Membrane Clustering and the Role of Rebinding in Biochemical Signaling

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ABSTRACT In many cellular signaling pathways, key components form clusters at the cell membrane. Although much work has focused on the mechanisms behind such cluster formation, the implications for downstream signaling remain poorly understood. Here, motivated by recent experiments, we use particle-based simulation to study a covalent modification network in which the activating component is either clustered or randomly distributed on the membrane. We find that whereas clustering reduces the response of a single-modification network, it can enhance the response of a double-modification network. The reduction is a bulk effect: a cluster presents a smaller effective target to a substrate molecule in the bulk. The enhancement, on the other hand, is a local effect: a cluster promotes the rapid rebinding and second activation of singly activated substrate molecules. As such, the enhancement relies on frequent collisions on a short timescale, leading to an optimal ratio of diffusion on the other hand, is a local effect: a cluster promotes the rapid rebinding and second activation of singly activated substrate molecules. As such, the enhancement relies on frequent collisions on a short timescale, leading to an optimal ratio of diffusion to association that agrees with typical measured rates. We complement simulation with analytic results at both the mean-field and first-passage distribution levels. Our results emphasize the importance of spatially resolved models, showing that significant effects of spatial correlations persist even in spatially averaged quantities such as response curves.

INTRODUCTION

Although cells are often modeled as well-mixed chemical reactors, they are highly spatially heterogeneous entities. Beyond merely providing the blueprint for space-dependent processes such as division or patterning, spatial heterogeneities in cellular components are frequently exploited by biochemical networks as additional degrees of freedom in signaling computations (1). The most direct example of this is compartmentalization, in which the same chemical component initiates different phenotypic responses depending on where it is localized within the cell (2,3). In a similar way, the localization of signaling components via scaffolding proteins has effects on signal amplification that depend nontrivially on the surrounding chemical conditions (4). In fact, the colocalization of just two components can have a dramatic response on the amplification properties of an enzyme-driven reaction network (5). Even in spatially uniform systems, spatial correlations between individual molecules can have significant effects on the mean response (6).

One of the most actively studied areas in which spatial heterogeneity is emerging as a key factor is signal transduction at the cell membrane. In addition to imposing a quasi-two-dimensional geometry, the membrane plays host to a large diversity of cellular components, and interactions among these components give rise to a complex spatial organization (7). A central theme of recent work in this field is the prevalence and role of membrane clusters, groups of colocalized molecules that often participate in the detection of external signals and subsequently drive responses within the cell. Perhaps the best-known example of this process is bacterial chemotaxis, in which clusters of receptors detect external ligands, triggering messenger molecules to modulate the activity of flagellar motors (8). Recent studies have also provided evidence for clustering in eukaryotic cell membranes: data from immunoelectron microscopy (9) and single-molecule fluorescence experiments (10) suggest that Ras, a protein that has been implicated in a variety of phenotypic responses (e.g., oncogenesis), forms membrane clusters on which the efficacy of its downstream signaling critically relies. Clustering may also be connected to the partitioning of the membrane itself into spatially segmented domains (11,12), e.g., via interaction with the cytoskeleton (13) or the formation of so-called lipid rafts (14).

Although much modeling work has been done to elucidate the possible mechanisms by which clusters form (15–17), insights into the role that clustering plays in downstream signaling remain largely speculative. Therefore, our primary goal in this study was to quantitatively assess the effect of clustering on the input-output response of a canonical signaling network, using a spatially resolved model. Recognizing its ubiquity in the systems in which clustering is observed (18), we focused on a covalent modification network (often called the push-pull network) in which a substrate is alternately activated and deactivated by two antagonistic components (Fig. 1 A). For example, in bacterial chemotaxis, the kinase CheA and the phosphatase CheZ phosphorylate and dephosphorylate the messenger protein CheY, respectively; CheA and CheZ therefore play the roles of the two antagonistic components, and CheY plays the role of the substrate.

Moreover, focusing on a push-pull network naturally permits extension to a double-modification process (Fig. 1 B),
which is a critical step in many membrane-signaling pathways. In eukaryotic cells, for example, active Ras molecules at the membrane initiate a mitogen-activated protein kinase (MAPK) cascade within the cell, each layer of which consists of a dual phosphorylation cycle. In general, dual phosphorylation can be carried out by one of two mechanisms. In a processive mechanism, an enzyme modifies both phosphorylation sites on a substrate molecule before releasing it. In a distributive mechanism, on the other hand, the enzyme must release the substrate after modifying the first site, before rebinding and modifying the second site. It has been shown experimentally that key kinases (19,20) and phosphatases (21) in the MAPK cascade act in a distributive manner, which makes the rebinding process critically important. Therefore, a second goal of this study was to investigate the interplay between clustering and rebinding, and its role in determining the input-output response of a distributive push-pull network.

We provide a spatially resolved description of the system by performing particle-based simulations on a lattice. In parallel, we gain important physical intuition from analytic results derived at both the mean-field and first-passage distribution levels. We find that the input-output response of the network changes depending on whether the activating component is clustered or randomly distributed on the membrane (Fig. 1 C). Specifically, whereas clustering reduces the response of a single-modification network, it can enhance the response of a double-modification network. We demonstrate that the reduction is a direct consequence of the fact that a cluster presents a smaller effective target to a substrate molecule in the bulk. By investigating in detail the stochastic nature of the rebinding process, we discover that the enhancement has an entirely different origin: clustering promotes the rapid rebinding and second activation of singly activated substrate molecules (Fig. 1 C). We find that such a rapid effect is only exploited when both the activating and deactivating components are sufficiently free to react, such that ultrasensitive networks (22) in which one or the other component is saturated by the substrate do not exhibit the enhancement.

Furthermore, we find that the enhancement with clustering becomes more pronounced as the diffusion coefficient is raised. Underlying this observation is the fundamental advantage that clustering affords as collisions occur more frequently: although the diffusion may be high enough to prevent a substrate molecule from rapidly re-binding an isolated enzyme molecule, it may be insufficient to enable the substrate molecule to escape an entire cluster. Clustering thus prolongs the possibility of rapid rebinding, effectively boosting the association rates of individual molecules, which are often limited by tight orientational constraints (23). Of course, this advantage reaches a limit: at infinite diffusion, all spatial arrangement is forgotten. We are thus led naturally to a ratio of diffusion to association at which the enhancement is optimal.

Together, our results provide a quantitative picture of the nontrivial effects that membrane clustering has on biochemical signaling, for a network that plays a critical role in systems in which clustering has been experimentally observed. More broadly, our results demonstrate the crucial role that spatial correlations play in cellular function, and the associated importance of considering spatial resolution in biophysical models.

**METHODS**

We consider both a single- and a double-modification push-pull network, which the activating enzyme is localized to the membrane (Fig. 1). To understand the effect of clustering, we compare the situation in which activating enzyme molecules are arranged randomly on the membrane with that in which they are localized at the same surface density to clusters of size N (Fig. 1 C). Because we are interested in the effect of clustering on downstream signaling, and not in the dynamics of cluster formation on the membrane, we take activating enzyme molecules to be fixed. Substrate and deactivating enzyme molecules diffuse freely in the cytoplasm with diffusion coefficient D.

**Chemical reactions, input-output relation, and sensitivity**

The single-modification network (Fig. 1 A) is described by the reactions

\[ E_a + S \stackrel{k_1}{\rightleftharpoons} E_aS \stackrel{k_2}{\rightarrow} E_a + S^* \]  
(1)

\[ E_d + S^* \stackrel{k_3}{\rightleftharpoons} E_dS^* \stackrel{k_4}{\rightarrow} E_d + S \]  
(2)

where S and S* denote the substrate in its inactive and active forms, respectively. Activation is catalyzed by the activating enzyme E_a, which first
forms a complex before releasing the substrate in its active state, and deactivating is performed similarly by the deactivating enzyme $E_a$. The double-modification network (Fig. 1 B) prescribes additional reactions identical to Eqs. 1 and 2, except with $S$ and $S^*$ replaced by $S$ and $S^*$, respectively. We restrict our analysis to networks whose first and second modification processes are identical (i.e., the rates $k_1, k_2, \ldots, k_6$ describing the first modification also describe the second). Furthermore, we assume negligible back reactions: $k_2 = k_5 = 0$. We tested the effects on the main results (Fig. 2) of systematically varying $k_2$ and $k_5$. The effects can be understood in terms of the insights provided in the Results section, and are discussed explicitly in the Supporting Material.

The input of the network is defined as the catalytic rate of the activating enzyme $k_1$, scaled by its counterpart $k_0$ for the deactivating enzyme: $\chi = k_1/k_0$ (in chemotaxis, for example, $k_3$ is typically set by the time-averaged ligand occupancy of the receptor cluster (24)). The output is the relative activity of the substrate, i.e., the fraction $\phi = [S^*/S]_T$, $[S^*/S]_T$ for the single- or double-modification network, respectively. In a deterministic, well-mixed description, in which rate equations determine the dynamics, the steady-state input-output relation is completely specified by the reaction rates and the conserved total concentrations of substrate $[S]_T$ and enzymes $[E]_T$ and $[E]_T$. In particular, for both the single- and double-modification networks, one may write the input-output relations entirely in terms of the dimensionless parameters (e.g., see Supporting Material):

$$\alpha = \frac{[E]_T}{[E]_T}, \beta = \frac{k_4}{k_1}, \gamma = \frac{K}{[S]_T}, \epsilon = \frac{[E]_T}{[S]_T},$$

where $K = k_0/k_4$ is the Michaelis-Menten concentration of the deactivation process, and $[E]_T/N$ is divided by the volume. The first two parameters determine the bias of the network toward deactivation; $\alpha = \beta = 1$ therefore corresponds to a symmetric network. The last two parameters characterize the sensitivity of the network: in the zero-order (or ultrasensitive) regime, the substrate saturates the enzymes and operates far beyond the Michaelis-Menten concentration ($\epsilon, \gamma \ll 1$), whereas in the linear regime, both substrate and enzymes operate in the linear regions of their response curves ($\epsilon^{-1}, \gamma^{-1} \ll 1$).

**Lattice model**

We perform spatially resolved simulations with excluded volume interactions on a regular three-dimensional lattice. We make the approximation that all molecules have equal diameter $d$, and we let this diameter define the lattice spacing, such that molecules neighboring each other on the lattice are in contact. Clustered molecules are placed in contact in a square arrangement on the membrane (we tested that a circular arrangement does not change the results). The membrane comprises the $x-y$ plane and extends for a length $L$ in each direction, beyond which periodic boundaries are imposed. The cytoplasm has depth $Z$, with reflective boundaries at both the membrane ($z = 0$) and the farthest point from it ($z = Z$). The Supporting Material provides a detailed account of how reactions and diffusion are implemented on the lattice, in particular such that detailed balance is obeyed. All source code, written in C++ and MATLAB (The MathWorks, Natick, MA), is freely available at http://rebind.sourceforge.net.

Spatial resolution introduces new parameters into the problem beyond those of the well-mixed system (Eq. 3), which are captured by the dimensionless quantities given below. In addition to the cluster size $N$, one has

$$\delta \equiv \frac{4 \pi D}{k_1}, \mu \equiv \frac{N \ell^2}{L^2}, \xi \equiv \frac{Z}{\ell}$$

The quantity $1/4\pi \delta = k_1/4\pi D$ is the ratio of the activating enzyme’s intrinsic association rate $k_1$, which is the association rate given that the molecules are in contact, to the corresponding diffusion-limited value $4\pi D$; as such, $\delta$ captures the strength of diffusion relative to association. The parameters $\mu$ and $\xi$ describe the surface density of activating enzymes and the cytoplasmic depth, respectively.

**Parameter selection**

In the following section, we discuss in detail the effects of varying the parameters that govern network symmetry ($\alpha, \beta$), sensitivity ($\gamma, \epsilon$), and diffusion ($\delta$). In all results that establish network characteristics (see Figs. 2, 3, and 5, and Fig. S1, Fig. S2, and Fig. S3), the surface density of activating enzymes and the cytoplasmic depth are set using estimates from experimentally studied systems. We arrive at the typical values $\mu = 0.01$ and $\xi = 25$, as described in the Supporting Material.

**RESULTS**

We begin by presenting and explaining the main difference between the single- and double-modification networks:
whereas clustering reduces the response of a single-modification network, it can enhance the response of a double-modification network. The magnitude of each effect scales with the cluster size $N$ (Fig. 2). The reduction for single-modification networks is generic and persists with changes in network symmetry ($\alpha$, $\beta$), sensitivity ($\gamma$, $\epsilon$), and diffusion ($\delta$). The enhancement for double-modification networks, on the other hand, is more specific and occurs in deactivation-biased linear-sensitivity networks with intermediate diffusion. Results presented later in this section will explain this specificity.

Fig. 2 also illustrates more generally the effect of localizing the activating enzymes to the membrane by comparing the spatially averaged response with the response in the well-mixed case. As shown in Fig. 2A, localization reduces the maximal response of a single-modification network (compare the well-mixed curve with the random curve). Such a reduction was seen in previous work (5) and is the result of the concentration gradients that form due to the asymmetric localization of activating and deactivating enzymes. As shown in Fig. 2B, the double-modification network can avoid this reduction and can in fact achieve an amplification beyond the well-mixed response instead.

**Clustering reduces the effective target size**

How does clustering reduce the response of a single-modification network? The key is that a cluster presents a smaller effective target to a molecule in the bulk. The target size reduction leads to substrate molecules spending more time in the inactive state, and thus to a reduced output (Fig. 2A). One can understand this target size reduction by imagining that each molecule possesses a local neighborhood, i.e., a volume into which another molecule can diffuse and eventually react (Fig. 3A). When the $E_a$ molecules are arranged in a random configuration at sufficiently low density, the total volume of such neighborhoods (the target size) is simply $N$ times an individual $E_a$ molecule’s neighborhood. However, when the $E_a$ molecules are clustered, the individual neighborhoods overlap and the target size is reduced (Fig. 3A).

To understand quantitatively the impact of the target size reduction on the response of the network, we consider the time it takes an $S$ molecule, released from the bulk, to bind an $E_a$ molecule on the membrane (the lifetime of the $S$ molecule). If the $E_a$ molecules are free with high probability (i.e., unoccupied by other substrate molecules), the lifetime is dominated by the search time $s$, the time to find and bind an $E_a$ molecule. The mean search time from the bulk can be estimated as the inverse of the association rate over the volume of the box. An excursion from the bulk is dominated by the diffusive trajectory, such that we can estimate the association rate by its diffusion-limited value: $\tau = (4\pi D/L^2)^{-1} = L^2/4\pi D$. A random distribution of $E_a$ molecules presents $N$ targets of diameter $\ell$, which reduces the mean search time by a factor of $N$: $\tau = s/N = L^2/4\pi N D$. A cluster, on the other hand, presents one target with effective diameter $\ell_{\text{eff}} \sim \sqrt{N \ell}$, making the mean search time $\tau = L^2/4\pi (\sqrt{N} \ell) D$. Scaling by the natural timescale $\ell^2/D$ (the time to diffuse approximately

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**FIGURE 3 Target size effect.** (A) Left: Cartoon depicting neighborhoods (solid lines) surrounding activating enzyme molecules (solid circles) in random and clustered configurations. The total neighborhood volume (the target size) is smaller in the clustered configuration due to the overlap of individual molecules’ volumes. Right: A smaller target admits fewer distinct paths from the bulk (dashed lines), thereby increasing the mean time required to find the target by a diffusive search. (B) The main plot shows the mean lifetime of $S$ molecules in the single-modification network; parameters are as in Fig. 2A. The inset shows that the value to which the mean lifetime asymptotes at high input $\chi$ in the clustered configuration (the mean search time $\tau$ ) grows as the square root of the cluster size $N$. 

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one molecular diameter), and recalling Eq. 4, these times read $\bar{\sigma} \equiv \bar{\sigma} / (d^2 / D) = \zeta / 4\pi\mu$ and $\bar{\sigma} \equiv \bar{\sigma} / (d^2 / D) = \sqrt{N\zeta / 4\pi\mu}$, which makes clear that at constant surface density the search time is independent of $N$ for the random configuration, but scales with $N^{1/2}$ for the clustered configuration.

Fig. 3 B shows the mean lifetime of $S$ molecules as a function of the input for a single-modification network with zero-order sensitivity. At high input, the mean lifetime asymptotes to the value corresponding to the search from the bulk, consistent with the above analysis. It is clear that for the clustered configuration, this asymptotic value depends on the cluster size $N$, and the inset shows that it indeed scales with $N^{1/2}$, as predicted.

Two important conditions of the above analysis are that the $E_a$ molecules are free and that the $S$ molecule is released randomly from the bulk (and not, say, still within the neighborhood of the cluster). The first condition is met at high input $(\chi \equiv k_f / k_b \gg 1)$, when the high catalytic rate of the activating enzymes leaves the $E_a$ molecules free with high probability. The second condition is also met at high input for networks with zero-order sensitivity, in which saturation of the deactivating enzymes leaves the $E_d$ molecules occupied with high probability. High occupation of $E_d$ molecules means that a typical $S^*$ molecule has ample time to randomize its position before ultimately binding a free $E_d$ molecule and being released as an $S$ molecule. Interestingly, we observe that in networks with linear sensitivity, in which $E_d$ molecules remain free even at high input (Supporting Material), the reduction in the output upon clustering persists (not shown), which leads us to conclude that the portion of $S$ molecules that do originate in the bulk continue to contribute to a reduction in the response.

It is important to emphasize that the target size effect is a bulk effect, not a local effect, in the sense that clustering not only reduces the number of neighboring sites from which a substrate molecule can bind, it more generally reduces the number of distinct paths that lead to the target from a point in the bulk (Fig. 3 A). This intuition is confirmed by a simple test: under an alternative implementation, in which a substrate molecule can only bind an $E_a$ molecule from the neighboring lattice site perpendicular to the membrane, we observe an increase in the search time with cluster size $N$ that is only slightly less pronounced than that in the inset of Fig. 3 B (not shown). Because this alternative implementation has the property that clustering the $E_a$ molecules does not change the number of available neighboring sites, the increase in search time with $N$ is strictly due to a reduction in the number of paths from which the target is accessible.

It is also important to point out that the target size effect is a generic property of diffusive random walks, and as such it is just as present for double-modification networks as it is for single-modification networks. However, as we will describe next, in a particular parameter regime the effects of rapid rebinding can overcome the target size effect, leading to an enhancement of the response rather than a reduction.

### Clustering promotes rapid rebinding

How does clustering enhance the response of a double-modification network? The key is that a cluster promotes rapid rebinding of singly activated substrate molecules. Rapid rebinding results in more doubly activated substrate and thus an enhanced output (Fig. 2 B). Rebinding only occurs in the double-modification network. In the single-modification network, once a substrate molecule is released by an enzyme, it can only bind to an enzyme of the opposite type. To clearly understand the rapid rebinding effect in double-modification networks, we first consider the distribution of rebinding times for a reduced system: a single $S^*$ molecule is released from one of $N E_a$ molecules, with no $E_d$ molecules present. It rebinds to any $E_a$ molecule in a time $r$, whose dimensionless analog we define as $\rho \equiv r / (d^2 / D)$. Previous studies considered rebinding to a planar geometry in the context of ligand-receptor binding (25–27) (even specifically for clustered chemoreceptors in bacteria (27)), and the analytic results derived below extend the results of those studies, particularly for double-modification networks.

As shown in Fig. 4, for both a random and a clustered configuration of $E_a$ molecules, the rebinding time distribution contains three regimes. Short times (the molecular regime) correspond to short excursions, after which the $S^*$ molecule rebinds to the same $E_a$ molecule (or cluster) from which it came. Intermediate times (the planar regime) correspond to excursions that are sufficiently far for the $S^*$ to see the membrane as a plane uniformly populated with $E_a$ molecules (or, due to the periodicity, with clusters), yet not far enough to see the reflective boundary. The granularity of individual $E_a$ molecules is thus lost, and the membrane appears as a uniform semiabsorbent plane. Long times (the bulk regime) correspond to long excursions, during which the $S^*$ molecule randomizes its position completely, returning as if from the bulk. These three regimes are discussed in detail below.

### Bulk regime

The bulk regime exhibits an exponential distribution because it describes the time required to find an $E_a$ molecule given a random starting position in the bulk. An exponential distribution is expected from a well-mixed system, in which reactions obey exponential waiting time statistics. Here, however, the molecular and planar regimes emerge due entirely to spatial correlations, affecting the network even at the level of the mean response (Fig. 2 B). Moreover, in the bulk regime, the substrate molecule has strayed far enough from the membrane that it effectively returns from the bulk. This return time is therefore equivalent to the search time defined above for $S$ molecules in single-modification
networks. Accordingly, one can see in Fig. 4 that the time constant characterizing the exponential is larger in the clustered case, precisely due to the target size effect discussed above. Finally, the onset of the bulk regime is determined by the time it takes the S' molecule to randomize its position, which is approximately the time required to diffuse the full cytoplasmic depth: \( r_m = z^2/2D \), or \( \rho_m = \xi^2/2 \).

**Planar regime**

In the planar regime, the substrate molecule has not diffused far enough to enter the bulk and lose memory of its starting position, but it has diffused far enough that the membrane appears as a uniform semiabsorptive plane. The problem can be reduced to an effectively one-dimensional one in the \( z \) direction with a radiation boundary at \( z = 0 \). The one-dimensional rate \( k_{\text{eff}} \) (with dimensions of length per time) describing association at the boundary follows from a renormalization of the three-dimensional rate \( k_1 \). Clearly, \( k_{\text{eff}} \) should scale with the surface density \( N/L^2 \), and we find good agreement with the simplest dimensionally consistent definition, \( k_{\text{eff}} = k_1 N/L^2 \).

As shown in the Supporting Material, the rebinding time distribution for this one-dimensional problem is readily obtained from the Green’s function and exhibits scalings of \( \rho^{\alpha-1/2} \) at short times, \( \rho^{\alpha-3/2} \) at long times, and a crossover time of \( \rho_p = (D/k_{\text{eff}})^2 = \delta^2/\mu^2 \). Short times comprise a collision-dominated subregime in which the excursion is dominated by many unsuccessful reflections, and thus inherits the \( \rho^{\alpha-1/2} \) scaling from the Gaussian Green’s function of a particle freely diffusing in one dimension. Long times comprise a search-dominated subregime in which after a long excursion the particle returns to an effectively absorbing boundary, producing the \( \rho^{\alpha-3/2} \) scaling characteristic of a one-dimensional random walker returning to an absorbing origin. Further details are provided in the Supporting Material.

The transition between the molecular and planar regimes occurs when the \( S' \) molecule diffuses far enough perpendicular to the membrane that it no longer detects the granularity of the \( E_a \) molecules, a distance roughly equal to half the mean spacing between \( E_a \) molecules in the random configuration, or between clusters in the clustered configuration. In the random configuration, the mean spacing between \( E_a \) molecules is set by the surface density, yielding a separating time of \( r_{\text{mp}}^r = \sqrt{(L^2/N)/2} \), or \( \rho_{\text{mp}}^r = 1/8 \mu \). In the clustered configuration, the spacing between clusters is \( L \), yielding a separating time of \( r_{\text{mp}}^c = (L/2)^2 \), or \( \rho_{\text{mp}}^c = N/8 \mu \).

**Molecular regime**

The molecular regime is defined by short excursions in which the substrate molecule rebinds to the \( E_a \) molecule or cluster from which it came. The molecular regime exhibits \( \rho^{\alpha-1/2} \) and \( \rho^{\alpha-3/2} \) scalings whose origins are the same as those in the planar regime: the scalings arise from a collision-dominated or search-dominated return, respectively, to a single molecule or cluster. For a return to single molecule, which applies to the random configuration, these scalings were described in previous work (6). The crossover time was derived to be

\[
\rho_m = \frac{\xi^2}{D} \left(1 + k/4\pi\ell D^2\right)^{-1/2},
\]

where \( k = 2k_1 \), with the factor of 2 arising from reflection of the \( E_a \) across the membrane. However, one can see from Fig. 4 that in the random configuration, the crossover time is obscured by alternations in the probability density at short times, which is an artifact of the lattice implementation. To be precise, an \( S' \) molecule starting next to an \( E_a \) molecule can only rebind in an odd number of time steps (assuming it moves diffusively every time step, which is true at short times for large \( \delta \)); the exception occurs when another \( E_a \) molecule is placed next to or very near the first \( E_a \) molecule, but at low surface densities such a placement occurs with low probability. We validated the distributions in Fig. 4.
using Green’s function reaction dynamics (6), verifying that lattice artifacts do not quantitatively change the probability densities.

In the clustered configuration, the crossover time within the molecular regime is indeed resolvable and can be described in terms of the previously considered results. A large, absorbent cluster (\(N \gg 1, \delta \ll 1\)) can be approximated as a plane with an effective one-dimensional association rate \(k_{\text{eff}} = k_1/\ell^2\), yielding a dimensionless crossover time of \((D/k_{\text{eff}})^2 = \ell^2\). In the opposite limit, a small, reflective cluster (\(N \sim 1, \delta \gg 1\)) can be approximated as a spherical object whose effective diameter is obtained by equating surface areas: \(4\pi (\ell_{\text{eff}}/2)^2 = 2N\ell^2\) (neglecting cluster edges). In the limit of large \(\delta\), the denominator in Eq. 5 approaches unity, making the crossover time approximately \(\ell_{\text{eff}}^2/D\), or \(2N/\pi\) in dimensionless units. Because the expressions in both the plane and sphere limits scale with parameters that are large in the opposite limits, we use the minimum as an estimate of the crossover time: \(\rho_m^c \approx \min(\delta^2, 2N/\pi)\).

Fig. 4 corroborates all scalings and crossover times derived above using an illustrative set of sample parameters. Because we have analytic estimates for the crossover times, they can be tuned to expand or contract the various regimes, a fact we used to confirm the validity of the scalings beyond the confidence implied by Fig. 4 alone.

Fig. 4 also directly displays the advantage that clustering affords in the rebinding problem: at short times, the probability of rebinding is enhanced, leading to a probability gap over the random configuration. In fact, the characteristic time that determines the extent of this gap, \(\rho_m^c\), reveals the parameter regimes that give rise to enhanced rebinding, and thus ultimately to an enhanced signal output for the network. Specifically, the gap increases as \(\rho_m^c\) increases, by either increasing the cluster size \(N\) or increasing diffusion relative to association, \(\delta\). Increasing the cluster size is a straightforward way of enhancing rebinding, and the associated enhancement of the output is demonstrated in Fig. 2 B.

The reason that increasing diffusion increases the probability gap is perhaps less straightforward but can be understood at the molecular level. Increasing diffusion induces more unsuccessful collisions before rebinding eventually occurs. Rapid rebinding to a single \(E_a\) molecule (which is the task when the \(E_a\) molecules are randomly distributed) therefore becomes unlikely. Rapid rebinding to a cluster, on the other hand, remains less unlikely, owing to the presence of neighbors. The number of collisions in the neighborhood of a cluster is simply larger than that for a single molecule by virtue of the former’s increased size. The probability of ultimately achieving a successful collision is thus higher for the clustered configuration than for the random configuration, by a factor that increases as diffusion increases.

At a more detailed mechanistic level, we may consider the fate of an \(S^*\) molecule that has just been released by an \(E_a\) molecule, and now resides at a neighboring lattice point. As diffusion increases, the probability increases that the \(S^*\) molecule will take a step away from the \(E_a\) molecule. In the random configuration, it is then increasingly likely for diffusion to carry the \(S^*\) molecule away from the immediate vicinity of the \(E_a\) molecule. In the clustered configuration, on the other hand, several of these diffusive paths will lead directly to another \(E_a\) molecule. Clustering therefore poses an advantage when immediate rebinding is unlikely but rebinding after several diffusive steps is more probable.

Deactivation connects rebinding to the network response

Interestingly, despite the probability gap elucidated above, we observe that the means of the two rebinding distributions in Fig. 4 are the same: the enhancement conferred to the clustered configuration in the molecular regime is compensated for by the target size effect in the bulk regime. The equivalence of means is a consequence of the fact that we isolated the rebinding process. Alone, the rebinding process is equivalent to one dissociation and subsequent association event of the equilibrium reaction \(A + B \rightleftharpoons C\). Because it is an equilibrium reaction, detailed balance implies that the occupancy (i.e., the fraction of time in which an \(A\) molecule is bound to a \(B\) molecule) is independent of the spatial configuration of \(A\) molecules (excluded volume effects can be safely neglected given the low densities). Because we know that the mean time that an \(A\) molecule is bound to a \(B\) molecule (the mean bound time) is the inverse of the dissociation rate and thus independent of the spatial configuration of \(A\) molecules, the mean unbound time must also be independent of the spatial configuration of \(A\) molecules. Therefore, although the push-pull network as a whole prescribes a nonequilibrium process, the rebinding process alone is effectively in equilibrium, and the mean rebinding time is the same for a random and a clustered configuration.

Why then does the probability gap translate to an enhancement with clustering at the level of the mean response, as in Fig. 2 B? Indeed, it is precisely because thus far in the discussion we have not reintroduced the \(E_d\) molecules. The effect of the \(E_d\) molecules is to bind and deactivate the \(S^*\) molecules with the longest excursion times, removing them from the rebinding problem and thereby truncating the rebinding distributions beyond a characteristic timescale, which we call the capture time. The truncation alleviates the target size effect, imparting the clustered configuration with a shorter mean rebinding time than the random configuration. The capture effect is discussed in more detail in the Supporting Material and illustrated in Fig. S1.

The capture effect also underlies the observation that only double-modification networks with linear sensitivity, not zero-order sensitivity, benefit from clustering, as demonstrated in Fig. S2. The reason is that the capture effect relies
on the $E_d$ molecules being free. The zero-order regime, on the other hand, corresponds to saturation of the enzymes by the substrate, such that at high input the $E_d$ molecules are not free and instead are occupied with high probability. The relationship between the capture effect and network sensitivity is also discussed in more detail in the Supporting Material.

Clustering leads to an optimal ratio of diffusion to association

In studying the rebinding distributions, we discovered that increasing diffusion relative to association enhances rapid rebinding to a cluster more strongly than to a random configuration, because it increases the probability of unsuccessful collisions. Therefore, one would expect the enhancement of the output with clustering to increase with the ratio of diffusion to association, $\delta$. However, we also know that at high diffusion, the network is well mixed and the spatial arrangement of the molecules is irrelevant; clustering should therefore confer no enhancement at high $\delta$. In fact, these competing effects lead to a value of $\delta$ at which the enhancement is optimal, as shown in Fig. 5.

Fig. 5 illustrates that as $\delta$ increases, the enhancement (i.e., the difference in maximal output between the clustered and the random configuration $\Delta \phi_{\text{max}} \equiv \phi_{\text{max}}^c - \phi_{\text{max}}^r$) first increases and then decreases. The inset shows this nonmonotonic behavior for several values of the cluster size $N$. The optimal enhancement increases with $N$; moreover, the value $\delta^*$ at which the optimum occurs also increases with $N$. These observations are consistent with the notion that a larger cluster more effectively confers the advantage associated with frequent unsuccessful collisions.

Quantitatively, $\delta^*$ approaches $\sim 10$ for the largest cluster size ($N = 100$), corresponding to an association rate $k_1 \sim cD/10$, roughly $10^4 - 10^5 = 125$ times less than the diffusion-limited value. In fact, this is precisely the regime in which many protein–protein association reactions operate within cells. Most measured association rates are on the order of $10^6 - 10^7 \text{M}^{-1}\text{s}^{-1}$ (23), whereas the diffusion-limited rate is typically $10^8 - 10^9 \text{M}^{-1}\text{s}^{-1}$. Our results suggest that for these reactions, clustering can optimally enhance the output. Therefore, we expect the mechanism identified here to have quite significant biological relevance.

DISCUSSION

We have provided a detailed view of the varied effects that membrane clustering can have on the signaling properties of a canonical biochemical network. The network under study and the values of relevant biophysical parameters were drawn from experimentally studied systems, both prokaryotic and eukaryotic, in which membrane clustering was recently observed. We implemented a spatially resolved model, appealing to both simulation and analytic results to demonstrate that spatial correlations can have nontrivial effects, even at the level of the mean input-output response. In particular, we have shown that spatial effects at both the bulk scale (in terms of a diffusive target search process) and the molecular scale (in terms of rapid stochastic rebinding events) affect the response of a network in ways that are not captured by a well-mixed, spatially uniform description.

Our results make it clear that the effect of clustering depends on both the network topology and the biochemical parameters. For example, we identify a general property of diffusive random walks, i.e., that clustering the target increases the search time from the bulk, which leads generically to a reduced response in a single-modification network. However, when the topology of the network is extended to double-modification, the reduction can be overcome by a local effect: clustering promotes rapid rebinding of singly activated substrate molecules to the activating enzyme molecules. When the concentration of free deactivating enzyme molecules is sufficiently high to isolate these rapid rebinding events, the result is an enhancement of the response. Importantly, this enhancement is specific to networks with linear sensitivity. Ultrasensitive networks, in which the deactivating enzyme molecules are saturated by the substrate at high input, do not confer the enhancement because the mechanism relies on the deactivating enzyme molecules being free. Moreover, the enhancement is most pronounced in the presence of unsuccessful collisions, when the probability to escape a single activating enzyme molecule is much higher than that to escape a cluster. The specificity of the enhancement to both linear sensitivity and a certain ratio of diffusion to association highlights the importance of biochemical parameters in predicting the effects of clustering.

The result that clustering is most beneficial in the presence of unsuccessful collisions has important functional
implications. We find that the diffusion coefficient at which the cluster-induced enhancement is optimal corresponds to an intrinsic association rate roughly 100 times smaller than its diffusion-limited value. Such a result is quantitatively consistent with intracellular conditions because typical protein–protein association rates are measured to be roughly two orders of magnitude lower than the diffusion limit. Indeed, intrinsic association rates are inherently constrained by the tight orientational precision required to achieve a successful binding event, because binding requires the alignment of small reactive patches. This fact naturally induces unsuccessful collisions. Clustering may therefore have evolved as a functional way around such tight molecular geometric constraints, allowing a cell to boost signal output despite the low intrinsic association rates of individual molecules.

The effect of molecular orientation on intrinsic association rates is a subtle but important issue, especially with regard to rapid rebinding events. During very rapid rebinding between a pair of molecules, one might expect rotational alignment to be preserved, leading to an increase in the effective intrinsic association rate. On the other hand, if the molecules diffuse only a molecular diameter away from each other, then orientational diffusion will already significantly reduce the effective intrinsic association rate. Moreover, many enzymes are only capable of rebinding a substrate molecule after a characteristic reactivation time because, for example, ADP/ATP exchange must take place before rebinding can occur. In this study we take a coarse-grained view, neglecting these opposing effects and simply assuming that orientation is randomized immediately upon dissociation, even for very rapid rebinding events. Although this assumption may not be accurate for rapid rebinding to the same particle, it is important to emphasize that the benefit provided by clustering relies on rebinding to neighboring particles, for which the assumption of random orientation is more reasonable. Nonetheless, the relationship between orientation and rebinding rates would surely benefit from further detailed study.

Finally, our findings emphasize the general importance of the role of the rebinding process in biochemical signaling. The importance of rebinding in related systems has been discussed, with interesting consequences for the mean response. For example, in studying how the diffusive motion of a repressor protein effects gene expression, investigators observed that rebinding boosts noise, leading to bursts in gene expression, and that this effect can be captured by renormalizing the on- and off-rates in a well-mixed model. On the other hand, simulations and experiments on signaling via a MAPK cascade revealed that spatial correlations due to rapid rebinding introduce qualitative changes in the mean response that cannot be captured by the well-mixed theory. Our results here are more resonant with the second case, because it is clear that membrane localization and subsequent clustering introduce changes to the rebinding statistics that go beyond the exponential distributions expected from a well-mixed description. More broadly, the importance of rebinding in explaining the potency of T-cell ligand binding, in which a long aggregated binding time arises from a sequence of many fast rebinding events, has been recognized.

This study represents a first step in using simulation and analytic techniques to understand the role of spatial organization in signaling. It is our view that spatially resolved models, as well as a sharp theoretical framework, can help formalize and make more quantitative the inferences that are being made from the wealth of experimental data on systems that exhibit clustering, colocalization, and other nontrivial spatial heterogeneity.

SUPPORTING MATERIAL

Five sections, including three figures, plus references (32–38) are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(12)00205-6.

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