ISI is reduced when the enzymes are used.

5.3 | Shape of transmitter node

Different topologies of the enzymatic MCvD channel are analyzed to decide whether to use a sphere Tx or a point Tx. We compared PT-ATx and ST-ATx to determine which is better in terms of ITR. It is clearly supported by Equation 1 that the hitting probability increases with increasing the receiver radius, so the Rx will remain as a sphere instead of a point. For the analysis we keep the r_r fixed and only focus on the Tx's shape. The sphere Tx system is modeled by changing the point Tx's simulation apt to the sphere Tx.

Figure 7 indicates that the received signals and ITRs for ST-ATx and PT-ATx differ from each other for distance 4μ m and 8μ m when each scenarios' other system parameters were kept identical. The signals for the PT-ATx cases have heavier tails than those of the ST-ATx signals. The ITR for ST-ATx is much lower than that of PT-ATx for both of the distances. Hence, using a sphere Tx shows better ISI mitigation



FIGURE 7 Received signals (top) and ITR (bottom) for comparing sphere and point source scenarios ($r_r = 5\mu$ m, $r_{enz} = 2\mu$ m, $t_s = 0.5s$, $t_{end} = 2.0s$, $\Lambda_{1/2}^{1\mu m} = 0.002s$). ITR, interference-to-total-received

performance, and we will use a sphere Tx node for the rest of the analysis. It is noticeable that even though the analytical derivation of the hitting probability is not separately solved for the sphere Tx, still the modified simulation using the analytical equation for the point Tx provides valid results for the received concentration of the sphere Tx case.

5.4 | Deployment location

After deciding that the sphere transmitter causes less ISI, the second decision parameter is about the enzyme deployment location. We analyze the performance of the following deployment locations, namely, ST-ARx, ST-ATx, and "everywhere (randomly spread)"[†]. With a limited amount of enzymes, whether enzymes should be densely deployed in a specific structure like ST-ARx and ST-ATX or just randomly spread around the entire channel like "everywhere" is unclear. In either case we use the same amount of limited enzymes. If randomly spreading the enzymes yields better ISI mitigation than the other densely deploying scenarios, then pre-deciding a specific structure and area for the enzyme deployment will be unnecessary. Results in Figure 8 show that allocating enzymes in a specific structure, that is, ST-ARx and ST-ATx, has lower ITR than just randomly spreading them everywhere. Spreading a certain amount of enzymes randomly around the channel has a received signal almost identical to that of using no enzymes. This implies that when the enzymes are spread out randomly throughout the channel, the amount of enzymes is so low compared to the entire volume of the channel that the



FIGURE 8 Received signals (top) and ITR (bottom) for comparing deployment schemes ($d = 4\mu$ m, $r_r = 5\mu$ m, $r_{enz} = 6\mu$ m, $t_s = 1.0s$, $t_{end} = 2.0s$, $\Lambda_{1/2}^{l\mu m} = 0.002s$). ITR, interference-to-total-received

channel is almost identical to that of "No Enzymes." Hence, when a limited amount of enzymes is used, allocating the enzymes in a specific structure has lower ITR than randomly allocating them. ST-ARx and ST-ATx exhibit similar performance with the given parameters. The specific allocation structure that has better ITR property between ST-ARx and ST-ATx is analyzed more thoroughly in the next section.

5.5 Deployment structure: around Rx/Tx

In general, ISI molecules are considered to accumulate closer to the Rx than the Tx after propagating some distance. ST-ARx may therefore be assumed to give better ISI mitigation. To evaluate this hypothesis, ST-ARx and ST-ATx are compared for 2 distances (4μ m and 8μ m) with different t_s and r_{enz} . Figure 9 shows the received signals for each of the scenarios. The difference is clear between the signals in terms of signal peak and the heaviness of the signal tail. For both distances, ST-ATx has the signal with lower peak and shorter, less-heavy tail than that of ST-ARx.

More analysis is done with more varied system parameters in the ITR graph in Figure 10. Figure 10 shows the ITR for ST-ARx and ST-ATx with different r_{enz} for $t_s = 0.3$ and 0.6 seconds. The ITR graphs show similar trends for both t_s values. Until the r_{enz} reaches a certain value, 4μ m in this case, ST-ATx has lower ITR than ST-ARx. Once that value is exceeded, ST-ARx starts to have lower ITR than ST-ATx and reaches the lowest ITR value. Once r_{enz} gets large enough, however, the ITR of ST-ARx and ST-ATx increases and gets almost identical as both channels become similar to the channel in which enzymes are randomly spread everywhere.

When the enzyme area is tight, ST-ATx is better in ISI mitigation than ST-ARx. Hence, with the selected parameters, if the enzyme deployment constraints do not allow r_{enz}

[†]Note that we need to use a limited enzyme area not to have 0 enzyme concentration. Hence, a considerably big area is used to refer to the case of spread randomly "everywhere." For instance we consider the enzyme radius 4 times the longest Tx-Rx distance.



FIGURE 9 Received signals of ST-ARx and ST-ATx scenarios for $d = 4\mu$ m (top) and 8μ m (bottom) ($r_r = 5\mu$ m, $r_{enz} = 2\mu$ m, $t_{end} = 2.0s$, $\Lambda_{1/2}^{1\mu m} = 0.002s$)



FIGURE 10 ITR of ST-ARx and ST-ATx scenarios for $t_s = 0.3$ (top) and 0.6 seconds (bottom) and different r_{enz} ($d = 4\mu m, 8\mu m, r_r = 5\mu m, t_{end} = 2.0s, \Lambda_{1/2}^{l\mu m} = 0.002s$). ITR, interference-to-total-received

to be greater than 4μ m, then deploying the enzymes around Tx should be selected. When, however, the enzyme area gets large to a certain value, around Rx is preferable. The 2 scenarios get nearly identical ITR when the enzyme area gets very large. The lowest ITR occurs for ST-ARx. Therefore when optimum ITR mitigation is necessary regardless of the enzyme area size, ST-ARx should be used.

The size of the enzyme area (ie, r_{enz}) that maximizes ISI mitigation for the ST-ARx scenario is also analyzed. Figure 11 shows the graph of the ITR for varying r_{enz} and



FIGURE 11 ITR of ST-ARx with varying r_{enz} and t_s for $d = 6\mu m$ (top) and $8\mu m$ (bottom) ($r_r = 5\mu m$, $t_{end} = 2.0s$, $\Lambda_{1/2}^{lum} = 0.002s$). ITR, interference-to-total-received

 t_s for $d = 6\mu$ m and 8μ m. Clearly, there is an optimum r_{enz} , namely, r_{enz}^* , where lowest ITR occurs for each d and t_s . Here, r_{enz}^* is defined as the r_{enz} when lowest ITR occurs for a specific channel. In cases of Figure 11, r_{enz}^* ranges from 6 to 12μ m depending on the distance and t_s . How r_{enz}^* is influenced by the channel parameters d, t_s , and $\Lambda_{1/2}^{1\mu m}$ is elaborated in the next section.

5.6 Relation of r_{enz}^* to channel parameters

This section analyzes how r_{enz}^* is related to the channel's distance, symbol period, and half-life. To specify in detail the relation between the distance and r_{enz}^* , Figure 12 presents the varying r_{enz}^* depending on the increasing *d* for different t_s . It is clear from the graph that there is a steady, upward trend relationship between the distance and the r_{enz}^* for all t_s . If the distance increases this will mean that the r_{enz}^* also is increased, implying an optimum ratio of distance to r_{enz}^* for maximized ISI mitigation. The slope of the fitting lines, Δ , for each of the t_s is also shown, suggesting that as t_s increases geometrically by a multiplication of 2, the increase of r_{enz}^* gets less steep. Therefore, for an increasing distance, r_{enz} also must be increased for optimizing ISI mitigation but the symbol period, t_s , should also be taken into consideration regarding to how steeply r_{enz}^* changes.

 r_{enz}^* 's dependence on the unit half-life, $\Lambda_{1/2}^{l\mu m}$, is also evaluated. The half-life of an enzyme is defined as the time required for the enzyme's target substrate concentration to fall to its



FIGURE 12 r_{enz}^* depending on the distance for different t_s for ST-ARx scenario with fitting lines ($r_r = 5\mu m$, $t_{end} = 2.0s$, $\Lambda_{1/2}^{1\mu m} = 0.002s$)



FIGURE 13 Heatmap of interference-to-total-received with varying r_{enz} and $\Lambda_{1/2}^{l \mu m}$ for ST-ARx scenario ($d = 6\mu m$, $r_r = 5\mu m$, $t_s = 0.1s$, $t_{end} = 2.0s$)

half value. Hence, the lower the half-life the faster the enzyme degrades the substrates. Figure 13 shows a heatmap of the ITR with r_{enz} , $\Lambda_{1/2}^{l\mu m}$ as the *x*-, *y*-axis, respectively. Four different half-lives are considered in this study: 2, 4, 6, and 8 milliseconds. Clearly, the lower the $\Lambda_{1/2}^{l\mu m}$ the lower the ITR because the degradation occurs faster. The r_{enz}^* , however, does not change according to the $\Lambda_{1/2}^{l\mu m}$. For all 4 $\Lambda_{1/2}^{l\mu m}$ the r_{enz}^* is $6\mu m$ in this case. Therefore, the $\Lambda_{1/2}^{l\mu m}$ affects only the rate of degradation but not the r_{enz}^* value.

5.7 | Shape of the enzyme region

In the previous analysis, we refined the shape of the enzyme region to only a spherical shape as an extension of the spherical Tx or Rx. In a different perspective, however, we can alter the shape of the enzyme region to a cube and see its effect on the ITR. Figure 14 shows the ITR comparison between spherical and cubical enzyme regions for 2 different sizes of the enzyme regions, large and small r_{enz} . Note that the vol-



FIGURE 14 ITR of ST-ARx for sphere and cube enzyme regions of different size ($d = 8\mu m$, $r_r = 5\mu m$, $t_{end} = 2.0s$, $\Lambda_{1/2}^{1\mu m} = 0.002s$). ITR, interference-to-total-received

umes of the 2 different shapes of enzyme regions must be identical for a fair comparison; having the same volume of the enzyme regions means having the same degradation power. Hence, the V_{totenz} for cube and sphere cases is equal, but the r_{enz} for each shape will vary. Here, small r_{enz} indicates the case when V_{totenz} is calculated for the spherical enzyme region when $r_{\text{enz}} = 4\mu$ m and the large r_{enz} is for the case when spherical r_{enz} is 6μ m. The corresponding r_{enz} s for the cubical cases are found to match the volumes of the spherical cases.

From Figure 14 we can see that the cubical enzyme region yields better ISI mitigation than the spherical enzyme region. Other than the ITR case for 0.2 second of large r_{enz} , which



FIGURE 15 Plot of the spherical and cubical enzyme region shapes with the same volume for small r_{enz} case

has almost identical ITR for cube and sphere, the cubical regions have lower ITRs. To analyze the reasons for such result, we plotted the actual enzyme regions for sphere and cube as in Figure 15. From the figure it is clear that while the cubical enzyme region has a higher probability of degrading molecules that are less directed toward the receiver, the spherical enzyme region is more likely to degrade the molecules that are directed toward the receiver. In other words, the cubical region is more efficient in degrading the molecules that are less likely to arrive at the receiver, so it can degrade the molecules while maintaining a higher signal power than the spherical region. This result opens the possibility of an entire different area of research on the shape of the enzyme region that can differently affect the extent of ISI mitigation.

6 | CONCLUSION

This paper analyzed the different system structures and parameters that can maximize ISI mitigation by using a limited amount of enzymes. In terms of topology, when the same amount of enzymes were used a sphere Tx was shown to yield more ISI mitigation than a point Tx. For the enzyme deploying location, randomly deploying the enzymes everywhere created more ISI molecules than deploying them in a specific structure. As to which specific structure is more preferable for ISI mitigation, when the enzyme area is small to a certain extent, ST-ATx had less ISI. For a larger enzyme area, however, ST-ARx had less ISI and the lowest ISI occurred for the ST-ARx scenario. Once the enzyme area got very large, the 2 different scenarios yielded almost identical results.

For the case of ST-ARx there proved to be an optimum size of the enzyme area that maximizes the ISI mitigation. This optimum enzyme area increased as the distance between the Rx and Tx increased, and the rate of increase decreased as the symbol period increased. The half-life, on the other hand, had no effect on the optimum enzyme area size, but a lower half-life meant less ISI.

Further research is possible on deriving the mathematical interpretations and expressions for the limited enzyme implementation with a spherical Tx and optimized system parameters. Although only spherical enzyme regions are mainly considered in this research, other options for the shape of the enzyme region are open for research such as the cubical enzyme regions we briefly presented in our analysis. Moreover, the research can be applied to molecular multiple-input multiple-output (MIMO)^{27,28} where the limited enzymes around Rx or Tx can be used as a methodology for mitigating interlink interference.

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REFERENCES

- Akyildiz IF, Jornet JM, Pierobon M. Nanonetworks: a new frontier in communications. *Commun ACM*. 2011;54(11):84–89.
- Nakano T, Eckford AW, Haraguchi T. Molecular Communication. New York: Cambridge University Press; 2013.
- Farsad N, Yilmaz HB, Eckford AW, Chae CB, Guo W. A comprehensive survey of recent advancements in molecular communication. *IEEE Commun* Surveys Tuts. 2016;18(3):1887–1919.
- Guo W, Mias C, Farsad N, Wu JL. Molecular versus electromagnetic wave propagation loss in macro-scale environments. *IEEE Trans Mol Bio Multi-Scale Commun.* 2015;1(1):18–25.
- Farsad N, Kim NR, Eckford AW, Chae CB. Channel and noise models for nonlinear molecular communication systems. *IEEE J Sel Areas Commun.* 2014;32(12):2392–2401.
- Tepekule B, Pusane AE, Yilmaz HB, Chae CB, Tugcu T. ISI mitigation techniques in molecular communication. *IEEE Trans Mol Bio Multi-Scale Commun.* 2015;1(2):202–216.
- Yilmaz HB, Chae CB. Simulation study of molecular communication systems with an absorbing receiver. *Simul Model Pract Th*. 2014;49:136–150.
- Kuran MŞ, Yilmaz HB, Tugcu T, Akyildiz IF. Interference effects on modulation techniques in diffusion based nanonetworks. *Nano Commun Netw.* 2012;3(1):65–73.
- Kilinc D, Akan OB. Receiver design for molecular communication. *IEEE J Sel Areas Commun.* 2013;31(12):705–714.
- Kuran MŞ, Yilmaz HB, Tugcu T, Akyildiz IF. Modulation techniques for communication via diffusion in nanonetworks. Proc. IEEE Int. Conf. on Commun. (ICC); 2011; Kyoto.
- Kim NR, Chae CB. Novel modulation techniques using isomers as messenger molecules for nano communication networks via diffusion. *IEEE J Sel Areas Commun.* 2013;31(12):847–856.
- Kim NR, Eckford AW, Chae CB. Symbol interval optimization for molecular communication with drift. *IEEE Trans NanoBiosci.* 2014;13(3):223–229.
- Lin WA, Lee YC, Yeh PC, Lee C. Signal detection and ISI cancellation for quantity-based amplitude modulation in diffusion-based molecular communications. Proc. IEEE GLOB. Telecomun. Conf. (GLOBECOM); 2012; California.
- Kuran MŞ, Yilmaz HB, Tugcu T. A tunnel-based approach for signal shaping in molecular communication. Proc. IEEE Int. Conf. on Commun. (ICC); 2013; Budapest.
- Noel A, Cheung KC, Schober R. Improving receiver performance of diffusive molecular communication with enzymes. *IEEE Trans NanoBiosci*. 2014;13(1):31–43.
- Heren AC, Yilmaz HB, Chae CB, Tugcu T. Effect of degradation in molecular communication: impairment or enhancement?. *IEEE Trans Mol Bio Multi-Scale Commun.* 2015;1(2):217–229.
- Yilmaz HB, Cho YJ, Guo W, Chae CB. Interference reduction via enzyme deployment for molecular communication. *IET Electronic Letters*. 2016;52(13):1094–1096.
- Nakano T, Okaie Y, Liu JQ. Channel model and capacity analysis of molecular communication with brownian motion. *IEEE Commun Lett.* 2012;16(6):797–800.
- Tyrrell HJV, Harris KR. Diffusion in Liquids, a Theoretical and Experimental Study. Stoneham, MA: Butterworth Publishers; 1984.
- Berg HC. Random Walks in Biology. Princeton, NJ, USA; Princeton University Press:1993.
- Redner S. A Guide to First-passage Processes. New York: Cambridge University Press; 2001.

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- Yilmaz HB, Heren AC, Tugcu T, Chae CB. Three-dimensional channel characteristics for molecular communications with an absorbing receiver. *IEEE Commun Lett.* 2014;18(6):929–932.
- 23. Yilmaz HB, Chae CB. Arrival modelling for molecular communication via diffusion. *IET Electron. Lett.* 2014;50(23):1667–1669.
- 24. Murray JD. Mathematical Biology: I. An Introduction. (3 ed). New York: Springer; 2002.
- Akkaya A, Yilmaz HB, Chae CB, Tugcu T. Effect of receptor density and size on signal reception in molecular communication via diffusion with an absorbing receiver. *IEEE Commun Lett.* 2015;19(2):155–158.
- Kern WF, Bland JR. Solid Mensuration: with Proofs. New York: J. Wiley & Sons, Incorporated; 1938.
- Lee C, Koo B, Farsad N, Eckford AW, Chae CB. Molecular MIMO communication link. Proc. IEEE Int. Conf. on Comput. Commun. (INFOCOM); 2015; Hong Kong.

 Koo B, Lee C, Farsad N, Eckford AW, Chae CB. Molecular MIMO: from theory to prototype. *IEEE J Sel Areas Commun.* 2016;34(3):600–614.

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